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COMPUTING

George Eleftherakis Petros Kefalas Gheorghe Păun

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Eighth Workshop on Membrane Computing (WMC8)

Thessaloniki, June 25 – June 28, 2007

George Eleftherakis, Petros Kefalas, Gheorghe Păun Editors

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Workshop organised by the SOUTH-EAST EUROPEAN RESEARCH CENTRE (SEERC)

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Preface

This volume contains the papers presented at the **Eighth Workshop on Membrane Computing, WMC8**, which has been organized in Thessaloniki, Greece, from June 25 to June 28, 2007. The first three workshops on membrane computing were organized in Curtea de Argeş, Romania – they took place in August 2000 (with the proceedings published in *Lecture Notes in Computer Science*, volume 2235), in August 2001 (with a selection of papers published as a special issue of *Fundamenta Informaticae*, volume 49, numbers 1–3, 2002), and in August 2002 (with the proceedings published in *Lecture Notes in Computer Science*, volume 2597). The next four workshops were organized in Tarragona, Spain, in July 2003, in Milan, Italy, in June 2004, in Vienna, Austria, in July 2005, and in Leiden, The Netherlands, in July 2006, with the proceedings published as volumes 2933, 3365, 3850, and 4361, respectively, of *Lecture Notes in Computer Science*.

The 2007 edition of WMC was organized at City College, Thessaloniki, by the South-East European Research Centre (SEERC), under the auspices of the European Molecular Computing Consortium (EMCC). As usual in the last years, a balanced attention was paid to both "theory" and "practice", to mathematical and theoretical computer science topics as well as to applications and implementations. This is especially visible in what concerns the five invited talks, all of them included in the present volume, delivered by Luca Bianco (Verona, Italy), Pierluigi Frisco (Edinburgh, UK), Alberto Leporati (Milano, Italy), Andrea Maggiolo-Schettini (Pisa, Italy), and Gheorghe Ştefan (USA).

The volume also contains the 32 accepted papers. Each of them was subject of two or three referee reports. The program committee consisted of Gabriel Ciobanu (Iaşi, Romania), Erzsébet Csuhaj-Varjú (Budapest, Hungary), Rudolf Freund (Vienna, Austria), Pierluigi Frisco (Edinburgh, UK), Marian Gheorghe (Sheffield, UK), Oscar H. Ibarra (Santa Barbara, CA, USA), Petros Kefalas (Thessaloniki, Greece) – Co-Chair, Vincenzo Manca (Verona, Italy), Giancarlo Mauri (Milano, Italy), Linqiang Pan (Wuhan, China), Gheorghe Păun (Bucharest, Romania) – Chair, Mario J. Pérez-Jiménez (Sevilla, Spain), Athina Vakali (Thessaloniki, Greece).

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The Organizing committee consisted of George Eleftherakis – Chair, Konstantinos Dimopoulos, Petros Kefalas, Ioanna Stamatopoulou, and Ognen Paunovski – Secretariat.

The invited papers and a selection of regular papers, improved according to the discussions held in Thessaloniki and additionally refereed, will be published in a special issue of *Lecture Notes in Computer Science*.

Details about membrane computing can be found at the area web site (maintained in Milano, Italy) from http://psystems.disco.unimib.it. The workshop web site was http://www.seerc.org/wmc8/.

The workshop was sponsored by City College, Thessaloniki, and the South-East European Research Centre (SEERC).

Finally, we would like to thank you all, the organizing and programme committee, the invited speakers, the authors of the papers, the lecturers, the reviewers, and all the participants. Your effort made the Eighth Workshop on Membrane Computing (WMC8) a very successful event.

> George Eleftherakis Petros Kefalas Gheorghe Păun Editors

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Psim: A Computational Platform for Metabolic P Systems

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Summary. Although born as unconventional models of computation, P systems can be conveniently adopted as modelling frameworks for biological systems simulations. This choice brings with it the advantage of producing easier to be devised and understood models than with other formalisms. Nevertheless the employment of P systems for modelling purposes demands for biologically meaningful evolution strategies as well as for complete computational tools to run simulations on. In previous papers an evolution strategy known as *metabolic algorithm* has been presented, here a simulation tool called *Psim* (current version 2.4) is discussed and a case study of its application is given as well.

1 Introduction

Membranes play a prominent role in living cells [1, 20]. In fact, membranes not only act as a separation barrier indispensable to create different environments within cells boundaries, but they can also physically constitute a kind of "working board" whereby enzymes can activate and perform their duties on substrates. Other examples of the crucial role of membranes within cells are their ability to perform selective uptakes and expulsion of chemicals as well as their being the interface of the cell with the surrounding environment allowing communication with neighboring cells.

P systems originate from the recognition of this important role of membranes and, by abstracting from the functioning and structure of living cells, they provide a novel computation model rooted in the context of formal language theory [33, 35].

P systems investigations are nowadays focused on several research lines that make the field "a fast Emerging Research Front" in computer science (as stated by the Institute for Scientific Information). In particular, theoretical investigations on the power of the computational model have been carried on and important results have been achieved so far in order to characterize the computational power of many elements of P systems (such as objects and membranes) and, from a complexity viewpoint, P systems have been employed as well in the solution of NP hard problems. For a constant up to date bibliography of P systems we refer the reader to [39].

Parallel to these lines some more practical investigations are under way too. These studies exploit the resemblance of P systems to biological membranes in order to develop computational models of interesting biological systems. P systems seem to be particularly suitable to model biological systems, due to their direct correspondence of many elements (namely membranes, objects-chemicals and rules-reactions), even in their basic formulation, with real biological entities building the system to be modelled. Moreover, many extensions have been proposed to the standard formulation of P systems, such as some biologically relevant communication mechanisms [28, 36, 11], energy account [37] and active membranes [34] among others, which show the flexibility of the model. In this way, discrete mathematical tools can be used to represent interesting biological realities to be investigated. A further step is that of simulating all systems described in this way to get more information about their internal regulatory mechanisms and deeper insights into their underlying elements.

Although born as a non-conventional model of computation inspired by nature, P systems can therefore be employed as a simulation framework in which to embed the in silico simulation of interesting biological systems. The strength of this choice is, as said, the advantage that P systems share with biological systems many of their features and this leads to easy-to-devise and easy-to-understand descriptions of the studied realities. In fact, the membrane construct in P systems has a direct counterpart into biological membranes: objects correspond to all chemicals, proteins and macromolecules swimming in the aqueous solution within the cell and, eventually, rewriting rules represent biochemical reactions taking place in the controlled cellular environment. Other formalisms have been employed as modelling and simulation frameworks too, such as Π calculus [29], Petri nets [38] and Ambient calculus [10], but in their case the very same notions of membranes, chemicals and reactions need to be reinterpreted and immersed in the particular representation formalism in a less immediate way.

Nevertheless, the employment of P systems as a modelling framework for biological systems posed, from a purely computational viewpoint, some new challenges to cope with, such as the identification of suitable, biologically meaningful, strategies for system evolution and the development of new automatic tools to describe, simulate and analyze the investigated system.

In previous works a novel strategy for systems evolution, called *metabolic algorithm* has been introduced [6, 27, 8], an hybrid (deterministic-stochastic) variant of which has been proposed as well [5]. Other strategies of evolution are known, such as *Dynamical Probabilistic P systems* [32, 31] and *Multi-compartmental Gillespie's algorithm* [30, 2].

Here we will focus on the metabolic algorithm in its deterministic version which has been confronted with the dynamics of several systems (a collection of case studies can also be found in [4]). Some examples of investigated systems by means of the metabolic algorithm are the Belousov-Zhabotinsky reaction (in the Brusselator formulation) [6, 8], the Lotka-Volterra dynamics [6, 27, 7, 14], the SIR (Susceptible-Infected-Recovered) epidemic [6], the leukocyte selective recruitment in the immune response [16, 6], the Protein Kinase C activation [8], circadian rhythms [12] and mitotic cycles in early amphibian embryos [26]. In order to cope with the demand of computational tools to simulate the dynamics of P systems, we developed a first simulator called *Psim* [6], which has now been extended with several new features as will be explained later on. The new version of the simulator is freely available for download at [15].

The remaining of the discussion will firstly introduce (section 2) some theoretical aspects of the simulation framework we developed and some recent advances will be mentioned too. Section 3 will then be devoted to the newer version of the simulator itself and a practical case study will be given as well in such a way to show to the reader how to set up a simulation with the interface of Psim.

2 MP systems

MP systems (Metabolic P systems) [21, 26, 24, 23] are a special class of P systems [33, 35], introduced for expressing the dynamics of metabolic (or, more generally speaking, biological) systems. Their dynamics is computed by means of a deterministic algorithm called *metabolic algorithm* which transforms populations of objects according to a *mass partition* principle, based on suitable generalizations of chemical laws.

A definition of MP systems follows, as given in [4].

Definition 1 (MP system). An MP system of level n - 1 (i.e., with $n \in \mathbb{N}$ membranes) is a construct:

$$\Pi = (T, \mu, Q, R, F, q_0)$$

in which:

- T is a finite set of symbols (or objects) called the alphabet;
- μ is the hierarchical membrane structure, constituted by n membranes, labeled uniquely from 0 to n - 1, or equivalently, associated in a one-to-one manner to labels from a set L of n - 1 distinct labels;
- Q is the set of the possible states reachable by the MP system. Each element q ∈ Q is a function q : T × L → ℝ, from couples objects-membranes to real values. A value q(X, l), with X ∈ T and l ∈ L gives the amount of objects X inside a membrane labeled l, with respect to a conventional unit measure (grams, moles, individuals, ...);
- R is the finite set of rewriting rules. Each r ∈ R is specified according to the boundary notation [3]. In other words, each rule r has the form α_r → β_r, where α_r, β_r are strings defined over the alphabet T enhanced with indexed parenthesis representing membranes. As an example, an hypothetical rule can have the form: α₁β → γ₁δ, with α, β, γ, δ ∈ T*, meaning that α and β are

respectively changed in γ and δ , where all objects within α and δ are outside membrane labeled 1, whereas elements of β and γ are placed inside membrane 1;

- *F* is the set of reaction maps, each $f_r \in F$ is a function uniquely associated to a rule $r \in R$, defined as $f_r : Q \longrightarrow \mathbb{R}$ and, given a certain state q, it produces $f_r(q)$ that is a real number specifying the strength of rule r in acquiring objects;
- $q_0 \in Q$ is the initial state of the system. It specifies the initial amount of all the species throughout the various compartments of the system.

Since encodings like [9] show that the membrane structure can be flattened by augmenting the alphabet size, the definition of the membrane structure μ is not very important in this context and the choice to employ 0-level MP systems in the remaining of the discussion is not limiting from a theoretical point of view. Moreover, dealing with 0-level MP systems ends up in a easier discussion, in fact all states $q \in Q$ do not need the specification of a membrane label and in this way they have the simpler form: $q: T \longrightarrow \mathbb{R}$. For this reason, in the following whenever the term MP system will be used, the more correct term 0-level MP system has to be implicitly assumed.

The dynamics of MP systems has been calculated, starting from the initial state q_0 by means of an evolution strategy called *metabolic algorithm* [6, 27, 8], which is substantially different from the well known *non-deterministic and maximally parallel* paradigm followed by standard P systems. More precisely, the perspective of MP systems is to model systems at a population level rather than at an objects level. In this way, nothing can be precisely said about individuals but the investigation is focused on the macroscopic dynamics which assumes a deterministic flow in spite of individual behaviors.

2.1 The metabolic algorithm: hints

Without entering into many details (which can be found anyway in [6, 27, 8, 25]), the metabolic algorithm is a deterministic strategy for MP systems evolution based on mass partition among rules of all elements in the alphabet T.

In very general terms, the metabolic algorithm can be summarized in the following main points [26]:

- Reactants are distributed among all the rules, as the system evolves, according to a "competition" strategy.
- If some rules compete for the same reactant, then each of them obtains a portion of the available substance that is proportional to its reaction strength (*reactivity*) in that state.
- The reactivity of a rule in a certain state measures the capability of the rule to acquire its reactants. It is calculated by the evaluation of the reaction map corresponding to the rule due and it depends on the state of the system, that is defined as the concentration and localization of all substances in the considered instant.

• The evolution strategy determines the reaction unit of all rules, that is, the unitary amount of substance which is dealt by the rule. The stoichiometry is used then to obtain the consumed and/or produced amount of substances for each rule.

An example may be useful to clarify the concepts yet introduced. Let us suppose that in a given instant, four rules, namely r_1, r_2, r_3 and $r_4 \in R$, need molecules of a species A (with A belonging to the alphabet T) as reactant (see Figure 1), then a partition strategy for A is necessary.



Fig. 1. Competition for object A between rules r_1, r_2, r_4 and r_5 .

A real number called *reactivity* represents the strength of the rule (i.e. the rule's capability of obtaining matter to work on), given by the value assumed by a function uniquely associated to the rule called *reaction map*, in the considered state. For example, with respect to Figure 1, if we denote with a, b and c the concentrations of species A, B, C respectively (in a state q not specified for the sake of simplicity), then the reactivities of rules r_1, r_2, r_4 and r_5 , which ask for A molecules, can be:

$$f_1 = 200 \cdot a, f_2 = 0.5 \cdot a^{1.25} \cdot b^{-1}, f_4 = a^{1.25} \cdot (b+c)^{-1}$$
 and $f_5 = 10$

where the choice of reaction maps f_i , $i = \{1, 2, 4, 5\}$ is completely arbitrary in this example.

We define the quantity

$$K_{A,q} = \sum_{i=1,2,4,5} f_i(q)$$

as the *total pressure* on A in the state q (the intuitive idea is that all reaction maps of rules competing for a certain species give the force that pushes that species to react).

Then, for each of the competing rules r_j we consider the *partial pressure* (or *weight*) of r_j on type A as

$$w_{A,q}(r_j) = \frac{f_j(q)}{K_{A,q}}$$

(again the idea behind this is that the strongest the force pushing an element to follow a particular reaction channel, compared to other reaction channels, the more matter will follow that path).

Note that, in general, the quantity $K_{X,q}$ is defined for each couple (X,q) where $X \in T$ and q is a possible state of the system, moreover a weight $w_{X,q}(r)$ has to be calculated for each triple (X,q,r) where X and q are respectively, as before, an element of the biological alphabet and a state of the system, while $r \in R$ is a rule in which the element X appears as a reactant (i.e., according to the terminology adopted above, $X \in \alpha_r$).

Getting back to the example discussed above, it should be easy to see that the partial pressure of r_1 on A is

$$w_{A,q}(r_1) = \frac{200a}{200a + 0.5a^{1.25}b^{-1} + a^{1.25}(b+c)^{-1} + 10}$$

while the same pressure due to r_2 results to be equal to

$$w_{A,q}(r_2) = \frac{0.5a^{1.25}b^{-1}}{200a + 0.5a^{1.25}b^{-1} + a^{1.25}(b+c)^{-1} + 10}$$

and the other weights can be calculated analogously. The weights calculated so far determine the partition factors of the amount of species A, available in the state q, among the rules which need objects A as reactants.

Now to calculate the reaction unit of a particular rule (i.e. the amount of reactant that can be dealt by the rule) we simply need to multiply the partial pressure of the rule on the reactant by the real amount of reactant present into the system at the considered state. For example, the reaction unit of rule r_1 (or equivalently, the amount of A that that can be assigned to rule r_1) turns out to be $w_{A,q}(r_1) \cdot a = \frac{0.5a^{2.25}b^{-1}}{200a+0.5a^{1.25}b^{-1}+a^{1.25}(b+c)^{-1}+10}$.

In this way, if r_1 is a rule of the form $A \to X$, no matter what element is represented by $X \in T$, then the amount of A associated to r_1 is exactly $u_1 = w_{A,q}(r_1) \cdot a$ and the effect of r_1 's application is the loss of u_1 units of A and the acquisition of the same number of units of X.

In the case of cooperative rules (i.e. rules with more than just one reactant) things are a little bit more complicated since we need to take into account the real availability of all reactants taking part to the reaction. That is, for each X belonging to the reactants of a certain rule r we need firstly to compute the quantities $w_{X,q}(r) \cdot x$ and, since we have to respect species availability, the reaction unit associated to the rule is then computed as the minimum of those quantities. If we suppose that a rule r_1 has the form $AAB \to X$, then the reaction unit

$$u_1 = min(rac{1}{2}w_{A,q}(r_1) \cdot a \;,\; w_{B,q}(r_1) \cdot b)$$

where the term $\frac{1}{2}$ in the first element of the minimum is due to the fact that A appears twice in the stoichiometry of the rule.

In general terms, the metabolic algorithm is an effective procedure for calculating in each state of the system a reaction unit u_i for all rules $r_i \in R$ by using a partition strategy that employs particular functions f_i associated in a 1-1 manner to rules. After this calculation, the evolution of the system can be obtained in a straightforward way by consuming and producing species in a quantity given by rules' reaction units and by following the stoichiometry of the system.

Assuming an ordering on objects and on rules, let us denote with M the $m \times n$ stoichiometric matrix associated to an MP system having m symbols and n rules (in which the $c_{i,j}$ element of M denotes the gain or the loss of the *i*th object due to rule j^1) and with U_q the $[u_1 \cdots u_n]^T$ vector of the reaction units in a state of the system q, calculated as mentioned above.

Then, as pointed out in [25] the transition from one state q to the following one is done by means of the *delta operator* $(\Delta_x(q))$ which is a *m*-sized vector giving the variation of each species in the transition from state q to the next state q'. In particular

$$\Delta_x(q) = M \times U_q$$

stating that the delta operator can be obtained as the product of the stoichiometric matrix M by the reaction units vector of state q, U_q .

Since each row i of $\Delta_x(q)$ gives the variation on the *i*th object, then if we think of a state q as a vector containing the concentration of all the m species at the corresponding instant, then the next state can easily be calculated as

$$q' = q + \Delta_x(q) = q + M \times U_q.$$

Just to exemplify the last concepts discussed, we can think about an alphabet $T = \{A, B, C\}$ and focus on a rule set comprising the following four rewriting rules:

$$\begin{array}{ccc} r_1 : A & B \longrightarrow C \\ r_2 : B & B \longrightarrow A \\ r_3 : C & \longrightarrow A \\ r_4 : C & \longrightarrow B \end{array}$$

then, assuming the lexicographic order on elements of the alphabet, we can obtain the following stoichiometric matrix:

$$M = \begin{bmatrix} -1 & 1 & 1 & 0\\ -1 & -2 & 0 & 1\\ 1 & 0 & -1 & -1 \end{bmatrix}$$

in which, the first row corresponds to the object A and states that we lose one conventional unit of A due to rule r_1 , we get one A both from rule r_2 and r_3 and finally r_4 does not affect A concentration at all.

¹ It is the difference between the number of occurrences of the *i*th symbol among products and among reactants of *j*th rule.

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Then, let us suppose to be in a state q, described by the vector of concentrations $q = [10 \ 32 \ 20]^T$ (i.e. we have 10 units of A, 32 of B and 20 of C) and that the corresponding reaction units vector $U_q = [7 \ 12 \ 5 \ 9]^T$ (i.e. reaction r_1 moves 7 mass units, $r_2 \ 12$, $r_3 \ 5$ and finally $r_4 \ 9$). In this way it is possible to calculate the next state q' which turns out to be described by the following vector:

$$q' = q + M \times U_q = \begin{bmatrix} 10\\32\\20 \end{bmatrix} + \begin{bmatrix} -1 & 1 & 1 & 0\\-1 & -2 & 0 & 1\\1 & 0 & -1 & -1 \end{bmatrix} \times \begin{bmatrix} 7\\12\\5\\9 \end{bmatrix} = \begin{bmatrix} 20\\10\\13 \end{bmatrix}$$

describing the amount of all species at that particular instant.

In previous papers [26] a convenient and intuitive formalism for representing MP systems called *MP graphs* has been proposed. In particular, MP graphs are bipartite graphs describing both the stoichiometry (i.e. the shape of the rules) and the regulative part of MP systems that need to be effectively calculated in order to obtain the dynamics of the system (i.e. the reaction maps). According to what said above, MP graphs represent all the information needed to simulate MP systems by means of the metabolic algorithm. An example of MP graphs, as produced by the simulator we developed, will be shown later on.

2.2 Generalizing the metabolic algorithm

According to the formulation of the dynamics given in the previous section, the metabolic algorithm is a strategy that given a particular state q provides the system with the corresponding reaction units vector U_q which is used to calculate the transition to the state q + 1. As discussed in [25], other strategies can be considered whose aim is to produce a reasonable mass partition among all rules of an MP system, or in other words that give a different U_q for each state q of the system.

This view leads to the definition of several *metabolic algorithms* instead of a single one and the definition of MP systems can be generalized accordingly.

Based on the definition given in [22], Definition 1 can be easily generalized, in very general terms, in the following way:

Definition 2 (Generalized MP systems). A 0-level (generalized) MP system is a 6-tuple:

$$\Pi = (T, Q, R, V, q_0, \phi)$$

in which:

- T is a finite set of symbols (or objects) called the alphabet;
- V is a finite set of variables;
- Q is the set of the possible states reachable by the MP system. Each element $q \in Q$ is a function $q: T \cup V \longrightarrow \mathbb{R}$;
- *R* is the finite set of rewriting rules;

- $q_0 \in Q$ is the initial state of the system;
- ϕ is the strategy of evolution, $\phi: Q \longrightarrow \mathbb{R}^n$ with |R| = n.

Note that nothing is said about the cardinality of the set of variables V and they are not necessarily associated in a one-one manner to rules of R. Moreover, the strategy of evolution ϕ , given a state q has to be defined in such a way that it outputs the *n*-tuple providing the reaction unit vector of the system, or following the terminology used above, $\phi(q) = U_q$.

Complete freedom is left in the implementation of the strategy of evolution, whose only constraints are that given a state it has to provide the reaction unit vector corresponding to that state, which will be used to calculate the evolution of the system by means of the matrix product recalled in the previous section. As will be mentioned in the following, the specification of a fully customizable strategy of evolution will be one of the prominent features of the new version of the simulator Psim that has been implemented within the MNC Group of the University of Verona.

3 Psim

Based on the theoretical framework expressed in previous sections, a simulator called *Psim* (P systems simulator) has been developed to cope with the problem of calculating the dynamics of biological systems. An early version of Psim has been developed previously [6], with which the newer version shares the same philosophy, though extending some of its concepts and enhancing the simulation environment with many features.

The present release of Psim (version 2.4) has been developed in response to the need of an effective and easy to use tool to calculate the dynamics of MP systems by means of the metabolic algorithm. Its implementation has moreover followed some flexibility and extensibility principles which led to a tool that can be easily extended and integrated with other tools. In this way Psim, thanks to its immediate setup (nothing needs actually to be done provided a Java virtual machine is installed on the computer that is meant to run Psim) and to the easy user interface, can be used by people without a strong background in programming and a deep knowledge in the field of computer science. On the other hand, the extendability provided, as we will see, by means of the plugins mechanism, allows people with stronger expertise in programming to build their own tools to complement and integrate the main core of Psim.

Some features of this tool, which is implemented by using the Java programming language, are listed below:

• friendly user interface which is born to be easy-to-use and to interact with people not necessarily having a strong computer science background. Its immediacy can be found in the input side, which can be specified by means of a transposition of the concept of MP graphs into a point and click graphical

interface. Moreover, the same simplicity principle holds for the output side as well, which is basically constituted by a graph containing the temporal evolution of all the species constituting the system (both on a temporal scale and on the phase plan space);

- plugins architecture: the interaction with the system can either be done man-. ually or by means of some specifically designed plugins which, thanks to the plugins support offered by the simulation engine, can interact with the simulation engine itself. More specifically, three different kinds of plugins can be devised and implemented in Java as well. Input plugins can be used to implement various sources for the data to run the simulation on (let us think to some specific pathways databases like KEGG for instance); output plugins conversely, can be used to observe and analyze in various ways the results obtained from a simulation and can therefore give some meaningful insights into the simulated dynamics. Moreover, they can be used to export simulation data into particular formats. Finally, experiment plugins can directly control and intimately interact with the simulation engine, by controlling the execution flow, checking some properties and changing some experimental conditions. This kind of plugins can be very useful in tasks like model optimizations and stability analysis;
- extreme flexibility. The simulation tool we propose is based on a simulation engine which is designed to accept the definition of new evolution strategies for the calculation of the systems dynamics. At the only price of the implementation of some specific interfaces, the developer has the chance to define his own simulation strategies and to design a customized library of *metabolic algorithms* to calculate the systems evolution;
- models portability has been implemented by using the standard XML language and some extensions towards the SBML language are being considered too;
- cross platform applicability, thanks to the choice of Java, Psim can be run on all platforms supporting the Java virtual machine architecture.

An aspect deserving a special emphasis here is the possibility offered by the simulation engine, to specify custom evolution strategies. Getting back to the definition of generalized MP systems, the architecture of the simulator allows the specification of a fully customized ϕ function. A set of evolution strategies can be devised by developing in Java a specific class implementing a particular interface provided by the main engine. Several different strategies can be handled simultaneously by the simulator that gives the chance to decide which simulation strategy employ in the simulation process. This gives the tool a very high level of flexibility and power as well as the plugins mechanism does. Since plugins can interact with the simulation engine at a different levels, such as input, output but also at the simulation level too, they can be used for various reasons within the simulator and this again gives users plenty of ways to improve the system and to extend its functionalities.

3.1 A case study

In this section we show an application of the Psim computational tool for the simulation of the well known mitotic oscillator as found in early amphibian embryos [18, 19, 26].

Mitotic oscillations are a mechanism exploited by nature to regulate the onset of mitosis, that is the process of cell division aimed at producing two identical daughter cells from a single parent. More precisely, mitotic oscillations concern the fluctuation in the activation state of a protein produced by cdc2 gene in fission yeasts or by homolog genes in other eukaryotes. The model here considered focuses on the simplest form of this mechanism, as it is found in early amphibian embryos. Here, the progressive accumulation of the cyclin protein leads to the activation of cdc2 kinase. This activation is achieved by a bound between cyclin and cdc2 kinase forming a complex known as M-phase promoting factor (or MPF). The complex triggers mitosis and degrades cyclin as well; the degradation of cyclin leads to the inactivation of the cdc2 kinase that brings the cell back to the initial conditions in which a new division cycle can take place.

Goldbeter [18, 19] proposed a minimal structure for the mitotic oscillator in early amphibian embryos in which the two main entities are cyclin and cdc2 kinase. According to this model, depicted in Figure 2, the signalling protein cyclin is produced at a constant rate v_i and it triggers the activation (by means of a dephosphorylation) of cdc2 kinase, passing from the inactive form labelled M^+ to the active one, denoted by M. This modification is reversible and the other way round is performed by the action of another kinase (not taken into account in the model) that brings M back to its inactive form M^+ . Moreover, active cdc2 kinase (M) elicits the activation of a protease X^+ that, when in the active (phosphorylated) form (X), is able to degrade the cyclin. This activation, as the previous one, is reversible as stated by the arrow connecting X to X^+ .

The set of differential equations devised by Goldbeter produces an oscillatory behavior in the activation of the three elements M, C, X that repeatedly go through a state in which cells enter in a mitotic cycle (see Figure 3).

The goal of the case study showed here is to obtain a description and a simulation of the very same model of mitotic oscillations by means of the simulator Psim.

In general, there is no unique way to translate a differential equation system in terms of a metabolic P system, therefore we choose to obtain it by the application of the MP-ODE transformation [13]. The resulting MP system is reported here:

$$\Pi = (T, \mu, \mathcal{R}, F, q_0)$$

where:

- The alphabet: $T = \{A, C, X, X^+, M, M^+\}$
- The membrane structure: $\mu = \begin{bmatrix} 0 \end{bmatrix}_0$;



Fig. 2. The mitotic oscillator model by A. Goldbeter, from [18].



Fig. 3. Dynamics of the mitotic oscillator from [18].

• The set of rules is $\mathcal{R} = \{r_1, r_2, ..., r_{10}\}$, where:

 $\begin{array}{rcl} r_1: & A & \rightarrow A \; C \\ r_2: & C & \rightarrow X \\ r_3: & X & \rightarrow \lambda \\ r_4: & C & \rightarrow \lambda \\ r_5: & C & \rightarrow M \; C \\ r_6: & M^+ & \rightarrow \lambda \\ r_7: & M & \rightarrow M^+ \\ r_8: & X^+ & \rightarrow X \; M \\ r_9: & M & \rightarrow \lambda \\ r_{10}: & X & \rightarrow X^+ \end{array}$

in which all symbols have the meaning described before (and A is a kind of well to draw substance C out from). Moreover, for every symbol in the system, we

have introduced an inertia rule (i.e., a rule having the form $Y \to Y$, for each $Y \in T$ to model the inertia of the system), omitted in this set of rules.

• The set of reaction maps is $F = \{Fr_1, Fr_2, ..., Fr_{10}\}$, where:

$$\begin{array}{ll} Fr_{1} &= k_{1} \\ Fr_{2} &= k_{2} \frac{x}{k_{3}+c} \\ Fr_{3} &= k_{2} \frac{c}{k_{3}+c} \\ Fr_{4} &= k_{3} \\ Fr_{5} &= k_{5} \frac{m^{+}}{(k_{6}+c)(k_{7}+m^{+})} \\ Fr_{6} &= k_{5} \frac{c}{(k_{6}+c)(k_{7}+m^{+})} \\ Fr_{7} &= k_{8} \frac{c}{k_{1}} \frac{m^{+}}{(k_{1}+m)} \\ Fr_{8} &= k_{10} \frac{x^{+}}{(k_{11}+x^{+})} \\ Fr_{9} &= k_{10} \frac{x^{+}}{(k_{11}+x^{+})} \\ Fr_{10} &= k_{12} \frac{1}{(k_{13}+x)} \end{array}$$

• The initial state q_0 of the single membrane system is defined by:

$$\begin{array}{l} q_0(A) = 1.3;\\ q_0(C) = 0.01;\\ q_0(X) = 0.01;\\ q_0(X^+) = 0.99;\\ q_0(M) = 0.01;\\ q_0(M^+) = 0.99; \end{array}$$

in which we deal with concentrations of species, rather than with objects, and in this way the initial amounts are real numbers;

where, for each element of T the reaction map of inertia rules is set to 1600.

We start the modeling session by opening the Psim's main interface showed in Figure 4. This window allows the user to manage all the experiment's stages. In particular the main possible choices involve:

- 1. modelling the system, setting substances, initial conditions, reaction maps and rules;
- 2. starting the simulation;
- 3. displaying output charts.

The first step to consider while setting up a system's simulation is the specification of the corresponding MP graph. In what follows, some steps towards the creation of an MP graph modelling the mitotic oscillator are presented.

After clicking on the *New Experiment* label of the *File* menu, a window like the one depicted in the Figure 5 appears. This is the main window of the graphical interface allowing the user to input in a easy way the MP graph components by simply dragging them from the upper toolbar to the bottom white panel. This task is performed by using the following toolbar icons:

• The *blue circle*: adds a new *type node* that stores the name of a substance, its initial number of molar units and its inertia value (as explained in previous papers, inertias are a way to represent the fact that not all reactants need to

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٤.	SIM v2.4								
File	Simulation	Engine	Evolution	Input Plugins	Experiment Plugins	Output	Plugins	?	
ſ	-System mod	lel informa	ation				-Model o	description	
	Model nam	ie: no na	me						
	Membrane	s: unkna	wn						
	Rules:	unkna	wn						
	Objects:	unkna	wn		Edit mode				
S	how								
	Obje	ects		Rules	Environment				
s	imulation stat	te							
					Results				

Fig. 4. The Psim's main interface

PSim Regulated Stoichiometric Network input GUI Fia. 5th. You	
₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽₽	00
New organism No description Environment measures Name Value Measure unit	
None none none	

Fig. 5. Psim's input interface

react at a certain instant, they are a sort of resistance opposed to species to performing reactions).

- The *black circle*: adds a new *metabolic reaction node* that represents a reaction channel between interacting substances and stores the name of a reaction rule.
- The *red rectangle*: adds a new *reactivity node* building the regulatory part of MP graphs. In the simulator, reactivity nodes store the reactivity map function corresponding to the connected rule and, if necessary, a boolean guard function for the rule activation.

• The green triangles: add input gates and output gates nodes that identify rules which respectively introduce new matter in the membrane or expel part of it from the system.

After the insertion of the nodes in the white panel the user can specify their internal parameters and connect the nodes by drawing arcs among them. The best way to accomplish this task is to start by defining the *type node* parameters and the *metabolic reaction node* parameters by double clicking on the corresponding nodes and filling in the window's field that automatically appears (Figure 6). Importantly enough, a parser has been implemented to check the consistency of inserted parameters and to alert the user if any irregularity arose.

TypeN_0 properties ed	litor		X
	Type node name:	c	
	Number of molar units:	0.01	
TIK	Inertia value:	1600	
	No errors	s found while checking user inputs	
	Submit	Cancel	

Fig. 6. Insertion of the type node parameters

At this point, one can connect the *type nodes* and the *metabolic reaction nodes* with each other by drawing arcs among them with the simple use of the mouse. This is a very important step because it allows to represent the stoichiometric part of the system by means of the MP graph topology (Figure 7).

As an example, let us consider the reaction $r_2 : C \to X$. Within the input graphical interface it is represented by the R2 black circle that is connected by means of black arcs to the C and the X blue circles, representing the corresponding substances; the direction of the arrow represents the substance flow of the reaction.

A further modeling step is needed to add the *reactivity nodes* describing the regulatory part of the system. This can be done by first linking every *type node*, that affects a reaction map, with the corresponding *reactivity nodes* (as showed in Figure 8). Finally the reactivity map function of every reactivity node is specified by using the linked *type nodes* and the *environmental measures* as variables or constants (as reported in Figure 9). Figure 8 represents the final mitotic oscillator MP graph as produced by the Psim GUI.

This completes the modelling stage and the next logical step is to start the simulation of the specified system. This is done by clicking on the rightmost icon of the upper toolbar (the rightward arrow). The click causes a small window to pop out, in which it is possible to set the number of steps the simulation will span (Figure 10). A possible choice for this system is to run 150000 steps. By click on the *Start* button the dynamics computation begins.

When the simulation is finished the system prompts that results are available and ready to be visualized by the *Psim chart visualization form* (Figure 11). Using

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Fig. 7. Adding type nodes, metabolic reaction nodes and drawing arches among them



Fig. 8. An MP graph that models the mitotic oscillator

Reactivity_node_7 prop	perties editor		X
	Reactivity node name:	Reactivity_node_7	
	Reactivity map:	(0.25*X)/(0.01+C)	
	Guard:	true	
	No errors	s found while checking user inputs	
	Submit	Cancel	

Fig. 9. Reactivity node input parameter window

Simulation	.4 simulation parameters
	New organism simulation
Engine: Steps	default 150000 flooring
	Start Cancel

Fig. 10. Set the number of steps for the simulation to 150000

the bottom panel check boxes it is possible to decide elements to be displayed. In the considered case oscillations of cyclin C (red line), active cdc2 kinase M (blue line) and active protease M (green line) are displayed but the phase space plot can be drawn as well.



Fig. 11. Oscillations of the mitotic systems as calculated by Psim.

We finally highlight an important mechanism of the Psim platform: plugins extensibility. As already mentioned, plugins allow the user to enhance the main Psim computing core with powerful functionalities for the data import, export, control and analysis. An example under development is the *experiment plugin* that stores the experiment state (concentration of the substances and environmental measures) every x steps, where x is a parameter set by the user before the computation starts. This plugin could save, for instance, an XML file for every state, allowing the user to export the experiment samples in an standard way.

A software developer generates the plugin code (basically some Java classes) relying on the Psim's *JavaDoc* documentation obtainable at [15] which lists the *experiment plugin* methods to be mandatorily implemented. Plugin classes are

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meant to be archived in a Jar file and placed in a proper *plugins* directory. Provided this, at the following start up Psim will automatically find and load all the plugins contained in the *plugins* directory. The user can find all available experiment plugin statements in the main interface *Experiment Plugin* menu (Figure 4) in the form of a list. By clicking on the relative label its possible to activate the plugin that will be run at each step of the subsequent simulation and will save the state every x steps chosen by the user filling in a plugin popup window.

The plugin just described yields a set of XML files but the same principles can be extended also to the other kinds of plugins (*input*, *output*, *engine plugins*).

A particular mention is deserved by *engine plugins* that allow to implement new simulation strategies which can be different from the *metabolic algorithm* described above. This gives the simulation tool a very high flexibility as well as extendability as discussed previously.

4 Conclusion and further work

P systems can be useful frameworks to embed biological systems models in. This demands for some modifications to the classical definition of P systems and particularly a biologically meaningful evolution strategy is needed. In previous papers an essentially deterministic strategy, called metabolic algorithm, for the calculation of biological systems dynamics has been provided as well as an extension of the classic model of P systems, known as MP systems, focused on the dynamics of bio-systems. Moreover, all data needed for the simulation of MP systems dynamics can be provided by means of a graphical formalism known as MP graphs.

The basics of MP systems have been briefly revisited in this paper and based on them a simulation tool called Psim has been highlighted, together with a case study of a well known and previously investigated model of mitotic cycles in early amphibians. Psim v. 2.4 is the latest release of the MP systems simulator developed within the MNC Group of the University of Verona and it has very interesting features such as the plugin mechanism and the meta-engine architecture which give the tool an high level of extendability and personalization. In particular, plugins can be useful to perform several tasks such as data import/export, control of the simulation flow, output of dynamics obtained and analysis of the results among others. Moreover, the meta-engine architecture of the simulator allows users to define their own evolution strategies by implementing some fixed interfaces of the simulator.

In the future we plan to enrich the core of this simulation tool by implementing a series of plugins such as the one described above to have a snapshot of the state of the system in particular instants. Other plugins under investigation involve some automatic procedures for parameter estimation given suitable observations of the reality to be modelled. Finally, we plan to employ this simulation tool for the calculation of the dynamics of systems not already modelled and in this respect the possibility to devise ad-hoc evolution strategies can be very important to tackle some specific issues related with the particular reality to be modelled.

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Advances in Modeling the Dynamics of HIV Infection with Conformon-P Systems

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Summary. Further results on the study of the dynamics of HIV infection with grids of conformon-P systems are reported. This study clearly shows a subdivision in two main phases, the first faster than the second, of the mechanism at the base of the considered dynamics.

1 Introduction

The infection by the human immuno-deficiency virus (HIV), the cause of acquired immunodeficiency syndrome (AIDS), has been widely studied both in the laboratory and with computer models in order to understand the different aspects that regulate the virus-host interaction.

Several mathematical models have been proposed (for example [12, 18, 10]) but all of them fail to describe some aspects of the infection. The recent model reported by Dos Santos & Coutinho in [14], based on cellular automata, clearly shows the different time scales of the infection and has a broad qualitative agreement to the density of healthy and infected cells observed *in vivo*. However, in [15] it is noted that this qualitative agreement is reached only if some parameters are chosen in a small interval. If some of the parameters are chosen outside this interval, then the model of [14] does not follow the dynamics of what is observed *in vivo*.

In the present paper we continue our study on the modeling of the dynamics of HIV infection with grids of conformon-P systems started in [2]. There our model proved to be robust than the cellular automata model of [14] to a wide range of conditions and parameters, with more reproducible qualitative agreement to the overall dynamics and to the densities of healthy and infected cells observed *in vivo*.

2 The Modeling Platforms

2.1 Cellular automata

Cellular Automata (CA) are a regularly used platform for modeling, and are increasingly explored as modeling tools in the context of natural phenomena that exhibit characteristic spatiotemporal dynamics [16, 3]. Of interest here, for example, are their use in modeling the spread of infection [1, 11, 14, 10, 17].

A CA consists of a finite number of cells (invariably arranged in a regular spatial grid), each of which can be in one of a finite (typically small) number of specific states. In the usual approach, at each time step t the status of the CA is characterized by its state vector; that is, the state of each of the cells. In the simplest type of CA, the state vector at time t + 1 is obtained from that at time t by the operation of a single rule applied in parallel (synchronously) to each cell. The rule specifies how the state of a cell changes as a function of its current state and the states of the cells in its neighborhood (see Figure 4). In many applications, including that addressed here, it is appropriate for the rule to be probabilistic.

The straightforward nature of the time evolution of a CA, combined with its emphasis on local interactions, has made it an accessible and attractive tool for modeling many biological processes.

2.2 Conformon-P systems

Conformon-P systems (cP systems) [5] have been introduced as a novel computational device (P systems are the chief systems arising in the emerging research area of Membrane Computing [13]) whose early inspiration comes from a theoretical model of the living cell.

CP systems are defined in an extremely simple way that does not limit either their computational power, or their modeling capabilities. As a variant of P systems, they capture the dynamics of interacting processes in a novel way, using constructs that characterize the flow of information between regions in a range of cell-like topological structures. Moreover, their definition allows them to model different kinds of process (a compartment defines locality in general, it is not necessarily a membrane compartment in a cell) and to integrate several degrees of abstraction in the same system.

P systems are well-defined models of parallel computational systems that have a rich and growing base [19] of theoretical understanding of their properties.

A cP system has conformons, a name-value pair, as objects. If V is an alphabet (a finite set of letters) and \mathbb{N}_0 is the set of natural numbers (with 0 included), then we can define a conformon as $[\gamma, a]$, where $\gamma \in V$ and $a \in \mathbb{N}_0$, we will say that γ is the *name* and a is the *value* of the conformon $[\gamma, a]$. If, for instance, V = A, B, C, \ldots, Z , then [A, 5], [C, 0], [Z, 14] are conformons, while [AB, 21], [C, -15], and [D, 0.5] are not.

Two conformons can interact according to an *interaction rule*. An interaction rule is of the form $\gamma \xrightarrow{n} \beta$, where $\gamma, \beta \in V$ and $n \in \mathbb{N}_0$, and it says that a conformon

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with name γ can give *n* from its value to the value of a conformon having name β . A rule can be applied only if the value of the conformon with name γ is greater or equal to *n*. If, for instance, there are conformons [G, 5] and [R, 9] and the rule $G \xrightarrow{3} R$, the application of *r* leads to [G, 2] and [R, 12].

The (membrane) compartments present in a cP system have a label (it is a name which makes it easier to refer to a compartment), every label being different. Compartments can be unidirectionally connected to each other and for each connection there is a *predicate*. A predicate is an element of the set $\{\geq n, \leq n \mid n \in \mathbb{N}_0\}$. Examples of predicates are: $\geq 5, \leq 2$, etc.. A connection and its predicate are referred as *passage rules*. If, for instance, there are two compartments (with labels) m_1 and m_2 and there is a passage rule from m_1 to m_2 having predicate ≥ 4 , then conformons having value greater or equal to 4 can pass from m_1 to m_2 . In a time unit any number of conformons can move between two connected membranes as long as the predicate of the passage rule is satisfied. Notice that we have *unidirectional passage rules* that is: m_1 connected to m_2 does not imply that m_2 is connected to m_1 . Moreover, each passage rule has its own predicate. If, for instance, m_1 is connected to m_2 and m_2 is connected to m_1 , the two connections can have different predicates.

A simple cP system is illustrated in Figure 1.



Fig. 1. A cP system

CP systems do not work under the requirement of maximal parallelism, typical to the majority of the models of P systems. When used as modeling platform cP systems can be classified as stochastic descriptive dynamic discrete model based on a discrete spatial heterogeneity. CP systems have been successfully used to model biological processes [7, 2].

A grid of cP systems (Figure 2) is composed by cells, each cell being a simple conformon-P system connected to some other cells, the neighborhood of the cell.

Ongoing research is establishing the computational properties of (models of) cP systems [8, 9, 5, 6, 4].

CP systems can contain *modules*: groups of membranes with conformons and interaction rules able to perform a specific task. The task performed by a module



Fig. 2. A grid of cP systems

can be considered atomic (i.e., completed in one time unit) in the context of the cP system containing it. Modules allow cP systems to be scalable.

Some modules are: Splitter, Separator, Decreaser/Increaser [5]. The combination of Separators and Decreaser/Increaser allows us to define strict interaction rule: $\gamma^{(a)} \stackrel{c}{\rightarrow} \beta_{(b)}$ where $\gamma, \beta \in V$, $a, b, c \in \mathbb{N}_0$, meaning that a conformon with name γ can interact with β passing just c only if the value of γ and β before the interaction is a and b respectively. Notice that in a strict interaction just c is passed even if the value of γ could be decreased by any multiple of c. Interactions of the kind $\gamma \stackrel{c}{\rightarrow} \beta_{(b)}$ (before the interaction γ can have any value while β has b as value) and $\gamma^{(a)} \stackrel{c}{\rightarrow} \beta$ (before the interaction γ has a as value while β can have any value) can be defined, too.

3 The Process and the Models

The dynamics observed in HIV infections can be divided into three phases. Initially the amount of virus in the host grows in an exponential way, then the viral load drops to what is known as the "set point". Finally the amount of virus in the host increases slowly, accelerating near the onset of AIDS. The first two phases occur in the first weeks following the infection; the third phase can last years. This is plotted in Figure 3 where each unit in the x axes represent a week in time.

In [14] this process was modeled with a CA in which each cell could be in any of four possible states: *healthy*, *A-infected*, *AA-infected*, and *dead*. In the (random) initial configuration a cell had probability p_{HIV} to be *A-infected*, otherwise it is *healthy*.

The rules used in [14] are:

1. if an *healthy* cell has at least one *A1-infected* neighbor, then it becomes *A1-infected*;

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Fig. 3. Typical dynamics of HIV infection.

- 2. if an *healthy* cell has not A1-infected neighbors but it has at least R A2-infected neighbors, then it becomes A1-infected;
- 3. an A1-infected cell becomes A2-infected after τ time steps;
- 4. A2-infected cells become dead cells;
- 5. dead cells can become (be replaced by) healthy cells with probability p_{repl} ;
- 6. newly introduced *healthy* cells can become A1-infected with probability p_{infec} .

The biological reasoning behind these rules is explained in [14]. Essentially, rules 1 and 2 model the basic spread of viral infection from cells to neighboring
cells; rules 3-5 model the short life of an infected cell, and rule 6 models the body's continual replenishment of new healthy cells but maintaining a small probability of infection.

In [14] the following parameters were chosen: $p_{HIV} = 0.05$, $p_{repl} = 0.99$, $p_{infec} = 10^{-5}$, R = 5 and $\tau = 4$. They experimented with grids of size ranging from 300×300 to 1000×1000 , and the averaged results of 500 simulations reported in [14] on toroidal grids ranging from 700×700 show a qualitative agreement to the density of healthy and infected cells observed *in vivo*.

In [15] it is shown that this qualitative agreement is reached only for values of the parameters close to the ones just indicated. If either $p_{HIV} < 10^{-2}$ or p_{infec} is chosen in the range 10^{-2} to 10^{-4} , then the CA model of [14] does not follow the dynamics of what is observed *in vivo*.

3.1 The CA model

3.2 The grid of cP system model

The main difference that our model has in respect to the one reported in [14] is that the interaction rules are divided in two subsets: *part 1* and *part 2* (see Appendix A). The rules in the two subsets differ in the probabilities associated to them.

Other differences as, for instance, the presence of *pre-dead* cells, exist in order to simulate in terms of operations in a cP system some instructions of the CA presented in [14].

Each cell can be in one of five states: 1-healthy, A-infected, AA-infected, predead, and dead (in respect to the rules in part 1) identified by the presence of the conformons: [H, 1], [A, 1], [AA, 1], [PD, 1], and [D, 1] respectively. If, for instance, a cell is in an healthy state, then it will contain [H, 1], [A, 0], [AA, 0], [PD, 0], and [D, 0] (similarly for the other cases). In the initial configuration, each cell contains the conformons [R, 1], [V, 10], [E, 0], and [W, 0] are present in an unbounded number of copies.

In the following we consider and describe the rules in *part 1*.

If a cell is A-infected, then it can generate [V, 11] (meaning: if a cell is A-infected it can generate a virus). This is performed by the rules:

1:
$$R \xrightarrow{1} A_{(1)}$$
 2: $A^{(2)} \xrightarrow{1} V_{(10)}$

Notice that [V, 10] does not represent a virus, but [V, 11] does.

[V, 11] conformons can pass from a cell to any other in its neighborhood (meaning: viruses can spread between cells).

An *1-healthy* cell can become *A-infected* if it contains a virus. This is performed by the rules:

3: $V \xrightarrow{11} H_{(1)}$ **4:** $H^{(12)} \xrightarrow{12} A_{(0)}$ **5:** $A^{(12)} \xrightarrow{11} W_{(0)}$

An AA-infected cell can generate [E, 1] conformons. These conformons can pass to other cells and interact such that [E, 4] conformons are created. When a [E, 4]conformon is present in an *healthy* cell, then it can become A-infected. This process mimics rule II in Section 3 and it is performed by:

and by the fact that [E, 1] can pass from one cell to any other in its neighborhood. From the rules 7, 8, and 9 it should be clear that only [E, 1], [E, 2], and [E, 4] can be present in the system. Because of rule 6 an *AA-infected* cell can generate [E, 1]. When two [E, 1] are present in the same cell they can interact to create [E, 2] (rule 8) and two [E, 2] present in the same cell can interact to create [E, 4] (rule 9). If the creation of [E, 4] took place in an *healthy* cell, then this cell can become *A-infected* (rules 10, 11 and 12).

An *A-infected* cell can become *AA-infected* by the application of the rule:

13:
$$A^{(1)} \xrightarrow{1} AA_{(0)}$$

An AA-infected cell can become dead. Before doing so it goes into the predead state in which the [V, 11], [E, 1], [E, 2], and [E, 4] conformons present in it are removed. This is performed by the rules:

14:
$$AA^{(11)} \xrightarrow{1} PD_{(0)}$$
 15: $V^{(11)} \xrightarrow{1} PD_{(1)}$ **16:** $E \xrightarrow{1} PD_{(1)}$ **17:** $E \xrightarrow{2} PD_{(1)}$
18: $E \xrightarrow{4} PD_{(1)}$ **19:** $PD^{(1)} \xrightarrow{1} D_{(0)}$ **20:** $PD^{(2)} \xrightarrow{1} W_{(0)}$ **21:** $PD^{(3)} \xrightarrow{2} W_{(0)}$
22: $PD^{(5)} \xrightarrow{4} W_{(0)}$

A *dead* cell can become 2-healthy cell by the application of the rule

23:
$$D^{(1)} \xrightarrow{1} H2_{(0)}$$

The R and W conformons do not have a direct relationship with any aspect of HIV infection. In broad terms, the R conformons can be regarded as 'food' molecules needed by a cell in a certain state to perform an action (for instance, if *A-infected* to generate a virus). The W conformons can be regarded as 'waste' molecules, to which some conformons can pass part of their value. As W conformons only receive values from other conformons, their initial value is not relevant for the simulation.

The state 2-healthy, together with A2-infected, AA2-infected, 2-pre-dead, and 2-dead are managed by the rules in part 2. The rules in part 2 are similar to the ones in part 1 but they have H2 instead of H, A2 instead of A, AA2 instead of AA, PD2 instead of PD, and D2 instead of D.

In the diagrams related to the grid of cP systems the curve of *healthy* cells is obtained adding up the number of H and H2 cells; the curve of *infected* cells is obtained adding up the number of A, AA, A2 and AA2 cells; the curve of *dead* cells is obtained adding up the number of D, PD, D2 and PD2 cells.

The interaction rules indicated in Appendix A can be logically divided in two sets: *state-change* and *internal dynamics*. The *state-change* rules allow the cells to pass from a state to another. For instance, rule 4 is a state change rule as when it is applied in a cell the state of the cell passed from *1-healthy* to *A-infected*. The *state-change* rules are: 4, 11, 13, 14, 19, 23, 27, 32, 34, 35, 40 and 44.

The remaining rules belong to *internal dynamics* as they do not directly effect the state of a cell.

Differently than what done in [2], in the present study the probabilities associated to the *internal dynamics* rules in phase 1 are equal to the ones in phase 2. The probabilities of the *state-change* rules in phase 1 are higher than then ones in phase 2.

Considering what we said in Section 3, rules in *part 1* model the behavior of the first two phases of the dynamics of HIV infection, while rules in *part 2* model the behavior of the third phase.

4 Experiments and Results

The simulations performed with the cP system were based on a toroidal 50×50 grid, using a Moore neighborhood (considering Figure 4 the black cell can pass conformons to any grey cell) and with $p_{HIV} = 0.05$.



Fig. 4. The Moore neighborhood.

All the 10 simulations (with different random number sequences) run for 16000 iterations and they all show a dynamics very similar to the one observed *in vivo*. A typical outcome is depicted in Figure 5.

This outcome (even if run for only one kind of neighborhood and one values of p_{HIV}) fits the dynamics observed *in vivo* better than the outcomes reported in [2]:

- the tempo of the dynamics is constant during the simulation. In [2] the dynamics was 'too fast' in the later years (or 'too slow' in the first weeks). In the present study 1 year corresponds to 1560 iterations. This means that phase I and phase II (both taking place in at most 10 weeks) should correspond to 300 iterations. In this way the 16000 iterations of out tests corresponds to a bit more than 10 years.
- the percentage of *healthy* and *infected* cells in phase III is closer to what observed *in vivo* than what reported in [2].
- the dynamics of *healthy* and *infected* cells in phase III is not flat as in [2] but shows a concavity similar to the one observed *in vivo*.



Fig. 5. Typical outcome for grids of cP systems.

There are two major differences between the dynamics obtained by us and the one observed *in vivo*:

- in phase III the number of *healthy* cells should become equal to the one of *dead* cells;
- the curves followed by the number of $healthy \ {\rm and} \ infected \ {\rm cells}$ in phase III do not change concavity.

5 Final Remarks

We consider the reported study still in its initial phases. In the future we will try to better fit the dynamics obtained with grids of cP system to what observer *in vivo* and we will run the tests on different neighborhoods and different values of p_{HIV} (as done in [2]).

Some results obtained by us indicates that the E conformons play a negligible role in the whole dynamics. On this base we will try to simply our model in the number of interaction rules and conformons present in it.

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	part 1			part 2	
label	rule	prob.	label	rule	prob.
1	$R \xrightarrow{1} A_{(1)}$	0.7071	24	$R \xrightarrow{1} A2_{(1)}$	0.7071
2	$A^{(2)} \xrightarrow{1} V_{(10)}$	0.7071	25	$A2^{(2)} \xrightarrow{1} V_{(10)}$	0.7071
3	$V \xrightarrow{11} H_{(1)}$	0.79	26	$V \xrightarrow{11} H2_{(1)}$	0.79
4	$H^{(12)} \xrightarrow{12} A_{(0)}$	0.79	27	$H2^{(12)} \xrightarrow{12} A2_{(0)}$	0.0001
5	$A^{(12)} \xrightarrow{11} W_{(0)}$	0.79	28	$A2^{(12)} \xrightarrow{11} W_{(0)}$	0.79
6	$R \xrightarrow{1} AA_{(1)}$	0.7071	29	$R \xrightarrow{1} AA2_{(1)}$	0.7071
7	$AA^{(2)} \xrightarrow{1} E_{(0)}$	0.7071	30	$AA2^{(2)} \xrightarrow{1} E_{(0)}$	0.7071
8	$E^{(1)} \xrightarrow{1} E_{(1)}$	0.07071			
9	$E^{(2)} \xrightarrow{2} E_{(2)}$	0.07071			
10	$E \xrightarrow{4} H_{(1)}$	0.79	31	$E \xrightarrow{4} H2_{(1)}$	0.79
11	$H^{(5)} \xrightarrow{5} A_{(0)}$	0.79	32	$H2^{(5)} \xrightarrow{5} A2_{(0)}$	0.0001
12	$A^{(5)} \xrightarrow{4} W_{(0)}$	0.79	33	$A2^{(5)} \xrightarrow{4} W_{(0)}$	0.79
13	$A^{(1)} \xrightarrow{1} AA_{(0)}$	0.04	34	$A2^{(1)} \xrightarrow{1} AA2_{(0)}$	0.0001
14	$AA^{(11)} \xrightarrow{1} PD_{(0)}$	0.1	35	$AA2^{(11)} \xrightarrow{1} PD2_{(0)}$	0.00075
15	$V^{(11)} \xrightarrow{1} PD_{(1)}$	0.7071	36	$V^{(11)} \xrightarrow{1} PD2_{(1)}$	0.7071
16	$E \xrightarrow{1} PD_{(1)}$	0.7071	37	$E \xrightarrow{1} PD2_{(1)}$	0.7071
17	$E \xrightarrow{2} PD_{(1)}$	0.7071	38	$E \xrightarrow{2} PD2_{(1)}$	0.7071
18	$E \xrightarrow{4} PD_{(1)}$	0.7071	39	$E \xrightarrow{4} PD2_{(1)}$	0.7071
19	$PD^{(1)} \xrightarrow{1} D_{(0)}$	0.2	40	$PD2^{(1)} \xrightarrow{1} D2_{(0)}$	0.001
20	$PD^{(2)} \xrightarrow{1} W_{(0)}$	0.7071	41	$PD2^{(2)} \xrightarrow{1} W_{(0)}$	0.7071
21	$PD^{(3)} \xrightarrow{2} W_{(0)}$	0.7071	42	$PD2^{(3)} \xrightarrow{2} W_{(0)}$	0.7071
22	$PD^{(5)} \xrightarrow{4} W_{(0)}$	0.7071	43	$PD2^{(5)} \xrightarrow{4} W_{(0)}$	0.7071
23	$D^{(1)} \xrightarrow{1} H2_{(0)}$	0.1	44	$D2^{(1)} \xrightarrow{1} H2_{(0)}$	0.001

A Rules, links, and probabilities

Links:

 $\left[V,11\right]$ can pass with probability 1 from any cell to any of its neighbors; $\left[E,1\right]$ can pass with probability 0.01 from any cell to any of its neighbors.

Quantum (UREM) P Systems: Background, Definition and Computational Power

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Summary. Quantum UREM P systems constitute an attempt to introduce in Membrane Computing notions and techniques deriving from quantum mechanics. As we will see, the approach we have adopted is different from what is usually done in Quantum Computing; in fact, we have been inspired by the functioning of some elementary operations that are used in quantum mechanics to exchange quanta of energy among quantum systems: creation and annihilation operators. In this paper we will provide the background which has led to the current definition of quantum UREM P systems, and we will recall some results concerning their computational power.

1 The Quest for Quantum P Systems

Membrane systems (also known as P systems) have been introduced by Gheorghe Păun in 1998 [27] as a new class of distributed and parallel computing devices, inspired by the structure and functioning of living cells. The basic model consists of a hierarchical structure composed by several membranes, embedded into a main membrane called the *skin*. Membranes divide the Euclidean space into *regions*, that contain some *objects* (represented by symbols of an alphabet) and *evolution rules*. Using these rules, the objects may evolve and/or move from a region to a neighboring one. A *computation* starts from an initial configuration of the system and terminates when no evolution rule can be applied. Usually, the result of a computation is the multiset of objects contained into an *output membrane* or emitted from the skin of the system.

In what follows we assume the reader is already familiar with the basic notions and the terminology underlying P systems. For a layman-oriented introduction to P systems see [29], whereas for a systematic introduction we refer the reader to [28]. The latest information about P systems can be found in [32].

At the beginning of 2004, the Membrane Computing community started to query about the possibility to define a quantum version of P systems, and hence we started to work on the subject. A first paper [21] was presented in Palma de

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Mallorca in November 2004. There, we proposed two options: either to follow the steps usually performed in Quantum Computing to define the quantum version of a given computation device, or to propose a completely new computation device which is based on the most elementary operation which can be conceived in physics: the exchange of a quantum of energy among two quantum systems. In the former case we would have obtained yet another quantum computation device whose computation steps are defined as the action of unitary operators, whose computations are logically reversible, and in which there are severe constraints on the amount of information which can be extracted from the system by measuring its state. In the latter case, instead, we felt that a new and interesting computation device could be introduced. Indeed, after a long and careful investigation, we decided to adopt creation and annihilation operators as the most elementary operations which can be performed by our computation device.

It was since 2001 that several authors introduced the notion of energy in P systems [1, 10, 31, 15, 22, 23]. Hence, we looked at the literature to find a model of P systems that was easily transformable in a quantum computation device. Our first choice, explored in [21], was to focus on energy-based P systems, in which a given amount of energy is associated to each object of the system. Moreover, instances of a special symbol e are used to denote free energy units occurring into the regions of the system. These energy units can be used to transform objects, using appropriate rules. The rules are defined according to conservativeness considerations. Indeed, in [21] we proposed two different versions of quantum P systems based on this classical model. Both versions were defined just like classical energy-based P systems, but for objects and rules. Objects were represented as pure states in the Hilbert space \mathbb{C}^d , $d \geq 2$, whereas the definition of rules differs between the two models. In the former, rules are defined as bijective functions — implemented as unitary operators — which transform the objects from the alphabet. In the latter, rules are defined as generic functions which map the alphabet into itself. Such functions are implemented using a generalization of the Conditional Quantum Control technique [3], and may yield to non unitary operators (a fact which is usually seen with suspect in traditional Quantum Computing).

However, several problems were pointed out in [21], the most serious being that it is difficult to avoid unwanted exchanges of energy among the objects, that yield the system to unintended states. Another difficulty was tied to the assignment of the amount of energy to every object of the system. In the original definition of energy–based P system, every object incorporated a different amount of energy; in other words, the amount of energy uniquely determined the kind of object and, by acquiring or releasing energy from the environment, one object was transformed to another kind of object. Under this definition, we were able in [22] to simulate a single Fredkin gate. However, in order to simulate an entire Fredkin circuit [23, 24] we were forced to relax the definition, and allow different kinds of objects to have the same amount of energy, otherwise the number of different kinds of objects would have become unmanageable. Last, but not the least, we have the problem of objects localization and control. How do we force an object to stay in a given region for a long time, or to move to the desired region? Indeed, one notable feature of quantum systems is the so called "tunnel effect", thanks to which at every moment we have a positive probability that the object spontaneously leaves the current region. Of course this is a problem which is generally found when trying to control the behavior of a quantum system, but in the case of energy– based P systems the problem is exacerbated by the fact that the objects should move (only) as the effect of the application of a rule. This problem does not occur with quantum UREM P systems: there the objects interact by exchanging some amounts of energy, which are stored in quantum harmonic oscillators, but never move nor cross any membrane.

Looking for some alternatives, we considered the model of P systems introduced in [11], in which a non-negative integer value is assigned to each membrane. Such a value can be conveniently interpreted as the *energy* of the membrane. In these P systems, rules are assigned to the membranes rather than to the regions of the system. Every rule has the form $(in_i : \alpha, \Delta e, \beta)$ or $(out_i : \alpha, \Delta e, \beta)$, where i is the number of the membrane in a one-to-one labeling, α and β are symbols of the alphabet and Δe is a (possibly negative) integer number. The rule $(in_i : \alpha, \Delta e, \beta)$ is interpreted as follows: if a copy of α is in the region immediately surrounding membrane i, then this object crosses membrane i, is transformed to β , and modifies the energy of membrane i from the current value e_i to the new value $e_i + \Delta e$. Similarly, the rule $(out_i : \alpha, \Delta e, \beta)$ is interpreted as follows: if a copy of α is in the region surrounded by membrane i, then this object crosses membrane i, is transformed to β , and modifies the energy of membrane *i* from the current value e_i to the new value $e_i + \Delta e$. Both kinds of rules can be applied only if $e_i + \Delta e$ is non-negative. Since these rules transform one copy of an object to (one copy of) another object, in [11] they are referred to as *unit* rules. Hence, for conciseness, this model of P systems with unit rules and energy assigned to membranes is usually abbreviated as UREM P systems. An important observation is that in [11] the rules of UREM P systems are applied in a sequential way: at each computation step, one rule is selected from the pool of currently active rules, and it is applied. In [11] it has been proved that if we assign some local (that is, affecting only the membrane in which they are defined) priorities to the rules then UREM P systems are Turing complete, whereas if we omit the priorities then we do not get systems with universal computational power: indeed, we obtain a characterization of $PsMAT^{\lambda}$, the family of Parikh sets generated by context-free matrix grammars (without occurrence checking and with λ -rules).

So, finally, in [20] a quantum version of UREM P systems has been introduced, and it has been shown that such a model of computation is able to compute every partial recursive function (that is, it reaches the computational power of Turing machines) without the need to assign any priority between the rules of the system. In quantum UREM P systems, the rules $(in_i : \alpha, \Delta e, \beta)$ and $(out_i : \alpha, \Delta e, \beta)$ are realized through (not necessarily unitary) linear operators, which can be expressed as an appropriate composition of a truncated version of creation and annihilation operators. The operators which correspond to the rules have the form $|\beta\rangle \langle \alpha| \otimes O$, where O is a linear operator which modifies the energy associated with the membrane (implemented as the state of a truncated quantum harmonic oscillator).

In [20] also Quantum Register Machines (QRMs, for short) have been introduced. It is our opinion that they could play the same role in proofs concerning the computational power of quantum computation devices as played by classical register machines for classical computing devices. Indeed, it has been shown in [20] that they are able to simulate any classical (deterministic) register machine, and hence they are (at least) Turing complete. The advantage of quantum UREM P systems over QRMs is that, due to the locality of interactions, the operators which correspond to the rules of the former are generally smaller than the operators corresponding to the instructions of the latter.

Finally, in [19] we have shown that, under the assumption that an external observer is able to discriminate a null vector from a non-null vector, the NP-complete problem 3-SAT can be solved using quantum (Fredkin) circuits, quantum register machines and quantum UREM P systems. Precisely, for each type of computation device we have proposed a *brute force* technique that exploits quantum parallelism (as well as the ability to alter quantum states by using creation and annihilation operators) to explore the whole space of assignments to the boolean variables of any given instance ϕ of 3-SAT, in order to determine whether at least one of such assignments satisfies ϕ . The solutions are presented in the so-called *semi-uniform* setting, which means that for every instance ϕ of 3-SAT a specific computation device (circuit, register machine or UREM P system) that solves it is built. Even if it is not formally proved, it is apparent that the proposed constructions can be performed in polynomial time by a classical deterministic Turing machine (whose output is a "reasonable" encoding of the machine, in the sense given in [16]).

In the rest of the paper we overview the basic notions of quantum mechanics which have led to the definition of quantum UREM P systems, and the results obtained so far about their computational power. Precisely, in section 2 we recall some basic notions on quantum computers, and we extend them to quantum systems which are able to assume a generic number $d \geq 2$ of base states. We also introduce some operators which can be used to operate on the states of such systems; for these operators, we first give a mathematical description and then we propose some possible physical interpretations. In sections 3 and 4 we give the precise definitions of both classical and quantum register machines and UREM P systems, respectively. In section 5 we prove that quantum UREM P systems are able to compute any partial recursive function, and hence they are (at least) as powerful as Turing machines. In section 6 we show how to build two families of quantum register machines and quantum UREM P systems, respectively, that solve (in the semi–uniform setting) the **NP**–complete decision problem 3-SAT. The conclusions, as well as some directions for future research, are given in section 7.

2 Quantum computers

From an abstract point of view, a quantum computer can be considered as made up of interacting parts. The elementary units (memory cells) that compose these parts are two-levels quantum systems called *qubits*. A qubit is typically implemented using the energy levels of a two-levels atom, or the two spin states of a $\text{spin}-\frac{1}{2}$ atomic nucleus, or a polarization photon. The mathematical description — independent of the practical realization — of a single qubit is based on the two-dimensional complex Hilbert space \mathbb{C}^2 . The boolean truth values 0 and 1 are represented in this framework by the unit vectors of the canonical orthonormal basis, called the *computational basis* of \mathbb{C}^2 :

$$|0\rangle = \begin{bmatrix} 1\\ 0 \end{bmatrix} \qquad \qquad |1\rangle = \begin{bmatrix} 0\\ 1 \end{bmatrix}$$

Qubits are thus the quantum extension of the classical notion of bit, but whereas bits can only take two different values, 0 and 1, qubits are not confined to their two basis (also *pure*) states, $|0\rangle$ and $|1\rangle$, but can also exist in states which are coherent superpositions such as $\psi = c_0 |0\rangle + c_1 |1\rangle$, where c_0 and c_1 are complex numbers satisfying the condition $|c_0|^2 + |c_1|^2 = 1$. Performing a measurement of the state alters it. Indeed, performing a measurement on a qubit in the above superposition will return 0 with probability $|c_0|^2$ and 1 with probability $|c_1|^2$; the state of the qubit after the measurement (*post-measurement state*) will be $|0\rangle$ or $|1\rangle$, depending on the outcome.

A quantum register of size n (also called an n-register) is mathematically described by the Hilbert space $\otimes^n \mathbb{C}^2 = \underbrace{\mathbb{C}^2 \otimes \ldots \otimes \mathbb{C}^2}_{n \text{ times}}$, representing a set of n

qubits labeled by the index $i \in \{1, \ldots, n\}$. An *n*-configuration (also pattern) is a vector $|x_1\rangle \otimes \ldots \otimes |x_n\rangle \in \otimes^n \mathbb{C}^2$, usually written as $|x_1, \ldots, x_n\rangle$, considered as a quantum realization of the boolean tuple (x_1, \ldots, x_n) . Let us recall that the dimension of $\otimes^n \mathbb{C}^2$ is 2^n and that $\{|x_1, \ldots, x_n\rangle : x_i \in \{0, 1\}\}$ is an orthonormal basis of this space called the *n*-register computational basis.

Computations are performed as follows. Each qubit of a given *n*-register is prepared in some particular pure state ($|0\rangle$ or $|1\rangle$) in order to realize the required *n*-configuration $|x_1, \ldots, x_n\rangle$, quantum realization of an input boolean tuple of length *n*. Then, a linear operator $G : \otimes^n \mathbb{C}^2 \to \otimes^n \mathbb{C}^2$ is applied to the *n*-register. The application of *G* has the effect of transforming the *n*-configuration $|x_1, \ldots, x_n\rangle$ into a new *n*-configuration $G(|x_1, \ldots, x_n\rangle) = |y_1, \ldots, y_n\rangle$, which is the quantum realization of the output tuple of the computer. We interpret such modification as a computation step performed by the quantum computer. The action of the operator *G* on a superposition $\Phi = \sum c^{i_1 \ldots i_n} |x_{i_1}, \ldots, x_{i_n}\rangle$, expressed as a linear combination of the elements of the *n*-register basis, is obtained by linearity: $G(\Phi) = \sum c^{i_1 \ldots i_n} G(|x_{i_1}, \ldots, x_{i_n}\rangle)$. We recall that linear operators which act on *n*-registers can be represented as order 2^n square matrices of complex entries. Usually (but not in this paper) such operators, as well as the corresponding matrices, are required to be unitary. In particular, this implies that the implemented operations are logically reversible (an operation is *logically reversible* if its inputs can always be deduced from its outputs).

All these notions can be easily extended to quantum systems which have d > 2 pure states. In this setting, the *d*-valued versions of qubits are usually called *qudits* [17]. As it happens with qubits, a qudit is typically implemented using the energy levels of an atom or a nuclear spin. The mathematical description — independent of the practical realization — of a single qudit is based on the *d*-dimensional complex Hilbert space \mathbb{C}^d . In particular, the pure states $|0\rangle$, $\left|\frac{1}{d-1}\rangle$, $\left|\frac{2}{d-1}\rangle$, \ldots , $\left|\frac{d-2}{d-1}\rangle$, $|1\rangle$ are represented by the unit vectors of the canonical orthonormal basis, called the *computational basis* of \mathbb{C}^d :

$$|0\rangle = \begin{bmatrix} 1\\0\\\vdots\\0\\0 \end{bmatrix}, \quad \left|\frac{1}{d-1}\right\rangle = \begin{bmatrix} 0\\1\\\vdots\\0\\0 \end{bmatrix}, \quad \cdots, \quad \left|\frac{d-2}{d-1}\right\rangle = \begin{bmatrix} 0\\0\\\vdots\\1\\0 \end{bmatrix}, \quad |1\rangle = \begin{bmatrix} 0\\0\\\vdots\\1\\0 \end{bmatrix}$$

As before, a quantum register of size n can be defined as a collection of n qudits. It is mathematically described by the Hilbert space $\otimes^n \mathbb{C}^d$. An n-configuration is now a vector $|x_1\rangle \otimes \ldots \otimes |x_n\rangle \in \otimes^n \mathbb{C}^d$, simply written as $|x_1, \ldots, x_n\rangle$, for x_i running on $L_d = \left\{0, \frac{1}{d-1}, \frac{2}{d-1}, \ldots, \frac{d-2}{d-1}, 1\right\}$. An n-configuration can be viewed as the quantum realization of the "classical" tuple $(x_1, \ldots, x_n) \in L_d^n$. The dimension of $\otimes^n \mathbb{C}^d$ is d^n and the set $\{|x_1, \ldots, x_n\rangle : x_i \in L_d\}$ of all n-configurations is an orthonormal basis of this space, called the n-register computational basis. Notice that the set L_d can also be interpreted as a set of truth values, where 0 denotes falsity, 1 denotes truth and the other elements indicate different degrees of indefiniteness.

Let us now consider the set $\mathcal{E}_d = \left\{\varepsilon_0, \varepsilon_{\frac{1}{d-1}}, \varepsilon_{\frac{2}{d-1}}, \ldots, \varepsilon_{\frac{d-2}{d-1}}, \varepsilon_1\right\} \subseteq \mathbb{R}$ of real values; we can think to such quantities as energy values. To each element $v \in L_d$ we associate the energy level ε_v ; moreover, let us assume that the values of \mathcal{E}_d are all positive, equispaced, and ordered according to the corresponding objects: $0 < \varepsilon_0 < \varepsilon_{\frac{1}{d-1}} < \cdots < \varepsilon_{\frac{d-2}{d-1}} < \varepsilon_1$. If we denote by $\Delta \varepsilon$ the gap between two adjacent energy levels then the following linear relation holds:

$$\varepsilon_k = \varepsilon_0 + \Delta \varepsilon \left(d - 1 \right) k \qquad \forall k \in L_d \tag{1}$$

Notice that it is not required that $\varepsilon_0 = \Delta \varepsilon$. As explained in [21, 19], the values ε_k can be thought of as the energy eigenvalues of the infinite dimensional quantum harmonic oscillator truncated at the (d-1)-th excited level (see Figure 1), whose Hamiltonian on \mathbb{C}^d is:

$$H = \begin{bmatrix} \varepsilon_0 & 0 & \dots & 0 \\ 0 & \varepsilon_0 + \Delta \varepsilon & \dots & 0 \\ \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & \dots & \varepsilon_0 + (d-1)\Delta \varepsilon \end{bmatrix}$$
(2)



Fig. 1. Energy levels of the infinite dimensional (on the left) and of the truncated (on the right) quantum harmonic oscillator

The unit vector $|H = \varepsilon_k\rangle = \left|\frac{k}{d-1}\right\rangle$, for $k \in \{0, 1, \dots, d-1\}$, is the eigenvector of the state of energy $\varepsilon_0 + k\Delta\varepsilon$. To modify the state of a qudit we can use creation and annihilation operators on the Hilbert space \mathbb{C}^d , which are defined respectively as:

$$a^{\dagger} = \begin{bmatrix} 0 & 0 & \cdots & 0 & 0 \\ 1 & 0 & \cdots & 0 & 0 \\ 0 & \sqrt{2} & \cdots & 0 & 0 \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ 0 & 0 & \cdots & \sqrt{d-1} & 0 \end{bmatrix} \qquad a = \begin{bmatrix} 0 & 1 & 0 & \cdots & 0 \\ 0 & 0 & \sqrt{2} & \cdots & 0 \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & 0 & \cdots & \sqrt{d-1} \\ 0 & 0 & 0 & \cdots & 0 \end{bmatrix}$$

It is easily verified that the action of a^{\dagger} on the vectors of the canonical orthonormal basis of \mathbb{C}^d is the following:

$$a^{\dagger} \left| \frac{k}{d-1} \right\rangle = \sqrt{k+1} \left| \frac{k+1}{d-1} \right\rangle \qquad \text{for } k \in \{0, 1, \dots, d-2\}$$
$$a^{\dagger} \left| 1 \right\rangle = \mathbf{0}$$

whereas the action of a is:

$$a \left| \frac{k}{d-1} \right\rangle = \sqrt{k} \left| \frac{k-1}{d-1} \right\rangle \qquad \text{for } k \in \{1, 2, \dots, d-1\}$$
$$a \left| 0 \right\rangle = \mathbf{0}$$

Using a^{\dagger} and a we can also introduce the following operators:

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$$N = a^{\dagger}a = \begin{bmatrix} 0 & 0 & 0 & \cdots & 0 \\ 0 & 1 & 0 & \cdots & 0 \\ 0 & 0 & 2 & \cdots & 0 \\ \vdots & \vdots & \vdots & \ddots & \vdots \\ 0 & 0 & 0 & \cdots & d - 1 \end{bmatrix} \qquad aa^{\dagger} = \begin{bmatrix} 1 & 0 & \cdots & 0 & 0 \\ 0 & 2 & \cdots & 0 & 0 \\ \vdots & \vdots & \ddots & \vdots & \vdots \\ 0 & 0 & \cdots & d - 1 & 0 \\ 0 & 0 & \cdots & 0 & 0 \end{bmatrix}$$

The eigenvalues of the self-adjoint operator N are $0, 1, 2, \ldots, d-1$, and the eigenvector corresponding to the generic eigenvalue k is $|N = k\rangle = \left|\frac{k}{d-1}\right\rangle$. This corresponds to the notation adopted in [17], where the qudit base states are denoted by $|0\rangle, |1\rangle, \ldots, |d-1\rangle$, and it is assumed that a qudit can be in a superposition of the d base states:

$$c_0 |0\rangle + c_1 |1\rangle + \ldots + c_{d-1} |d-1\rangle$$

with $c_i \in \mathbb{C}$ for $i \in \{0, 1, \dots, d-1\}$, and $|c_0|^2 + |c_1|^2 + \dots + |c_{d-1}|^2 = 1$.

One possible physical interpretation of N is that it describes the number of particles of physical systems consisting of a maximum number of d-1 particles. In order to add a particle to the k particles state $|N = k\rangle$ (thus making it switch to the "next" state $|N = k + 1\rangle$) we apply the creation operator a^{\dagger} , while to remove a particle from this system (thus making it switch to the "previous" state $|N = k - 1\rangle$) we apply the annihilation operator a. Since the maximum number of particles that can be simultaneously in the system is d-1, the application of the system, and returns as a result the null vector. Analogously, the application of the annihilation operator to a a full d-1 particle system does not affect the system and returns the null vector as a result.

Another physical interpretation of operators a^{\dagger} and a, by operator N, follows from the possibility of expressing the Hamiltonian (2) as follows:

$$H = \varepsilon_0 \,\mathbb{I} + \Delta \varepsilon \, N = \varepsilon_0 \,\mathbb{I} + \Delta \varepsilon \, a^{\dagger} a$$

In this case a^{\dagger} (resp., a) realizes the transition from the eigenstate of energy $\varepsilon_k = \varepsilon_0 + k \Delta \varepsilon$ to the "next" (resp., "previous") eigenstate of energy $\varepsilon_{k+1} = \varepsilon_0 + (k+1) \Delta \varepsilon$ (resp., $\varepsilon_{k-1} = \varepsilon_0 + (k-1) \Delta \varepsilon$) for any $0 \leq k < d-1$ (resp., $0 < k \leq d-1$), while it collapses the last excited (resp., ground) state of energy $\varepsilon_0 + (d-1) \Delta \varepsilon$ (resp., ε_0) to the null vector.

The collection of all linear operators on \mathbb{C}^d is a d^2 -dimensional linear space whose canonical basis is:

$$\{E_{x,y} = |y\rangle \langle x| : x, y \in L_d\}$$

Since $E_{x,y} |x\rangle = |y\rangle$ and $E_{x,y} |z\rangle = 0$ for every $z \in L_d$ such that $z \neq x$, this operator transforms the unit vector $|x\rangle$ into the unit vector $|y\rangle$, collapsing all the other vectors of the canonical orthonormal basis of \mathbb{C}^d to the null vector. Each of the operators $E_{x,y}$ can be expressed, using the whole algebraic structure of the associative algebra of operators, as a suitable composition of creation and annihilation operators, as follows:

$$E_{\frac{i}{d-1},\frac{j}{d-1}} = \begin{cases} \frac{\sqrt{j!}}{(d-1)!} A_{a^{\dagger},a^{\dagger}}^{d-2,d-1-j,0} & \text{if } i = 0\\ \frac{\sqrt{j!}}{(d-1)!} A_{a,a^{\dagger}}^{d-1,d-1-j,0} & \text{if } i = 1 \text{ and } j \ge 1\\ \frac{\sqrt{i!}}{(d-1)!\sqrt{j!}} A_{a^{\dagger},a^{\dagger}}^{d-2-i,d-1,j} & \text{if } (i = 1, j = 0 \text{ and } d \ge 3) \text{ or }\\ (1 < i < d-2 \text{ and } j \le i)\\ \frac{\sqrt{j!}}{(d-1)!\sqrt{j!}} A_{a,a}^{i-1,d-1,d-1-j} & \text{if } (i = d-2, j = d-1 \text{ and } d \ge 3)\\ 0 & \text{or } (1 < i < d-2 \text{ and } j > i)\\ \frac{1}{\sqrt{(d-1)!j!(d-1)}} A_{a^{\dagger},a}^{d-1,j,0} & \text{if } i = d-2 \text{ and } j \le d-2\\ \frac{1}{\sqrt{(d-1)!j!}} A_{a,a}^{d-2,j,0} & \text{if } i = d-1 \end{cases}$$

Here we just recall, in order to keep the length of the paper under a reasonable size, that an alternative interpretation of qudits is possible, based on the values which can be assumed by the z component of the angular momentum of semi-integer spin quantum systems. Also with this interpretation every linear operator, and in particular operators $E_{x,y}$, can be realized as appropriate compositions of spin-rising (J_+) and spin-lowering (J_-) operators, similarly to what we have done with creation and annihilation operators. For the details, we refer the reader to [21, 19].

3 Classical and Quantum Register Machines

A (classical, deterministic) n-register machine is a construct $M = (n, P, l_0, l_h)$, where n is the number of registers, P is a finite set of instructions injectively labeled with a given set lab(M), l_0 is the label of the first instruction to be executed, and l_h is the label of the last instruction of P. Registers contain non-negative integer values. Without loss of generality, we can assume $lab(M) = \{1, 2, \ldots, m\}, l_0 = 1$ and $l_h = m$. The instructions of P have the following forms:

- j: (INC(r), k), with $j, k \in lab(M)$ This instruction increments the value contained in register r, and then jumps to instruction k.
- j: (DEC(r), k, l), with $j, k, l \in lab(M)$ If the value contained in register r is positive then decrement it and jump to instruction k. If the value of r is zero then jump to instruction l (without altering the contents of the register).

• m: Halt

Stop the machine. Note that this instruction can only be assigned to the final label m.

Register machines provide a simple universal computational model. Indeed, the results proved in [12] (based on the results established in [25]) as well as in [13] and [14] immediately lead to the following proposition.

Proposition 1. For any partial recursive function $f : \mathbb{N}^{\alpha} \to \mathbb{N}^{\beta}$ there exists a deterministic $(\max\{\alpha,\beta\}+2)$ -register machine M computing f in such a way

that, when starting with $(n_1, \ldots, n_{\alpha}) \in \mathbb{N}^{\alpha}$ in registers 1 to α , M has computed $f(n_1, \ldots, n_{\alpha}) = (r_1, \ldots, r_{\beta})$ if it halts in the final label l_h with registers 1 to β containing r_1 to r_{β} , and all other registers being empty; if the final label cannot be reached, then $f(n_1, \ldots, n_{\alpha})$ remains undefined.

A quantum *n*-register machine is defined exactly as in the classical case, as a four-tuple $M = (n, P, l_0, l_h)$. Each register of the machine can be associated to an infinite dimensional quantum harmonic oscillator capable to assume the base states $|\varepsilon_0\rangle$, $|\varepsilon_1\rangle$, $|\varepsilon_2\rangle$, ..., corresponding to its energy levels, as described in section 2. The program counter of the machine is instead realized through a quantum system capable to assume *m* different base states, from the set $\{|x\rangle : x \in L_m\}$. For simplicity, the instructions of *P* are denoted in the usual way:

$$j:(INC(i),k)$$
 and $j:(DEC(i),k,l)$

This time, however, these instructions are appropriate linear operators acting on the Hilbert space whose vectors describe the (global) state of M. Precisely, the instruction j: (INC(r), k) is defined as the operator

$$O_{j,r,k}^{INC} = \left| p_k \right\rangle \left\langle p_j \right| \otimes \left(\otimes^{r-1} \mathbb{I} \right) \otimes a^{\dagger} \otimes \left(\otimes^{n-r} \mathbb{I} \right)$$

with I the identity operator on \mathcal{H} (the Hilbert space in which the state vectors of the infinite dimensional quantum harmonic oscillators associated with the registers exist), whereas the instruction j : (DEC(r), k, l) is defined as the operator

$$\begin{split} O_{j,r,k,l}^{DEC} &= \left| p_l \right\rangle \left\langle p_j \right| \otimes \left(\otimes^{r-1} \mathbb{I} \right) \otimes \left| \varepsilon_0 \right\rangle \left\langle \varepsilon_0 \right| \otimes \left(\otimes^{n-r} \mathbb{I} \right) + \\ & \left| p_k \right\rangle \left\langle p_j \right| \otimes \left(\otimes^{r-1} \mathbb{I} \right) \otimes a \otimes \left(\otimes^{n-r} \mathbb{I} \right) \end{split}$$

Hence the program P can be formally defined as the sum O_P of all these operators:

$$O_P = \sum_{j,r,k} O_{j,r,k}^{INC} + \sum_{j,r,k,l} O_{j,r,k,l}^{DEC}$$

Thus O_P is the global operator which describes a computation step of M. The Halt instruction is simply executed by doing nothing when the program counter assumes the value $|p_m\rangle$. For such a value, O_P would produce the null vector as a result; however, in what follows we will add a term to O_P that allows us to extract the solution of the problem from a prefixed register when the program counter assumes the value $|p_m\rangle$.

A configuration of M is given by the value of the program counter and the values contained in the registers. From a mathematical point of view, a configuration of M is a vector of the Hilbert space $\mathbb{C}^m \otimes (\otimes^n \mathcal{H})$. A transition between two configurations is obtained by executing one instruction of P (the one pointed at by the program counter), that is, by applying the operator O_P to the current configuration of M.

As shown in [20], QRMs can simulate any (classical, deterministic) register machine, and thus they are (at least) computationally complete.

4 Classical and Quantum UREM P Systems

We are now ready to focus our attention to P systems. As stated in the introduction, quantum UREM P systems have been introduced in [20] as a quantum version of UREM P systems. Hence, let us start by recalling the definition of the classical model of computation.

A UREM P system [11] of degree d + 1 is a construct Π of the form:

$$\Pi = (A, \mu, e_0, \dots, e_d, w_0, \dots, w_d, R_0, \dots, R_d)$$

where:

- A is an alphabet of *objects*;
- μ is a *membrane structure*, with the membranes labeled by numbers $0, \ldots, d$ in a one-to-one manner;
- e_0, \ldots, e_d are the initial energy values assigned to the membranes $0, \ldots, d$. In what follows we assume that e_0, \ldots, e_d are non-negative integers;
- w_0, \ldots, w_d are multisets over A associated with the regions $0, \ldots, d$ of μ ;
- R_0, \ldots, R_d are finite sets of *unit rules* associated with the membranes $0, \ldots, d$. Each rule has the form $(\alpha : a, \Delta e, b)$, where $\alpha \in \{in, out\}, a, b \in A$, and $|\Delta e|$ is the amount of energy that — for $\Delta e \geq 0$ — is added to or — for $\Delta e < 0$ — is subtracted from e_i (the energy assigned to membrane *i*) by the application of the rule.

By writing $(\alpha_i : a, \Delta e, b)$ instead of $(\alpha : a, \Delta e, b) \in R_i$, we can specify only one set of rules R with

$$R = \{ (\alpha_i : a, \Delta e, b) : (\alpha : a, \Delta e, b) \in R_i, 0 \le i \le d \}$$

The *initial configuration* of Π consists of e_0, \ldots, e_d and w_0, \ldots, w_d . The transition from a configuration to another one is performed by non-deterministically choosing one rule from some R_i and applying it (observe that here we consider a *sequential* model of applying the rules instead of choosing rules in a maximally parallel way, as it is often required in P systems). Applying $(in_i : a, \Delta e, b)$ means that an object a (being in the membrane immediately outside of i) is changed into b while entering membrane i, thereby changing the energy value e_i of membrane iby Δe . On the other hand, the application of a rule ($out_i : a, \Delta e, b$) changes object a into b while leaving membrane i, and changes the energy value e_i by Δe . The rules can be applied only if the amount e_i of energy assigned to membrane i fulfills the requirement $e_i + \Delta e \geq 0$. Moreover, we use some sort of local priorities: if there are two or more applicable rules in membrane i, then one of the rules with max $|\Delta e|$ has to be used.

A sequence of transitions is called a *computation*; it is *successful* if and only if it halts. The *result* of a successful computation is considered to be the distribution of energies among the membranes (a non-halting computation does not produce a result). If we consider the energy distribution of the membrane structure as

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the input to be analysed, we obtain a model for accepting sets of (vectors of) non-negative integers.

The following result, proved in [11], establishes computational completeness for this model of P systems.

Proposition 2. Every partial recursive function $f : \mathbb{N}^{\alpha} \to \mathbb{N}^{\beta}$ can be computed by a UREM P system with (at most) $\max\{\alpha, \beta\} + 3$ membranes.

It is interesting to note that the proof of this proposition is obtained by simulating register machines. In the simulation, a P system is defined which contains one subsystem for each register of the simulated machine. The contents of the register are expressed as the energy value e_i assigned to the *i*-th subsystem. A single object is present in the system at every computation step, which stores the label of the instruction of the program P currently simulated. Increment instructions are simulated in two steps by using the rules $(in_i : p_j, 1, \tilde{p}_j)$ and $(out_i : \tilde{p}_j, 0, p_k)$. Decrement instructions are also simulated in two steps, by using the rules $(in_i : p_j, 0, \tilde{p}_j)$ and $(out_i : \tilde{p}_j, -1, p_k)$ or $(out_i : \tilde{p}_j, 0, p_l)$. The use of priorities associated to these last rules is crucial to correctly simulate a decrement instruction. For the details of the proof we refer the reader to [11].

On the other hand, by omitting the priority feature we do not get systems with universal computational power. Precisely, in [11] it is proved that P systems with unit rules and energy assigned to membranes without priorities and with an arbitrary number of membranes characterize the family $PsMAT^{\lambda}$ of Parikh sets generated by context-free matrix grammars (without occurrence checking and with λ -rules).

In quantum UREM P systems, all the elements of the model (multisets, the membrane hierarchy, configurations, and computations) are defined just like the corresponding elements of the classical P systems, but for objects and rules. The objects of A are represented as pure states of a quantum system. If the alphabet contains $d \ge 2$ elements then, recalling the notation introduced in section 2, without loss of generality we can put $A = \left\{ |0\rangle, \left| \frac{1}{d-1} \right\rangle, \left| \frac{2}{d-1} \right\rangle, \dots, \left| \frac{d-2}{d-1} \right\rangle, |1\rangle \right\}$, that is, $A = \{|a\rangle : a \in L_d\}$. As stated above, the quantum system will also be able to assume as a state any superposition of the kind:

$$c_0 |0\rangle + c_{\frac{1}{d-1}} \left| \frac{1}{d-1} \right\rangle + \ldots + c_{\frac{d-2}{d-1}} \left| \frac{d-2}{d-1} \right\rangle + c_1 |1\rangle$$

with $c_0, c_{\frac{1}{d-1}}, \ldots, c_{\frac{d-2}{d-1}}, c_1 \in \mathbb{C}$ such that $\sum_{i=0}^{d-1} |c_{\frac{i}{d-1}}|^2 = 1$. A multiset is simply a collection of quantum systems, each in its own state.

In order to represent the energy values assigned to membranes we must use quantum systems which can exist in an infinite (countable) number of states. Hence we assume that every membrane of the quantum P system has an associated infinite dimensional quantum harmonic oscillator whose state represents the energy value assigned to the membrane. To modify the state of such harmonic oscillator we can use the infinite dimensional version of creation (a^{\dagger}) and annihilation (a) operators¹ described in section 2, which are commonly used in quantum mechanics. The actions of a^{\dagger} and a on the state of an infinite dimensional harmonic oscillator are analogous to the actions on the states of truncated harmonic oscillators; the only difference is that in the former case there is no state with maximum energy, and hence the creation operator never produces the null vector. Also in this case it is possible to express operators $E_{x,y} = |y\rangle \langle x|$ as appropriate compositions of a^{\dagger} and a.

As in the classical case, rules are associated to the membranes rather than to the regions enclosed by them. Each rule of R_i is an operator of the form

$$|y\rangle \langle x| \otimes O, \quad \text{with } x, y \in L_d$$

$$\tag{3}$$

where O is a linear operator which can be expressed by an appropriate composition of operators a^{\dagger} and a. The part $|y\rangle\langle x|$ is the guard of the rule: it makes the rule "active" (that is, the rule produces an effect) if and only if a quantum system in the basis state $|x\rangle$ is present. The semantics of rule (3) is the following: If an object in state $|x\rangle$ is present in the region immediately outside membrane i, then the state of the object is changed to $|y\rangle$ and the operator O is applied to the state of the harmonic oscillator associated with the membrane. Notice that the application of O can result in the null vector, so that the rule has no effect even if its guard is satisfied; this fact is equivalent to the condition $e_i + \Delta e \geq 0$ on the energy of membrane *i* required in the classical case. Differently from the classical case, no local priorities are assigned to the rules. If two or more rules are associated to membrane i, then they are summed. This means that, indeed, we can think to each membrane as having only one rule with many guards. When an object is present, the inactive parts of the rule (those for which the guard is not satisfied) produce the null vector as a result. If the region in which the object occurs contains two or more membranes, then all their rules are applied to the object. Observe that the object which activates the rules never crosses the membranes. This means that the objects specified in the initial configuration can change their state but never move to a different region. Notwithstanding, transmission of information between different membranes is possible, since different objects may modify in different ways the energy state of the harmonic oscillators associated with the membranes.

The application of one or more rules determines a *transition* between two configurations. A *halting configuration* is a configuration in which no rule can be applied. A sequence of transitions is a *computation*. A computation is *successful* if and only if it *halts*, that is, reaches a halting configuration. The *result* of a successful computation is considered to be the distribution of energies among the membranes in the halting configuration. A non-halting computation does not produce a result. Just like in the classical case, if we consider the energy distribution of the membrane structure as the input to be analyzed, we obtain a model for accepting sets of (vectors of) non-negative integers.

¹ We recall that an alternative formulation that uses spin-rising (J_+) and spin-lowering (J_-) operators instead of creation and annihilation is also possible.

5 Computational Completeness of Quantum UREM P Systems

In this section we prove that quantum P systems with unit rules and energy assigned to membranes are computationally complete, that is, they are able to compute any partial recursive function $f: \mathbf{N}^{\alpha} \to \mathbf{N}^{\beta}$. As in the classical case, the proof is obtained by simulating register machines.

Theorem 1. Every partial recursive function $f : \mathbf{N}^{\alpha} \to \mathbf{N}^{\beta}$ can be computed by a quantum P system with unit rules and energy assigned to membranes with (at most) max{ α, β } + 3 membranes.

Proof. Let M = (n, P, 1, m) be a deterministic *n*-register machine that computes f. Let m be the number of instructions of P. The initial instruction of P has the label 1, and the halting instruction has the label m. Observe that, according to Proposition 1, $n = \max\{\alpha, \beta\} + 2$ is enough.

The input values x_1, \ldots, x_α are expected to be in the first α registers, and the output values are expected to be in registers 1 to β at the end of a successful computation. Moreover, without loss of generality, we may assume that at the beginning of a computation all the registers except (eventually) the registers 1 to α contain zero.

We construct the quantum P system

$$\Pi = (A, \mu, e_0, \dots, e_n, w_0, \dots, w_n, R_0, \dots, R_n)$$

where:

- $A = \{|j\rangle \mid j \in L_m\}$ $\mu = [0[1]_1 \cdots [\alpha]_\alpha \cdots [n]_n]_0$ $e_i = \begin{cases} |\varepsilon_{x_i}\rangle & \text{for } 1 \le i \le \alpha \\ |\varepsilon_0\rangle & \text{for } \alpha + 1 \le i \le n \\ 0 & \text{(the null vector) for } i = 0 \end{cases}$
- $w_0 = |0\rangle$
- $w_0 = |0\rangle$ $w_i = \emptyset$ for $1 \le i \le n$
- $R_0 = \emptyset$ $R_i = \sum_{j=1}^m O_{ij}$ for $1 \le i \le n$

where the O_{i_i} 's are local operators which simulate instructions of the kind j:(INC(i),k) and j:(DEC(i),k,l) (one local operator for each increment or decrement operation which affects register i). The details on how the O_{i_i} 's are defined are given below.

The value contained into register $i, 1 \leq i \leq n$, is represented by the energy value $e_i = |\varepsilon_{x_i}\rangle$ of the infinite dimensional quantum harmonic oscillator associated with membrane i. Figure 2 depicts a typical configuration of Π . The skin contains one object of the kind $|j\rangle$, $j \in L_m$, which mimics the program counter of machine M. Precisely, if the program counter of M has the value $k \in \{1, 2, \ldots, m\}$ then



Fig. 2. A configuration of the simulating P system

the object present in region 0 is $\left|\frac{k-1}{m-1}\right\rangle$. In order to avoid cumbersome notation, in what follows we denote by $|p_k\rangle$ the state $\left|\frac{k-1}{m-1}\right\rangle$ of the quantum system which mimics the program counter.

The sets of rules R_i depend upon the instructions of P. Precisely, the simulation works as follows.

Increment instructions j: (INC(i), k) are simulated by a guarded rule of the kind |p_k⟩ ⟨p_j| ⊗ a[†] ∈ R_i.
 If the object |p_j⟩ is present in region 0, then the rule transforms it into object |p_k⟩ and increments the energy level of the harmonic oscillator contained into

 $|p_k\rangle$ and increments the energy level of the harmonic oscillator contained into membrane *i*.

2. Decrement instructions j : (DEC(i), k, l) are simulated by a guarded rule of the kind:

$$|p_l\rangle \langle p_j| \otimes |\varepsilon_0\rangle \langle \varepsilon_0| + |p_k\rangle \langle p_j| \otimes a \in R_i$$

In fact, let us assume that the object $|p_j\rangle$ is present in region 0 (if $|p_j\rangle$ is not present then the above rule produces the null operator), and let us denote by O the above rule. The harmonic oscillator may be in the base state $|\varepsilon_0\rangle$ or in a base state $|\varepsilon_x\rangle$ with x a positive integer.

If the state of the harmonic oscillator is $|\varepsilon_0\rangle$ then the rule produces:

$$O(|p_{j}\rangle \otimes |\varepsilon_{0}\rangle) = \\ = (|p_{l}\rangle \langle p_{j}| \otimes |\varepsilon_{0}\rangle \langle \varepsilon_{0}|)(|p_{j}\rangle \otimes |\varepsilon_{0}\rangle) + (|p_{k}\rangle \langle p_{j}| \otimes a)(|p_{j}\rangle \otimes |\varepsilon_{0}\rangle) = \\ = |p_{l}\rangle \otimes |\varepsilon_{0}\rangle + |p_{k}\rangle \otimes \mathbf{0} = |p_{l}\rangle \otimes |\varepsilon_{0}\rangle$$

that is, the state of the oscillator is unaltered and the program counter is set to $|p_l\rangle$.

If the state of the harmonic oscillator is $|\varepsilon_x\rangle$, for a positive integer x, then the rule produces:

$$O(|p_j\rangle \otimes |\varepsilon_x\rangle) = = (|p_l\rangle \langle p_j| \otimes |\varepsilon_0\rangle \langle \varepsilon_0|)(|p_j\rangle \otimes |\varepsilon_x\rangle) + (|p_k\rangle \langle p_j| \otimes a)(|p_j\rangle \otimes |\varepsilon_x\rangle) = = |p_l\rangle \otimes \mathbf{0} + |p_k\rangle \otimes a |\varepsilon_x\rangle = |p_k\rangle \otimes |\varepsilon_{x-1}\rangle$$

that is, the energy level of the harmonic oscillator is decremented and the program counter is set to $|p_k\rangle$.

The set R_i of rules is obtained by summing all the operators which affect (increment or decrement) register *i*. The Halt instruction is simply simulated by doing nothing with the object $|p_m\rangle$ when it appears in region 0.

It is apparent from the description given above that after the simulation of each instruction each energy value e_i equals the value contained into register i, with $1 \leq i \leq m$. Hence, when the halting symbol $|p_m\rangle$ appears in region 0, the energy values e_1, \ldots, e_β equal the output of the program P.

Let us conclude this section by observing that, in order to obtain computational completeness, it is not necessary that the objects cross the membranes. This fact avoids one of the problems raised in [21]: the existence of a "magic" quantum transportation mechanism which is able to move objects according to the target contained into the rule. In quantum P systems with unit rules and energy assigned to membranes, the only problem is to keep the object $|p_j\rangle$ localized in region 0, so that it never enters into the other regions. In other words, the major problem of this kind of quantum P systems is to oppose the tunnel effect.

It should also be evident that the proof of Theorem 1 can be modified to show that quantum P systems are able to simulate quantum register machines. Indeed, the notable difference between the quantum P systems described above and quantum register machines is that in the latter model we modify the values contained into registers using *global* operators (if a given register must not be modified then the identity operator is applied to its state) whereas in the former model we operate locally, on a smaller Hilbert space. Hence, as it happens in classical P systems, membranes are used to divide the site where the computation occurs into independent local areas. The effect of each rule is local, in the sense that the rule affects only the state of one subsystem. Due to the simulations mentioned above, we can order these computational models with respect to their computational power, as follows:

deterministic	/	quantum	/	quantum P with unit r	systems ules and
register machines	\geq	machines	\geq	energy assi	gned to
maoninos				membranes	

Quantum register machines can thus be used as a tool to study the computational power of other quantum models of computation, just like it happens in the classical case.

6 Solving 3-SAT with QRMs and with Quantum UREM P Systems

Quantum UREM P systems are not only able to compute all partial recursive functions, like Turing machines, but they can also be very efficient computation devices. Indeed, in this section we show how we can solve in polynomial time the **NP**–complete decision problem 3-SAT by quantum register machines and by quantum UREM P systems. As we will see, the solution provided by quantum UREM P systems will be even more efficient that the one obtained with QRMs.

It is important to stress that our solutions assume that a specific non–unitary operator, built using the truncated version of creation and annihilation operators, can be realized as an instruction of quantum register machines and as a rule of quantum UREM P systems, respectively. The construction relies also upon the assumption that an external observer is able to discriminate, as the result of a measurement, a null vector from a non–null vector.

6.1 The 3-SAT problem

A boolean variable is a variable which can assume one of two possible truth values: TRUE and FALSE. As usually done in the literature, we will denote TRUE by 1 and FALSE by 0. A *literal* is either a directed or a negated boolean variable. A *clause* is a disjunction of literals, whereas a 3-clause is a disjunction of exactly three literals. Given a set $X = \{x_1, x_2, \ldots, x_n\}$ of boolean variables, an *assignment* is a mapping $a : X \to \{0, 1\}$ that associates to each variable a truth value. The number of all possible assignments to the variables of X is 2^n . We say that an assignment *satisfies* the clause C if, assigned the truth values to all the variables which occur in C, the evaluation of C (considered as a boolean formula) gives 1 as a result.

The 3-SAT decision problem is defined as follows.

Problem 1. NAME: 3-SAT.

- INSTANCE: a set $C = \{C_1, C_2, \dots, C_m\}$ of 3-clauses, built on a finite set $\{x_1, x_2, \dots, x_n\}$ of boolean variables.
- QUESTION: is there an assignment of the variables x_1, x_2, \ldots, x_n that satisfies all the clauses in C?

Notice that the number m of possible 3-clauses is polynomially bounded with respect to n: in fact, since each clause contains exactly three literals, we can have at most $(2n)^3 = 8n^3$ clauses.

In what follows we will equivalently say that an instance of 3-SAT is a boolean formula ϕ_n , built on *n* free variables and expressed in conjunctive normal form, with each clause containing exactly three literals. The formula ϕ_n is thus the conjunction of the above clauses.

It is well known [16] that 3-SAT is an **NP**–complete problem.

6.2 Solving 3-SAT with quantum register machines

Let ϕ_n be an instance of 3-SAT containing *n* free variables. We will first show how to evaluate ϕ_n with a classical register machine; then, we will initialize the input registers with a superposition of all possible assignments, we will compute the corresponding superposition of output values into an output register, and finally we will apply the linear operator $2^n |1\rangle \langle 1|$ to the output register to check whether ϕ_n is a positive instance of 3-SAT.

The register machine that we use to evaluate ϕ_n is composed by n+1 registers. The first *n* registers correspond (in a one-to-one manner) to the free variables of ϕ_n , while the last register is used to compute the output value. The *structure* of the program used to evaluate ϕ_n is the following:

```
\phi = 0

if C_1 = 0 then goto end

if C_2 = 0 then goto end

:

if C_m = 0 then goto end

\phi = 1

end:
```

where ϕ denotes the output register, and C_1, C_2, \ldots, C_m are the clauses of ϕ_n . Let $X_{i,j}$, with $j \in \{1, 2, 3\}$, be the literals (director or negated variables) which occur in the clause C_i (hence $C_i = X_{i,1} \lor X_{i,2} \lor X_{i,3}$). We can thus write the above structure of the program, at a finer grain, as follows:

$$\phi = 0$$
if $X_{1,1} = 1$ then goto end₁
if $X_{1,2} = 1$ then goto end₁
if $X_{1,3} = 1$ then goto end₁
goto end
end₁: if $X_{2,1} = 1$ then goto end₂
if $X_{2,2} = 1$ then goto end₂
if $X_{2,2} = 1$ then goto end₂
goto end
end₂:
$$\vdots$$
end_{m-1}: if $X_{m,1} = 1$ then goto end
if $X_{m,2} = 1$ then goto end
if $X_{m,3} = 1$ then goto end
end:
end:

In the above structure it is assumed that each literal $X_{i,j}$, with $1 \leq i \leq m$ and $j \in \{1, 2, 3\}$, is substituted with the corresponding variable which occurs in it; moreover, if the variable occurs negated into the literal then the comparison must be done with 0 instead of 1:

if
$$X_{i,j} = 0$$
 then goto end_i

Since the free variables of ϕ_n are bijectively associated with the first *n* registers of the machine, in order to evaluate ϕ_n we need a method to check whether a given register contains 0 (or 1) without destroying its value. Let us assume that, when the program counter of the machine reaches the value k, we have to execute the following instruction:

k: if
$$X_{i,j} = 1$$
 then goto end_i

We translate such instruction as follows (where, instead of $X_{i,j}$, we specify the register which corresponds to the variable indicated in $X_{i,j}$):

$$k: DEC(X_{i,j}), k+1, k+2$$

 $k+1: INC(X_{i,j}), \text{end}_i$

The instruction:

k: if
$$X_{i,j} = 0$$
 then goto end_i

is instead translated as follows:

$$k: DEC(X_{i,j}), k+1, \text{ end}_i$$
$$k+1: INC(X_{i,j}), k+2$$

Notice that the only difference with the previous sequence of instructions is in the position of "end_i" and "k + 2". Moreover, the structure of the program is always the same. As a consequence, given an instance ϕ_n of 3-SAT, the program P of a register machine which evaluates ϕ_n can be obtained in a very straightforward (mechanical) way.

On a *classical* register machine, this program computes the value of ϕ_n for a given assignment to its variables x_1, x_2, \ldots, x_n . On a *quantum* register machine we can initialize the registers with the following state:

$$\otimes^{n-1}H_1|0\rangle\otimes|0\rangle$$

which sets the output register ϕ to 0 and the registers corresponding to x_1, x_2, \ldots, x_n to a superposition of all possible assignments. Then, we apply the global operator O_P which corresponds to the program P until the program counter reaches the value $|p_{\text{end}}\rangle$, thus computing in the output register a superposition of all classical results. The operator O_P is built as described in section 3, with the only difference that now it contains also the term:

$$|p_{\text{end}}\rangle \langle p_{\text{end}}| \otimes \text{ID}_n \otimes 2^n |1\rangle \langle 1| = |p_{\text{end}}\rangle \langle p_{\text{end}}| \otimes \text{ID}_n \otimes \left[\underbrace{(|1\rangle \langle 1| + |1\rangle \langle 1|) \circ \dots \circ (|1\rangle \langle 1| + |1\rangle \langle 1|)}_{n \text{ times}}\right]$$

which extracts the result from the output register when the program counter assumes the value $|p_{end}\rangle$. The number of times we have to apply O_P is equal to the length of P, that is, $2 \cdot 3m + 2 = 6m + 2$: two instructions for each literal in every clause, plus two final instructions.

Now, if ϕ_n is not satisfiable then the contents of the output register is $|0\rangle$, and when the program counter reaches the value $|p_{\text{end}}\rangle$ the operator O_P transforms it to the null vector. On the other hand, if ϕ_n is satisfiable then the contents of the output register will be a superposition $\alpha_0 |0\rangle + \alpha_1 |1\rangle$, with $\alpha_1 \neq 0$. By applying the operator O_P we obtain (here $|\psi_n\rangle$ denotes the state of the *n* input registers):

$$O_{P}(|p_{end}\rangle \otimes |\psi_{n}\rangle \otimes (\alpha_{0} |0\rangle + \alpha_{1} |1\rangle)) =$$

$$= (|p_{end}\rangle \langle p_{end}| \otimes ID_{n} \otimes 2^{n} |1\rangle \langle 1|) \cdot \cdot (|p_{end}\rangle \otimes |\psi_{n}\rangle \otimes (\alpha_{0} |0\rangle + \alpha_{1} |1\rangle)) =$$

$$= |p_{end}\rangle \langle p_{end} |p_{end}\rangle \otimes ID_{n} |\psi_{n}\rangle \otimes 2^{n} |1\rangle \langle 1| (\alpha_{0} |0\rangle + \alpha_{1} |1\rangle) =$$

$$= |p_{end}\rangle \otimes |\psi_{n}\rangle \otimes (2^{n}\alpha_{0} |1\rangle \langle 1|0\rangle + 2^{n}\alpha_{1} |1\rangle \langle 1|1\rangle) =$$

$$= |p_{end}\rangle \otimes |\psi_{n}\rangle \otimes (\mathbf{0} + 2^{n}\alpha_{1} |1\rangle) =$$

$$= |p_{end}\rangle \otimes |\psi_{n}\rangle \otimes 2^{n}\alpha_{1} |1\rangle$$

that is, a non–null vector.

We can thus conclude that if an external observer is able to discriminate between a null vector and a non-null vector, and it is possible to build and apply the operator $2^n |1\rangle \langle 1| = E_{1,1} = N = a^{\dagger}a$ to the output register of a QRM, then we have a family of QRMs that solve 3-SAT in polynomial time. This solution is given in a *semi-uniform* setting: in particular, the program P executed by the QRM depends upon the instance ϕ_n of 3-SAT we want to solve.

6.3 Solving 3-SAT with Quantum UREM P Systems

In this section we finally show how to build a (semi–uniform) family of quantum UREM P systems that solves 3-SAT. Let ϕ_n be an instance of 3-SAT containing n free variables. The structure and the initial configuration of the P system that determines whether ϕ_n is satisfiable is similar to what shown in Figure 2, the only difference being that there are n + 1 subsystems instead of n.

As we have done with quantum register machines, let us start by showing how to evaluate ϕ_n for a given assignment of truth values to its variables x_1, \ldots, x_n . The input values are set as the energies $|\varepsilon_{x_i}\rangle$ of the harmonic oscillators associated with the membranes from 1 to n. The energy (eventually) associated with the skin membrane is not used. The (n + 1)-th membrane, whose harmonic oscillator will contain the output at the end of the computation, is initialized with $|\varepsilon_0\rangle$. The alphabet A consists of all the possible values which can be assumed by the program counter of the QRM that evaluates ϕ_n . In the initial configuration the P system contains only one copy of the object $|p_1\rangle$, corresponding to the initial value of the program counter, in the region enclosed by the skin membrane.

The evaluation of ϕ_n could be performed by simulating the QRM obtained from ϕ_n as explained in the previous section. However, we can obtain a slightly more efficient P system as follows. We start from the program structure (4), which can be obtained from ϕ_n in a straightforward way. Now, let us suppose we must execute the following instruction:

k: if
$$X_{i,j} = 1$$
 then goto end_i

As told above, this instruction is performed as follows in a register machine:

$$k: DEC(X_{i,j}), k+1, k+2$$

 $k+1: INC(X_{i,j}), end_i$

If we had to simulate these two instructions using a quantum UREM P system, we should use the following sum of rules:

$$\underbrace{\left(\left|p_{\mathrm{end}_{i}}\right\rangle\left\langle p_{k+1}\right|\otimes a^{\dagger}\right)}_{k+1:\ INC(X_{i,j}),\ \mathrm{end}_{i}}+\underbrace{\left(\left|p_{k+2}\right\rangle\left\langle p_{k}\right|\otimes\left|\varepsilon_{0}\right\rangle\left\langle\varepsilon_{0}\right|+\left|p_{k+1}\right\rangle\left\langle p_{k}\right|\otimes a\right)}_{k:\ DEC(X_{i,j}),\ k+1,\ k+2}\in R_{\ell}$$

where $\ell = \langle i, j \rangle$ is the index of the variable (in the set $\{x_1, x_2, \ldots, x_n\}$) which occurs in literal $X_{i,j}$. As we can see, this operator produces the vector $|p_{k+2}\rangle \otimes |\varepsilon_0\rangle$ if the harmonic oscillator of membrane ℓ is in state $|\varepsilon_0\rangle$; otherwise, it produces the vector $|p_{\text{end}_i}\rangle \otimes |\varepsilon_1\rangle$. Hence we can simplify the above expression as follows:

$$\begin{split} |p_{\mathrm{end}_i}\rangle \langle p_k| \otimes |\varepsilon_1\rangle \langle \varepsilon_1| + |p_{k+2}\rangle \langle p_k| \otimes |\varepsilon_0\rangle \langle \varepsilon_0| = \\ &= |p_{\mathrm{end}_i}\rangle \langle p_k| \otimes a^{\dagger}a + |p_{k+2}\rangle \langle p_k| \otimes aa^{\dagger} \end{split}$$

We denote this operator by $O_{i,j,k}^{(1)}$. Analogously, if the instruction to be executed is:

k: if $X_{i,j} = 0$ then goto end_i

then we use the operator

$$O_{i,j,k}^{(0)} = \left| p_{\mathrm{end}_i} \right\rangle \left\langle p_k \right| \otimes a a^{\dagger} + \left| p_{k+2} \right\rangle \left\langle p_k \right| \otimes a^{\dagger} a \in R_{\ell}$$

which produces the vector $|p_{k+2}\rangle \otimes |\varepsilon_1\rangle$ if the harmonic oscillator of membrane ℓ is in state $|\varepsilon_1\rangle$, otherwise it produces the vector $|p_{\text{end}_i}\rangle \otimes |\varepsilon_0\rangle$.

Since the value $|p_{k+1}\rangle$ is no longer used, we can "compact" the program by redefining the operators $O_{i,j,k}^{(0)}$ and $O_{i,j,k}^{(1)}$ respectively as:

$$\begin{aligned} O_{i,j,k}^{(0)} &= |p_{\text{end}_i}\rangle \left\langle p_k \right| \otimes aa^{\dagger} + |p_{k+1}\rangle \left\langle p_k \right| \otimes a^{\dagger}a \\ O_{i,j,k}^{(1)} &= |p_{\text{end}_i}\rangle \left\langle p_k \right| \otimes a^{\dagger}a + |p_{k+1}\rangle \left\langle p_k \right| \otimes aa^{\dagger} \end{aligned}$$

The "goto end" instructions in (4) can be executed as if they were if statements whose condition is the negation of the condition given in the previous if. Hence the two instructions:

7: if $X_{2,3} = 1$ then goto end₂ 8: goto end

can be thought of as:

7: if X_{2,3} = 1 then goto end₂
8: if X_{2,3} = 0 then goto end

which are realized by the operators $O_{2,3,7}^{(1)}$ and $O_{2,3,8}^{(0)}$ (to be added to membrane $\langle 2,3\rangle$). The last instruction ($\phi = 1$) of the program can be implemented as follows:

$$|p_{\mathrm{end}}\rangle \langle p_{\mathrm{end}-1}|\otimes a^{\dagger}$$

to be added to membrane n + 1.

For each membrane $i \in \{1, 2, ..., n\}$, the set of rules R_i is obtained by summing all the operators which concern variable x_i .

Note that the formulation given in terms of quantum P systems is simpler than the one obtained with QRMs. As usual, if we consider a single assignment to the variables of ϕ_n then at the end of the computation we will obtain the result of the evaluation of ϕ_n as the energy of the output membrane. Instead, if we initialize the harmonic oscillators of the *n* input membranes with a uniform superposition of all possible classical assignments to x_1, x_2, \ldots, x_n , then at the end of the computation the harmonic oscillator of membrane n + 1 will be in one of the following states:

- $|0\rangle$, if ϕ_n is not satisfiable;
- a superposition $\alpha_0 |0\rangle + \alpha_1 |1\rangle$, with $\alpha_1 \neq 0$, if ϕ_n is satisfiable.

Once again, we add the rule:

$$|p_{\text{end}}\rangle \langle p_{\text{end}}|\otimes 2^n |1\rangle \langle 1| \in R_{n+1}$$

to membrane n + 1 to extract the result.

We have thus obtained a family of quantum UREM P systems which solves 3-SAT in polynomial time. Also this scheme works in the *semi-uniform* setting: in fact, it is immediately verified that the rules of the system depend upon the instance ϕ_n of 3-SAT to be solved.

7 Conclusions and Directions for Future Research

In this paper we have overviewed the state of the art concerning quantum UREM P systems. Starting from basic notions of quantum computers and quantum mechanics, we have seen how quantum register machines and quantum UREM P systems can be defined.

Subsequently, we have proved that such quantum models of computation are computationally complete, that is, they are able to compute any partial recursive function $f : \mathbf{N}^{\alpha} \to \mathbf{N}^{\beta}$. This result has been obtained by simulating classical

deterministic register machines. Moreover, we have shown a family of QRMs and a family of quantum UREM P systems that solve (in the semi–uniform setting) the 3-SAT **NP**–complete decision problem in polynomial time. Their construction relies upon the following assumptions: (1) an external observer is able to discriminate, as the result of a measurement, a null vector from a non–null vector, and (2) a specific non–unitary operator, which can be expressed using creation and annihilation operators, can be realized as an instruction of the quantum register machine, and as a rule of the quantum P system, respectively.

One possible direction for future research is to study the computational properties of quantum P systems which contain and process entangled objects. Another line of research is to study the limits of the computational power of quantum P systems by attacking harder than NP-complete problems. In particular, we conjecture that EXP-complete problems can be solved in polynomial time with quantum P systems.

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The Calculus of Looping Sequences for Modeling Biological Membranes

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Summary. We survey the formalism Calculus of Looping Sequences (CLS) and a number of its variants from the point of view of their use for describing biological membranes. The formalism CLS is based on term rewriting and allows describing biomolecular systems. A first variant of CLS, called Stochastic CLS, extends the formalism with stochastic time, another variant, called LCLS (CLS with links), allows describing proteins interaction at the domain level. A third variant is introduced for easier description of biological membranes. This extension can be encoded into CLS as well as other formalisms capable of membrane description such as Brane Calculi and P Systems. Such encodings allow verifying and simulating descriptions in Brane Calculi and P Systems by means of verifiers and simulators developed for CLS.

1 Introduction

Cell biology, the study of the morphological and functional organization of cells, is now an established field in biochemical research. Computer Science can help the research in cell biology in several ways. For instance, it can provide biologists with models and formalisms capable of describing and analyzing complex systems such as cells. In the last few years many formalisms originally developed by computer scientists to model systems of interacting components have been applied to Biology. Among these, there are Petri Nets [16], Hybrid Systems [1], and the π -calculus [9, 25]. Moreover, new formalisms have been defined for describing biomolecular and membrane interactions [2, 7, 8, 11, 21, 23]. Others, such as P Systems [17, 18], have been proposed as biologically inspired computational models and have been later applied to the description of biological systems.

The π -calculus and new calculi based on it [21, 23] have been particularly successful in the description of biological systems, as they allow describing systems in a compositional manner. Interactions of biological components are modeled as communications on channels whose names can be passed; sharing names of private channels allows describing biological compartments. However, these calculi offer very low-level interaction primitives, and this causes models to become very large and difficult to read. Calculi such as those proposed in [7, 8, 11] give a more abstract description of systems and offer special biologically motivated operators. However, they are often specialized to the description of some particular kinds of phenomena such as membrane interactions or protein interactions. Finally, P Systems have a simple notation and are not specialized to the description of a particular class of systems, but they are still not completely general. For instance, it is possible to describe biological membranes and the movement of molecules across membranes, and there are some variants able to describe also more complex membrane activities. However, the formalism is not so flexible to allow describing easily new activities observed on membranes without extending the formalism to model such activities.

Therefore, we conclude that there is a need for a formalism having a simple notation, having the ability of describing biological systems at different levels of abstraction, having some notions of compositionality and being flexible enough to allow describing new kinds of phenomena as they are discovered, without being specialized to the description of a particular class of systems. For this reason in [3] we have introduced the Calculus of Looping Sequences (CLS).

CLS is a formalism based on term rewriting with some features, such as a commutative parallel composition operator, and some semantic means, such as bisimulations, which are common in process calculi. This permits to combine the simplicity of notation of rewriting systems with the advantage of a form of compositionality. Actually, in [4] we have defined bisimilarity relations on CLS terms which are congruences with respect to the operators. The bisimilarity relation may be used to verify a property of a system by assessing its bisimilarity with a system one knows to enjoy that property. The fact that bisimilarity is a congruence is very important for a compositional account of behavioral equivalence.

In [5, 6], we have defined two extensions of CLS. The first, Stochastic CLS, allows describing quantitative aspects of the modeled systems such as the time spent by occurrences of chemical reactions. The second, CLS with links, allows describing protein interaction more precisely at a lower level of abstraction, namely at the domain level.

In this paper, after recalling CLS and the two mentioned extensions, we focus on the modeling of biological membranes by means of CLS. Now, CLS does not offer an easy representation for membranes whose nature is fluid and for proteins which consequently move freely on membrane surfaces. For this reason, in [15] we have defined a CLS variant, called CLS+, which introduces a new operator allowing commutativity on membrane surfaces. We show how CLS+ can be encoded into CLS.

In [3, 15] we have shown how Brane Calculi [7] and P Systems [18] can be translated into CLS. Here we recall the ideas on which the translations are based.

CLS appears to allow description and manipulation of biological membranes and, moreover, offers, via translations, verification and simulation tools to other formalisms for membrane description.



Fig. 1. (i) represents $(a \cdot b \cdot c)^{L}$; (ii) represents $(a \cdot b \cdot c)^{L} \perp (d \cdot e)^{L}$; (iii) represents $(a \cdot b \cdot c)^{L} \perp ((d \cdot e)^{L} \mid f \cdot g)$.

2 The Calculus of Looping Sequences (CLS)

In this section we recall the Calculus of Looping Sequences (CLS) and we give some guidelines for the modeling of biological systems. CLS is essentially based on term rewriting, hence a CLS model consists of a term and a set of rewrite rules. The term is intended to represent the structure of the modeled system, and the rewrite rules to represent the events that may cause the system to evolve.

2.1 Formal Definition

We start with defining the syntax of terms. We assume a possibly infinite alphabet \mathcal{E} of symbols ranged over by a, b, c, \ldots

Definition 1 (Terms). Terms T and sequences S of CLS are given by the following grammar:

Т	::= S	$(S)^L \ \ T$	$T \mid T$
S	$::= \epsilon$	$a \qquad S \cdot S$	

where a is a generic element of \mathcal{E} , and ϵ represents the empty sequence. We denote with \mathcal{T} the infinite set of terms, and with \mathcal{S} the infinite set of sequences.

In CLS we have a sequencing operator $_...$, a looping operator $(_.)^{L}$, a parallel composition operator $_|_$ and a containment operator $_|_$. Sequencing can be used to concatenate elements of the alphabet \mathcal{E} . The empty sequence ϵ denotes the concatenation of zero symbols. A term can be either a sequence or a looping sequence (that is the application of the looping operator to a sequence) containing another term, or the parallel composition of two terms. By definition, looping and containment are always applied together, hence we can consider them as a single binary operator $(_)^{L} _$ which applies to one sequence and one term. Brackets can be used to indicate the order of application of the operators,

Brackets can be used to indicate the order of application of the operators, and we assume $(_)^L _$ to have precedence over $_ | _$. In Figure 1 we show some examples of CLS terms and their visual representation.

In CLS we may have syntactically different terms representing the same structure. We introduce a structural congruence relation to identify such terms. **Definition 2 (Structural Congruence).** The structural congruence relations \equiv_S and \equiv_T are the least congruence relations on sequences and on terms, respectively, satisfying the following rules:

$$S_{1} \cdot (S_{2} \cdot S_{3}) \equiv_{S} (S_{1} \cdot S_{2}) \cdot S_{3} \qquad S \cdot \epsilon \equiv_{S} \epsilon \cdot S \equiv_{S} S$$

$$S_{1} \equiv_{S} S_{2} \text{ implies } S_{1} \equiv_{T} S_{2} \text{ and } (S_{1})^{L} \downarrow T \equiv_{T} (S_{2})^{L} \downarrow T$$

$$T_{1} \mid T_{2} \equiv_{T} T_{2} \mid T_{1} \qquad T_{1} \mid (T_{2} \mid T_{3}) \equiv_{T} (T_{1} \mid T_{2}) \mid T_{3} \qquad T \mid \epsilon \equiv_{T} T$$

$$(\epsilon)^{L} \downarrow \epsilon \equiv_{T} \epsilon \qquad (S_{1} \cdot S_{2})^{L} \downarrow T \equiv_{T} (S_{2} \cdot S_{1})^{L} \downarrow T$$

Rules of the structural congruence state the associativity of \cdot and |, the commutativity of the latter and the neutral role of ϵ . Moreover, axiom $(S_1 \cdot S_2)^L \, \rfloor \, T \equiv_T (S_2 \cdot S_1)^L \, \rfloor \, T$ says that looping sequences can rotate. In the following, for simplicity, we will use \equiv in place of \equiv_T .

Rewrite rules will be defined essentially as pairs of terms, with the first term describing the portion of the system in which the event modeled by the rule may occur, and the second term describing how that portion of the system changes when the event occurs. In the terms of a rewrite rule we allow the use of variables. As a consequence, a rule will be applicable to all terms which can be obtained by properly instantiating its variables. Variables can be of three kinds: two of these are associated with the two different syntactic categories of terms and sequences, and one is associated with single alphabet elements. We assume a set of term variables TV ranged over by X, Y, Z, \ldots , a set of sequence variables SV ranged over by $\tilde{x}, \tilde{y}, \tilde{z}, \ldots$, and a set of element variables \mathcal{X} ranged over by x, y, z, \ldots . All these sets are possibly infinite and pairwise disjoint. We denote by \mathcal{V} the set of all variables, $\mathcal{V} = TV \cup SV \cup \mathcal{X}$, and with ρ a generic variable of \mathcal{V} . Hence, a pattern is a term that may include variables.

Definition 3 (Patterns). Patterns P and sequence patterns SP of CLS are given by the following grammar:

$$P ::= SP \mid (SP)^{L} \rfloor P \mid P \mid P \mid X$$

$$SP ::= \epsilon \mid a \mid SP \cdot SP \mid \tilde{x} \mid x$$

where a is a generic element of \mathcal{E} , and X, \tilde{x} and x are generic elements of TV, SVand \mathcal{X} , respectively. We denote with \mathcal{P} the infinite set of patterns.

We assume the structural congruence relation to be trivially extended to patterns. An *instantiation* is a partial function $\sigma : \mathcal{V} \to \mathcal{T}$. An instantiation must preserve the type of variables, thus for $X \in TV, \tilde{x} \in SV$ and $x \in \mathcal{X}$ we have $\sigma(X) \in \mathcal{T}, \sigma(\tilde{x}) \in \mathcal{S}$ and $\sigma(x) \in \mathcal{E}$, respectively. Given $P \in \mathcal{P}$, with $P\sigma$ we denote the term obtained by replacing each occurrence of each variable $\rho \in \mathcal{V}$ appearing in P with the corresponding term $\sigma(\rho)$. With Σ we denote the set of all the possible instantiations and, given $P \in \mathcal{P}$, with Var(P) we denote the set of variables appearing in P. Now we define rewrite rules.

Biomolecular Entity	CLS Term
Elementary object	Alphabet symbol
(genes, domains,	
other molecules, etc)	
DNA strand	Sequence of elements repr. genes
RNA strand	Sequence of elements repr. transcribed genes
Protein	Sequence of elements repr. domains
	or single alphabet symbol
Molecular population	Parallel composition of molecules
Membrane	Looping sequence

Table 1. Guidelines for the abstraction of biomolecular entities into CLS.

Definition 4 (Rewrite Rules). A rewrite rule is a pair of patterns (P_1, P_2) , denoted with $P_1 \mapsto P_2$, where $P_1, P_2 \in \mathcal{P}$, $P_1 \not\equiv \epsilon$ and such that $Var(P_2) \subseteq Var(P_1)$. We denote with \Re the infinite set of all the possible rewrite rules.

A rewrite rule $P_1 \mapsto P_2$ states that a term $P_1\sigma$, obtained by instantiating variables in P_1 by some instantiation function σ , can be transformed into the term $P_2\sigma$. We define the semantics of CLS as a transition system, in which states correspond to terms, and transitions correspond to rule applications.

Definition 5 (Semantics). Given a set of rewrite rules $\mathcal{R} \subseteq \mathfrak{R}$, the semantics of *CLS* is the least transition relation \rightarrow on terms closed under \equiv , and satisfying the following inference rules:

$$\frac{P_1 \mapsto P_2 \in \mathcal{R} \quad P_1 \sigma \neq \epsilon \quad \sigma \in \Sigma}{P_1 \sigma \to P_2 \sigma} \qquad \frac{T_1 \to T_2}{T \mid T_1 \to T \mid T_2} \qquad \frac{T_1 \to T_2}{\left(S\right)^L \mid T_1 \to \left(S\right)^L \mid T_2}$$

where the symmetric rule for the parallel composition is omitted.

A model in CLS is given by a term describing the initial state of the system and by a set of rewrite rules describing all the events that may occur.

2.2 Modeling Guidelines

We describe how CLS can be used to model biomolecular systems analogously to what done by Regev and Shapiro in [24] for the π -calculus. An abstraction is a mapping from a real-world domain to a mathematical domain, which may allow highlighting some essential properties of a system while ignoring other, complicating, ones. In [24], Regev and Shapiro show how to abstract biomolecular systems as concurrent computations by identifying the biomolecular entities and events of interest and by associating them with concepts of concurrent computations such as concurrent processes and communications. In particular, they give some guidelines for the abstraction of biomolecular systems to the π -calculus, and give some simple examples.
Biomolecular Event	Examples of CLS Rewrite Rule
State change	$a \mapsto b$
	$\widetilde{x} \cdot a \cdot \widetilde{y} \hspace{0.2cm} \mapsto \hspace{0.2cm} \widetilde{x} \cdot b \cdot \widetilde{y}$
Complexation	$a \mid b \mapsto c$
	$\widetilde{x} \cdot a \cdot \widetilde{y} \mid b \mapsto \widetilde{x} \cdot c \cdot \widetilde{y}$
Decomplexation	$\begin{array}{ccc} c & \mapsto & a \mid b \\ \widetilde{m} & c & \widetilde{n} & \downarrow \end{pmatrix} & \widetilde{m} & c & \widetilde{n} \mid b \end{array}$
Catalysis	$\begin{array}{cccccccccccccccccccccccccccccccccccc$
Catarysis	where $P_1 \mapsto P_2$ is the catalyzed event
State change	$(a \cdot \widetilde{x})^L \mid X \mapsto (b \cdot \widetilde{x})^L \mid X$
on membrane	
Complexation	$(a \cdot \widetilde{x} \cdot b \cdot \widetilde{y})^L \ ig X \mapsto (c \cdot \widetilde{x} \cdot \widetilde{y})^L \ ig X$
on membrane	$a \mid (b \cdot \widetilde{x})^L \mid X \mapsto (c \cdot \widetilde{x})^L \mid X$
	$(b \cdot \widetilde{x})^L \downarrow (a \mid X) \mapsto (c \cdot \widetilde{x})^L \downarrow X$
Decomplexation	$ig ig(c\cdot\widetilde{x}ig)^L\ ig \ X \ \mapsto \ ig(a\cdot b\cdot\widetilde{x}ig)^L\ ig \ X$
on membrane	$\left(c \cdot \widetilde{x}\right)^{L} \ ig X \mapsto a \mid \left(b \cdot \widetilde{x}\right)^{L} \ ig X$
	$\left(c \cdot \widetilde{x} ight)^L ig X \mapsto \left(b \cdot \widetilde{x} ight)^L ig (a \mid X)$
Catalysis	$egin{array}{cccccccccccccccccccccccccccccccccccc$
on membrane	where $SP_1 \mapsto SP_2$ is the catalyzed event
Membrane crossing	$a \mid \left(\widetilde{x}\right)^{L} \ \ X \mapsto \left(\widetilde{x}\right)^{L} \ \ (a \mid X)$
	$\left(\widetilde{x} ight)^{L} ot (a \mid X) \mapsto a \mid \left(\widetilde{x} ight)^{L} ot X$
	$\widetilde{x} \cdot a \cdot \widetilde{y} \mid (\widetilde{z})^L \perp X \mapsto (\widetilde{z})^L \perp (\widetilde{x} \cdot a \cdot \widetilde{y} \mid X)$
	$\left(\widetilde{z}\right)^{L} \ \ (\widetilde{x} \cdot a \cdot \widetilde{y} \mid X) \mapsto \widetilde{x} \cdot a \cdot \widetilde{y} \mid \left(\widetilde{z}\right)^{L} \ \ \ X$
Catalyzed	$a \mid \left(b \cdot \widetilde{x}\right)^L \rfloor X \; \; \mapsto \; \; \left(b \cdot \widetilde{x}\right)^L \downarrow (a \mid X)$
membrane crossing	$(b \cdot \widetilde{x})^L \downarrow (a \mid X) \mapsto a \mid (b \cdot \widetilde{x})^L \downarrow X$
	$\widetilde{x} \cdot a \cdot \widetilde{y} \mid (b \cdot \widetilde{z})^L \perp X \mapsto (b \cdot \widetilde{z})^L \perp (\widetilde{x} \cdot a \cdot \widetilde{y} \mid X)$
	$(b \cdot \widetilde{z})^{L} \downarrow (\widetilde{x} \cdot a \cdot \widetilde{y} \mid X) \mapsto \widetilde{x} \cdot a \cdot \widetilde{y} \mid (b \cdot \widetilde{z})^{L} \downarrow X$
Membrane joining	$\left(\widetilde{x}\right)_{L}^{L} \mid (a \mid X) \mapsto \left(a \cdot \widetilde{x}\right)^{L} \mid X$
	$(\widetilde{x})^{L} \sqcup (\widetilde{y} \cdot a \cdot \widetilde{z} \mid X) \mapsto (\widetilde{y} \cdot a \cdot \widetilde{z} \cdot \widetilde{x})^{L} \sqcup X$
Catalyzed	$\left(b \cdot \widetilde{x}\right)^{L} \mid (a \mid X) \mapsto \left(a \cdot b \cdot \widetilde{x}\right)^{L} \mid X$
membrane joining	$\left(\widetilde{x}\right)^{L} \downarrow (a \mid b \mid X) \mapsto \left(a \cdot \widetilde{x}\right)^{L} \downarrow (b \mid X)$
	$\left(b \cdot \widetilde{x}\right)^{L} \downarrow \left(\widetilde{y} \cdot a \cdot \widetilde{z} \mid X\right) \mapsto \left(\widetilde{y} \cdot a \cdot \widetilde{z} \cdot \widetilde{x}\right)^{L} \downarrow X$
	$\left \left(\widetilde{x} \right)^L \mid \left(\widetilde{y} \cdot a \cdot \widetilde{z} \mid b \mid X \right) \mapsto \left(\widetilde{y} \cdot a \cdot \widetilde{z} \cdot \widetilde{x} \right)^L \mid \left(b \mid X \right) \right $
Membrane fusion	$\left(\widetilde{x}\right)^{L} \downarrow (X) \mid \left(\widetilde{y}\right)^{L} \downarrow (Y) \mapsto \left(\widetilde{x} \cdot \widetilde{y}\right)^{L} \downarrow (X \mid Y)$
Catalyzed	$\left(a \cdot \widetilde{x}\right)^{L} \perp (X) \mid \left(b \cdot \widetilde{y}\right)^{L} \perp (Y) \mapsto $
membrane fusion	$ig(a \cdot \widetilde{x} \cdot b \cdot \widetilde{y} ig)^L ig (X \mid Y)$
Membrane division	$\left \begin{pmatrix} \widetilde{x} \cdot \widetilde{y} \end{pmatrix}^L \mid (X \mid Y) \mapsto \left(\widetilde{x} \right)^L \mid (X) \mid \left(\widetilde{y} \right)^L \mid (Y) \right $
Catalyzed	$\left[\left(a\cdot\widetilde{x}\cdot b\cdot\widetilde{y}\right)^{L}\right](X\mid Y) \mapsto$
membrane division	$(a \cdot \widetilde{x})^{L} \mid (X) \mid (b \cdot \widetilde{y})^{L} \mid (Y)$

Table 2. Guidelines for the abstraction of biomolecular events into CLS.

The use of rewrite systems, such as CLS, to describe biological systems is founded on a different abstraction. Usually, entities (and their structures) are abstracted by terms of the rewrite system, and events by rewriting rules. We have already introduced the biological interpretation of CLS operators in the previous section. Here we want to give more general guidelines.

First of all, we select the biomolecular entities of interest. Since we want to describe cells, we consider molecular populations and membranes. Molecular populations are groups of molecules that are in the same compartment of the cell. Molecules can be of many types: we classify them as DNA and RNA strands, proteins, and other molecules. DNA and RNA strands and proteins can be seen as non-elementary objects. DNA strands are composed by genes, RNA strands are composed by parts corresponding to the transcription of individual genes, and proteins are composed by parts having the role of interaction sites (or domains). Other molecules are considered as elementary objects, even if they are complexes. Membranes are considered as elementary objects, in the sense that we do not describe them at the level of the lipids they are made of. The only interesting properties of a membrane are that it may contain something (hence, create a compartment) and that it may have molecules on its surface.

Now, we select the biomolecular events of interest. The simplest kind of event is the change of state of an elementary object. Then, we may have interactions between molecules: in particular complexation, decomplexation and catalysis. These interactions may involve single elements of non–elementary molecules (DNA and RNA strands, and proteins). Moreover, we may have interactions between membranes and molecules: in particular a molecule may cross or join a membrane. Finally, we may have interactions between membranes: in this case there may be many kinds of interactions (fusion, division, etc...).

The guidelines for the abstraction of biomolecular entities and events into CLS are given in Table 1 and Table 2, respectively. Entities are associated with CLS terms: elementary objects are modeled as alphabet symbols, non-elementary objects as CLS sequences and membranes as looping sequences. Biomolecular events are associated with CLS rewrite rules. In the figure we give some examples of rewrite rules for each type of event. The list of examples is not complete: one could imagine also rewrite rules for the description of complexation/decomplexation events involving more than two molecules, or catalysis events in which the catalyzing molecule is on a membrane and the catalyzed event occurs in its content, or more complex interactions between membranes. We remark that in the second example of rewrite rule associated with the complexation event we have that one of the two molecules which are involved should be either an elementary object or a protein modeled as a single alphabet symbol. As before, this is caused by the problem of modeling protein interaction at the domain level. This problem is solved by the extension of CLS with links, called LCLS, we shall describe in the following.

2.3 Examples

A well-known example of biomolecular system is the epidermal growth factor (EGF) signal transduction pathway [26, 19]. If EGF proteins are present in the environment of a cell, they should be interpreted as a proliferation signal from the environment, and hence the cell should react by synthesizing proteins which stimulate its proliferation. A cell recognizes the EGF signal because it has on its membrane some EGF receptor proteins (EGFR), which are transmembrane proteins (they have some intra-cellular and some extra-cellular domains). One of the extra-cellular domains binds to one EGF protein in the environment, forming a signal-receptor complex on the membrane. This causes a conformational change on the receptor protein that enables it to bind to another one signal-receptor complex. The formation of the binding of the two signal-receptor complexes (called dimerization) causes the phosphorylation of some intra-cellular domains of the dimer. This, in turn, causes the internal domains of the dimer to be recognized by a protein that is inside the cell (in the cytoplasm), called SHC. The protein SHC binds to the dimer, enabling a long chain of protein-protein interactions, which finally activate some proteins, such as one called ERK, which bind to the DNA and stimulate synthesis of proteins for cell proliferation.

Now, we use CLS to build a model of the first steps of the EGF signaling pathway up to the binding of the signal-receptor dimer to the SHC protein. In the following we shall refine the model by using the LCLS extension to describe interactions at the domain level.

We model the EGFR, EGF and SHC proteins as the alphabet symbols EGFR, EGF and SHC, respectively. The cell is modeled as a looping sequence (representing its external membrane), initially composed only by EGFR symbols, containing SHC symbols and surrounded by EGF symbols. The rewrite rules modeling the first steps of the pathway are the following:

$$EGF \mid \left(EGFR \cdot \widetilde{x}\right)^{L} \mid X \quad \mapsto \quad \left(CMPLX \cdot \widetilde{x}\right)^{L} \mid X \tag{R1}$$

$$\left(DIM\cdot\widetilde{x}\right)^{L} \ \ X \quad \mapsto \quad \left(DIMp\cdot\widetilde{x}\right)^{L} \ \ X$$
 (R3)

$$\left(DIMp\cdot\widetilde{x}\right)^{L} \mid (SHC \mid X) \mapsto \left(DIMpSHC\cdot\widetilde{x}\right)^{L} \mid X$$
 (R4)

Rule R1 describes the binding of a EGF protein to a EGFR receptor protein on the membrane surface. The result of the binding is a signal-receptor complex denoted CMPLX. Rule R2 describes the dimerization of two signal-receptor complex, the result is denoted DIM. Rule R3 describes the phosphorylation (and activation) of a signal-receptor dimer, that is the replacement of a DIM symbol with a DIMp symbol. Finally, rule R4 describes the binding of an active dimer DIMp with a SHC protein contained in the cytoplasm. The result is a DIMpSHCsymbol placed on the membrane surface.

A possible initial term for the model in this example is given by a looping sequence composed by some EGFR symbols, containing some SHC symbols and

with some EGF symbols outside. A possible evolution of such a term by means of application of the given rewrite rules is the following (we write on each transition the name of the rewrite rule applied):

$$EGF \mid EGF \mid (EGFR \cdot EGFR \cdot EGFR \cdot EGFR)^{L} \mid (SHC \mid SHC)$$

$$\xrightarrow{(R1)} EGF \mid (EGFR \cdot CMPLX \cdot EGFR \cdot EGFR)^{L} \mid (SHC \mid SHC)$$

$$\xrightarrow{(R1)} (EGFR \cdot CMPLX \cdot EGFR \cdot CMPLX)^{L} \mid (SHC \mid SHC)$$

$$\xrightarrow{(R2)} (EGFR \cdot DIM \cdot EGFR)^{L} \mid (SHC \mid SHC)$$

$$\xrightarrow{(R3)} (EGFR \cdot DIMp \cdot EGFR)^{L} \mid (SHC \mid SHC)$$

$$\xrightarrow{(R4)} (EGFR \cdot DIMpSHC \cdot EGFR)^{L} \mid SHC$$

We show another example of modeling of a biomolecular system with CLS, that is the modeling of a simple gene regulation process. This kind of processes are essential for cell life as they allow a cell to regulate the production of proteins that may have important roles for instance in metabolism, growth, proliferation and differentiation.

The example we consider is as follows: we have a simple DNA fragment consisting of a sequence of three genes. The first, denoted p, is called *promoter* and is the place where a *RNA polymerase* enzyme (responsible for translation of DNA into RNA) binds to the DNA. The second, denoted o, is called *operator* and it is the place where a *repressor* protein (responsible for regulating the activity of the RNA polymerase) binds to the DNA. The third, denoted as g, is the gene that encodes for the protein whose production is regulated by this process.

When the repressor is not bound to the DNA, the RNA polymerase can scan the sequence of genes and transcribe gene g into a piece of RNA that will be later translated into the protein encoded by g. When the repressor is bound to the DNA, it becomes an obstacle for the RNA polymerase that cannot scan any more the sequence of genes.

The CLS model of this simple regulation process is a follows. The sequence of genes is represented as the CLS sequence $p \cdot o \cdot g$, the RNA polymerase enzyme as *polym*, the repressor protein as *repr*, and the piece of RNA obtained by the translation of gene g as *rna*. The rewrite rules describing the process are the following:

$$polym \mid p \cdot \widetilde{x} \quad \mapsto \quad pp \cdot \widetilde{x} \tag{R1}$$

$$repr \mid \widetilde{x} \cdot o \cdot \widetilde{y} \quad \mapsto \quad \widetilde{x} \cdot ro \cdot \widetilde{y} \tag{R2}$$

$$pp \cdot o \cdot \widetilde{x} \mapsto p \cdot po \cdot \widetilde{x}$$
 (R3)

- $\widetilde{x} \cdot po \cdot g \mapsto \widetilde{x} \cdot o \cdot pg$ (R4)
- $\widetilde{x} \cdot pg \mapsto polym \mid rna \mid \widetilde{x} \cdot g$ (R5)

Rules R1 and R2 describe the binding of the RNA polymerase and of the repressor to the corresponding genes in the DNA sequences. The results of these

bindings are that the symbols representing the two genes are replaced by pp and ro, respectively. Rules R3, R4 and R5 describe the activity of the RNA polymerase enzyme in the absence of the repressor: it moves from gene p to gene o in rule R3, then it moves from gene o to gene g in rule R4, and finally it produces the RNA fragment and leaves the DNA in rule R5. Note that, in order to apply rule R3, the repressor must be not bound to the DNA.

The only possible evolution of a term representing an initial situation in which no repressors are present is

$$\begin{array}{cccc} polym \mid p \cdot o \cdot g & \xrightarrow{(R1)} & pp \cdot o \cdot g & \xrightarrow{(R3)} & p \cdot po \cdot g \\ \\ \xrightarrow{(R4)} & p \cdot o \cdot pg & \xrightarrow{(R5)} & polym \mid rna \mid p \cdot o \cdot g \end{array}$$

that represent the case in which the RNA polymerase enzyme can scan the DNA sequence and transcribe gene g into a piece of RNA. When the repressor is present, instead, a possible evolution is

$$polym \mid p \cdot o \cdot g \quad \xrightarrow{(R1)} \quad pp \cdot o \cdot g \quad \xrightarrow{(R2)} \quad pp \cdot ro \cdot g$$

and it corresponds to a situation in which the repressor stops the transcription of the gene by hampering the activity of the RNA polymerase.

3 Two Extensions of CLS

In this section we describe two extensions of CLS. The first, Stochastic CLS, allows describing quantitative aspects of the modeled systems, such as the time spent by occurrences of chemical reactions. The second, CLS with links, allows describing protein interaction more precisely at a lower level of abstraction, namely at the domain level.

3.1 Stochastic CLS

In CLS only qualitative aspects of biological systems are considered, such as their structure and the presence (or the absence) of certain molecules. As a consequence, on CLS models it is only possible to verify properties such as the reachability of particular states or causality relationships between events. It would be interesting to verify also properties such as the time spent to reach a particular state, or the probability of reaching it. To face this problem, in [6] we have developed a stochastic extension of CLS, called *Stochastic CLS*, in which quantitative aspects, such as time and probability are taken into account.

The standard way of extending a formalism to model quantitative aspects of biological systems is by incorporating the stochastic framework developed by Gillespie with its simulation algorithm for chemical reactions [12] in the semantics of the formalism. This has been done, for instance, for the π -calculus [20, 22]. The



Fig. 2. Simulation result of the regulation process: number of RNA molecules over time.

idea of Gillespie's algorithm is that a rate constant is associated with each chemical reaction that may occur in the system. Such a constant is obtained by multiplying the kinetic constant of the reaction by the number of possible combinations of reactants that may occur in the system. The resulting rate constant is then used as the parameter of an exponential distribution modeling the time spent between two occurrences of the considered chemical reaction.

The use of exponential distributions to represent the (stochastic) time spent between two occurrences of chemical reactions allows describing the system as a Continuous Time Markov Chain (CTMC), and consequently it allows verifying properties of the described system by means of analytic means and by means of stochastic model checkers.

In Stochastic CLS, incorporating Gillespie's stochastic framework is not a simple exercise. The main difficulty is counting the number of possible reactant combinations of the chemical reaction described by a rewrite rule. This means counting the number of different positions where the rewrite rule can be applied, by taking into account that rules may contain variables. We have defined the Stochastic CLS in [6], and showed how to derive a CTMC from the semantics of a system modeled in Stochastic CLS. This allows performing simulation and verification of properties of the described systems, for instance by using stochastic model checkers, such as PRISM [13].

Let us consider the simple regulation process we modeled with CLS in Section 2.3. We now extend the CLS model by including a kinetic constant in each rewrite rule. The result is a Stochastic CLS model. In order to make the model a little more realistic we add two rewrite rules describing the unbinding of the RNA polymerase and of the repressor from the DNA. Hence, the rewrite rules of the Stochastic CLS model are the following:

$$polym \mid p \cdot \widetilde{x} \stackrel{0.1}{\longmapsto} pp \cdot \widetilde{x} \tag{R1}$$

$$pp \cdot \widetilde{x} \xrightarrow{2} polym \mid p \cdot \widetilde{x}$$
 (R1')

$$repr \mid \widetilde{x} \cdot o \cdot \widetilde{y} \stackrel{1}{\longmapsto} \widetilde{x} \cdot ro \cdot \widetilde{y} \tag{R2}$$

$$\widetilde{x} \cdot ro \cdot \widetilde{y} \stackrel{10}{\longmapsto} repr \mid \widetilde{x} \cdot o \cdot \widetilde{y} \tag{R2'}$$

$$pp \cdot o \cdot \widetilde{x} \xrightarrow{100} p \cdot po \cdot \widetilde{x}$$
 (R3)

$$\widetilde{x} \cdot po \cdot g \xrightarrow{100} \widetilde{x} \cdot o \cdot pg$$
 (R4)

$$\widetilde{x} \cdot pg \xrightarrow{30} polym \mid rna \mid \widetilde{x} \cdot g$$
 (R5)

We developed a simulator based on Stochastic CLS, and we used it to study the behavior of the regulation process. In particular, we performed simulations by varying the quantity of repressors and we observed the production of RNA fragments in each case. The initial configuration of the system is given by the following term

$$\underbrace{repr \mid \ldots \mid repr}_{n} \mid \underbrace{polym \mid \ldots \mid polym}_{100} \mid p \cdot o \cdot g$$

and we performed simulations with n = 0, 10, 25 and 50. The results of the simulations are shown in Figure 2. By varying the number of repressors from 0 to 50 the rate of transcription of the DNA into RNA molecules decreases.

3.2 CLS with Links (LCLS)

A formalism for modeling proteins interactions at the domain level was developed in the seminal paper by Danos and Laneve [11], and extended in [14]. This formalism allows expressing proteins by a node with a fixed number of domains; binding between domains allow complexating proteins. In this section we extend CLS to represent proteins interaction at the domain level. Such an extension, called Calculus of Linked Looping Sequences (LCLS), is obtained by labeling elementary components of sequences. Two elements with the same label are considered to be linked.

To model a protein at the domain level in CLS it would be natural to use a sequence with one symbol for each domain. However, the binding between two domains of two different proteins, that is the linking between two elements of two different sequences, cannot be expressed in CLS. To represent this, we extend CLS by labels on basic symbols. If in a term two symbols appear having the same label, we intend that they represent domains which are bound to each other. If in a term there is a symbol with a label and no other symbol with the same label, we intend that the term represents only a part of a system we model, and that the symbol will be linked to some other symbol in another part of the term representing the full model.

As membranes create compartments, elements inside a looping sequence cannot be linked to elements outside. Elements inside a membrane can be linked either to other elements inside the membrane or to elements of the membrane itself. An element can be linked at most to another element.

As an example, we model in LCLS the first steps of the EGF pathway described before. We model the EGFR protein as the sequence $R_{E1} \cdot R_{E2} \cdot R_{I1} \cdot R_{I2}$ where R_{E1} and R_{E2} are two extra-cellular domains and R_{I1} and R_{I2} are two intra-cellular domains. The membrane of the cell is modeled as a looping sequence which could contain EGFR proteins. Outside the looping sequence (i.e. in the environment) there could be EGF proteins, and inside (i.e. in the cytoplasm) there could be SHC proteins. Rewrite rules modeling the pathway are the following:

$$EGF \mid (R_{E1} \cdot \widetilde{x})^L \mid X \mapsto (sR_{E1} \cdot \widetilde{x})^L \mid X$$
 (R1)

$$\left(sR_{E1} \cdot R_{E2} \cdot x \cdot y \cdot \widetilde{x} \cdot sR_{E1} \cdot R_{E2} \cdot z \cdot w \cdot \widetilde{y} \right)^{L} \ \rfloor \ X \quad \mapsto \\ \left(sR_{E1} \cdot R_{E2}^{1} \cdot x \cdot y \cdot sR_{E1} \cdot R_{E2}^{1} \cdot z \cdot w \cdot \widetilde{x} \cdot \widetilde{y} \right)^{L} \ \rfloor \ X$$
 (R2)

$$\left(R_{E2}^{1} \cdot R_{I1} \cdot \widetilde{x} \cdot R_{E2}^{1} \cdot R_{I1} \cdot \widetilde{y} \right)^{L} \downarrow X \quad \mapsto \\ \left(R_{E2}^{1} \cdot PR_{I1} \cdot \widetilde{x} \cdot R_{E2}^{1} \cdot R_{I1} \cdot \widetilde{y} \right)^{L} \downarrow X$$
 (R3)

$$\left(R_{E2}^{1} \cdot PR_{I1} \cdot \widetilde{x} \cdot R_{E2}^{1} \cdot R_{I1} \cdot \widetilde{y} \right)^{L} \downarrow X \mapsto \left(R_{E2}^{1} \cdot PR_{I1} \cdot \widetilde{x} \cdot R_{E2}^{1} \cdot PR_{I1} \cdot \widetilde{y} \right)^{L} \downarrow X$$
 (R4)

$$\begin{pmatrix} R_{E2}^{1} \cdot PR_{I1} \cdot R_{I2} \cdot \widetilde{x} \cdot R_{E2}^{1} \cdot PR_{I1} \cdot R_{I2} \cdot \widetilde{y} \end{pmatrix}^{L} \mid (SHC \mid X) \mapsto \\ \begin{pmatrix} R_{E2}^{1} \cdot PR_{I1} \cdot R_{I2}^{2} \cdot \widetilde{x} \cdot R_{E2}^{1} \cdot PR_{I1} \cdot R_{I2} \cdot \widetilde{y} \end{pmatrix}^{L} \mid (SHC^{2} \mid X)$$
 (R5)

Rule R1 represents the binding of the EGF protein to the receptor domain R_{E1} with sR_{E1} as a result. Rule R2 represents that when two EGFR proteins activated by proteins EGF occur on the membrane, they may bind to each other to form a dimer (shown by the link 1). Rule R3 represents the phosphorylation of one of the internal domains R_{I1} of the dimer, and rule R4 represents the phosphorylation of the other internal domain R_{I1} of the dimer. The result of each phosphorylation is pR_{I1} . Rule R5 represents the binding of the protein SHC in the cytoplasm to an internal domain R_{I2} of the dimer. Remark that the binding of SHC to the dimer is represented by the link 2, allowing the protein SHC to continue the interactions to stimulate cell proliferation.

Let us denote the sequence $R_{E1} \cdot R_{E2} \cdot R_{I1} \cdot R_{I2}$ by EGFR. By starting from a cell with some EGFR proteins on its membrane, some SHC proteins in the cytoplasm and some EGF proteins in the environment, a possible evolution is the following:

$$\begin{split} & EGF \mid EGF \mid \left(EGFR \cdot EGFR \cdot EGFR\right)^{L} \mid (SHC \mid SHC) \\ \hline (R1) & EGF \mid \left(sR_{E1} \cdot R_{E2} \cdot R_{I1} \cdot R_{I2} \cdot EGFR \cdot EGFR\right)^{L} \mid (SHC \mid SHC) \\ \hline (R1) & \left(sR_{E1} \cdot R_{E2} \cdot R_{I1} \cdot R_{I2} \cdot EGFR \cdot sR_{E1} \cdot R_{E2} \cdot R_{I1} \cdot R_{I2}\right)^{L} \mid (SHC \mid SHC) \\ \hline (R2) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot R_{I1} \cdot R_{I2} \cdot sR_{E1} \cdot R_{E2}^{1} \cdot R_{I1} \cdot R_{I2} \cdot EGFR\right)^{L} \mid (SHC \mid SHC) \\ \hline (R3) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2} \cdot sR_{E1} \cdot R_{E2}^{1} \cdot R_{I1} \cdot R_{I2} \cdot EGFR\right)^{L} \mid (SHC \mid SHC) \\ \hline (R4) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2} \cdot sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2} \cdot EGFR\right)^{L} \mid (SHC \mid SHC) \\ \hline (R5) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2}^{2} \cdot sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2} \cdot EGFR\right)^{L} \mid (SHC \mid SHC) \\ \hline (R5) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2}^{2} \cdot sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2} \cdot EGFR\right)^{L} \mid (SHC^{2} \mid SHC) \\ \hline (R5) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2}^{2} \cdot sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2} \cdot EGFR\right)^{L} \mid (SHC^{2} \mid SHC) \\ \hline (R5) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2}^{2} \cdot sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2} \cdot EGFR\right)^{L} \mid (SHC^{2} \mid SHC) \\ \hline (R5) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2}^{2} \cdot sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2} \cdot EGFR\right)^{L} \mid (SHC^{2} \mid SHC) \\ \hline (R5) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2}^{2} \cdot sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2} \cdot EGFR\right)^{L} \mid (SHC^{2} \mid SHC) \\ \hline (R5) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2}^{2} \cdot sR_{E1} \cdot R_{E2}^{1} \cdot pR_{I1} \cdot R_{I2} \cdot EGFR\right)^{L} \mid (SHC^{2} \mid SHC) \\ \hline (R5) & \left(sR_{E1} \cdot R_{E2}^{1} \cdot pR_{E1} \cdot R_{E2}^{2} \cdot$$

4 CLS and Membranes

What could seem strange in CLS is the use of looping sequences for the description of membranes, as sequencing is not a commutative operation and this do not correspond to the usual fluid representation of membranes in which objects can move freely. What one would expect is to have a multiset or a parallel composition of objects on a membrane. In the case of CLS, what could be used is a parallel composition of sequences. To address this problem, we define an extension of CLS, called CLS+, in which the looping operator can be applied to a parallel composition of sequences, and we show that we can translate quite easily CLS+ models into CLS ones.

4.1 Definition of CLS+

Terms in CLS+ are defined as follows.

Definition 6 (Terms). Terms T, branes B, and sequences S of CLS+ are given by the following grammar:

$$T ::= S | (B)^{L} \rfloor T | T | T$$
$$B ::= S | S | S$$
$$S ::= \epsilon | a | S \cdot S$$

where a is a generic element of \mathcal{E} . We denote with \mathcal{T} the infinite set of terms, with \mathcal{B} the infinite set of branes, and with \mathcal{S} the infinite set of sequences.

The structural congruence relation of CLS+ is a trivial extension of the one of CLS. The only difference is that commutativity of branes replaces rotation of looping sequences.

Definition 7 (Structural Congruence). The structural congruence relations \equiv_S , \equiv_B and \equiv_T are the least congruence relations on sequences, on branes and on terms, respectively, satisfying the following rules:

$$S_1 \cdot (S_2 \cdot S_3) \equiv_S (S_1 \cdot S_2) \cdot S_3 \qquad S \cdot \epsilon \equiv_S \epsilon \cdot S \equiv_S S$$

$$S_{1} \equiv_{S} S_{2} \text{ implies } S_{1} \equiv_{B} S_{2}$$

$$B_{1} \mid B_{2} \equiv_{B} B_{2} \mid B_{1} \qquad B_{1} \mid (B_{2} \mid B_{3}) \equiv_{B} (B_{1} \mid B_{2}) \mid B_{3} \qquad B \mid \epsilon \equiv_{B} B$$

$$S_{1} \equiv_{S} S_{2} \text{ implies } S_{1} \equiv_{T} S_{2}$$

$$B_{1} \equiv_{B} B_{2} \text{ implies } (B_{1})^{L} \mid T \equiv_{T} (B_{2})^{L} \mid T$$

$$T_{1} \mid T_{2} \equiv_{T} T_{2} \mid T_{1} \qquad T_{1} \mid (T_{2} \mid T_{3}) \equiv_{T} (T_{1} \mid T_{2}) \mid T_{3} \qquad T \mid \epsilon \equiv_{T} T \quad (\epsilon)^{L} \mid \epsilon \equiv \epsilon$$

Now, to define patterns in CLS+ we consider an additional type of variables with respect of CLS, namely brane variables. We assume a set of brane variables BV ranged over by $\overline{x}, \overline{y}, \overline{z}, \dots$

Definition 8 (Patterns). Patterns P, brane patterns BP and sequence patterns SP of CLS+ are given by the following grammar:

$$P ::= SP | (BP)^{L} \rfloor P | P | P | X$$

$$BP ::= SP | SP | SP | \overline{x}$$

$$SP ::= \epsilon | a | SP \cdot SP | \widetilde{x} | x$$

where a is a generic element of \mathcal{E} , and $X, \overline{x}, \widetilde{x}$ and x are generic elements of TV, BV, SV and \mathcal{X} , respectively. We denote with \mathcal{P} the infinite set of patterns.

As usual, rewrite rules are pairs of patterns.

Definition 9 (Rewrite Rules). A rewrite rule is a pair of patterns (P_1, P_2) , denoted with $P_1 \mapsto P_2$, where $P_1, P_2 \in PP$, $P_1 \not\equiv \epsilon$ and such that $Var(P_2) \subseteq Var(P_1)$. We denote with \Re the infinite set of all the possible rewrite rules.

Now, differently from CLS, we have that a rule such as $a \mid b \mapsto c$ could be applied to elements of a looping sequence. For instance, $a \mid b \mapsto c$ can be applied to the term $(a \mid b)^L \mid d$ so to obtain the term $(c)^L \mid d$. However, a rule such as $(a)^L \mid b \mapsto c$ still cannot be applied to elements of a looping sequences, as $((a)^L \mid b)^L \mid c$ is not a CLS+ term.

The rules that can be applied to elements of a looping sequence are those having the form (B_1, B_2) with $B_1, B_2 \in \mathcal{B}$. We call these rules brane rules and we denote as $\Re_{\mathcal{B}} \subset \Re$ their infinite set. Now, in the semantics of CLS+ we have to take into account brane rules and allow them to be applied also to elements of looping sequences. Hence, we define the semantics as follows.

Definition 10 (Semantics). Given a set of rewrite rules $\mathcal{R} \subseteq \mathfrak{R}$, and a set of brane rules $\mathcal{R}_{\mathcal{B}} \subseteq \mathcal{R}$, such that $(\mathcal{R} \setminus \mathcal{R}_{\mathcal{B}}) \cap \mathfrak{R}_{\mathcal{B}} = \emptyset$, the semantics of CLS is the least transition relation \rightarrow on terms closed under \equiv , and satisfying the following inference rules:

$$\frac{(P_1, P_2) \in \mathcal{R} \quad P_1 \sigma \neq \epsilon \quad \sigma \in \Sigma}{P_1 \sigma \to P_2 \sigma} \quad \frac{T_1 \to T_2}{T \mid T_1 \to T \mid T_2} \quad \frac{T_1 \to T_2}{(B)^L \mid T_1 \to (B)^L \mid T_2} \\
\frac{(BP_1, BP_2) \in \mathcal{R}_B \quad BP_1 \sigma \neq \epsilon \quad \sigma \in \Sigma}{BP_1 \sigma \to_{\mathcal{B}} BP_2 \sigma} \\
\frac{B_1 \to_{\mathcal{B}} B_2}{B \mid B_1 \to_{\mathcal{B}} B \mid B_2} \quad \frac{B_1 \to_{\mathcal{B}} B_2}{(B_1)^L \mid T \to (B_2)^L \mid T}$$

where $\rightarrow_{\mathcal{B}}$ is a transition relation on branes, and where the symmetric rules for the parallel composition of terms and of branes are omitted.

In the definition of the semantics of CLS+ we use an additional transition relation $\rightarrow_{\mathcal{B}}$ on branes. This relation is used to describe the application of a brane rule to elements of a looping sequence. As usual, a CLS+ model is composed by a term, representing the initial state of the modeled system, and a set of rewrite rules.

In the following section we show that CLS+ models can be translated into CLS models. The translation into CLS is compositional and preserves the semantics of the model.

4.2 Translating CLS+ into CLS

The first step of the translation of a CLS+ model into CLS is a preprocessing procedure. For each brane rule (BP_1, BP_2) in the CLS+ model, we add to the set of rules of the model a new rule, namely $((BP_1 | \overline{x})^L \rfloor X, (BP_2 | \overline{x})^L \rfloor X)$. This new rule is redundant in the model, as every time it can be applied to a CLS+ term, also the original one can be applied with the same result. However, the translation we are going to define will translate the original rule into a CLS rule that will be applicable only inside looping sequences, or at the top level of the term, and will translate the new rule into a CLS rule applicable only to elements that compose a looping sequence.

Now, the translation of CLS+ into CLS consists mainly of an encoding function, denoted {[·]}, which maps CLS+ patterns into CLS patterns. This encoding function will be used to translate each rewrite rule of the CLS+ model into a rewrite rule for the corresponding CLS model, and to translate the term representing the initial state of the system in the CLS+ model into a CLS term for the corresponding CLS model.

The encoding function for CLS+ patterns is defined as follows. We assume a total and injective function from brane variables into a subset of term variables

that are never used in CLS models. More easily, we assume brane variables to be a subset of the term variables of CLS. Moreover, we assume *in* and *out* to be symbols of the alphabet \mathcal{E} never used in CLS models.

The encoding follows the "ball-bearing" technique described by Cardelli in [7]. Intuitively, every CLS+ looping sequence is translated into a couple of CLS looping sequences, one contained in the other, with the brane patterns of the CLS+ looping sequence between the two corresponding CLS looping sequences.

Definition 11 (Encoding Function). The encoding function $\{[\cdot]\}$ maps CLS+ patterns into CLS patterns, and is given by the following recursive definition:

$$\{[SP]\} = SP$$

$$\{[X]\} = X$$

$$\{[(BP)^{L} \ \]P]\} = (out)^{L} \ \ (BP \ \ (in)^{L} \ \ \{[P]\})$$

$$\{[P_{1} \ \ P_{2}]\} = \{[P_{1}]\} \ \ \{[P_{2}]\}$$

A CLS rewrite rule is obtained from each CLS+ rewrite rule of the translated model by applying the encoding function to the two patterns of the rule. More precisely, given a CLS+ rule $P_1 \mapsto P_2$, the corresponding CLS rule is $(in)^L \ | \ (\{P_1\}\} | X) \mapsto (in)^L \ | \ (\{P_2\}\} | X)$ where X is a term variable that does not occur in P_1 and P_2 . For example, by applying the encoding to the two patterns of the CLS+ rewrite rule

$$R = b \cdot x \mid c \mapsto b \cdot x$$

we obtain

$$R_{\{\!\!\left[\cdot\right]\!\!\}} = \left(in\right)^L \, \left| \, \left(b \cdot x \mid c \mid X\right) \right. \mapsto \left. \left(in\right)^L \, \right| \left(b \cdot x \mid X\right).$$

The encoding of a CLS+ term into a CLS term is as follows: given a CLS+ term T the corresponding CLS term is $(in)^L \downarrow \{[T]\}$. In this case we have that the encoding function never encounters variables. Consider, as an example, the following CLS+ term:

$$T = a \mid \left(c \mid d \mid b \cdot b \mid d\right)^{L} \rfloor d$$

the corresponding CLS term is as follows:

$$T_{\{\!\left[\cdot\right]\!\}} = \left(in\right)^L \, \rfloor \, \left(a \mid \left(out\right)^L \, \rfloor \, \left(c \mid d \mid b \cdot b \mid d \mid \left(in\right)^L \, \rfloor \, d\right)\right)$$

Now, it is easy to see that R can be applied to T, because parallel components in the looping sequence can be commuted, and the result of the application is

$$T' = a \mid (b \cdot b \mid d \mid d)^{L} \rfloor d$$

but the corresponding CLS rewrite rule $R_{\{\![\cdot]\!]}$ cannot be applied to $T_{\{\![\cdot]\!]}$. However, we have that $R \in \mathcal{R}_{\mathcal{B}}$, and hence, by the preprocessing phase described above, we have that also

$$R' = \left(b \cdot x \mid c \mid \overline{x}\right)^L \, \rfloor \, X \quad \mapsto \quad \left(b \cdot x \mid \overline{x}\right)^L \, \rfloor \, X$$

is a rule of the CLS+ model. By translating rule R' we obtain

$$R'_{\{\!\!\{\cdot\}\!\}} = (in)^L \, \rfloor \, ((out)^L \, \rfloor \, (b \cdot x \mid c \mid \overline{x} \mid (in)^L \, \rfloor \, X) \mid Y) \quad \mapsto \\ (in)^L \, \rfloor \, ((out)^L \, \rfloor \, (b \cdot x \mid \overline{x} \mid (in)^L \, \rfloor \, X) \mid Y)$$

that can be applied to $T_{\{\![\cdot]\!]}$. The result of the application is

$$(in)^{L} \rfloor (a \mid (out)^{L} \rfloor (b \cdot b \mid d \mid d \mid (in)^{L} \rfloor d))$$

that corresponds exactly to the encoding of T'.

4.3 CLS, Brane Calculi and P Systems

Brane Calculi are a family of process calculi specialized in the description of membrane activity, and they allow associating processes with membranes of a membrane structure. Each process is composed by actions whose execution has an effect on the membrane structure. Some examples of actions are phagocytosis (a membrane engulfs another one), exocytosis (a membrane expels another one), and pinocytosis (a new membrane is created inside another one). These three actions are enough to define the simplest of Brane Calculi, namely the PEP calculus. Other actions, such as fusion of membranes and mitosis can be used to define different calculi of the family. Moreover, extensions of Brane Calculi allow describing interactions with molecules and complexes, such as letting them enter and exit membranes.

We have given a sound and complete encoding of the PEP calculus into CLS in [3, 15]. Here, to recall shortly the encoding technique, we give a very simple example of PEP system and we show its translation into CLS. The PEP system we consider is the following

$$\phi(\diamond) \circ \phi^{\perp}(\mathbf{0})(\diamond)$$

representing two adjacent membranes $\phi_n(|\diamond|)$ and $\phi_n^{\perp}(\mathbf{0})(|\diamond|)$ (\diamond denotes juxtaposition) both containing nothing of relevant (what is between brackets (|) is the content of the membrane and \diamond is the null system). The processes associated with the two membranes are ϕ and $\phi^{\perp}(\mathbf{0})$, respectively, representing two complementary phagocytosis actions: the first says that the membrane it is associated with can be engulfed by another membrane, and the second that the membrane it is associated with can engulf another membrane, that will be surrounded by another new membrane whose associated process is the parameter of the action (in this case it is the idle process $\mathbf{0}$). Hence, in accordance with the semantics of the PEP calculus, we have that the only transition that can be performed by the system is the following, leading to a system that is equivalent to the null system \diamond :

$$\phi(|\diamond|) \circ \phi^{\perp}(\mathbf{0})(|\diamond|) \rightarrow \mathbf{0}(|\mathbf{0}||\diamond|)|) \equiv \diamond$$

By applying the encoding to the system we obtain the following CLS term T:

$$act \cdot circ \cdot e \cdot brane \cdot b \cdot \phi \cdot a \cdot \mathbf{0} \cdot a \cdot b \cdot \mathbf{0} \cdot e \cdot brane \cdot d \cdot \phi^{\perp} \cdot c \cdot \mathbf{0} \cdot c \cdot \mathbf{0} \cdot c \cdot d \cdot \mathbf{0}$$

where act is a sort of program counter that precedes the symbol representing the next action to be executed, symbol circ represents \circ , symbol brane represents a membrane (|), symbols ϕ and ϕ^{\perp} represent the corresponding actions, symbol **0** represents the idle process and symbols a, b, c, d and e are used as separators of actions and parameters. The translation consists also of a set of CLS rewrite rules to be applied to terms obtained by the encoding of PEP systems. Such a set of rewrite rules does not depend on the encoded PEP system, hence it is always the same. By applying rewrite rules, the long sequence obtained from the encoding is transformed into a hierarchy of looping sequences corresponding to the membrane hierarchy in the original PEP system, then rewrite rules are applied that correspond to the semantics of the actions occurring in the processes associated with membranes.

Hence, by means of application of rewrite rules, the result of the encoding of the PEP system may evolve as follows (where \rightarrow^* represents a sequence of rewrite rule applications):

$$T \rightarrow act \cdot brane \cdot b \cdot \phi \cdot a \cdot \mathbf{0} \cdot a \cdot \mathbf{b} \cdot \mathbf{0} \mid act \cdot brane \cdot d \cdot \phi^{\perp} \cdot c \cdot \mathbf{0} \cdot c \cdot \mathbf{0} \cdot c \cdot d$$

$$\rightarrow^{*} (act \cdot \phi \cdot a \cdot \mathbf{0} \cdot a)^{L} \mid act \cdot \mathbf{0} \mid (act \cdot \phi^{\perp} \cdot c \cdot \mathbf{0} \cdot c \cdot \mathbf{0} \cdot c)^{L} \mid act \cdot \mathbf{0}$$

$$\rightarrow (act \cdot \mathbf{0})^{L} \mid (act \cdot \mathbf{0}) (act \cdot \mathbf{0})^{L} \mid (act \cdot \mathbf{0})^{L} \mid act \cdot \mathbf{0})$$

$$\rightarrow^{*} act \cdot \mathbf{0}$$

Differently from Brane Calculi, P Systems (in their most common formulation) do not allow describing complex membrane activities such as phagocytosis and exocytosis. However, they are specialized in the description of reactions between molecules which are placed in a compartment of a complex membrane structure.

A P System is a membrane structure (a nesting of membranes) in which there could be multisets of objects representing molecules. A set of multiset rewrite rules is associated with each membrane, and describe the reactions that may occur between the molecules contained in the membrane. The result of the application of a rewrite rule can either remain in the same membrane, or exit the membrane, or enter an inner membrane. Priorities can be imposed on rewrite rules, meaning that some rules can be applied only if some others cannot, and it is possible for a membrane to dissolve and release its content to in the environment.

A peculiarity of P Systems is that rewrite rules are applied in a fully–parallel manner, namely in one step of evolution of the system all rules are applied as many

times as possible (to different molecules), and this is one of the main differences with respect to CLS in which at each step one only rewrite rule is applied. We show that P Systems can be translated into CLS, and that the execution of a (fully parallel) step of a P System is simulated by a sequence of steps in CLS. A variant of P Systems, called Sequential P Systems, in which rules are applied sequentially is described in [10]. We do not consider the translation of this variant into CLS as it would be quite trivial and of little interest.



Fig. 3. A simple example of P System.

To recall the encoding technique, we give a simple example of P System and we show its translation into CLS. We focus on the translation of multiple parallelism, hence we consider a P System (depicted in Figure 3) consisting of a single membrane with only two rules, without priorities and without membrane dissolutions. We give a simplified translation: more details can be found in [15].

The alphabet of objects in the considered P System is $\{a, b, c\}$. A multiset of objects from this alphabet is represented by a CLS term as follows: let n_a, n_b and n_c be the number of occurrences of a, b and c in the multiset, respectively, then

$$a \cdot \overbrace{1 \cdot \ldots \cdot 1}^{n_a} | b \cdot \overbrace{1 \cdot \ldots \cdot 1}^{n_b} | c \cdot \overbrace{1 \cdot \ldots \cdot 1}^{n_c}$$

is the term representing the multiset. We choose this representation as it allows us checking whether an object is absent, by checking whether the corresponding symbol if followed by zero 1s. An empty multiset is represented as $a \mid b \mid c$.

The CLS term obtained by the translation of the considered P System is the following:

$$(1)^{L} \perp (Check \mid a \cdot 1 \mid b \cdot 1 \cdot 1 \mid c \mid r_{1} \mid r_{2} \mid (next)^{L} \perp (a \mid b \mid c))$$

where the membrane of the P System is represented by a looping sequence composed by the membrane label (in this case 1). Inside the looping sequence there is a *Check* symbol representing the current state of the system, the translation of the multiset of objects of the membrane, two symbols r_1 and r_2 corresponding to the evolutionary rules of the membrane, and an empty multiset surrounded by a looping sequence *next*. This empty multiset is used to store temporary information on the result of the application of evolutionary rules. The CLS rewrite rules obtained by the encoding of the considered P System are the following:

$$(1)^{L} \downarrow (X \mid Check \mid a \cdot 1 \cdot \widetilde{x} \mid r_{1}) \mapsto (1)^{L} \downarrow (X \mid Check \mid a \cdot 1 \cdot \widetilde{x} \mid r_{1} \cdot 1)$$
(C1)

$$(1)^{L} \rfloor (X \mid Check \mid a \mid r_{1}) \mapsto (1)^{L} \rfloor (X \mid Check \mid a \mid r_{1} \cdot 0)$$
(C2)

$$(1)^{L} \downarrow (X \mid Check \mid a \mid r_{1} \cdot z \mid r_{2}) \mapsto (1)^{L} \downarrow (X \mid Run \mid a \mid r_{1} \cdot z \mid r_{2} \cdot 0)$$
(C4)

$$(1)^{L} \downarrow (X \mid Check \mid b \mid r_{1} \cdot z \mid r_{2}) \mapsto (1)^{L} \downarrow (X \mid Run \mid b \mid r_{1} \cdot z \mid r_{2} \cdot 0)$$
(C5)

$$(1)^{L} \downarrow (X \mid Run \mid a \cdot 1 \cdot \widetilde{x} \mid b \cdot 1 \cdot \widetilde{y} \mid r_{2} \cdot 1 \mid (next)^{L} \downarrow (Y \mid c \cdot \widetilde{z})) \mapsto$$

$$(1)^{L} \downarrow (X \mid Run \mid a \cdot \widetilde{x} \mid b \cdot \widetilde{y} \mid r_{2} \cdot 1 \mid (next)^{L} \downarrow (Y \mid c \cdot 1 \cdot \widetilde{z}))$$

$$(R2)$$

$$(1)^{L} \downarrow (X \mid Run \mid a \mid r_{1} \cdot 1) \mapsto (1)^{L} \downarrow (X \mid Run \mid a \mid r_{1} \cdot 0)$$
(R3)

$$(1)^{L} \downarrow (X \mid Run \mid a \mid r_{2} \cdot 1) \mapsto (1)^{L} \downarrow (X \mid Run \mid a \mid r_{2} \cdot 0)$$
(R4)

$$(1)^{L} \rfloor (X \mid Run \mid b \mid r_{2} \cdot 1) \mapsto (1)^{L} \rfloor (X \mid Run \mid b \mid r_{2} \cdot 0)$$
(R5)

$$(1)^{L} \rfloor (X \mid Run \mid r_{1} \cdot 0 \mid r_{2} \cdot 0) \mapsto (1)^{L} \rfloor (X \mid Update \mid r_{1} \cdot 0 \mid r_{2} \cdot 0)$$
(R6)

$$(1)^{L} \rfloor (X \mid Update \mid (next)^{L} \rfloor (a \mid b \mid c)) \mapsto$$

$$(1)^{L} \rfloor (X \mid Check \mid (next)^{L} \rfloor (a \mid b \mid c))$$

$$(U2)$$

Rules (C1)–(C5) describe the steps performed by the system while it is in *Check* state: the objective of this phase is to test whether each evolutionary rule is applicable or not. When all rules have been tested, the systems moves into a state called *Run*, whose steps are given by the application of rules (R1)–(R6). In this second phase, evolutionary rules previously identified as applicable are actually applied, and the result of the application is stored inside the looping sequence *next*. Finally, when no evolutionary rule is further applicable, the system moves

into a state called Update, in which the content of the looping sequence next is used to reset the multiset of objects of the membrane by applying rule (U1)–(U2). When this update operation has been performed, the system moves back to the *Check* state.

5 Conclusions

We have surveyed the formalism CLS and a number of its variants from the point of view of its use for describing biological membranes. Verification and simulation tools have been developed for CLS and its variants and can be used to study properties of membrane systems. Via translations, these tools can be used to study systems described by other formalisms such as Brane Calculi and P Systems, capable of describing biological membranes.

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Membrane Computing in Connex Environment

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Summary. The Connex technology is presented as a possible way to implement efficiently membrane computations in Silicon environment. The opportunity is offered by the recent trend of promoting the parallel computation as a real competitor on the consumer market. The Connex environment has an integral parallel architecture, which is introduced and its main performances are presented. Some suggestions are provided about how to use the Connex environment as accelerator for membrane computation.

1 Introduction

The computation model of membrane computing can be supported by a specific physical environment or by non-specific, on-Silicon parallel architectures. The second way is investigated from the view point of the Connex technology: a highly integrated parallel machine.

Membrane computing summary:

The membrane computing model is based on multi-set rewriting rules applied on a membrane structure populated with objects belonging to a finite alphabet. The potential degree of parallelism is very high in P systems because in each step all possible rules are applied (see details in [Păun '02]).

Connex environment summary:

The Connex technology has an intensive integral parallel architecture [Stefan '06d]. The first embodiment of this technology (see [Stefan '06c]) targets the high definition TV market, but the chip CA1024 can be used also as a general purpose machine for data intensive computing. Application in graphics, data mining, neural network [Andonie '07], and communication are efficiently supported by the Connex technology. Then, why not for membrane computing!

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The Intel study:

Because, since 2002 the clock speed of the processor has improved less than 20%/year, after a long period characterized by around 50%/year, the promise of parallel computing starts to fascinate in a special way. Intel published seminal studies (see [Dubey '05], [Borkar '05]) about the next generation of parallel computers. The future processors will contain multi- or maybe many-processors optimized for the magic triad of **Recognition** – **Mining** – **Synthesis** (RMS). The main problem for this promised development is to find the way to program efficiently the next generation of parallel machines. New programming languages or more sophisticated computation models are needed to fructify the opportunities offered by the new coming parallel computation technologies. In this context membrane computing could play a very promising role.

The Berkeley study:

Rather than starting from the market opportunities, as Intel did with the RMS domains, the Berkeley approach [Asanovic '06] starts from their "13 dwarfs" (dense linear algebra, sparse linear algebra, spectral methods, ... finite state machines) identified as **parallel computational patterns** able to cover almost all the applications for the next few decades. While Intel takes into consideration a continuous transition from multi- to many-processors, the Berkeley approach is oriented from the start toward the many-processor systems working on data-intensive computation applications. Here is also the place for membrane computation if a good representation will be developed.

Application oriented vs. functionally oriented parallel architectures:

A complex, intense and general purpose application means usually a multithreaded approach. In contrast with it, there are functions involving data intense computations. By the rule, multi-processors are involved in the first case (because they are able to exploit thread-level parallelism), and many-processors are needed in the second case. A multi-processor has usually a MIMD architecture, rarely a SIMD architecture, and never a MISD type one. For a many-processor machine the architecture must be shaped starting from a functional approach, and by the rule involves all the special forms of possible parallelism.

Our functional approach:

The **integral parallel architecture** (IPA) is a parallel architecture derived starting from the computational model of *partial recursive functions* [Kleene '36]. The Turing machine model has been successfully used to ground various sequential computing architectures. Because the functional approach of Kleene is more related with circuits (which are intrinsic parallel structures) we consider there is a better fit between the functional recursive model and the parallel computation. The composition rule provides the best starting point to develop parallel architectures able to support efficiently the other two rules: the primitive recursion and the minimalization. If the 13 dwarfs will be able to cover the RMS domains, maybe then an IPA will be enough powerful to cover efficiently the 13 computational patterns emphasized by the seminal work done at Berkeley. A three level hierarchy results. It is topped by application domains (RMS), mediated by computational patterns (the 13 dwarfs), and grounded on various IPAs.

In the following sections the idea of IPA and the Connex environment are introduced in order to offer various suggestions for a *membrane computing accelerator*. Membrane computing being an intrinsic parallel computational model has the chance to open new ways toward the efficient use of parallel machines.

2 Integral Parallel Architecture (IPA)

Various taxonomies were proposed for parallel computations (see [Flynn '72] [Xavier & Iyengar '98]). All of them tell us about different forms of parallelism. We can discuss about many forms only when we use the parallel approach to accelerate specific computations. But, when a real complex and intensive computation must pe done, sometimes we can not use only one form of parallelism. Actual computations involve usually all possible forms. For example, using Flynn's taxonomy, MIMD or SIMD machines can be defined, but it is not so easy to define MIMD or SIMD application domains.

General purpose or even application domain oriented parallel machines must be able to perform all the forms of parallelism, no matter how these forms are segregated. We propose in the following a new taxonomy and a way to put together all the resulting forms of parallelism in order to solve efficiently data intensive computations.

2.1 Parallelism and Partial Recursiveness

We claim that the most suggestive classic computational model for defining parallel architectures is the model of partial recursive functions, because the rules defining it have direct correspondences in circuits – the intrinsic parallel support for computation.

Composition & basic parallel structures

The first rule, of composition, provides the basic parallel structures to be used in defining all the forms of parallelism. Let be m n-ary functions $h_i(x_0, \ldots x_{n-1})$, for $i = 0, 1, \ldots m - 1$, and an *m*-ary function $g(y_0, \ldots y_{m-1})$. Using them the the composition rule is defined as computing the following function:

$$f(x_0, \dots, x_{n-1}) = g(h_0(x_0, \dots, x_{n-1}), \dots, h_{m-1}(x_0, \dots, x_{n-1}))$$

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The associated physical structure (containing simple circuits or simple programmable machines) is represented in Figure 1.



Fig. 1. The physical structure associated to the composition rule. The composition of the function g with the functions h_0, \ldots, h_{m-1} implies a two-level system. The first level, performing in **parallel** m computations, is **serially** connected with the second level which performs a reduction function.

The following four particular, but meaningful forms (see Figure 2) can be emphasized:

1. data parallel composition: with n = m, each function $h_i = h$ depends on a single input variable x_i , for i = 0, 1, ..., n - 1, and g performs the identity function (see Figure 2a). Being given an input vector containing n scalars:

$$\mathbf{X} = \{x_0, x_1, \dots, x_{n-1}\}$$

the result is another vector:

$${h(x_0), h(x_1), \ldots, h(x_{n-1})}$$

2. speculative composition: with n = 1, i.e. $x_0 = x$, (see Figure 2b), and g performs the identity function. It computes a vector of functions:

$$\mathbf{H} = [h_0(x), \dots h_{n-1}(x)]$$

on the same scalar input x, generating a vector of results:

$$\mathbf{H}(x) = \{h_0(x), h_1(x), \dots, h_{n-1}(x)\}\$$

3. serial composition: with n = m = 1 (see Figure 2c). A "pipe" of two different machines receives a stream of n scalars as input:

$$\langle \mathbf{X} \rangle = \langle x_0, x_1, \dots, x_{n-1} \rangle$$



Fig. 2. The four simple forms of composition. a. Data parallel composition. b. Speculative composition. c. Serial composition. d. Reduction composition.

and provides another stream of scalars

$$< f(x_0), f(x_1), \ldots, f(x_{n-1}) > .$$

In the general case the function f(x) is a composition of more than two functions h and g. Thus, the function f can be expressed as a vector of functions **F** receiving as input a data stream $\langle \mathbf{X} \rangle$:

$$\mathbf{F} = [f_0, \dots f_{p-1}]$$

(in Figure 2c $\mathbf{F} = [h, g]$)

4. reduction composition: for each h_i performing the identity function (see Figure 2d), receives a vector $\{x_0, \ldots, x_{n-1}\}$ as input and provides the scalar, $g(x_0, \ldots x_{n-1})$ (it transforms a stream of vectors into a stream of scalars).

Concluding, the composition rule provides the context of defining computation using the following basic concepts:

scalar : xvector : $\mathbf{X} = \{x_0, x_1, \dots, x_{n-1}\}$ stream : $\langle \mathbf{X} \rangle = \langle x_0, x_1, \dots, x_{n-1} \rangle$ function : f(x)vector of functions :

- F = [f₀,...f_{p-1}] applied on streams
 F(x) = [f₀(x),...f_{p-1}(x)] applied on scalars.

Using the previous features all the requirement for the next two rules (primitive recursion, minimalization) are fulfilled.

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Primitive recursive rule

There are two ways to implement in parallel the primitive recursive rule. In both cases a lot of data is supposed available to be computed, i.e. there are vector or streams of data as inputs for a primitive recursive function.

The primitive recursive rule computes the function f(x, y) using the rule:

$$f(x,y) = h(x, f(x, y-1))$$

where: f(x, 0) = g(x). This rule can be translated in the following serial composition:

$$f(x,y) = h(x, h(x, h(x, \dots, h(x, g(x)) \dots)))$$

If the function f(x, y) must be computed for the vector of scalars $\mathbf{X} = \{y_0, y_1, \ldots, y_{n-1}\}$, then a data parallel structure is used. Each machine will compute, using a local *data loop*, the function $f(x, y_i)$ in $max(y_0, y_1, \ldots, y_{n-1})$ "cycles".

If the function f(x, y) must be computed for a stream of scalars, a time parallel structure is used. A "pipe" of n machines will receive in each "cycle" a new scalar from the stream of scalars. If y > n, then a *data loop* can be closed from the output of the pipe to its input.

Minimalization

Minimalization has also two kinds of parallel solutions: one using data parallel structures and another using time parallel structures.

The minimalization rule assumes

$$f(x) = min(y)[m(x, y) = 0]$$

i.e., the value of f(x) is the minimum y for which m(x, y) = 0.

The first, "brute force" implementation uses the speculative structure represented in Figure 2b, where each block computes a function which returns a pair containing a predicate and a scalar:

$$h_i = \{(m(x,i) = 0), i\}$$

after which reduction step (using a structure belonging to the class represented in Figure 2d) selects the smallest *i* from all pairs having the form $\{1, i\}$, if any, that were generated on the previous speculative composition level (all pairs of the form $\{0, i\}$ are ignored).

The second implementation occurs in time-parallel environments where speculation can be used to speed-up the pipe processing. **Reconfigurable pipes** can be conceived and implemented using special reduction features distributed along a pipe. Let be a pipe of functions described by the function vector:

$$\mathbf{P} = [f_0(x), \dots f_{p-1}(x)]$$

where $y_i = f_i(x)$, for i = 0, ..., p - 1. The associated reconfigurable pipe means to transform the original pipe characterized by:

$$\mathbf{P} = [\dots f_i(y_{i-1}), \dots]$$

into a pipe characterized by:

$$\mathbf{P} = [\dots f_i(y_{i-1}, \dots y_{i-s}), \dots]$$

where: $f_i(y_{i-1}, \ldots, y_{i-s})$ is a function or a program which decides in each step the variable to be involved in the current computation, selecting (which is one of the simplest reduction functions) one variable of $\{y_{i-1}, \ldots, y_{i-s}\}$. The maximum *degree* of speculation is s.

2.2 Functional Taxonomy of Parallel Computing

According to the previously identified simple form of compositions (see Figure 2) we propose a functional taxonomy of parallel computation. We will consider the following types of parallel processing:

- data-parallel computing : uses operators that take vectors as arguments and returns vectors, scalars (by reduction operations) or streams (input values for time-parallel computations); it is very similar to a SIMD machine
- time-parallel computing : uses operators that take streams as arguments and returns streams, scalars, or vectors (input values for data-parallel computations): it is a kind of MIMD machine which works to compute only one function (while a true MIMD performs multi-threading)
- speculative-parallel computing : with operators that take scalars as arguments and return vectors reduced to scalars using selection (used mainly to speed up timeparallel computations); this contains a true MISD-like structure (completely ignored in the current multi-processing environments).

An IPA is a parallel architecture featured with all kinds of parallelism.

2.3 IPA & Market Tendencies

The market tendencies emphasized in the Intel approach and based on Berkeley's dwarfs demand for an IPA. IPA is a many-core (not multi-core) architecture designed to support data intensive computations. It is supposed to work as an accelerator in a mono- or multi-core environment. For all the computational patterns emphasized in the Berkeley's view an IPA provides efficient solutions. Even for the 13th dwarf – Finite State Machine – the speculative- & time-parallel aspects of an IPA provides a solution. (Berkeley's view claims that "nothing helps".)

The need for solving real hard applications promotes IPA as an efficient actual solution.

3 The Connex System

3.1 Structural description

The first embodiment of an IPA is the Connex System. It is part of CA1024 chip produced by Connex Technology Inc¹. The Connex System contains mainly an array of 1024 PEs working as a **data parallel sub-system**, DPS, a stream accelerator machine containing 8 PEs (the **time parallel sub-system**, TPS). DPS is driven by an instruction *sequencer*, S, used to broadcast in each clock cycle the same instruction toward each PE from DPS. An *input output controller*, IOC, feeds DPS with data and sends out the results from it. An interconnection fabric allows DPS and TPS to communicate with each other and with the other components of the chip. S and IOC interact using interrupts. They are both simple stack machines with their own data and program memory.

The Connex System uses also other components on the chip to be interfaced with the external world. They are: a MIPS processor acting as a local host, PCI interface to the external host, and a DDR interface to the external memory.

TPS receives streams of data under the control of the local host, and sends the results into the external memory. DPS receives the data vectors from the external memory and sends back the results in the same place. Thus, the two parallel machines communicate usually through the content of the external memory. A data stream is converted into a vector of data, and vice versa, by the programs, run by *Host* and IOC, used to control the buffers organized in the external memory.

3.2 General performances

The first embodiment of the Connex Architecture is designed for 130nm standard process technology, and has the following general performances:

- clock frequency: $f_{CK} = 200 MHz$
- area for the Connex System: $\sim 70 \ mm^2$
- 200 GOPS (OP is a 16-bit simple operation; no multiplication, division or floating point)
- $> 60 \ GOPS/Watt$
- $> 2 GOPS/mm^2$
- internal bandwidth: $400 \ GB/sec$
- external bandwidth: 3.2 GB/sec, involving an additional 2 Watt

3.3 Specific performances

The first application domain investigated in the Connex environment is of **High Definition TV** (HDTV). We estimated 80% of the computational power of the Connex System is necessary to decode in real time two H.264 HDTV streams. Some figures referring to specific functions in HDTV domains follow:

¹ From the moment the title of this paper was announced the name of the company changed in **BrightScale Inc.**.



Fig. 3. The Connex System.

- 8 × 8 DCT: 4.2 clock_cycle (0.066 clock_cycle/pixel)
- 8×8 IDCT: 4.9 *clock_cycle* (0.077 *clock_cycle/pixel*)
- 4 × 4 SAD: 0.04 clock_cycle (0.0025 clock_cycle/pixel)

Graphics is another application domain. A preliminary investigation for an image having the complexity characterized by:

- dynamic images having 10,000 triangles, each covering an average of 100 pixels, one-half being obscured by other triangles
- ambient and diffuze illumination model
- 1920 x 1080 display screen, at 30 frames per second

provides the following figures:

- uses 6.6 GOPS = 3.3% of the total computational power of the Connex System
- and $390 \ MB/sec = 12.2\%$ of the total external bandwidth of the CA1024 chip.

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For **linear algebra** domain we present here only the computation of the dot product for vectors of up to 1024 components. Two cases are estimated:

- for vectors having as components 32-bit floats: 150 clock_cycle (> 1.3 MDot_Product/sec)
- for vectors having as components 16-bit signed integer: 28 clock_cycle (~ 7 MDot_Product/sec)

The **neural network** domain is also targeted as an application domain. A preliminary estimation is done in [Andonie '07]: 5 Giga Connection Updates per Second (about 17 times faster than the fastest *specialized* chip on the market: *Hitachi WSI*).

All these estimations are very encouraging for those who are looking for using the Connex environment as an accelerator for membrane computation.

4 An IPA: The Connex Architecture

The IPA of the Connex System is described in the following two subsections. The vector section describes the architecture of the data parallel sub-system, and the stream section is devoted to describe the time parallel sub-system.

4.1 Vector section

The main physical resources of the Connex System are represented in Figure 4 and are described also in the following pseudo-Verilog form:

```
// Scalar vectors & the index veector
   reg [15:0] svec_000[0:1023],
               svec_001[0:1023],
                . . .
               svec_255[0:1023],
               ixVect[0:1023]
   initial
       ixVect = {0, 1, 2, ... 1023}; // 16-bit scalars
// Boolean vectors
               bvec_0[0:1023]
   reg
               bvec_1[0:1023]
                . . .
               bvec_7[0:1023]
               selVect[0:1023] ; // it is used only as variable
// Scalars
   reg [31:0]
               scalar[0:1023]
                               ;
// Flag vectors
               cryFlag[0:1023] ,
   wire
               zeroFlag[0:1023],
```



Fig. 4. The vector variables of the data parallel subsystem. If the execution is conditioned by AND(bvector0, bvector1), then only column1, ... column1022 of scalars can be involved in computation.

```
eqFlag[0:1023] ,
gtFlag[0:1023] ,
... ;
```

The Boolean vectors are used to select the active components of the scalar vectors. The where construct is a sort of "spatial if".

```
// 'where' construct
where BooleanOP(booleanVect_i, booleanVect_j, ...) {
    svect_k = ScalarOP(svect_p, svect_q, ...),
    bvect_r = xxxFlag;
elsew {
    ...
    }
    }
```

Here is an example of how this construct can be used:

```
where AND(bvect_2, OR(bvect_0, bvect_5)) {
    svect_034 = ADD(svect_012, svect_078, svect_002),
    bvect_3 = cryVect;
    }
elsew {
    svect_034 = ADD(svect_022, svect_222),
```

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bvect_3 = cryVect;
}

It is executed by the Connex System as follows:

```
selVect = OR(bvect_0, bvect_5);
selVect = AND(bvect_2, selVect);
for(i=0; i<1024; i=i+1)
if (selVect[i]) {
    svect_034[i] = ADD(svect_012[i], svect_078[i]),
        bvect_3[i] = cryVect[i];
        svect_034[i] = ADD(svect_034[i], svect_002[i]),
        bvect_3[i] = OR(cryVect[i], bvect_3[i]);
    }
else {svect_034[i] = ADD(svect_022[i], svect_222[i]),
        bvect_3[i] = cryVect[i];
    }
```

There are two distinct ways to generate selections. One is pattern based. It starts from the index vector. Here is an example:

```
/* Pattern based selection example (each other of 4 bit in selVect
will be set on 1)*/
   svect_000 = ixVect;
   svect_000 = AND(svect_000, 16'b11);
   svect_000 = XOR(svect_000, 16'b11), selVect = zeroFlag;
```

The second way to make a selection is to start from the data contained in the scalar vector.

```
// Patternless (data dependent) selection example:
    svect_070 = SUB(svect_070, 16'b10011001), selVect = gtFlag;
```

Usually, any operation specified by one line, having the form:

```
svect_xyz = ScalarOp(...), bvect_q = BooleanOP(...);
```

is executeable in one clock cycle. (Exceptions are specified. For example MULT(...) is executed in 9 clock cycles for 16-bit signed integers, and in 10 clock cycles for unsigned integers.)

4.2 Stream section

The stream section of the Connex System receives the input stream $\langle \mathbf{X} \rangle$ and sends back the output stream $\langle \mathbf{Z} \rangle$, where:

<X> = <x_0, ... x_(p-1)> <Z> = <z_0, ... z_(q-1)> with p = q or $p \neq q$.

The function of the two-dimension pipe $(n \times w)$ is specified by the function vector F, as follows:

F = [func_0, ... func_7]; func_i(y_(i-1), y_(i-2), ... y_(i-w)) = y_i;

where: func_i is the program executed by PE_i . It could be a one instruction looping program, if the pipe "advances" in each clock cycle, or a *s*-instruction loop for pipe propagation executed at each *s* clock cycles. Each PE can have the associated program using variables generated by the previous *w* PEs. The degree of speculation is *w*.

Let be, as an example, the following partially defined computation:

```
 \begin{array}{l} \dots \\ x = \dots \\ y = y[15] ? y + (x + c1) : y + (x + c2); \\ \dots \end{array}
```

Where c1 and c2 are constants. The associated function vector is:

F = [... func_i(...),
 func_(i+1)(y_i),
 func_(i+2)(y_i),
 func_(i+3)(y_(i-1), y_(i-2)), ...];

where:

The output of the processing element PE_i works as input for both, PE_{i+1} and PE_{i+2} . The processing element PE_{i+3} receives the input variables from the previous two machines PE_{i+1} and PE_{i+2} . The second constant dimension of the pipe allows these "shortcuts" which accelerate the computation.

4.3 Putting together the vector section and the stream section

The two sections of the IPA interact through the content of the external memory. In the external memory a vector or a stream have the same representation. Thus, depending on the source or on the destination, an array of data can be interpreted as a vector or as a stream.

Data exchange between the vector section (DPS) and the stream section (TPS) is done by executing one of the two operation:

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X <= <Y>; // stream to vector transfer <X> <= Y; // vector to stream transfer

where:

 $X = \{x_0, \dots x_{(n-1)}\};$ $\langle X \rangle = \langle x_0, \dots x_{(n-1)} \rangle;$ $Y = \{y_0, \dots y_{(n-1)}\};$ $\langle Y \rangle = \langle y_0, \dots y_{(n-1)} \rangle;$

because the destination and the source must have the same dimension n.

5 How to Use Connex to Accelerate Membrane Computing

The key is the representation. The big amount of parallel resources of the Connex architecture can be activated only if an appropriate representation of membrane is adopted. Follow some simple suggestions. The functionality used in these proposals are described in Appendix A.

The first suggestion:

Using the formal definition from [Păun '0x] (see pag. 11), the content of a membrane system can have associated an *n*-component vector which contains an *m*component list (with $n \ge m$). For example (see Fig. 3 in [Păun '0x]):

[[[<w_3>]<w_2>]<w_1>]... = [[[a f c]]]...

where each symbol is represented by a 2-byte word, (index, ASCII_code), as follows:

```
(1,[) (2,[) (3,[) (0,a) (0,f) (0,c) (3,]) (2,]) (1,]) \dots
```

For this first suggestion, only the square parenthesis are indexed, and all the objects are represented with the index having the value 0.

The sets of rules $(R_1, R_2, ...)$ are represented inside the program run by the sequencer S. Thus, the list representing the membrane system will evolve as follows:

(00) [[[a f c]]]... =>
(01) [[[a b f f c]]]... => // in 11 clock cycles
(02) [[[a b b f f f f c]]]... => // in 15 clock cycles
(03) [[b b b f f f f f f f f c]]... => // in 27 clock cycles
(04) [[d d d f f f f c]]... => // in 10 clock cycles
(05) [[d e d e d e f f c]]... => // in 10 clock cycles
(06) [d e d e d e d f c]... => // in 10 clock cycles
(07) [d d d d f c] e e e... => // in 15 clock cycles

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The degree of parallelism is not big enough in each cycle during the previously described computation. Only in step (04) all the three *b*s were substituted by *d*s in parallel (in 2 clock cycles). The parallelism is also involved in searching different symbols such as [,], a, f. On the other hand, all the insertions asked by the evolution rules are performed sequentially.

The performance of the implementation can be increased only by changing the representation of the membrane system.

The second suggestion:

Another way to represent the membrane system is to use indexes also for objects. The most significant byte of each vector component is used to tell us how many objects of the kind indicated by the other byte are represented. The same membrane system have now the following content:

(1,[) (2,[) (3,[) (1,a) (1,f) (1,c) (3,]) (2,]) (1,]) ...

For the same rules applied results the following evolution of the system:

[[[1a 1f 1c]]] =>	
[[[1a 1b 2f 1c]]] =>	<pre>// in 5 clock cycles</pre>
[[[1a 2b 4f 1c]]] =>	<pre>// in 5 clock cycles</pre>
[[3b 8f 1c]] =>	<pre>// in 10 clock cycles</pre>
[[3d 4f 1c]] =>	<pre>// in 7 clock cycles</pre>
[[3d 3e 2f 1c]] =>	<pre>// in 8 clock cycles</pre>
[4d 3e 1f 1c] =>	<pre>// in 8 clock cycles</pre>
[4d 1f 1c] 3e =>	<pre>// in 5 clock cycles</pre>

Now applying the rule $f \to ff$ is executed by simply doubling the index associated to **f**. The same for the rule $d \to de$. For the rule $ff \to f$ the index is divided. The main effects are: the representation is kept smaller and the execution time is reduced more than two times.

The degree of parallelism remains small because the application supposes to work only in one membrane at a time. It will be improved if many membranes having the same rules are processed in the same time.

The third suggestion:

The degree of parallelism increases if on the lowest level more similar membranes are defined. Let us make a little more complex the example presented in [Păun 0x] (see Fig. 3). Suppose on the lowest level there are two membranes ([1a 1f 1c] and [2a 1f 1c]) with the initial content a little different, but working governed by the same rules. Results the following evolution:

```
[[[1a 1f 1c] [2a 1f 1c] ] ]... =>
[[[1a 1b 2f 1c] [2a 2b 2f 1c] ] ]... => // in 5 clock cycles
[[[1a 2b 4f 1c] [2a 4b 4f 1c] ] ]... => // in 5 clock cycles
```

[[3b 8f 1c 6b 8f 1c]] =>	<pre>// in 10 clock cycles</pre>
[[3d 4f 1c 6d 4f 1c]] =>	<pre>// in 7 clock cycles</pre>
[[3d 3e 2f 1c 6d 6e 2f 1c]] =>	<pre>// in 8 clock cycles</pre>
[4d 3e 1f 1c 7d 6e 1f 1c] =>	<pre>// in 8 clock cycles</pre>
[4d 1f 1c 7d 1f 1c] 9e =>	// in 10 clock cycles

The execution time has very little increased (only in the last step). It is obvious that having 3 or more low level membranes the degree of parallelism will increase correspondingly.

The performance can be increased more if the rules are integrated in or as a vector representation. In the previous examples the rules were applied sequentially because they were "known" only by the program issued by the sequencer S. The sequencer must know only to apply rules defined inside the Connex Array in an appropriate manner.

6 Concluding Remarks

Functional vs. Flynn's taxonomy

Our functional taxonomy works better in many-processor environment, while Flynn's taxonomy fits better the multi-processor environment. The functional taxonomy supposes three different types equally involved in defining a high performance architecture, while Flynn's taxonomy proposes also three kinds of parallel machines, only one of them being (MIMD) considered as an effective efficient solution for real machines (see [Hennessy '07]).

Limited non-determinism

The physical resources added for the speculative mechanism are used to support a sort of limited non-deterministic computation inside an IPA.

Can we accelerate molecular computing in vector environment?

The vector section of an IPA can be used to accelerate molecular computation if appropriate representations are imagined. Molecular computing has a huge potential for data parallelism and vector processing is a special kind of data parallel computation. The main problem is to reformulate the molecular approach to fit with the restrictions and promises imposed/offered by the vector computation. The Connex System has also some additional features helping the implementation of specific search functions, very helpful for rewriting rule based processing. Various insert and delete capabilities can be used for the same purpose.

An efficient P-Architecture is slightly different from the current Connex Architecture

Although the Connex environment is helpful for investigating molecular computing based applications, there are needed few specific features in order to obtain a market efficient environment.

Why not a P-language?

A very useful intermediary step toward the definition of a marketable environment for this new computation model is providing a P-language. Working with the basic definition of P-systems is not enough flexible for solving real and complex problems. Using a high level type language and developing for it a specific environment will speed-up the work for a specific membrane platform.

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Skin Output in P Systems with Minimal Symport/Antiport and Two Membranes^{*}

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Summary. It is known that symport/antiport P systems with two membranes and minimal cooperation can generate any recursively enumerable sets of natural numbers using exactly one superfluous object in the output membrane, where the output membrane is an elementary membrane. In this paper we consider symport/antiport P systems where the output membrane is the skin membrane. In this case we prove an unexpected characterization: symport/antiport P systems with two membranes and minimal cooperation generate exactly the recursively enumerable sets of natural numbers. The question about power of purely symport P systems with two membranes and minimal cooperation where the output membrane is the skin membrane is still open.

1 Introduction

P systems with symport/antiport rules, i.e., P systems with pure communication rules assigned to membranes, first were introduced in [21]; symport rules move objects across a membrane together in one direction, whereas antiport rules move objects across a membrane in opposite directions. These operations are very powerful, i.e., P systems with symport/antiport rules have universal computational power with only one membrane, e.g., see [12], [15], [13].

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A comprehensive overview of the most important results obtained in the area of P systems and tissue P systems with *symport/antiport* rules, with respect to the development of computational completeness results improving descriptional complexity parameters as the number of membranes and cells, respectively, the weight of the rules and the number of objects can be found in [1].

For instance, in [3] one obtains the exact characterization of $\mathbb{N}RE$ for symport/antiport P systems with three membranes and minimal cooperation and for corresponding purely symport P systems.

In [5] one shows that if some P system with two membranes and with minimal cooperation, i.e., a P system with symport/antiport rules of weight one or a P system with symport rules of weight two, generates a set of numbers *containing zero*, then this set is **finite**. After that one proves that P systems with symport/antiport rules of weight one can generate any *recursively enumerable* set of natural numbers without zero (i.e., they are computationally complete with just **one superfluous object** remaining in the output membrane at the end of a halting computation). The same result is true also for purely symport P systems of weight two. Therefore, one superfluous object is both necessary and sufficient in case of two membranes.

The question about precise characterization of computational power of symport/antiport P systems (purely symport P systems) with two membranes and minimal cooperation is still open.

Interpreting the result of the computation as the sequence of terminal symbols sent to the environment, one shows that P systems with two membranes and symport rules of weight two or symport/antiport rules of weight one generate all recursively enumerable languages [6].

In this paper we show that P systems with minimal symport/antiport with two membranes characterize $\mathbb{N}RE$ when we consider the **output in the skin membrane** rather than the elementary membrane.

2 Basic Notations and Definitions

For the basic elements of formal language theory needed in the following, we refer to [26]. We just list a few notions and notations: \mathbb{N} denotes the set of natural numbers (i.e., of non-negative integers). V^* is the free monoid generated by the alphabet V under the operation of concatenation and the empty string, denoted by λ , as unit element; by $\mathbb{N}RE$, $\mathbb{N}REG$, and $\mathbb{N}FIN$ we denote the family of recursively enumerable sets, regular sets, and finite sets of natural numbers, respectively. For $k \geq 1$, by $\mathbb{N}_k RE$ we denote the family of recursively enumerable sets of natural numbers excluding the initial segment 0 to k - 1. Particularly, $\mathbb{N}_1 RE = \{N \in \mathbb{N}RE \mid 0 \notin N\}$. The families of recursively enumerable sets of vectors of natural numbers are denoted by PsRE.

2.1 Counter Automata

A non-deterministic *counter automaton* (see [11], [1]) is a construct

$$M = (d, Q, q_0, q_f, P)$$
, where

- d is the number of counters, and we denote $D = \{1, ..., d\};$
- Q is a finite set of states, and without loss of generality, we use the notation $Q = \{q_i \mid 0 \le i \le f\}$ and $F = \{0, 1, ..., f\},$
- $q_0 \in Q$ is the initial state,
- $q_f \in Q$ is the final state, and
- *P* is a finite set of instructions of the following form:
- 1. $(q_i \rightarrow q_l, k+)$, with $i, l \in F$, $i \neq f$, $k \in D$ ("increment" -instruction). This instruction increments counter k by one and changes the state of the system from q_i to q_l .
- 2. $(q_i \rightarrow q_l, k-)$, with $i, l \in F$, $i \neq f$, $k \in D$ ("decrement" -instruction). If the value of counter k is greater than zero, then this instruction decrements it by 1 and changes the state of the system from q_i to q_l . Otherwise (when the value of counter k is zero) the computation is blocked in state q_i .
- 3. $(q_i \to q_l, k = 0)$, with $i, l \in F$, $i \neq f$, $k \in D$ ("test for zero" -instruction). If the value of counter k is zero, then this instruction changes the state of the system from q_i to q_l . Otherwise (the value stored in counter k is greater than zero) the computation is blocked in state q_i .
- 4. halt. This instruction stops the computation of the counter automaton, and it can only be assigned to the final state q_f .

A transition of the counter automaton consists in updating/checking the value of a counter according to an instruction of one of the types described above and by changing the current state to another one. The computation starts in state q_0 with all counters being equal to zero. The result of the computation of a counter automaton is the value of the first k counters when the automaton halts in state $q_f \in Q$ (without loss of generality we may assume that in this case all other counters are empty). A counter automaton thus (by means of all computations) generates a set of k-vectors of natural numbers. If k = 1, then by N(M) we denote the corresponding numeric set generated by M.

2.2 P Systems with Symport/Antiport Rules

The reader is supposed to be familiar with basic elements of membrane computing, e.g., from [23]; comprehensive information can be found in the P systems web page, [30].

A P system with symport/antiport rules is a construct

$$\Pi = (O, \mu, w_1, \dots, w_k, E, R_1, \dots, R_k, i_0),$$
 where

- 1. *O* is a finite alphabet of symbols called *objects*;
- 2. μ is a *membrane structure* consisting of k membranes that are labelled in a one-to-one manner by $1, 2, \ldots, k$;

- 3. $w_i \in O^*$, for each $1 \le i \le k$, is a finite multiset of objects associated with the region *i* (delimited by membrane *i*);
- 4. $E \subseteq O$ is the set of objects that appear in the environment in an infinite number of copies;
- 5. R_i , for each $1 \le i \le k$, is a finite set of symport/antiport rules associated with membrane *i*; these rules are of the forms (x, in) and (y, out) (symport rules) and (y, out; x, in) (antiport rules), respectively, where $x, y \in O^+$;
- 6. i_0 is the label of a membrane of μ that identifies the corresponding output region.

A P system with symport/antiport rules is defined as a computational device consisting of a set of k hierarchically nested membranes that identify k distinct regions (the membrane structure μ), where to each membrane i there are assigned a multiset of objects w_i and a finite set of symport/antiport rules R_i , $1 \le i \le k$. A rule $(x, in) \in R_i$ permits the objects specified by x to be moved into region i from the immediately outer region. Notice that for P systems with symport rules the rules in the skin membrane of the form (x, in), where $x \in E^*$, are forbidden. A rule $(x, out) \in R_i$ permits the multiset x to be moved from region i into the outer region. A rule (y, out; x, in) permits the multisets y and x, which are situated in region i and the outer region of i, respectively, to be exchanged. It is clear that a rule can be applied if and only if the multisets involved by this rule are present in the corresponding regions. The weight of a symport rule (x, in) or (x, out) is given by |x|, while the weight of an antiport rule (y, out; x, in) is given by $max\{|x|, |y|\}$.

As usual, a computation in a P system with symport/antiport rules is obtained by applying the rules in a non-deterministic maximally parallel manner. Specifically, in this variant, a computation is restricted to moving objects through membranes, since symport/antiport rules do not allow the system to modify the objects placed inside the regions. Initially, each region i contains the corresponding finite multiset w_i , whereas the environment contains only objects from E that appear in infinitely many copies.

A computation is successful if starting from the initial configuration, the P system reaches a configuration where no rule can be applied anymore. The result of a successful computation is a natural number that is obtained by counting all objects present in region i_0 . Given a P system Π , the set of natural numbers computed in this way by Π is denoted by $N(\Pi)$. If the multiplicity of each object is counted separately, then a vector of natural numbers is obtained, denoted by $Ps(\Pi)$, see [23].

By $\mathbb{N}OP_m(sym_s, anti_t)$ we denote the family of sets of natural numbers generated by P systems with symport/antiport rules with at most m > 0 membranes, symport rules of size at most $s \ge 0$, and antiport rules of size at most $t \ge 0$. In the papers on P systems, following [23], i_0 is assumed to be an elementary membrane. In this paper we will write $\mathbb{N}^{skin}OP_m(sym_s, anti_t)$ if i_0 is the skin membrane. Any unbounded parameter m, s, t is replaced by *. If t = 0, then we may omit $anti_t$.

3 Main result

Theorem 1. $\mathbb{N}^{skin}OP_2(sym_1, anti_1) = \mathbb{N}RE.$

Proof. We simulate a counter automaton $M = (d, Q, q_0, q_f, P)$. Recall that M starts with empty counters. We also suppose that all instructions from P are labeled in a one-to-one manner with elements of $\{1, \ldots, n\} = I$, n is a label of the *halt* instruction and $I' = I \setminus \{n\}$. We denote by I_+, I_- , and $I_{=0}$ the set of labels for the "increment" -, "decrement" -, and "test for zero" -instructions, respectively. We also use the following notation: $C = \{c_k\}, k \in D$ and $Q' = Q \setminus \{q_0\}$.

We construct the P system Π_1 as follows:

$$\begin{split} \Pi_1 &= (O, \begin{bmatrix} 1 & 2 & 1 \\ 2 & 1 \end{bmatrix}_1, w_1, w_2, E, R_1, R_2, 1), \\ O &= E \cup \{L, T_1, T_2, P_2, J_1, J_2, J_3\} \cup \{b_j \mid j \in I\} \cup \{d_j \mid j \in I'\}, \\ E &= Q' \cup C \cup \{a_j \mid j \in I\} \cup \{a'_j, e_j \mid j \in I'\} \cup \{J_0, P_1\} \cup \{F_i \mid 0 \le i \le 9\}, \\ w_1 &= q_0 L J_1 J_2 J_3, \\ w_2 &= T_1 T_2 P_2 \prod_{j \in I} b_j \prod_{j \in I'} d_j, \\ R_i &= R_{i,s} \cup R_{i,r} \cup R_{i,f}, \quad i = 1, 2. \end{split}$$

We code the counter automaton as follows:

Region 1 will hold the current state of the automaton, represented by a symbol $q_i \in Q$ and also the value of all counters, represented by the number of occurrences of symbols $c_k \in C$, $k \in D$, where $D = \{1, ..., d\}$.

We split our proof into several parts that depend on the logical separation of the behavior of the system. We will present the rules and the initial symbols for each part, but we remark that the system we present is the union of all these parts. The rules R_i are given by three phases:

- 1. START: preparation of the system for the computation.
- 2. RUN: simulation of instructions of the counter automaton.
- 3. END: terminating the computation.

The parts of the computations illustrated in the following describe different phases of the evolution of the P system. For simplicity, we focus on explaining a particular phase and omit the objects that do not participate in the evolution at that time. Each rectangle represents a membrane, each variable represents a copy of an object in a corresponding membrane (symbols outside of the outermost rectangle are found in the environment). In each step, the symbols that will evolve (will be moved) are written in **boldface**. The labels of the applied rules are written above the symbol \Rightarrow .

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1. START.

We use the following idea: in our system we have a symbol L which moves from region 1 to the environment and back in an infinite loop. This loop may be stopped only if all stages are completed correctly.

$$\begin{split} R_{1,s} &= \{\texttt{1s1}: (L, out), \texttt{1s2}: (L, in)\}.\\ R_{2,s} &= \emptyset. \end{split}$$

Notice that some rules are never executed during a correct simulation: applying them would lead to an infinite computation. To help the reader, we will <u>underline</u> the labels of such rules in the description below.

2. RUN.

$$\begin{split} R_{1,r} &= \{ \mathbf{1r1} : (q_i, out; a_j, in) \mid (j : q_i \to q_l, c_k \gamma) \in P, \gamma \in \{+, -, = 0\} \} \\ &\cup \{ \mathbf{1r2} : (q_f, out; a_n, in) \} \\ &\cup \{ \mathbf{1r3} : (b_j, out; a'_j, in) \mid j \in I' \} \\ &\cup \{ \mathbf{1r4} : (a_j, out; J_0, in), \ \mathbf{1r5} : (J_1, out; b_j, in) \mid j \in I \} \\ &\cup \{ \mathbf{1r4} : (a_j, out; J_1, in) \} \\ &\cup \{ \mathbf{1r6} : (J_0, out; J_1, in) \} \\ &\cup \{ \mathbf{1r7} : (a'_j, out; c_k, in) \mid (j : q_i \to q_l, c_k +) \in P \} \\ &\cup \{ \mathbf{1r8} : (a'_j, out) \mid j \in I_+ \cup I_{=0} \} \\ &\cup \{ \mathbf{1r9} : (d_j, in) \mid j \in I_+ \cup I_{=0} \} \\ &\cup \{ \mathbf{1r10} : (c_k, out; d_j, in) \mid (j : q_i \to q_l, c_k -) \in P \} \\ &\cup \{ \mathbf{1r11} : (J_3, out; d_j, in) \mid \in I_- \} \\ &\cup \{ \mathbf{1r12} : (J_3, out; d_j, in) \mid j \in I' \} \\ &\cup \{ \mathbf{1r13} : (d_j, out; e_j, in) \mid j \in I' \} \\ &\cup \{ \mathbf{1r14} : (e_j, out, q_l, in) \mid (j : q_i \to q_l, c_k \gamma) \in P, \gamma \in \{+, -, = 0\} \} \\ &\cup \{ \mathbf{1r15} : (b_n, out; F_0, in) \} \\ &\cup \{ \mathbf{1r16} : (\#, out), \mathbf{1r17} : (\#, in) \}. \end{split}$$

$$\begin{aligned} R_{2,r} &= \{ 2\mathbf{r}\mathbf{1} : (b_j, out; a_j, in), 2\mathbf{r}\mathbf{2} : (a_j, out; J_2, in) \mid j \in I \} \\ &\cup \{ \underline{2\mathbf{r}\mathbf{3}} : (a_j, out; J_1, in) \mid j \in I \} \\ &\cup \{ 2\mathbf{r}\mathbf{4} : (d_j, out; a'_j, in) \mid j \in I' \} \\ &\cup \{ \underline{2\mathbf{r}\mathbf{5}} : (a'_j, out; c_k, in) \mid (j : q_i \to q_l, c_k = 0) \in P \} \\ &\cup \{ 2\mathbf{r}\mathbf{6} : (a'_j, out; e_j, in) \mid j \in I_{=0} \} \\ &\cup \{ \underline{2\mathbf{r}\mathbf{7}} : (a'_j, out; J_1, in) \mid j \in I_{=0} \} \\ &\cup \{ 2\mathbf{r}\mathbf{8} : (e_j, out; d_j, in) \mid j \in I_{=0} \} \end{aligned}$$

 $\cup \{\underline{2r9}: (e_j, out; J_1, in) \mid j \in I_{=0} \} \\ \cup \{2r10: (d_j, in) \mid j \in I_+ \cup I_- \} \\ \cup \{2r11: (a'_j, out) \mid j \in I_+ \cup I_- \} \\ \cup \{2r12: (J_2, out; b_j, in) \mid j \in I' \} \\ \cup \{2r13: (J_2, out; J_1, in), 2r14: (\#, out; J_0, in) \}.$

First of all, we mention that if during the phase RUN object J_3 comes to the environment by rules <u>1r11</u>, <u>1r12</u> (Scenario 0), it remains there forever and cannot move object L to region 2 (during the phase END), thus to stop the infinite loop. So, the computation never halts.

Let us explain the synchronization of a_j coming to the environment and b_j leaving the environment: the first one brings J_0 into region 1 while the latter brings J_1 into the environment; then rule **1r6** returns J_0 and J_1 to their original locations.

If a_j comes to the environment without b_j leaving it or b_j is in region 1 or 2 at that moment (it is possible after applying rules <u>2r3</u>, <u>2r7</u>, <u>2r13</u>), J_1 remains in region 1 (or 2) and J_0 comes to region 1 and after that in region 2 by rules **1r4**, <u>2r14</u> (Scenario 1), thus causing an endless computation since <u>1r16</u> and <u>1r17</u> are always applicable.

If b_j leaves the environment without a_j coming there, J_0 remains in the environment and J_1 comes there (Scenario 2), so <u>1r12</u> is applied and J_3 comes to the environment. The computation never halts, see scenario 0.

Scenario 3 takes place when two symbols a_j and symbol $b_j, j \in I$ appear in region 1 and in the environment accordingly. In this case rules 1r4, 1r5 will be applied, and rule 1r4 two times. Thus, two symbols J_0 appear in region 1 and rule 2r14 will be applied eventually. The computation never halts, see scenario 1.

We also mention that applying rule <u>1r11</u> causes scenario 0 (this is a case of modeling a "decrement"-instruction, there is no c_k in region 1); applying <u>2r5</u> leads to scenario 3 (this is a case of modeling a "test for zero"-instruction, there is some c_k in region 1), and applying <u>2r7</u> and <u>2r9</u> eventually causing scenario 1. Therefore, in order for a computation to halt, no underlined rules should be applied.

We will now consider the "main" line of computation. We explain the behavior of simulating the instruction $(j : q_i \rightarrow q_l, c_k \gamma)$. Index s stands for any possible instruction associated to state q_l .

"Increment" -instruction:

$$q_{l}\mathbf{a}_{j}a_{s}a'_{j}e_{j}c_{k}J_{0}\left[\mathbf{q}_{i}J_{1}J_{2}J_{3}\overline{b_{j}d_{j}\#}\right] \Rightarrow^{\mathbf{1r1}} q_{l}q_{i}a_{s}a'_{j}e_{j}c_{k}J_{0}\left[\mathbf{a}_{j}J_{1}J_{2}J_{3}\overline{\mathbf{b}_{j}d_{j}\#}\right] \Rightarrow^{\mathbf{2r1}} q_{l}q_{i}a_{s}a'_{j}e_{j}c_{k}J_{0}\left[\mathbf{b}_{j}J_{1}\mathbf{J}_{2}J_{3}\overline{\mathbf{a}_{j}d_{j}\#}\right] \Rightarrow^{\mathbf{1r3},\mathbf{2r2}} q_{l}q_{i}a_{s}\mathbf{b}_{j}e_{j}c_{k}\mathbf{J}_{0}\left[\mathbf{a}'_{j}\mathbf{J}_{1}\mathbf{a}_{j}J_{3}\overline{J_{2}\mathbf{d}_{j}\#}\right] \Rightarrow^{\mathbf{1r4},\mathbf{1r5},\mathbf{2r4}} q_{l}q_{i}a_{j}a_{s}\mathbf{e}_{j}c_{k}\mathbf{J}_{1}\left[\mathbf{b}_{j}\mathbf{d}_{j}\mathbf{J}_{0}J_{3}\overline{\mathbf{J}_{2}\mathbf{a}'_{j}\#}\right]$$
(A)

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$$\Rightarrow^{\mathbf{1r6},\mathbf{1r13},\mathbf{2r11},\mathbf{2r12}} \mathbf{q}_{\mathbf{l}}q_{i}a_{j}a_{\mathbf{s}}\mathbf{d}_{\mathbf{j}}\mathbf{c}_{\mathbf{k}}J_{0} \begin{bmatrix} J_{1}J_{2}\mathbf{a}_{\mathbf{j}}'\mathbf{e}_{\mathbf{j}}J_{3} \begin{bmatrix} b_{j} \# \end{bmatrix} \Rightarrow^{\mathbf{1r7},\mathbf{1r9},\mathbf{1r14}} \\ q_{i}a_{j}\mathbf{a}_{\mathbf{s}}a_{j}'e_{j}J_{0} \begin{bmatrix} \mathbf{q}_{\mathbf{l}}\mathbf{d}_{\mathbf{j}}J_{1}J_{2}J_{3}c_{k} \begin{bmatrix} b_{j} \# \end{bmatrix} \end{bmatrix} \Rightarrow^{\mathbf{1r1},\mathbf{2r10}} q_{l}q_{i}a_{j}a_{j}'e_{j}J_{0} \begin{bmatrix} \mathbf{a}_{\mathbf{s}}J_{1}J_{2}J_{3}c_{k} \begin{bmatrix} b_{j} \# \end{bmatrix} \end{bmatrix}$$

In that way, q_i is replaced by q_l and c_k is moved from the environment into region 1. Notice that symbols a_j , b_j , a'_j , d_j , e_j , J_0 , J_1 , J_2 have returned to their original positions. Symbol d_j returns to region 2 in the first step of the simulation of the next instruction (the last step of the illustration).

"Decrement" -instruction:

(i) There is some c_k in region 1:

We consider configuration (A) above with symbol c_k in region 1.

$$\begin{aligned} q_{l}q_{i}a_{j}a_{s}\mathbf{e_{j}J_{1}}\mathbf{b_{j}d_{j}J_{0}}J_{3}c_{k}\mathbf{J_{2}a_{j}'\#} & \Rightarrow^{\mathrm{1r6},\mathrm{1r13},2\mathrm{r11},2\mathrm{r12}} \\ \mathbf{q}_{l}q_{i}a_{j}a_{s}\mathbf{d}_{j}J_{0}\mathbf{J}_{1}J_{2}\mathbf{a_{j}'e_{j}J_{3}c_{k}}\mathbf{b}_{j}\#} & \Rightarrow^{\mathrm{1r8},\mathrm{1r10},\mathrm{1r14}}q_{i}a_{j}a_{s}a_{j}'e_{j}c_{k}J_{0}\mathbf{q}_{l}J_{1}J_{2}J_{3}\mathbf{d}_{j}\mathbf{b}_{j}\#} \\ & \Rightarrow^{\mathrm{1r1},2\mathrm{r10}}q_{l}q_{i}a_{j}a_{j}'e_{j}c_{k}J_{0}\mathbf{a}_{s}J_{1}J_{2}J_{3}\mathbf{b}_{j}d_{j}\#} \end{aligned}$$

In the way described above, q_i is replaced by q_l and c_k is removed from region 1 to the environment. Notice that symbols a_j , a'_j , b_j , d_j , e_j , J_0 , J_1 , J_2 have returned to their original positions. Symbol d_j returns to region 2 in the first step of the simulation of the next instruction (the last step of the illustration).

(ii) There is no c_k in region 1:

Again we start with configuration (A).

$$q_l q_i a_j a_s \mathbf{e_j J_1} \mathbf{b_j d_j J_0} J_3 \mathbf{J_2 a'_j \#}$$

$$\Rightarrow^{\mathbf{1r6},\mathbf{1r13},\mathbf{2r11},\mathbf{2r12}} \mathbf{q_l} q_i a_j a_s \mathbf{d_j} J_0 \overline{J_1 J_2 \mathbf{a'_j e_j J_3} b_j \#} \Rightarrow^{\mathbf{1r8},\mathbf{1r11},\mathbf{1r14}}$$

Now rule <u>1r11</u> will be applied, leading to an infinite computation (see scenario 0).

"Test for zero" -instruction:

 q_i is replaced by q_l if there is no c_k in region 1, otherwise a'_j in region 2 exchanges with c_k in region 1 and the computation will never stop.

(i) There is no c_k in region 1:

We consider configuration (A) above.

$$\begin{aligned} q_l q_i a_j a_s \mathbf{e_j J_1} \mathbf{b_j d_j J_0} J_3 \overline{\mathbf{J_2}a'_j \#} & \Rightarrow^{\mathrm{ir6},\mathrm{ir13},2\mathrm{r12}} q_l q_i a_j a_s \mathbf{d_j} J_0 \overline{J_1 J_2 \mathbf{e_j} J_3 \mathbf{a'_j b_j \#}} \\ \Rightarrow^{\mathrm{ir9},2\mathrm{r6}} q_l q_i a_j a_s J_0 \mathbf{d_j} J_1 J_2 J_3 \mathbf{a'_j e_j b_j \#} & \Rightarrow^{\mathrm{ir8},2\mathrm{r8}} \mathbf{q_l} q_i a_j a_s a'_j J_0 \mathbf{e_j} J_1 J_2 J_3 \overline{b_j d_j \#} \\ \Rightarrow^{\mathrm{ir14}} q_i a_j \mathbf{a_s} a'_j e_j J_0 \mathbf{q_l} J_1 J_2 J_3 \overline{b_j d_j \#} \end{aligned}$$

In this case, q_i is replaced by q_l . Notice that symbols a_j , a'_j , b_j , d_j , e_j , J_0 , J_1 , J_2 have returned to their original positions.

(ii) There is some c_k in region 1:

Consider configuration (A) with object c_k in region 1:

$$q_l q_i a_j a_s \mathbf{e_j J_1} \underbrace{\mathbf{b_j d_j J_0 J_3 c_k J_2 a'_j \#}} \Rightarrow^{\mathrm{tr6}, \mathrm{tr13}, \underline{2r5}, 2r12}$$

Now applying rule 2r5 leads to an infinite computation.

$$\begin{aligned} \mathbf{q}_{\mathbf{l}}q_{i}a_{j}a_{s}a_{s}a'_{s}\mathbf{d}_{\mathbf{j}}J_{0}J_{0} & J_{1}J_{2}\mathbf{a}'_{\mathbf{j}}\mathbf{e}_{\mathbf{j}}J_{3} & b_{j}b_{s}c_{k}\# \end{aligned} \Rightarrow^{\mathbf{1}\mathbf{r}\mathbf{8},\mathbf{1}\mathbf{r}\mathbf{9},\mathbf{1}\mathbf{r}\mathbf{1}\mathbf{4}} \\ q_{i}a_{j}a_{s}\mathbf{a}_{\mathbf{s}}a'_{j}a'_{\mathbf{s}}\mathbf{e}_{\mathbf{j}}J_{0}J_{0} & \mathbf{q}_{\mathbf{l}}\mathbf{d}_{\mathbf{j}}J_{1}J_{2}J_{3} & b_{j}b_{s}c_{k}\# \end{aligned} \Rightarrow^{\mathbf{1}\mathbf{r}\mathbf{1},\mathbf{1}\mathbf{r}\mathbf{1}\mathbf{3}} \\ \mathbf{q}_{\mathbf{l}}q_{i}a_{j}a_{s}a'_{j}a'_{\mathbf{s}}\mathbf{d}_{\mathbf{j}}J_{0}J_{0} & \mathbf{a}_{\mathbf{s}}\mathbf{e}_{\mathbf{j}}J_{1}J_{2}J_{3} & b_{j}\mathbf{b}_{\mathbf{s}}\# \end{aligned} \Rightarrow^{\mathbf{1}\mathbf{r}\mathbf{1},\mathbf{1}\mathbf{r}\mathbf{1}\mathbf{3}} \\ q_{i}a_{j}\mathbf{a}_{s}a'_{j}a'_{\mathbf{s}}e_{j}J_{0}J_{0} & \mathbf{a}_{\mathbf{s}}\mathbf{e}_{\mathbf{j}}J_{1}J_{2}J_{3} & \mathbf{b}_{j}\mathbf{b}_{\mathbf{s}}\# \end{aligned} \Rightarrow^{\mathbf{1}\mathbf{r}\mathbf{1},\mathbf{1}\mathbf{r}\mathbf{3},\mathbf{2}\mathbf{r}\mathbf{2}} \\ q_{l}q_{i}a_{j}a'_{\mathbf{j}}\mathbf{b}_{\mathbf{s}}e_{j}\mathbf{J}_{0}\mathbf{J}_{0} & \mathbf{a}_{\mathbf{s}}\mathbf{a}_{\mathbf{s}}a'_{\mathbf{s}}\mathbf{J}_{\mathbf{1}}J_{2}J_{3} & \mathbf{d}_{j}\# \end{aligned}$$

So, scenario 3 takes place and the computation never halts.

3. END.

$$\begin{split} R_{1,f} &= \{ \texttt{lf1}: (T_1, out; F_1, in) \} \cup \{ \texttt{lf2}: (F_i, out; F_{i+1}, in) \mid 1 \leq i \leq 8 \} \\ &\cup \{ \texttt{lf3}: (T_2, out; P_1), \ \texttt{lf4}: (P_2, out), \ \texttt{lf5}: (F_0, out; P_2, in) \}. \\ R_{2,f} &= \{ \texttt{2f1}: (T_1, out; F_0, in), \texttt{2f2}: (F_0, out), \texttt{2f3}: (T_2, out; F_0, in) \} \\ &\cup \{ \texttt{2f4}: (P_1, in), \texttt{2f5}: (P_1, out; J_1, in), \texttt{2f6}: (P_1, out; J_2, in) \} \\ &\cup \{ \texttt{2f7}: (P_1, out; J_3, in), \texttt{2f8}: (J_3, out; L, in), \texttt{2f9}: (P_2, out; F_9, in) \}. \end{split}$$

Once the counter automaton reaches the final state, q_f is in region 1 and it exchanges with object a_n (rule 1r2) and object F_0 will be moved to region 1 in several steps (rules 1r15).

It takes T_1 and T_2 to region 1, in either order. The duty of T_2 is to bring P_1 from the environment to region 2, where P_1 pumps objects J_1, J_2, J_3 from region 1 to region 2. If on the previous steps of simulation of counter automaton M object

 J_3 was moved to the environment (by rules <u>1r11</u>, <u>1r12</u>), scenario 0 takes place and the computation never halts, as there is only one possibility to stop an infinite loop with object L, i.e. to move it to region 2 by rule 2f8.

 T_1 starts a chain of exchanges of objects F_i , as a result object F_9 will be moved to region 1 and then object P_2 will be moved to the environment, where it pumps object F_0 to the environment. So, at the end of the computation there are only objects $c_k, k \in D$ in region 1. The entire simulation shows the inclusion $N(\Pi_1) \supseteq N(M)$.

The converse inclusion also holds because the system may only halt if it has correctly simulated a computation of the counter automaton (according to the design of the system) from state q_0 to state q_f , while if behavior of M is not simulated correctly, then the computation never halts and hence does not contribute to $N(\Pi_1)$. This shows that P systems with two membranes and symport/antiport rules of weight one with the output in the skin membrane generate all recursively enumerable sets of natural numbers. Since the power of such systems cannot exceed that of Turing machines, the statement of the theorem is an equality. \Box

4 Conclusions

In this paper we prove the new result that any recursively enumerable set of natural numbers is generated by symport/antiport P systems with two membranes and minimal cooperation where the output membrane is the skin membrane. It contrasts with the previous result where an elementary membrane is used as the output membrane, where at least one superfluous object is necessary in the output membrane in order to get universality. Thus we answered the question of Francesco Bernardini about computational power of symport/antiport P systems with two membranes and minimal cooperation where the output membrane is the skin membrane. The question about power of purely symport P systems with two membranes and minimal cooperation where the output membrane is the skin membrane is still open.

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On the Reachability Problem in P Systems with Mobile Membranes

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Summary. We investigate the problem of reaching a configuration from another configuration in mobile membranes, and prove that the reachability can be decided by reducing it to the reachability problem of a version of pure and public ambient calculus without the capability **open**.

1 Introduction

Membrane systems (called also P systems) are introduced by Gh.Păun in [8, 9] as a class of parallel computing devices inspired by biology. The definition of this computing model starts from the observation that any biological system is a complex hierarchical structure, with a flow of materials and information which underlies their functioning. The membrane computing deals with the evolution of systems composed by objects, rules and membranes nested in other membranes. The P systems with mobile membranes [5] is a model which expresses mobility by the movement of membranes in such a system. The movement is given mainly by two operations: exocytosis and endocytosis.

Ambient calculus is a formalism introduced in [3] to describe concurrent and mobile computation. In contrast with other formalisms for mobile processes such as the π -calculus [7] (whose computational model is based on the notion of **communication**), the ambient calculus is based on the notion of **movement**. An ambient is a named location, and represents a unit of movement. Ambients mobility is controlled by the capabilities in, out, and open; the mobile ambients describe the migration of processes between certain boundaries.

The membrane systems and mobile ambients have similar structures and common concepts. Both have a hierarchical structure, work mainly with a notion of location, and are used to model various aspects on the distributed systems. The distributed features of mobile ambients are described in [3], and distributed algorithms for membrane systems are presented in [4]. In this paper we investigate the problem of reaching a certain configuration in mobile membranes starting from a given configuration. We prove that reachability in mobile membranes can be decided by reducing it to the reachability problem of a version of pure and public ambient calculus from which the **open** capability has been removed. In [1] it is proven that the reachability for this fragment of ambient calculus is decidable by reducing it to marking reachability for Petri nets, which is proven to be decidable in [6].

The structure of the paper is as follows. In Section 2 we present the mobile membrane systems, whereas in Section 3 we present a version of pure and public mobile ambients without the capability **open**. The core of the paper is represented by Section 4, where we investigate the reachability problem for mobile membranes. Conclusions and references end the paper.

2 Mobile Membranes Systems

- **Definition 1.** A mobile membrane system is a construct $\prod = (V \cup \overline{V}, H \cup \overline{H}, \mu, w_1, \dots, w_n, R), \quad where:$
 - 1. $n \ge 1$ (the initial degree of the system);
 - 2. $V \cup \overline{V}$ is an alphabet (its elements are called objects), where $V \cap \overline{V} = \emptyset$;
 - 3. $H \cup \overline{H}$ is a finite set of labels for membranes, where $H \cap \overline{H} = \emptyset$;
 - 4. μ is a membrane structure, consisting of n membranes, labeled (not necessarily in a one-to-one manner) with elements of H;
 - 5. w_1, w_2, \ldots, w_n are multisets of objects placed in the n membranes of the system;
 - 6. R is a finite set of developmental rules, of the following forms:
 - a) ā↓→ ā↓ a↓, for a↓∈ V, ā↓∈ V; replication rule
 The objects ā↓ are used to create new objects a↓ without being consumed.
 b) ā↑→ ā↑ a↑, for a↑∈ V, ā↑∈ V; replication rule

The objects $\overline{a}\uparrow$ are used to create new objects $a\uparrow$ without being consumed.

- c) $[a\downarrow]_h []_a \rightarrow [[]_h]_a$, for $a, h \in H, a \downarrow \in V$; endocytosis An elementary membrane labeled h enters the adjacent membrane labeled a, under the control of object $a\downarrow$. The labels h and a remain unchanged during this process; however the object $a\downarrow$ is consumed during the operation. Membrane a is not necessarily elementary.
- d) [[a↑]_h]_a → []_h []_a, for a, h ∈ H, a↑∈ V; exocytosis
 An elementary membrane labeled h is sent out of a membrane labeled a, under the control of object a↑. The labels of the two membranes remain unchanged; the object a↑ of membrane h is consumed during the operation. Membrane a is not necessarily elementary.
- e) $[]_{\overline{h}} \to []_{\overline{h}}[]_h$ for $h \in H$, $\overline{h} \in \overline{H}$ division rules An elementary membrane labeled \overline{h} is divided into two membranes labeled by \overline{h} and h having the same objects.

In 3, $H \cap \overline{H} = \emptyset$ states that the membranes having labels from the set \overline{H} can participate only in rules of type (e). Similarly, $V \cap \overline{V} = \emptyset$ in 2 states that the objects from \overline{V} can participate only in rules of type (a) and (b)

The rules are applied using the following principles:

- 1. In biological systems molecules are divided into classes of different types. We make the same decision here and split the objects into four classes: $a \downarrow$ objects that control the *endocytosis*, $a \uparrow$ objects that control the *exocytosis*, and $\overline{a} \downarrow$, $\overline{a} \uparrow$ objects that produce new objects from the first two classes without being consumed.
- 2. All the rules of type (c), (d) are applied in parallel, non-deterministically choosing the rules, the membranes, and the objects, but in such a way that the parallelism is maximal; this means that in each step we apply a set of rules such that no further rule of type (c), (d) can be added to the set, no further membranes and objects can evolve at the same time.
- 3. The membrane a from each rules of type (c), (d) is said to be **passive**, while the membrane h is said to be **active**. In any step of a computation, any object and any active membrane can be involved in at most one rule, but the passive membranes are not considered involved in the use of rules (hence they can be used by several rules at the same time as passive membranes).
- 4. When a membrane is moved across another membrane, by endocytosis or exocytosis, its whole content (its objects) are moved.
- 5. If a membrane is divided, then its content is replicated in the two new copies.
- 6. The skin membrane can never be divided.
- 7. Not all the rules of type (a), (b), (e) are applied whenever it is possible; we choose non-deterministically whether the rules of these types are applied.

According to these rules, we get transitions among the configurations of the system. For two mobile membrane systems M and N we say that M reduces to N if there is a sequence of rules applicable in the membrane system M in order to obtain the membrane system N.

3 Mobile Ambients

We describe a variant of pure and public mobile ambients (mobile ambients in which communication and name restriction are omitted); more details can be found in [1]. Given an infinite set of names \mathcal{N} (ranged over by m, n, \ldots), we define the set \mathcal{A} of mobile ambients (denoted by A, A', B, \ldots) together with their capabilities (denoted by C, C', \ldots) as follows:

 $C ::= in n \mid out n$

Capabilities

 $A ::= A \mid B \mid C.A \mid n[A] \mid !A$ Processes

A movement C.A is provided by the capability C, followed by the execution of process A. An entry capability in n instructs the surrounding ambient to enter a

sibling ambient labeled by n, while an exit capability *out* n intructs the surrounding ambient to exit its parent ambient labeled by n. An ambient n[A] represents a bounded place labeled by n in which a process A is executed. $A \mid B$ is a parallel composition of processes A and B. The process !A denotes the unbounded replication of process A.

Processes of this calculus are grouped into equivalence classes, up to trivial syntactic restructuring, by the following structural congruence relation, \equiv , which is the least congruence satisfying the following requirements:

 $\begin{array}{ll} A \mid B \equiv B \mid A & A \equiv B \text{ implies } A \mid A' \equiv B \mid A' \\ (A \mid B) \mid A' \equiv A \mid (B \mid A') & A \equiv B \text{ implies } !A \equiv !B \\ A \equiv A & A \equiv B \text{ implies } B \equiv A & A \equiv B \text{ implies } n[A] \equiv n[B] \\ A \equiv B, B \equiv A' \text{ implies } A \equiv A' & \end{array}$

The operational semantics of the mobile ambients is defined in terms of a reduction relation \Rightarrow by the following axioms and rules:

Axioms:

(In) $n[in m. A | A'] | m[B] \Rightarrow m[n[A | A'] | B];$ (Out) $m[n[out m. A | A'] | B] \Rightarrow n[A | A'] | m[B];$ (Repl) $!A \Rightarrow A | !A$. Rules: (Comp) $\frac{A \Rightarrow A'}{A | B \Rightarrow A' | B}$ (Amb) $\frac{A \Rightarrow A'}{n[A] \Rightarrow n[A']}$ (Struc) $\frac{A \equiv A', A' \Rightarrow B', B' \equiv B}{A \Rightarrow B}$.

The **Axioms** represent the one-step reductions for *in* and *out*, and the unfolding of replication. The **Rules** propagate reduction across ambient nesting, parallel composition and allow the use of equivalence during reduction. The **In** and **Out** rules are applied as soon as possible in a maximal parallel manner. We denote by \Rightarrow^* the reflexive and transitive closure of the binary relation \Rightarrow .

4 Reachability Problem

In this section we prove that the problem of reaching a configuration (membranes and objects) starting from a certain configuration is decidable for the special class of mobile membranes systems introduced in Section 2.

Theorem 1. For two arbitrary mobile membranes M_1 and M_2 , it is decidable whether M_1 reduces to M_2 .

The main steps of the proof are as follows:

1. we reduce mobile membranes systems to pure and public mobile ambients without the capability *open*.

- 2. we show that the reachability problem for two arbitrary mobile membranes can be expressed as the reachability problem for the corresponding mobile ambients.
- 3. we use the result that the reachability problem is decidable for a fragment of pure and public mobile ambients without the capability *open*.

The following subsections are devoted to the proof of Theorem 1.

4.1 From Mobile Membranes to Mobile Ambients

We use the following translation steps:

- 1. any object $a \downarrow$ is translated into a capability in a;
- 2. any object $a\uparrow$ is translated into a capability *out* a;
- 3. any object $\overline{a} \downarrow$ is translated into a replication !in a
- 4. any object \overline{a} is translated into a replication !*out* a
- 5. a membrane h is translated into an ambient h
- 6. an elementary membrane \overline{h} is translated into a replication !h[] where all the objects inside membrane h are translated into capabilities in ambient h using the above steps.

A correspondence exists between the rules from mobile membranes and the reduction rules from mobile ambients as follows:

- rule (c) corresponds to rule (In)
- rule (d) corresponds to rule (Out)
- rules (a), (b), (e) correspond to instances of rule (**Repl**)

If we start with a mobile membrane system M, we denote by $\mathcal{T}(M)$ the mobile ambient obtained using the above translation steps. For example, starting from the membrane system $M = [m \downarrow m \uparrow]_n []_m$ we obtain $\mathcal{T}(M) = n[in \ m \ | \ out \ m] \ | \ m[]$.

Proposition 1. For mobile membrane systems M and N, M reduces to N by applying one rule if and only if $\mathcal{T}(M)$ reduces to $\mathcal{T}(N)$ by applying only one reduction rule.

Proof (Sketch). Since M reduces N by applying one rule, then one of the rules of type $(a), \ldots, (e)$ is applied. We treat only the case when a rule of type (a) is applied, the others being treated in a similar manner.

If a rule $\overline{a} \downarrow \rightarrow \overline{a} \downarrow a \downarrow$ is applied, only one object from the membrane system M is used, namely $\overline{a} \downarrow$, to create a new object $a \downarrow$, thus obtaining the membrane system N. By translating the membrane system M into $\mathcal{T}(M)$ we have that $\overline{a} \downarrow$ is translated in $!in \ a$. By applying the reduction rule corresponding to (a) (namely the rule **(Repl)**) to $!in \ m$, then we have that $!in \ a \Rightarrow !in \ a \mid in \ a$, namely a new capability in a is created. We observe that $\mathcal{T}(a \downarrow) = in \ a$, which means that the obtained mobile ambient is in fact $\mathcal{T}(N)$.

According to Proposition 1 the reachability problem for mobile membranes can be reduced to a similar problem for mobile ambients.

4.2 From Mobile Ambients to Petri Nets

After translating the mobile membranes into a fragment of mobile ambients known to be decidable, we present for our fragment of mobile ambients the algorithm used in [1] to translate mobile ambients into a fragment of Petri nets, which is known to be decidable from [6]. The fragment of mobile ambients used here is a subset of the fragment of mobile ambients used in [1] and the difference is provided by the extra-rule $!A \Rightarrow !A \mid !A$ used in [1].

We observe that applying a reduction rule over a process either increases the number of ambients or leaves it unchanged. The only reduction rule that increases the number of ambients when applied is the rule (**Repl**), while the other reduction rules leave the number of ambients unchanged. If we reach process B starting from process A, then the number of ambients of process B is known. Therefore, we can use this information to know how many times the reduction rule (**Repl**) is applied to replicate ambients. A similar argument does not hold for capabilities as they can be consumed by the reduction rules (**In**) and (**Out**).

An ambient context C is a process in which may occur some holes (denoted by \Box). Starting from this observations we split a process into two parts: one will be a context containing ambients, whereas the other one will be a process without ambients.

In order to uniquely identify all the occurrences of replication, ambient, capability or hole \Box within an ambient context or a process, we introduce a labeling system. Using a countable set of labels, we say that a process A or an ambient context C is well-labeled if any label occurs at most once in A or C. We denote by Amb(C) the multiset of ambients occurring in an ambient context C.

I) labeled Transition System

For the reachability problem $A \Rightarrow^* B$, we denote by C_A a well-labeled ambient context and with θ_A a mapping from the set of holes in C_A to some labeled processes without replicable ambients such that $\theta_A(C_A)$ is well-labeled, and $\theta_A(C_A) = A$ ignoring labels.

A labeled transition system $L_{A,B}$ describes all possible reductions for the context C_A : this includes reductions of replications and capabilities contained in C_A and in the processes associated with the holes of the context. The states of the labeled transition system $L_{A,B}$ are associative-commutative equivalent classes of ambient contexts, and for simplicity, we often identify a state as one of the representatives of its class.

We define a mapping $\theta_{L_{A,B}}$ which extends the mapping θ_A . Initially, $L_{A,B}$ contains (the equivalence class of) C_A as a unique state and we have $\theta_{L_{A,B}} = \theta_A$. We present in what follows the construction steps of $\theta_{L_{A,B}}$, where cap stands for in or out:

1. for any ambient context C from $L_{A,B}$ and for any labeled capability $cap^w n$ in C if this capability can be executed using one of the rules (In) or (Out) leading

to some ambient context \mathcal{C}' , then the state \mathcal{C}' and a transition from \mathcal{C} to \mathcal{C}' labeled by $cap^w n$ are added to $L_{A,B}$.

- 2. for any ambient context \mathcal{C} from $L_{A,B}$ and for any labeled replication $!^w$ in \mathcal{C} such that the reduction rule **(Repl)** is applied, we define the ambient context \mathcal{C}' as follows: \mathcal{C}' is identical to \mathcal{C} except that the subcontext $!^w\mathcal{C}_a$ in \mathcal{C} is replaced by $!^w\mathcal{C}_a \mid \gamma(\mathcal{C}_a)$ in \mathcal{C}' ; the mapping γ relabels \mathcal{C}_a with fresh labels, such that \mathcal{C}' is well-labeled. If $Amb(\mathcal{C}') \subseteq Amb(B)$ then the state \mathcal{C}' and a transition from \mathcal{C} to \mathcal{C}' labeled by $!^w$ is added to $L_{A,B}$. Additionally, we define $\theta'_{L_{A,B}}$ as an extension of $\theta_{L_{A,B}}$ such that for all $\Box^{w'}$ in \mathcal{C}_a we have:
 - (i) $\theta'_{L_{A,B}}(\gamma(\Box^{w'}))$ and $\theta_{L_{A,B}}(\Box^{w'})$ are identical ignoring labels,
 - (ii) labels in $\theta'_{L_{A,B}}(\gamma(\Box^{w'}))$ are fresh in the currently built transition system $L_{A,B}$
 - (iii) $\theta'_{L_{A,B}}(\gamma(\Box^{w'}))$ is well-labeled.

As a final step, we set $\theta_{L_{A,B}}$ to be $\theta'_{L_{A,B}}$.

- 3. for any ambient context \mathcal{C} from $L_{A,B}$ and for any labeled hole \Box^w in \mathcal{C} and for any capability $\operatorname{cap}^w \mathbf{n}$ in the process $\theta_{L_{A,B}}(\Box^w)$, we consider the ambient context \mathcal{C}_m identical to \mathcal{C} except that \Box^w in \mathcal{C} has been replaced by $\Box^w | \operatorname{cap}^w n$ in \mathcal{C}_m . If this capability $\operatorname{cap}^w n$ can be consumed in \mathcal{C}_m using one of the rules (In) or (Out) leading to some ambient context \mathcal{C}' , then the state \mathcal{C}' and a transition from \mathcal{C} to \mathcal{C}' labeled by $\operatorname{cap}^w n$ are added to transition system $L_{A,B}$.
- 4. for any ambient context C from $L_{A,B}$ and for any labeled hole \Box^w in C associated by $\theta_{L_{A,B}}$ with a process of the form $!^{w'}A'$, if the replication $!^{w'}$ can be reduced in the process $\theta_{L_{A,B}}(C)$ using the rule (**Repl**), then for any replication $!^{w''}$ in $\theta_{L_{A,B}}(\Box^w)$, a transition from C to itself, labeled by $!^{w''}$ is added to $L_{A,B}$.

In step 2 the reduction of a replication contained in the ambient context by means of the rule **(Repl)** is done only when the number of ambients in the resulting process is smaller than the number of ambients in the target process B, namely $Amb(\mathcal{C}') \subseteq Amb(B)$. This requirement is crucial as it implies that the transition system $L_{A,B}$ has only finitely many states.

As an example, we give in Figure 1 the labeled transition system associated with the process $n[!^{1}in \ m.!^{2}out \ m] \mid m[]$ (we omit in this process unnecessary labels). We use the labeled replications $!^{1}$ and $!^{2}$ to distinguish between different replication operators which appear in the process.

We observe that the labeled transitions in $L_{A,B}$ for replications and capabilities from an ambient context correspond to reductions performed on processes. As shown in steps 3 and 4 the transitions applied for any capabilities or replications associated with the holes are independently of the fact that they are effectively at this point available to perform a transition.



Fig. 1. A labeled transition system for the process $n[!^{1} \text{ in } m.!^{2} \text{ out } m] \mid m[$

II) From Processes Without Ambients to Petri Nets

In what follows we show how to build a Petri net from a labeled process without ambients. We denote by $\mathcal{E}(E)$ the set of all multisets that can be built with elements from the set E.

We recall that a Petri net is given by a 5-tuple $(\mathcal{P}, \mathcal{P}_i, \mathcal{T}, Pre, Post)$ with

- \mathcal{P} a finite set of places;
- $\mathcal{P} \subseteq \mathcal{P}_i$ a set of initial places;
- \mathcal{T} a finite set of transitions;
- $Pre, Post: \mathcal{T} \to \mathcal{E}(\mathcal{P})$ mappings from transitions to multisets of places.

We say that an ambient-free process is **rooted** if it is of the form $cap^w n.A'$ or of the form $!^w A'$. We define $PN_{A'}$ the Petri net associated with some rooted process A' as follows: places for $PN_{A'}$ are precisely rooted subprocesses of A', and A' itself is the unique initial place. Transitions are defined as the set of all capabilities $in^w n$, $out^{w'}n$ and replications $!^w$ occurring in A'. Finally, Pre and Post are defined for all transitions as follows:

- $Pre(cap^w n) = \{cap^w n\}$ and $Post(cap^w n) = \emptyset$ if $cap^w n$ is a place in $PN_{A'}$.
- $Pre(cap^w n) = \{cap^w n.(A_1 \mid \ldots \mid A_k)\}$ and $Post(cap^w n) = \{A_1 \mid \ldots \mid A_k\}$ if $cap^w n.(A_1 \mid \ldots \mid A_k)$ is a place in $PN_{A'}$ $(A_1 \mid \ldots \mid A_k)$ being rooted processes).
- $Pre(!^w) = Pre(!^w) = \{!^w A'\}, Post(!^w) = \{!^w A', A'\} \text{ and } Post(!^w) = \{!^w A'\} \text{ if } !^w A' \text{ is a place in } PN_{A'}.$

For $!^{1}in \ m.!^{2}out \ m$, we obtain the Petri net given in Figure 2.



Fig. 2. A Petri net for the process $!^1$ in $m.!^2$ out m

We will denote by PN_{\square^w} the Petri net $PN(\theta_{L_{A,B}}(\square^w))$, that is, the Petri net corresponding to the rooted ambient-free process associated with \square^w by $\theta_{L_{A,B}}$. In what follows we show how to combine the transition system $L_{A,B}$ and the Petri nets PN_{\square^w} into one single Petri net.

III) Combining the Transition System and Petri Nets

We first turn the labeled transition system $L_{A,B}$ into a Petri net $PN_L = (\mathcal{P}_L, \mathcal{P}_L^i, \mathcal{T}_L, Pre_L, Post_L)$ with:

- \mathcal{P}_L a set of states of $L_{A,B}$;
- \mathcal{P}_L^i a singleton set containing the state corresponding to \mathcal{C}_A , the ambient context of A;
- T_L the set of transitions of the form (s, l, s'), with
 - s and s' states from $L_{A,B}$;
 - l transition from s to s' in $L_{A,B}$.
- Pre(t) = s and $Post(t) = \{s'\}$, for all transitions t = (s, l, s').

We define the Petri net $PN_{A,B} = (\mathcal{P}_{A,B}, \mathcal{P}_{A,B}^i, \mathcal{T}_{A,B}, Pre_{A,B}, Post_{A,B})$ as:

- places (resp. initial places) from $PN_{A,B}$ are the union of places (resp. initial places) of PN_L and of each of the Petri nets PN_{\square^w} (for \square^w occurring in one of the states of $L_{A,B}$).
- transitions of $PN_{A,B}$ are precisely the transitions of PN_L .
- The mappings Pre_{A,B} and Post_{A,B} are defined for all transitions t = (a, f, b):
 (i) Pre_{A,B}(t) = {a} and Post_{A,B}(t) = {b}, if f does not occur as a transition in any PN_{□^w} (for □^w occurring in one of the states of L_{A,B})
 - (ii) $Pre_{A,B}(t) = \{a\} \cup Pre_{\square^w}(f)$ and $Post_{A,B}(t) = \{b\} \cup Post_{\square^w}(f)$, if f is a transition of PN_{\square^w} ($Pre_{\square^w}(\text{resp. }Post_{\square^w})$ being the mapping Pre (resp. Post) of PN_{\square^w}).

4.3 Deciding Reachability

We recall that for a Petri net $PN = (\mathcal{P}, \mathcal{P}^i, \mathcal{T}, Pre, Post)$, a marking m is a multiset from $\mathcal{E}(P)$. A transition t is enabled by a marking m if $Pre(t) \subseteq m$. Executing an enabled transition t for a marking m gives a marking m' defined as $m' = (m \setminus Pre(t)) \cup Post(t)$ (where \setminus stands for the multiset difference). A marking m' is reachable from m if there exists a sequence m_0, \ldots, m_k of markings such that $m_0 = m, m_k = m'$ and for each m_i, m_{i+1} , there exists an enabled transition for m_i whose execution gives m_{i+1} .

Theorem 2 ([6]). For all Petri nets P, for all markings m, m' for P, one can decide whether m' is reachable from m.

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For the reachability problem $A \Rightarrow^* B$ over ambients, we consider the Petri net $PN_{A,B}$ and the initial marking m_A defined as $m_A = \mathcal{P}^i_{A,B}$. In Figure 3 is depicted the initial marking for the process $n[!^{1}in \ m.!^{2}out \ m] \mid m[]$ as a combination of the labeled transition system from Figure 1 and the Petri net from Figure 2.



Fig. 3. The Petri net for the labeled process $n[!^{1}$ in $m.!^{2}$ out $m \mid m[$

It should be noticed that for any marking m reachable from m_A , m contains exactly one occurrence of a place from \mathcal{P}_L . Roughly speaking, to any reachable marking corresponds exactly one ambient context. Moreover, the execution of one transition in the Petri net $PN_{A,B}$ simulates a reduction from \Rightarrow .

We define now \mathcal{M}_B , the set of markings of $PN_{A,B}$ corresponding to B. Intuitively, a marking m belongs to \mathcal{M}_B if m contains exactly one occurrence C of a place from \mathcal{P}_L (that is, representing some ambient context) and in the context C, the holes can be replaced with processes without ambients to obtain B. Each of the processes without replication must correspond to a marking of the sub-Petri net associated with the hole it fills up. \mathcal{M}_B is defined as the set of markings m for $PN_{A,B}$ satisfying:

- (i) in *m* there exists exactly one ambient context C_m ;
- (ii) ignoring labels, $\sigma_m(\mathcal{C}_m)$ is equal to B modulo associative-commutative, for the substitution σ_m from holes \Box^w occurring in \mathcal{C}_m to processes without ambients defined as: $\sigma_m(\Box_m) = P_1 \mid \ldots \mid P_k$ for $\{P_1, \ldots, P_k\}$ the multiset corresponding to the restriction of m to the places of PN_{\Box^w}
- (iii)for all holes \Box^w occurring in some state of the transition system $L_{A,B}$ but not in \mathcal{C}_m , the restriction of m to places of PN_{\Box^w} is precisely the set of initial places from PN_{\Box^w} .

We adapt the results presented in [1] to a restricted fragment of mobile ambients.

Proposition 2. For a Petri net $PN_{A,B}$, there are only finitely many markings corresponding to the process B, and their set \mathcal{M}_B can be computed.

The translation correctness is ensured by the following result.

Proposition 3. For all processes A, B we have that $A \Rightarrow B$ if and only if there exists a marking from \mathcal{M}_B such that m_B is reachable from m_A in $PN_{A,B}$.

Using the Proposition 3 and Theorem 2, we can decide whether an ambient A can be reduced to an ambient B.

Theorem 3. For two arbitrary ambients A and B from our restricted fragment, it is decidable whether A reduces to B.

5 Conclusions

In this paper we have investigated the problem of reaching a configuration in mobile membranes starting from another configuration. In order to do this we use [1] where the reachability problem for a fragment of ambient calculus is proven to be decidable, namely the pure and public ambient calculus except the *open* capability. The same problem is tackled in [2], but the authors do not take into account the replication of ambients, which is used to simulate the division rules in mobile membranes. We proved that the reachability can be decided by reducing this problem to the reachability problem of a version of pure and public ambient calculus from which the **open** capability has been removed.

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Modeling Symport/Antiport P Systems with a Class of Hierarchical Petri Nets^{*}

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Summary. A model of P systems with symport / antiport rules is given in terms of hypernets, a generalization of a class of hierarchical Petri nets introduced for modeling mobility inside the nets-within-nets paradigm. The hierarchical structure of a P system is reflected by the associated hypernet, where molecules are modeled by unstructured agents (simple tokens) and membranes by agents. Each agent is modeled by a net which may contain in its places unstructured agents or other agents. Agents can exchange tokens with their sub- or super-agents and thus the hierarchy may change. The main result of the paper shows a correspondence between reachable configurations of the P system and reachable hypermarkings of the related hypernet, in such a way that if the P system can evolve from one configuration to another one then in the hypernet there exists a corresponding transformation of hypermarkings.

1 Introduction

In recent years the notion of *system of mobile agents* has gained importance in computer science and engineering. These systems are formed by *agents* which move around a space, interacting with each other. Often, these agents are pieces of software traveling across a network of hosts, where they can be executed in a local environment. Such a development has led to envisage formal models in which one can represent mobile agents, their environment, and their interactions. Since agents move and run in parallel with others, concurrency theory is a natural framework in which to look for adequate models.

In 1986, Valk proposed a kind of Petri nets in which tokens can be nets, which can be moved across the places of a hosting net, possibly interacting with it (see [15]). Building on this idea, *hypernets* were defined in [1]. A hypernet is formed by agents, each modeled by a Petri net. In a given configuration, each agent, except one, is also a token residing in a place of another agent (the exception consists in the highest level agent, which acts as an environment for all others). The relation

^{*} Partially supported by MIUR and CNR - IPI PAN.

of containment can dynamically change as an effect of firing transitions; agents can exchange their sub-agents by forming so called *consortia*.

The hierarchy of agents in a hypernet resembles the hierarchy of membranes in a P system, and the mechanism of consortia can be seen as a way to exchange molecules across a membrane. This idea is the subject of the present paper, where we define a translation from P systems with symport/antiport rules to a class of hypernets. Such class is a generalization of the class defined in [1]. The main idea of this translation is quite simple: each membrane and each individual molecule in the P system is represented by an agent in the hypernet. Molecule agents are unstructured, that is, they are simple tokens, like in usual nets, and can only be passively moved by the active components. Membrane agents, viceversa, are nets, with places that can contain molecule agents, and places that can contain other membrane agents. Consortia correspond to rules of the P system, whereby molecules can be exchanged across a membrane.

It should be noted that hypernets would allow, in themselves, movement of membrane agents, so that the hierarchical structure of membranes could change. This capability is not exploited here, since we deal with P systems where only molecules move around, but might be useful in modeling more general kinds of systems.

In this paper, we are not interested in the computational aspects of the theory of P systems, but rather focus on modeling aspects. Consequently, we compare the two models on the basis of their reachable configurations.

After recalling the basic definitions related to the class of P systems with symport/antiport rules (Section 2), we define hypernets in Section 3. Section 4 shows how to build a hypernet from a P system, and states in which sense the two models can be considered as equivalent. Finally, in Section 5, we draw some considerations, and suggest possible developments.

2 P Systems with Symport/Antiport Rules

Many kinds of membrane systems have been investigated during the last years. One of the most studied variant of the general model of P systems was introduced in [10] under the name of systems with symport/antiport rules. Those terms came from two membrane transport mechanisms. Whereas the term symport stands for the biological process by which two molecules pass together across a membrane, when the two molecules pass simultaneously, but in the opposite direction, the process is called antiport.

The class of membrane systems with symport/antiport rules is a class of purely communicating P systems, where the objects involved in the computation only pass through membranes. This means that the objects involved never change and a sort of conservation law for objects is observed during the entire evolution of the system. Many results on this kind of P systems, especially about their computational power, can be found in [11],[7],[8],[4]. Here we provide a simplified version of the definition of P system with symport/antiport rules supplied by Păun [12].

2.1 Formal definition

Formally, we define a P system with symport/antiport rules (of degree m), as a construct of the form

$$\Pi = (O, \mu, w_1, w_2, \dots, w_m, R_1, R_2, \dots, R_m),$$

where:

- *O* is the (finite and non empty) alphabet of objects
- the membrane structure $\mu = (N, E, i)$ is a rooted tree underlying Π , where $N = \{1, 2, \ldots, m\}$ is the set of nodes and each node in N defines a membrane of Π . The set $E \subseteq N \times N$ defines the edges. For each node $j \in N$, the membrane associated to the node j contains all the membranes associated to the children of j. i is the root of the tree and hence the skin membrane (the outermost membrane of the system)
- w_1, w_2, \ldots, w_m are multisets over O representing the objects present in the regions $1, 2, \ldots, m$ of the membrane structure μ in the initial configuration of the system (in the following, multisets will be described either by strings, with exponents denoting the multiplicity of elements, or by the usual characteristic function of multisets) where the
- R_1, R_2, \ldots, R_m are finite sets of evolution rules associated with the membranes of μ . Moreover we impose $R_i = \emptyset$, where *i* is the skin of the membrane structure. This clause ensures that the external membrane is impermeable and hence the total number of objects involved in the computation is finite (and constant); this is required if we want to build hypernets with a finite number of agents

In the following we often use the term *molecule* when referring to an object in a membrane of the P system.

As said above, each rule governs the communication through a specific membrane and can be of two kinds, symport rule or antiport rule. A symport rule is of the form (u, in) or (u, out), where u is a multiset over O, stating that all the objects of u pass together through a membrane, entering in the former case and exiting in the latter. For example, in a membrane i, after the application of the symport rule (u, in), the multiset associated to this membrane will contain all the objects previously present, plus the objects present in u. The multiset associated to the membrane that contains i, will contain all the objects previously present, minus those in u. Similarly, an antiport rule is of the form (u, out; v, in), where uand v are multisets over O, stating that when u exits, at the same time, a multiset v must enter the membrane.

The P system described above evolves from configuration to configuration by the application of a multiset of rules in each membrane. Formally, a configuration is a tuple $C = (v_1, v_2, \ldots, v_m)$ and $C \stackrel{\hat{R}}{\Rightarrow} C'$ denotes that C evolves into C' due to the application of \hat{R} , where $\hat{R} = (\bar{R}_1, \bar{R}_2, \ldots, \bar{R}_m)$ is a multi-rules vector applicable to C and \bar{R}_j is a multiset over R_j .

The evolution of the system is non-deterministic and maximally parallel: at each step, the configuration changes by applying a maximal multiset of rules, chosen in a non deterministic way; the rules must be all applicable without mutual interferences in the current configuration.



Fig. 1. Fragment of a symport/antiport P system

2.2 Example

Fig. 1 shows a fragment of a P system with symport rules. The system depicted here consists of two nested membranes: j, the inner membrane, and i, the outer one, which we assume to be a membrane contained in a larger membrane structure. The set of rules of i is $R_i = \{(b, out), (a^2, out)\}$, and the set of rules of j is $R_j = \{(ab^2, in), (a, out)\}$. In the same way we define the initial multisets of objects $w_i = ab^3$ and $w_j = a^2$.

In this configuration the rules r_1, r_3, r_4 are enabled and a multi-rules vector can be built with this rules in a maximally parallel manner, i.e.: the multi-rules vector $\hat{R} = (\{r_1\}, \{r_3, r_4, r_4\})$ is applicable to the initial configuration. Note that other multi-rules vectors can be applied to the same configuration. The application of \hat{R} leads to a new state where the objects in the membrane *i* are a^2 and the objects in the membrane *j* are ab^2 .

3 Hypernets

In this section we introduce a generalization of Petri hypernets [1] that for simplicity we call also here *hypernets*. A hypernet is defined by a fixed set of agents, each

agent is modeled by a net and can manipulate other agents as tokens, while being manipulated as token by another agent at the same time. This yields a hierarchy of agents. The highest level agent acts as an environment for all other agents, these latter are located each one in some place of another agent. Agents can exchange tokens with their sub- or super-agents and thus the hierarchy may change.

In what follows we first define the structure of hypernets giving the definition of agent and of hypernet, then we define the behavior of hypernets, and at the end of the section we illustrate hypernets on an example modeling the P system given in Subsection 2.2 as it will be discussed in Section 4.

3.1 Structure of hypernets

An agent is modeled by a Petri net, a bipartite oriented graph, whose nodes are of two types: places and transitions. Places are partitioned into two disjoint sets: the set of local places, which are locations in which other agents can stay, and the set of virtual places, which represent communication channels along which agents exchange tokens each others. Places and transitions are interconnected by weighted oriented arcs, which define how many tokens are taken away from an input place and how many are put into an output place, when a transition fires. For each transition the sum of the weights of the input arcs must be equal to the sum of the weights of the output arcs. In this way the amount of tokens will not change while transitions fire. Moreover, to each triple of interconnected elements place-transition-place it is assigned, by a function ϕ_A , a value which defines, in a way compatible with the arc weights, the number of tokens which flow along the path identified by the triple. In other words, ϕ_A defines how the tokens taken away from an input place of a transition will be distributed into the output places, when the transition will fire; and this distribution will be the same for each occurrence of the same transition. For basic definitions and notions on Petri nets, see, for example, [14].

Definition 1. An agent is a tuple $A = (P_A \cup V_A, T_A, F_A, \phi_A)$, where $(P_A \cup V_A, T_A, F_A)$ is a, possibly empty, finite Petri net in which :

- P_A is the set of local places and V_A is the set of virtual places, (or communication places), with $P_A \cap V_A = \emptyset$;
- T_A is the set of transitions;
- the function $F_A : ((V_A \cup P_A) \times T_A) \cup (T_A \times (V_A \cup P_A)) \longrightarrow \mathbb{N}$ defines the flow, assigning a weight to each arc identified by the pair of elements x, y such that: $F_A(x, y) > 0$, in such a way that $: \forall t \in T_A, \sum_{p \in \bullet t} F_A(p, t) = \sum_{p \in t^\bullet} F_A(t, p)$, where $p \in \bullet t$ iff $F_A(p, t) > 0$ and $p \in t^\bullet$ iff $F_A(t, p) > 0$;

and the function $\phi_A : (V_A \cup P_A) \times T_A \times (V_A \cup P_A) \longrightarrow \mathbb{N}$ defines the paths, *i.e.*: the triples (p, t, q) such that: $\phi_A(p, t, q) > 0$, by assigning a weight to them in such a way that:

$$\begin{aligned} \forall p \in {}^{\bullet}t, \ F_A(p,t) &= \sum_{q \in t^{\bullet}} \phi_A(p,t,q) \\ \forall q \in t^{\bullet}, \ F_A(t,q) &= \sum_{p \in {}^{\bullet}t} \phi_A(p,t,q) \end{aligned}$$

In the following $(p, t, q) \in \phi_A$ iff $\phi_A(p, t, q) > 0$, moreover, given a subset of agents $X \subseteq \mathcal{N}$, we use the following notation: $P_X = \bigcup_{A \in X} P_A$, $V_X = \bigcup_{A \in X} V_A$, $T_X = \bigcup_{A \in X} T_A$, $\phi_X = \bigcup_{A \in X} \phi_A$.

A hypernet is defined by a set of agents and by a relation Δ . Agents have disjoint sets of places. A transition may belong to different agents, modeling synchronous interaction among them. Transitions connected to virtual places model interchanges of tokens among sub-/super-agents. Said *output paths* the path ending with a virtual place and *input paths* the ones starting with a virtual place, the relation Δ identifies communication channels by defining, for a given transition belonging to different agents, a correspondence (output path - input path) in a way compatible with path weights.

Definition 2. Let $\mathcal{N} = \{A_1, A_2, \ldots, A_n\}$ be a family of agents, and let $S^o = \{(p, t, v) \in \phi_A | A \in \mathcal{N} \text{ and } v \in V_A\}$ and $S^i = \{(v, t, q) \in \phi_A | A \in \mathcal{N} \text{ and } v \in V_A\}$ be the sets of output paths and input paths, respectively. (Note that a path can be both an output and an input path.)

A hypernet is a pair $H = (\mathcal{N}, \Delta)$, where

• The agents in N have disjoint sets of places:

 $\forall A_i, A_j \in \mathcal{N}, \ (P_{A_i} \cup V_{A_i}) \cap (P_{A_i} \cup V_{A_i}) = \emptyset;$

 and Δ ⊆ S^o × Sⁱ is a relation which associates, for a given transition, output paths to input paths with the same weight and belonging to different agents, i.e.:

 $\begin{array}{l} \forall t \in T_{\mathcal{N}}, \, \forall (p,t,q) \in \phi_{A_i} \, and \, \forall (p',t,q') \in \phi_{A_j} \, such \, that \, A_i, A_j \in \mathcal{N} \colon \\ ((p,t,q),(p',t,q')) \in \varDelta \Rightarrow A_i \neq A_j \, and \, \phi_{A_i}(p,t,q) = \phi_{A_j}(p',t,q'). \end{array}$

Definition 3. Let $\mathcal{N} = \{A_1, A_2, \ldots, A_n\}$ be a family of agents. A map $\mathcal{M} : \{A_2, \ldots, A_n\} \longrightarrow P_{\mathcal{N}}$, assigning to each agent different from A_1 the local place in which is located, is a hypermarking of \mathcal{N} iff, considering the relation $\uparrow_{\mathcal{M}} \subseteq \mathcal{N} \times \mathcal{N}$ defined by : $A_i \uparrow_{\mathcal{M}} A_j \Leftrightarrow \mathcal{M}(A_i) \in P_{A_j}$, then the graph $\langle \mathcal{N}, \uparrow_{\mathcal{M}} \rangle$ is a tree with root A_1 .

Definition 4. A marked hypernet is a pair (H, \mathcal{M}) where H is a hypernet and \mathcal{M} is a hypermarking defining the initial configuration.

In a configuration the system results hierarchically structured. The highest level agent A_1 , the root of the tree describing the hierarchy, plays the role of the environment containing all the other agents. The relation of containment between agents, and then the hierarchical structure, can change as an effect of firing transitions as formalized in the following subsection.

3.2 Behaviour of hypernets

Let $H = (\mathcal{N}, \Delta)$, with $\mathcal{N} = \{A_1, A_2, \dots, A_n\}$, be a hypernet.

A consortium is a set of interconnected active agents, cooperating in performing a transition t, moving other passive agents along the paths containing t.

Definition 5. A consortium is a tuple $\Gamma = (t, \tau, \delta, \gamma)$ where:

- $t \in T_{\mathcal{N}}$ is the name of the consortium,
- $\tau \subseteq \{A \in \mathcal{N} | t \in T_A\}, \tau \neq \emptyset$, is the non empty set of active agents. To this set we can associate $\phi_{\tau_t} = \{(p, t, q) \in \phi_\tau \mid p, q \in P_\tau \cup V_\tau\}$, the set of paths of the agents τ containing the transition t.
- δ defines a bijective correspondence between output paths containing t and input paths containing t of active agents, without contradicting the relation Δ . Let $\phi_{o,\tau_t} = \phi_{\tau_t} \cap S^o$ and $\phi_{i,\tau_t} = \phi_{\tau_t} \cap S^i$. If $\phi_{o,\tau_t} \neq \emptyset$, $\delta : \phi_{o,\tau_t} \longrightarrow \phi_{i,\tau_t}$ is a bijection such that: $\forall s \in \phi_{o,\tau_t}$, $\delta(s) = s' \Rightarrow (s, s') \in \Delta$, while if $\phi_{o,\tau_t} = \emptyset$, then δ is the empty map. Note that δ relates paths belonging to different agents.
- The passive agents which are moved when the consortium occurs are selected through the map γ . Let $C \subseteq \mathcal{N} \setminus A_1$ be a chosen set of passive agents, then $\gamma: C \longrightarrow \phi_{\tau_t} \setminus S^i$ is surjective and associates as many passive agents to each path containing t and belonging to an active agent as the weight of the path itself, i.e.: $\forall s \in \phi_{\tau_t} \setminus S^i$, $|\gamma^{-1}(s)| = \phi_{\mathcal{N}}(s)$. Note that an agent can be active and passive at the same time.

Moreover the following conditions must be satisfied:

- the set of active agents τ is a minimal one, in the sense that the agents in τ must be each other interconnected through the interaction t, i.e.: the undirected graph $G_1 = (\tau, E_1)$ is connected, where $E_1 = \{(A_i, A_j) \mid A_i, A_j \in \tau \text{ and } \exists s_i \in \phi_{A_i}, \exists s_j \in \phi_{A_j} : \delta(s_i) = s_j\}$ and
- the undirected graph $G_2 = (\tau \cup C, E_2)$ is acyclic, where E_2 connects A_i to A_j if A_i will be put inside A_j through t, i.e.: considered the recursively defined map $\delta^* : \phi_{\tau_t} \longrightarrow \phi_{\tau_t}$ such that

$$\delta^*(s) = \begin{cases} s & \text{if } s \notin \phi_{o,\tau_i} \\ \delta^*(\delta(s)) & \text{otherwise} \end{cases}$$
$$E_2 = \{ (A_i, A_j) | \delta^*(\gamma(A_i)) \in \phi_{A_j}, A_i \in C, A_j \in \tau \}.$$

The intuition behind the last condition of the previous definition is the following. By subsequent applications of the map δ it is possible to construct chains of paths interrelated through paths with only virtual places. However, the meaningful chains are the one which starts with a path with a real input place, the one from which an agent will be taken out, and ends with a path with a real output place, the one in which the agent will be put into. The last condition requires that these chains are not closed.

In [2] it is proven that chains containing a real place can be prolonged to finite chains containing at most two real places, one in an input path and one in an output path. **Definition 6.** Let $H = (\mathcal{N}, \Delta)$ be a generalized hypernet and \mathcal{M} be a hypermarking.

A consortium $\Gamma = (t, \tau, \delta, \gamma)$ is enabled in \mathcal{M} , denoted $\mathcal{M}[\Gamma\rangle$, iff the following two conditions hold

- $\forall A \in C, \ \gamma(A) = (p, t, q) \Rightarrow \mathcal{M}(A) = p$
- $\forall A_i, A_j \in \tau, \forall s \in S^o \cap \phi_{A_i}, \, \delta(s) \in \phi_{A_j} \Rightarrow A_i \uparrow_{\mathcal{M}} A_j \lor A_j \uparrow_{\mathcal{M}} A_i$

If $\mathcal{M}[\Gamma\rangle$, then the occurrence of Γ leads to the new hypermarking \mathcal{M}' , denoted $\mathcal{M}[\Gamma\rangle\mathcal{M}'$, such that $\forall A \in \mathcal{N}$:

$$\mathcal{M}'(A) = \begin{cases} \mathcal{M}(A) \text{ if } A \notin C; \\ q \quad \text{if } A \in C \text{ and } \delta^*(\gamma(A)) = (p, t, q). \end{cases}$$

It is possible to prove [2] that \mathcal{M}' is a hypermarking, i.e.: that the class of hypermarkings of a hypernet is closed under the occurrence of a consortium.

Two consortia $\Gamma_1 = (t_1, \tau_1, \delta_1, \gamma_1)$ and $\Gamma_2 = (t_2, \tau_2, \delta_2, \gamma_2)$ are *independent* iff the maps γ_1 and γ_2 select two different sets of passive agents, i.e.: iff $C_1 \cap C_2 = \emptyset$

If two independent consortia are both enabled in a hypermarking \mathcal{M} then they can *concurrently* occur in \mathcal{M} .

Let Γ_H be the set of possible consortia in H. A set of consortia $U \subseteq \Gamma_H$ is a *step* enabled in a hypermarking \mathcal{M} , denoted $\mathcal{M}[U\rangle$, iff

- $\forall \Gamma_i, \Gamma_j \in U, \ \Gamma_i \text{ and } \Gamma_j \text{ are independent},$
- $\forall \Gamma_i \in U, \mathcal{M}[\Gamma_i)$

If $\mathcal{M}[U\rangle$, then the occurrence of the step U leads to the new hypermarking \mathcal{M}' , denoted $\mathcal{M}[U\rangle\mathcal{M}'$, such that $\forall A \in \mathcal{N}$:

$$\mathcal{M}'(A) = \begin{cases} \mathcal{M}(A) \text{ if } \forall \Gamma_i \in U, A \notin C_i; \\ q \quad \text{if } \exists \Gamma_i \in U : (A \in C_i \text{ and } \delta_i^*(\gamma_i(A)) = (p, t_i, q)). \end{cases}$$

U is a maximal step enabled in \mathcal{M} , and its occurrence yields \mathcal{M}' , iff $\mathcal{M}[U\rangle\mathcal{M}'$ and $\forall U' \supset U : not(\mathcal{M}[U'\rangle).$

In [2] it is shown how it is possible to associate to each hypernet a 1-safe net in such a way that there is a strict correspondence between their behaviors, i.e., in terms of Petri net theory, in such a way that the case graph of the 1-safe net is isomorphic to the transition system generated by the reachable hypermarkings of the hypernet.

Since 1-safe nets are a basic class model in Petri net theory, this translation shows that hypernets are well rooted inside the theory of Petri nets.

3.3 Example

The Fig. 2 shows the structure of two hypernet's agents. The unfilled circles are local places while the filled ones are virtual places. The agent A_i is nested in the

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Fig. 2. Fragment of a hypernet

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agent A_i , in fact $\mathcal{M}(A_j) = a_j^i$, so $A_j \uparrow_{\mathcal{M}} A_i$. Moreover we assume u_1, u_2, u_3, u_4 to be unstructured agents such that $\mathcal{M}(u_1) = \mathcal{M}(u_2) = \mathcal{M}(u_3) = b^i$ and $\mathcal{M}(u_4) = a^i$. Now consider the consortium $\Gamma = (r_3, \tau, \delta, \gamma)$ where

- the set of active agents is $\tau = \{A_i, A_j\},\$
- the bijection δ builds two communication channels between A_i and A_j gluing two pair of paths: $\delta(a^i, r_3, \bar{a}^i) = (\bar{a}^j, r_3, a^j)$ and $\delta(b^i, r_3, \bar{b}^i) = (\bar{b}^j, r_3, b^j)$,
- the set of passive agents is $C = \{u_1, u_2, u_4\}$ and $\gamma(u_1) = \gamma(u_2) = (b^i, r_3, \overline{b}^i)$ and $\gamma(u_4) = (a^i, r_3, \overline{a}^i)$.

The consortium Γ is valid and enabled in the initial hypermarking. When Γ occurs the system reaches a new hypermarking \mathcal{M}' where $\mathcal{M}'(u_1) = \mathcal{M}'(u_2) = b^j$ and $\mathcal{M}'(u_4) = a^j$. Note that the agents u_1, u_2, u_4 pass through the communication channels established by δ from the agent A_i to the agent A_j .

4 Membrane Systems as Hypernets

Our goal in this section is to show how a P system with symport/antiport rules and with an impermeable external membrane can be modeled as a hypernet.

In the following, we write $i \triangleleft j$ to mean that membrane *i* is directly contained in membrane *j*.

Let $\Pi = (O, \mu, w_1, \ldots, w_m, R_1, \ldots, R_m)$ be a P system of degree m, with symport/antiport rules. We assume that 1 is the outer membrane, with no rules, so that $R_1 = \emptyset$.

The hypernet associated to Π will be denoted by $H = (\mathcal{N}, \Delta)$. The hypernet H contains one agent for each membrane, and one agent for each individual molecule in Π . Notice that, in the P-systems we handle, molecules are neither created nor deleted.

Let $W = \sum_{i=1}^{m} w_i$. W is a multiset giving the total number of objects for each type in the system. Define

$$MOL = \{(x, i) | x \in O \land 1 \le i \le W(x)\}$$

For each (x, i) in MOL, we define an unstructured agent in the hypernet H.

$$\mathcal{N} = \{A_1, A_2, \dots, A_m\} \cup \text{MOL}$$

Agent A_i corresponds to membrane *i* of the P system. It has one place for each membrane directly contained in *i*, and one for each type of molecule; moreover, it has one virtual place for each type of molecule, to be used in exchanging tokens.

$$P_i = \{a_j^i | j \triangleleft i\} \cup \{x^i | x \in O\}$$
$$V_i = \{\bar{x}^i | x \in O\}$$

The set of transitions of agent A_i has one transition for each rule in membrane i, and one for each rule in membranes directly contained in i.

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$$T_i = \{r | r \in R_i\} \cup \{r | r \in R_j \land j \triangleleft i\}$$

We now turn to define the flow function and the paths for agent A_i .

• For each rule $r = (u, in) \in R_i$, and for each rule $r = (u, in; v, out) \in R_i$:

$$F(\bar{x}^{i}, r) = F(r, x^{i}) = \phi((\bar{x}^{i}, r, x^{i})) = u(x)$$

• For each rule $r = (v, out) \in R_i$, and for each rule $r = (u, in; v, out) \in R_i$:

$$F(x^{i}, r) = F(r, \bar{x}^{i}) = \phi_{i}((x^{i}, r, \bar{x}^{i})) = v(x)$$

Let $j \triangleleft i$. Then,

• For each rule $r = (u, in) \in R_j$, and for each rule $r = (u, in; v, out) \in R_j$:

$$F(x^{i}, r) = F(r, \bar{x}^{i}) = \phi_{i}((x_{i}, r, \bar{x}^{i})) = u(x)$$

• For each rule $r = (v, out) \in R_j$, and for each rule $r = (u, in; v, out) \in R_j$:

$$F(\bar{x}^{i}, r) = F(r, x_{i}) = \phi_{i}((\bar{x}^{i}, r, x_{i})) = v(x)$$

Define now the Δ relation. For all i, j such that $i \triangleleft j$:

$$\begin{aligned} \forall r &= (u, in) \in R_i, \forall x \in O : u(x) > 0, (x^j, r, \bar{x}^j) \Delta(\bar{x}^i, r, x^i) \\ \forall r &= (u, out) \in R_i, \forall x \in O : u(x) > 0, (x^i, r, \bar{x}^i) \Delta(\bar{x}^j, r, x^j) \\ \forall r &= (u, in; v, out) \in R_i, \forall x \in O : u(x) > 0, (x^j, r, \bar{x}^j) \Delta(\bar{x}^i, r, x^i) \\ \forall r &= (u, in; v, out) \in R_i, \forall x \in O : v(x) > 0, (x^j, r, \bar{x}^j) \Delta(\bar{x}^i, r, x^i) \end{aligned}$$

The initial hypermarking \mathcal{M} reflects the initial configuration of Π . Membrane agents are placed according to the hierarchical structure of Π :

$$\forall i \in \{2, \dots, m\} : \mathcal{M}(A_i) = a_i^j \text{ iff } i \triangleleft j$$

All agents (x, k) corresponding to molecules are initially distributed in the corresponding places x^i in membrane agents so that a place x^i contains $w_i(x)$ unstructured agents of type (x, k).

In order to state the exact relation between the dynamics of a P system Π and the dynamics of the corresponding hypernet H, we need to define two relations. The first defines a correspondence between configurations of Π and hypermarkings of H. The other defines a correspondence between steps of Π and maximal steps of H. Define

Conf = {
$$(v_1, \dots, v_m) | \sum_{1}^{m} v_i = \sum_{1}^{m} w_i }$$

as the set of all potential configurations of Π with the same number and type of molecules as the initial configuration. Define **HM** as the set of all hypermarkings of H.
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Let $\mathcal{M} : \{A_1, \ldots, A_m\} \cup \text{MOL} \to P$ be an element of **HM**, where P is the set of all local places of H, and $C = (v_1, \ldots, v_m) \in \text{Conf}$, where $v_i : O \to \mathbb{N}$. We also need some auxiliary definition. By $I(x, i, \mathcal{M})$ we denote the set of agents representing molecules of type x hosted in the corresponding place of agent A_i in \mathcal{M} .

 $I(x, i, \mathcal{M}) = \{(x, i) | (x, i) \in \text{MOL} \land \mathcal{M}((x, i)) = x^i\}$

Definition 7. The hypermarking \mathcal{M} simulates configuration C (denoted by $\mathcal{M} \sim C$) iff

1. $\mathcal{M}(A_i) \in P_j$ iff $i \triangleleft j$, for $i \in \{2, \ldots, m\}$ 2. $|I(x, i, \mathcal{M})| = v_i(x)$

Notice that \sim is a partial surjective function: each configuration of Π has at least one corresponding hypermarking. The hypermarkings corresponding to one given configuration differ only for the distribution of molecules of the same kind in membrane agents. These molecules are identical in the P system, while their corresponding agents are distinguished.

We now define a correspondence between maximal steps in the P system and maximal steps of consortia in the hypernet. This correspondence is based on another one, associating single rules and consortia.

Let r be a rule of membrane i in Π . By construction, the associated hypernet has two transitions labeled by r, one in the agent corresponding to i, and one in the agent corresponding to the membrane containing i; assume it is j. A consortium simulating the execution of r involves i and j as active agents, and a number of passive agents taken from MOL.

We consider here an antiport rule r = (u, in; v, out), where u and v are multiset on O. Symport rules can be seen as special cases where either u or v is the empty multiset.

Definition 8. Let $\Gamma = (r, \tau, \delta, \gamma)$ be a consortium. Then $\Gamma \sim r$ iff the following conditions hold.

1. $\tau = \{A_i, A_j\}$

- 2. The output paths involved in Γ are either of the form (y^i, r, \bar{y}^i) if v(y) > 0, or of the form (x^j, r, \bar{x}^j) if u(x) > 0.
- 3. The function δ is defined by

$$\begin{split} \delta((y^i,r,\bar{y}^i)) &= (\bar{y}^j,r,y^j) \\ \delta((x^j,r,\bar{x}^j)) &= (\bar{x}^i,r,x^i) \end{split}$$

4. Let $Z = \{z \in \text{MOL} | z = (x, k) \land \gamma(z) = (x^j, r, \bar{x}^j)\}$; then |Z| = u(x)5. Let $Z = \{z \in \text{MOL} | z = (y, k) \land \gamma(z) = (y^i, r, \bar{y}^i)\}$; then |Z| = v(y)

A transition in a P system is a multiset of independently executable rules. Let $R = \bigcup_{i=1}^{m} R_i$ be the set of all rules of Π , and $\rho : R \to \mathbb{N}$ be a multiset of rules. A set U of consortia in H simulates ρ (denoted by $U \sim \rho$) if, for each $r \in R$, U contains $\rho(r)$ consortia which simulate r, and the consortia in U are pairwise independent.

We are now ready to state the main result of this section. The following lemma states that any change of configuration in Π can be simulated by a set of mutually independent consortia in H. Let Π be a P-system with symport and antiport rules, such that 1 is the outer membrane with $R_1 = \emptyset$, and $H = (\mathcal{N}, \Delta)$ be the associated hypernet, with initial hypermarking \mathcal{M} .

Lemma 1. Let C be a configuration of Π , and ρ be a multiset of rules, enabled at C, with $C \stackrel{\rho}{\Rightarrow} C'$. Then, for all $\mathcal{M} \in \mathbf{HM}$,

$$\mathcal{M} \sim C \Rightarrow \exists U \subseteq \Gamma_H : U \sim \rho, \ \mathcal{M}[U\rangle \mathcal{M}', \ \mathcal{M}' \sim C'$$

Notice that the consortia forming U can always be chosen to be pairwise independent. From this lemma, one can prove, by induction from the initial configuration, that the evolution of the P system can be simulated by the hypernet.

4.1 Example

Fig. 2, already discussed above (Section 3) as a generic hypernet, shows the fragment of the hypernet corresponding to the P system of Fig. 1. The two membranes i and j are modeled by the agents A_i (Fig. 2(a)) and A_j (Fig. 2(b)). The local place $a_j^i \in P_i$, which contains (as token) the agent A_j , reflects the fact that the membrane j is nested inside i, while the local places a^i, b^i represent the presence of molecules a and b respectively, inside the agent A_i (this is also true for a^j, b^j and the membrane j). Then $\{r_1, r_2, r_3, r_4\} \subseteq T_N$ are transitions built from the evolution rules of the membrane system. The initial hypermarking matches the initial configuration of the P system.

5 Conclusions

In this paper we have considered P systems with symport/antiport rules and we have shown how they can be modeled by a class of hierarchical Petri net systems, a generalization of hypernets [1].

The hierarchical structure of the P system is reflected by the agents's hierarchy of the hypernet, where molecules are modeled by unstructured agents (hence empty nets or simple tokens) and membranes by agents, nets which may contain in their places unstructured agents or other agents.

The exchange of molecules through a membrane of a P system, as defined by a symport or an antiport rule, corresponds to a consortium involving two active agents, that represent the two nested membranes which exchange each other passive unstructured tokens (molecules).

The main result, as given in Section 4, states a correspondence between reachable configurations of the P system and reachable hypermarkings of the related 136 L. Bernardinello et al.

hypernet. If the P system can evolves from a configuration to another one as the result of the application of a multi-rules vector, then in the hypernet exists an associated set of consortia transforming a hypermarking, corresponding to the first configuration, and another corresponding to the second one.

A translation, that takes a hypernet and returns a 1-safe Petri net (one of the basic models in Petri net theory) such that the case graphs of the latter is isomorphic to the transition system generated by the execution of consortia of the former, has been shown in [2]. This transformation proves that hypernets are well rooted in net theory. In [9] a definition of non sequential processes for hypernets was given. This can be used to derive an alternative semantics for P systems based on a purely causal dependency notion.

In the literature other works have investigated the relation between P system and Petri nets [6], [13], [5]. It is a matter of future work a deeper comparison with these approaches and with other computational models, inspired by biological membranes and derived from calculi of concurrency and mobility, as for example those proposed by Cardelli [3].

Hypernets allow movement of structured agents from one level to another one, so that the hierarchy of agents may change. In terms of P systems, this means to consider movements of membrane agents. This capability is not exploited here, however it would be interesting in future to study the modeling of P systems with active membranes [12].

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A Hybrid Approach to Modelling Biological Systems

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Summary. This paper investigates a hybrid approach to modelling molecular interactions in biology. Some computer science models are presented, namely, P systems, π -calculus and Petri nets, and two tools, Daikon, used initially in reverse-engineering, and PRISM, a probabilistic model checker. All these approaches are investigated for their complementary role in modelling which is illustrated through a simple case study.

1 Introduction

In the last decade there has been a great interest in using theoretical computer science models in biology, based on different paradigms (process algebras, cellular automata, Lindenmayer systems, Petri nets, Boolean functions, P systems, etc.) with the aim of providing an understandable, extensible and computable modelling framework while keeping the needed formalisation to perform mathematical analysis. Every such model covers certain aspects of a system and combining two or more leads to obtaining a better and more powerful modelling approach. In order to include quantitative and qualitative aspects, there have been suggested various variants of certain models with new features like: Petri nets [10, 22], stochastic π -calculus [28] and stochastic P systems [19].

In this paper we investigate the concerted use of different methods that will reveal a new vision on modelling biological systems by combining different complementary approaches. This is quite different from the hybrid approach discussed by [1] where it is shown how to switch between deterministic and stochastic behaviour.

Section 2 introduces the three modelling approaches used in the paper: P systems, pi-calculus and Petri nets, as well as Daikon tool and a simple example involving a regulatory network that will be modelled within each approach. Section 3 presents Daikon's findings and the analysis of the invariants provided. The following two sections show how PRISM and a Petri net tool are used in order to confirm some of the properties suggested by Daikon analysis. The final section summarises our findings.

2 Modelling Paradigms

In this section we will present three modelling approaches, namely P systems, π -calculus, Petri nets. A simple case study will be used to illustrate the approach. This example will be written directly into the three modelling paradigms mentioned above, P systems, π -calculus and Petri nets and will be executed with a simulator dedicated to P systems. In this way we will show how the same problem is modelled with different paradigms and consequently will be able to point out to specific characteristics of these approaches that are used for the same purpose. A system of differential equations will be also associated to this example and the results obtained will be compared to the stochastic behaviour exhibited by the P system simulator. In the next three sections we will be using a tool called Daikon to reveal certain properties of our models as they appear through data sets generated by simulators, and two tools, namely PRISM and PIPE, that are used to analyse and verify properties identified by Daikon.

The aim of this investigation is not to study the relationships between the results produced by using differential equations and those generated by P systems. This has been already considered for a special class of P systems working in a deterministic manner according to a metabolic algorithm [7]. In this study we will be using differential equations only as a substitute for real data in order to illustrate our approach that allows us to "guess" certain properties of the model and then to verify whether they hold or not as general properties or just only happens to be true for the instances generated by simulation.

Nowadays ordinary differential equations (ODE) constitute the most widely used approach in modelling molecular interaction networks in cell systems. They have been used successfully to model kinetics of conventional macroscopic chemical reactions. Nevertheless the realisation of a reaction network as a system of ODEs is based on two assumptions. First, cells are assumed to be well stirred and homogeneous volumes so that concentrations do not change with respect to space. Whether or not this is a good approximation depends on the time and space scales involved. In bacteria it has been shown that molecular diffusion is sufficiently fast to mix proteins. This is not the case in eukaryotic cells where the volume is considerably bigger and it is structured in different compartments like nucleus, mitochondria, golgi body, etc. The second basic assumption is that chemical concentrations vary continuously over time. This assumption is valid if the number of molecules of each species in the reaction volume (the cell or the subcellular compartment) are sufficiently large and the reactions are fast. A sufficiently large number of molecules is considered to be at least thousands of molecules; for hundreds or fewer molecules the continuous approach is questionable.

Writing and solving numerically a system of ODE describing a chemical reaction network can be largely automated. Each species is assigned a single variable X(t) which represents the concentration of the species at time t. Then, for each molecular species, a differential equation is written to describe its concentration change over time due to interactions with other species in the system. The rate of each reaction is represented using a kinetic rate law, which commonly depends on one or more rate constants. Exponential decay law, mass action law and Michaelis-Menten dynamic are the most widely used kinetic mechanisms. Finally in order to solve the system of ODEs we must impose a set of initial condition representing the initial concentration of the species.

Due to the limitations of ODEs to handle cellular systems with low number of molecules and spatial heterogeneity, some computational approaches have recently been proposed. In what follows we disccuss three different approaches, P systems, π -calculus and Petri nets.

2.1 P systems

Membrane computing is an emergent branch of natural computing introduced by Gh. Păun in [18]. The models defined in this context are called P systems. In the sequel we will use membrane computing and P systems with the same meaning. Roughly speaking, a P system consists of a cell-like membrane structure, in the compartments of which one places multisets of objects and strings which evolve according to given rules. Recently P systems have been used to model biological phenomena within the framework of computational systems biology presenting models of oscillatory systems [6], signal transduction [19], gene regulation control [20], quorum sensing [27] and metapopulations [21]. In this respect, P systems present a formal framework for the specification and simulation of cellular systems which integrates structural and dynamic aspects in a comprehensive and relevant way while providing the required formalisation to perform mathematical and computational analysis.

In the original approach of P systems the rules are applied in a maximally parallel way. This produces two inaccuracies: the reactions represented by the rules do not take place at the correct rate, and all time steps are equal and do not represent the time evolution of the real system. In order to solve these two problems stochastic P systems were introduced in [19].

Definition 1. A stochastic P system is a construct

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$$\Pi = (O, L, \mu, M_1, M_2, \dots, M_n, R_1, \dots, R_n)$$

where:

- O is a finite alphabet of symbols representing objects;
- *L* is a finite alphabet of symbols representing labels for the compartments;
- *μ* is a membrane structure containing n ≥ 1 membranes labelled with elements from L;
- M_i = (l_i, w_i, s_i), for each 1 ≤ i ≤ n, is the initial configuration of membrane i with l_i ∈ L, the label of this membrane, w_i ∈ O^{*} a finite multiset of objects and s_i a finite set of strings over O;
- R_i, for each 1 ≤ i ≤ n, is a finite set of rewriting rules associated with membrane i, of one of the following two forms:
 - Multiset rewriting rules:

$$obj_1 [obj_2]_l \xrightarrow{k} obj'_1 [obj'_2]_l$$

$$\tag{1}$$

with $obj_1, obj_2, obj'_1, obj'_2 \in O^*$ some finite multisets of objects and l a label from L. A multiset of objects, obj is represented as $obj = o_1 + o_2 + \cdots + o_m$ with $o_1, \ldots, o_m \in O$.

These rules are multiset rewriting rules that operate on both sides of membranes, that is, a multiset obj_1 placed outside a membrane labelled by l and a multiset obj_2 placed inside the same membrane can be simultaneously replaced with a multiset obj'_1 and a multiset obj'_2 , respectively.

• String rewriting rules:

$$[obj_1 + str_1; \dots; obj_p + str_p]_l \xrightarrow{k} [obj'_1 + str'_{1,1} + \dots str'_{1,i_1}; \dots; obj'_p + str'_{p,1} + \dots str'_{p,i_p}]_l$$
(2)

A string str is represented as follows str = $\langle s_1.s_2.\dots.s_i \rangle$ where $s_1,\dots,s_i \in O$. In this case each multiset of objects obj_j and string str_j , $1 \leq j \leq p$, are replaced by a multiset of objects obj'_j and strings $str'_{j,1}...str'_{j,i_j}$.

The stochastic constant k is used to compute the propensity of the rule by multiplying it by the number of distinct possible combinations of the objects and substrings present on the left-side of the rule with respect to the current contents of membranes involved in the rule. The propensity associated with each rule is used to compute the probability and time needed to apply it.

Cellular systems consisting of molecular interactions taking place in different locations of living cells are specified using stochastic P systems as follows. Different regions and compartments are specified using membranes. Each molecular species is represented by an object in the multiset associated with the region or compartment where the molecule is located. The multiplicity of each object represents the number of molecules of the molecular species represented by the object. Strings are used to specify the genetic information encoded in DNA and RNA. Molecular interactions, compartment translocation and gene expression are specified using rewriting rules on multisets of objects and strings - see Table 1.

In stochastic P systems [19] constants are associated with rules in order to compute their probabilities and time needed to be applied according to Gillespie algorithm. This approach is based on a Monte Carlo algorithm for stochastic simulation of molecular interactions taking place inside a single volume [8]. In contrast to this, in P systems we have a membrane structure delimiting different compartments (volumes), each one with its own set of rules (molecular interactions) and multiset of objects and strings (molecules). In this respect, a scheduling algorithm called the Multicompartmental Gillespie algorithm [19] is used so that each compartment evolves according to a different Gillespie algorithm. In this point our approach differs from other computational approaches which run a single Gillespie algorithm across the whole system without taking into account the compartmentalised cellular structure [10, 28].

Biochemistry	P System
Compartment	Region defined by a membrane
Molecule	Object
Molecular Population	Multiset of objects
Biochemical Transformation	Rewriting rule
Compartment Translocation	Boundary rule

 Table 1. Modelling Principles in P Systems

We illustrate our approach with a biomolecular system consisting in positive, negative and constitutive expression of a gene. Our model includes the specification of a gene, its transcribed RNA, the corresponding translated protein and activator and repressor molecules which bind to the gene producing an increase in transcription rate or prevent the gene from being transcribed, respectively. The bacterium where the system is located is represented using a membrane. The stochastic constants used in our model are taken from the gene control system in the lac operon in E. coli [2, 13, 14]. In this case transcription and translation have been represented using rewriting rules on multisets of objects, a more detailed description of the concurrent processes of transcription and translation using rewriting rules on strings is presented in [20]. The P systems model is formally defined in Figure 1. It consists of one single compartment labelled b, with no strings, and consequently using only multiset rewriting rules. The model refers to three distinct initial conditions, denoted by multisets $M_{0,i}$, and corresponding to constitutive expression, positive and negative regulations, respectively. Simulations of constitutive expression and positive regulation case studies are presented in Figure 2 using a tool available at [30]. A set of ordinary differential equations and their associated graphs, modelling the same examples, are provided in Figure 3. The ODE model is not used here to show its relationship to the previous P systems

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approach, but to provide a set of data that normally is taken through biological experiments. This will only be used to provide data measurements that will help identifying and validating properties of the P systems model.

 $\Pi = (\{gene, rna, protein, act, rep, act-gene, rep-gene\}, \{b\}, []_b, (b, M_i, \emptyset), \{r_1, \ldots, r_9\})$

Initial multisets: $M_{0,1} = gene; M_{0,2} = gene + act... + act and M_{0,3} = gene + rep... + rep where act and rep occur 10 times each.$ Rules: $<math>r_1: [gene]_b \stackrel{c_1}{\longrightarrow} [gene + rna]_b \quad c_1 = 0.347 \ min^{-1}$ $r_2: [rna]_b \stackrel{c_2}{\longrightarrow} [rna + protein]_b \quad c_2 = 0.174 \ min^{-1}$ $r_3: [rna]_b \stackrel{c_3}{\longrightarrow} []_b \quad c_3 = 0.347 \ min^{-1}$ $r_4: [protein]_b \stackrel{c_4}{\longrightarrow} []_b \quad c_4 = 0.0116 \ min^{-1}$ $r_5: [act + gene]_b \stackrel{c_5}{\longrightarrow} [act - gene]_b \quad c_5 = 6.6412087 \ molec^{-1}min^{-1}$ $r_6: [act - gene]_b \stackrel{c_7}{\longrightarrow} [act - gene + rna]_b \quad c_7 = 3.47 \ min^{-1}$ $r_8: [rep + gene]_b \stackrel{c_8}{\longrightarrow} [rep - gene]_b \quad c_8 = 6.6412087 \ molec^{-1}min^{-1}$

Fig. 1. P system model of gene expression.



Fig. 2. Constitutive expression and positive regulation.

2.2 π -calculus

The π -calculus approach was introduced as a formal language to describe mobile concurrent processes [17]. It is now a widely accepted model for interacting systems with dynamically evolving communication topology. The π -calculus has a $\frac{dr}{dt} = c_1 - c_3 r + c_7 \frac{act}{act + K} \quad \text{where } K \text{ is the Michaelis-Menten constant}$ $\frac{dp}{dt} = c_2 r - c_4 p$

Fig. 3. Constitutive and positive expression using ODE model.

simple semantics and a tractable algebraic theory. Starting with atomic actions and simpler processes, complex processes can be then constructed. The process expressions are defined by guarded processes, parallel composition P|Q, nondeterministic choice P + Q, replication !P, and a restriction operator $(\nu x)P$ creating a local fresh channel x for a process P. Different variants have been used to model molecular interactions [28]. A π -calculus specification of our system is provided by Figure 4. As usual for this type of modelling approach, each chemical element will be represented as a process and its definition will refer to all possible interactions of it. The initial process may be any of $S_{0,i}$. The process called *gene* defines all possible interactions of a constitutive reaction, producing messenger RNA, a positive regulation, leading to a complex denoted by *act-gene*, or negative regulation, that gets the complex *rep-gene*. This process definition corresponds in a P systems model to rules r_1 , r_5 and r_8 . In this way we can see, at least syntactically, similarities and differences between the two modelling approaches for expressing chemical interactions. More about the use of both P systems and π -calculus to model chemical interactions is provided by [26].

Biochemistry	π -calculus
Compartment	Private communication channel
Molecule	Process
Molecular Population	Systems of communicating processes
Biochemical Transformation	Communication channel
Compartment Translocation	Extrusion of a private channel's scope

Table 2. Modelling Principles in π -calculus

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 \begin{array}{l} \mbox{Initial processes: } S_{0,1} = gene; \ S_{0,2} = gene \mid act \mid \hdots \mid act \mbox{ and } S_{0,3} = gene \mid rep \mid \hdots \mid rep \\ \mbox{Processes: } \\ gene := \tau_{c_1}.(gene \mid rna ) + a_{c_5}?.act-gene + r_{c_8}?.rep-gene \\ rna := \tau_{c_2}.(rna \mid protein ) + \tau_{c_3}.0 \\ protein := \tau_{c_4}.0 \\ act := a_{c_5}!.0 \\ act-gene := \tau_{c_6}.(act \mid gene ) + \tau_{c_7}.(act-gene \mid rna ) \\ rep := r_{c_8}!.0 \\ rep-gene := \tau_{c_9}.(rep \mid gene ) \end{array}
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Fig. 4. π -calculus model of gene expression.

2.3 Petri nets

Petri nets are a mathematical and computational tool for modelling and analysis of discrete event systems typically with a concurrent behaviour. Petri nets offer a formal way to represent the structure of a discrete event system, simulate its behaviour, and prove certain properties of the system. Petri nets have applications in many fields of system engineering and computer science. Here we only recall some basic concepts of Petri nets and refer to the current literature [9, 23, 24, 25] for details regarding the theory and applications of Petri nets. In particular, we focus only on a specific class of Petri nets called place-transition nets or PT-nets, for short.

Informally, a PT-net is a directed graph formed by two kinds of nodes called places and transitions respectively. Directed edges, called arcs, connect places to transitions, and transitions to places; each arc has associated a weight. Thus, for each transition, one identifies a set of input places, the places which have at least one arc directed to that transition, and a set of output places, the places which the outgoing arcs of that transitions are directed to. Then, a non-negative integer number of tokens is assigned to each place; these numbers of tokens define the state of the PT-net also called the marking of the PT-net. In a PT-net, a transition is enabled when the number of tokens in each input place is greater than or equal to the weight of the arc connecting that place to the transition. An enabled transition can fire by consuming tokens from its input places and producing tokens in its output places; the number of tokens produced and consumed are determined by the weights of the arcs involved. The firing of a transition can be understood as the movement of tokens from some input places to some output places.

More precisely, we give the following definition.

Definition 2. A PT-net is a construct $N = (P, T, W, M_0)$ where: P is a finite set of places, T is a finite set of transitions, with $P \cap T = \emptyset$, $W : (P \times T) \cup (T \times P) \rightarrow \mathbf{N}$ is the weight function, M_0 is a multiset over P called the initial marking, and L is a location mapping.

PT-nets are usually represented by diagrams where places are drawn as circles, transitions are drawn as squares, and an arc (x, y) is added between x and y if $W(x, y) \ge 1$. These arcs are then annotated with their weight if this is 2 or more.

Given a PT-net N, the pre- and post-multiset of a transition t are respectively the multiset $pre_N(t)$ and the multiset $post_N(t)$ such that, for all $p \in P$, $|p|_{pre_N(t)} = W(p,t)$ and $|p|_{post_N(t)} = W(t,p)$. A configuration of N, which is called a marking, is any multiset over P; in particular, for every $p \in P$, $|p|_M$ represents the number of tokens present inside place p. A transition t is enabled at a marking M if the multiset $pre_N(t)$ is contained in the multiset M. An enabled transition t at marking M can fire and produce a new marking M' such that $M' = M - pre_N(t) + post_N(t)$ (i.e., for every place $p \in P$, the firing transition t consumes $|p|_{pre_N(t)}$ tokens and produces $|p|_{post_N(t)}$ tokens).

In order to reason about some basic properties, it is convenient to introduce a matrix-based representation for PT-nets. Specifically, let $N = (P, T, W, M_0)$ be a PT-net and let $\pi : P \to |P|$ and $\tau : T \to |T|$ be two bijective functions. We call place j the place p with $\pi(p) = j$, and we call transition i the transition t with $\tau(t) = i$. Then, a marking M is represented as a |P| vector which contains in each position j the number of tokens currently present inside place j. The *incidence matrix* of N is the $|T| \times |P|$ matrix A such that, for every element a_{ij} of A, $a_{ij} = |\pi^{-1}(j)|_{post_N(\tau^{-1}(i))}| - |\pi^{-1}(j)|_{pre_N(\tau^{-1}(i))}|$ (i.e., a_{ij} denotes the change in the number of tokens in place j due to the firing of transition i). A control vector u is a |T| vector containing 1 in position i to denote the firing of transition i, 0 otherwise. Thus, if a particular marking M_n is reached from the initial marking M_0 through a firing sequence u_1, u_2, \ldots, u_n of enabled transitions, we obtain

$$M_n = M_0 + A^T \cdot \sum_{k=1}^n u_k$$

which represents the reachable-marking equation.

The aforementioned representation of a PT-net N allows us to introduce the notions of P-invariants and T-invariants. P-invariants are the positive solutions of the equation $A \cdot y = 0$; the non-zero entries of a solution y represents the set of places whose total number of tokens does not change with any firing sequence from M_0 . T-invariants instead are the positive solutions of the equation $A^T \cdot x = 0$; a solution vector x represents the set of transitions which have to fire from some marking M to return to the same marking M. Then, a PT-net is said to be *bounded* if there exists a |P| vector B such that, for all marking M reachable from M_0 , we have $M \leq B$; a PT-net is said to be *alive* if, for all marking M.

As pointed out in [10, 22], a PT-net model for a system of molecular interactions can be obtained by representing each molecular species as a different place and each biochemical transformation as a different transition. Tokens inside a place can then be used to indicate the presence of a molecule in certain proportions. This modelling approach is summarised in Table 3. Thus, a biochemical system is represented as a discrete event system whose structural properties are useful for

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Biochemistry	PT-net
Molecule	Place
Molecular Population	Marking
Biochemical Transformation	Transition
Reactant	Input Place
Product	Output Place

 Table 3. Modelling Principles in PT-nets.

drawing conclusions about the behaviour and structure of the original biochemical system [22]. For instance, P-invariants determine the set of molecules whose total net concentrantions remain unchanged during the application of certain biochemical transformations; T-invariants instead indicate the presence of cyclic reactions which lead to a condition where some reactions are in a state of continuous operation. Also, the property of liveness is useful to determine the absence of metabolic blocks which may hinder the progress of the biochemical system.

Finally, we recall that it was shown in [4, 15, 16] how to transform a P system into a corresponding PT-net. This is done by considering a transition for each rule in the P system that has the left-hand side of the rule as pre-multiset and the right-hand side of the rule as post-multiset. In particular, in order to model the localisation of rules and objects inside the membranes, one considers in the corresponding PT-net a distinct place for each object possibly present inside a membrane. Thus, the transformation of objects inside the membranes and the communication of objects between membranes is mapped into the movement of tokens between places of a PT-net. This translation is briefly illustrated in Table 4. Thus, we have a direct way for obtaining a PT-net representation of a given P

P System	PT-net
Object a inside membrane i	Place a_i
Multiplicity of an object	Number of tokens inside a place
Rule	Transition
Left-hand side of a rule	Input places
Right-hand side of a rule	Output places

Table 4. Translation of a P system into a PT-net.

system model that offers us the possibility of analysing the P system model in terms of PT-net properties. We illustrate this approach by showing in Figure 5 the PT-net translation of the P system model of Figure 1.

Clearly, we have that transition r_i corresponds to rule r_i , $1 \le i \le 9$. For the PT-net of Figure 5, if we set M_0 as initial marking, where M_0 contains one token in the place *gene*, then we have only consitutive expression; if we set M_0 as having one token in *gene* and n in *act* with $n \ge 1$, then we have positive regulation; if we



Fig. 5. PT-net representation of gene positive and negative regulation.

set M_0 with one token in gene and m in rep with $m \ge 1$, then we have negative regulation. The incidence matrix for PT-net of Figure 5 is reported in Appendix 1 together with its P-invariants and T-invariants. The relevance of these invariants with respect to this specific case study is discussed in Section 5. These are obtained by using PIPE [29], a freely available Petri net tool. As well as this, PIPE allows us to check for the properties of boundedness and liveness (i.e., absence of deadlock).

2.4 Daikon tool

Daikon [5] is a tool that was initially developed to reverse-engineer specifications from software systems. The specifications are in terms of invariants, which are rules that must hold true at particular points as the program executes. To detect invariants the program is executed multiple times and the values of the variables are recorded at specific points (e.g., the start and end of a program function). Daikon infers the invariants by attempting to fit sets of predefined rules to the values of program variables at every recorded program point. Usually the most valuable invariants are preconditions, postconditions and system invariants. These specify the conditions that must hold between variables before a function is executed, after a function has finished executing, and throughout the program execution respectively. As a trivial example, a precondition for the function div(a, b) that divides a by b would be $b \ge 0$. Daikon provides about 70 predefined invariants [5], such as x > y, a < x < b, y = ax + b, and can also be extended to check for new user-supplied invariants.

The idea of using a set of executions to infer rules that govern system behaviour, as espoused by Daikon, is particularly useful in the context of biological models. The ability to automatically infer invariants from model simulations is useful for the following reasons: (1) obvious invariants will confirm that the model is behaving as it should, (2) anomalous invariants can indicate a fault in the model and its parameter values or (3) could even suggest novel, latent relationships between model variables.

3 Finding functional relationships in raw (wet real) data: Data analysis using Daikon

This section demonstrates the use of Daikon to discover relationships between variables in the output from P system simulations of the gene regulation model. The aim is to identify invariants that govern model behaviour for negative, constitutive and positive gene regulation. Here we select a sample of the generated invariants and show how they relate to the high-level functionality of the system, and how they can be of use for further model analysis. Invariants are only useful if they are representative of a broad range of model behaviour. A single simulation can usually not be considered to be representative, especially if the model is non-trivial and contains stochastic behaviour. For this analysis the model was simulated 30 times, ten times for negative, constitutive and positive regulation respectively.

Using Daikon to generate invariants from simulation output is relatively straightforward. It takes as input two files, one of which declares the types of invariants that are of interest, along with the set of relevant variables for each type of invariant. The other file contains the variable values from model simulations, and lists them under their respective declared invariants. In our case the output is in the form of a linear time series, where variable values are provided for t = 0...n time points in the simulation. To analyse this with Daikon, the declaration file contains the three invariant types described above (preconditions, postconditions and system invariants), along with the key model variables under each type (gene, act-gene, rep-gene, rna, prot, rep, act). The data trace file maps any variable values at t = 0 to the preconditions, the relationship between every pair of variables t and t-1 to the postconditions, and the variable values for every value of t to the system invariants. To guarantee accurate invariants, the data trace has to be constructed from a set of simulations that can be deemed to be sufficiently representative of the system's behaviour.

Figure 6 contains a sample of the invariants that were discovered. These provide a number of insights into the behaviour of the system that would be difficult to

	Positive	Negative	Constitutive
Pre-conditions	gene = 1 rna = prot = rep-gene = act-gene = 0 act = 10	gene = 1 rna = prot = rep-gene = act = act-gene = 0 rep = 10	gene = 1 rna = prot = rep-gene = act-gene = act = 0
Post-condtions	$\begin{array}{ll} (prot = 0) & \rightarrow \\ (orig(prot) = 0) \\ (rna = 0) \rightarrow (gene = 0) \\ (orig(rna) = 0) \rightarrow (gene = 0) \\ gene \leq rna \\ (prot = 0) \rightarrow (gene = 0) \\ (rna = 0) \rightarrow (orig(act-gene = 0)) \end{array}$	act = orig(act) = orig(act-gene) rna < orig(rep) rep > orig(prot)	$\begin{array}{ll} (prot &= & 0) & \rightarrow \\ (orig(prot) &= & 0) \\ gene &= & orig(gene) \\ rep &= & orig(rep) \end{array}$
Invariants	$gene = one of \{0, 1\}$ $rep = rep-gene$ $= 0$ $0 \le rna \le 24$ $0 \le prot \le 205$ $act = one of \{9, 10\}$ $act-gene = one of \{0, 1\}$ $(gene \land act-gene) = 0$ $(rna = 0) \rightarrow (prot = 0)$	$gene = one of \{0, 1\}$ $act = act-gene$ $= 0$ $rna = one of \{0, 1\}$ $rep-gene = one of \{0, 1\}$ $prot = one of \{0, 1, 2, 3\}$ $(gene \land rep-gene) = 0$ $rna < rep$ $rep > prot$	$\begin{array}{l} rep = rep-gene \\ = act \\ = act-gene \\ = 0 \\ gene = 1 \\ 0 \leq rna \leq 7 \\ 0 \leq prot \leq 32 \\ rna \geq rep \end{array}$

Fig. 6. Invariants discovered by Daikon

ascertain from passively observing the simulations. Here we provide an overview of some of these results.

The preconditions show precisely which model parameters are altered for the positive, negative and constitutive sets of simulations. For all simulations, *gene* starts off as active, and all other variables are zero, apart from the activators variable *act* for positive and the repressors variable *rep* for negative regulation.

The postconditions provide a number of insights into the dynamics of the model because they summarise the rules that govern the change in variable values for every single time-step. For positive regulation we learn that the number of proteins will never decrease back to zero throughout the simulation (protein can only be zero if it was already zero at the previous value of t), and that the gene must become active to produce rna, which can only happen when *act-gene* becomes 1. For negative regulation it shows that the amount of activators remain constantly zero. For constitutive regulation, similarly to positive regulation, the number of proteins can never decrease to zero.

The invariants are rules that hold throughout the entire simulation. These usually cover the range of values a variable can hold, e.g., *gene* can either be on or off for positive or negative regulation, but is constantly on for constitutive regulation, or the number of proteins is always between zero and 205 for positive regulation. It also points out rules that can sometimes be fundamental to the behaviour of the model. For example, in positive regulation *act-gene* and *gene* can never be on at the same time, which makes sense because *act-gene* is responsible for activating the *gene* when it is not active. The same holds for *rep-gene* and *gene* in negative regulation. In positive regulation it also points out that, without any *rna*, there can be no proteins.

These rules provide a number of useful insights into the behaviour of the model, many of which are expected, but some of which may either be anomalous or might identify relationships that had not been previously considered. As an example, the preconditions, which simply summarise the input parameters, are obviously expected, but in practice identified that a small number of our experimental simulations had been mistakenly executed with the wrong parameter values (the precondition for positive regulations stated $act = one of \{9, 10, 19, 20\}$ instead of just 9 and 10). Rules such as $rna \ge rep$ in constitutive regulation and rep > prot in positive regulation are obviously statistically justified by the simulations, but had not been considered explicitly. New rules like these are useful seeds for further experimentation and analysis, and the following section will show how we have investigated these novel properties with the PRISM model checker.

4 PRISM analysis of the system

Most research in systems biology focuses on the development of models of different biological systems in order to be able to simulate them, accurately enough such as to be able to reveal new properties that can be difficult or impossible to discover through direct experiments. One key question is what one can do with a model, other than just simulate trajectories. This question has been considered in detail for deterministic models, but less for stochastic models. Stochastic systems defy conventional intuition and consequently are harder to conceive. The field is widely open for theoretical advances that help us to reason about systems in greater detail and with finer precision. An attempt in this direction consists of using model checking tools to analyse in an automatic way various properties of the model. Probabilistic model checking is a formal verification technique. It is based on the construction of a precise mathematical model of a system which is to be analysed. Properties of this system are then expressed formally using temporal logic and analysed against the constructed model by a probabilistic model checker. Our current attempt uses a probabilistic symbolic model checking approach based on PRISM (Probabilistic and Symbolic Model Checker) [11, 12]. PRISM supports three different types of probabilistic models, discrete time Markov chains (DTMC), Markov decision processes (MDP) and continuous time Markov chains (CTMC). PRISM supports systems specifications through two temporal logics, PCTL (probabilistic computation tree logic) for DTMC and MDP and CSL (continuous stochastic logic) for CTMC.

In order to construct and analyse a model with PRISM, it must be specified in the PRISM language, a simple, high level, state-based language. The fundamental components of the PRISM language are modules, variables and commands. A model is composed of a number of modules which can interact with each other. A module contains a number of local variables and commands.

The values of these variables at any given time constitute the states of the module. The space of reachable states is computed using the range of each variable and its initial value. The global state of the whole model is determined by the local state of all modules.

The behaviour of each module is described by a set of commands. A command takes the form:

 $[\quad \texttt{action} \quad] \quad \texttt{g} \rightarrow \lambda_{1}: \mathbf{u_{1}} + \dots + \lambda_{n}: \mathbf{u_{n}};$

The guard g is a predicate over all the variables of the model. Each update $\mathbf{u_i}$ describes the new values of the variables in the module specifying a transition of the module. The expressions λ_i are used to assign probabilistic information, rates, to transitions.

The label action placed inside the square brackets are used to synchronise the application of different commands in different modules. This forces two or more modules to make transitions simultaneously. The rate of this transition is equal to the product of the individual rates, since the processes are assumed to be independent.

The main components of a P system are a membrane structure consisting of a number of membranes that can interact with each other, an alphabet of objects and a set of rules associated to each membrane. These components can easily be mapped into the components of the PRISM language using modules to represent membranes, variables to describe the alphabet and commands to specify the rules.

A PRISM specification of our system is provided in Appendix 2. Appendix 3 shows the probability that some molecules concentrations will reach certain values at steady state. The ranges of values provided by Daikon represent an indication of possible levels for various molecular concentrations, but in order to know the likely values around steady states, PRISM provides a set of properties that help in this respect. For example, for positive regulation, Daikon provides the range 0 to 24, for *rna* molecules, but PRISM shows that values between 0 and 15 are more likely to be obtained than values greater than 15, and values over 20 are very unlikely to be reached. These values are also confirmed by the graphs provided by differential equations and P system simulator.

Other properties, suggested by Daikon analysis, like rna < rep, prot < rep, are also validated by PRISM by showing they take place with a higher probability for values of rep less than 5 - see Appendix 4. The average or expected behaviour of the stochastic system is also provided and this is very close to ODE behaviour.

5 Petri net analysis of the system

In this section we will show how different invariants will emerge from the analysis of the Petri net and how Daikon hypotheses are formally verified or new problems are formulated.

T-invariants in Appendix 1 show that:

- If we fire transition r1 and then transition r3, the current marking of the newtork remains unchanged because we first produce a molecule of rna and then we consume it; the same happens if we first fire transition r7 and than transition r3, or if we first fire transition r2 and then transition r4 (i.e., we first produce a molecule of *protein* and then we consume it);
- The operation of binding the activator to the gene and its de-binding are one the reverse of the other, hence firing transition r5 followed by transition r6 (or vice versa) has no effect on the current marking; these two transitions consitute a continuous loop;
- The operation of binding the repressor to the gene and its de-binding are one the reverse of the other, hence firing transition r8 followed by transition r9 (or vice versa) has no effect on the current marking; these two transitions consitute a continuous loop.

P-invariants computed by PIPE, in Appendix 1, for some initial marking with one element in gene, n in act and m in rep, where $n, m \ge 0$ show that:

- the gene is always present and it can assume three different states: gene, actgene, and rep-gene; these three states are mutually exclusive; in the case of constitutive expression (i.e, n = m = 0), we have M(gene) = 1 indicating that the gene is always present - this confirms Daikon invariant gene = 1; in the case of positive regulation (i.e., $n \ge 1$ and m = 0), we have M(gene) + M(actgene) = 1 indicating two mutually exclusive states - this confirms Daikon invariant gene $\land act-gene = 0$; in the case of negative regulation (i.e., $m \ge 1$ and n = 0, we have M(gene) + M(rep-gene) = 1 indicating two mutually exclusive states - this confirms Daikon invariant gene $\land rep-gene = 0$;
- for positive regulation, the number of activator molecules cannot be increased but can be decreased only by 1 similar invariant is found by Daikon;
- for negative regulation, the number of repressor molecules cannot be increased but can be decreased only by 1 similar invariant is found by Daikon.

PIPE also shows that the network is not bounded but it is alive. In fact, gene is always present and we can keep firing transition r1 to increase indefinitely the amount of rna. The liveness here comes from the above invariant and shows that the system will be working forever. The boundedness instead produces a result that apparently contradicts PRISM findings, where the probability that the number of rna's is greater than 7 is almost 0! This comes from the fact that PIPE uses a non-deterministic system instead of a probabilistic one considered by PRISM and P system simulator. It will be interesting to check this property with a probabilistic Petri net tool.

6 Conclusions

In this paper we have investigated the concerted use of different methods, and shown how these can provide complementary insights into different facets of biological system behaviour. Individual modelling techniques have their own respective benefits and usually excel at reasoning about a system from a particular perspective. This paper shows how these benefits can be leveraged by using different modelling techniques in concert.

As a case study, we have constructed a P system model of a small gene expression system and produced equivalent specifications using Petri net and π -calculus approaches. Simulations of the P system model were analysed by Daikon (to identify potential rules that govern model output), and some of the most interesting suggested rules were checked using the PRISM probabilistic model checker. The Petri net model was analysed with PIPE, a general Petri net analysis tool. The results show how analysis results from different models of the same system are useful for the purposes of both validating and improving each other.

The gene expression model was chosen because it is manageable, and thus forms a useful basis for a case study to compare different modelling techniques. Our future work will apply the techniques shown in this paper to a larger and more realistic case study. This should provide further insights into the benefits that arise in modelling increasingly complex systems when the modeller is increasingly reliant upon the use of various automated tools to study the model behaviour.

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APPENDIX 1

Transpose of the incidence matrix for PT-net of Figure 5:

	r1	r2	r4	r3	r5	r7	r6	r8	r9
gene	0	0	0	0	-1	0	1	-1	1
rna	1	0	0	-1	0	1	0	0	0
protein	0	1	-1	0	0	0	0	0	0
act	0	0	0	0	-1	0	1	0	0
act-gene	9 0	0	0	0	1	0	-1	0	0
rep	0	0	0	0	0	0	0	-1	1
rep-gene	e 0	0	0	0	0	0	0	1	-1

which shows the variations on the number of tokens determined by each transition. T-invariants obtained in PIPE:

r1	1	0	0	0	0
r2	0	1	0	0	0
r4	0	1	0	0	0
r3	1	0	0	1	0
r5	0	0	1	0	0
r7	0	0	0	1	0
r6	0	0	1	0	0
r8	0	0	0	0	1
r9	0	0	0	0	1

P-invariants computed by PIPE for some initial marking contains one token in gene n in act and m in rep, with $n, m \ge 0$:

gene	1	0	0	
rna	0	0	0	M(gene) + M(act-gene) + M(rep-gene) = 1
protein	0	0	0	M(act) + M(act-gene) = n
act	0	1	0	M(rep) + M(rep-gene) = m
act-gene	1	1	0	
rep	0	0	1	
rep-gene	1	0	1	

APPENDIX 3

Ranges of molecules

P = ? [true U <= T rna > bound]
P = ? [true U <= T protein > bound]

APPENDIX 2

```
// Gene expression control
 // Model is stochastic
 stochastic
 // Bounds to the number of molecules
 const int rna_bound;
 const int protein_bound;
 const int number_activators;
const int number_repressors;
 const int initact:
 const int initrep;
// Stochastics constants associated with each command/rule/molecular interaction
// Stochastics constants associated with each command/ule/molecular inter:

const double c1 = 0.347; // [gene ]_b -c1> [gene + rna ]_b

const double c2 = 0.174; // [rna ]_b -c2> [rna + protein ]_b \\

const double c3 = 0.347; // [rna ]_b -c3> [ ]_b

const double c3 = 0.347; // [protein ]_b -c4> [ ]_b

const double c5 = 0.6412087; // [act + gene ]_b -c6> [act gene ]_b

const double c6 = 0.6; // [actgene ]_b -c6> [act + gene ]_b

const double c7 = 3.47; // [actgene ]_b -c6>> [actgene ]_b

const double c7 = c3.47; // [actgene ]_b -c6> [cat + gene ]_b

const double c7 = c3.47; // [actgene ]_b -c7> [actgene + rna ]_b
const double c8 = 6.6412087; // [rep + gene ]_b -c8-> [repgene ]_b
const double c9 = 0.6; // [repgene ]\_b -c9-> [rep + gene ]_b
 // Module representing a bacterium
 module bacterium
        gene : [ 0 .. 1 ] init 1;
       actgene : [ 0 .. 1 ] init 0;
repgene : [ 0 .. 1 ] init 0;
        act : [ 0 .. 1 ] init initact;
       rep : [ 0 .. 1 ] init initrep;
rna : [ 0 .. rna_bound ] init 0;
       ina . [0 ... protein_bound ] init 0;
// [gene ]_b -cl-> [gene + rna ]_b
[] gene = 1 & rna < rna_bound -> cl : (rna' = rna + 1);
// [rna ]_b -c2-> [ rna + protein ]_b
        [] rna > 0 & protein < protein_bound -> c2*rna : (protein' = protein + 1);
        // [ rna ]_b -c3-> [ ]_b
[ ] rna > 0 -> c3*rna : (rna' = rna - 1);
// [ protein ]_b -c4-> [ ]_b
        [] protein > 0 -> c4*protein : (protein' = protein - 1);
        // [ act + gene ] _b -c5-> [ actgene ] _b
[ ] act = 1 & gene = 1 -> c5*number_activators : (gene' = 0) & (act' = 0) & (actgene' = 1);
        // [ actgene ]_b -c6-> [ act + gene ]_b
        [ ] actgene = 1 & act = 0 -> c6 : (actgene' = 0) & (act' = 1) & (gene' = 1);
// [ actgene ]_b -c7-> [ actgene + rna ]_b
        [] actgene = 1 & rna < rna_bound -> c7 : (rna' = rna + 1);
        [] degene = 1 & rind < find question = 1 & (find = 1 m + 1),
// [rep + gene ]_b - c3-> [repgene ]_b
[] rep = 1 & gene = 1 -> c3*number_repressors : (gene' = 0) & (rep' = 0) & (repgene' = 1);
// [repgene ]_b - c3-> [rep + gene ]_b
[] repgene = 1 & rep = 0 -> c9 : (repgene' = 0) & (rep' = 1) & (gene' = 1);
```

Constitutive regulation

endmodule

```
rna <= 7
rna >= 0
prot <= 32
prot >= 0
```



Positive regulation

rna <= 24

rna >= 0

prot <= 205

prot >= 0



Negative regulation

rna one of $\{0, 1\}$

prot one of { 0, 1, 2, 3 }





APPENDIX 4

Relationship between the number of repressors and rna and protein molecules.









 $\mathbf{R}= ?~[~\mathbf{I}=\mathbf{T}~]$





Information Theory over Multisets^{*}

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Summary. Starting from Shannon theory of information, we present the case of producing information in the form of multisets, and encoding information using multisets. We rewiew the entropy rate of a multiset information source and we derive a formula for the information content of a multiset. We then study the encoder and channel part of the system, obtaining some results about multiset encoding length and channel capacity.

1 Motivation

The attempt to study information sources which produce multisets instead of strings, and ways to encode information on multisets rather than strings, originates in observing new computational models like membrane systems which employ multisets [5]. Membrane systems have been studied extensively and there are plenty of results regarding their computing power, language hierarchies and complexity. However, while any researcher working with membrane systems (called also P systems) would agree that P systems process information, and that living cells and organisms do this too, we are unaware of any attempt to precisely describe natural ways to encode information on multisets or to study sources of information which produce multisets instead of strings. One could argue that, while some of the information in a living organism is encoded in a sequential manner, like in DNA for example, there might be important molecular information sources which involve multisets (of molecules) in a non-trivial way.

A simple question: given a P system with one membrane and, say, 2 objects a and 3 objects b from a known vocabulary V (suppose there are no evolution rules), how much information is present in that system? Also, many examples of P systems perform various computational tasks. Authors of such systems encode the input (usually numbers) in various ways, some by superimposing a string-like structure on the membrane system [1], some by using the natural encoding or unary numeral system, that is, the natural number n is represented with n objects, for example,

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 a^n . However, just imagine a gland which uses the bloodstream to send molecules to some tissue which, in turn, sends back some other molecules. There is for sure an energy and information exchange. How to describe it? Another, more general way to pose that question is: what are the natural ways to encode numbers, and more generally, information on multisets, and how to measure the encoded information?

If membrane systems, living cells and any other (abstract or concrete) multiset processing machines are understood as information processing machines, then we believe that such questions should be investigated. According to our knowledge, this is the first attempt of such an investigation. We start from the idea that a study of multiset information theory might produce interesting, useful results at least in systems biology; if we understand the *natural* ways to encode information on multisets, there is a chance that *Nature* might be using similar mechanisms.

Another way in which this investigation seems interesting to us is that there is more challenge in efficiently encoding information on multisets, because they constitute a poorer encoding media compared to strings. Encoding information on strings or even richer, more organized and complex structures are obviously possible and have been studied. Removing the symbol order, or their position in the representation as strings can lead to multisets carrying a certain penalty, which deserves a precise description. Order or position do *not* represent essential aspects for information encoding; symbol multiplicity, a native quality of multisets, is *enough* for many valid purposes. We focus mainly on such "natural" approaches to information encoding over multisets, and present some advantages they have over approaches that superimpose a string structure on the multiset. Then we encode information using multisets in a similar way as it is done using strings.

There is also a connection between this work and the theory of numeral systems. The study of number encodings using multisets can be seen as a study of a class of purely non-positional numeral systems.

2 Entropy rate of an Information Source

Shannon's information theory represents one of the great intellectual achievements of the twentieth century. Information theory has had an important and significant influence on probability theory and ergodic theory, and Shannon's mathematics is a considerable and profound contribution to pure mathematics.

Shannon's important contribution comes from the invention of the sourceencoder-channel-decoder-destination model, and from the elegant and general solution of the fundamental problems which he was able to pose in terms of this model. Shannon has provided significant demonstration of the power of coding with delay in a communication system, the separation of the source and channel coding problems, and he has established the fundamental natural limits on communication. As time goes on, the information theoretic concepts introduced by Shannon become more relevant to day-to-day more complex process of communication.

2.1 Short Review of Shannon Information Theory

We use the notions defined in the classical paper [6] where Shannon has formulated a general model of a communication system which is tractable to a mathematical treatment.

Consider an information source modelled by a discrete Markov process. For each possible state *i* of the source there is a set of probabilities $p_i(j)$ associated to the transitions to state *j*. Each state transition produces a symbol corresponding to the destination state, e.g. if there is a transition from state *i* to state *j*, the symbol x_j is produced. Each symbol x_i has an initial probability $p_{i\in\overline{1..n}}$ corresponding to the transition probability from the initial state to each state *i*.

We can also view this as a random variable X with x_i as events with probabilities $p_i, X = \begin{pmatrix} x_1 & x_2 & \cdots & x_n \\ p_1 & p_2 & \cdots & p_n \end{pmatrix}$.

There is an entropy H_i for each state. The entropy rate of the source is defined as the average of these H_i weighted in accordance with the probability P_i of occurrence of the states:

$$H(X) = \sum_{i} P_{i}H_{i} = -\sum_{i,j} P_{i}p_{i}(j)\log p_{i}(j)$$
(1)

Suppose there are two symbols x_i, x_j and p(i, j) is the probability of the successive occurrence of x_i and then x_j . The entropy of the joint event is

$$H(i,j) = -\sum_{i,j} p(i,j) \log p(i,j)$$

The probability of symbol x_j to appear after the symbol x_i is the conditional probability $p_i(j)$.

Remark 1. The quantity H is a reasonable measure of choice or information.

String Entropy

Consider an information source X which produces sequences of symbols selected from a set of n independent symbols x_i with probabilities p_i . The entropy formula for such a source is given in [6]:

$$H(X) = \sum_{i=1}^{n} p_i log_b \frac{1}{p_i}$$

2.2 Multiset Entropy

We consider a discrete information source which produces multiset messages (as opposed to string messages). A message is a multiset of symbols, and a multiset is a string equivalence class. The entropy rate of such a source is proved to be zero in [7]:

$$H(X_{multiset}) = \lim_{n \to \infty} \frac{1}{n} H(\{X_i\}_{i=1}^n) = 0$$

Information content

The information content of an outcome (multiset) x is $h(x) = \log \frac{1}{P(x)}$. See [4].

Let be $k \in \mathbb{N}$ and $X = \begin{pmatrix} x_1 & x_2 \dots & x_n \\ p_1 & p_2 \dots & p_n \end{pmatrix}$ a random variable and $x = x_1^{m_1} x_2^{m_2} \dots x_n^{m_n}$ a multiset over symbols from X, with $\sum_{i=1}^n m_i = k$, then the probability of the outcome x is given by the multinomial distribution

 $\binom{k}{m_1, m_2, \dots, m_n} \prod_{i=1}^n p_i^{m_i}$:

$$P[x = (m_1, m_2, \dots, m_n)] = \frac{(\sum_{i=1}^n m_i)!}{\prod_{i=1}^n m_i!} \prod_{i=1}^n p_i^{m_i}$$

So, the information content of the multiset x is:

$$\begin{aligned} h(x = x_1^{m_1} x_2^{m_2} \dots x_n^{m_n}) &= \log \frac{1}{P[x]} = \log \left(1 / \frac{(\sum_{i=1}^n m_i)!}{\prod_{i=1}^n m_i!} \prod_{i=1}^n p_i^{m_i} \right) = \\ &= \log \frac{\prod_{i=1}^n m_i!}{(\sum_{i=1}^n m_i)! \prod_{i=1}^n p_i^{m_i}} \end{aligned}$$

3 Multiset Encoding and Channel Capacity

After exploring the characteristics of a multiset generating information source, we move to the channel part of the communication system. Properties of previously developed multiset encodings are analyzed in [2, 3]. The capacity of multiset communication channel is derived based on Shannon's definition and also on the capacity theorem. Please note that one can have a multiset information source and a usual sequence-based encoder and channel. All the following combinations are possible:

3.1 String Encoding

We shortly review the results concerning the string encoding.

Source/Encoder	Sequential	Multiset
Sequential	[6]	this paper
Multiset	this paper	this paper

Table 1	1.	Source/	Encoder	types
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Encoding Length

We have a set of symbols X to be encoded, and an alphabet A. We consider the uniform encoding. Considering the length l of the encoding, then $X = \{x_i = a_1 a_2 \dots a_l | a_j \in A\}$.

If $p_i = P(x_i) = \frac{1}{n}$, then we have

$$H(X) = \sum_{i=1}^{n} \frac{1}{n} log_b(n) = log_b(n) \le l$$

It follows that $n \leq b^l$. For $n \in \mathbb{N}$, $n - b^x = 0$ implies $x_0 = \log_b n$ and so $l = \lceil x_0 \rceil = \lceil \log_b n \rceil$.

Channel Capacity

Definition 1. [6] The capacity C of a discrete channel is given by

$$C = \lim_{T \to \infty} \frac{\log N(T)}{T}$$

where N(T) is the number of allowed signals of duration T.

Theorem 1. [6] Let $b_{ij}^{(s)}$ be the duration of the sth symbol which is allowable in state i and leads to state j. Then the channel capacity C is equal to $\log W$ where W is the largest real root of the determinant equation:

$$\left|\sum_{s} W^{-b_{ij}^{(s)}} - \delta_{ij}\right| = 0$$

where $\delta_{ij} = 1$ if i = j, and zero otherwise.

3.2 Multiset Encoding

We present some results related to the multiset encoding.
Encoding Length

We consider a set X of N symbols, an alphabet A, and the length of encoding l, therefore:

 $X = \{x_i = a_1^{n_1} a_2^{n_2} \dots a_b^{n_b} \mid \sum_{j=1}^b n_j = l, \, a_j \in A\}$

Proposition 1. Non-uniform encodings over multisets are shorter than uniform encodings over multisets.

Proof. Over multisets we have

1. for an uniform (all the encoding representation have the same length l) encoding: $N \leq N(b,l) = \left\langle \begin{array}{c} b \\ l \end{array} \right\rangle = \left(\begin{array}{c} b+l-1 \\ l \end{array} \right) = \frac{(b+l-1)!}{l!(b-1)!} = \frac{\prod_{i=1}^{b-1}(l+i)}{(b-1)!}$. If x_0 is the real root of $n - \frac{\prod_{i=1}^{b-1}(x+i)}{(b-1)!} = 0$ then $l = \lceil x_0 \rceil$.

2. for non-uniform encoding: $N \le N(b+1, l-1) = \left\langle \begin{array}{c} b+1\\ l-1 \end{array} \right\rangle = \left(\begin{array}{c} b+l-1\\ l-1 \end{array} \right) = \left(\begin{array}{c} \frac{(b+l-1)!}{l-1} \\ \frac{(b+l-1)!b!}{l-1} \\ \frac{\prod_{i=0}^{b-1}(l+i)}{b!} \\ \frac{\prod_{i=0}^{b-1}(l+i)}{(b-1)!} \\ \frac{\prod_{i=0}^{b-1}(x+i)}{(b-1)!} \\ \frac{(b+l)!}{(b-1)!} \\ \frac{(b+l)!}{(b+1)!} \\ \frac{(b+l)$

From $n - N(b, x_0) = 0$ and $n - \frac{x'_0}{b}N(b, x'_0) = 0$ we get $N(b, x_0) = \frac{x'_0}{b}N(b, x'_0)$. In order to prove $l > l' \iff x_0 > x'_0$, let suppose that $x_0 \le x'_0$. We have $x'_0 > b$ (for sufficiently large numbers), and this implies that $N(b, x_0) \le N(b, x'_0) < \frac{x'_0}{b}N(b, x'_0)$. Since this is false, it follows that $x_0 > x'_0$ implies $l \ge l'$.

Channel Capacity

We consider that a sequence of multisets is transmitted along the channel. The capacity of such a channel is computed for base 4, then some properties of it for any base are presented.

Multiset channel capacity in base 4

In Figure 1 we have a graph G(V, E) with 4 vertices $V = \{S_1, S_2, S_3, S_4\}$ and $E = \{(i, j) \mid i, j = \overline{1..4}, i \leq j\} \cup \{(i, j) \mid i = 4, j = \overline{1..3}\}$

In Theorem 1 we get $b_{ij}^{(a_k)} = t_k$ because we consider that the duration to produce a_k is the same for each $(i, j) \in E$. The determinant equation is

$$\begin{vmatrix} W^{-t_1} - 1 & W^{-t_2} & W^{-t_3} & W^{-t_4} \\ 0 & W^{-t_2} - 1 & W^{-t_3} & W^{-t_4} \\ 0 & 0 & W^{-t_3} - 1 & W^{-t_4} \\ 0 & 0 & 0 & W^{-t_4} - 1 \end{vmatrix} = 0$$

If we consider $t_k = t$, then the equation becomes $\left(1 - \frac{1}{W^t}\right)^4 = 0$, and $W_{real} = 1$. Therefore $C = \log_4 1 = 0$.



Fig. 1. Multiset channel capacity

Multiset channel capacity in base b

Theorem 2. The multiset channel capacity is zero, C = 0.

Proof. First approach

The first method for computing the capacity is using the definition from [6].

$$\begin{split} C &= \lim_{T \to \infty} \frac{\log N(T)}{T} = \lim_{T \to \infty} \frac{\log N(b,T)}{T} = \\ &= \lim_{T \to \infty} \frac{\log \left\langle \begin{array}{c} b \\ T \end{array} \right\rangle}{T} = \lim_{T \to \infty} \frac{1}{T} \log \frac{(b+T-1)!}{T!(b-1)!} \end{split}$$

Using Stirling's approximation $\log n! \approx n \log n - n$ we obtain

$$C = \lim_{T \to \infty} \frac{1}{T} \left(\log(b + T - 1)! + \log T! + \log(b - 1)! \right) =$$

=
$$\lim_{T \to \infty} \frac{1}{T} \left((b + T - 1) \log(b + T - 1) - T \log T - (b - 1) \log(b - 1) \right) =$$

=
$$\lim_{T \to \infty} \frac{b - 1}{T} \log \left(1 + \frac{T}{b - 1} \right) + \lim_{T \to \infty} \log \left(1 + \frac{b - 1}{T} \right) = 0$$

Second approach

Using 1, the determinant equation for a multiset encoder is:

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$$\begin{vmatrix} W^{-t_1} - 1 & W^{-t_2} & W^{-t_3} & \cdots & W^{-t_b} \\ 0 & W^{-t_2} - 1 & W^{-t_3} & \cdots & W^{-t_b} \\ 0 & 0 & W^{-t_3} - 1 & \cdots & W^{-t_b} \\ \vdots & \vdots & \vdots & \vdots & \vdots \\ 0 & \cdots & 0 & W^{-t_{b-1}} - 1 & W^{-t_b} \\ 0 & 0 & 0 & \cdots & W^{-t_b} - 1 \end{vmatrix} = 0$$

Proposition 2. If $t_k = t$, then the determinant equation becomes

$$\left(1 - \frac{1}{W^t}\right)^b = 0. \tag{2}$$

The capacity C is given by $C = \log_b W$, where W is the largest real root of the equation (2). Considering $x = W^{-t}$, then we have

$$W = \frac{1}{\sqrt[t]{x}} \Rightarrow C = -\frac{1}{t} \log_b x.$$
(3)

Since we need the largest real root W then we should find the smallest positive root x of the equation $(1-x)^b = 0 \Rightarrow x = 1 \Rightarrow C = 0$.

4 Conclusion

Based on Shannon's classical work, we derive a formula for the information content of a multiset. Using the definition and the determinant capacity formula, we compute the multiset channel capacity. As future work we plan to further explore the properties of multiset-based communication systems, and compare these to similar results for string-based communication systems.

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Causality in Membrane Systems

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Summary. P systems are a biologically inspired model introduced by Gheorghe Păun with the aim of representing the structure and the functioning of the cell. P systems are usually equipped with the maximal parallelism semantics; however, since their introduction, some alternative semantics have been proposed and investigated.

We propose a semantics that describes the causal dependencies occurring between the reactions of a P system. We investigate the basic properties that are satisfied by such a semantics. The notion of causality turns out to be quite relevant for biological systems, as it permits to point out which events occurring in a biological pathway are necessary for another event to happen.

1 Introduction

Membrane computing is a branch of natural computing, initiated by Gheorghe Păun with the definition of P systems in [22, 23, 24]. The aim is to provide a formal modeling of the structure and the functioning of the cell, making use especially of automata, languages and complexity theoretic tools.

Membrane systems are based upon the notion of *membrane structure*, which is a structure composed by several cell-membranes, hierarchically embedded in a main membrane called the *skin membrane*. A plane representation of a membrane structure can be given by means of a Venn diagram, without intersected sets and with a unique superset. The membranes delimit *regions* and we associate with each region a set of *objects*, described by some symbols over an alphabet, and a set of *evolution rules*.

In the basic variant, the objects evolve according to the evolution rules, which can modify the objects to obtain new objects and send them outside the membrane or to an inner membrane. The evolution rules are applied in a maximally parallel manner: at each step, all the objects which can evolve should evolve.

A computation device is obtained: we start from an initial configuration, with a certain number of objects in certain membranes, and we let the system evolve. If a computation *halts*, that is no further evolution rule can be applied, the result of the computation is defined to be the number of objects in a specified membrane (or expelled through the skin membrane). If a computation never halts (i.e. one or more object can be rewritten forever), then it provides no output.

Since their introduction, plenty of variants of P systems have been introduced, and a lot of research effort has been carried out, expecially concerned with the study of the expressivity and the universality of the proposed models and with the ability to solve NP-complete problems in polynomial time.

The aim of this work is to start an investigation of the causal dependencies arising in among reactions occurring in P systems. The main motivation for this work comes from system biology, as the understanding of the causal relations occurring between the events of a complex biological pathway could be of precious help, e.g., for limiting the search space in the case some unpredicted event occurs.

In this paper we concentrate on P systems with cooperative rules, namely systems whose evolution rules are of the form $u \to v$, representing the fact that the objects in u are consumed and the objects in v are produced.

The study of causal semantics in concurrency theory is quite old. For example, the study of a causal semantics for process algebras dates back to the early nineties for CCS [20] (see, e.g., [13, 11, 18]), and to the mid nineties for the π -calculus [21] (see, e.g., [3, 5, 14, 15]).

To the best of our knowledge, the only other works that deal with causality in bio-inspired calculi with membranes and compartments are the following. In [7] a causal semantics for the Mate/Bud/Drip Brane Calculus [9] is proposed. In [17] a causal semantics for Beta Binders [26, 27] – based on the π -calculus semantics and on the enhanced operational semantics approach of [15] – is defined. One of the main differences between Beta Binders on one side, and Brane Calculi and P systems on the other side, is that the membrane structure in Beta Binders is flat, whereas in Brane Calculi and in P systems the membranes are nested to form a hierarchical structure.

The paper is organized as follows. After providing some basic definitions in Section 2, in Section 3 we define (cooperative) P systems. Section 4 recalls a detailed definition of maximal parallelism semantics that will be used in the following to provide a comparison between the causal and the maximal parallelism semantics. Section 5 is devoted to the definition of the causal semantics; after an informal introduction, a formal definition is provided, and finally some result on the properties enjoyed by the causal semantics are given. Section 6 reports some conclusive remarks.

2 Basic Definitions

In this section we provide some definitions that will be used throughout the paper.

Definition 1. Given a set S, a finite multiset over S is a function $m : S \to \mathbb{N}$ such that the set $dom(m) = \{s \in S \mid m(s) \neq 0\}$ is finite. The multiplicity of an element s in m is given by the natural number m(s). The set of all finite

multisets over S, denoted by $\mathcal{M}_{fin}(S)$, is ranged over by m. A multiset m such that $dom(m) = \emptyset$ is called empty. The empty multiset is denoted by \emptyset .

Given the multisets m and m', we write $m \subseteq m'$ if $m(s) \leq m'(s)$ for all $s \in S$ while \oplus denotes their multiset union, i.e., $m \oplus m'(s) = m(s) + m'(s)$. The operator \setminus denotes multiset difference: $(m \setminus m')(s) = if m(s) \geq m'(s)$ then m(s) - m'(s)else 0. The scalar product, $j \cdot m$, of a number j with m is $(j \cdot m)(s) = j \cdot (m(s))$. The cardinality of a multiset is the number of occurrences of elements contained in the multiset: $|m| = \sum_{s \in S} m(s)$.

The powerset of a set S is defined as $\mathcal{P}(S) = \{X \mid X \subseteq S\}.$

Definition 2. Let m be a finite multiset over S and $X \subseteq S$. The multiset $m|_X$ is defined as follows: for all $s \in S$, $m|_X(s) = m(s)$ if $s \in X$, and $m|_X(s) = 0$ otherwise.

Definition 3. A string over S is a finite (possibly empty) sequence of elements in S. Given a string $u = x_1 \dots x_n$, the length of u is the number of occurrences of elements contained in u and is defined by |u| = n.

With S^* we denote the set of strings over S, and u, v, w, \ldots range over S. Given $n \ge 0$, with S^n we denote the set of strings of length n over S.

Given a string $u = x_1 \dots x_n$ and i such that $1 \leq i \leq n$, with $(u)_i$ we denote the *i*-th element of u, namely, $(u)_i = x_i$.

Given a string $u = x_1 \dots x_n$, the multiset corresponding to u is defined as follows: for all $s \in S$, $m_u(s) = |\{i \mid x_i = s \land 1 \le i \le n\}|$. With abuse of notation, we use u to denote also m_u .

Definition 4. With $S \times T$ we denote the Cartesian product of sets S and T, with $\times_n S$, $n \ge 1$, we denote the Cartesian product of n copies of set S and with $\times_{i=1}^n S_i$ we denote the Cartesian product of sets S_1, \ldots, S_n , i.e., $S_1 \times \ldots \times S_n$. The ith projection of $(x_1, \ldots, x_n) \in \times_{i=1}^n S_i$ is defined as $\pi_i(x) = x_i$, and lifted to subsets $X \subseteq \times_{i=1}^n S_i$ as follows: $\pi_i(X) = {\pi_i(x) \mid x \in X}$.

Given a binary relation R over a set S, with R^n we denote the composition of n instances or R, with R^+ we denote the transitive closure of R, and with R^* we denote the reflexive and transitive closure of R.

3 P Systems

We recall the definition of catalytic P systems without priorities on rules. For a thorough description of the model, motivation, and examples see, e.g., [8, 12, 22, 23, 24]. To this aim, we start with the definition of *membrane structure*:

Definition 5. Given the alphabet $\{[,]\}$, the set MS is the least set inductively defined by the following rules:

• $[] \in MS$

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• *if* $\mu_1, \mu_2, \dots, \mu_n \in MS$, $n \ge 1$, then $[\mu_1 \dots \mu_n] \in MS$

We define the following relation over $MS: x \sim y$ iff the two strings can be written as $x = [1 \dots [2 \dots]2 \dots [3 \dots]3 \dots]1$ and $y = [1 \dots [3 \dots]3 \dots [2 \dots]2 \dots]1$ (i.e., if two pairs of parentheses that are neighbors can be swapped together with their contents).

The set \overline{MS} of membrane structures is defined as the set of equivalence classes w.r.t. the relation \sim^* .

We call a *membrane* each matching pair of parentheses appearing in the membrane structure. A membrane structure μ can be represented as a Venn diagram, in which any closed space (delimited by a membrane and by the membranes immediately inside) is called a *region* of μ .

Definition 6. A P system (of degree d, with $d \ge 1$) is a construct

$$\Pi = (V, \mu, w_1^0, \dots, w_d^0, R_1, \dots, R_d, i_0), \text{ where:}$$

- 1. V is a finite alphabet whose elements are called objects;
- 2. μ is a membrane structure consisting of d membranes (usually labelled with i and represented by corresponding brackets $[i \text{ and }]_i$, with $1 \leq i \leq d$);
- 3. w_i^0 , $1 \le i \le d$, are strings over V associated with the regions 1, 2, ..., d of μ ; they represent multisets of objects present in the regions of μ at the beginning of computation (the multiplicity of a symbol in a region is given by the number of occurrences of this symbol in the string corresponding to that region);
- 4. R_i , $1 \le i \le d$, are finite sets of evolution rules over V associated with the regions $1, 2, \ldots, d$ of μ ; these evolution rules are of the form $u \to v$, where u and v are strings from $(V \times \{here, out, in\})^*$;
- 5. $i_0 \in \{1, \ldots, d\}$ specifies the output membrane of Π .

The membrane structure and the multisets represented by w_i^0 , $1 \le i \le d$, in Π constitute the *initial state*¹ of the system. A transition between states is governed by an application of the evolution rules which is done in parallel; all objects, from all membranes, which *can be* the subject of local evolution rules *have to* evolve simultaneously.

The application of a rule $u \to v$ in a region containing a multiset m results in subtracting from m the multiset identified by u, and then in adding the multiset defined by v. The objects can eventually be transported through membranes due to the targets *in* and *out* (we usually omit the target *here*).

The system continues parallel steps until there remain no applicable rules in any region of Π ; then the system halts. We consider the number of objects from V contained in the output membrane i_0 when the system halts as the *result* of the underlying computation of Π .

¹ Here we use the term *state* instead of the classical term *configuration* because we will define a (essentially equivalent but syntactically) different notion of configuration in Section 5.

We introduce a couple of functions on membrane structures that will be useful in the following:

Definition 7. Let μ be a membrane structure consisting of d membranes, labelled with $\{1, \ldots, d\}$.

Given two membranes i and j in μ , we say that i is contained in j if the surface delimited by the perimeter of i in the Venn diagram representation of μ is contained inside the perimeter of j.

We say that i is the father of j (and j is a child of i) if the membrane j is contained in i, and no membrane exists that contains j and is contained in i.

The partial function father : $\{1, \ldots, d\} \rightarrow \{1, \ldots, d\}$ returns the father of a membrane *i*, or is undefined if *i* is the external membrane.

The function children : $\{1, \ldots, d\} \rightarrow \mathcal{P}(\{1, \ldots, d\})$ returns the set of children of a membrane.

For example, take $\mu = [1[2[3]_3]_2 [4]_4]_1$; then, father(2) = father(4) = 1, father(3) = 2 and father(1) is undefined; moreover, $children(4) = \emptyset$ and $children(1) = \{2, 4\}$.

4 Maximal Parallelism Semantics for P Systems

In order to compare the classical maximal parallelism semantics with the causal semantics, in this section we recall a detailed definition of the computation of a P system, proposed in [4], where a maximal parallelism evolution step is represented as a (maximal) sequence of simple evolution steps, which are obtained by the application of a single evolution rule.

Throughout this section, we let $\Pi = (V, \mu, w_1^0, \dots, w_d^0, R_1, \dots, R_d, i_0)$ be a P system.

To represent the states of the system reached after the execution of a non maximal sequence of simple evolution rules, we introduce the notion of *partial configuration* of a system. In a partial configuration, the contents of each region is represented by two multisets:

- The multiset of *active objects* contains the objects that were in the region at the beginning of the current maximal parallelism evolution step. These objects can be used by the next simple evolution step.
- The multiset of *frozen objects* contains the objects that have been produced in the region during the current maximal parallelism evolution step. These objects will be available for consumption in the next maximal parallelism evolution step.

Definition 8. A partial configuration of Π is a tuple $((w_1, \bar{w}_1), \ldots, (w_d, \bar{w}_d)) \in \times_d (V^* \times V^*).$

We use $\times_{i=1}^{d}(w_i, \bar{w}_i)$ to denote the partial configuration above.

The set of partial configurations of Π is denoted by $Conf_{\Pi}$. We use $\gamma, \gamma', \gamma_1, \ldots$ to range over $Conf_{\Pi}$.

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In the above definition, w_1, \ldots, w_d represent the active multisets, whereas $\bar{w}_1, \ldots, \bar{w}_d$ represent the frozen multisets.

A *configuration* is a partial configuration containing no frozen objects; configurations represent the states reached after the execution of a maximal parallelism computation step.

Definition 9. A configuration of Π is a partial configuration $\times_{i=1}^{d}(w_i, \bar{w}_i)$ satisfying the following: $\bar{w}_i = \emptyset$ for $i = 1, \ldots, d$.

The initial configuration of Π is the configuration $\times_{i=1}^{d}(w_i^0, \emptyset)$.

The size of a partial configuration is the number of active objects contained in the configuration.

Definition 10. Let $\gamma = \times_{i=1}^{d} (w_i, \bar{w}_i)$ be a partial configuration. The size of γ is $\#(\gamma) = \sum_{i=1}^{d} |w_i|$.

The execution of a simple evolution rule is formalized by the notion of reaction relation, defined as follows:

Definition 11. The reaction relation \mapsto over $Conf_{\Pi} \times Conf_{\Pi}$ is defined as follows:

 $\times_{i=1}^{d}(w_i, \bar{w}_i) \mapsto \times_{i=1}^{d}(w'_i, \bar{w}'_i)$ iff there exist k, with $1 \leq k \leq d$, an evolution rule $u \to v \in R_k$ and a migration string $\rho \in \{1, \ldots, d\}^{|v|}$ such that

- $u \subseteq w_k$
- $w'_k = w_k \setminus u$
- $\forall i: 1 \leq i \leq d \text{ and } i \neq k \text{ implies } w'_i = w_i$
- $\forall j: 1 \leq j \leq |v|$ the following holds:

- if
$$\pi_2((v)_j) = here \ then \ (\rho)_j = k$$

- if
$$\pi_2((v)_j) = out \ then \ (\rho)_j = father(k)^2$$

- if $\pi_2((v)_j) = in$ then $(\rho)_j \in children(k)^3$
- $\forall i, 1 \leq i \leq d$: $\bar{w}'_k = \bar{w}_k \oplus \bigoplus_{1 \leq j \leq |v|, (\rho)_j = k} (v)_j$

Note that the size of a configuration represents an upper bound to the length of the sequences of reactions starting from that configuration. Hence, infinite sequences of reactions are not possible.

Proposition 1. Let γ be a configuration. If $\gamma \mapsto^n \gamma'$ then $n \leq \#(\gamma)$.

The *heating function heated* transforms the frozen objects of a configuration in active objects, and will be used in the definition of the maximal parallelism computation step.

Definition 12. Let $\times_{i=1}^{d}(w_i, \bar{w}_i)$ be a partial configuration of Π . The heating function heated : $Conf_{\Pi} \to Conf_{\Pi}$ is defined as follows: $heated(\times_{i=1}^{d}(w_i, \bar{w}_i)) = \times_{i=1}^{d}(w_i \oplus \bar{w}_i, \emptyset)$

² As $\rho \in \{1, \ldots, d\}^{|v|}$, this implies that father(k) is defined.

³ This implies that children(k) is not empty.

Now we are ready to define the maximal parallelism computational step \Rightarrow :

Definition 13. The maximal parallelism computational step \Rightarrow over (nonpartial) configurations of Π is defined as follows: $\gamma_1 \Rightarrow \gamma_2$ iff there exists a partial configuration γ' such that $\gamma_1 \mapsto^+ \gamma'$, $\gamma' \not\mapsto$ and $\gamma_2 = heated(\gamma')$.

An operational semantics for P systems with maximal parallelism semantics has been defined for P systems in [1, 2, 10]. The main difference w.r.t. our approach is concerned with the fact that, while in this Section a maximal parallelism computational step is defined as a maximal sequence of reactions, in [1, 2, 10] no notion of reaction is provided, and the notion equivalent to the maximal parallelism computational step is defined directly by SOS rules [25]. A detailed comparison of the two approaches is beyond the scope of the present paper and deserves further investigation.

5 A Causal Semantics for P Systems

In this section we provide a causal semantics for cooperative P systems. To define a causal semantics, we follow the approach used in [18] for CCS, and in [3] for the π -calculus.

5.1 An informal explanation

The idea consists in decorating the reaction relation with two pieces of information:

- a fresh name k, that is associated to the reaction and it is taken from the set of causes K;
- a set $H \subseteq \mathcal{K}$, containing all the names associated to the already occurred reactions, that represent a cause for the current reaction.

To keep track of the names of the already occurred reactions that may represent a cause for the reactions that may happen in the future, we introduce a notion of causal configuration that associates to each object an information on its causal dependencies. As in [3], for the sake of clarity we only keep track of the so called immediate causes, as the set of general causes can be reconstructed by transitive closure of the immediate causal relation. We will provide more explanation on this point with an example in the following part of the paper.

Now we start with an informal introduction of causality in P systems. Consider the following system with a unique membrane:

$$\Pi_1 = (\{a, b, c, d, e, f\}, [1]_1, ae, \{a \to bc, c \to d, e \to f\}, 1).$$

If we consider the reaction relation \mapsto defined in the previous section, we have that the system Π_1 can perform either a reaction obtained by the application of the rule $a \to bc$ followed by a reaction obtained by the application of rule $e \to f$, or a sequence of two reactions where the application the rule $e \to f$ is followed by the application of $a \to bc$. The applications of the two rules are independent, in the sense that all the objects consumed by both the rules are already present in the initial configuration. Hence, the two rules can be applied in the same maximal parallelism step, and no one of the rules is causally dependent on the other one.

Consider now the system

$$\Pi_2 = (\{a, b, c, d, e, f\}, [1]_1, a, \{a \to bc, c \to d, e \to f\}, 1),$$

obtained from Π_1 by removing object e from the initial state. In this case, only rule $a \to bc$ can be applied. After the application of such a rule, an instance of object c is created by the application of rule $a \to bc$. Now, a further reduction step can be performed, consisting in applying rule $c \to d$. However, the applications of the two rules $a \to bc$ and $c \to d$ cannot be swapped, and the two rules cannot be applied in the same maximal parallelism computational step. This is because the object c consumed by rule $c \to d$ has been produced by rule $a \to bc$. In this case, we say that the reduction step consisting in the application of rule $c \to d$ causally depends on the reduction step consisting in the application of rule $a \to bc$.

If we consider again system Π_1 , we have that, after the application of the two rules $a \to bc$ and $e \to f$, the rule $c \to e$ can be applied, and it is caused by the application of rule $a \to bc$.

We would like to note that the causal semantics is in some sense "finer" than the maximal parallelism step semantics, as it permits to identify exactly which rule(s) represent a cause for the execution of another rule. Consider, e.g., the system

$$\Pi_3 = (\{a, b, c, d, e, f\}, [1]_1, ae, \{a \to bc, cf \to d, e \to f\}, 1).$$

According to the maximal parallelism semantics, the two systems cannot be distinguished, as both can perform a maximal parallelism step containing two rules (i.e., $\{a \to bc, e \to f\}$), followed by a maximal parallelism step containing a single rule (resp. $\{c \to d\}$ for Π_1 and $\{cf \to d\}$ for Π_3). On the other hand, if we consider the causal semantics, we have that the application of rule $c \to d$ in Π_1 causally depends only on one of the two rules applied in the previous maximal parallelism step, i.e., $a \to bc$, whereas the application fo the rule $cf \to d$ in Π_3 causally depends on both the rules applied in the previous maximal parallelism step.

5.2 The formal definition of causal semantics

In this section we provide a formal definition of the notions introduced in the previous section.

Let \mathcal{K} be a denumerable set of cause names, disjoint from the set V of objects. Throughout this section, we let $\Pi = (V, \mu, w_1^0, \dots, w_d^0, R_1, \dots, R_d, i_0)$ be a P system.

To be able to define the set of causes of a reaction, we proceed in the following way: we associate a fresh (i.e., never used before) cause name to each reaction performed in the system. Then, each instance of object in a configuration of the system is decorated with the causal name of the reaction that produced it, or with \emptyset if the object is already present in the initial configuration.⁴ To keep track of such causal information, we introduce the notion of *causal configuration* fo a system.

Definition 14. A causal configuration of Π is a tuple z_1, \ldots, z_d , where $z_i \in (V \times \mathcal{P}(\mathcal{K}))^*$ for $i = 1, \ldots, d$.

We use $\times_{i=1}^{d} z_i$ to denote the causal configuration above.

The set of causal configurations of Π is denoted by $CConf_{\Pi}$.

We use $\gamma, \gamma', \gamma_1, \ldots$ to range over $CConf_{II}$.⁵

Let $w_i^0 = o_{i,1}o_{i,2}\dots o_{i,n_i}$ for $i = 1, \dots, d$. The initial causal configuration of Π is the configuration $\times_{i=1}^d (o_{i,1}, \emptyset)(o_{i,2}, \emptyset) \dots (o_{i,n_i}, \emptyset)$.

For example, $((a, \emptyset)(e, \emptyset))$ represents the initial causal configuration of the P system Π_1 in the previous subsection, and $((b, k_1)(c, k_1)(e, \emptyset))$ represents another configuration of Π_1 , reached after the firing of rule $a \to bc$ (for the sake of clarity we omit the surrounding braces if the set of causes is a singleton).

Now we are ready to define the causal semantics for P systems. We write $\gamma \xrightarrow{h;H} \gamma'$ to denote the fact that system Π in configuration γ performs an action – to which we associate the cause name h – that is caused by the (previously occurred) actions whose action names form the set H. The cause name h is a fresh name: this means that it has not been used yet in the current computation.

The execution of an evolution rule is formalized by the notion of causal reaction relation.

Before providing the definition of causal reaction relation, we need some auxiliary definitions.

Definition 15. The function drop : $(V \times \mathcal{P}(\mathcal{K}))^* \to V^*$ removes the causality information:

$$drop(\varepsilon) = \varepsilon$$

$$drop((o, H)w) = o \ drop(w)$$

The function drop is extended to configurations in the obvious way:

$$drop(\times_{i=1}^{d} z_i) = \times_{i=1}^{d} drop(z_i)$$

The function causes $: V \times \mathcal{P}(\mathcal{K}))^* \to \mathcal{P}(\mathcal{K})$ produces the set of causal labels in a string:

$$causes(\varepsilon) = \emptyset$$

$$causes((o, H)w) = H \cup causes(w)$$

⁴ For homogeneity with other classes of P systems, actually we decorate each object with a – possibly empty – set of cause names, even if, in the class of P systems considered in this paper, a single cause name is sufficient.

⁵ With abuse of notation, we use $\gamma, \gamma', \gamma_1, \ldots$ to denote both partial configurations and causal configurations. It will be clear from the context to which kind of configuration we are referring to.

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The function deco: $V^* \to V \times \mathcal{P}(\mathcal{K}))^*$ decorates each object in a string with a given set of causal labels:

$$deco(\varepsilon, H) = \emptyset$$

$$deco(ow, H) = (o, H)deco(w, H)$$

Definition 16. The causal reaction relation $\xrightarrow{h;H}$ over $CConf_{\Pi} \times CConf_{\Pi}$ is defined as follows:

 $\times_{i=1}^{d} z_i \xrightarrow{h;H} \times_{i=1}^{d} z'_i \text{ iff there exist } k, \text{ with } 1 \leq k \leq d, \text{ a string } w \in (V \times \mathcal{P}(\mathcal{K}))^*,$ an evolution rule $u \to v \in R_k$ and a migration string $\rho \in \{1, \ldots, d\}^{|v|}$ such that

- u = drop(w)
- H = causes(w)
- $w \subseteq z_k$
- $z'_k = z_k \setminus w \oplus deco(v, \{h\})$
- $\forall j: 1 \leq j \leq |v|$ the following holds:
 - if $\pi_2((v)_i) = here \ then \ (\rho)_i = k$
 - if $\pi_2((v)_i) = out$ then $(\rho)_i = father(k)^6$
- $\forall i, 1 \leq i \leq d \text{ and } i \neq k: z'_i = z_i \oplus \bigoplus_{1 \leq j \leq |v|, (\rho)_j = i} ((v)_j, h)$

5.3 Properties of the causal semantics

The causal semantics for the class of P systems considered in this paper enjoys some nice properties.

The first property is the *retrievability* of the maximal parallelism step semantics from the causal semantics. According to such a property, there is no loss of information when moving from the maximal parallelism to the causal semantics, as we can reconstruct the maximal parallelism semantics of a system by looking at its causal execution:

Theorem 1. $\times_{i=1}^{d}(w_i, \emptyset) \Rightarrow \times_{i=1}^{d}(w'_i, \emptyset)$ is a maximal parallelism computational step if and only if there exist $\gamma, \gamma' \in CConf(\Pi), h_1, \ldots, h_n, H_1, \ldots, H_n$ such that

- $drop(\gamma) = \times_{i=1}^{d} (w_i, \emptyset)$ $drop(\gamma') = \times_{i=1}^{d} (w'_i, \emptyset)$

•
$$\gamma \xrightarrow{h_1;H_1} \dots \xrightarrow{h_n;H_n} \gamma$$

- $h_i \notin H_j$ for all $i, j: 1 \le i, j \le n$
- if there exist h, H such that $\gamma' \xrightarrow{h;H}$ then there exists i such that $1 \leq i \leq n$ and $h_i \in H$

The other property is the so-called diamond property, stating that if two noncausally related actions can happen one after the other, then they can happen also in the other order, and at the end they reach the same system.

⁶ As $\rho \in \{1, \ldots, d\}^{|v|}$, this implies that father(k) is defined.

⁷ This implies that children(k) is not empty.

Theorem 2. If $\gamma \xrightarrow{h_1;H_1}_{r_1} \gamma' \xrightarrow{h_2;H_2}_{r_2} \gamma''$ and $h_1 \notin H_2$ then there exists a causal configuration γ''' such that $\gamma \xrightarrow{h_2;H_2}_{r_2} \gamma''' \xrightarrow{h_1;H_1}_{r_1} \gamma''$.

6 Conclusion

In this paper we tackled the problem of defining a causal semantics for a basic class of P systems. We think that the study of the causal dependencies that arise between the actions performed by a system is of primary importance for models inspired by the biology, because of its possible application to the analysis of complex biological pathways.

This paper represents a first step in this direction, but a lot of work remains to be done. For example, if we move to other classes of membrane systems, such as, e.g., P systems with promoters and inhibitors, we have to deal with more involved causal relations among reactions, and it could happen that some of the properties enjoyed by the causal semantics for basic P systems presented in this work no longer hold. Another interesting research topic is the investigation of the causal semantics for classes of P systems whose membrane structure is dynamically evolving (e.g., we can consider dissolution rules, duplication, gemmation or either brane-like operations). Once we have completed the definition of a causal semantics for systems with an evolving structure, we will start investigating the causal dependencies arising in biological pathways involving membranes, such as, e.g., the LDL Cholesterol Degradation Pathway [19], that was modeled in P systems in [6].

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Hierarchical Clustering with Membrane Computing

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Summary. In this paper we approach the problem of the hierarchical clustering through membrane computing. A specific P system with external output is designed for each boolean matrix associated with a finite set of individuals. The computation of the system allows us to obtain one of the possible classifications in a non-deterministic way. The amount of resources required in the constructions is polynomial in the number of individuals and the number of characteristics analyzed.

1 Introduction

Researchers develop a lot of investigation that depend on many factors and this makes their study very complex. In order to simplify and make the problems more tractable it is necessary to group individuals with similar characteristics. The individuals are characterized by a high number of properties so the grouping is not a simple task. The clustering methods appear with the purpose of establishing a methodology with a statistical base in order to obtain the groupings of the individuals according to their degree of similarity.

There are different methods of ranking the groups of individuals. In order to simplify it we can consider two types, the nonhierarchical clustering and the hierarchical clustering. In a nonhierarchical clustering homogenous groups are formed without establishing relations among them; in the hierarchical clustering the individuals are grouped in levels. The inferior levels are contained in the superior levels. The hierarchical clustering is the most used and it is dealt with in this paper. 186 M. Cardona et al.

Hierarchical clustering refers to the formation of a recursive clustering of the individuals by means of the partitions P_0, P_1, \ldots, P_m of the set of N individuals with $1 \leq m \leq N-1$. The partition P_0 consists of N groups each one of them formed by a single individual. The groups that form this partition join progressively until arriving at the last partition, P_m , that consists of a single group formed by all the individuals. In each step the two most similar groups are joined according to a previously established criterion.

Researchers use the clustering to characterize and to order a vast amount of information on the variability of population of individuals. These populations are grouped in more or less homogenous clusters based on their properties. This methodology has been applied in fields as diverse as Medicine, Biology, classification of words, of the fingerprints, artificial intelligence... Recently the clustering has been applied to the classification of musical genre [13], to predict essential hypertension [12], in the classification of material planning and control systems [9], in the classification of the ocean color [1], in the classification of the plants gens [14].

The different groups obtained by means of the classification are characterized by different levels of the measured variables. These values allow us to give common properties of the individuals belonging to the same group. To have established groups allows us to identify the most similar cluster of a new individual. The characteristics measured of the individuals can be qualitative variables or quantitative variables. In most cases we are only interested in the presence or absence of certain qualitative characteristics. So in this paper we make a hierarchical clustering using dichotomizing variables by means of membrane computing.

In this paper the problem of hierarchical clustering is approached with the framework of cellular computing with membranes. It is interesting because allows us treated some statistics topics with this new models of computation. The amount of used resources is polynomial in the number of individuals and the number of characterizes analyzed without increasing the complexity of the classical clustering algorithms.

In the following, we assume that the reader is familiar with the basic notions of P systems, and we refer, for details, to [7], [15], [8], [6].

2 Overview

2.1 Hierarchical Clustering

In order to obtain a hierarchical clustering we need a set of observations or individuals that we define as follows:

Definition 1. A k-set Ω over a metric space (E,d), with $d(E \times E) \subseteq \mathbb{N}$, is a subset of E^k .

The hierarchical clustering needs a finite k-set Ω with N elements, $\Omega = \{\omega_1, \ldots, \omega_N\}$. The elements of the set Ω are called individuals or observations

and their components in the k-tuple are denoted characteristics or variables. The values of the individuals can be represented in matrix form:

$$P_{Nk} = \begin{pmatrix} \omega_{11} & \omega_{12} & \cdots & \omega_{1k} \\ \omega_{21} & \omega_{22} & \cdots & \omega_{2k} \\ & \ddots & & \\ \omega_{N1} & \omega_{N2} & \cdots & \omega_{Nk} \end{pmatrix}$$

where ω_{ij} is the value of the *j*-th variable of the individual *i*.

The objective of any clustering is to group the individuals in similar groups whose members are all close to one another with various dimensions being measured. It will be necessary to establish criteria in order to measure the similarity between individuals and similarity between groups. Evidently, the clustering that is obtained will depend on the similarity function that is chosen. This function is called similarity and it is defined as follows [10].

Definition 2. A similarity over a finite k-set $\Omega = \{\omega_1, \ldots, \omega_N\}$ is a function s of $\Omega \times \Omega$ in \mathbb{R}^+ that verifies

- s is symmetric, that is $\forall (\omega_i, \omega_j) \in \Omega \times \Omega$: $s(\omega_i, \omega_j) = s(\omega_j, \omega_i)$
- $\forall \omega_i, \omega_j \in \Omega \text{ with } i \neq j : \quad s(\omega_i, \omega_i) = s(\omega_j, \omega_j) \ge s(\omega_i, \omega_j)$

In this paper we work with dichotomizing variables, in particular their values are denoted by 0 and 1. One of the similarities most used for dichotomizing variables is the similarity of Sokal and Michener [2] and it is defined by:

$$\forall \omega_i, \omega_j \in \Omega: \quad s'(\omega_i, \omega_j) = \frac{1}{k} \cdot \sum_{r=1}^k (1 - |\omega_{ir} - \omega_{jr}|) \tag{1}$$

where $\omega_i = \{\omega_{i1}, \ldots, \omega_{ik}\}.$

In this paper the similarity that we use is a modification of the previous one. This similarity represents the number of coincidences in the number of total characteristics and it is defined as follow:

$$\forall (\omega_i, \omega_j) \in \Omega \times \Omega : \quad s(\omega_i, \omega_j) = \sum_{r=1}^k (1 - |\omega_{ir} - \omega_{jr}|)$$
(2)

We use this similarity because it is easier to implement with P systems and the result obtained is the same as we obtain with the similarity of Sokal and Michener.

In the case of the hierarchical clustering the groupings follow a hierarchy formed by partitions. The partitions are formed in a recursive manner. We start with as many clusters as individuals, in each iteration the partition is obtained joining the two closest clusters. This process is done until we obtain a single set formed by all the individuals. The partitions obtained P_0, P_1, \ldots, P_m verify $P_0 \subseteq P_1 \subseteq P_2 \subseteq$ $\ldots \subseteq P_m$ with $1 \le m \le N-1$ and the sets that form the partitions are called clusters.

Next we define the necessary mathematical concepts in the hierarchical clustering [11].

Definition 3. Let $\Omega = \{\omega_1, \ldots, \omega_N\}$ the k-set of N individuals to classify. A subset H of the parts of Ω , $H \subseteq \mathcal{P}(\Omega)$, is a hierarchy over Ω if it verifies:

- $\Omega \in H$
- $\{\omega\} \in H \quad (\forall \omega \in \Omega)$ •
- If $h \cap h' \neq \emptyset \Rightarrow h \subset h'$ or $h' \subset h$ $(\forall h, h' \in H)$
- $\bigcup \{h' \mid h' \in H, h' \subseteq h\} \in \{h, \emptyset\} \quad (\forall h \in H)$

The elements of H are called clusters. If $h_1, \ldots, h_p \in H$ with $\Omega = h_1 \cup \ldots \cup h_p$ then the set $\{h_1, \ldots, h_p\}$ is a clustering.

In order to construct a hierarchy it is necessary to have a similarity between individuals and another function that measures the similarity between clusters. The second function is called the aggregation index.

Definition 4. A symmetrical and nonnegative application $\delta: \mathcal{P}(\Omega) \times \mathcal{P}(\Omega) \to \mathbf{R}$ is called aggregation index between clusters if it verifies:

- $\forall h_1, h_2 \in \mathcal{P}(\Omega) : \quad \delta(h_1, h_2) \ge 0$ $\forall h_1, h_2 \in \mathcal{P}(\Omega) : \quad \delta(h_1, h_2) = \delta(h_2, h_1)$

There are several aggregation indices [4] that depend on the similarity s chosen. In this paper we use the aggregation index based on the minimum [5] defined by:

$$\delta(h_1, h_2) = \min\{s(\omega_i, \omega_j) \mid \omega_i \in h_1, \ \omega_j \in h_2\}$$
(3)

If a hierarchy has associated an index that measures the homogeneity degree between the individuals belonging to the same cluster it is called indexed hierarchy. We refer to this index by the hierarchical index.

Definition 5. An indexed hierarchy is a pair (H, f) where H is a hierarchy and f is an application H over \mathbf{R}^+ such that:

- $f(\{\omega\}) = k \; (\forall \omega \in \Omega)$
- $\forall h' \in H : h \subsetneq h' \Rightarrow f(h) > f(h')$

The hierarchical index is always obtained by means of the aggregation index. In this paper we define the hierarchical index of a new cluster h obtained from the union of two clusters $h = h_1 \cup h_2$, by means of $f(h) = \delta(h_1, h_2)$.

An algorithm for the construction of an indexed hierarchy

The algorithms that are used to obtain an indexed hierarchy have the same structure, the only differences in them is the way to compute the similarities between clusters [3].

The input of this algorithm is the k-set Ω and the aggregation index δ . The output is an indexed hierarchy (H, f).

1 Place each individual of Ω in its own cluster (singleton), creating the list of clusters $L = P_0$

$$L = P_0 = \{S_1 = \{\omega_1\}, S_2 = \{\omega_2\}, \dots, S_N = \{\omega_N\}\}$$

In this moment $\delta(\{\omega_i\}, \{\omega_j\}) = s(\omega_i, \omega_j)$ and $f(\{\omega_i\}) = k \ (1 \le i < j \le N)$

- 2 Find the two closest clusters S_i, S_j with $1 \le i < j \le N$, which will form a new class $S_i = S_i \cup S_j$.
- 3 Remove S_j from L.
- 4 Compute the aggregation index, by equation (3), between all the pair of clusters in L.
- 5 Go to step 2 until there is only one set remaining.

Remark: If at step 2 there are more than one possibility, then one of them is chosen at random so the hierarchy obtained is not unique.

3 Hierarchical Clustering of a Group of Individuals

3.1 Designing a P System

The goal of this paper is to obtain one hierarchical clustering of a k-set Ω , of N different individuals by means of the cellular computing with membranes. We considered each individual $\omega_i \in \Omega$ by a k-tuple of dichotomizing variables, $\Omega \subseteq \{0, 1\}^k$ which is denoted by $\omega_i = (\omega_{i1}, \omega_{i2}, \ldots, \omega_{ik})$. The similarity between individuals that we use is the following:

$$s(\omega_i, \omega_j) = \sum_{t=1}^k (1 - |\omega_{it} - \omega_{jt}|)$$

This similarity measures the number of equal components between two individuals.

Let $P_{Nk} = (\omega_{ij})_{1 \le i \le N, 1 \le j \le k}$ be the matrix formed by the k values of N individuals to classify. We define the P system of degree N with external output,

$$\Pi(P_{Nk}) = (\Gamma(P_{Nk}), \mu(P_{Nk}), \mathcal{M}_1, \mathcal{M}_2, \dots, \mathcal{M}_{N-1}, \mathcal{M}_N, R, \rho)$$

associated with the matrix P_{Nk} , as follows:

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• Working alphabet:

$$\begin{split} \varGamma(P_{Nk}) &= \{e_{js}, \quad d_{js}: \ 1 \leq j \leq N, 1 \leq s \leq k\} \ \cup \{a_{s}, \quad b_{s}: \ 1 \leq s \leq k\} \ \cup \\ \{S_{ij}, \quad C_{ij}: \ 1 \leq i < j \leq N\} \ \cup \{\beta_{i}: 0 \leq i \leq k-2\} \ \cup \\ \{\alpha_{ijt}, \quad X_{ijt}: \ 1 \leq i < j \leq N, \ 1 \leq t \leq k-1\} \ \cup \{\gamma_{i}: \ 1 \leq i \leq N\} \ \cup \\ \{\epsilon_{i}: \ 0 \leq i \leq 3k-2\} \ \cup \{\eta_{i}: \ 0 \leq i \leq (N-1)(3k-1)\} \ \cup \{\sharp\} \end{split}$$

- Membrane structure: $\mu(P_{Nk}) = [N \ [1 \]_1 \ [2 \]_2 \ \dots \ [N-1 \]_{N-1} \]_N.$
- Initial multisets:

$$\mathcal{M}_{i} = \{a_{s}^{(N-i)\omega_{is}}: 1 \leq s \leq k \land 1 \leq i \leq N-1\} \cup \\ \{b_{s}^{(N-i)(1-\omega_{is})}: 1 \leq s \leq k \land 1 \leq i \leq N-1\} \cup \\ \{e_{js}^{\omega_{js}}: 1 \leq s \leq k \land i \leq j \leq N\} \cup \\ \{d_{js}^{(1-\omega_{js})}: 1 \leq s \leq k \land i \leq j \leq N\}; 1 \leq i \leq N-1\}$$

 $\mathcal{M}_N = \{\gamma_N, \quad \epsilon_0, \quad \eta_0\};$

• The set *R* of evolution rules consists of the following rules: – Rules in the skin membrane labeled *N*:

$$r_0 = \{\epsilon_0 \to \epsilon_1 \beta_0\} \cup \{\epsilon_i \to \epsilon_{i+1} : 1 \le i \le 3k - 2 \land i \ne k\} \cup \{\eta_i \to \eta_{i+1} : 0 \le i \le (N-1)(3k-1) - 1\}$$

$$r_u = \{\beta_{u-1} S_{ij}^{k-u} \to \alpha_{ij(k-u)} : 1 \le i < j \le N\} \quad 1 \le u \le k-1$$

$$r'_u = \{\beta_{u-1} \to \beta_u\} \quad 1 \le u \le k-1$$

$$r_{k-1}' = \{\eta_{(N-1)(3k-1)} \rightarrow (\sharp, out)\}$$

$$r_k = \{ \epsilon_k \gamma_q \alpha_{ijt} \to \epsilon_{k+1} X_{ijt}^{q-2} \gamma_{q-1}(X_{ijt}, out) : 2 \le q \le N, \\ 1 \le i < j \le N, \ 1 \le t \le k-1 \}$$

$$r'_k = \{\epsilon_k \to \epsilon_{k+1}\}$$

$$\begin{aligned} r_{k+1} &= \{ X_{ijt} S_{ip} S_{jp} \to C_{ip} X_{ijt} : \ 1 \leq i < j < p \leq N, \ 1 \leq t \leq k-1 \} \cup \\ \{ X_{ijt} S_{ip} S_{pj} \to C_{ip} X_{ijt} : \ 1 \leq i < p < j \leq N, \ 1 \leq t \leq k-1 \} \cup \\ \{ X_{ijt} S_{pi} S_{pj} \to C_{pi} X_{ijt} : \ 1 \leq p < i < j \leq N, \ 1 \leq t \leq k-1 \} \end{aligned}$$

$$\begin{aligned} r_{k+2} &= \{ X_{ijt}S_{ip} \to X_{ijt} : \ 1 \leq i$$

$$\begin{aligned} r_{k+3} &= \{ C_{ij} \to S_{ij} : \ 1 \leq i < j \leq N \} \cup \\ \{ \epsilon_{3k-1} X_{ijt}^{q-2} \gamma_{q-1} \to \epsilon_1 \beta_0 \gamma_{q-1} : \ 1 \leq i < j \leq N, \ 1 \leq t \leq k-1 \} \end{aligned}$$

$$r'_{k+3} = \{\epsilon_{3k-1} \to \epsilon_1 \beta_0\}$$

- Rules in the membrane labeled
$$i \{1 \le i \le N-1\}$$
:
 $r_{k+4} = \{a_s e_{js} \to (S_{ij}, out): 1 \le s \le k, i+1 \le j \le N\}$
 $r_{k+5} = \{b_s d_{js} \to (S_{ij}, out): 1 \le s \le k, i+1 \le j \le N\}$

- The partial order relation ρ over R consists of the following priority relations:
 - Priority relation on the membrane labeled *i* with $1 \le i \le N 1$: $\rho_i = \emptyset$ - Priority relation on the skin membrane labeled *N*:
 - $\rho_N = \{r_1 > r'_1 > r_2 > r'_2 > \dots > r_{k-1} > r'_{k-1}\} \cup \{r_k > r'_k\} \cup \{r_{k+1} > r_{k+2} > r_{k+3} > r'_{k+3}\}$

3.2 An Overview of Computations

At the beginning of a computation the membrane labeled i, with $1 \leq i \leq N-1$, contains the objects a_s , b_s , e_{js} , d_{js} with $1 \leq s \leq k$ and $i+1 \leq j \leq N$. In this membrane the presence or absence of the objects a_s , b_s encode the values of the individual ω_i . If the value of ω_{is} is equal to 1 we have the object a_s and if the value ω_{is} is equal to 0 we have the object b_s .

The objects e_{js} , d_{js} with $a \leq s \leq k$ and $i < j \leq N$ are also in this membrane, and they codify the values of the k components of the individuals ω_j . If the value of the component s is 1, i.e. $\omega_{js} = 1$ then the membrane i contains the object e_{js} , if $\omega_{js} = 0$ then the membrane i contains the object d_{js} .

Initially, the skin membrane contains the objects γ_N , ϵ_0 and η_0 . The evolution of the object γ_N allows us to know the number of clusters in any configuration of the P system: when the object γ_i appears, then the individuals are grouped in *i* clusters. We use the object ϵ_0 in order to synchronize in 3k - 1 steps the loop, that allows us to unite two clusters with maximum similarity. The object η_0 is a counter used to stop the P system in the configuration (3k - 1)(N - 1) sending in the environment the object \sharp .

In the initial configuration the only rules that can be applied in membrane labeled i with $1 \leq i \leq N-1$ are r_{k+4} , r_{k+5} , that send the objects S_{ij} with $1 \leq i < j \leq N$ to the skin membrane. The multiplicity of these objects allows us to know the similarity between individuals of the set Ω , that is the number of equal components between these individuals. In this configuration the rule r_0 constructs the object β_0 .

¿From this configuration the computation of the P system is formed by loops of 3k-1 steps. Each one of these loops is formed by two very differentiated stages. The first stage is formed by k steps and begins with the object β_0 . In these steps the object S_{ij} with maximum multiplicity is selected encoding the maximum similarity between the clusters i and j. In the k-th step of the loop the rule r_k creates the objects X_{ijt} in the skin membrane and sends a copy to the environment. This object represents the clusters that have the highest similarity, t, that can be joined to form a new cluster. Moreover, in this step the object γ_q is transformed in the object γ_{q-1} , encoding the fact that two clusters have been joined.

The second stage is formed by 2k - 1 steps. In the skin membrane there are the objects X_{ijt} meaning that a new cluster *i* is formed by the union of the previous clusters *i*, *j*. The rules $r_{k+1}, r_{k+2}, r_{k+3}$ calculate the similarities between new cluster *i* and the other clusters, this information is kept in the multiplicity of the objects S_{ip} .

In the (3k-1)-th step of the loop the rule r_{k+3} transforms the object ϵ_{3k-1} in the objects β_0 and ϵ_1 that allow us to go to the top of the loop.

The first partition consist of N singletons; in each loop two clusters are joined so it is necessary N-1 loops to obtain the last partition that consists of a cluster containing all N individuals. Therefore the loop repeats N-1 times and the rule r'_{k-1} is applied finalizing the P system.

3.3 Formal Verification

In this section we are going to show that the P system $\Pi(P_{Nk})$ is non-deterministic, but, in spite of this for any computation we will obtain a solution of the clustering problem.

First of all, let us list the necessary resources to construct the P system $\Pi(P_{Nk})$ from the matrix P_{Nk} .

- Size of the alphabet: $\Theta(N^2 \cdot k)$.
- Sum of the sizes of initial multisets: $\Theta(N \cdot k)$.
- Maximum of rules' lengths: $\Theta(N)$.
- Number of rules: $\Theta(k \cdot N^3)$.
- Number of priority relations: $\Theta(k^2 \cdot N^6)$.
- Cost of time: $\Theta(N \cdot k)$.

Bearing in mind the recursive description of the rules and that the amount of resources is polynomial in N, k, it is possible to construct the system $\Pi(P_{Nk})$ from the matrix P_{Nk} by means of a Turing machine working in polynomial time.

Given a computation C of the P system $\Pi(P_{Nk})$, for each $p \in \mathbb{N}$ we denote by C_p the configuration of the P system obtained after the execution of p steps. For each level $l \in \{1, 2, \ldots, N\}$, we denote by $C_p(l)$ the multiset of objects contained in the membrane labeled l in the configuration C_p .

The following result proves that in the configuration C_1 , the multiplicity of the object S_{ij} , $\forall 1 \leq i < j \leq N$, represents the similarity between the individual $\omega_i = (\omega_{i1}, \ldots, \omega_{ik})$ and the individual $\omega_j = (\omega_{j1}, \ldots, \omega_{jk})$.

Proposition 1. Let C an arbitrary computation of the P system. If $t_{ij}^{(1)} = \max\{t : S_{ij}^t \in C_1(N)\} \quad \forall i, j, t \quad (1 \le i < j \le N, \quad 1 \le t \le k-1) \text{ then } t_{ij}^{(1)} = \sum_{s=1}^k (1 - |\omega_{is} - \omega_{js}|).$

Proof. In the initial configuration we have

$$\mathcal{C}_0(i) = \{a_s^{(N-i)\omega_{is}}, b_s^{(N-i)(1-\omega_{is})}, e_{js}^{\omega_{js}}, d_{js}^{(1-\omega_{js})} | \ i \le j \le N, \omega_{is} \in \{0, 1\}\}$$

with $1 \leq i \leq N - 1$.

The only rules that can be applied are r_{k+4} and r_{k+5} . The rule r_{k+4} is only possible to apply when the component s of the individuals ω_i and ω_j is equal 1. The rule r_{k+5} is only applied when the component s of the individuals ω_i and ω_j is equal 0.

Whenever one of these rules is applied the object S_{ij} goes out to the skin membrane. Then in $C_1(N)$ the multiplicity of the objects S_{ij} will coincide with the number of equal components between the individuals ω_i and ω_j , i.e. $|\omega_{is} - \omega_{js}| = 0$. Therefore the multiplicity of the objects S_{ij} is

$$t_{ij}^{(1)} = \sum_{s=1}^{k} (1 - \mid \omega_{is} - \omega_{js} \mid)$$

that is, $t_{ij}^{(1)}$ corresponds to the similarity between the individuals ω_i and ω_j . \Box

From now on we denote the maximum multiplicity of the objects S_{ij} in the step one of the *n*-th loop of the computation by

$$t_{ij}^{(n)} = \max\{t : S_{ij}^t \in \mathcal{C}_{1+(n-1)(3k-1)}(N)\}$$

In the following proposition we prove that each 3k - 1 steps is constructing the object β_0 so this object is in the skin of all the configurations of the type 1 + n(3k - 1) with $1 \le n \le N - 2$. Moreover we prove in what configuration the object ϵ_j with $1 \le j \le 3k - 1$ appears.

The objects β_0 and ϵ_1 determine the moment that the loop starts and the object ϵ_{3k-1} determines when the loop finishes.

Proposition 2. For each $n (0 \le n \le N - 2)$, we have:

a)
$$\beta_0 \in \mathcal{C}_{1+n(3k-1)}(N)$$

b) If
$$1 \le j \le 3k - 1$$
 then $\epsilon_j \in \mathcal{C}_{1+n(3k-1)+(j-1)}(N)$

Proof. We prove this proposition by induction on n.

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• For n = 0, it is necessary to verify that $\beta_0 \in C_1(N)$, and $\epsilon_j \in C_j(N)$, $\forall \ 1 \le j \le 3k - 1$.

In the initial configuration we have $\epsilon_0 \in \mathcal{C}_0(N)$ that allows us to apply one of the rules r_0 in order to obtain $\epsilon_1, \beta_0 \in \mathcal{C}_1(N)$, so a) is proved for n = 0.

In the following k-1 steps the rules r_0 will be applied transforming the object ϵ_1 until we obtain the object $\epsilon_k \in C_k(N)$. In this configuration if there are the objects $\alpha_{ijt}, \gamma_q \in C_k(N)$ the rule r_k will be applied, or the rule r'_k will be applied. In both cases ϵ_k evolves to $\epsilon_{k+1} \in C_{k+1}(N)$.

In the successive configurations the rule r_0 transforms the objects $\epsilon_j \in \mathcal{C}_j(N)$, $k+1 \leq j \leq 3k-2$ until we obtain the object $\epsilon_{3k-1} \in \mathcal{C}_{3k-1}(N)$.

• Let us suppose the hypothesis for $0 \le n < N-2$. Then, we will show that $\epsilon_j \in \mathcal{C}_{1+(n+1)(3k-1)+(j-1)}(N), \quad \forall \ 1 \le j \le 3k-1 \text{ and } \beta_0 \in \mathcal{C}_{1+(n+1)(3k-1)}(N).$ By induction hypothesis $\epsilon_{3k-1} \in \mathcal{C}_{1+n(3k-1)+(3k-1-1)}(N) = \mathcal{C}_{(n+1)(3k-1)}(N).$ If in this configuration there is some object X_{ijt} the rules r_{k+3} will be applied and in the other case the rule r'_{k+3} will be applied. In both cases the object ϵ_{3k-1} is transformed in $\epsilon_1, \ \beta_0 \in \mathcal{C}_{1+(n+1)(3k-1)}(N)$. So that a) is proved.

Applying k - 1 times the rules r_0 we obtain $\forall 1 \leq j \leq k$ that the object $\epsilon_j \in C_{1+(n+1)(3k-1)+(j-1)}(N)$. In the configuration $C_{1+(n+1)(3k-1)+(k-1)}(N)$ the object ϵ_k is transformed in the object $\epsilon_{k+1} \in C_{1+(n+1)(3k-1)+k}(N)$ by means of one of the rules r_k or r'_k . Then applying the rules r_0 successively we obtain $\forall k+1 \leq j \leq 3k-1$ that the object $\epsilon_j \in C_{1+(n+1)(3k-1)+(j-1)}(N)$.

Remark: According to Proposition 2 we have: $\epsilon_k \in \mathcal{C}_{1+n(3k-1)+k-1}(N) = \mathcal{C}_{k+n(3k-1)}(N) \quad \forall n \quad 0 \le n \le N-2$

Corollary 1. The objects X_{ijt} are sent to the environment at moments of the type $C_{1+n(3k-1)+k}$ with $0 \le n \le N-2$.

Proof. The only rule that sends some object X_{ijt} to the environment is the rule r_k . In order to be able to apply this rule the object ϵ_k is necessary, that verifies $\epsilon_k \in \mathcal{C}_{1+n(3k-1)+k-1}(N)$ with $0 \le n \le N-2$ by Proposition 2.

Therefore the objects X_{ijt} can only be sent to the environment in the following configuration, that is $X_{ijt} \in \mathcal{C}_{1+n(3k-1)+k}(env)$.

Proposition 3. The configuration $C_{(N-1)(3k-1)}$ sends to the environment the halt object \sharp .

Proof. Applying (N-1)(3k-1) times the rule r_0 the object $\eta_0 \in C_0(N)$ is transformed to $\eta_{(N-1)(3k-1)} \in C_{(N-1)(3k-1)}(N)$. In this configuration the rule r'_{k-1} sends the halt object \sharp to the environment.

In the following result we prove that it is only possible to modify the environment in the k-th step of the loop.

Corollary 2. Let C be an arbitrary computation of the P system. For each $0 \le n \le N - 2$ the following assertions hold:

a) For each r (1 + n(3k - 1) < r < 1 + n(3k - 1) + k) we have:

$$\mathcal{C}_r(env) = \mathcal{C}_{1+n(3k-1)}(env)$$

b) For each r (1 + n(3k - 1) + k < r < 1 + n(3k - 1) + 3k - 1) we have:

$$\mathcal{C}_r(env) = \mathcal{C}_{1+n(3k-1)+k}(env)$$

Proof. ¿From Proposition 3 the only rule that sends some objects to the environment before the halting configuration is the rule r_k . From Corollary 1 this rule sends the objects X_{ijt} to the environment in the configuration $C_{1+n(3k-1)+k}$. Therefore, for each $r \forall r = 1 + n(3k - 1) < r < 1 + n(3k - 1) + k$, $C_r(env) = C_{1+n(3k-1)+k}(env)$ and $\forall r = 1 + n(3k - 1) + k < r < 1 + n(3k - 1) + 3k - 1$ $C_r(env) = C_{1+n(3k-1)+k}(env)$ concluding the proof of a) and b).

In the following results we will prove that in each loop one object X_{ijt} is sent to the environment. If a loop exists that doesn't send any object X_{ijt} to the environment in all the following loops no more objects are sent to the environment. Therefore a configuration always exists from which any object X_{ijt} is not sent to the environment.

Firstly we prove that if in the k-th step of the loop the rule r_k is not possible to be applied then in the following loop it is not possible to apply this rule either. This is because the objects S_{ij} do not exist in the skin membrane.

Proposition 4. For each n $(0 \le n \le N-2)$ if the rule r_k cannot be applied in the configuration $C_{1+n(3k-1)+k-1}$, then it cannot be applied in the configuration $C_{1+(n+1)(3k-1)+k-1}$.

Proof. In order to apply the rule r_k it is necessary to have the objects ϵ_k , γ_q and α_{ijt} . According to Proposition 2 $\epsilon_k \in \mathcal{C}_{1+n(3k-1)+k-1}(N)$ for any n.

With the object γ_q only the rules r_k and r_{k+3} are applied, this object never disappears, so it always remains in the skin membrane.

The object α_{ijt} is produced by means of the rule r_u $(1 \le u \le k-1)$. In order to apply this rule it is necessary to have the object β_{u-1} and some object S_{ij} . The object β_{u-1} is produced by means of the rules $r'_1, r'_2, \ldots, r'_{u-1}$.

Therefore if in the configuration $C_{1+n(3k-1)+k}$ the rule r_k cannot be applied, it is because the object α_{ijt} does not exist, then the objects $S_{ij} \in C_{1+n(3k-1)}(N)$ do not exist.

¿From the configuration $C_{1+n(3k-1)+k-1}$ to the configuration $C_{1+(n+1)(3k-1)+k-1}$ in the skin membrane the only rule that can produce the objects S_{ij} is the rule r_{k+3} . To apply this rule, the objects C_{ij} are necessary, that are produced in the rule r_{k+1} from the objects S_{ij} . As the objects S_{ij} do not exist the rule r_{k+1} cannot be applied. 196 M. Cardona et al.

The following result proves that if the environment in the k-th step of the loop n + 1 is equal to the environment in the k-th step of the loop n then the environment is the same until the halting configuration.

Corollary 3. For each $n (0 \le n \le N - 2)$ if

$$\mathcal{C}_{1+n(3k-1)+k}(env) = \mathcal{C}_{1+(n+1)(3k-1)+k}(env)$$

then for each n' $(n \le n' \le N-2)$ we have

$$\mathcal{C}_{1+n(3k-1)+k}(env) = \mathcal{C}_{1+n'(3k-1)+k}(env)$$

Proof. We prove by induction that

$$\mathcal{C}_{1+(n+j)(3k-1)+k}(env) = \mathcal{C}_{1+(n+j+1)(3k-1)+k}(env) \quad \forall j (0 \le j \le N-n-3)$$

- The case base, j = 0, corresponds to the hypothesis of the corollary, so $C_{1+n(3k-1)+k}(env) = C_{1+(n+1)(3k-1)+k}(env).$
- We suppose true for the cases $0 \le j < N n 3$. Let us show that the result is true for j + 1.

By induction hypothesis we have

$$\mathcal{C}_{1+(n+j)(3k-1)+k}(env) = \mathcal{C}_{1+(n+j+1)(3k-1)+k}(env)$$

that is, in the previous configuration to these it has not been able to send any object to the environment, that is the rule r_k has not been possible to apply. By Proposition 4 if in the configuration $C_{1+(n+j+1)(3k-1)+k-1}$ cannot be applied the rule r_k in the configuration $C_{1+(n+j+2)(3k-1)+k-1}$ cannot be applied either. Therefore it is not possible to send any object to the environment and $C_{1+(n+j+1)(3k-1)+k}(env) = C_{1+(n+j+2)(3k-1)+k}(env)$.

We are going to prove that a loop always exists from any object X_{ijt} is sent to the environment, so it is not possible to apply the rule r_k .

Corollary 4. For each computation C there exists an unique object ν_C $(1 \le \nu_C \le N-2)$ such that in the configuration $C_{1+(\nu_C-1)(3k-1)+k}$ the rule r_k is applicable and in the configuration $C_{1+(\nu_C)(3k-1)+k}$ the rule r_k is not applicable.

Proof. By Proposition 4 and by Corollary 3 if in the configuration $C_{1+(\nu_{\mathcal{C}})(3k-1)+k-1}$ the rule r_k is not applicable, then for each j ($\nu_{\mathcal{C}} \leq j \leq N-2$) we have $C_{1+\nu_{\mathcal{C}}(3k-1)+k}(env) = C_{1+j(3k-1)+k}(env).$

Therefore, the rule r_k is not applicable in any configuration of the type $C_{1+j(3k-1)+k-1}$, $\forall j \quad \nu_{\mathcal{C}} \leq j \leq N-2$.

The following result allows us to give a meaning to the value t of the object X_{ijt} .

Proposition 5. Let C be an arbitrary computation of the P system and let the object $X_{i_n j_n t_{i_n j_n}}$ that is sent to the environment by the rule r_k in the configuration $C_{(k+1)+n(3k-1)-1}$. Then, we have

$$t_{i_n j_n}^{(n)} = \max\{t \mid S_{ij}^t \in \mathcal{C}_{1+n(3k-1)}(N), \ 1 \le i < j \le N\}$$

Proof. As the rule r_k is applicable in the configuration $C_{(k+1)+n(3k-1)-1}$ then $\alpha_{i_n j_n t_{i_n j_n}^{(n)}} \in C_{(k+1)+n(3k-1)-1}$. The object $\alpha_{i_n j_n t_{i_n j_n}^{(n)}}$ is obtained from the application of one of the rules $r_{k-t_{i_n j_n}^{(n)}}$ over the object $S_{ij}^{t_{i_n j_n}^{(n)}}$, where $t_{i_n j_n}^{(n)}$ is the maximum of the multiplicities of the objects S_{ij} . If another $t' > t_{i_n j_n}^{(n)}$ exists then the rule $r_{k-t'}$ will be applied and so the rule $r_{k-t_{i_n j_n}^{(n)}}$ has not been applied.

The following result proves that the maximum multiplicity to the objects S_{ij} pertaining to the skin membrane in any loop n is always greater or equal to the multiplicity the objects S_{ij} of the following loop n + 1.

Proposition 6. Let $w_n = \max\{t : S_{ij}^t \in C_{1+n(3k-1)}(N), 1 \le i < j \le N\}$, with $0 \le n \le N-2$. Then $w_n \ge w_{n+1}$, for each n.

Proof. If $w_n = \max\{t : S_{ij}^t \in \mathcal{C}_{1+n(3k-1)}(N)\}$, in the following configurations $\mathcal{C}_{1+n(3k-1)}$ the rule r_0 is applied successively and the rules with priority $r'_1, r'_2, \ldots, r'_{w_n-1}, r_{w_n}$ until arriving at the configuration $\mathcal{C}_{1+n(3k-1)+w_n}$. The object S_{ij} is not used in the rules $r'_1, r'_2, \ldots, r'_{w_n-1}$. For Proposition 5 in the rule r_{w_n} is used the object S_{ij} that have the maximum multiplicity equal to w_n . By this rule the object S_{ij} is eliminated by the membrane labeled by N, therefore $w_n \geq \max\{t : S_{ij}^t \in \mathcal{C}_{1+n(3k-1)+w_n}(N)\}.$

From this configuration the rule r_0 is applied $k - w_n$ times until we obtain the object ϵ_k . In these configurations the objects S_{ij} do not evolve.

• If $w_n \neq 0$, then in the configuration $C_{1+n(3k-1)+k-1}$ when the rule r_{k+1} is applied the information of some objects S_{ij} is sent to the object C_{ij} and later this information is transformed in the object S_{ij} by means of the rule r_{k+3} . The rule r_{k+2} deletes some objects S_{ij} , so the multiplicity of these objects never increases, it is only possible to decrease. After that the rule r_0 is applied since to arrive at the configuration $C_{1+(n+1)(3k-1)}$.

So, $w_{n+1} = \max\{t : S_{ij}^t \in \mathcal{C}_{1+(n+1)(3k-1)}(N)\} \le w_n$.

• If $w_n = 0$, then the objects S_{ij} do not belong to the skin membrane and by Proposition 4 it is not possible to produce any object S_{ij} , so: $w_{n+1} = \max\{t : S_{ij}^t \in \mathcal{C}_{1+(n+1)(3k-1)}(N)\} = 0.$

Remark: According to Proposition 6 we obtain $t_1 \ge t_2 \ge \ldots \ge t_n$.

By the following result we show that if a loop goes out to the environment an object of the type X_{ijt} , then in the following loop the objects $S_{ij}, S_{i'j}, S_{ji'}$,

 $i' \notin \{i, j\}$ disappears from the skin membrane. That is, at the moment that two clusters $\{i, j\}$ are joined a new class i is formed and all the objects $S_{i'j'}$ that have subscript j disappear.

Proposition 7. Let C be an arbitrary computation of the P system. Let

$$X_{i_{1}j_{1}t_{i_{1}j_{1}}^{(1)}}, X_{i_{2}j_{2}t_{i_{2}j_{2}}^{(2)}}, \dots, X_{i_{n}j_{n}t_{i_{n}j_{n}}^{(n)}} \in \mathcal{C}_{1+n(3k-1)}(env), \text{ with } 1 \le n \le \nu_{\mathcal{C}}$$

If $S_{ij} \in \mathcal{C}_{1+n(3k-1)}(N)$ then

a) $(i, j) \notin \{(i_1, j_1), \dots, (i_n, j_n)\}.$ b) $\{i, j\} \in \{1, \dots, N\} - \{j_1, \dots, j_n\}.$

Proof. We prove the result by induction on n.

• For n=1.

If in the configuration C_k the rule r_k sends the object $X_{i_1j_1t_{i_1j_1}^{(1)}}$ to the environment, then by Proposition 6 in the configuration $C_{k-t_{i_1j_1}^{(1)}}$ with $1 \le t_{i_1j_1}^{(1)} < k$ the rule $r_{k-t_{i_1j_1}^{(1)}}$ has had to apply so the objects $S_{i_1j_1}$ have disappeared. Therefore the objects $S_{ij} \in C_{1+(3k-1)}(N)$ verify that $(i,j) \notin \{(i_1,j_1)\}$.

In the following configurations when we apply the rule r_{k+1} the pairs of objects $(S_{i_1p}, S_{j_1p}), (S_{i_1p}, S_{pj_1}), (S_{pi_1}, S_{pj_1})$ are transformed respectively to the objects $C_{i_1p}, C_{i_1p}, C_{pi_1}$, these objects do not have the subscript j_1 . After that the rule r_{k+2} is applied in order to eliminate the objects

 $S_{i_1p}, S_{j_1p}, S_{pi_1}, S_{pj_1} \in C_1$ that have not been eliminated in the previous configurations. By these rules all the objects S_{ij} with $\{i, j\} \cap \{i_1, j_1\} \neq \emptyset$ have disappeared.

After that when we apply the rule r_{k+3} the objects C_{ij} , $i_1 \in \{i, j\}$ are transformed in the objects S_{ij} , $i_1 \in \{i, j\}$. Therefore if the objects $S_{ij} \in C_{1+(3k-1)}$ then $(i, j) \notin \{(i_1j_1)\}, \{i, j\} \in \{1, \ldots, N\} - \{j_1\}$.

• Let us suppose the proposition holds for $1 \le n < \nu_{\mathcal{C}}$. Let us show that the the result is held for n + 1.

If the object $X_{i_{n+1}j_{n+1}t_{i_{n+1}j_{n+1}}^{(n+1)}} \in \mathcal{C}_{k+n(3k-1)}(env)$ by Proposition 6 in the configuration $\mathcal{C}_{k-t_{i_{n+1}j_{n+1}}^{(n+1)}+n(3k-1)}$ with $1 \leq t_{i_{n+1}j_{n+1}}^{(n+1)} < k$ the rule $r_{k-t_{i_{n+1}j_{n+1}}^{(n+1)}}$ has had to apply and the objects $S_{i_{n+1}j_{n+1}}$ have disappeared. Therefore the objects $S_{ij} \in \mathcal{C}_{1+(n+1)(3k-1)}(N)$ verify that $(i,j) \notin \{(i_{n+1},j_{n+1})\}$, by induction hypothesis $(i,j) \notin \{(i_1,j_1),\ldots,(i_n,j_n)\}$ then $(i,j) \notin \{(i_1,j_1),\ldots,(i_{n+1},j_{n+1})\}$.

In the following configurations when we apply the rule r_{k+1} the pairs of objects $(S_{i_{n+1}p}, S_{j_{n+1}p}), (S_{i_{n+1}p}, S_{pj_{n+1}}), (S_{pi_{n+1}}, S_{pj_{n+1}})$ are transformed respectively in the objects $C_{i_{n+1}p}, C_{i_{n+1}p}, C_{pi_{n+1}}$, these objects do not have the subscript j_{n+1} . After that the rule r_{k+2} is applied in order to eliminate the objects

 $S_{i_{n+1}p}, S_{j_{n+1}p}, S_{pi_{n+1}}, S_{pj_{n+1}} \in \mathcal{C}_{1+n(3k-1)}$ that have not been eliminated in the previous configurations, with these two rules all the objects S_{ij} with $\{i, j\} \cap$ $\{i_{n+1}, j_{n+1}\} \neq \emptyset$ have disappeared. When we apply the rule r_{k+3} the objects C_{ij} , $i_{n+1} \in \{i, j\}$ are transformed in the objects $S_{ij}, i_{n+1} \in \{i, j\}$. So if $S_{ij} \in \mathcal{C}_{1+(n+1)(3k-1)}(N)$ then $(i,j) \notin \{(i_{n+1}j_{n+1})\}, \{i,j\} \in \{1,\ldots,N\} - \{j_{n+1}\}.$ And by the induction hypothesis $S_{ij} \in \mathcal{C}_{1+(n+1)(3k-1)}(N)$ $(i,j) \notin \{(i_1j_1), \dots, (i_nj_n)\}, \{i,j\} \in \{1, \dots, N\} - \{j_1, \dots, j_n\}$ therefore $(i,j) \notin \{(i_1j_1), \dots, (i_{n+1}j_{n+1})\}, \{i,j\} \in \{1,\dots,N\} - \{j_1\dots, j_{n+1}\}$ concluding the proof of b). \square

In the following proposition we study how the multiplicities of the objects S_{ij} changed at the moment that two clusters joined.

Proposition 8. Let C be an arbitrary computation of the P system. Let us supposse that $X_{i_1j_1t_{i_1j_1}^{(1)}}, X_{i_2j_2t_{i_2j_2}^{(2)}}, \dots, X_{i_nj_nt_{i_nj_n}^{(n)}} \in \mathcal{C}_{1+n(3k-1)}(env)$ with $1 \le n \le \nu_{\mathcal{C}}$, and $t_{ii}^{(n)} = max\{t: S_{ii}^t \in \mathcal{C}_{1+n(3k-1)}\}$. Then,

- If i_n ∉ {i, j} then t⁽ⁿ⁺¹⁾_{ij} = t⁽ⁿ⁾_{ij}. That is, the multiplicity of the objects S_{ij} is the same in the configurations C_{1+n(3k-1)} and C_{1+(n+1)(3k-1)}.
 If 1 ≤ i_n < j_n (n+1)</sup>_{i_np} = min{t⁽ⁿ⁾_{i_np}, t⁽ⁿ⁾_{j_np}}. That is, the multiplicity of the objects S_{in}p corresponds to the minimum multiplicity of the objects $S_{i_np}, S_{j_np}.$
- If $1 \le i_n then <math>t_{i_n p}^{(n+1)} = \min\{t_{i_n p}^{(n)}, t_{p j_n}^{(n)}\}$. That is, the multiplicity of the object $S_{i_n p}$ corresponds to the minimum multiplicity of the objects $S_{i_n p}, S_{p j_n}.$
- If $1 \le p < i_n < j_n \le N$ then $t_{pi_n}^{(n+1)} = \min\{t_{pi_n}^{(n)}, t_{pj_n}^{(n)}\}$. That is, the multiplicity of the object S_{pi_n} corresponds to the minimum multiplicity of the objects $S_{pi_n}, S_{pj_n}.$

Proof. For the proof of Proposition 7 when we apply the rule r_{k+1} in the configuration $\mathcal{C}_{k+(n-1)(3k-1)}$ the pairs of objects $(S_{i_np}, S_{j_np}), (S_{i_np}, S_{pj_n}), (S_{pi_n}, S_{pj_n})$ are transformed respectively in the objects $C_{i_np}, C_{i_np}, C_{pi_n}$. Therefore the number of objects $C_{i_n p}, C_{i_n p}, C_{p i_n}$ are the same respectively of the pairs of the objects $(S_{i_np}, S_{j_np}), (S_{i_np}, S_{pj_n}), (S_{pi_n}, S_{pj_n})$. After that the rule r_{k+2} is applied in order to eliminate the objects $S_{i_np}, S_{j_np}, S_{pi_n}, S_{pj_n} \in \mathcal{C}_{1+(n-1)(3k-1)}$ that have not been eliminated in the previous configurations. So the multiplicity of the objects $C_{i_np}, C_{i_np}, C_{pi_n}$ is respectively equal to $\min\{t_{i_np}^{(n)}, t_{j_np}^{(n)}\}, \min\{t_{i_np}^{(n)}, t_{pj_n}^{(n)}\}$, and $\min\{t_{p_{i_n}}^{(n)}, t_{p_{j_n}}^{(n)}\}$. When we apply the rule r_{k+3} the objects C_{ij} are transformed in the objects S_{ij} .

Next, we define how the partition of the individuals is formed from the objects sent to the environment.

Definition 6. Given a computation C of the P system we denote the succession of partitions of the set of the individuals by Δ_0^C , Δ_1^C , ..., Δ_{θ}^C . These partitions are constructed recursively as follows:

The initial partition is formed by the initial individuals, $\Delta_0^{\mathcal{C}} = \{B_{a_1^1}^0, \ldots, B_{a_N^N}^0\} \text{ with } q_i^0 = i \ y \ B_i^0 = \{\omega_i\} \equiv \{i\}$

The partition $\Delta_1^{\mathcal{C}}$ is constructed from the object $X_{i_1j_1t_{i_1j_1}^{(1)}} \in \mathcal{C}_{k+1}(env)$ with $1 \leq i_1 < j_1 \leq N$ as follows:

As $\{q_0^1, \ldots, q_0^N\} = \{1, \ldots, N\}$ then $\{i_1, j_1\} \subseteq \{q_0^1, \ldots, q_0^N\}$. If $i_1 = q_0^u$, $j_1 = q_0^s$ with $1 \le u < s \le N$, then the new cluster is $B_{q_1^u}^1 = B_{q_0^u}^0 \cup B_{q_0^s}^0$ with $q_1^u = q_0^u$ and

$$B^{0}_{q^{s}_{0}} \notin \Delta^{\mathcal{C}}_{1} B^{1}_{l} = B^{0}_{l} \text{ for } l \in \{q^{1}_{0}, \dots, q^{N}_{0}\} - \{q^{u}_{0}, q^{s}_{0}\}$$

Then the new partition obtained is $\Delta_1^{\mathcal{C}} = \{B_{q_1}^1, \ldots, B_{q_1}^{N-1}\}$

In a recursive manner we obtain the partition $\Delta_{n+1}^{\mathcal{C}}$ as follows: *jFrom the objects* $X_{i_1j_1t_{i_1j_1}^{(1)}}, \ldots X_{i_nj_nti_nj_n^{(n)}} \in \mathcal{C}_{(k+1)+(n-1)(3k-1)}(env)$ the partition $\Delta_n^{\mathcal{C}} = \{B_{q_n^1}^n, \ldots, B_{q_n^{N-n}}^n\}$ have been obtained and in the configuration $\mathcal{C}_{(k+1)+n(3k-1)}$ the object $X_{i_{n+1}j_{n+1}t_{i_{n+1}j_{n+1}}^{(n+1)}}$ with $1 \leq i_{n+1} < j_{n+1} \leq N$ is sent to the environment.

By Proposition 7, $\{i_{n+1}, j_{n+1}\} \in \{1, \ldots, N\} - \{j_1, j_2, \ldots, j_n\}$ therefore $i_{n+1} < j_{n+1}$ and $\{i_{n+1}, j_{n+1}\} \subseteq \{q_n^1, \ldots, q_n^{N-n}\}$. By construction of the partition it is verified $\{q_n^1, \ldots, q_n^{N-n}\} \subset \{1, \ldots, N\}$ and we have $\{j_1, j_2, \ldots, j_n\} \cap \{q_n^1, \ldots, q_n^{N-n}\} = \emptyset$.

Let $i_{n+1} = q_n^u$, $j_{n+1} = q_n^s$ with $1 \le u < s \le N$, then the new cluster is $B_{q_{n+1}^u}^{n+1} = B_{q_n^u}^n \cup B_{q_n^s}^n$ with $q_{n+1}^u = q_n^u$ $B_{q_n^s}^n \notin \Delta_{n+1}^{\mathcal{C}}$ $B_l^{n+1} = B_l^n$ with $l \in \{q_n^1, \dots, q_n^{N-n}\} - \{q_n^u, q_n^s\}$ Then $\Delta_{n+1}^{\mathcal{C}} = \{B_{q_{n+1}^1}^{n+1}, \dots, B_{q_{n+1}^{N-n-1}}^{n+1}\}$

Theorem 1. Let C be an arbitrary computation of the P system. Let us suppose that $S_{ij}^{t_{ij}^{(n)}} \in C_{1+n(3k-1)}(N)$ and $S_{ij}^{t_{ij}^{(n)}+1} \notin C_{1+n(3k-1)}(N)$, with $1 \le n \le \nu_{\mathcal{C}}$. Then, $t_{ij}^{(n)}$ is the minimum similarity between any pair of individuals pertaining to $B_i^{n-1} \cup B_j^{n-1}$. That is, $t_{ij}^{(n)}$ corresponds to the aggregation index between the groups B_i^{n-1} and B_j^{n-1} :

$$t_{ij}^{(n)} = \min\{t': S_{i'j'}^{t'} \in \mathcal{C}_1(N), \quad S_{i'j'}^{t'+1} \notin \mathcal{C}_1(N): \ i', j' \in B_i^n \cup B_j^n\}$$

Proof. We prove the theorem by induction on n.

- For n = 0. By Proposition 1 if $S_{ij}^{t_{ij}^{(0)}} \in C_1(N)$ then $t_{ij}^{(0)}$ corresponds to the similarity between individuals i, j.
- Let us suppose it is certain for n with $1 \le n < \nu_{\mathcal{C}}$.
- Let us show that the theorem is held for n + 1. In the configuration $C_{(k+1)+(n-1)(3k-1)}$ the partition $\Delta_n^{\mathcal{C}} = \{B_{q_1}^n, \ldots, B_{q_n}^n, N\}$ is obtained and in the configuration $C_{(k+1)+n(3k-1)}$ the object $X_{i_{n+1}j_{n+1}t_{i_{n+1}j_{n+1}}}$ with $1 \leq i_{n+1} < j_{n+1} \leq N$ is sent to the environment, then:
 - For Proposition 8 if $i_{n+1} \notin \{i, j\}$ then $t_{ij}^{(n+1)} = t_{ij}^{(n)}$. By the induction hypothesis $t_{ij}^{(n)}$ is the aggregation index between the groups i, j and for the definition 6, $B_i^{n+1} = B_i^{n+1}$, $B_j^{n+1} = B_j^{n+1}$, then the theorem is true.
 - For Proposition 8 we have $t_{i_{n+1j}}^{(n+1)} = \min\{t_{i_{n+1j}}^{(n)}, t_{j_{n+1j}}^{(n)}\}$ and by induction hypothesis $t_{i_{n+1j}}^{(n)}, t_{j_{n+1j}}^{(n)}$ corresponds to the minimum similarity between any pair of individuals pertaining respectively to $B_{i_{n+1}}^{n-1} \cup B_j^{n-1}, B_{j_{n+1}}^{n-1} \cup B_j^{n-1}$. Therefore $t_{i_{n+1j}}^{(n+1)}$ is the minimum of these two similarities and $B_{i_{n+1}}^n = B_{i_{n+1}}^{n-1} \cup B_{j_{n+1}}^{n-1}$ then it is verified that the minimum similarity between any pair of individuals pertaining to $B_{i_{n+1}}^n \cup B_j^n$.

The following result proves that if in the configuration $C_{(k+1)+(n-1)(3k-1)}$ the object $X_{i_n j_n t_{i_n j_n}^{(n)}}$ goes out to the environment, then when we form the partition $\Delta_n^{\mathcal{C}}$ the similarities between two individuals obtained in each set of $\Delta_n^{\mathcal{C}}$ are always greater or equal than $t_{i_n j_n}^{(n)}$.

Proposition 9. Let C be an arbitrary computation of the P system. Let us suppose that in the configuration $C_{(k+1)+(n-1)(3k-1)}$, with $0 \le n \le \nu_{\mathcal{C}} - 1$, the object $X_{i_n j_n t_{i_n j_n}}^{(n)}$ is sent to the environment, and the new partition is $\Delta_n^{\mathcal{C}} = \{B_{q_n}^n, \ldots, B_{q_n}^n\}$. Then, the hierarchical index of the cluster $B_{i_n}^n$ is $t_{i_n j_n}^{(n)}$, and the hierarchical index of the rest of clusters $\Delta_n^{\mathcal{C}} - \{B_{i_n}^n\}$ is greater or equal to $t_{i_n j_n}^{(n)}$.

Proof. We prove this result by induction on n. For Definition 6, $\Delta_0^{\mathcal{C}} = \{\{\omega_1\}, \ldots, \{\omega_N\}\}$ and $f(\{\omega_i\}) = k$ with $1 \le i \le N$.

• For n = 1. $\Delta_1^{\mathcal{C}} = \{B_{q_1}^1, \ldots, B_{q_1}^{1_{n-1}}\}$ with $B_{i_1} = \{\omega_{i_1}, \omega_{j_1}\}$ and $B_j = \{\omega_j\}, \quad \forall j \neq i_1$. Then $f(B_j) = k$ with $j \neq i_1$ and $f(B_{i_1}) = t_{i_1j_1}^{(1)}$ because $s(\omega_{i_1}, \omega_{j_1}) = t_{i_1j_1}^{(1)} \leq k - 1$. 202 M. Cardona et al.

• We suppose that is true for any $1 \le n \le \nu_{\mathcal{C}}$. Let us show that the result holds for n + 1. In the configuration $\mathcal{C}_{(k+1)+n(3k-1)}$ the rule r_k sends the object $X_{i_{n+1}j_{n+1}t_{i_{n+1}j_{n+1}}}^{(n+1)}$ to the environment, where $t_{i_{n+1}j_{n+1}}^{(n+1)}$ is the similarity between the clusters i_{n+1} and j_{n+1} . By construction of the P system $t_{i_{n+1}j_{n+1}}^{(n+1)} \le t_n$ and the partition is $\Delta_{n+1}^{\mathcal{C}} = \{B_{q_{n+1}}^{n+1}, \dots, B_{q_{n+1}}^{n+1}\}$ for Definition 6: - If $i_{n+1}, j_{n+1} \in B_{q_n}^n$ then $B_{q_{n+1}}^{n+1} = B_{q_n}^n$ and for induction hypothesis $f(B_{q_{n+1}}^{n+1}) = t_{i_nj_n}^{(n)} \ge t_{i_{n+1}j_{n+1}}^{(n+1)}$. - If $B_{i_{n+1}}^{n+1} = B_{i_{n+1}}^n \cup B_{j_{n+1}}^n$ and $\delta(B_{i_{n+1}}^n, B_{j_{n+1}}^n) = t_{i_{n+1}j_{n+1}}^{(n+1)}$, then $f(B_{i_{n+1}}^{n+1}) = t_{i_{n+1}j_{n+1}}^{(n+1)}$.

Proposition 10. The P system $\Pi(P_{Nk})$ allows us to find a hierarchical clustering.

Proof. From the partition $\Delta_0^{\mathcal{C}}$, $\Delta_1^{\mathcal{C}}$, ..., $\Delta_{\theta}^{\mathcal{C}}$ according to Definition 6 we can obtain an indexed hierarchy P_0, P_1, \ldots, P_m with $1 \le m \le N-1$.

By Proposition 9 all partitions $\Delta_n^{\mathcal{C}}$ have a hierarchical index $t_n = t_{i_n j_n}^{(n)}$ and we denoted by $(\Delta_0^{\mathcal{C}}, t_0), (\Delta_1^{\mathcal{C}}, t_1), \dots, (\Delta_{\theta}^{\mathcal{C}}, t_{\theta})$ with

 $t_0 > t_1 = t_2 = \dots t_{p_1} > t_{p_1+1} = \dots = t_{p_2} > \dots > t_{p_m} = \dots = t_{\theta}.$

In order to construct the partition P_0, P_1, \ldots, P_m we do the following steps:

- $P_0 = \Delta_0 = \{\{\omega_1\}, \{\omega_2\}, \dots, \{\omega_N\}\}.$
- The partitions $\Delta_1, \ldots, \Delta_{p_1}$ have associated a hierarchical index equal to t_1 so $P_1 = \Delta_{p_1}$.
- The partitions $\Delta_{p_1+1}, \ldots, \Delta_{p_2}$ have associated a hierarchical index equal to t_{p_2} so $P_2 = \Delta_{p_2}$.
- And successively until we have one of the following situations:
 - if Δ_{θ} has a hierarchical index $t_{\theta} = k 1$ then $P_m = \Delta_{\theta} = \Omega$.
 - if Δ_{θ} has a hierarchical index $t_{\theta} < k 1$ then $P_{m-1} = \Delta_{\theta}$ and $P_m = \Omega$.

4 Conclusions

One of the central issues when we have a set of individuals characterized by a k-tuple is to obtain a cluster that allows us to group similar individuals.

In this paper we propose a non-deterministic P system with external output to obtain a hierarchical clustering. This P system gives one of the possible solutions to the problem. We give an efficient (semi-uniform) solution to the problem of clustering in the framework of the cellular computing with membranes. The solution is semi-uniform because for each matrix formed by the values of the individuals, a specific P system with external output is designed. The solution is efficient, because it is polynomial in order of the number of N individuals and of the number

of k characteristics. The amount of resources initially required to construct the system is quadratic in N and k.

The mechanisms of the formal verification of P systems are often a very hard task. So to have new examples is always interesting, in order to find systematic processes of formal verification in a model of computation oriented to machines, like the cellular model. The paper also provides a new example of formal verification of P systems designed to solve a problem, following a specific methodology valid in some cases as those considered in the paper.

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Simulating the Bitonic Sort on a 2D-mesh with P Systems

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Summary. This paper gives a version of the parallel bitonic sorting algorithm of Batcher, which can sort N elements in time $O(\log^2 N)$. When applying it to the 2D mesh architecture, two indexing functions are considered, row-major and shuffled row-major. Some properties are proved for the later, together with a correctness proof of the proposed algorithm. Two simulations with P systems are proposed and discussed. The first one uses dynamic communication graphs and follows the guidelines of the mesh version of the algorithm. The second simulation requires only symbol rewriting rules in one membrane.

1 Introduction

P systems, introduced in [19], are powerful computational models, with nondeterministic as well as parallel features. Deterministic P systems can be also considered, and the power of their parallel features compared against the power of other computational models which enjoy parallelism. Along this line we refer to previous work, which relates P systems with parallel networks of processors, functioning according to the SIMD paradigm (Single Instruction Multiple Data machines), in [8], [9], for shuffle-exchange networks, and in [6] for 2D mesh networks. The comparison was approached by designing P systems which simulate the functioning of a specific architecture, when solving a specific problem. In [7] the general features of this type of approach were abstracted, giving a "blueprint" for the design of a class of deterministic P sytems, with *dynamic communication* graphs, which simulate a given parallel architecture, functioning to implement a given algorithm.

Among the choices to be made for the problem to solve, the *static sorting* imposes itself, being a central theme in computer science. Although it is well known that comparison-based sorting algorithms require at least $O(N \log N)$ comparisons
to sort N items, performing many comparisons in parallel can reduce the sorting time. This paper analyses the bitonic sorting algorithm, one of the fastest parallel sorting algorithms where the sequence of comparisons is not data-dependent. The bitonic sorting network was discovered by Batcher [3], who also discovered the network for odd-even sort. These were the first networks capable of sorting N elements in time $O(\log^2 N)$. Stone [24] maps the bitonic sort onto a perfect-shuffle interconnection network, sorting N elements by using N processors in time $O(\log^2 N)$. Siegel [23] shows that bitonic sort can also be performed on the hypercube in time $O(\log^2 N)$. The shuffled row-major indexing formulation of bitonic sort on a mesh-connected computer is presented by Thompson and Kung [25]. They also show how the odd-even merge sort can be used with snakelike row-major indexing. Nassimi and Sahni [16] present a row-major indexed bitonic sort formulation for a mesh with the same performance as shuffled row-major indexing.

Static sorting algorithms have been developped and proposed also in the P systems area. Among the first approaches, made independently, we mention [2] and [4], [5]. The problem of sorting with P systems occupies Chapter 8, [1], of the monograph [10].

We analyze in this paper a version of the bitonic sorting algorithm of Batcher, and its implementation on the 2D mesh architecture. Section 2 introduces the mesh topology and the model of computation. In 2.2 we begin the formal study of indexing functions, and we stress their importance for the passing to a network architecture. (Some similar work has been done in [12] and [13], but we develop our own formalism.) In 2.3 we present the algorithm, and the main result, Theorem 1, whose Corrolary is the correctness proof of the algorithm. Other results in this subsection, like Lemma 3, and the Remarks, are subsequently used to prove assertions about the algorithm, and, in Section 3, about the simulations with P systems.

Section 3 is devoted to the presentation of two different simulations of the algorithm with P systems. The first simulation uses dynamic communication graphs, as in [7]. A generative approach to the sequence of graphs used to communicate values between the membranes is a novel feature. The second simulation, uses only one membrane, and symbol rewriting rules.

2 Preliminaries: The bitonic sort on the 2D-mesh

2.1 Model of Computation

The presentation of the bitonic sort on the 2D-mesh architecture is made here based mainly on the paper [25]. It is the same algorithm as in [21], but with more emphasis on the routings necessary to compare elements situated at greater distances on the mesh. Also, some restrictions imposed in [25], will be eliminated, or re-examined, since they were dictated by their explicit connection to the ILLIAC IV-type parallel computer. In general, our references to parallel machines/architectures will be at the level of generalization to be found for instance in [21]. We also found it useful to formalize properly some aspects related to the indexing function.

Let us assume as in [25] that we have a parallel computer with $N = n \times n$ identical processors, disposed in a 2D-mesh structure. A processor is connected to all of its four vertical or horizontal neighbors, except for the processors situated on the perimeter, which have at most two or three neighbors, as no "wrap-around connections" are permitted.

Another assumption is that it is a SIMD (Single Instruction Multiple Data) machine. During each time unit, a single instruction is executed by a set of processors. In what follows, only two processor registers and two instructions are needed. For inter-processor data moves, we will use a routing instruction which copies the value of a register to a register of a neighbor processor. The second instruction is the internal comparison between the values of the two registers of a processor.

We define t_R the time for one-unit distance routing step, and t_C the time required for one comparison step. Concurrent data movement is allowed, as long as it is in the same direction; also any number of parallel comparisons can be made simultaneously.

2.2 The sorting problem and indexing functions

We assume to have an indexing function on the processors that is a one-to-one mapping from $\{0, 1, \ldots, n-1\} \times \{0, 1, \ldots, n-1\}$ onto $\{0, 1, \ldots, N-1\}$ and that initially N integers are loaded in the N processors. Therefore the sorting problem is defined as moving the *j*th smallest element to the processor indexed by *j*, for all $j \in \{0, 1, \ldots, N-1\}$.

Let $I : \{0, 1, \dots, n-1\} \times \{0, 1, \dots, n-1\} \longrightarrow \{0, 1, \dots, N-1\}$ denote an indexing function. Two indexing schemes are the following:

- (i) Row-major indexing. This is illustrated in Figure 1, and we denote it by I = RM.
- (ii) Shuffled row-major indexing. This is illustrated in Figure 2, and we denote it by I = sRM.

In order not to make the notation cumbersome, we let the same letter, say i, stand for an integer in $\{0, 1, \ldots, n-1\}$, and for its binary representation as a string. For $n = 2^k$, as the case will be, i will be a binary string of length k. Whenever necessary, we complete with zeroes (obviously, to the left) to obtain strings of the same length. (When we refer to bits of such a string, we count from 1 to n, starting from right to left, such that the "first" bit will be that of the least significant digit, and so forth. However, when we write such a string, we will write it with bits numbered from right to left.)

Consider the following definitions:

Definition 1. The row-major indexing function RM is defined by RM(i, j) = ij, where in the right hand-side we have denoted by ij the string concatenation of the

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Fig. 1. Row-major indexing scheme for a 4×4 mesh

binary representations of the integers i and j, after they have been brought to the same length.

More precisely, for every k, we have the bijections

$$RM_k: \{0, 1, \dots, 2^k - 1\} \times \{0, 1, \dots, 2^k - 1\} \longrightarrow \{0, 1, \dots, 2^{2k} - 1\},\$$

defined by

$$RM_k(i_1i_2\cdots i_k, j_1j_2\cdots j_k) = i_1i_2\cdots i_kj_1j_2\cdots j_k$$

Let sh (from 'shuffle') stand generically for the family of bit-shuffle functions $sh_k: \{0, 1, \ldots, 2^{2k} - 1\} \longrightarrow \{0, 1, \ldots, 2^{2k} - 1\}$, defined by

$$sh_k(i_1i_2\cdots i_kj_1j_2\cdots j_k)=i_1j_1i_2j_2\cdots i_kj_k$$

This is a bijection, with the obvious inverse

$$ush_k(i_1j_1i_2j_2\cdots i_kj_k) = i_1i_2\cdots i_kj_1j_2\cdots j_k,$$

where ush comes from 'un-shuffle'. In the following we will drop the index k whenever it is clear from the context.

Definition 2. The shuffled row-major indexing function sRM is defined by sRM(i, j) = sh(ij), where in the right hand-side we have denoted by ij the string concatenation of the binary representations of the integers i and j, after they have been brought to the same length, and sh is the appropriate bit-shuffle function.

More precisely, for every k, we have the bijections

$$sRM_k: \{0, 1, \dots, 2^k - 1\} \times \{0, 1, \dots, 2^k - 1\} \longrightarrow \{0, 1, \dots, 2^{2k} - 1\},\$$

defined by

$$sRM_k(i_1i_2\cdots i_k, j_1j_2\cdots j_k) = sh_k(i_1i_2\cdots i_kj_1j_2\cdots j_k) =$$
$$= i_1j_1i_2j_2\cdots i_kj_k.$$



Fig. 2. Shuffled row-major indexing scheme for a 4×4 mesh

Lemma 1. For every $i, j_1, j_2 \in \{0, 1, ..., 2^k - 1\}$ with $j_1 < j_2$, we have that $sRM(i, j_1) < sRM(i, j_2)$. Analogously, for every $i_1, i_2, j \in \{0, 1, ..., 2^k - 1\}$ with $i_1 < i_2$, we have that $sRM(i_1, j) < sRM(i_2, j)$.

Proof. The proof is immediate, from the definition of sRM. \Box

Let us consider the following general definition:

Definition 3. We call indexing function on the $2^k \times 2^k$ mesh a bijection

 $I_k: \{0, 1, \dots, 2^k - 1\} \times \{0, 1, \dots, 2^k - 1\} \longrightarrow \{0, 1, \dots, 2^{2k} - 1\}.$

The problem of sorting on a $2^k \times 2^k$ mesh is obviously related to an indexing function *I*: given a family of values $(P_{ij})_{ij}$, to sort them means to sort the corresponding 'linear' family $(P_{I(i,j)})_{I(i,j)}$, i.e., to find the permutation σ_k of $\{0, 1, \ldots, 2^{2k} - 1\}$ such that $(P_{\sigma I(i,j)})_{\sigma I(i,j)}$ is ascending (or descending).

Furthermore, when designing or implementing sorting algorithms on the 2D mesh, through an indexing function I they will be translated into algorithms for sorting a linear set, $(P_{I(i,j)})_{I(i,j)}$. But, the linear version of the algorithm, for sorting say an array $\langle P_0, \ldots, P_{2^{2k}-1} \rangle$, has to be such that its translation into 2D-mesh operations be admissible. By this last word we mean to obey certain rules for the functioning of the 2D mesh as a network of processors. One such rule is the possibility of a processor to communicate only with its neighbours. Other rules may involve simultaneous communication with neighbours: a processor may communicate with two neighbours simultaneously, provided they are on the same line or column, and that, if the same register is involved, the old value is read and communicated while the new value is written. Still further rules may involve the parallel functioning of the network: communications in parallel may be allowed only if either only lines or only columns are involved at one parallel step.

The above mentioned restrictions can be formulated in a formal manner, leading to a notion of *good* indexing function, but this is beyond the scope of this paper. Let us just say for now, that, if the linear version of the algorithm performs a comparison between P_r and P_s , then $P_{I^{-1}(r)}$ and $P_{I^{-1}(s)}$ must be neighbours in the 2D mesh topology. Since the linear version of the algorithm is also parallel, whole pairs of families must be mapped by I^{-1} into adjacent families, and the last ones are naturally the adjacent lines or columns of the mesh. This justifies to a certain degree results present in this paper such as Lemma 3.

The choice of the indexing function $I_k = sRM_k$ provides the connection between the bitonic sorting algorithm as presented in [21] and the bitonic sorting network of [14] and Figure 3.

Let us also note that both RM_k^{-1} and sRM_k^{-1} are easy to compute. For a linear index $r = i_1 i_2 \cdots i_k j_1 j_2 \cdots j_k$,

$$RM_k^{-1}(r) = RM_k^{-1}(i_1i_2\cdots i_kj_1j_2\cdots j_k) =$$

= $(i_1i_2\cdots i_k, j_1j_2\cdots j_k) = (r \operatorname{div} 2^k, r \operatorname{mod} 2^k),$

and similarly for sRM_k^{-1} .

2.3 The bitonic sorting algorithm

In Batcher's bitonic sorting network [3] of order n, the input is a bitonic sequence a of n/2 increasing elements followed by n/2 decreasing elements. These two sequences are merged by first applying n/2 comparators to a_0 and $a_{n/2}$, a_1 and $a_{(n/2)+1}$, ... $a_{n/2}$ and a_{n-1} . This first-phase partitions the elements into two bitonic sequences of n/2 smaller elements and of n/2 larger elements. These two bitonic sequences are further sorted by applying two bitonic merging networks of size n/2 to each sequence. A bitonic sorting network for 16 elements appears in Figure 3.

Lemma 2. [3] Given a bitonic sequence $\langle a_1, a_2, \ldots, a_{2n} \rangle$ the following hold.

- 1. $d = \langle \min\{a_i, a_{n+i}\}_{i=1}^n \rangle = \langle \min\{a_1, a_{n+1}\}, \min\{a_2, a_{n+2}\}, \dots, \min\{a_n, a_{2n}\} \rangle$ is bitonic.
- 2. $e = \langle \max\{a_i, a_{n+i}\}_{i=1}^n \rangle = \langle \max\{a_1, a_{n+1}\}, \max\{a_2, a_{n+2}\}, \dots, \max\{a_n, a_{2n}\} \rangle$ is bitonic.
- 3. $\max(d) < \min(e)$.

By an abuse of notations, we shall refer to a sequence of processors as the sequence of integers stored in one designated register A of the processors at a certain moment. Similarly, we shall use min $/\max\{P_i, P_j\}$ meaning min $/\max\{P_i[A], P_j[A]\}$ and refer to such operations as a comparison and interchange of values between processors P_i and P_j .

We shall give a generic algorithm for Batcher's bitonic sorter on an array $\langle P_0, \ldots, P_{2^{2k}-1} \rangle$ of processors, independent of the indexing function used. The algorithm (as illustrated in Figure 3 for k = 2) will consist of 2k stages, numbered from 1 to 2k. After each Stage *i*, the sequence $\langle P_{2^i j}, \ldots, P_{2^i j+2^i-1} \rangle$ with



Fig. 3. A bitonic sorting network of size 16

 $0 \leq j \leq 2^{k-i}-1$ will be an ascending sequence for all j even, and a descending sequence, for all j odd.

Input: an array $\langle P_0, \ldots, P_{2^{2k}-1} \rangle$ of processors Output: the sequence $\langle P_0, \ldots, P_{2^{2k}-1} \rangle$ is ascending Stage(*i*) for $t \leftarrow i$ downto 1 do // compare processors with indices differing on bit tforall $j \leftarrow 0$ to $2^{2k-t} - 1$ in parallel do | if $2^t j$ div 2^i is even then order = ascending else order = descending Merge($2^t j, 2^t j + 2^t - 1, order$) end Bitonic-Sort

```
 \begin{vmatrix} \text{for } i \leftarrow 1 \text{ to } 2k \text{ do} \\ \lfloor \text{ Stage}(i) \end{vmatrix}
```

end

Algorithm 1: Bitonic sort on an array of 2^{2k} processors

Given a bitonic sequence of processors $\langle P_1, P_2, \ldots, P_{2n} \rangle$, by **Merge**(1, 2n, ascending) we mean an operation which yields the sequence:

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 $\langle \min\{P_1, P_{n+1}\}, \min\{P_2, P_{n+2}\}, \dots, \min\{P_n, P_{2n}\},$

 $\max\{P_1, P_{n+1}\}, \max\{P_2, P_{n+2}\}, \dots, \max\{P_n, P_{2n}\}\rangle.$

Analogously, a call to Merge(1, 2n, descending) produces

$$(\max\{P_1, P_{n+1}\}, \max\{P_2, P_{n+2}\}, \dots, \max\{P_n, P_{2n}\}, \dots)$$

$$\min\{P_1, P_{n+1}\}, \min\{P_2, P_{n+2}\}, \dots, \min\{P_n, P_{2n}\}\rangle$$

Theorem 1. After each Stage *i*, the sequence $\langle P_{2^ij}, \ldots, P_{2^ij+2^i-1} \rangle$, $0 \le j \le 2^{k-i} - 1$ will be an ascending sequence for all *j* even, and a descending sequence, for all *j* odd.

Proof. We shall reason by induction on i. For the base case i = 1 it is immediate that the statement holds. Now let the statement be true for i and show that is it also true for i + 1.

First, t = i + 1 and $0 \le j \le 2^{k-i-1} - 1$. The sequence S for the i + 1 case can be written as

$$S = \langle P_{2^{i+1}j}, \dots, P_{2^{i+1}j+2^{i-1}} \rangle =$$

$$\langle P_{2^{i}2j}, \dots, P_{2^{i}2j+2^{i-1}}, P_{2^{i}(2j+1)}, \dots, P_{2^{i}(2j+1)+2^{i-1}} \rangle$$

From the induction hypothesis, we have that the sub-sequence

$$S_1 = \langle P_{2^i 2j}, \dots, P_{2^i 2j+2^i-1} \rangle$$

is ascending as 2j is even for any j, and that

$$S_2 = \langle P_{2^i(2j+1)}, \dots, P_{2^i(2j+1)+2^i-1} \rangle$$

is descending as 2j+1 is odd for any j. Therefore, the whole sequence S is bitonic.

At this point we apply the **Merge** operation on S, and get $S' = S'_1 S'_2$. By Lemma 2 we have that S'_1 and S'_2 are both bitonic. Moreover, when doing an ascending merge, $\max(S'_1) < \min(S'_2)$ and when doing a descending merge, $\min(S'_1) > \max(S'_2)$. This ensures that the two sequences are relatively ordered and can be sorted independently in parallel.

For $1 \le t < i+1$ the **Merge** operations are the same as in a merging network. We note that for all $2^{i+1}j \le l < 2^{i+1}(j+1)$, l div $2^{i+1} = j$ and therefore all subsequent **Merge** operations for t < i+1 on these processors will have the same order as when t = i+1. \Box

Corollary 1. Given a sequence $\langle P_0, \ldots, P_{2^{2k}-1} \rangle$ of processors, Algorithm 1 is correct.

Proof. The proof is immediate by Theorem 1. At Stage k we have j = 0 and hence the sequence $\langle P_0, \ldots, P_{2^{2k}-1} \rangle$ is ascending. \Box

Lemma 3. Given a $2^k \times 2^k$ 2D-mesh indexed with the function sRM and using Algorithm 1, for any two processors $x = 2^t j + l$ and $y = 2^t j + l + 2^{t-1}$, with $0 \le l \le 2^{t-1} - 1$, $1 \le t \le i$, and $0 \le j \le 2^{k-t} - 1$, which compare and interchange values inside a call of the form Merge $(2^t j, 2^t j + 2^t - 1, order)$, the following hold:

- (i) the binary representations of x and y differ only on bit t;
- (ii) if t is even then x and y reside on the same vertical line of the mesh; if t is odd they are on the same horizontal line;
- (iii) the distance on the mesh between x and y is $2^{\lceil t/2 \rceil 1}$;
- (iv)all processors situated on the same line between x and y are involved in the same Merge operation (i.e., have indices between $2^t j$ and $2^t j + 2^t 1$).

Proof. (i) Since $x = 2^t j + l$ and $l \le 2^{t-1} - 1$, we have that l contributes to bits 1 to t-1 and that $2^t j$ contributes to bits t+1 to 2k. Therefore bit t of x is 0. Similarly, since $y = x + 2^{t-1}$, bit t of y is 1, and all other bits are the same as those of x.

(ii) We apply the 'un-shuffle' function to x and y and get $ush(x) = i_1j_1$ and $ush(y) = i_2j_2$. By the definition of sRM we have that the i is the row index, while j is the column index. From i) we have that x and y differ on bit t, and hence the following two cases hold: t is even and $i_1 \neq i_2$, $j_1 = j_2$, or t is odd and $i_1 = i_2$, $j_1 \neq j_2$. In the first case x and y are on the same column, and in the latter, they are on the same row.

(*iii*) Using the notations above, let us assume that t is even and $i_1 \neq i_2$, $j_1 = j_2$. If x and y differ on bit t, then i_1 and i_2 will differ on bit t/2, and therefore $|i_1 - i_2| = 2^{t/2-1}$. From *ii*) x and y are on the same line of the mesh and the distance between them is $|i_1 - i_2| = 2^{t/2-1}$. Similarly, when t is odd and $i_1 = i_2$, $j_1 \neq j_2$, we have that j_1 and j_2 differ on bit $\lceil t/2 \rceil$. As before, the distance between x and y is $2^{\lceil t/2 \rceil - 1}$.

(*iv*) Consider again the case t even and $i_1 \neq i_2$, $j_1 = j_2$. We have to show that for all numbers i with $i_1 \leq i \leq i_2$, we have $2^t j \leq sRM(i, j_1) \leq 2^t j + 2^t - 1$. But since $i_1 \leq i \leq i_2$, form Lemma 1, we have that $x \leq sRM(i, j_1) \leq y$, which concludes our proof as $2^t j \leq x$ and $y \leq 2^t j + 2^t - 1$. Analogously for t odd. \Box

2.4 Applying the bitonic sorting algorithm to the 2D-mesh

Thompson and Kung [25], and Orcutt [18] showed that Batcher's bitonic sorting algorithm can be applied to sorting on a mesh-connected parallel computer, once the indexing function is chosen. In [25] it is noted that a necessary condition for optimality is that a comparison-interchange on the *j*th bit be no more expensive than the (j + 1)th bit, for all *j*. From (*iii*) of Lemma 3 we have that the "shuffled row-major" indexing scheme satisfies such condition, and leads to a complexity of $(14(n-1) - 8\log n)t_R + (2\log^2 n + \log n)t_C$.

The algorithm for a 4×4 is illustrated below and in Figure 4, where by "well ordered" we reffer to the corresponding comparison directions from Figure 3.





t= 2





Stage 4



t= 4











Result



Fig. 4. Bitonic sorting algorithm applied on a 4×4 2D-mesh, using shuffled row-major indexing

Stage 1 Bitonic sort on pairs of adjacent 1×1 matrices by the comparison interchange indicated, result: "well ordered" 1×2 matrices. Time: $2t_R + t_C$.

Stage 2 Bitonic sort on 1×2 matrices, result: 2×2 matrices. Time: $4t_R + 2t_C$. Stage 3 Bitonic sort on 2×2 matrices, result: 2×4 matrices. Time: $8t_R + 3t_C$. Stage 4 Bitonic sort on the two 2×4 matrices. Time: $12t_R + 4t_C$.

At each stage of Algorithm 1, we have a comparison and interchange of values between two processors. We have seen in Lemma 3 that using the sRM indexing function, these two processors will sit on the same vertical or horizontal line of the mesh. In the cases when they are not directly connected, they will have to route their values through neighbour processors, residing on the shortest path between (i.e., the line of the mesh on which they are placed). At each Stage *i*, we will have a comparison and interchange between processors whose indices differs only on bit *t*, with $1 \le t \le i$. Keeping in mind the parallel structure of our machine, the merging operation becomes a merging of square or rectangular portions of the mesh. Therefore, using the sRM indexing, the **Merge** operation defined previously becomes a **Merge** operation on sub-arrays of processors situated on the same line of the mesh. We denote such operation **compare-interchange**.

For a better understanding of the way a call Merge $(2^t j, 2^t j + 2^t - 1, order)$ $(1 \leq i \leq 2^k, 1 \leq t \leq i, 0 \leq j \leq 2^{k-t} - 1)$ is translated to the $2^k \times 2^k$ mesh topology, we shall make the following observations:

Remark 1. The portion of the mesh will have dimensions $2^{t-\lceil t/2 \rceil} \times 2^{\lceil t/2 \rceil}$ (i.e., $2^{t-\lceil t/2 \rceil}$ rows and $2^{\lceil t/2 \rceil}$ columns). This is true since 2^t processors are involved in the **Merge** and from Lemma 3 the maximal length of the sub-arrays involved in the **Merge** situated on the same line is $2 \cdot 2^{\lceil t/2 \rceil - 1}$.

Remark 2. For t even, we have a merging of square portions of the mesh of size $2^{t/2} \times 2^{t/2}$ and the compare interchange operations are done between processors residing on the same column of the mesh, For t odd we have a merging of rectangular portions of the mesh of size $2^{\lceil t/2 \rceil - 1} \times 2^{\lceil t/2 \rceil}$, and the compare interchange operation are done between processors residing on the same row of the mesh.

Let us see what are the necessary routings for the case for t = 1 (the processors are directly connected since $2^{\lceil t/2 \rceil - 1} = 1$). Consider a call of the form **Merge**(x, x + 1, order) Let processors P_x and P_{x+1} have the two registers denoted by A and B. Then the first instruction performed is a routing from $P_x[A]$ to $P_{x+1}[B]$. Next, perform a comparison operation in processor P_{x+1} , and store the minimum/maximum in register B. Finally, route back to P_x the value of the B register of P_{x+1} , with a total time is $2t_R + t_C$. The pseudo-code is written below, where by **compare** $(P_{x+1}, ascending|descending)$ we understand an internal comparison in processor P_{x+1} , which places the minimal/maximal value in register B.

Input: index x and sorting order order **Output**: the sequence $\langle P_x, P_{x+1} \rangle$ is ordered w.r.t. order

route $(P_x[A], P_{x+1}[B])$ compare $(P_{x+1}, \text{ order})$ route $(P_{x+1}[B], P_x[A])$

Algorithm 2: Compare-interchange operation for adjacent processors

Let us now see what is the case when we have to merge an array a of 2^i processors situated on the same line of the mesh, indexed from 0 to $2^i - 1$, and such that $P_{a[j]}$ is neighbour with $P_{a[j+1]}$ for all $0 \leq j < 2^i - 1$. The basic idea is that we have to shift the values of the first half of the array in the B registers of the second half, perform a comparison operation in parallel in these processors, and then shift back the minimal/maximal values. Hence a total time of $2^i t_R + t_C$.

Input: array of indices a, integer i, and sorting order order **Output**: the sequence $\langle P_{a[0]}, P_{a[1]}, \ldots, P_{a[2^i-1]} \rangle$ is ordered w.r.t. order

```
compare-interchange(a, i, order)
    forall j \leftarrow 0 to 2^{i-1} - 1 in parallel do
       // route left one unit in the B registers
       route(P_{a[j]}[A], P_{a[j+1]}[B])
   for k \leftarrow 1 to 2^{i-1} - 1 do
        // shift the values to the second half of the array
        forall j \leftarrow 0 to 2^{i-1} - 1 in parallel do
        | route(P_{a[j+k]}[B], P_{a[j+1+k]}[B])
   forall j \leftarrow 2^{i-1} to 2^i - 1 in parallel do
        // compare internally
     compare(P_{a[j]}, order)
    for k \leftarrow 2^{i-1} - 1 downto 1 do
        // shift back the results
        forall j \leftarrow 0 to 2^{i-1} - 1 in parallel do
         [ route(P_{a[j+k+1]}[B], P_{a[j+k]}[B]) ]
    forall j \leftarrow 0 to 2^{i-1} - 1 in parallel do
       // final routing back in the A registers
       \mathbf{route}(P_{a[j+1]}[B], P_{a[j]}[A])
```

end

Algorithm 3: Compare-interchange operation for an array of neighbour processors situated on the same line of the mesh

3 Modeling with membranes

Given the embedded parallel structure of a P system, modeling a 2D-mesh is a natural and straightforward approach. In what follows, we will present two such systems.

3.1 A P system with dynamic communication of 2D-mesh type

The first P system we introduce is along the same general lines as the model proposed in [6]. For each processors P_i , $i \in \{0, 1, \ldots, 2^{2k}-1\}$ we will have an associated membrane which we denote i. The two registers A and B of each processors are coded by two different symbols, say a and b. The number of occurrences of a represents the value of the A register, and analogously for b. Similarly to tissue-like P systems, we will have a collection of elementary membranes, connected by certain graphs, at certain moments of their evolution in time. The graphs we will consider will be sub-graphs of the total graph of the 2D-mesh network, also sub-graphs of the identity graph of the 2D-mesh network.

Basically, we have to model:

- Patterns of specific internal processing in each processor: these will be modeled by symbol rewriting rules.
- Patterns of communication between processors.

In a slightly different manner from [8] or [9], we shall refer to the communication graph associated to a given architecture with the following conventions: the vertices of the graph are the processors, and the edges (in our case not oriented as communications between processors are both ways) are the network connections characteristic of the architecture.

In the case of the $2^k \times 2^k$ 2D-mesh with the sRM indexing function, let G_{total} be the underlying communication graph composed of all edges necessary to the architecture. We introduce the following notation for the set of vertices of G_{total} :

$$V(G_{total}) = \{0, 1, \dots, 2^{2k} - 1\}.$$

Hence, the set of edges is

$$E(G_{total}) = \{(sRM(i,j), sRM(i,j+1),) \mid 0 \le i \le 2^k - 1, 0 \le j \le 2^k - 2\} \bigcup$$
$$\{(sRM(i,j), sRM(i+1,j)) \mid 0 \le i \le 2^k - 2, 0 \le j \le 2^k - 1\}.$$

Note that at a certain step of the sorting algorithm not all edges are involved in communication. Therefore we shall call *active sub-graphs* of G_{total} those graphs containing only such edges. We introduce also the *identity* graph, with

$$V(Id) = \{0, 1, \dots, 2^{2k} - 1\},\$$
$$E(Id) = \{(sRM(i, j), sRM(i, j)) \mid 0 \le i \le 2^k - 1, 0 \le j \le 2^k - 1\}$$

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for modeling internal processing steps.

As in [6], the P system which we shall consider in the sequel, departs from the classical P systems in two respects:

- The connections between individual membranes of a P system, μ , which was a tree-like structure of membranes (see [19]), and which in tissue-like P systems becomes a graph structure, is now, a sequence of graphs.
- The rules of a P system, usually associated to *membranes*, will now be associated to *communication graphs* between membranes.
 - a) We simulate the internal computations performed by a subset of processors by the action of symbol or object rewriting rules, at work simultaneously inside the corresponding subset of membranes. We will associate such rules to the corresponding active subsets of *Id*.
 - b) We simulate the exchange of data performed by the processors with communication rules (symport/antiport rules) between membranes. The communication rules will be associated to the active sub-graphs of G_{total} .

In order to describe the evolution of a P system which simulates the behavior of the bitonic sorting algorithm in the 2D-mesh architecture, we will use pairs [graph, rules]. We have graph a sub-graph of G_{total} or Id and rules a mapping from the set of all edges of graph, E(graph), to the set of all symbol/object rewriting rules for routing or comparison operations.

Let R_{μ} be the finite sequence of pairs [graph, rules] which simulates Algorithm 1, such that: (i) if $E(graph) \subseteq E(Id)$ then its rules are rewriting rules; (ii) if $E(graph) \subseteq E(G_{total})$ then its rules are communication rules.

In order to give such a sequence, we have to closely follow Algorithm 1. In a very intuitive manner, for every $\mathbf{Stage}(i)$, $1 \le i \le 2k$, and for every comparison on bit $t, i \ge t \ge 1$ we will have a sequence of graphs. From Lemma 3, the **Merge** operations executed in parallel in Algorithm 1 involve disjoint sub-matrices of the mesh and have the same length, therefore they can also be executed in parallel when implementing them on a 2D-mesh or P system.

To be more precise, let us analyse a call of the form $\mathbf{Merge}(2^t j, 2^t j + 2^t - 1, order)$. From Remarks 1-2 we know that the dimensions of the sub-matrix of the mesh involved in the **Merge** are $2^{\lceil t/2 \rceil} \times 2^{t-\lceil t/2 \rceil}$. Hence the maximal sequence of processors situated on the same line which compare and interchange values in a **Merge** operation has length $2^{\lceil t/2 \rceil}$. Using the observations made on Algorithm 3, for each **Merge** operation, we will need a sequence of $2^{\lceil t/2 \rceil} + 1$ graphs. The first $2^{\lceil t/2 \rceil - 1}$ route values in the destination membranes for comparison, then we have an application of the identity graph Id for internal comparisons, and another sequence of $2^{\lceil t/2 \rceil - 1}$ graphs to route back the results. Another important aspect is that for a comparison on bit t, the processors which compare values are the same at every stage, only that the order is different. Therefore, we will have the same communication graphs for routing operations, only that the pair [Id, rules] will be different at each **Stage**(i).

Let us denote as below the projection of the first and second argument of sRM^{-1} . These represent the row and column indices, respectively, of a processors indexed with $r \in \{0, 1, \ldots, 2^{2k} - 1\}$.

$$sRM_{row}^{-1}, sRM_{col}^{-1}: \{0, 1, \dots, 2^{2k} - 1\} \to \{0, 1, \dots, 2^k - 1\}$$

Then, the right / down neighbors of r (defined whenever possible) are:

$$\begin{aligned} right(r) &= sRM(sRM_{row}^{-1}(r), sRM_{col}^{-1}(r) + 1) \\ down(r) &= sRM(sRM_{row}^{-1}(r) + 1, sRM_{col}^{-1}(r)) \end{aligned}$$

In order to give an algorithm independent of the parity of bit t, denote (whenever possible):

$$next_t(r) = \begin{cases} right(r), & \text{if } t \text{ odd;} \\ down(r), & \text{if } t \text{ even.} \end{cases}$$

Remark 3. The indices of the first processors on every line l (i.e., the smallest indices on every line l) in a **Merge** $(2^t j, 2^t j+2^t-1, order)$ are $sRM(sRM_{row}^{-1}(2^t j)+l, sRM_{col}^{-1}(2^t j))$, with $0 \le l \le 2^{t-\lceil t/2 \rceil} - 1$.

Consider two adjacent processors P_x and P_y which need to interchange values. The three possible routing operations are: **route** $(P_x[A], P_y[B])$, **route** $(P_x[B], P_y[B])$, **route** $(P_x[B], P_y[A])$. The implementation with rewriting and communication rules of the first operation follows the lines: rewrite $a \to a^*$ into membrane x, apply the communication rule (a^*, out) along the edge (x, y), which transports all the a^* symbols from membrane x into y, and then in membrane y rewrite a^* back to the desired symbol, in this case $a^* \to b$. We give below a specification of a sequence [graph, rules] accomplishing this routing operation.

$$[Id_1, rules_1], [G, rules], [Id_2, rules_2], \text{ such that}$$
(rAB)

$$Id_1 \subseteq Id, (x, x) \in E(Id_1), rules_1((x, x)) = \{a \to a^*\},$$

$$G \subseteq G_{total}, (x, y) \in E(G), rules((x, y)) = \{(a^*, out)\},$$

$$Id_2 \subseteq Id, (y, y) \in E(Id_2), rules_2((y, y)) = \{a^* \to b\}.$$

Similarly, an operation $\mathbf{route}(P_x[B], P_y[A])$ is specified as:

$$[Id_1, rules_1], [G, rules], [Id_2, rules_2], \text{ such that}$$
(rBA)

$$Id_1 \subseteq Id, (x, x) \in E(Id_1), \ rules_1((x, x)) = \{b \to b^*\},$$

$$G \subseteq G_{total}, (x, y) \in E(G), \ rules((x, y)) = \{(b^*, out)\},$$

$$Id_2 \subseteq Id, (y, y) \in E(Id_2), \ rules_2((y, y)) = \{b^* \to a\}.$$

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In the case of a **route** $(P_x[B], P_y[B])$, only one communication graph is needed. The reason for not having supplementary rewritings is that such routings are done in parallel. The value from $P_x[B]$ is routed to $P_y[B]$ in parallel with the routing of $P_y[B]$ to a *B* register of a neighbor processors. Hence the number of symbols *b* in membrane *y* is the desired one $P_x[B]$.

$$[G, rules], \text{ such that}$$
(rBB)
$$G \subseteq G_{total}, (x, y) \in E(G), \ rules((x, y)) = \{(b, out)\}.$$

Consider now an internal comparison operation in processor P_x , **compare** $(P_x$, order) which places $\max(P_x[A], P_x[B])$ in register B if the order is ascending, or in register A if the order is descending. This can be formalised as:

$$[Id', rules], \text{ such that } Id' \subseteq Id, (x, x) \in E(Id'),$$
(C)
$$rules((x, x)) = \begin{cases} \{ab \to ab, a \to b, b \to b\}, & \text{if order is ascending,} \\ \{ab \to ab, a \to a, b \to a\}, & \text{if order is descending.} \end{cases}$$

For all $s = 0, 2^{\lceil t/2 \rceil - 1} - 1$ denote with G_s^t (sub-graphs of G_{total}) the communication graphs which simulate all parallel routing operations when comparing of bit t, in Algorithm 1.

From the above considerations and the steps illustrated in Algorithm 3, we introduce the following algorithms: Algorithm 4 to generate the edges of a communication graph, and Algorithm 5 to generate sub-graphs of the Id where comparisons are to be performed:

```
Input: integers k, t

Output: communication graphs G_s^t, for all s = 0, 2^{\lceil t/2 \rceil} - 1

set all E(G_s^t) \leftarrow \emptyset

for j \leftarrow 0 to 2^{2k-t} - 1 do

// for every Merge operation

for l \leftarrow 0 to 2^{t-\lceil t/2 \rceil} - 1 do

// for every line in the Merge operation

for s \leftarrow 0 to 2^{\lceil t/2 \rceil - 1} - 1 do

// for each communication graph

node = sRM(sRM_{row}^{-1}(2^tj) + l, sRM_{col}^{-1}(2^tj) + s)

for q \leftarrow 1 to 2^{\lceil t/2 \rceil - 1} do

// add the 2^{\lceil t/2 \rceil - 1} edges

E(G_s^t) \leftarrow E(G_s^t) \cup \{(\text{node, } next_t(\text{node}))\}

node = next_t(\text{node})
```

Algorithm 4: Generating all communication graphs G_s^t to compare on bit t

```
 \begin{array}{l} \textbf{Input: integers } k,t \\ \textbf{Output: internal processing graphs } Id^t \\ \textbf{set all } E(Id^t) \leftarrow \emptyset \\ \textbf{for } j \leftarrow 0 \textbf{ to } 2^{2k-t} - 1 \textbf{ do} \\ \hline // \textbf{ for every Merge operation} \\ \textbf{for } l \leftarrow 0 \textbf{ to } 2^{t-\lceil t/2 \rceil} - 1 \textbf{ do} \\ \hline // \textbf{ for every line in the Merge operation} \\ \textbf{node} = sRM(sRM_{row}^{-1}(2^tj) + l, sRM_{col}^{-1}(2^tj) + 2^{\lceil t/2 \rceil - 1}) \\ \textbf{for } q \leftarrow 1 \textbf{ to } 2^{\lceil t/2 \rceil - 1} \textbf{ do} \\ \hline E(Id^t) \leftarrow E(Id^t) \cup \{(\textbf{node, node})\} \\ \textbf{node} = next_t(\textbf{node}) \end{array}
```

Algorithm 5: Generating internal processing graphs Id^t to compare on bit t

Let us denote with G_{μ} the sequence of graphs simulating the algoritm. Then G_{μ} can be obtained also algorithmically, using Algorithm 6:

```
set G_{\mu} \leftarrow \lambda

for i \leftarrow 1 to 2k do

// compare interchange on bit t

for t \leftarrow i downto 1 do

// route to the second half

for s \leftarrow 0 to 2^{\lceil t/2 \rceil - 1} - 1 do

\lfloor G_{\mu} \leftarrow G_{\mu} \cdot G_{s}^{t}

// compare internally

G_{\mu} \leftarrow G_{\mu} \cdot Id^{t}

// route back to the first half

for s \leftarrow 2^{\lceil t/2 \rceil - 1} - 1 downto 0 do

\lfloor G_{\mu} \leftarrow G_{\mu} \cdot G_{s}^{t}
```

Algorithm 6: Generating the sequence of graphs G_{μ} for simulating the bitonic sorting algoritm on the $2^k \times 2^k$ 2D mesh

where by λ we denote the empty sequence, and by "." we denote the concatenation of two sequences.

The above algorithms could be easily modified to produce a the finite sequence R_{μ} of pairs [graph, rules] which simulates Algorithm 1. Keeping in mind the way routing and comparison operations are transformed into communication and rewriting rules of a P system (rAB, rBA, rBB, C), every time when adding an edge (x, y) to a graph G (subgraph of G_{total} or Id), the appropriate image rules((x, y)) should be specified.

3.2 Bitonic sorting in one membrane

We propose here a simulation of the bitonic sorting, which uses only one membrane. We will use (cooperative) symbol rewriting rules. The cooperation will be 'minimal', i.e., of degree two, since we follow closely the algorithm, and thus the whole process is based on comparators.

Consider an alphabet with 2^{2k} symbols, $V = \{v_0, v_1, \cdots, v_{2^{2k}-1}\}$. We will call it the *primary alphabet*.

We will consider also *auxiliary* alphabets, which we will specify in the sequel, in order to achieve sorting by rewritings.

We want to sort in ascending order the sequence of *distinct* integers

$$\langle x_0, x_1, \cdots x_{2^{2k}-1} \rangle,$$

codified over V as the multiset

$$w = v_0^{x_0} v_1^{x_1} \cdots v_{2^{2k}-1}^{x_{2^{2k}-1}}$$

We want to design a P system which, by rewritings acting in a maximal parallel manner and competing for objects, produces, from the initial configuration w, the configuration

$$w_f = v_0^{\sigma(x_0)} v_1^{\sigma(x_1)} \cdots v_{2^{2k}-1}^{\sigma(x_{2^{2k}-1})},$$

where σ is the permutation which yields the total order, i.e., such that $\sigma(x_0) < \sigma(x_1) < \cdots < \sigma(x_{2^{2k}-1})$.

Consider the alphabet V as ordered, by the natural order given by the indices, and let $v = v_0v_1 \cdots v_{2^{2k}-1}$ be the *alphabet word* (see [1]), i.e., the word obtained by concatenating the letters of V in their natural order. We call *extended alphabet words* over V, all words in V^{*} in which all the letters appear in their natural order. Note that both w and w_f , the initial and the final configuration of our P system, are extended alphabet words. Actually, all the intermediate configurations over V will be of this type.

Let $M_i(u)$ denote the multiplicity of letter v_i in a word $u \in V^*$. Then

$$w = v_0^{x_0} v_1^{x_1} \cdots v_{2^{2k}-1}^{x_{2^{2k}-1}} = v_0^{M_0(w)} \cdots v_{2^{2k}-1}^{M_{2^{2k}-1}(w)}.$$

Consider first the case n = 2 (k = 0). We have 2 integers codified over $\{v_0, v_1\}$ as an extended alphabet word. Consider the auxiliary alphabets

- $\{a, b\}$, for writing sources of a comparator
- $\{c^+, d^+\}$, for writing targets of a \oplus -comparator
- $\{c^-, d^-\}$, for writing targets of a \ominus -comparator

Consider the rules:

$$C_{\oplus} = \{v_0 \to a, v_1 \to b\} \cup \{ab \to c^+d^+, a \to d^+, b \to d^+\} \cup \{c^+ \to v_0, d^+ \to v_1\}.$$

The first group rewrites all v_0 s to *a*s and v_1 s to *b*s, the second group performs the comparison and produces the ascending order, and the last group rewrites back into the original alphabet. We have the sequence of configurations

$$v_0^{x_0}v_1^{x_1} \to a^{x_0}b^{x_1} \to c^{+\min(x_0,x_1)}d^{+\max(x_0,x_1)} \to v_0^{\min(x_0,x_1)}v_1^{\max(x_0,x_1)}$$

Similarly, the rules:

$$C_{\ominus} = \{v_0 \to a, v_1 \to b\} \cup \{ab \to c^-d^-, a \to c^-, b \to c^-\} \cup \{c^- \to v_0, d^- \to v_1\},$$

achieve a descending comparator, generating the sequence of configurations

$$v_0^{x_0}v_1^{x_1} \to a^{x_0}b^{x_1} \to c^{-\max(x_0,x_1)}d^{-\min(x_0,x_1)} \to v_0^{\max(x_0,x_1)}v_1^{\min(x_0,x_1)}.$$

Lemma 4. On a two-letter alphabet, starting from an initial configuration $w = v_0^{x_0}v_1^{x_1}$, by applying rules in C_{\oplus} we obtain w_f such that $(M_i(w_f))_i$ is ascending, and by applying rules in C_{\oplus} we obtain w_f such that $(M_i(w_f))_i$ is descending. \Box

Note that rules C_{\oplus} simulate a **Merge**(0, 1, +), and C_{\ominus} a **Merge**(0, 1, -).

We now want to simulate a whole family of merge operations done in parallel.

We take 2 auxiliary alphabets, S^+ and S^- to codify sources of + or - comparators, and another pair, T^+ and T^- , to codify outputs (targets) of + or - comparators. We label them in a bijective correspondence with V.

$$S^{+} = \{s_0^{+}, \cdots s_{2^{2k}-1}^{+}\},$$
$$T^{+} = \{t_0^{+}, \cdots t_{2^{2k}-1}^{+}\},$$

and similarly for -. (For the time being, only 4 copies of the initial alphabet. We will probably need 4 different copies for every stage, in order to keep them independent.)

At Stage (1) we have to simulate Merge(2j, 2j + 1, order), for all $0 \le j \le 2^{2k-1} - 1$, where order = + for all j even, and order = - for all j odd.

This is equivalent to:

• Rewrite all symbols of V into start symbols for appropriate comparators, using the sets of rules

$$\{ v_{2j} \to s_{2j}^+, v_{2j+1} \to s_{2j+1}^+ \mid 0 \le j \le 2^{2k-1} - 1 , j \text{ even} \} \cup \\ \cup \{ v_{2j} \to s_{2j}^-, v_{2j+1} \to s_{2j+1}^- \mid 0 \le j \le 2^{2k-1} - 1 , j \text{ odd} \}.$$

• Apply in parallel the rewritings of symbols which correspond to the simulations of the comparators:

$$\{ s_{2j}^{+} s_{2j+1}^{+} \to t_{2j}^{+} t_{2j+1}^{+}, s_{2j}^{+} \to t_{2j+1}^{+}, s_{2j+1}^{+} \to t_{2j+1}^{+} \mid 0 \le j \le 2^{2k-1} - 1, j \text{ even} \} \bigcup \\ \cup \{ s_{2j}^{-} s_{2j+1}^{-} \to t_{2j}^{+} t_{2j+1}^{-}, s_{2j}^{-} \to t_{2j}^{-}, s_{2j+1}^{-} \to t_{2j}^{-} \mid 0 \le j \le 2^{2k-1} - 1, j \text{ odd} \}.$$

• Rewrite back all symbols of T's into V.

$$\{ v_{2j} \leftarrow t_{2j}^+, v_{2j+1} \leftarrow t_{2j+1}^+ \mid 0 \le j \le 2^{2k-1} - 1 \text{, j even} \} \cup \\ \cup \{ v_{2j} \leftarrow t_{2j}^-, v_{2j+1} \leftarrow t_{2j+1}^- \mid 0 \le j \le 2^{2k-1} - 1 \text{, j odd} \}.$$

The general scheme is as follows:

Input: an extended alphabet word w over V

Output: the extended alphabet word w_f over V, such that $\langle M_i(w_f) \rangle_i$ is ascending

Sim-Stage(i)

for $t \leftarrow i$ downto 1 do Take 4 extra copies of the start and the terminal alphabets, $S_t^+, S_t^-, T_t^+, T_t^+$, different for each value of t. For t's smaller than i we can re-use the alphabets of previous stages. forall $j \leftarrow 0$ to $2^{2k-t} - 1$ in parallel do if $2^t j$ div 2^i is even then order = ascending else order = descending // Simulate the calls Merge $(2^t j, 2^t j + 2^t - 1, order)$ (WF) Rewrite all symbols in V with the appropriate symbol in $S_t^+ \cup S_t^-$. (C) Apply the rewritings which simulate the appropriate comparators. (WB) Rewrite back all symbols in $T_t^+ \cup T_t^-$ to symbols of V.

 \mathbf{end}

```
Sim-Bitonic-Sort
```

```
for i \leftarrow 1 to 2k do
```

```
\_ Sim-Stage(i)
```

\mathbf{end}

Algorithm 7: Simulating bitonic sort on an alphabet of 2^{2k} letters V

The calls to $\mathbf{Merge}(2^t j, 2^t j + 2^t - 1, order)$ are equivalent to parallel calls to $\mathbf{Merge}(x, y, order)$, where x and y are like in Lemma 3. The same result ensures us that, both the rewritings which feed the comparators, and the rewritings which implement the comparators can be done in parallel. For $\mathbf{Merge}(x, y, order = -)$, we use

$$\{s_x \ \bar{s}_y \ \to t_x \ \bar{t}_y \ , s_x \ \to t_x \ , s_y \ \to t_x \]\}.$$

We propose the following sets of rules for simulating iteration t at **Sim-Stage**(i):

$$(WR) ewritings to S's, with * = \begin{cases} +, & \text{if } 2^t j \text{ div } 2^i \text{ is even,} \\ -, & \text{if } 2^t j \text{ div } 2^i \text{ is odd,} \end{cases}$$

$$\{ v_x \to s_x^* \in S_t^* \mid x \in [2^t j, 2^t j + 2^{t-1}), 0 \le j \le 2^{2k-t+1} - 1 \}.$$

(C)Rewritings which simulate the comparators, for appropriate pairs of indices:

$$\begin{split} &\{s_x^+ s_y^+ \to t_x^+ t_y^+, s_x^+ \to t_y^+, s_y^+ \to t_y^+ \mid \\ &x \in [2^t j, 2^t j + 2^{t-1}), \; y = x + 2^{t-1}, \; 0 \leq j \leq 2^{2k-t} - 1\}, \\ &\{s_x^- s_y^- \to t_y^- t_x^-, s_x^- \to t_x^-, s_y^- \to t_x^- \mid \\ &x \in [2^t j, 2^t j + 2^{t-1}), \; y = x + 2^{t-1}, \; 0 \leq j \leq 2^{2k-t} - 1\}. \end{split}$$

(WR) ewritings from T's:

$$\{v_x \leftarrow t_x^* \in T_t^* \mid x \in [2^t j, 2^t j + 2^{t-1}), 0 \le j \le 2^{2k-t+1} - 1\}.$$

4 Conclusions and open problems

We have presented a bitonic sorting algorithm which can be implemented on a 2D mesh of processors. The dependence between its performance and the choice of the indexing function still remains to be fully explored. However, we believe that we have proved some results which explain the choice of sRM as a "good" indexing function.

We have not yet found in the literature a formal proof of the correctness of bitonic sorting, an equivalent, or an analogue of our Theorem 1.

Much work remains to be done concerning the proposed simulations with P systems. The first simulation, derived in a "straightforward" manner from the functioning of the algorithm on the mesh, is inspired from work in [6], [8], [9], and [7], where the general framework was abstracted. It introduces a generative approach to the sequence of communication graphs, a feature to be explored in subsequent work. The second one is at the opposite pole: it requires no routings of values at all, just an appropriate codification of the symbols. It is in this area that other versions of the algorithm could be implemented, independent of the topology of a given structure, and the parallel features of the P systems can be compared against those of other computational devices.

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On the Number of Agents in P Colonies

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Summary. We continue the investigation of P colonies introduced in [7], a class of abstract computing devices composed of independent agents, acting and evolving in a shared environment.

We decrease the number of agents needed to computational completeness of P colonies with one and two objects inside each agent, respectively, owing some special restrictions to the type of programs. We characterize the generative power of the partially blind machine by the generative power of special P colonies.

1 Introduction

P colonies were introduced in paper [7] as formal models of a computing device inspired by membrane systems and by grammar systems called colonies. This model is inspired by structure and functioning of a community of living organisms in a shared environment.

The independent organisms living in a P colony are called agents. Each agent is represented by a collection of objects embedded in a membrane. The number of objects inside the agent is the same for each one of them. The environment contains several copies of a basic environmental object denoted by e. The number of the copies of e is unlimited.

A set of programs is associated with each agent. The program determines the activity of the agent by rules. In every moment all the objects inside of the agent are being evolved (by an evolution rule) or transported (by a communication rule). The third type of the rules used is a checking rule. This type of the rules sets the priority between two rules.

The computation starts in the initial configuration specified in the definition. Using their programs the agents change themselves and by the environment they can affect the behavior of the other agents. In each step of the computation, each agent with at least one applicable program nondeterministically chooses one of them and executes it. The computation halts when no agent can apply any of its programs. The result of the computation is the number of some specific objects present at the environment at the end of the computation.

There are several different ways how to define the beginning of the computation. (1) At the beginning of computation the environment and all agents contain only copies of object e. (2) All the agents can contain various objects at the beginning of computation - the agents are in different initial states.

(3) The initial state of the environment is nonempty - the environment contains initial "parameters" for future computation, the agents start with e-s.

In [4, 6, 7] the authors study P colonies with two objects inside the agents. In this case programs consist of two rules, one for each object. If the former of these rules is an evolution and the latter is a communication or checking, we talk about restricted P colonies. If we allow also another combination of the types of the rules, we obtain non-restricted P colonies. The restricted P colonies with the checking rules are computationally complete [3, 4]. Activities carried out in the field of membrane computing are currently numerous and they are available also at [11].

In the present paper we start with definitions in Section 2.

In Section 3 we will deal with P colonies with one object inside each agent. In recent paper [1] there was shown, that at most seven programs for each agent as well as five agents guarantees the computational completeness of these P colonies. In the preset paper we look for the generative power of P colonies with less than five agents. Two results are achieved in this direction. First, we show, that four agents are enough for computational completeness of P colonies. The second result gives a lower bound for the generative power the P colonies with two agents. Even a restricted variant of these P colonies is at least as powerful as the partially blind register machines.

Restricted P colonies are studied in Section 4. It is known that one agent is sufficient to obtain computational completeness of restricted P systems with checking rules. If no checking rules are used in the restricted P colonies then we need two agents to prove the universal computational power of those P colonies.

2 Definitions

Throughout the paper we assume the reader to be familiar with the basics of the formal language theory.

We use NRE to denote the family of the recursively enumerable sets of natural numbers. Let Σ be the alphabet. Let Σ^* be the set of all words over Σ (including the empty word ε). We denote the length of the word $w \in \Sigma^*$ by |w| and the number of occurrences of the symbol $a \in \Sigma$ in w by $|w|_a$.

A multiset of objects M is a pair M = (V, f), where V is an arbitrary (not necessarily finite) set of objects and f is a mapping $f : V \to N$; f assigns to each object in V its multiplicity in M. The set of all multisets with the set of objects V is denoted by V° . The set V' is called the support of M and denoted by supp(M) if for all $x \in V' f(x) \neq 0$ holds. The cardinality of M, denoted by |M|, is defined by $|M| = \sum_{a \in V} f(a)$. Any multiset of objects M with the set of objects $V' = \{a_1, \ldots a_n\}$ can be represented as a string w over alphabet V' with $|w|_{a_i} = f(a_i); \ 1 \leq i \leq n$. Obviously, all words obtained from w by permuting the letters can also represent the same M, and ε represents the empty multiset.

2.1 P colonies

We briefly recall the notion of P colonies. A P colony consists of agents and environment. Both the agents and the environment contain objects. With every agent the set of program is associated. There are two types of rules in the programs. The first type, called the evolution, is of the form $a \rightarrow b$. It means that object a inside of the agent is rewritten (evolved) to the object b. The second type of rules, called a communication, is in the form $c \leftrightarrow d$. When this rule is performed, the object c inside the agent and the object d outside of the agent change their places, so, after execution of the rule d appears inside the agent and c is placed outside of the agent.

In [6] the ability of agents is extended by checking rules. These rules give to the agents an opportunity to opt between two possibilities. They have form r_1/r_2 . If the checking rule is performed, the rule r_1 has higher priority to be executed as the rule r_2 has. It means that the agent checks the possibility to use rule r_1 . If it can be executed, the agent has to use it. If the first rule cannot be applied, the agent uses the second one.

Definition 1. The P colony of the capacity k is a construct $\Pi = (A, e, f, V_E, B_1, \dots, B_n)$, where

- A is an alphabet of the colony, its elements are called objects,
- $e \in A$ is the basic object of the colony,
- $f \in A$ is the final object of the colony,
- V_E is a multiset over $A \{e\}$,
- B_i, 1 ≤ i ≤ n, are agents, each agent is a construct B_i = (O_i, P_i), where
 O_i is a multiset over A, it determines the initial state (content) of the agent, |O_i| = k,
 - $P_i = \{p_{i,1}, \dots, p_{i,k_i}\}$ is a finite multiset of programs, where each program contains exactly k rules, which are in one of the following forms each:
 - $\cdot \quad a \rightarrow b, \ called \ an \ evolution \ rule,$
 - · $c \leftrightarrow d$, called a communication rule,
 - r_1/r_2 , called a checking rule; r_1, r_2 are an evolution or a communication rules.

An initial configuration of the P colony is an (n+1)-tuple of strings of objects present in the P colony at the beginning of the computation, it is given by O_i for $1 \leq i \leq n$ and by V_E . Formally, the configuration of P colony Π is given by (w_1, \ldots, w_n, w_E) , where $|w_i| = k$, $1 \leq i \leq n$, w_i represents all the objects placed inside the *i*-th agent and $w_E \in (A - \{e\})^*$ represents all the objects in the environment different from the object e.

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In the paper parallel model of P colonies will be studied. At each step of the parallel computation each agent tries to find one program to use. If the number of applicable programs is higher than one, the agent nondeterministically chooses one of them. At one step of computation the maximal possible number of agents are active.

Let the programs of each P_i be labeled in a one-to-one manner by labels in a set $lab(P_i)$ in such a way that $lab(P_i) \cap lab(P_j) = \emptyset$ for $i \neq j, 1 \leq i, j \leq n$.

To express derivation step formally we introduce following four functions for the agent using the rule r of program $p \in P$ with objects w in the environment:

For rule r being $a \to b, c \leftrightarrow d$ and $c \leftrightarrow d/c' \leftrightarrow d'$, respectively, and for multiset $w \in V^{\circ}$ we define:

$left \left(a \to b, w \right) = a$	$left\left(c\leftrightarrow d,w\right) =\varepsilon$
$right (a \to b, w) = b$	$right \left(c \leftrightarrow d, w \right) = \varepsilon$
$export \ (a \to b, w) = \varepsilon$	$export\left(c\leftrightarrow d,w\right) =c$
$import(a \to b, w) = \varepsilon$	$import\left(c\leftrightarrow d,w\right) =d$

$$\begin{array}{l} left \left(c \leftrightarrow d/c' \leftrightarrow d', w \right) = \varepsilon \\ right \left(c \leftrightarrow d/c' \leftrightarrow d', w \right) = \varepsilon \\ export \left(c \leftrightarrow d/c' \leftrightarrow d', w \right) = c \\ import \left(c \leftrightarrow d/c' \leftrightarrow d', w \right) = d \end{array} \right\} \text{ for } |w|_d \ge 1$$

 $\begin{array}{l} export\left(c\leftrightarrow d/c'\leftrightarrow d',w\right)=c'\\ import\left(c\leftrightarrow d/c'\leftrightarrow d',w\right)=d' \end{array} \right\} \text{for } |w|_{d}=0 \text{ and } |w|_{d'}\geq 1 \end{array}$

For a program p and any $\alpha \in \{left, right, export, import\}$, let $\alpha(p, w) = \bigcup_{r \in p} \alpha(r, w).$

A transition from a configuration to another is denoted as

 $(w_1, \ldots, w_n; w_E) \Rightarrow (w'_1, \ldots, w'_n; w'_E)$, where the following conditions are satisfied:

- There is a set of program labels P with $|P| \leq n$ such that
 - $p, p' \in P, p \neq p', p \in lab(P_j)$ implies $p' \notin lab(P_j)$,
 - for each $p \in P$, $p \in lab(P_j)$, $left(p, w_E) \cup export(p, w_E) = w_j$, and $\bigcup_{p \in P} import(p, w_E) \subseteq w_E$.
- Furthermore, the chosen set P is maximal, that is, if any other program $r \in \bigcup_{1 \leq i \leq n} lab(P_i), r \notin P$, is added to P, then the conditions above are not satisfied.

Now, for each $j, 1 \leq j \leq n$, for which there exists a $p \in P$ with $p \in lab(P_j)$, let $w'_j = right(p, w_E) \cup import(p, w_E)$. If there is no $p \in P$ with $p \in lab(P_j)$ for some $j, 1 \leq j \leq n$, then let $w'_j = w_j$ and moreover, let $w'_E = w_E - \prod import(p, w_E) \cup \prod export(p, w_E)$.

$$w'_{E} = w_{E} - \bigcup_{p \in P} import(p, w_{E}) \cup \bigcup_{p \in P} export(p, w_{E}).$$

A configuration is halting if the set of program labels P satisfying the conditions above cannot be chosen to be other than the empty set. A set of all possible halting configurations is denoted by H. With a halting computation we can associate a result of the computation. It is given by the number of copies of the special symbol f present in the environment. The set of numbers computed by a P colony Π is defined as

$$N(\Pi) = \left\{ |v_E|_f \mid (w_1, \dots, w_n, V_E) \Rightarrow^* (v_1, \dots, v_n, v_E) \in H \right\},$$

where (w_1, \ldots, w_n, V_E) is the initial configuration, (v_1, \ldots, v_n, v_E) is a halting configuration, and \Rightarrow^* denotes the reflexive and transitive closure of \Rightarrow .

Given a P colony $\Pi = (A, e, f, V_E, B_1, \ldots, B_n)$ the maximal number of programs associated with the agents in P colony Π is called the height of P colony Π . The degree of P colony Π is the number of agents in P colony Π . The third parameter characterizing a P colony is the capacity of P colony Π describing the number of the objects inside each agent.

Let us use the following notations:

 $NPCOL_{par}(k, n, h)$ for the family of all sets of numbers computed by P colonies working in parallel, using no checking rules and with:

- the capacity at most k,
- the degree at most n and
- the height at most h.

If we allow checking rules the family of all sets of numbers computed by P colonies is denoted by $NPCOL_{par}K$. If the P colonies are restricted, we replace the denotation to $NPCOL_{par}R$ and $NPCOL_{par}KR$, respectively.

2.2 Register machines

In this paper we want to characterize the size of the families $NPCOL_{par}(k, n, h)$ comparing them with the recursively enumerable sets of numbers. To achieve this aim we use the notion of a register machine.

Definition 2. [8] A register machine is the construct $M = (m, H, l_0, l_h, P)$ where: - m is the number of registers,

- H is the set of instruction labels,
- l_0 is the start label, l_h is the final label,
- *P* is a finite set of instructions injectively labeled with the elements from the set *H*.

The instruction of the register machine are of the following forms:

- $l_1: (ADD(r), l_2, l_3)$ Add 1 to the content of the register r and proceed to the instruction (labeled with) l_2 or l_3 .
- $l_1: (SUB(r), l_2, l_3)$ If the register r stores the value different from zero, then subtract 1 from its content and go to instruction l_2 , otherwise proceed to instruction l_3 .
- $l_h: HALT$ Stop the machine. The final label l_h is only assigned to this instruction.

Without loss of generality, one can assume that in each ADD-instruction l_1 : $(ADD(r), l_2, l_3)$ and in each conditional SUB-instruction l_1 : $(SUB(r), l_2, l_3)$ the labels l_1, l_2, l_3 are mutually distinct.

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The register machine M computes a set N(M) of numbers in the following way: it starts with all registers empty (hence storing the number zero) with the instruction labeled l_0 and it proceeds to apply the instructions as indicated by the labels (and made possible by the contents of registers). If it reaches the halt instruction, then the number stored at that time in the register 1 is said to be computed by Mand hence it is introduced in N(M). (Because of the nondeterminism in choosing the continuation of the computation in the case of ADD-instructions, N(M) can be an infinite set.) It is known (see e.g.[8]) that in this way we can compute all sets of numbers which are Turing computable.

Moreover, we call a register machine partially blind [5], if we interpret a subtract instruction in the following way: $l_1 : (SUB(r); l_2; l_3)$ - if in register r there is value different from zero, then subtract one from its contents and go to instruction l_2 or to instruction l_3 ; if in register r there is stored zero when attempting to decrement register r, then the program ends without yielding a result.

When the register machine reaches the final state, the result obtained in the first register is only taken into account if the remaining registers store value zero. The family of sets of non-negative integers generated by partially blind register machines is denoted by NRM_{pb} . The partially blind register machine accepts a proper subset of NRE.

3 P colonies with one object inside the agent

In this Section we analyze the behavior of P colonies with only one object inside each agent of P colonies. This gives that every program is formed by only one rule, either an evolution or a communication.

If all the agents have their programs with evolution rules, the agents "live only for themselves" and do not communicate with the environment.

In [1] following results was proved:

 $-NPCOL_{par}K(1,*,7) = NRE.$

 $-NPCOL_{par}K(1,5,*) = NRE$

The number of agents in the second result can be decreased:

Theorem 1. $NPCOL_{par}K(1, 4, *) = NRE$

Proof. We construct a P colony simulating the computation of the register machine. Because there are only copies of e in the environment and inside the agents, we have to initialize a computation by generating initial label l_0 . After generating symbol l_0 this agent stops and it can start its activity only by using a program with communicating rule. Two agents will cooperate in order to simulate the ADD and SUB instructions.

Let us consider an *m*-register machine $M = (m, H, l_0, l_h, P)$ and present the content of the register *i* by the number of copies of a specific object a_i in the environment. We construct the P colony $\Pi = (A, e, f, \emptyset, B_1, \ldots, B_4)$ with:

$$\begin{array}{l} - \text{ alphabet } A = \{l, l' | l \in H\} \cup \\ \cup \{E_i, E'_i, F_i, F'_i, F''_i \mid \text{ for each } l_i \in H\} \cup \\ \cup \{a_i | 1 \leq i \leq m\} \cup \{e, d, m, C\}, \\ - f = a_1, \\ - B_i = (e, P_i), \ 1 \leq i \leq 4. \end{array}$$

(1) To initialize simulation of computation of M we take agent $B_1 = (e, P_1)$ with a set of programs:

 P_1 :

 $\frac{P_1:}{1:\langle e \to l_0 \rangle, 2:\langle l_0 \leftrightarrow d \rangle;}$

(2) We need one more agent to generate some special object d. In every pair of steps the agent B_2 places one copy of d to the environment.

$$P_2$$
:

 $\overline{3:\langle e \to d \rangle, 4:\langle d \leftrightarrow C/d \leftrightarrow e \rangle;}$

The P colony Π starts its computation in the initial configuration $(e, e, e, e, \varepsilon)$. In the first subsequence of steps of P colony Π only agents B_1 , B_2 can apply its programs.

		conf	iguration	ı of П					
step	B_1	B_2	B_3	B_4	Env	P_1	P_2	P_3	P_4
1.	e	e	e	e		1	3		
2.	l_0	d	e	e			4		
3.	l_0	e	e	e	d	2	3		
4.	d	d	e	e	l_0				

(3) To simulate the ADD-instruction $l_1 : (ADD(r), l_2, l_3)$ there are two agents B_3 and B_4 in P colony Π . These agents help each other to add one copy of object a_r and object l_2 or l_3 to the environment.

P_3	P_3	P_4	P_4
$5: \langle e \leftrightarrow l_1 \rangle,$	$11: \left\langle E_1' \to l_2' \right\rangle,$	$15: \langle e \leftrightarrow E_1 \rangle,$	$21:\left\langle e\leftrightarrow l_{2}^{\prime}\right\rangle ,$
$6:\left\langle l_{1}\rightarrow E_{1}\right\rangle ,$	$12: \langle E_1' \to l_3' \rangle,$	$16:\left\langle E_1\to E_1'\right\rangle,$	$22:\left\langle e\leftrightarrow l_{3}^{\prime}\right\rangle ,$
$7:\left\langle E_1\leftrightarrow d\right\rangle,$	$13: \langle l'_2 \leftrightarrow e \rangle,$	$17: \langle E'_1 \leftrightarrow e \rangle,$	$23:\left\langle l_{2}^{\prime}\rightarrow l_{2}\right\rangle ,$
$8:\left\langle d\to L_1\right\rangle,$	$14: \langle l'_3 \leftrightarrow e \rangle,$	$18: \langle e \leftrightarrow L_1 \rangle,$	$24:\left\langle l_3'\to l_3\right\rangle,$
$9: \langle L_1 \leftrightarrow E_1'/L_1 \to m \rangle$,	$19: \langle L_1 \leftarrow a_r \rangle,$	$25:\left\langle l_{2}\leftrightarrow e\right\rangle ,$
$10: \langle m \to d \rangle,$		$20:\left\langle a_{r}\leftrightarrow e\right\rangle ,$	$26: \langle l_3 \leftrightarrow e \rangle;$

The agent B_3 consumes the object l_1 , changes it to E_1 and places it to the environment. The agent B_4 borrows E_1 from the environment and gives a little altered (to E'_1) back. B_3 rewrites the object d to some L_i . If this L_i has the same index as E'_i placed in the environment, the computation can go to the next phase. If indices of L_i and E_i are different the agent B_3 tries to generate another L_i . If the computation gets over this checking step, B_3 generates the helpful object l'_2 or l'_3 and places it to the environment. The agent B_4 exchanges it for "valid label" l_2 or l_3 .

An instruction l_i : $(ADD(r), l_j, l_k)$ is simulated by the following sequence of steps. Let the content of the agent B_2 be d.

		con	figuratio	on of Π					
step	B_1	B_2	B_3	B_4	Env	P_1	P_2	P_3	P_4
1.	d	d	e	e	$l_i a_r^u d^v$		4	5	
2.	d	e	l_i	e	$a_r^u d^{v+1}$		3	6	
3.	d	d	E_i	e	$a_r^u d^{v+1} d$		4	7	
4.	d	e	d	e	$E_i a_r^u d^{v+1}$		3	8	15
		cor	nfigurati	ion of I	I				
step	B_1	B_2	B_3	B_4	Env	P_1	P_2	P_3	P_4
5.	d	d	L_i	E_i	$a_r^u d^{v+1}$		4		16
6.	d	e	L_i	E'_i	$a_r^u d^{v+2}$		3		17
7.	d	d	L_i	e	$E'_i a^u_r d^{v+2}$		4	9	
8.	d	e	E'_i	e	$L_i a_r^u d^{v+3}$		3	11 or 12	2 18
9.	d	d	l'_i	L_i	$a_r^u d^{v+3}$		4	13	19
10.	d	e	$\overset{g}{e}$	a_r	$l'_i a^u_r d^{v+4}$		3		20
11.	d	d	e	e	$l'_i a^{u+1}_r d^{v+i}$	4	4		21
12.	d	e	e	l'_i	$a_r^{u+1}d^{v+5}$		3		23
13.	d	d	e	l_j	$a_r^{u+1}d^{v+5}$		4		25
14.	d	e	e	$\overset{\circ}{e}$	$l_j a_r^{u+1} d^{v+0}$	5			

(4) For each SUB-instruction $l_1 : (SUB(r), l_2, l_3)$, the next programs are introduced in the sets P_1 , P_3 and in the set P_4 :

P_3	P_3	P_1	P_4
$27: \left\langle e \leftrightarrow l_1 \right\rangle,$	$33:\left\langle F_{1}^{\prime\prime}\rightarrow l_{3}^{\prime}\right\rangle ,$	$36: \langle d \leftrightarrow F_1 \rangle,$	$41: \left\langle e \leftrightarrow l_2' \right\rangle,$
$28:\left\langle l_{1}\rightarrow F_{1}\right\rangle ,$	$34: \langle l'_2 \leftrightarrow e \rangle,$	$37: \langle F_1 \to F_1' \rangle,$	$42:\left\langle e\leftrightarrow l_{3}^{\prime}\right\rangle ,$
$29:\left\langle F_1\leftrightarrow d\right\rangle,$	$35: \langle l'_3 \leftrightarrow e \rangle;$	$38: \left\langle F_1' \leftrightarrow a_r / F_1' \to F_1'' \right\rangle,$	$43:\left\langle l_{2}^{\prime}\rightarrow l_{2}\right\rangle ,$
$30: \langle d \leftrightarrow F_1' \rangle,$		$39: \langle a_r \to d \rangle,$	$44:\left\langle l_{3}^{\prime}\rightarrow l_{3}\right\rangle ,$
$31: \left\langle F_1' \to l_2' \right\rangle,$		$40:\left\langle F_{1}^{\prime\prime}\leftrightarrow d\right\rangle ,$	$45:\left\langle l_2\leftrightarrow e\right\rangle,$
$32:\left\langle d\leftrightarrow F_{1}^{\prime\prime}\right\rangle ,$			$46: \langle l_3 \leftrightarrow e \rangle$

Agent B_4 starts simulation of executing SUB-instruction l_1 , the agent B_1 checks whether there is a copy of the object a_r in the environment or not and gives this information (F'_1 - there is some a_r ; F''_1 - there is no object a_r in the environment) to the environment.

An instruction l_i : $(SUB(r), l_j, l_k)$ is simulated by the following sequence of steps. When the value in counter r is zero:

		cont	figuratio	n of Π		a	pplicab	le progr	ams	
step	B_1	B_2	B_3	B_4	Env	P_1	P_2	P_3	P_4	
1.	d	d	e	e	$l_i d^v$		4	27		
2.	d	e	l_i	e	d^{v+1}		3	28		
3.	d	d	F_i	e	$d^{v+1}d$		4	29		
4.	d	e	d	e	$F_i d^{v+1}$	36	3			
5.	F_i	d	d	e	d^{v+2}	37	4			
6.	F'_i	e	d	e	d^{v+3}	38	3			
7.	$F_i^{\prime\prime}$	d	d	e	d^{v+3}	40	4			
8.	d	e	d	e	$F_i'' d^{v+3}$		3	32		
9.	d	d	F_i''	e	d^{v+4}		4	33		
10.	d	e	l'_k	e	d^{v+5}		3	35		
11.	d	d	e	e	$l'_k d^{v+5}$		4		42	
12.	d	e	e	l'_k	d^{v+6}		3		44	
13.	d	d	e	l_k^n	d^{v+6}		4		46	
14.	d	e	e	e	$l_{k}d^{v+7}$					

When register r stores value different from zero:

		cor	nfigurat	ion of I	Ι				
step	B_1	B_2	B_3	B_4	Env	P_1	P_2	P_3	P_4
1.	d	d	e	e	$l_i a_r^u d^v$		4	27	
2.	d	e	l_i	e	$a_r^u d^{v+1}$		3	28	
3.	d	d	F_i	e	$a_r^u d^{v+1} d$		4	29	
4.	d	e	d	e	$F_i a_r^u d^{v+1}$	36	3		
5.	F_i	d	d	e	$a_r^u d^{v+2}$	37	4		
6.	F'_i	e	d	e	$a_r^u d^{v+3}$	38	3		
7.	a_r	d	d	e	$F_i a_r^{u-1} d^{v+3}$	39	4	30	
8.	d	e	F'_i	e	$a_r^{u-1}d^{v+5}$		3	31	
9.	d	d	l'_i	e	$a_r^{u-1} d^{v+5}$		4	34	
10.	d	e	$\overset{j}{e}$	e	$l'_{i}a^{u-1}_{r}d^{v+6}$		3		41
11.	d	d	e	l'_i	$a_r^{u-1}d^{v+6}$		4		43
12.	d	e	e	l_j	$a_r^{u-1}d^{v+7}$		3		45
13.	d	d	e	$\overset{\circ}{e}$	$l_j a_r^{u-1} d^{v+7}$				

(5) The halting instruction l_h is simulated by agent B_3 with subset of programs: <u> P_3 </u>

$$\frac{1}{47}: \left\langle e \leftrightarrow l_h \right\rangle, \, 48: \left\langle l_h \to C \right\rangle, \, 49: \left\langle C \leftrightarrow e \right\rangle.$$

The agent consumes the object l_h and in the environment there is no other object l_m . This agent places one copy of the object C to the environment and stops working. In the next step the object C is consumed by the agent B_3 . No agent can start its work and computation halts. The execution of halting instruction l_h stops all agents in P colony Π :

		cont	figuratio	on of Π					
step	B_1	B_2	B_3	B_4	Env	P_1	P_2	P_3	P_4
1.	d	d	e	e	$l_h d^v$		4	47	
2.	d	e	l_h	e	d^{v+1}		3	48	
3.	d	d	C	e	$d^{v+1}d$		4	49	
4.	d	e	e	e	Cd^{v+1}		3		
5.	d	d	e	e	Cd^{v+2}		4		
6.	d	C	e	e	d^{v+3}				

P colony Π correctly simulates computation in the register machine M. The computation of Π starts with no object a_r placed in the environment in the same way as the computation in M starts with zeros in all the registers. The computation of Π stops if the symbol l_h is placed inside the corresponding agent in the same way as M stops by executing the halting instruction labeled l_h . Consequently, $N(M) = N(\Pi)$ and because the number of agents equals four, the proof is complete. \Box

Theorem 2. $NRM_{pb} \subseteq NPCOL_{par}(1, 2, *).$

Proof. Let us consider a partially blind register machine M with m registers. We construct a P colony $\Pi = (A, e, f, V_E, B_1, B_2)$ simulating a computation of the register machine M with:

 $- A = \{J, J', V, Q\} \cup \{l_i, l'_i, l''_i, L_i, L'_i, L''_i, E_i \mid l_i \in H\} \cup \{a_r \mid 1 \le r \le m\},$ - $f = a_1,$

- $B_i = (O_i, P_i), O_i = \{e\}, i = 1, 2$

The sets of programs are as follows:

(1) For initializing the simulation:

$P_1:$	P_1 :	P_2 :
$1: \langle e \to J \rangle,$	$3: \langle J \to l_0 \rangle,$	$5: \langle e \leftrightarrow J \rangle,$
$2: \langle J \leftrightarrow e \rangle,$	$4: \langle Q \to Q \rangle,$	$6: \langle J \to J' \rangle,$
		7: $\langle J' \leftrightarrow e \rangle$;

At the beginning of the computation the first agent generates the object l_0 (the label of starting instruction of M). It generates some copies of object J. The agent B_2 exchange them by J'.

	conf	iguration	n of Π		
	B_1	B_2	Env	P_1	P_2
1.	e	e		1	-
2.	J	e		2 or 3	-
3.	e	e	J	1	5
4.	J	J		2 or 3	6
5.	l_0	J'		8 or 24 or 34	7
6.	?	e	J'		

(2) For every ADD-instruction $l_1 : (ADD(r), l_2, l_3) P_1$ and P_2 contain:

$P_1:$	P_1 :	P_2 :
$8: \langle l_1 \to l_1' \rangle,$	$14: \langle L_1 \leftrightarrow E_1 \rangle,$	$18: \langle e \leftrightarrow l_1' \rangle,$
9: $\langle l'_1 \leftrightarrow J' \rangle$,	$15: \langle L_1 \to Q \rangle,$	$19: \langle l_1' \to E_1 \rangle,$
$10: \langle l'_1 \to Q \rangle,$	$16: \langle E_1 \rightarrow l_2 \rangle$	$20: \langle E_1 \leftrightarrow e \rangle,$
11 : $\langle J' \to L_1'' \rangle$,	$17: \langle E_1 \rightarrow l_3 \rangle$	$21: \langle e \leftrightarrow L_1 \rangle$
$12: \langle L_1'' \to L_1' \rangle,$		$22: \langle L_1 \to a_r \rangle$
$13: \langle L'_1 \to L_1 \rangle,$		$23: \langle a_r \leftrightarrow e \rangle$

When there is object l_1 inside agent B_1 , the agent rewrites it to one copy of l'_1 and the agent sends it to the environment. The agent B_2 borrows E_1 from the environment and returns E'_1 back.

The agent B_1 rewrites the object J' to some L_i . The first agent has to generate it in three steps to wait till the second agent generates the symbol E'_i and places it to the environment. If this L_i has the same index as E'_i placed in the environment, the computation can go to the next phase. If the indices of L_i and E_i are different, the agent B_1 generates Q and the computation never stops. If the computation gets over this checking step, B_1 generates object l_2 or l_3 .

	cont	figuration	n of Π		
	B_1	B_2	Env	P_1	P_2
1.	l_1	e	J'	8	—
2.	l'_1	e	J'	9 or 10	_
3.	J'	e	l'_1	11	18
4.	L_1''	l'_1		12	19
5.	L'_1	E_1		13	20
6.	L_1	e	E_1	14 or 15	—
7.	E_1	e	L_1	16 or 17	21
8.	l_2	L_1		8 or 24 or 34	22
9.	?	a_r		9 or 25 or 35	23
10.	?	e	a_r		

(3) For every SUB-instruction $l_1 : (SUB(r), l_2, l_3)$ there are subsets of programs in P_1 and P_2 :

P_1 :	$P_1:$	P_2 :
$24: \langle l_1 \to l_1'' \rangle,$	$28: \langle V \leftrightarrow l_1^{\prime\prime\prime} \rangle,$	$31: \langle l_1'' \leftrightarrow e \rangle,$
$25: \langle l_1'' \leftrightarrow a_r \rangle,$	$29: \langle l_1^{\prime\prime\prime} \to l_2 \rangle,$	$32: \langle l_1'' \to l_1''' \rangle,$
$26: \langle l_1'' \to Q \rangle,$	$30: \langle l_1^{\prime\prime\prime} \to l_3 \rangle$	$33: \langle l_1''' \leftrightarrow e \rangle,$
$27: \langle a_r \to V \rangle,$		

In the first step the agent checks if there is any copy of a_r in the environment (for zero in register r). In the positive case it rewrites a_r to V, in the other case l''_1 is rewritten to Q and the computation will never halt. At the end of this simulation the agent B_1 generates object l_2 or l_3 .

	confi	guratio	n of Π				confi	n of Π	
	B_1	B_2	Env	P_1	P_2		B_1	B_2	Env
1.	l_1	e	a_r	24	—	1.	l_1	e	
2.	l_1''	e	a_r	25 or 26	—	2.	l_1''	e	
3.	a_r	e	l_1''	27	31	3.	Q	e	
4.	V	l_1''		-	32	4.	Q	e	
5.	V	$l_1^{\prime\prime\prime}$		-	33	<u> </u>			
6.	V	e	$l_1^{\prime\prime\prime}$	28	—				
7.	$l_1^{\prime\prime\prime}$	e		29 or 30	—				
8.	l_2	e							

(4) For halting instruction l_h there are programs in sets P_1 and P_2 :

P_1 :	P_2 :	P_2 :
$34: \langle l_h \leftrightarrow J' \rangle,$	$39: \langle e \leftrightarrow l_h \rangle,$	$43: \left\langle L_h \leftrightarrow a_r \right\rangle, 1 < r \le m$
$35: \langle J' \to L_h \rangle,$	$40: \left\langle l_h \to \overline{l_h} \right\rangle,$	$44: \langle a_r \leftrightarrow e \rangle$
$36: \langle l_h \to Q \rangle,$	$41: \langle \overline{l_h} \leftrightarrow e \rangle,$	
$37: \langle L_h \to L_h \rangle,$	$42: \langle e \leftrightarrow L_h \rangle$	
$38: \left\langle L_h \leftrightarrow \overline{l_h} \right\rangle,$		

 P_1

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 P_2

By using these programs, the P colony finishes the computation in the same way as the partially blind register machine halts its computation. Programs with labels 43 and 44 in P_2 check value zero stored in all except the first one registers.

all counters $r, 1 < r \le m$ store zero				content of some counter $r, 1 < r \le m$ is							
configuration of Π					different from zero						
	B_1	B_2	Env	P_1	P_2		confi	guratio			
1.	l_h	e	J'	34 or 36	_		B_1	B_2	Env	P_1	P_2
2.	J'	e	l_{h}	35	39	1.	l_h	e	$J'a_r$	34 or 36	_
3.	L_h	l_h	10	37	40	2.	J'	e	$l_h a_r$	35	39
4.	L_{h}	$\frac{l}{l_h}$		37	41	3.	L_h	l_h	a_r	37	40
5.	L_{H}^{n}	e	$\overline{l_h}$	38	_	4.	L_h	$\overline{l_h}$	a_r	37	41
6.	$\frac{1}{l_h}$	e	L_{h}	_	42	5.	L_H	e	$\overline{l_h}a_r$	38	—
7.	$\frac{l_h}{l_h}$	L	10	_	_	6.	$\overline{l_h}$	e	$L_h a_r$	_	42
Ľ	-11	-11				7.	$\overline{l_h}$	L_h	a_r	_	43
						8.	$\overline{l_h}$	a_r	L_h	-	44
						9.	$\overline{l_h}$	L_h	a_r	_	43

P colony Π correctly simulates any computation of the partially blind register machine M. \Box

4 On computational power of restricted P colonies without checking

For restricted P colonies Following results are known from the literature:

- $NPCOL_{par}KR(2, *, 5) = NRE \text{ in } [2, 7],$
- $NPCOL_{par}R(2, *, 5) = NPCOL_{par}KR(2, 1, *) = NRE$ in [4].

Theorem 3. $NPCOL_{par}R(2,2,*) = NRE.$

Proof. Let us consider a register machine M with m registers. We construct a P colony $\Pi = (A, e, f, V_e, B_1, B_2)$ simulating the computations of register machine M with:

$$\begin{array}{ll} -A &= \{G\} \cup \{l_i, l'_i, l''_i, l'''_i, \overline{l_i}, \overline{l_i}, \overline{l_i}, \underline{l_i}, \underline{l_i}, L_i, L'_i, L''_i, F_i \mid l_i \in H\} \cup \\ \cup \{a_r \mid 1 \le r \le m\}, \\ -f = a_1, \\ -B_j = (O_j, P_j), \ O_j = \{e, e\}, j = 1, 2 \end{array}$$

At the beginning of the computation the first agent generates the object l_0 (the label of starting instruction of M). Then it starts to simulate instruction labeled l_0 and it generates the label of the next instruction. The sets of programs are as follows:

(1) For initializing of the simulation there is one program in P_1 :

 $\frac{P_1}{1: \langle e \to l_0; e \leftrightarrow e \rangle}$

The initial configuration of Π is (ee, ee, ε) . After the first step of computation (only the program 1 is applicable) the system enters configuration (l_0e, ee, ε) . (2) For every *ADD*-instruction $l_1 : (ADD(r), l_2, l_3)$ we add to P_1 the programs: P_1

 $\begin{array}{l} \frac{1}{2} : \langle e \to a_r; l_1 \leftrightarrow e \rangle \,, & 3: \langle e \to G; a_r \leftrightarrow l_1 \rangle \,, \\ 4: \langle l_1 \to l_2; G \leftrightarrow e \rangle \,, & 5: \langle l_1 \to l_3; G \leftrightarrow e \rangle \end{array}$

When there is object l_1 inside the agent, it generates one copy of a_r , puts it to the environment and generates the label of the next instruction (it nondeterministically chooses one of the last two programs 4 and 5)

	conf				
	B_1	B_2	Env	P_1	P_2
1.	$l_1 e$	ee	a_r^x	2	—
2.	$a_r e$	ee	$l_1 a_r^x$	3	—
3.	Gl_1	ee	a_r^{x+1}	4 or 5	—
4.	$l_2 e$	ee	$a_r^{x+1}G$		

(3) For every SUB-instruction $l_1 : (SUB(r), l_2, l_3)$, the next programs are added to sets P_1 and P_2 :

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At the first phase of the simulation of the SUB instruction the first agent generates object l'_1 , which is consumed by the second agent. The agent B_2 generates symbol L_1 and tries to consume one copy of symbol a_r . If there is any a_r , the agent sends to the environment object L''_1 and consumes L_1 . After this step the first agent consumes L''_1 or L_1 and rewrites it to l_2 or l_3 . The objects \underline{x} , \overline{x} and $\overline{\overline{x}}$ are used for a synchronization of the computation in both agents and for storing information about the state of the computation.

Instruction $l_1: (SUB(r), l_2, l_3)$ is simulated by the following sequence of steps.

	configuration of \varPi						confi	guratio	n of Π		
	B_1	B_2	Env	P_1	P_2		B_1	B_2	Env	P_1	P_2
1.	$l_1 e$	ee	a_r^x	6	-	1.	$l_1 e$	ee		6	-
2.	$l_1'e$	ee	a_r^x	7	_	2.	$l_1'e$	ee		7	_
3.	$l_1''e$	ee	$l_1'a_r^x$	8	18	3.	$l_1''e$	ee	l_1'	8	18
4.	$l_1^{\prime\prime\prime}e$	$L_1 l_1'$	$l_1^{\prime\prime}a_r^x$	9	19	4.	$l_1^{\prime\prime\prime}e$	$L_1 l_1'$	l_1''	9	19
5.	$l_1^{\prime\prime\prime\prime}e$	$L_1^\prime l_1^{\prime\prime}$	$L_1 a_r^x$	10	20	5.	$l_1^{\prime\prime\prime\prime}e$	$L_1^\prime l_1^{\prime\prime}$	L_1	10	
6.	$\overline{l_1}e$	$L_1''a_r$	$L_1 L_1' a_r^{x-1}$	11	21	6.	$\overline{l_1}e$	$L_1^\prime l_1^{\prime\prime}$	L_1	11	
7.	$\overline{\overline{l_1}}e$	eL_1	$L_1'' a_r^{x-1}$	12	22	7.	$\overline{\overline{l_1}}e$	$L_1^\prime l_1^{\prime\prime}$	L_1	13	
8.	$\underline{l_2}L_1''$	ee	a_r^{x-1}	14	-	8.	$\underline{l_3}L_1$	$L_1' l_1''$		15	—
9.	$l_2 e$	ee	$a_r^{x-1}\underline{l_2}$			9.	F_3e	$L_1'l_1''$	$\underline{l_3}$	16	—
						10.	$l_{3}l_{3}$	$L_1^\prime l_1^{\prime\prime}$	F_3	17	23
						11.	$\overline{l_3}e$	F_3e	$\underline{l_3}L_1'$	2 or 6	24
									_	or none	
						12.	??	ee	$\underline{\underline{l}_3}L'_1$		

If the register r stores value zero : If the register r stores nonzero value:

(4) For halting instruction l_h no program is added to the sets P_1 and P_2 .

P colony Π correctly simulates all computations of the register machine M and the number contained on the first register of M corresponds to the number of copies of the object a_1 present in the environment of Π . \Box

5 Conclusions

We have shown that the P colonies with capacity k = 2 and without checking programs with height at most 2 are computationally complete. In Section 3 we have shown that the P colonies with capacity k = 1 and with checking/evolution programs and 4 agents are computationally complete.

We have verified also that partially blind register machines can be simulated by P colonies with capacity k = 1 without checking programs with two agents. The generative power of $NPCOL_{par}K(1, n, *)$ for n = 2, 3 remains open.

In Section 4 we have studied P colonies with capacity k = 2 without checking programs. Two agents guarantee the computational completeness in this case.

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Networks of Mealy Multiset Automata

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Summary. We introduce the networks of Mealy multiset automata, and study their computational power. The networks of Mealy multiset automata are computationally complete.

1 Learning from Molecular Biology

Systems biology represents a new cross-disciplinary approach in biology which has only recently been made possible by advances in computer science and technology. As it is mentioned in [10], it involves the application of experimental, theoretical, and modelling techniques to the study of biological organisms at all levels. Adding new abstractions, discrete models and methods able to help our understanding of the biological phenomena, systems biology may provide predictive power, useful classifications, new paradigms in computing and new perspectives on the dynamics of various biological systems.

Recent promising work [1] employs automata theory as an efficient tool of describing and controlling gene expression (a small automaton is encoded by DNA strands and then it is used in logical control of gene expression).

In [2], we present a way of interaction between gene machine and protein machine, namely the process of making proteins, in abstract terms of Mealy automata, transformation semigroup and abstract operations.

The Mealy automaton proposed as a formal model of the genetic message translation is a minimal one that accepts the mRNA messages and terminates the translation process (according to [9], there are no appropriate formalism for the process of translation).

However molecular biology "deals" not only with sequences, but also with multisets. The biological cells are "smart" enough to put together at work sequences and multisets of atoms and molecules, so if we try to get models from their functioning, we should not restrict ourselves in dealing only with sequential machines (like classical automata). To deal with multisets, the main approach is given by membrane systems [11]. There is also introduced and studied an automata-like machine to work with multisets [7], i.e. multiset automaton. At a first glance, it seams that they are nothing else but weighted automata with weights in the semiring of positive integers. In [8] it is proven that such automata (weighted automata with weights in the semiring of positive integers) has the same power as finite automata (accepts only regular languages). In fact, a careful reader should remark that a multiset automaton is not a sequential machine, and it is not working with sequences of multisets as in the case of weighted automata. A multiset automaton accepts, together with a sequence of multisets, an entire class, namely the class of that sequence obtained by "abelianization" (as an example, together with the sequence, say *aab*, it accepts *aba* and *baa*). In this manner, multiset automata become very powerful (see [7], for details). Mealy multiset automata, [3], can be viewed as the corresponding Mealy machine. We study some of their (co)algebraic properties in [3] and [4] and we connect this properties with various aspects of their behaviour. In [5], we organize them in a P machine, in order to simulate a P system. However the biological systems are not always organized in an hierarchical manner. This means that we also have to organize sets of Mealy multiset automata in networks. In order to obtain the computing power for networks of such automata, we relate them to neural P systems, proving that networks of Mealy multiset automata are computationally complete.

2 Networks of Mealy Multiset Automata

In order to define networks of Mealy multiset automata, we can connect these automata in many ways, having both parallel and serial connections. In [3] we define the *restricted direct product* of MmA for the parallel case, and the *cascade product* for a serial connection. See, also, the appendix for details.

2.1 Mealy Multiset Automata

Roughly speaking, an *Mealy multiset automata* (MmA) consists of a storage location (a box for short) in which we place a multiset over an input alphabet and a device to translate the multiset into a multiset over an output alphabet. We have a detection head that detects whether or not a given multiset appears in the multiset available in the box. The multiset is removed from the box whenever it is detected, and the automaton inserts a multiset over the output alphabet (or a marked symbol if the output alphabet is the same) that can not be viewed by the detection head. This automaton stops when no further move is possible. We say that the sub-multiset read by the head was translated to a multiset over the output alphabet. We give here only the definitions and the properties that we need for networks of MmA. For more informations see [3] or [4]. From the formal point of view, a *Mealy multiset automaton* is a construct $\mathcal{A} = (Q, V, O, f, g, q_0)$ where

1. Q is a finite set, the set of *states*;

2. $q_0 \in Q$ is special state, which is both initial and final;

- 3. V is a finite set of objects, the *input alphabet*;
- 4. *O* is a finite set of objects, the *output alphabet*, such that $O \cap V = \emptyset$;
- 5. $f: Q \times \mathbb{N} \langle V \rangle \to \mathcal{P}(Q)$ is the state-transition (partial) mapping;
- 6. $g: Q \times \mathbb{N} \langle V \rangle \to \mathcal{P}(\mathbb{N} \langle O \rangle)$ is the *output (partial) mapping*.

If $|f(q, a)| \leq 1$ we say that \mathcal{A} is Q- deterministic and if $|g(q, a)| \leq 1$ our automaton is O-deterministic.

An MmA \mathcal{A} receives a multiset in its box, and processing this multiset it passes through different *configurations*. It starts with a multiset from $\mathbb{N} \langle V \rangle$ and ends with a multiset from $\mathbb{N} \langle V \cup O \rangle$. A *configuration* of \mathcal{A} is a triple $(q, \alpha, \overline{\beta})$ where $q \in Q, \alpha \in \mathbb{N} \langle V \rangle, \overline{\beta} \in \mathbb{N} \langle O \rangle$. We say that a configuration $(q, \alpha, \overline{\beta})$ passes to $(s, \alpha - a, \overline{\beta} + \overline{b})$ (or, that we have a *transition* between those configurations) if there is $a \subseteq \alpha$ such that $s \in f(q, a), \overline{b} \in g(q, a)$. We denote this by $(q, \alpha, \overline{\beta}) \vdash$ $(s, \alpha - a, \overline{\beta} + \overline{b})$. We also denote by \vdash^* the reflexive and transitive closure of \vdash . We could alternatively define a configuration to be a pair (q, α) where $\alpha \in \mathbb{N} \langle V \cup O \rangle$ and the transition relation is $(q, \alpha) \vdash (s, \alpha - a + \overline{b})$, with the same conditions as above.

2.2 Networks of automata

The formal description of a network of Mealy multiset automata is not intuitive. On the other hand, these networks could be very powerful, so we think that they deserve our attention. We can consider several variants of such networks. Some of them can have no inter-communication and, in this case, the network is, in fact, a bigger MmA. The same remark can be done if we have only MmA connected in a serial manner, without any ramifications (as we have seen in the previous subsections). The case that we consider in this paper is inspired by the definition of neural P systems (nP systems). Neural P systems are defined in [11] as a computing model inspired by the network of cells. Each cell has a finite state memory, and processes multisets of symbol (impulses); it can send some impulses (called excitations) to the neighbouring cells. It is proved that such networks are rather powerful: they can simulate Turing machine using a small number of cells, every cell having in a small number of states. It is also proved that, in appropriate organization, such a network can solve in linear time the Hamiltonian Path Problem.

We consider a set of MmA that can communicate by means of some *commu*nication channels. All of them have the same input alphabet V, and their boxes contain an input multiset over V (they can also have an empty multiset ε as input). The output alphabet has a "real" part O of output alphabet, and a "specific" part used for communication. The specific part is, in fact, a Cartesian product between the input alphabet V and the set of targets T (the set of the indexes of the MmA forming the net). We can also have a special MmA to collect in its box the result of the computation (i.e. a multiset over O) for such a network. Alternatively, we can consider as result of the computation the tuple of multisets obtained in the box of every MmA of the net.

Definition 1. A network of Mealy multiset automata (shortly, nMmA) is a construct $\mathcal{N} = (V, O, \{\mathcal{A}_i\}_{i=\overline{1,n}}, \{\Lambda_i\}_{i=\overline{1,n}}, B)$ where:

- $V = \{a_1, a_2, ..., a_m\}$ is a finite set of objects, the *input alphabet*;
- *O* is the *output alphabet* such that $O \cap V = \emptyset$;
- $\mathcal{A}_i = (Q_i, V, \overline{O}, f_i, g_i, s_{0,i})$ are MmA's connected in the network. Their output alphabets are of the form $\overline{O} = O \cup (V \times T)$, where $T = \{1, 2, ..., n\}$;
- B is a box where \mathcal{N} "receives" the output multiset. Depending on features that we consider for the net, B can be a specific box of a specific MmA in the network, or B can be the Cartesian product of all the boxes.
- $\Lambda_i : \mathbb{N} \langle \overline{O} \rangle \to (\mathbb{N} \langle O \rangle \cup \mathbb{N} \langle V \rangle)^n$ are the communication mappings associated to all the $\mathcal{A}'_i, i \in T$.

A computation starts with some input multisets $w_{0,i}$ in the boxes of the MmA's that are in their initial states, $s_{0,i}$; then we have a big step given by a translation (made by the MmA's, in fact by their restricted direct product $\bigwedge_{i=1}^{n} \mathcal{A}_i$) and a communication (done by $\{A_i\}$) - a kind of "parallel cascade product", since every MmA is in cascade with the restricted direct product of itself and the other MmA.

A configuration of the network is of the form (s, w), where $s = (s_1, s_2, ..., s_n)$ with $s_i \in Q_i$ is the global state, and $w = (w_1, ..., w_n)$ where $w_i \in \mathbb{N} \langle O \rangle \cup \mathbb{N} \langle V \rangle$.

A transition between configurations is denoted by $(s, w) \vdash (s', w')$ and is defined in the following manner:

 $s' = (s'_1, s'_2, ..., s'_n)$, where $s'_i \in f_i(s_i, a_i)$ with $a_i \in \mathbb{N} \langle V \rangle$; we allow some of the a's to be ε if in the corresponding MmA there is no transition.

 $w' = \Lambda_1(b_1) + \Lambda_2(b_2) + \dots + \Lambda_n(b_n) + (w_1 - a_1, w_2 - a_2, \dots, w_n - a_n)$, where $b_i \in g_i(s_i, a_i)$.

A network of MmA can be used in various modes. We can use it as a generative system, looking to the number of output objects that we find in the boxes (without considering the final state for the MmA). It can be used also to compute functions from $\mathbb{N}\langle V \rangle$ to $\mathbb{N}\langle O \rangle$. An example of such a network used as a generative system could clarify these aspects:

Example 1. Let

$$\mathcal{N} = (V, O, \{\mathcal{A}_i\}_{i=\overline{1,3}}, \{\Lambda_i\}_{i=\overline{1,3}}, B_1)$$

where:

- $V = \{a\}$ is the input alphabet;
- $O = \{b\}$ is the output alphabet $b \neq a$;
- $\mathcal{A}_i = (\{s_i\}, V, \overline{O}, f_i, g_i, s_i)$, are the MmA's connected in the network. Their output alphabet is $\overline{O} = \{b, (a, 1), (a, 2), (a, 3)\}$
- B_1 is the box where \mathcal{N} "receives" the output multiset.
- $\Lambda_i : \mathbb{N} \langle \overline{O} \rangle \to (\mathbb{N} \langle O \rangle \cup \mathbb{N} \langle V \rangle)^3$ are the communication mappings associated to all the $\mathcal{A}_i, i \in T = \{1, 2, 3\}.$

We describe now the mappings. The transition mappings are

•
$$f_i(s_i, a) = s_i, i = \overline{1, 3};$$

The output mappings:

- $g_1(s_1, a) \in \{b, (a, 2) + (a, 3)\}$, so is a nondeterministic mapping;
- $g_2(s_2, a) = (a, 1);$
- $g_3(s_3, a) = (a, 1)$

The communication mappings:

- $\Lambda_1(nb+k_1(a,1)+k_2(a,2)+k_3(a,3))=(nb+k_1a, k_2a, k_3a))$
- $\Lambda_2(nb+k_1(a,1)+k_2(a,2)+k_3(a,3))=(k_1a, \varepsilon, \varepsilon))$
- $\Lambda_3(nb+k_1(a,1)+k_2(a,2)+k_3(a,3))=(k_1a, \varepsilon, \varepsilon))$

Since \mathcal{A}_1 has a nondeterministic output mapping, the behaviour of our network is nondeterministic. We denote the global state by $s = (s_1, s_2, s_3)$. We start our computation from $(s, (a, \varepsilon, \varepsilon))$ Applying the restricted direct product we can obtain $(s, (b, \varepsilon, \varepsilon))$ or $(s, (\varepsilon, a, a))$.

In the first case we obtain one b, so we generate 1. In the second case, the computation continues with communication, and we obtain $(s, (a + a, \varepsilon, \varepsilon) = (s, (2a, \varepsilon, \varepsilon))$, and, again, we have various possibilities to choose. Anyway, it should be clear now that we can generate any number of b's, so \mathcal{N} can generate every positive integer.

In order to study the computational power of nMmA, we are trying to simulate neural-like P systems. To be more specific, we try to simulate the neural P systems working in minimal mode and replicative manner. To keep the paper self-contained, we remember some facts about neural P systems and adapt the notations from [11].

3 Neural P Systems

The former tissue P systems were called neural-like P systems in [11]. We start with the classical definition, and later we adapt the notation to our needs. We consider a class of networks of membranes inspired by the way the neurons cooperate to process impulses in the complex net established by synapses. A possible model of this symbol processing machinery can be given by a network of membranes, each of them containing a multiset of objects and a state according to which the objects are processed. The membranes can communicate along "axons" channels. We make some minor modifications to the original notations, having in mind that in the Mealy multiset automata we distinguish between multisets and strings that could represent them (since we can deal with two kinds of behaviours, a global one and a sequential one). We also restrict our presentation of neural P systems working in *minimal mode and replicative manner*. **Definition 2.** A neural P system (*nP system*) of degree $m \ge 1$ is a construct

$$\Pi = (V, \sigma_1, \sigma_2, ..., \sigma_m, syn, i_{out}),$$

where

1. V is a finite non-empty alphabet (of *objects*);

- 2. $syn \subseteq \{1, 2, ..., m\} \times \{1, 2, ..., m\}$ (synapses among cells);
- 3. $i_{out} \in \{1, 2, ..., m\}$ indicates the *output cell*; we can put $i_{out} = 1$;
- 4. $\sigma_1, \sigma_2, ..., \sigma_m$ are *cells* of the form $\sigma_i = (Q_i, s_{i,0}, w_{i,0}, R_i), 1 \le i \le m$,

where:

- Q_i is a finite set (of *states*);
- R_i is a finite set of rules of the form $sw \to s'(x + y_{go} + z_{out})$, where $s, s' \in Q_i$, $w, x \in \mathbb{N} \langle V \rangle, y_{go} \in \mathbb{N} \langle V \times \{go\} \rangle, z_{out} \in \mathbb{N} \langle V \times \{out\} \rangle$, with the restriction that $z_{out} = \varepsilon$ for all *i* different from 1.

The objects that appears in the left hand multiset w of the rule $sw \to s'w'$ are called *impulses*, while those from w' are called *excitations*.

An *m*-tuple of the form $(s_1w_1, s_2w_2, ..., s_mw_m)$ is called a *configuration* of Π . Using the rules defined above, we can define *transitions* among the configurations of the system. To this end, there are considered three modes of processing the *impulse-objects* and three modes of transmitting *excitation-objects* from one cell to another one. As we already mention, we restrict ourselves to the *minimal processing mode*.

Notations: $V_{go} = \{(a; go) \mid a \in V\}, V_{out} = \{(a; out) \mid a \in V\}, \text{ and } V_{tot} = V \cup V_{go} \cup V_{out}.$ For $s, s' \in Q_i, x \in \mathbb{N} \langle V \rangle$ and $y \in \mathbb{N} \langle V_{tot} \rangle$, we write

 $sx \Rightarrow_{min} s'y \text{ iff } sw \to s'w' \in R_i, w \subseteq x \text{ and } y = (x - w) \cup w'.$

In this case, only one occurrence of the multiset from the left-hand side of a rule is processed, being replaced by the multiset from the right-hand of the rule, and at the same time changing the state of the cell.

We also write $sx \Rightarrow_{min} sx$ for $s \in Q_i$ and $x \in \mathbb{N} \langle V \rangle$ whenever there is no rule $sw \to s'w' \in R_i$ such that $w \subseteq x$. This encodes the case when a cell cannot process the current objects in a given state (it can be "unblocked" after receiving new impulses from the cells which are active and can send objects to it).

Now, recall that the multiset w' from a rule $sw \to s'w'$ contains symbols from V, but also symbols of the form (a, go) (or, in the case of the cell 1, of the form (a, out)). Such symbols are sent to the cells related by synapses to the cell σ_i where the rule $sw \to s'w'$ is applied, according to various manners of communication. As we already mention it, we choose the *replicative manner*, i.e. each symbol a from (a, go) appearing in w', it is sent to each of the cells σ_j such that $(i; j) \in syn$.

In order to formally define the transition among the configurations of Π , some further notations are needed. For a multiset w over V_{tot} , we consider the projections on V, V_{go} and V_{out} , namely $pr_V(w)$; $pr_{V_{go}}(w)$, and $pr_{V_{out}}(w)$ (see [11] for details). For a node i in the graph defined by syn, the ancestors and the successors of node *i* are denoted by $anc(i) = \{j \mid (j,i) \in syn\}$ and $succ(i) = \{j \mid (i,j) \in syn\}$, respectively.

Each transition lasts one time unit, and the network is synchronized: a global clock define the passage of time for all the cells.

For two configurations $C_1 = (s_1w_1, ..., s_mw_m)$ and $C_2 = (s'_1w''_1, ..., s'_mw''_m)$ we write $C_1 \Rightarrow C_2$ if there are $w'_1, ..., w'_m$ in $\mathbb{N} \langle V_{tot} \rangle$ such that

$$s_i w_i \Rightarrow s'_i w'_i, 1 \le i \le m$$

and

$$w"_i = pr_V(w'_i) + \sum_{j \in anc(i)} pr_{V_{go}}(w'_j)$$

Obviously, objects are always sent to a cell i only from its ancestors, namely from cells j such that a direct synapse exists from j to i. In the case of the cell 1, we remove from w'_1 all the symbols $a \in V$ which appear in w'_1 in the form (a, out). If during a transition a cell does nothing (no rule is applicable to the available multiset of objects in the current state), then the cell waits until new objects are sent to it from its ancestor cells.

A sequence of transitions among the configurations of Π is called a *computation* of Π . A computation ending in a configuration where no rule in no cell can be used is called a halting computation. The result of a halting computation is the number of objects in the output cell 1 (or sent to the environment from the output cell 1). We denote by $N(\Pi)$ the set of all natural numbers computed in this way by a system Π . We denote by $NOnP_{m,r}(coo)$ the family of sets $N(\Pi)$ computed by all cooperative neural-like P systems with at most $m \geq 1$ cells, each of them using at most $r \geq 1$ states. When non-cooperative systems are used, we write $NOnP_{m,r}(ncoo)$ for the corresponding family of sets $N(\Pi)$.

3.1 Computational power

We denote by *NRE* the family of Turing computable sets of natural numbers.

Following [11], we mention that the minimal mode of using the rules turns out to be computationally universal. If we consider the apparently weak neural-like P systems, then the fact that we obtain universality even in the non-cooperative case when using the mode min of applying the rules is rather unexpected. The same result holds true also when using cooperative rules. Among the results presented in [11] we mention here only those for minimal mode and for replicative manner.

Theorem 1. $NOnP_{2,5}(ncoo) = NRE$.

For the cooperative rules, the number of states can be decreased.

Theorem 2. $NOnP_{2,2}(coo) = NRE$.

4 Universality of the Networks

In order to obtain the generative power of a network of MmA, we give the following result.

Theorem 3. Any *nP* System working in min mode and replicative manner can be simulated by a network of MmA (possibly nondeterministic).

Proof. Let $\Pi = (V, \sigma_1, \sigma_2, ..., \sigma_m, syn, 1)$ be an nP system with its components described as in the previous section. We remind that $\sigma_1, \sigma_2, ..., \sigma_m$ are cells of the form $\sigma_i = (Q_i, s_{i,0}, w_{i,0}, R_i)$ $(1 \le i \le m)$, where Q_i is a finite set (of *states*) and R_i is a finite set of rules of the form $sw \to s'(x + y_{go} + z_{out})$ with $s, s' \in Q_i$, $w, x \in \mathbb{N} \langle V \rangle$, $y_{go} \in \mathbb{N} \langle V \times \{go\} \rangle$, $z_{out} \in \mathbb{N} \langle V \times \{out\} \rangle$ (with the restriction that $z_{out} = \varepsilon$ for all *i* different from 1).

We can build a nMmA $\mathcal{N} = (V, O, \{\mathcal{A}_i\}_{i=\overline{1,m}}, \{\Lambda_i\}_{i=\overline{1,m}}, B_1)$ where:

- the output alphabet O is V_{out} ;

- $\mathcal{A}_i = (Q_i, V, \overline{O}, f_i, g_i, s_{0,i})$ is the MmA simulating the activity of cell σ_i . The output alphabets are of the form $\overline{O} = O \cup (V \times T)$, where $T = \{1, 2, ..., m\}$;

- B_1 is the box where \mathcal{N} collects the output multisets;

- $\Lambda_i : \mathbb{N} \langle \overline{O} \rangle \to (\mathbb{N} \langle O \rangle \cup \mathbb{N} \langle V \rangle)^n$ are the communication mappings associated to $\mathcal{A}_i, i \in T$.

Consider a rule $sw \to s'(x + y_{go} + z_{out})$ from R_i . We can simulate this rule with f_i and g_i by defining them in the following manner:

- $f_i(s,w) = s';$

 $-g_i(s,w) = (z_{out} + x + k_1(y,1) + k_2(y,2) + \dots + k_m(y,m)),$

with the following restrictions in g_i :

* if $i \neq 1$, then $z_{out} = \varepsilon$;

* if there is no synapse from σ_i to σ_j , we define $k_j = 0$, else $k_j = 1$.

In this manner we can also simulate the replicative manner of applying the rules, since y is marked to be send to all the cells having synapse from σ_i .

It is easy to see that we have a transition $(s_1w_1, ..., s_mw_m) \Rightarrow (s'_1w''_1, ..., s'_mw''_m)$ in Π if and only if $((s_1, s_2, ..., s_n), (w_1, ..., w_n)) \vdash ((s'_1, s'_2, ..., s'_n), (w''_1, ..., w''_n))$

As an immediate consequence of this result we get the following

Theorem 4. Nondeterministic networks of Mealy multiset automata are universal.

Proof. We already know that $NOnP_{2,2}(coo) = NRE$. Applying the previous theorem we obtain that the generative power of a nondeterministic network of MmA is NRE. Therefore the nondeterministic network of MmA is universal.

Therefore a network of Mealy multiset automata is able to simulate Turing machines, and so it is computationally complete. The number of cells and states sufficient to characterize the power of Turing machines is rather small.

P systems are simulated on a network of computers [6]. It would be interesting to see whether such an implementation can be related to the network of Mealy multiset automata.

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Appendix

Behaviour of MmA

For some of the categorical properties of MmA's, as well as *behaviour* and *bisimulation relation*, we refer to [3] and [4]. Behaviour is often appropriately viewed as consisting of both dynamics and observations, which have to do with change of states and partial access to states, respectively. The main advantage of an MmA is that it has an output function that can play the main role in observability, i.e. we do not have to construct an other machine to describe the MmA's behaviour.

Definition 3. Let $\mathcal{A} = (Q, V, O, f, g)$ be a Mealy multiset automaton. The general behaviour of a state $q \in Q$ is a function $\mathbf{beh}(q)$ assigning to every multiset $\alpha \in \mathbb{N} \langle V \rangle$ the output multiset obtained after consuming α starting from q.

When talking about the behaviour, we consider a specific order of consuming multisets, i.e. in terms of strings of multisets.

A certain feature for MmA is that the behaviour is always finite because we can not go further after consuming the given multiset. On the other hand, since the outputs go back into the box, it is possible that we can not track the sequence of intermediate states. If we are interested only on the outcome of the machine, then we should not take care of the intermediate states, and if the input multiset is partially consumed, it should be of interest the state where the MmA arrives in order to (possible) provide the box with a supplementary multiset in order to make

the initial one a consumed multiset. These considerations lead us to the following definition.

Definition 4. Let $\mathcal{A} = (Q, V, O, f, g)$ be a Mealy multiset automaton. The sequential behaviour of a state $q \in Q$ is a function $\operatorname{seqbeh}(q)$ that assigns to every multiset $\alpha \in \mathbb{N} \langle V \rangle$ all the sequences of the output multisets obtained after consuming α starting from q.

Example 2. Suppose that we have the following sequence of transitions $(q, \alpha, \varepsilon) \vdash (q_1, \alpha - a_1, b_1) \vdash (q_2, \alpha - a_1 - a_2, b_1 + b_2) \vdash ... \vdash (q_n, \alpha - a_1 - ... - a_n, b_1 + ... + b_n)$ and MmA stops. Then **beh** $(q)(\alpha) = b_1 + ... + b_n$ and **seqbeh** $(q)(\alpha) \ni b_1...b_n$. Moreover, $b_1 + ... + b_n$ belongs to $\mathbb{N}\langle O \rangle$, while $b_1...b_n$ belongs to $(\mathbb{N}\langle O \rangle)^*$.

Consider the canonical inclusion $i : \mathbb{N} \langle O \rangle \to (\mathbb{N} \langle AO \rangle)^*$ and the identity map $id : \mathbb{N}(O) \to \mathbb{N}(O)$. By the universal property of the free monoid, we know that there exists a unique homomorphism of monoids $\mathbf{I}_O : (\mathbb{N} \langle O \rangle)^* \to \mathbb{N} \langle O \rangle$ defined by $\mathbf{I}_O(b_1...b_n) = b_1 + ... + b_n$ such that $\mathbf{I}_O \circ i = id$. Since id is onto, it follows that \mathbf{I} is onto, and so, applying the isomorphism theorem for monoids, we obtain that $(\mathbb{N} \langle O \rangle)^*/ker \mathbf{I}_O \simeq \mathbb{N} \langle O \rangle$.

Proposition 1. For all the states q of a Mealy multiset automaton we have $I_O \circ \operatorname{seqbeh}(q) = \operatorname{beh}(q).$

If q, q' are two bisimilar states (see [3] for details), they have the same sequential behaviour $\mathbf{seqbeh}(q) = \mathbf{seqbeh}(q')$. This implies that they also have the same behaviour $\mathbf{beh}(q) = \mathbf{beh}(q')$. We can define (since the reciprocal it is not true) a weaker equivalence relation:

$$q \approx q' \Leftrightarrow \mathbf{beh}(q) = \mathbf{beh}(q')$$

Proposition 2. We have $q \approx q' \Leftrightarrow (\forall) \alpha \in \mathbb{N} \langle V \rangle$:(seqbeh(q), seqbeh(q')) $\in ker \mathbf{I}_O$.

Since this relation is independent of the order of consuming resources from the box, we call it *output conservative equivalence*. The importance of this equivalence is given mainly by the idea of consuming and producing *resources*. The resource problem appears to be of interest when we consider the parallel/concurrent processes. In this manner we overpass the sequential framework previously represented by **seqbeh**.

Restricted direct product of Mealy multiset automata

Let $\mathcal{A}_i = (Q_i, V, O, f_i, g_i)$, and B_i their corresponding boxes, $i = \overline{1, n}$, a finite family of Mealy multiset automata. We can connect them in *parallel* in order to obtain a new MmA defined by $\mathcal{A} = \bigwedge_{i=1}^n \mathcal{A}_i = (\times_{i=1}^n Q_i, V, O^n, f, g)$, called the *restricted direct product* of \mathcal{A}_i , where:

- $f((q_1, q_2, ..., q_n), a) = (f_1(q_1, a), f_2(q_2, a), ..., f_n(q_n, a)),$
- $g((q_1, q_2, ..., q_n), a) = (g_1(q_1, a), g_2(q_2, a), ..., g_n(q_n, a)),$
- box of \mathcal{A} is the disjoint union $\bigsqcup_{i=1}^{n} B_i$ of $\{B_i \mid i = \overline{1, n}\},\$
- a configuration of \mathcal{A} is a triple $(q, \alpha, \overline{\beta})$, where $q = (q_1, q_2, ..., q_n)$, $\alpha = (\alpha_1, \alpha_2, ..., \alpha_n)$, and $\overline{\beta} = (\overline{\beta}_1, \overline{\beta}_2, ..., \overline{\beta}_n)$,
- the transition relation of \mathcal{A} : $(q, \alpha, \overline{\beta}) \vdash (s, \alpha a, \overline{\beta} + \overline{b})$ iff $s_i \in f_i(q_i, a_i)$ and $\overline{b}_i \in g_i(q_i, a_i)$ for all $i \in \overline{1, n}$.

The cascade product of Mealy multiset automata

The cascade product is useful to describe a serial connection, and provide also some results in decompositions of such machines in irreducible ones.

Let $\mathcal{A} = (Q, V, O, f, g)$, $\mathcal{A}' = (Q', V', O', f', g')$ be two Mealy multiset automata. In order to connect them, we need a multiset mapping linking the output of one of them to the input of the other. This can be done using a \mathbb{N} -homomorphism from $\mathbb{N} \langle O' \rangle$ to $\mathbb{N} \langle V \rangle$ (this homomorphism can be obtained by using a mapping from O' to V). We denote by $\Lambda : \mathbb{N} \langle O' \rangle \to \mathbb{N} \langle V \rangle$ this homomorphism. Then we can define a mapping $\Omega : Q' \times \mathbb{N} \langle V' \rangle \to \mathbb{N} \langle V \rangle$ by $\Omega(q', a') = \Lambda(g'(q', a'))$.

• This mapping gives us the cascade product induced by Ω :

$$\mathcal{A}\Omega\mathcal{A}' = (Q \times Q', V', O, f^{\Omega}, g^{\Omega})$$

where $f^{\Omega}((q,q'),a') = (f(q,\Omega(q',a')), f'(q',a'))$, and $g^{\Omega}((q,q'),a') = g(q,\Omega(q',a'))$, for all $a' \in \mathbb{N} \langle V' \rangle, (q,q') \in Q \times Q'$.

• The transition relation becomes $((q,q'), \alpha', \overline{\beta}) \vdash ((s,s'), \alpha'-a', \overline{\beta}+\overline{b})$ if there is $a' \subseteq \alpha'$ such that $(s,s') = f^{\Omega}((q,q'), a')$ and $\overline{b} = g^{\Omega}((q,q'), a')$, where $a', \alpha' \in \mathbb{N} \langle V' \rangle, (q,q') \in Q \times Q'$, and $\overline{\beta} \in \mathbb{N} \langle O \rangle$.

We can alternatively define the transition relation by

$$((q,q'),\alpha',\bar{\beta}) \vdash ((s,s'),\alpha'-a',\bar{\beta}+\bar{b})$$

if there is $a' \subseteq \alpha'$ such that $s=f(q, \Lambda(g'(q', a'))), s'=f'(q', a'), \bar{b}=g(q, \Lambda(g'(q', a'))),$ where $a', \alpha' \in \mathbb{N} \langle V' \rangle, (q, q') \in Q \times Q', \bar{\beta} \in \mathbb{N} \langle O \rangle$. The graphical representation of the cascade product is given in the following figure:



In order to obtain the behaviour of a network of MmA's, we should also consider the behaviour of the cascade product. Roughly speaking, the two types of behaviour depends mainly on the corresponding behaviours of \mathcal{A}' . On the other

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hand, when we have a cascade product, the observable part is strongly connected with the observations that could be made after we pass through \mathcal{A} . We may also emphasize the decisive role played by the connection homomorphism given by Λ . We have the following result:

Theorem 5. Let $\mathcal{A} = (Q, V, O, f, g)$, $\mathcal{A}' = (Q', V', O', f', g')$ be two MmA's, $\mathcal{A}\Omega\mathcal{A}'$ their cascade product, and (q, q') a state of this product. The behaviour of (q, q') is $\mathbf{beh}((q, q')) = \mathbf{beh}(q) \circ \Lambda \circ \mathbf{beh}(q')$.

If we want to get the sequential behaviour starting from $\mathbf{beh}((q,q')) = \mathbf{beh}(q) \circ \Lambda \circ \mathbf{beh}(q')$, then $(\mathbf{I}, \mathbf{I}') \circ \mathbf{seqbeh}((q,q')) = (\mathbf{I} \circ \mathbf{seqbeh}(q)) \circ \Lambda \circ (\mathbf{I}' \circ \mathbf{seqbeh}(q'))$.

What is an Event for Membrane Systems?*

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Summary. Event structures are formal models for parallel and nondeterministic systems. The use of the event structures defines formally where and how the parallelism and nondeterminism appear in systems. We study the event structure for membrane systems, considering both the causality and the conflict relations. We investigate how an event structures can be associated to an parallel evolution step. Two cases are considered: first the rules are applied over strings, and then rules are applied over multisets. We show that, if we apply the standard procedure of extracting event structures from labelled transition systems, then the meanings for event and causality are different for the two cases. The commutativity in the case of multisets introduces some "false" causalities which must be removed in order to capture the right parallelism of the membrane systems.

1 Introduction

Membrane systems [5] describe a computation mechanism inspired by cellular biology. Parallelism and nondeterminism represent the essential features of the membrane computation. Membrane computation is strongly parallel and nondeterministic mainly due to its way of applying the rules of a membrane to its objects. The rules associated with a compartment are applied to the objects from that compartment in a (maximally) parallel way, and the rules are chosen in a non-deterministic manner. Moreover, all compartments of the system evolve at the same time, and so we have two levels of parallelism.

In this paper we study the nature of parallelism and nondeterminism of the membrane systems in terms of a widely recognized formal model for parallelism and nondeterminism, namely in terms of event structures [7].

According to [6], the models for concurrency are classified according to the following criteria: whether they can represent the structure of systems or just their behaviours; whether they can faithfully take into account the difference between parallel and sequential computation (interleaving or non-interleaving model); and

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whether they can represent the branching structure of processes related to nondeterministic choices (linear or branching time model). Event structures are chosen to represent the non-interleaving and branching-time models, and so they model the true concurrency (parallelism) and nondeterminism.

2 Event Structures

In event-based models, a system is represented by a set of events (action occurrences) together with some structure on this set, determining the causality relations between the events. The causality between actions is expressed by a partial order, and the nondeterminism is expressed by a conflict relation on actions. For every two events d and e it is specified either whether one of them is a prerequisite for the other, whether they exclude each other, or whether they may happen in parallel. The behaviour of an event structure is formalized by associating to it a family of configurations representing sets of events which occur during (partial) runs of the system.

A parallel (concurrent) step is simultaneously executing several rules, each of them producing events which end up in the resulting event configuration. These steps are presumably cooperating to achieve a goal, and so they are not totally independent. They synchronize at certain points, and this is reflected in the events produced.

There are many levels of granularity at which one might describe the events that occur as a process executes. At the level of the components from which the system is composed, computation consists of events which reflect the rules. At a higher level, the system might be viewed in terms of parallel executions of its components. Thus, when we talk about the events which occur within a system, it is understood that we know the granularity of the representation, and that events are encoded to this granularity (degree of precision).

Definition 1 (Event Structure).

An event structures is a triple $(E, \leq, \#)$ where

- E is a set of events,
- $\leq \subseteq E \times E$ is a partial order, the causality relation,
- $\# \subseteq E \times E$ is an irreflexive and symmetric relation, the conflict relation, satisfying the principle of conflict heredity:

$$\forall e_1, e_2, e_3 \in E. \ e_1 \leq e_2 \land e_1 \# e_3 \Rightarrow e_2 \# e_3.$$

A more detailed presentation of event structures can be found in [8].

2.1 Event Structure Associated to a Labelled Transition System

There are many cases when the operational semantics of a system is given by means of a labelled transition system (lts) describing all possible sequential computations. In order to study the concurrency properties, we must determine the event structure defined by such a labelled transition system. Let (S, \to, L, s_0) be a transition system, where S is the set o states, \to is the transition relation consisting of triples $(s, \ell, s') \in S \times L \times S$, often written as $s \xrightarrow{\ell} s'$, L is the set of labels (actions), and s_0 is the initial state. A (sequential) computation is a sequence $s_0 \xrightarrow{\ell_1} s_1 \dots \xrightarrow{\ell_n} s_n$ such that $(s_{i-1}, \ell_i, s_i) \in \to$.

Definition 2 (Events in a lts).

Let \sim be the smallest equivalence satisfying: if (s, ℓ_1, s_1) , (s, ℓ_2, s_2) , (s_1, ℓ_2, s_3) , (s_2, ℓ_1, s_3) , $\in \rightarrow$ and $(\ell_1 \neq \ell_2 \text{ or } s_1 \neq s_2)$, then $(s, \ell_1, s_1) \sim (s_2, \ell_1, s_3)$. An event is a \sim -equivalence class written as $[s, \ell, s']$.

Intuitively, two transitions are equivalent iff they are occurrences of the same event. The relation \sim can be easier understood from the following picture:



We have two events which may occur in any order, i.e., the two events are concurrent.

Definition 3 (Configuration in a lts).

A configuration is a multiset of events $[s_{i-1}, \ell_i, s_i]$ corresponding to a computation $s_0 \xrightarrow{\ell_1} s_1 \dots \xrightarrow{\ell_n} s_n$.

Since all computations start from s_0 , each prefix of a configuration is also a configuration.

Theorem 1 (Event Structure of a lts). [4]

 (S, \rightarrow, L, s_0) can be organized as an event structure.

Proof (Sketch). The event structure (E, <, #) is defined by:

- *E* is the set of events as defined in Definition 2;
- $e_1 < e_2$ if every configuration which contains e_2 also contains e_1 ;
- $e_1 \# e_2$ if there is no configuration containing both e_1 and e_2 .

Example 1. The events defined by the lts in Figure 1 are:

$$E_{1} = \{(s_{0}, \ell_{1}, s_{1})\} \qquad E_{4} = \{(s_{1}, \ell_{2}, s_{4}), (s_{3}, \ell_{2}, s_{5})\}$$
$$E_{2} = \{(s_{0}, \ell_{2}, s_{2})\} \qquad E_{5} = \{(s_{2}, \ell_{1}, s_{6})\}$$
$$E_{3} = \{(s_{1}, \ell_{3}, s_{3}), (s_{4}, \ell_{3}, s_{5})\} \qquad E_{6} = \{(s_{4}, \ell_{1}, s_{6})\}$$

A configuration describe a (partial) computation expressed in terms of events. This lts defines the following configurations:

E_1	$E_2 E_5$
E_2	$E_1 E_3 E_4$
$E_1 E_3$	$E_1 E_4 E_3$
$E_1 E_4$	$E_1 E_4 E_6$

We have $E_1 < E_3$ because any configuration containing E_3 also contains E_1 . Since any occurrence of E_3 is always after an occurrence of E_1 , it follows that there is causal relationship between the two events. We get $E_1 < E_4 < E_6$ and $E_2 < E_5$ in a similar way.

Since there is no a configuration containing both E_1 and E_2 , it follows that there is a conflict between the two events, i.e., $E_1 \# E_2$. We get $E_1 \# E_5$, $E_2 \# E_3$, $E_2 \# E_4$, $E_2 \# E_6$, $E_5 \# E_3$, $E_5 \# E_4$, and $E_5 \# E_6$ in a similar way.



Fig. 1. A lts example

3 Event Structure of an Evolution Step

We assume that the reader is familiar with membrane systems (see [5] for a detailed presentation). A membrane structure and their contents, represented as multisets of objects, identify a configuration of a P system. By a nondeterministic and parallel use of rules, the system can pass to another configuration; such a step is called a transition. A sequence of transitions constitutes a computation. Because of the nondeterminism of the application of rules, starting from an initial configuration, we can get several successful computations.

Membrane computation is strongly parallel and nondeterministic mainly due to its way of applying the rules of a membrane to its objects. The rules associated with a compartment are applied to the objects from that compartment in a (maximally) parallel way, and the rules are chosen in a non-deterministic manner. We consider here only membrane systems with a single membrane.

Example 2. Let us consider a simple membrane consisting of the following three rules

$$\ell_1 : a \to b$$

$$\ell_2 : b \to a$$

$$\ell_3 : ab \to d$$

and having the content *aabc*. We investigate the space of all sequential rewritings corresponding to the application of rules in the evolution step $aabc \stackrel{mpr}{\Rightarrow} bbac$ in order to discover the events of this step. The exact definitions for $\stackrel{mpr}{\Rightarrow}$ is given in the next subsections.

3.1 Non-commutative Case

We first assume that the sequential rewritings are executed over non-commutative words (strings).

A context is a string of the form $w \bullet w'$, where w, w' are strings of objects, and • is a special symbol. Each rewriting step $wuw' \to wvw'$ is uniquely determined by the context $w \bullet w'$ and the rule $\ell : u \to v$. Therefore the transition $(wuw', \ell, wvw') = wuw' \xrightarrow{\ell} wvw'$ is denoted by $(w \bullet w', \ell)$.

The maximal parallel rewriting over strings is defined as follows: $w \stackrel{mpr}{\Rightarrow} w'$ if and only if there are $\ell_1, \ldots, \ell_n, \ell_i : u_i \to v_i$ $(i = \overline{1, n})$ such that $w = w_0 u_1 w_1 \ldots u_n w_n$, $w' = w_0 v_1 w_1 \ldots v_n w_n$ and $w_0 w_1 \ldots w_n$ irreducible (no rule can be applied).

We consider first an example. The space of all sequential rewritings for the evolution step of Example 2 is represented in Figure 2.a.

By Definition 4, we have the following three independent events:

$$\begin{split} A &= \{(\bullet a b c, \ell_1), (\bullet b b c, \ell_1), (\bullet b a c, \ell_1), , (\bullet a a c, \ell_1)\}\\ B &= \{(a \bullet b c, \ell_1), (b \bullet b c, \ell_1), (b \bullet a c, \ell_1), , (a \bullet a c, \ell_1)\}\\ C &= \{(a a \bullet c, \ell_2), (a b \bullet c, \ell_2), (b b \bullet c, \ell_2), , (b a \bullet c, \ell_2)\} \end{split}$$



Fig. 2. Lts corresponding to *aabc* $\stackrel{mpr}{\Rightarrow}$ *bbac*

Each event corresponds to the application of a certain evolution rule at a certain position in the string. The resulting event structure corresponds very well to the following description: distribute the object to the rules and then apply the evolutions rules in parallel. The parallel execution of all involved evolution rules is possible because the corresponding events are independent (no causalities, no conflicts). Figure 2.b represents the fact that *bbac* is obtained from *aabc* using the computation space described by the event structure ($\{A, B, C\}, \emptyset, \emptyset$). Any permutation of A, B, C is a computation (in terms of event structures) in this space.

We give now the formal definition for the event structure associated to a mprstep over strings.

Definition 4. The labelled transition system associated to $w \stackrel{\text{mpr}}{\Rightarrow} w'$ is given by all the sequential rewritings starting from w and ending in w'. The event structure ES(w, w') associated to $w \stackrel{\text{mpr}}{\Rightarrow} w'$ is the event structure associated to its labelled transition system.

Theorem 2. The event structure ES(w, w') = (E, <, #) associated to $w \stackrel{mpr}{\Rightarrow} w'$ consists of only independent events, i.e., $< = \emptyset$ and $\# = \emptyset$.

Proof. We have $w \stackrel{mpr}{\Rightarrow} w'$ iff $w = w_0 u_1 w_1 \dots u_n w_n$, $w = w_1 v_1 w_2 \dots v_n w_n$, $\ell_i : u_i \to v_i$ is an evolution rule, for $i = 1, \dots, n$, and $w_1 \dots w_0 w_1 \dots w_n$ is irreducible. The conclusion of the theorem follows by the fact that $[w_0 \dots \bullet w_i \dots w_n, \ell_i]$ is a configuration (any of events can occur first).

3.2 Commutative Case

We assume now that the sequential rewritings are executed over commutative words (multisets).

We write $w =_c w'$ if and only if w' is obtained from w by a permutation of the objects, i.e., w and w' are equal modulo commutativity. Let [w] denote the $=_c$ -equivalence class of w, i.e., $[w] = \{w' \mid w =_c w'\}$.

A context is a string of the form $[\bullet w]$, where w is a multiset of objects, and • is a special symbol. It is easy to see now that the position of • in a context is not important, and therefore we write • at the beginning. Each rewriting step $[uw] \to [vw]$ is uniquely determined by the context $[\bullet w]$ and the rule $\ell : u \to v$. Therefore the transition $([uw], \ell, [vw]) = [uw] \stackrel{\ell}{\to} [vw]$ is denoted by $([\bullet w], \ell)$.

The maximal parallel rewriting over multisets is defined as follows: $[w] \stackrel{mpr}{\Rightarrow} [w']$ iff there are $\ell_1, \ldots, \ell_n, \ell_i : u_i \to v_i \ (i = \overline{1, n})$ such that $[w] = [u_1 \ldots u_n r], [w'] = [v_1 \ldots v_n r$ and r is irreducible (no rule can be applied).

Again we start with an example. The space of all rewritings for the evolution step in Example 2 is:



We also have three events, but they are not totally independent:

$$\overline{A} = \{([\bullet abc], \ell_1), ([\bullet aac], \ell_1)\}$$
$$\overline{B} = \{([\bullet bbc], \ell_1), ([\bullet bac], \ell_1)\}$$
$$\overline{C} = \{([\bullet abc], \ell_2), ([\bullet bac], \ell_2), ([\bullet bbc]\ell_2)\}$$
$$\overline{A} < \overline{B}$$

An event corresponds now to the application of an evolution rule at an arbitrary position. The position in strings cannot be used anymore to distinguish between events. Moreover, between the events \overline{A} and \overline{B} we have a causal dependency: \overline{B} may occur only after \overline{A} . In fact, \overline{A} can be read as "the first application of the evolution rule ℓ_1 " and \overline{B} as "the second application of the evolution rule ℓ_1 ". We notice that the use of commutativity law changes dramatically the meaning of an event.

Definition 5. The labelled transition system associated to $[w] \stackrel{mpr}{\Rightarrow} [w']$ is given by all the sequential rewritings starting from [w] and ending in [w'].

Theorem 3. The event structure (E, <, #) associated to the labelled transition system defined by $[w] \stackrel{\text{mgr}}{\Rightarrow} [w']$ has the following properties:

- $[[\bullet w_1], \ell_1] < [[\bullet w_2], \ell_2]$ if and only if $\ell_1 = \ell_2$, $[[\bullet w_1], \ell_1]$ corresponds to the *i*-th application of the rule ℓ_1 , $[[\bullet w_2], \ell_1]$ corresponds to the *j*-th application of the rule ℓ_1 , and i < j;
- $\# = \emptyset$.

"Maximal parallel" means "no causal dependency" between the application of the rules. We may conclude either that working with multisets is not a good solution at this granularity, namely it is not possible to determine all the parallel rules applied in a mpr-step, or the procedure which determines the event structure from a lts finds "false" causalities for the particular case when the states are given by multisets. We believe that the later one is true; the causality relation given by -th application of a rule (when it is applied more than once) is artificial. Therefore we remove the false causal dependency in the definition of the event structure associated to a mpr-step.

Definition 6. The event structure associated to $[w] \stackrel{\text{mgr}}{\Rightarrow} [w']$ is $ES([w], [w']) = (E, \emptyset, \emptyset)$, where $(E, <, \emptyset)$ is the event structure associated to the labelled transition system defined by $[w] \stackrel{\text{mgr}}{\Rightarrow} [w']$.

4 Event Structure of a Membrane

In this section we determine the event structure given by a membrane. Since the definition for the event concept is different for the two algebraic structures used for contents, we get two definitions for the event structure of a membrane.

We first note that the notation for events is not longer suitable for the case of membranes. We assume that we have a membrane with two evolution rules: ℓ : $a \to b$ and $\ell': b \to a$. Let us consider the computation $aa \stackrel{mpr}{\Rightarrow} bb \stackrel{mpr}{\Rightarrow} aa \stackrel{mpr}{\Rightarrow} bb$. Since the events of $aa \stackrel{mpr}{\Rightarrow} bb$ occur always before the events of $bb \stackrel{mpr}{\Rightarrow} aa$, and the events of $bb \stackrel{mpr}{\Rightarrow} aa$ occur before the events of $aa \stackrel{mpr}{\Rightarrow} bb$, then we get $[a \bullet, \ell] < [b \bullet, \ell'] < [a \bullet, \ell]$, i.e., the causality relation < is cyclic. Therefore each event $[c, \ell]$ in ES(w, w') is denoted with a new fresh name e, and we define $action(e) = [c, \ell]$. In this way, the event sets corresponding to different computation steps are disjoint.

4.1 Non-commutative Case

If the commutativity law is not suitable at this level of granularity, then a question must be answered: who (and how) computes the correct permutation of the objects such that the maximal parallel rewriting is possible? For instance, if the membrane in Example 2 has the content *bbac*, then this content must be permuted to *babc* in order to have $bbac =_c babc \stackrel{\text{mgr}}{\Rightarrow} acc$.

The answer is given by the nondeterministic allocation of the available resources to the rules before the execution of a mpr-step. Such an allocation of resources is considered as an event appearing before a mpr-step. This event consists in the nondeterministic choice of a permutation of available resources, and reflect the nondeterminism of the membrane computation. Once such a permutation is selected, then the next mpr-step is applied over the corresponding string corresponding to this permutation.

We informally describe the event structure of a membrane M. A computation in M is of the form $w_0 =_c w'_0 \stackrel{\text{mpr}}{\Rightarrow} w_1 =_c w'_1 \stackrel{\text{mpr}}{\Rightarrow} \cdots$, where w_0 is the initial contents of M. A string (contents) w is *reachable* in M iff there is a computation from w_0 to w. A reachable string w is *ready-to-fire* if there is a w' such that $w \stackrel{\text{mpr}}{\Rightarrow} w'$.

The event structure $ES(M) = (E_M, <_M, \#_M)$ associated to a membrane M is defined as follows:

- 1. for each reachable w and each w' such that $w \stackrel{mpr}{\Rightarrow} w', ES(w, w') \subseteq ES(M)$ (we recall that the events in ES(w, w') are renamed);
- 2. for each reachable w and each ready-to-fire permutation w' of $w, w \neq w'$, we consider in E_M a distinct event e with action(e) equal to $w =_c w'$, and
 - a) for each reachable w_1 such that $w_1 \stackrel{mpr}{\Rightarrow} w$ and for each event e_1 in $ES(w_1, w)$ we have $e_1 < e$, and
 - b) for each w_2 such that $w' \stackrel{mpr}{\Rightarrow} w_2$ and for each event e_2 in $ES(w', w_2)$ we have $e < e_2$;
- 3. if $e_1, e_2 \in E_M$ such that $action(e_1)$ is $w =_c w_1$ and $action(e_2)$ is $w =_c w_2$ with $w_1 \neq w_2$, then we have $e_1 \# e_2$;
- 4. if $w \stackrel{\text{mpr}}{\Rightarrow} w_1, w \stackrel{\text{mpr}}{\Rightarrow} w_2$ and $w_1 \neq w_2$, then we have $e_1 \# e_2$ in ES(M) for each e_1 in $ES(w, w_1)$ and e_2 in $ES(w, w_2)$.

Example 3. Let M be the membrane presented in Example 2. We consider the computation subspace given by $aabc \stackrel{mpr}{\Rightarrow} bbac =_c babc \stackrel{mpr}{\Rightarrow} acc$ and $aabc \stackrel{mpr}{\Rightarrow} bcc \stackrel{mpr}{\Rightarrow} acc$. We denote by D the event corresponding to $bbac =_c babc$, by $(\{E, F\}, \emptyset, \emptyset)$ the event structure corresponding to $babc \stackrel{mpr}{\Rightarrow} acc$, by $(\{A', B'\}, \emptyset, \emptyset)$ the event structure corresponding to $bcc \stackrel{mpr}{\Rightarrow} acc$. The whole event structure corresponding to the computation subspace is represented in Figure 3.

The events given by item 2 are essential for the definition of the causality relationship. Such an example is the event D in Example 3. We have A, B, C < Dbecause D is causally dependent on A, B, C. The non-deterministic allocation of



Fig. 3. Event structure corresponding to a membrane computation subspace over strings

the resources to rules is given only after all the evolution rules from the previous mpr-step are completely applied. In this way, all ready-to-fire strings obtained from *bbac* have the same probability. However, it is debatable whether we have to distinguish between ready-to-fire strings producing essentially the same computations, e.g., the computation starting from *babc*, *abbc*, *bcab*, *abcb*, *cbab*, and *cabb* are bisimilar. We also have D < E, F because the evolution rules of the next step can be applied only after the resources are allocated to rules. We assume for the moment that $action(E) = [\bullet abc, \ell_2]$ and $action(F) = [b \bullet c, \ell_3]$. By transitivity, we get A < F even if the application of the rule ℓ_3 in F does not use resources produced by the application of the rule ℓ_1 in A.

The conflict relation # is given by the nondeterministic allocation of the available resources to the rules before the execution of a mpr-step (item 3), and the causality relation is given by the repeating evolution steps (item 4). Again, it is debatable if the events corresponding to $bbac =_c babc$, $bbac =_c abbc$, $bbac =_c bcab$, $bbac =_c cabb$, $and bbac =_c cabb$ are really in conflict.

4.2 Commutative Case

In terms of multisets, a computation in M is of the form $[w_0] \stackrel{mpr}{\Rightarrow} [w_1] \stackrel{mpr}{\Rightarrow} \cdots$, where $[w_0]$ is the initial contents of M. A multiset (contents) w is *reachable* in M iff there is a computation from $[w_0]$ to [w].

The construction of the event structure $\overline{ES}(M) = (E_M, <_M, \#_M)$ associated to a membrane M is simpler than that for the case of strings:

- 1. for each reachable [w] and each [w'] such that $[w] \stackrel{mpr}{\Rightarrow} [w'], ES([w], [w']) \subseteq \overline{ES}(M);$
- 2. if $[w] \stackrel{\text{mpr}}{\Rightarrow} [w_1], [w] \stackrel{\text{mpr}}{\Rightarrow} [w_2]$ and $[w_1] \neq [w_2]$, then we have $e_1 \# e_2$ in ES(M) for each e_1 in $ES([w], [w_1])$ and e_2 in $ES([w], [w_2])$;
- 3. if $[w] \stackrel{\text{mpr}}{\Rightarrow} [w_1], [w_1] \stackrel{\text{mpr}}{\Rightarrow} [w_2]$, then we have $e_1 < e_2$ in ES(M) for each e_1 in $ES([w], [w_1])$ and e_2 in $ES([w_1], [w_2])$.

Example 4. We consider the computation subspace given by $[aabc] \stackrel{mpr}{\Rightarrow} [bbac] \stackrel{mpr}{\Rightarrow} [acc]$ and $[aabc] \stackrel{mpr}{\Rightarrow} [bcc] \stackrel{mpr}{\Rightarrow} [acc]$. We denote by $(\{\overline{E}, \overline{F}\}, \emptyset, \emptyset)$ the event structure corresponding to $[babc] \stackrel{mpr}{\Rightarrow} [acc]$, by $(\{\overline{A'}, \overline{B'}\}, \emptyset, \emptyset)$ the event structure corresponding to $[aabc] \stackrel{mpr}{\Rightarrow} [bcc]$, and by $(\{\overline{E'}\}, \emptyset, \emptyset)$ the event structure corresponding to [bcc]. The whole event structure corresponding to the computation subspace is represented in Figure 4.



Fig. 4. Event structure corresponding to a membrane computation space over strings

The allocation of the resources to the rules is not longer given by an explicit event. It is given by the causality relation. For instance, in Example 4 we have A, B, C < E, F. The conflict relation is generated only by the mpr-steps starting from the same multiset (contents) and producing different new multisets (contents).

5 Conclusion

In this paper we study the event structure for membrane systems, considering both the causality and the conflict relations. We investigate how an event structures can be associated to an parallel evolution step. We found that the meaning of an event depends on the algebraic structure used for the contents of membranes: string or multiset.

The construction of the event structures for the cases of priorities and promoters can be reduced to the case of maximal parallel rewriting. A computation step $w \stackrel{pri}{\Rightarrow} w'$ ($[w] \stackrel{pri}{\Rightarrow} [w']$) in the presence of priorities is the same with $w \stackrel{mpr}{\Rightarrow} w'$ ($[w] \stackrel{mpr}{\Rightarrow} [w']$) by taking into account only the rules of maximal priority applicable on w. Similarly, a computation step $w \stackrel{prom}{\Rightarrow} w'$ ($[w] \stackrel{prom}{\Rightarrow} [w']$) in the presence of promoters is the same with $w \stackrel{mpr}{\Rightarrow} w'$ ($[w] \stackrel{mpr}{\Rightarrow} [w']$) where the rules requiring promoters not in w are not considered.

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P Systems with String Objects and with Communication by Request

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Summary. In this paper we study P systems using string-objects where the communication between the regions is indicated by the occurrence of so-called query symbols in the string. We define two variants of communication and prove that these systems with both types of communication are computationally complete, even having a number of membranes limited with relatively small constants.

1 Introduction

In this paper we continue our investigations on P systems with string objects and with communication by request. In [2], the authors studied tissue-like P systems over string objects where the evolution rules of the objects are represented by context-free rewriting rules which also describe the communication between the membranes by the help of communication symbols, called query symbols, one such symbol corresponding to each region of the system.

Membrane systems, or P systems, are distributed and parallel computing devices inspired by the functioning of the living cell [5]. A P system consists of a hierarchically embedded structure of membranes. Each membrane encloses a region that contains objects and might also contain other membranes. There are rules associated to the regions describing the evolution and the movement of the objects which together correspond to a computation. For details on membrane systems, see the monograph [6] and consult the web-page http://psystems.disco.unimib.it.

While in the standard case a P system consists of a hierarchically embedded structure of membranes, tissue-like P systems are organized in another manner [4]. Instead of an individual cell, these correspond to groups of cells, like tissues or organs, interacting with each other either directly or with the use of the environment, but in any case, having the common property that the membrane structures are not necessarily described by a tree as the ones corresponding to individual cells.

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Communication in tissue-like P systems with string objects and with communication by request was defined as follows: When one or more query symbols are introduced in a string, then the rewriting of that string stops and the queries are satisfied by replacing the query symbols with strings which do not contain further query symbols from the region indicated by the query symbol, in all possible combinations. If no query symbol free string exists in the queried region, then the string containing the query disappears.

This model has some biological resemblance: if the strings are considered as descriptions of simple organisms, the query symbols as their "weak points", possibly infected or attacked by another organism, then the communication mimics some features of an infection or parasitism. Inspired by these resemblances, we call a communication of type i (infection) if after communicating the copies of the strings, the strings themselves remain in the region, while the communication is called of type p (parasitism), if after communication the communicated string itself disappears from its original region. The model is called an MPC system in short.

MPC systems can also be considered as modified variants of parallel communicating (PC) grammar systems defined over multisets of strings. PC grammar systems are networks of grammars organized in a communicating system to generate a single language. The reader interested in the theory of grammar systems is referred to [1, 7].

In [2], the authors proved that MPC systems with 7 membranes and working with *i*-communication are able to describe all recursively enumerable languages. The computational completeness of these systems working with *p*-communication holds as well, even for a subclass consisting of systems having only 9 membranes.

In this paper we define the two types of communication for standard P systems and examine the computational power and the size complexity of these models. We call the new constructs RPC systems in short. In this case, the requested string can only be communicated either to the parent membrane or to one of the child membranes, depending on the issued query symbol. Thus, query symbols refer only to the neighboring regions. According to the above mentioned biological resemblance, both infection and parasitism are very local phenomena regarding their spread, i.e., in one step only the neighbors can be infected and parasitism can be developed only between two closely related, i.e. neighbor components. As for MPC systems, the computational completeness can be proved for RPC systems with both types of communication: in the case of i-communication systems with 10 membranes and in the case of p-communication systems with 30 membranes are enough for demonstrating the power of the Turing machines. The reader can observe that both MPC systems and P systems are able to obtain the computational completeness even with relatively small number of membranes. Moreover, in the case of *i*-communication the difference between the two numbers is very small, i.e. the difference in the underlying structure of the membrane system has not too much influence on the computational power of the system.

2 Preliminaries and Definitions

We first recall the notions and the notations we use. The reader is assumed to be familiar with the basics of formal language theory, for details see [7]. Let Σ be an alphabet and let Σ^* be the set of all words over Σ , that is, the set of finite strings of symbols from Σ , and let $\Sigma^+ = \Sigma^* - \{\varepsilon\}$ where ε denotes the empty word. For $w \in \Sigma^*$ and $S \subseteq \Sigma$, let $|w|_S$ denote the number of occurrences of symbols from S in the string w (if $S = \{a\}$ is a singleton set, we may write $|w|_a$ instead of $|w|_{\{a\}}$).

Let V be a set of objects, and let \mathbb{N} denote the set of non-negative integers. A multiset is a mapping $M : V \to \mathbb{N}$ which assigns to each object $a \in V$ its multiplicity M(a) in M. The support of M is the set $supp(M) = \{a \mid M(a) \ge 1\}$. If supp(M) is a finite set, then M is called a finite multiset. The set of all finite multisets over the set V is denoted by V° .

We say that $a \in M$ if $M(a) \geq 1$. For two multisets $M_1, M_2 : V \to \mathbb{N}, M_1 \subseteq M_2$ if for all $a \in V, M_1(a) \leq M_2(a)$. The union of M_1 and M_2 is defined as $(M_1 \cup M_2) : V \to \mathbb{N}$ with $(M_1 \cup M_2)(a) = M_1(a) + M_2(a)$ for all $a \in V$, the difference is defined for $M_2 \subseteq M_1$ as $(M_1 - M_2) : V \to \mathbb{N}$ with $(M_1 - M_2)(a) = M_1(a) - M_2(a)$ for all $a \in V$, and the intersection is $(M_1 \cap M_2) : V \to \mathbb{N}$ with $(M_1 \cap M_2)(a) = min(M_1(a), M_2(a))$ for $a \in V$, where min(x, y) denotes the minimum of $x, y \in \mathbb{N}$. We say that M is empty, denoted by ϵ , if its support is empty, $supp(M) = \emptyset$.

In the following we sometimes list elements a_1, \ldots, a_n of a multiset as $M = \{\{a_1, \ldots, a_n\}\}$, by using double brackets to distinguish from the usual set notation.

A P system is a structure of hierarchically embedded membranes, each having a label and enclosing a region containing a multiset of objects and possibly other membranes. The out-most membrane which is unique, is called the skin membrane. The membrane structure is denoted by a sequence of matching parentheses where the matching pairs have the same label as the membranes they represent. If membrane l_i of a given membrane structure μ contains membrane l_j , and there is no other membrane, l_k , such that l_k contains l_j and l_i contains l_k , then we say that membrane l_i is the parent membrane of l_j , denoted as $parent_{\mu}(l_j) = l_i$, and l_j is one of the child membranes of l_i , denoted as $l_j \in child_{\mu}(l_i)$. We also define for any region l_i the set of regions $neighbor_{\mu}(l_i) = \{l_j \mid parent_{\mu}(l_i) = l_j \text{ or } l_j \in child_{\mu}(l_i)\}$.

The evolution of the contents of the regions of a P system is described by rules associated to the regions. Applying the rules synchronously in each region, the system performs a computation by passing from one configuration to another one. Several variants of the basic notion have been introduced and studied proving the power of the framework, see the monograph [6] for a summary of notions and results of the area.

In the following we focus on systems where the objects are represented with strings, object evolution is modeled by context-free string rewriting rules, and communication is performed by dynamically emerging requests with the use of query symbols appearing in the string objects.

Definition 1 A string rewriting P system with communication by request or an RPC system (of degree $m \ge 1$) is a construct

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$$\Pi = (V, \mu, (M_1, R_1), \dots, (M_m, R_m), i_o),$$

where:

- $V = N \cup T \cup K$ where N, T, K are pairwise disjoint alphabets of noterminals, terminals, and *query symbols*, respectively, with $K = \{Q_1, \ldots, Q_m\}$ (one query symbol is associated to each region of Π);
- μ is a membrane structure of *m* membranes;
- M_1, \ldots, M_m are finite multisets over $(N \cup T)^*$;
- R_1, \ldots, R_m are finite sets of context-free rewriting rules of the form $A \to u$, with $A \in N$ and $u \in V^*$;
- $i_o \in \{1, 2, \dots, m\}$ is the index of the *output* membrane of Π .

The work of such a system starts from the initial configuration (M_1, \ldots, M_m) . It passes from a configuration (M'_1, \ldots, M'_m) , consisting of multisets of strings over $N \cup T \cup K$ placed in the *m* regions of the system, to another configuration (M''_1, \ldots, M''_m) in the following way. If no query symbol is present in the strings contained by the system, then each string from each multiset M'_i is rewritten which can be rewritten according to the rules from $R_i, 1 \leq i \leq m$. This means the use of one rule from R_i , non-deterministically chosen, for each string. The strings which cannot be rewritten (no rule can be applied to them) remain unchanged. The resulting multisets of strings are $M''_i, 1 \leq i \leq m$. Note that the rewritten must be rewritten, and that the process is non-deterministic, the choice of rules and the places where the rules are applied can lead to several possible new multisets of strings.

If any query symbol is present in any of the strings contained by M'_i , $1 \le i \le i$ n, then a communication is performed: Each symbol Q_j introduced in a string present in region i (that is, in the multiset M'_i), where j is the index of one of the neighboring regions, is replaced with all strings from this neighboring region j which do not contain query symbols. If in region j there are several strings without query symbols, then each of them is used, hence the string from region i is replicated (with the occurrence of Q_j replaced with strings from region j). If there are several query symbols in the same string from component i, then all of them are replaced (we also say that they are *satisfied*) at the same time, in all possible combinations. If a query symbol Q_i cannot be satisfied (region j contains no string without query symbols), then the string containing Q_j is removed (it is like replacing it with the strings from an empty language). We call such a system *i*-communicating if copies of the requested strings are communicated to the requesting components, and *p*-communicating if after replacing the query symbols with the requested strings, these strings are removed from the multiset associated to the queried region.

In this way, all query symbols introduced by the rewriting rules disappear, they are either satisfied (replaced by strings without query symbols) or they disappear together with the string which contain them (in the case when they cannot be satisfied). The multisets obtained in this way in one communication step are M''_1, \ldots, M''_m , constituting the next configuration of the system.

We give now the formal definition of the transition.

Definition 2 Let $\Pi = (V, \mu, (M_1, R_1), \ldots, (M_m, R_m), i_o)$ be an RPC system as above, and let (M'_1, \ldots, M'_m) and (M''_1, \ldots, M''_m) be two configurations of Π . We say that (M'_1, \ldots, M'_m) directly derives (M''_1, \ldots, M''_m) , if one of the following two cases holds.

- 1. There is no string containing query symbols, that is, $x \in (N \cup T)^*$ for all $x \in \bigcup_{i=1}^m M_i$. In this case, if $M'_i = \{\{x_{i,1}, \ldots, x_{i,t_i}\}\}$, then $M''_i = \{\{y_{i,1}, \ldots, y_{i,t_i}\}\}$ where either $x_{i,j} \Rightarrow y_{i,j}$ according to a context-free rule of R_i , or $y_{i,j} = x_{i,j}$ if there is no rule in R_i which can be applied to $x_{i,j}$, $1 \le j \le t_i$, $1 \le i \le m$.
- 2. There is at least one $x \in \bigcup_{i=1}^{m} M_i$ such that $|x|_K > 0$. In this case, rewriting is stopped and a communication step must be performed as follows. Let

$$M_i^{req} = \begin{cases} \{ x \in M_i' \mid |x|_K = 0 \} \} \text{ if there is a } j \in neighbor_{\mu}(i), \text{ such that } y \in M_j' \text{ with } |y|_{Q_i} > 0, \\ \emptyset \qquad \qquad \text{otherwise,} \end{cases}$$

 let

$$M_i^{avail} = \{ \{ x \in M_i' \mid |x|_K = 0 \} \}$$

for all $i, 1 \le i \le m$, and let for an $x = x_1 Q_{i_1} x_2 Q_{i_2} \dots Q_t x_{t+1}, x_j \in (N \cup T)^*, Q_{i_j} \in K, 1 \le j \le t+1,$

$$Sat(x) = \begin{cases} \{ \{x_1y_{i_1}x_2y_{i_2}\dots y_{i_t}x_{t+1} \mid y_{i_j} \in M_{i_j}^{avail} \} \} \text{ if } M_{i_j}^{avail} \neq \emptyset \text{ for} \\ \\ \emptyset & \text{all } i_j, \ 1 \le j \le t, \\ \emptyset & \text{otherwise.} \end{cases}$$

Now, for all $i, 1 \leq i \leq m$,

$$M_i'' = M_i' - M_i^{req} - \{\{x \in M_i' \mid |x|_K > 0\}\} + \bigcup_{x \in M_i', |x|_K > 0} Sat(x)$$

in the *p*-communicating mode, and

$$M_i'' = M_i' - \{\{x \in M_i' \mid |x|_K > 0\}\} + \bigcup_{x \in M_i', |x|_K > 0} Sat(x)$$

in the *i*-communicating mode.

Let us denote the transitions from one configuration to another, (M'_1, \ldots, M'_m) to (M''_1, \ldots, M''_m) , by $(M'_1, \ldots, M'_m) \Rightarrow_X (M''_1, \ldots, M''_m)$ with X = i and X = p for *i*-communicating and *p*-communicating systems, respectively.

The language generated by the RPC system consists of all terminal strings produced in region i_o during any possible computation in Π .

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$$L_X(\Pi) = \{ x \in T^* \mid (M_1, \dots, M_m) \Rightarrow^*_X (M'_1, \dots, M'_m) \text{ and } x \in M'_{i_o} \}$$

for $X \in \{i, p\}$, where \Rightarrow_X^* denote the reflexive and transitive closure of \Rightarrow_X .

The families of all languages generated in this way by RPC systems of degrees at most $m \ge 1$ with *i*-communication or *p*-communication, is denoted by $iRPC_mCF$ and $pRPC_mCF$, respectively. If we use systems of an arbitrary degree, then we replace the subscript *m* with *. Let us also denote the class of recursively enumerable languages by *RE*.

RPC systems are computationally universal; they characterize the class of recursively enumerable languages, even with a limited number of components.

Before proving this result, we recall the notion of a two-counter machine from [3]. A two-counter machine $TCM = (T \cup \{Z, B\}, E, R)$ is a 3-tape Turing machine where T is an alphabet, E is a set of internal states with two distinct elements $q_0, q_F \in E$, and R is a set of transition rules. The machine has a read-only input tape and two semi-infinite storage tapes (the counters). The alphabet of the storage tapes contains only two symbols, Z and B (blank), while the alphabet of the input tape is $T \cup \{B\}$. R contains transition rules of the form $(q, x, c_1, c_2) \rightarrow (q', e_1, e_2)$ where $x \in T \cup \{\varepsilon\}$ corresponds to the symbol scanned on the input tape in state $q \in E$, and $c_1, c_2 \in \{Z, *\}$ correspond to the symbols scanned on the storage tapes. If $c_i = Z$, then the symbol scanned on the *i*th counter tape is Z, if $c_i = *$, then the symbol scanned is either Z or B. By a rule of this form, M enters state $q' \in E$, and the counters are modified according to $e_1, e_2 \in \{-1, 0, +1\}$. If $x \in T$, then the machine was scanning x on the input tape, and the head moves one cell to the right; if $x = \varepsilon$, then the machine performs the transition irrespective of the scanned input symbol, and the reading head does not move.

The symbol Z appears initially on the cells scanned by the storage tape heads and may never appear on any other cell. An integer t can be stored by moving a tape head t cells to the right of Z. A stored number can be incremented or decremented by moving the tape head right or left. The machine is capable of checking whether a stored value is zero or not by looking at the symbol scanned by the storage tape heads. If the scanned symbol is Z, then the value stored in the corresponding counter is zero. Note that although we do not allow to explicitly check the non-emptiness of the counters which is allowed in [3], this feature can be simulated: After successfully decrementing and incrementing a counter, the stored value is not altered, but the machine can be sure that the scanned symbol is B. A word $w \in T^*$ is accepted by the two counter machine if the input head has read the last non-blank symbol on the input tape, and the machine is in the accepting state q_F . Two-counter machines are computationally complete; they are just as powerful as Turing-machines, see [3].

Theorem 1. $iRPC_{10}CF = RE$.

Proof. We only give the proof of the inclusion $RE \subseteq iRPC_{10}CF$. The reverse inclusion follows from the Church thesis. To this aim, let us consider a recursively enumerable language $L \subseteq T^*$ and a two-counter machine TCM =

 $(T \cup \{Z, B\}, E, R)$, in the above form, accepting the language L. We construct an RPC system L. Let

$$\Pi = (V, \mu, C_{sel}, C_{gen}, C_{ch_1}, C_{S_4}, C_{c_1}, C_{ind_1}, C_{ch_{2,1}}, C_{c_2}, C_{ind_2}, C_{ch_{2,2}}, sel)$$

where $C_{\alpha} = (M_{\alpha}, R_{\alpha})$ for $\alpha \in \{sel, gen, c_1, c_2, ind_1, ind_2, ch_1, S_4, ch_{2,1}, ch_{2,2}\}, \mu = [[[]_{S_4} []_{ind_1} []_{ch_{2,1}}]_{c_1} []_{ind_2} []_{ch_{2,2}}]_{c_2}]_{ch_1}]_{gen}]_{sel}$ and $V = N \cup K \cup T$.

Let $D = \{[q, x, c_1, c_2, q', e_1, e_2] \mid (q, x, c_1, c_2) \rightarrow (q', e_1, e_2) \in R\}$, and let us define for any $\alpha = [q, x, c_1, c_2, q', e_1, e_2] \in D$, the following notations: $State(\alpha) = q$, $Read(\alpha) = x$, $NextState(\alpha) = q'$, and $Store(\alpha, i) = c_i$, $Action(\alpha, i) = e_i$ for i = 1, 2.

The general idea of the simulation is to represent the states and the transitions of TCM with nonterminals of D and the values of the counters by strings of nonterminals containing as many A symbols as the value stored in the given counter. Let

 $N = \{S'_i, F_i, B_i \mid 1 \le i \le 6\} \cup \{S_i, C_i \mid 1 \le i \le 7\} \cup \{\alpha_i, H_i \mid \alpha \in D, 1 \le i \le 8\} \cup \{\alpha'_1, \alpha''_1, \bar{\alpha}'_1, \bar{\alpha}''_1, \bar{\alpha}'_1 \mid \alpha \in D\} \cup \{F_{1,i} \mid 1 \le i \le 5\} \cup \{J_i \mid 1 \le i \le 4\} \cup \{A, F'_1, \underline{F}'_1, \underline{F}'_1, F''_1, E, E_1, I, J, S''_6\} \text{ and let the rules be defined as follows.}$

$$\begin{split} M_{sel} &= \{\{I\}\},\\ R_{sel} &= \{I \to \alpha_1 \mid \alpha \in D, State(\alpha) = q_0\} \cup\\ &\{\alpha_8 \to \beta_1 \mid \alpha, \beta \in D, NextState(\alpha) = State(\beta)\} \cup\\ &\{\alpha_8 \to F_1 \mid \alpha \in D, NextState(\alpha) = q_F\} \cup\\ &\{\alpha_i \to \alpha_{i+1} \mid \alpha \in D, 1 \le i \le 7\} \cup \{F_i \to F_{i+1} \mid 1 \le i \le 5\} \cup\\ &\{F_6 \to Q_{gen}, S'_6 \to S'_6, S'_6 \to Q_{gen}, E \to \varepsilon, \bar{\alpha}''_1 \to \varepsilon\}. \end{split}$$

This region keeps track of the current state of the simulated two-counter machine and also selects the transition to be simulated. The symbol I is used to initialize the system by introducing one of the initial transition symbols of the form $[q_0, x, c_1, c_2, q', e_1, e_2]_1$ where q_0 is the initial state. It also produces the result of the computation when after simulating the entering of the counter machine into the final state (that is, after the appearance of the nonterminal F_1), it receives the strings produced in the lower regions and erases the occurrences of the nonterminals E which, if the simulation was successful, produces a terminal word accepted by the two-counter machine.

$$\begin{split} M_{gen} &= \{\{S_1, S_1'\}\},\\ R_{gen} &= \{S_1 \to S_2, S_2 \to Q_{sel}, S_1' \to Q_{sel}\} \cup \\ &\{\alpha_1 \to \alpha_1', \alpha_2 \to Q_{sel}, \alpha_i \to \alpha_{i+1} \mid \alpha \in D, 3 \leq i \leq 6\} \cup \\ &\{\alpha_1' \to \alpha_1'', \alpha_1'' \to S_2', \alpha_7 \to xS_1 \mid \alpha \in D, Read(\alpha) = x\} \cup \\ &\{F_1 \to F_{1,1}, F_{1,i} \to F_{1,i+1}, F_{1,5} \to Q_{ch_1} \mid 1 \leq i \leq 4\} \cup \\ &\{S_i' \to S_{i+1}' \mid 2 \leq i \leq 4\} \cup \{S_5' \to S_1', A \to \varepsilon, J \to \varepsilon\} \cup \\ &\{F_2 \to Q_{sel}, F_i \to F_{i+1} \mid 3 \leq i \leq 4\} \cup \{F_5 \to Q_{ch_1}\} \cup \\ &\{S_6' \to S_6'', H_7 \to \varepsilon\}. \end{split}$$

This region generates the string accepted by the counter machine by adding the symbol $Read(\alpha)$ for each $\alpha \in D$ chosen in the selector region. After the appearance of the nonterminal F_1 in the system, this region will append the words from the checking region ch_1 to its own string and send this string which also contains the generated word to the region *sel*. Then it will receive the word from region ch_2 and erase all A and J symbols before forwarding it also to region *sel*.

$$\begin{split} M_{ch_{1}} &= \{\{S_{1}, S_{1}'\}\},\\ R_{ch_{1}} &= \{S_{1} \rightarrow S_{2}, S_{2} \rightarrow S_{3}, S_{3} \rightarrow Q_{gen}, S_{1}' \rightarrow S_{2}', S_{2}' \rightarrow Q_{gen}\} \cup \\ &\{\alpha_{1}'' \rightarrow \delta_{1}\delta_{2}Q_{S_{4}} \mid \alpha \in D, \delta_{j} = Q_{c_{j}} \text{ if } Store(\alpha, j) = Z,\\ &\text{ or } \delta_{j} = \varepsilon \text{ otherwise}\} \cup \{S_{i} \rightarrow S_{i+1} \mid 4 \leq i \leq 6\} \cup \\ &\{\alpha_{1}' \rightarrow \bar{\alpha}_{1}', \bar{\alpha}_{1}' \rightarrow \underline{\bar{\alpha}}_{1}', \underline{\bar{\alpha}}_{1}' \rightarrow S_{3}', S_{3}' \rightarrow S_{4}', S_{4}' \rightarrow S_{5}'\} \cup \\ &\{F_{1,1} \rightarrow \bar{F}_{1}', \bar{F}_{1}' \rightarrow \underline{\bar{F}}_{1}', \underline{\bar{F}}_{1}' \rightarrow \underline{\bar{F}}_{1}', \underline{\bar{F}}_{1}' \rightarrow Q_{c_{1}}Q_{c_{2}}\} \cup \\ &\{S_{7} \rightarrow S_{1}, S_{5}' \rightarrow S_{1}', F_{1,2} \rightarrow Q_{S_{4}}\}, \text{ and} \\ M_{S_{4}} &= \{\{S_{4}\}\},\\ R_{S_{4}} = \emptyset. \end{split}$$

The region ch_1 checks whether the counter contents are zero when they should be zero by collecting the counter strings from regions c_1, c_2 when necessary. At the end of the simulation, the collected string is forwarded to the region *gen* and then to region *sel*, where a terminal string can only be produced if the word originating in region ch_1 contains no A symbols.

For j = 1, 2, let

$$\begin{split} M_{c_j} &= \{\{J, C_1\}\}, \\ R_{c_j} &= \{J \rightarrow J_1, J_1 \rightarrow J_2, J_2 \rightarrow Q_{ch_1}, A \rightarrow Q_{ind_j}, J_3 \rightarrow Q_{ind_j}, J_4 \rightarrow J\} \cup \\ &\{\bar{\alpha}'_1 \rightarrow \bar{\alpha}''_1, \bar{\alpha}''_1 \rightarrow \delta_{\alpha} J_3, \underline{\bar{\alpha}}'_1 \rightarrow C_5 \mid \alpha \in D, \delta_{\alpha} = A \text{ if } Action(\alpha, j) = 0, \\ &\delta_{\alpha} = AA \text{ if } Action(\alpha, j) = +1, \delta_{\alpha} = \varepsilon \text{ if } Action(\alpha, j) = -1\} \cup \\ &\{C_i \rightarrow C_{i+1}, C_4 \rightarrow Q_{ch_1}, C_7 \rightarrow C_1 \mid i \in \{1, 2, 3, 5, 6\} \} \cup \\ &\{\bar{F}'_1 \rightarrow \bar{F}''_1, \underline{\bar{F}}'_1 \rightarrow Q_{ch_{2,j}}, E \rightarrow E_1, H_6 \rightarrow H_7\}. \end{split}$$

These regions maintain strings representing the contents of the two counters. After the selection of a transition symbol in the region corresponding to C_{sel} , they execute the action required by the chosen transition symbol by adding AA, A, or ε to the counter string and then deleting one A and J_3 by rewriting it to Q_{ind_j} . The simulation can only be successful if exactly one A and the symbol J_3 is rewritten. This is ensured by region C_{ind_j} . If there is a string obtained after the two queries which contain only a number of A or E symbols and one J_4 symbol, then the simulation of the actions required by the chosen transition was successful. If a counter is empty, this construction also forbids the successful execution of the decrement instruction since this would introduce E_1 in the counter strings.

The rules of the region supporting the work of the counters C_{c_j} , j = 1, 2, are defined as follows.

Table 1. Components of Π in the proof of Theorem 1, simulating an instruction with $\alpha = [q, x, Z, B, q', +1, -1]$. The region C_{S_4} is omitted since it always contains the string S_4 .

	C_{sel}	C_{gen}	C_{ch_1}	C_{c_1}	C_{ind}	$C_{ch_{2,1}}$
0	β_8	wS_1, S_1'	ES_1, S'_1	EJ, C_1	B_1	$AEJH_1$
1	α_1	wS_2, Q_{sel}	ES_2, S'_2	EJ_1, C_2	B_2	$AEJH_2$
1C	α_1	wS_2, α_1	ES_2, S'_2	EJ_1, C_2	B_2	$AEJH_2$
2	α_2	wQ_{sel}, α'_1	ES_3, Q_{gen}	EJ_2, C_3	B_3	$AEJH_3$
2C	α_2	$w\alpha_2, \alpha_1'$	ES_3, α'_1	EJ_2, C_3	B_3	$AEJH_3$
3	α_3	wQ_{sel}, α_1''	$EQ_{gen}, \bar{\alpha}'_1$	EQ_{ch_1}, C_4	B_4	$AEJH_4$
3C	α_3	$w\alpha_3, \alpha_1''$	$E\alpha_1'', \bar{\alpha}_1'$	$E\bar{\alpha}_1', C_4$	B_4	$AEJH_4$
4	α_4	$w\alpha_4, S'_2$	$EQ_{c_1}Q_{S_4}, \underline{\bar{\alpha}}_1'$	$E\bar{\alpha}_1'', Q_{ch_1}$	B_5	$AEJH_5$
[4C]	α_4	$w\alpha_4, S'_2$	$E\bar{\alpha}_1''S_4, \underline{\bar{\alpha}}_1'$	$E\bar{\alpha}_1'', \underline{\bar{\alpha}}_1'$	B_5	$AEJH_5$
5	α_5	$w\alpha_5, S'_3$	$E\bar{\alpha}_1''S_5, S_3'$	$E\delta_{\alpha}J_3, C_5$	B_6	$AEJH_6$
6	α_6	$w\alpha_6, S'_4$	$E\bar{\alpha}_1''S_6, S_4'$	$EQ_{ind}J_3, C_6$	E	$AEJH_7$
[6C]	α_6	$w\alpha_6, S'_4$	$E\bar{\alpha}_1''S_6, S_4'$	EEJ_3, C_6	E	$AEJH_7$
7	α_7	$w\alpha_7, S_5'$	$E\bar{\alpha}_1''S_7, S_5'$	EEQ_{ind}, C_7	J_4	$AEJH_8$
[7C]	α_7	$w\alpha_7, S_5'$	$E\bar{\alpha}_1''S_7, S_5'$	EEJ_4, C_7	J_4	$AEJH_8$
8	α_8	wxS_1, S_1'	$E\bar{\alpha}_1''S_1, S_1'$	EEJ, C_1	B_1	$AEJQ_{c_1}H_1$
8C	α_8	wxS_1, S'_1	$E\bar{\alpha}_1''S_1, S_1'$	EEJ, C_1	B_1	$AEJH_1$

$$M_{ind_j} = \{\{B_1\}\},\$$

$$R_{ind_j} = \{B_i \to B_{i+1} \mid 1 \le i \le 5\} \cup \{B_6 \to E, E \to J_4, J_4 \to B_1\},\$$

and

$$\begin{split} M_{ch_{2,j}} &= \{\{H_1\}\},\\ R_{ch_{2,j}} &= \{H_i \to H_{i+1} \mid 1 \le i \le 7\} \cup \{H_8 \to Q_{c_j}H_1\} \end{split}$$

Instead of giving a detailed proof of the correctness of our construction, we demonstrate the work of the system in Table 1 and Table 2 by indicating a possible transition sequence of Π while simulating an instruction of the two-counter machine TCM, and by presenting the terminating part of the simulation. Note that the cells of the tables contain only some of the strings produced by the regions, those which are interesting from the point of view of the simulation.

Let us first look at Table 1. The simulated instruction is represented by a nonterminal $\alpha_1 = [q, x, Z, *, q', +1, -1]_1$ chosen in region C_{sel} in the first step. This indicates that the first counter should be empty which requirement is satisfied since region C_{c_1} contains a string containing zero A symbols. In the following few steps, the indexed versions of α reach the regions $C_{gen}, C_{ch_1}, C_{c_j}, j \in \{1, 2\}$, and each of these regions executes its part of the simulation. C_{gen} generates the letter read by the two-counter machine, C_{ch_1} queries the regions simulating the counters in the case when their contents should be zero, and this way collects a "checker"

string. If this string contains the nonterminal A, then the simulation is not correct. C_{c_j} maintain the contents of the counters by adding or deleting A-s. Its work is aided by C_{ind} and $C_{ch_{2,j}}$. The region $C_{ch_{2,j}}$ collects the counter strings at the end of each simulating cycle. The simulation was successful if and only if this collected string only contains A, E or J symbols.

The terminating phase of the simulation is presented on Table 2. When G_{sel} selects the symbol F, the system prepares to finish its work. The variously indexed versions of F travel through the system and result in the transfer of the word generated in G_{gen} and the checker string of region C_{ch_1} to region C_{sel} . There the symbol A cannot be erased, so a terminal word can only be produced if the checker string does not contain this symbol. Meanwhile, the other checker strings are transferred from $C_{ch_{2,j}}$ to region C_{gen} where the A and J symbols can be erased, but nothing else, so when later also this string is transferred to C_{sel} , a terminal string can only be produced if the behavior of the counter simulating regions were correct in each step of the simulation. The last row of the table represents the situation when the erasing process begins. When all A and J have disappeared from the string in region C_{gen} , then S'_6 can be changed to a query symbol transferring the result to C_{sel} , where all remaining symbols can be erased, in the case when the simulation was correct.

Next we prove that any RPC system using i-communication can be simulated with an RPC system using p-communication.

Theorem 2. $iRPC_nCF \subseteq pRPC_{3n}CF$, for any $n \ge 1$.

Proof. Let $\Pi = (V, \mu, (M_1, R_1), \dots, (M_n, R_n), 1)$ be a system of degree n with $V = N \cup K \cup T$. We construct $\Pi' = (V', \mu', (M'_1, R'_1), \dots, (M'_{3n}, R'_{3n}), 1)$ of degree 3n, such that $L_p(\Pi') = L_i(\Pi)$.

Let μ' be defined by adding two new regions $[[]_{2n+i}]_{n+i}$ inside every region *i*. This way, $n+i \in neighbor_{\mu'}(i)$, and $2n+i \in neighbor_{\mu'}(n+i)$ for all $1 \leq i \leq n$.

Let $V' = N' \cup K' \cup T$ where $N' = N \cup \{S_1, S_2, S_3, S_4\}$, and let the rules of Π' be defined as

$$M'_{i} = M_{i} \cup \{\{S_{1}, S_{2}\}\},\$$

$$R'_{i} = R_{i} \cup \{S_{1} \to Q_{i}, S_{2} \to Q_{n+i}\},\$$

and

$$M'_{n+i} = \{\{S_1, S_2, S_3, S_4\}\}, \ R'_{n+i} = \{S_3 \to Q_{n+i}, S_4 \to Q_{2n+i}\}, \\ M'_{2n+i} = \{\{S_1, S_2, S_3, S_4\}\}, \ R'_{2n+i} = \{S_1 \to Q_{n+i}, S_2 \to Q_{2n+i}\}$$

for all $1 \leq i \leq n$.

The additional membranes of Π' work as "suppliers" of symbols. In each step, each region *i* rewrites S_1 and S_2 to query itself, and the region n+i. From itself it "receives" the strings it contains besides S_1, S_2 , from region n+i it receives S_1, S_2 , so the same behavior can be repeated in the next step. This self query mechanism

	Csel	C_{gen}	C_{ch_1}	C_{c_1}	C_{ind}	$C_{ch_{2,1}}$
0	β_8	wS_1, S_1'	ES_1, S'_1	EJ, C_1	B_1	$AEJH_1$
1	F_1	wS_2, Q_{sel}	ES_2, S'_2	EJ_1, C_2	B_2	$AEJH_2$
$1\mathrm{C}$	F_1	wS_2, F_1	ES_2, S'_2	EJ_1, C_2	B_2	$AEJH_2$
2	F_2	$wQ_{sel}, F_{1,1}$	ES_3, Q_{gen}	EJ_2, C_3	B_3	$AEJH_3$
2C	F_2	$wF_2, F_{1,1}$	$ES_3, F_{1,1}$	EJ_2, C_3	B_3	$AEJH_3$
3	F_3	$wQ_{sel}, F_{1,2}$	EQ_{gen}, \bar{F}'_1	EQ_{ch_1}, C_4	B_4	$AEJH_4$
3C	F_3	$wF_3, F_{1,2}$	$EF_{1,2}, \bar{F}'_1$	$E\bar{F}'_1, C_4$	B_4	$AEJH_4$
4	F_4	$wF_4, F_{1,3}$	$EQ_{S_4}, \underline{\bar{F}}_1'$	$E\bar{F}_1'', Q_{ch_1}$	B_5	$AEJH_5$
$4\mathrm{C}$	F_4	$wF_4, F_{1,3}$	$ES_4, \underline{\bar{F}}_1'$	$E\bar{F}_1^{\prime\prime}, \underline{\bar{F}}_1^\prime$	B_5	$AEJH_5$
5	F_5	$wF_5, F_{1,4}$	$ES_5, \underline{\bar{F}}'_1$	$E\bar{F}_1^{\prime\prime}$	B_6	$AEJH_6$
				$Q_{ch_{2,1}}$		
5C	F_5	$wF_5, F_{1,4}$	$ES_5, \underline{\bar{F}}'_1$	$E\bar{F}_1^{\prime\prime}$	B_6	$AEJH_6$
				$AEJH_6$		
6	F_6	wQ_{ch_1}	ES_{6}	$E\bar{F}_{1}^{\prime\prime}$	E	$AEJH_7$
		$F_{1,5}$	$Q_{c_1}Q_{c_2}$	$AEJH_7$		
6C	F_6	wES_6	ES_6	$E\bar{F}_1''$	E	$AEJH_7$
		$F_{1,5}$	$AEJH_7$	$AEJH_7$	E	$AEJH_7$
7	Q_{gen}	wES_6	ES_7	$E\bar{F}_{1}''$	J_4	$AEJ\overline{H_8}$
		Q_{ch_1}	$AEJH_7$	$AEJH_7$		
7C	wES_6	wES_6'	ES_{7}	$E\overline{F}_{1}^{\prime\prime}$	J_4	$AEJH_8$
		$AEJH_7$	$AEJH_7$	$AEJH_7$		

Table 2. Components of Π in the proof of Theorem 1, simulating the terminating phase of the system. The region C_{S_4} is omitted since it always contains the string S_4 .

is used in each region to keep a copy of its contents even in the case when it is requested by some other region. This way, Π' simulates the communication behavior of Π .

By Theorems 1 and 2, we obtain the immediate corollary.

Corollary 3 $iRPC_{10} = pRPC_{30}CF = RE$.

3 Closing Remarks

We proved that in the case of string rewriting P systems communication according to dynamically emerging requests leads to computational completeness both for standard P systems and tissue-like P systems, even in the case of systems with bounded size parameters. There have remained several open problems for further study. For example, it is not known whether the obtained size bounds are sharp or not, and whether or not the sharp bounds are different for MPC systems and RPC systems.
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On the Dynamics of PB Systems with Volatile Membranes

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Summary. We investigate decision problems like reachability and boundedness for extensions of PB systems with volatile membranes. Specifically, we prove that reachability and boundedness are decidable for PB systems extended with rules for membrane dissolution. For PB systems extended with membrane creation, reachability is still decidable whereas boundedness becomes undecidable. Furthermore, we show that both problems are undecidable for PB systems extended with both dissolution and creation rules. Finally, we prove that reachability and boundedness become decidable for PB systems with dissolution rules and in which only one instance of each type of membrane can be created during a computation. Our work extends previous results obtained for PB systems by Dal Zilio and Formenti in a paper appeared in the proceedings of WMC 2003 [5].

1 Introduction

The PB systems of Bernardini and Manca [2] are a variant of P-systems [16] in which rules can operate on the boundary of a membrane. A *boundary rule* can be used here to move multisets of objects across a membrane. In biological modeling, PB are very useful for expressing complex interactions among biological membranes [8]. For this reason, it seems important to develop methods for qualitative and quantitative analysis of models specified in this formalism. In this paper we focus our attention on theoretical issues for the qualitative analysis of PB systems. Some preliminary results on decision problems for PB systems have been obtained in [5]. Specifically, in [5] Dal Zilio and Formenti proved that the reachability problem is decidable for PB systems with symbol objects. The *reachability problem* consists in checking if a given system can evolve into a fixed a priori configuration. The decidability proof in [5] is based on an encoding of PB systems into Petri nets [17], an infinite-state model of concurrent systems for which the reachability problem is decidable [14, 11]. A Petri net is a collection of places that contain tokens,

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and of transitions that define how tokens move from one place to another. The current configuration of a net is called marking. A marking specifies the current number of tokens in each place. A PB system can be encoded as a Petri net in which membranes are modelled as places, symbol objects as tokens, configurations as markings, and internal/boundary rules as transitions. The execution of a rule is simulated then by the firing of the corresponding Petri net transition. The Petri net encoding shows that the reachability problem is decidable in PB systems. The same reduction can be used to decide other properties like *boundedness* [6]. In [5] the authors observe that the aforementioned encoding can be extended to more sophisticated Petri net models so as to deal with dynamically changing membrane structures. As an example, Petri net transitions extended with transfer arcs naturally model the dissolution of a membrane. Indeed, a transfer arc can be used to atomically transfer all tokens from one place to another. This operation can be applied to move the content of a dissolved membrane to its father. Unfortunately, as pointed out in [5], this connection cannot be exploited in order to extend the decidability results obtained for PB systems. Indeed, problems like reachability and boundedness become undecidable in presence of transfer or reset arcs [6]. The decidability of reachability and boundedness for PB systems with volatile or moving membrane seems to be still an open problem. In this paper we focus our attention on decision problems for extensions of PB systems with dissolution and creation rules. More specifically, our technical results are as follows.

We first show that reachability is decidable in PB systems with dissolution rules (PBD systems). Dissolution rules are a peculiar feature of P-systems. Thus, PBD systems represent a natural extension of PB system. Our decidability proof is still based on a reduction to a Petri net reachability problem. Our construction extends the Petri net encoding of [5] in order to weakly simulate the original PBD system. More precisely, from a PBD reachability problem we compute a Petri net that may contain executions that do not correspond to real computations of the corresponding PBD system. Spurious computations can however be eliminated by enforcing special conditions (e.g. requiring a special set of places to be empty) on the initial and target markings used to encode a PBD reachability problem. It is important to notice that our reduction does not require the additional power provided by Petri nets with transfer arcs.

As a second result, we show that reachability is decidable in PB systems extended with creation rules (PBC systems). We consider here creation rules inspired to those proposed by Martín-Vide, Pãun, and Rodriguez-Paton in the context of P-systems [13]. Our proof exploits structural properties of PBC systems that allow us to reduce the reachability of a target configuration c, to a reachability problem in a Petri net extracted from both the original PBC system and the configuration c. As for PBD systems, this decidability result is a conservative extension of the result for PB systems obtained in [5].

We consider then a model with both dissolution and creation rules (PBDC systems). For this model, we first give a general negative result for the decidability of reachability, and then study a non-trivial subclass in which reachability becomes decidable. Specifically, we first show that it is sufficient to consider three membranes with the same name to encode a reachability problem for a two counter machine [15] as reachability of a PBDC system. We define then a restricted semantics for PBDC systems in which at most only one copy of each type of membrane can be created during a computation. This semantics is inspired to a view of membrane names/types as bounded resources. Under this semantics, we prove the decidability of PBDC reachability via a reduction to Petri net reachability. In the encoding we use special places to identify the membrane structure of the current configuration. The encoding is exponentially more complex than the encoding used for PBD and PBC systems, since it requires the construction of a Petri net where the number of places is equal to number of tree structures that can be built upon a finite and fixed a priori set of membrane names.

As a last analysis, we study the boundedness problem for the aforementioned extensions of PB systems. Specifically, we first show that boundedness is decidable for PBDC systems with restricted semantics. The proof exploits the theory of well-quasi ordering [7]. As a consequence, we obtain the decidability of boundedness for PBD systems. Finally, we prove that boundedness is undecidable in PBC systems. This result is obtained by encoding counter machines as PBC systems. The encoding exploits the possibility of creating several instances of the same membrane to simulate a counter (i.e. the same encoding cannot be applied in the restricted semantics of PBDC systems).

To our current knowledge, these are the first decidability/undecidability results obtained for reachability and boundedness in extensions of PB systems with dissolution and creation rules. Decision problems for qualitative analysis of subclasses of P-systems have been studied, e.g., in [12, 9]. It is important to notice that the decidability of reachability in PBC systems is not in contradiction with the undecidability of boundedness in the same model. Indeed, several other examples of universal models for which the reachability problem is decidable has recently been discovered in the field of process algebra, see e.g. [3, 4].

Plan of the Paper

In Section 2 we recall the main definitions of PB systems, Petri nets, and counter machines. In Section 3, 4, and 5 we study the reachability problem for extensions of PB systems resp. with dissolution, creation, and both dissolution and creation rules. In Section 6 we study the boundedness problem for the aforementioned extensions of PB systems. Finally, in Section 7 we address some conclusion and future work.

2 Preliminaries

In this section we recall the main definitions for PB systems with symbol objects taken from [2, 5], Petri nets [17], and counter machines [15]. We first need some

preliminary notions. Let \mathbb{N} be the set of positive integers. Consider a finite alphabet Γ of symbols. A multiset over Γ is a mapping $u : \Gamma \rightsquigarrow \mathbb{N}$. For any $a \in \Gamma$, the value u(a) denotes the multiplicity of a in u (the number of occurrences of symbol a in u). We often use a multiset as a string $a_1 \cdot \ldots \cdot a_n$ of symbols, i.e., $a_i \in \Gamma$. Furthermore, we use ϵ to denote the empty multiset, i.e., such that $\epsilon(a) = 0$ for any $a \in \Gamma$. As an example, for $\Gamma = \{a, b, c, d\}$, $a \cdot b \cdot c \cdot c$ represents the multiset u such that u(a) = u(b) = 1, u(c) = 2, u(d) = 0. We use Γ^{\otimes} to denote the set of all possible multisets over the alphabet Γ . Given two multisets u, v over Γ , we write $u \doteq v$ if u(a) = v(a) for all $a \in \Gamma$, and $u \preceq v$ if $u(a) \leq v(a)$ for all $a \in \Gamma$. Furthermore, we use \oplus and \ominus to denote multiset union and difference, respectively. Specifically, for any $a \in \Gamma$ we have that $(u \oplus v)(a) = u(a) + v(a)$, and $(u \oplus v)(a) = u(a) - v(a)$. We are ready now to formally define a PB system.

2.1 P-systems with Boundary Rules

A PB system [2] with symbol object is a tuple $\Pi = (\Gamma, M, R, \mu_0)$, where

- Γ is a finite alphabet of symbols;
- M is a finite tree representing the *membrane structure* with membrane names taken from a set N,
- *R* is a finite set of rules,
- μ_0 is the *initial configuration*, i.e., a mapping from membranes (nodes in M) to multisets of objects from Γ .

Rules can be of the following two forms:³

(1) Internal : $[_i \ u \to [_i \ v$ (2) Boundary : $u \ [_i \ v \to u' \ [_i \ v'$

where $i \in N$, and $u, u', v, v' \in \Gamma^{\otimes}$ and we assume that at least one between u and u' is not empty.

A configuration μ of a PB system Π is a distribution of objects in Γ in the membranes in M, i.e., a mapping from M to Γ^{\otimes} . A rule of the form (1) is enabled at μ , if i is a membrane in M and $u \leq \mu(i)$. Its application leads to a new configurations ν' such that $\nu'(i) = (\nu(i) \ominus u) \oplus v$ and $\nu'(j) = \nu(j)$ for any $j \in N$ s.t. $j \neq i$.

Suppose now that membrane j contains as immediate successor in M membrane i. A rule of the form (2) is enabled at μ , if $u \leq \mu(j)$ and $v \leq \mu(i)$. Its application leads to a new configurations ν' such that $\nu'(j) = (\nu(j) \ominus u) \oplus u'$ and $\nu'(i) = (\nu(i) \ominus v) \oplus v'$ and $\nu'(k) = \nu(k)$ for any $k \in N$ s.t. $k \neq i, j$. We say that there is a transition $\mu \Rightarrow \mu'$ if μ' can be obtained from μ by applying a rule in R. A computation with initial configuration μ_0 is a sequence of transitions $\mu_0 \Rightarrow \mu_1 \Rightarrow \ldots \Rightarrow \mu$

³ We consider here a slight generalization of the model in [5] in which we allow any kind of transformation between two membranes.

Definition 1 (Reachability). Given a PB system Π with initial configuration μ_0 and a configuration μ , the reachability problem consists in checking if μ is reachable from μ_0 .

Definition 2 (Boundedness). Given a PB system Π with initial configuration μ_0 , the boundedness problem consists in deciding if the set of configurations reachable from μ_0 is finite.

Reachability and boundedness are decidable for PB systems with symbol objects. The proof is based on an encoding PB systems into Petri nets defined in [5].

2.2 Petri nets

A Petri net [17, 18] is a pair (P, T, m_0) where P is a finite set of places, T is a finite set of transitions $(T \subseteq P^{\otimes} \times P^{\otimes})$, and m_0 is the initial marking. A transition tis defined by the pre-set ${}^{\bullet}t$ and by the post-set t^{\bullet} , two multisets of places in P (we consider here multisets of places instead of adding weights to arcs). A marking is just a multiset over P. Given a marking m and a place p, we say that the place pcontains m(p) tokens. A transition t is enabled at the marking m if ${}^{\bullet}t \preceq m$. If it is the case, t can fire and produces a marking m' (usually written $m \stackrel{t}{\longrightarrow} m'$) defined as $(m \ominus^{\bullet} t) \oplus t^{\bullet}$. A firing sequence is a sequence of markings $m_0m_1\ldots$ such that m_i is obtained from m_{i-1} by firing a transition in T at m_i . Finally, we say that m' is reachable from m_0 if there exists a firing sequence from m_0 passing through marking m'. The Petri net reachability problem has been proved to be decidable in [14, 11].

2.3 Counter Machines

A counter machine [15] consists of a finite set of locations/control states ℓ_1, \ldots, ℓ_k , a finite set of counters c_1, \ldots, c_m , and a finite set of instructions. The instruction set consists of the increment, decrement, and zero-test, and the nonzero-test operations on each counter. Each operation has three parameters: the current location, the successor location, and the counter on which it operates. When executed in location ℓ , the increment operation on c_i adds one unit to the current value of c_i and then moves to the successor location ℓ' . When executed in location ℓ , the decrement operation on c_i removes one unit from the current value of c_i and then moves to the successor location ℓ' . When executed in location ℓ , the zero-test on c_i moves to the successor location ℓ' only if c_i has value zero. When executed in location ℓ , the nonzero-test on c_i moves to the successor location ℓ' only if c_i has a value strictly greater than zero. For a fixed initial location ℓ_0 , the initial configuration has location ℓ_0 and all counters set to zero. A computation is a sequence of configurations obtained by applying instructions associated to locations. The *if-then-else* instruction ℓ : *if* $c_i = 0$ *goto* ℓ_1 *else goto* ℓ_2 typically found in the definition of counter machines can be simulated here by two rules associated to the same location ℓ : a zero-test ℓ : $c_i = 0$ goto ℓ_1 and a nonzero-test. ℓ : $c_i > 0$ goto ℓ_2 . It

is well-known that the model of *two counter machines* can simulate a Turing machine, i.e., properties like termination, reachability of a location, and boundedness are all undecidable in this model [15].

3 PB Systems with Dissolution Rules

A PB system with dissolution rules (PBD) provides, in addition to internal and boundary rules, a third kind of rules of the following form:

(3) Dissolution :
$$[_i \ u \to [_i \ v \cdot \delta]$$

where δ is a symbol not in Γ . The intuitive meaning of this rule is that after applying the rule $[i \ u \to [i \ v$ the membrane i is dissolved and its content (including its sub-membranes) is moved to the membrane j that contains i as immediate successor in the current membrane structure. To make the semantics formal, we make the membrane structure part of the current configuration, M_0 being the initial tree. Thus, a configuration is now a pair $c = (M, \mu)$, where M is a tree, and μ is a mapping from nodes of M to Γ^{\otimes} . Rules of type (1) and (2) operate on a configuration $c = (M, \mu)$ without changing the tree structure M and changing μ as specified in the semantics of PB systems. A dissolution rule like (3) operates on a configuration $c = (M, \mu)$ as follows. For simplicity, we assume that membrane i is not the root of M. Suppose now that i is an immediate successor of j in M. The rule is enabled if $u \leq \mu(i)$. Its application leads to a new configurations $c' = (M', \nu')$ such that

- *M'* is the tree obtained by removing node *i* and by letting all successor nodes of *i* become successors of *j*;
- ν' is the mapping defined as $\nu'(j) = \nu(j) \oplus (\nu(i) \oplus u) \oplus v$ and $\nu'(k) = \nu(k)$ for any $k \in M$ s.t. $k \neq i, j$.

Notice that rules of type (1-3) are enabled at $c = (M, \mu)$ only if the membrane *i* is in current tree *M*. The definition of sequences of transitions and of reachability problems can naturally be extended to the new type of rules.

3.1 Decidability of Reachability in PBD Systems

In this section we prove that the reachability problem is decidable in PB systems with dissolution rules. We assume here that names of membranes are all different. However, the construction we present can be extended to the general case. The starting point of our construction is the reduction of reachability for PB systems to reachability in Petri nets given in [5]. Let $\Pi = (\Gamma, M, R, \mu_0)$ be a PB system. For each membrane *i* in *M* and each symbol $a \in \Gamma$, the Petri net \mathcal{N} associated to Π makes use of place a^i to keep track of the number of occurrences (multiplicity) of objects of type *a* in *i*. Transitions associated to internal rules redistribute tokens in the set of places associated to the corresponding membrane. As an example, a rule like $[i \ a \cdot b \rightarrow [i \ c \ is encoded by a Petri net transition that removes one token$ $from place <math>a^i$ and one token from place b^i and adds one token to place c^i . Boundary rules are modelled by Petri net transitions that work on places associated to pairs of membranes. As an example, if membrane j contains i, a rule like $a \ [i \ b \rightarrow b \ [i \ a$ is encoded by a Petri net transition that removes one token from place a^j and one token from place b^i and adds one token to place b^j and one token to place a^i . For a membrane structure M, a configuration $\mu : M \rightsquigarrow \Gamma^{\otimes}$ is represented by a marking m_{μ} such that for every node i in M, a^i has k tokens in m iff a has koccurrences in $\mu(i)$. Reachability of a configuration μ is reduced the to reachability of the marking m_{μ} starting from m_{μ_0} in \mathcal{N} .

How to Cope with Dissolution Rules

The Petri net encoding of [5] exploits the property that the membrane structure of a PB system is never changed by the application of a rule. This property does not hold anymore for dissolution rules, since they removes nodes from the current membrane structure. Thus, the size of the membrane structure may decrease in a sequence of transitions. Our decidability proof is still based on a reduction to a Petri net reachability problem. Our reduction exploits the property that the number of applications of dissolution rules is bounded a priori by the size of the initial membrane structure M_0 .

Specifically, suppose that in M_0 there is an absolute path i_0, i_1, \ldots, i_k, i where i_0 is the root of M_0 . In the construction of the Petri net associated to the membrane i we must take into consideration the possibility that each one of the membranes i_1, \ldots, i_k can dissolve during the execution. This means that boundary rules associated to the membrane i should be redirected to the membrane i_j in the path i_0, i_1, \ldots, i_k such that all membranes i_{j+1}, \ldots, j_k are no more present in the current membrane structure. To achieve this, each membrane i comes with an associated flag $present_i/dissolved_i$. Transitions that encode boundary rules on membrane i are conditioned then by the present/dissolved flags of the membranes i_0, i_1, \ldots, i_k in the path leading to i. In other words we need to implement a sort of switch that redirects boundary rules to membrane i_j whenever membranes i_{j+1}, \ldots, j_k are all dissolved.

Another problem to solve is related to the transfer of the contents of a membrane to its immediate ancestor in the membrane structure. To simulate this process, our Petri net operates in two different modes. In the *normal* mode the Petri net simulates boundary and internal rules. Suppose now that j is the membrane that contains i in the current tree structure, and that the dissolution rule $[_iu \rightarrow [_iv \cdot \delta$ is enabled. We first execute the internal rule $[_iu \rightarrow [_iv.$ The Petri net then switches to the *dissolving_i* mode. In this mode all normal operations are blocked. This is achieved by conditioning all transitions associated to normal operations with the *normal* flag. After this step, we activate a set of transitions that move tokens (one-by-one) from i to j. Since in a Petri net it is not possible to test if a place is empty, instead of testing if all objects in i have been moved to j, we add a rule that non-deterministically stops the transfer process. As a last step, we marked membrane i as dissolved and reactivate the normal operating mode. The last step automatically disable internal/boundary/dissolve rules operating on membrane i (they are conditioned by the *present*_i flag) and redirects boundary rules of any membrane i' contained in i (in the current tree structure) to membrane j.

The non-deterministic termination of the transfer process may lead to incorrect simulations of the original PBD system. Indeed, we could restart in normal mode with a configuration in which a membrane is marked dissolved and some of its objects have not been transferred to its father. This problem can be solved by the following key observation. We first recall that we are interested in reachability of a configuration $c = (M, \mu)$. Thus, by looking at the structure of M_0 and M we can extract the set of nodes D that must be dissolved in a sequence of transitions leading from c_0 to c. Thus, we can tell good simulations from bad ones by imposing the constraint that, in the marking m_{μ} modelling the target configuration μ , every place associated to a membrane in D is empty. As a final remark, notice that all information needed in the encoding (nodes and paths in M_0) can statically be extracted from the description of the PBD system and from the initial configuration c_0 .

Formal Definition of the Petri Net Encoding

Assume a PBD system $\Pi = (\Gamma, M_0, R, \mu_0)$, where $\Gamma = \{a_1, \ldots, a_m\}$, and M_0 has the membranes with names in $N = \{n_0, n_1, \ldots, n_k\}$, $n_0 \in N$ being the root node. Given a membrane *i*, let path(i) be the sequence of nodes in the (unique) path from n_0 to *i* in M_0 .

We define the Petri net \mathcal{N} encoding Π in several steps. First of all we assume that \mathcal{N} has at least the places *normal*, $dissolving_1, \ldots, dissolving_k$ that we use to determine the simulation mode as described in the previous paragraphs. We assume here that *normal* contains one token iff $dissolving_i$ is empty for all $i : 1, \ldots, k$, and $dissolving_i$ contains one token iff *normal* as well $dissolving_j$ are empty for any $j \neq i$. Furthermore, for each membrane i, the Petri net \mathcal{N} has a place $present_i$ and a place $dissolved_i$ and, for any $a \in \Gamma$, a place a^i .

Notation: In the rest of the paper given a multiset of objects u and a membrane i we use $\pi_i(u)$ to denote the multiset of places in which, for each $a \in \Gamma$, a^i has the same number of occurrences as those of a in u.

Internal Rules

An internal rule $r = [_i u \rightarrow]_i v$ is encoded by a transition t_r that satisfies the following conditions. The pre-set of t_r contains place normal (normal mode), present_i (membrane *i* is still present), and the multiset of places $\pi_i(u)$. The post-set of t_r contains normal, present_i and the multiset of places $\pi_i(v)$. Thus, the only difference with the encoding of PB system is the condition on the normal and present_i flags (in normal mode internal rules are enabled only when the membrane is not dissolved).

Boundary Rules

Let $path(i) = (n_0, n_1, \ldots, n_q, i)$ with $q \ge 0$. A boundary rule $r = u[_iv \to u']_iv'$ is encoded by a set $B_r = \{b_r^{n_0}, \ldots, b_r^{n_q}\}$ of transitions. The pre-set of transition $b_r^{n_j}$ contains places normal and present_i together with the set of places $D^{n_j} = \{present_{n_j}, dissolved_{n_{j+1}}, \ldots, dissolved_{n_q}\}$, and the multisets $\pi_{n_j}(u)$ and $\pi_i(v)$. The post-set contains places normal and present_i together with the set of places D^{n_j} defined for the pre-set and the multisets $\pi_{n_j}(u')$ and $\pi_i(v')$. The pre-condition D^{n_j} allows us to select the membrane that is the immediate ancestor of i in the current configuration, i.e., a membrane $n_j \in path(i)$ that is not dissolved and such that all the intermediate membranes between n_j and i in path(i) are dissolved. Notice that, by the assumptions we made on the normal/dissolving and present/dissolved flags, in normal mode one and only one rule in B_r can be enabled at a given configuration (represented by a marking).

Dissolution Rules

Let $path(i) = (n_0, n_1, \ldots, n_q, i)$ with $q \ge 0$. Consider a dissolution rule $r = [iu \rightarrow [iv \cdot \delta]$. We first model the internal rule by the transition s_r . The pre-set of s_r contains the places *normal* and *present*_i and the multiset $\pi_i(u)$. The post-set contains the place dissolving_i and the multiset $\pi_i(v)$.

We then model the transfer of the contents of membrane *i* to its current immediate ancestor via a set of transitions $S_r^a = \{s_a^{n_0}, \ldots, s_a^{n_q}\}$ for each $a \in \Gamma$. The pre-set of transition $s_a^{n_j}$ contains places dissolving_i (*i* is dissolving) and a^i (the source of a token to be transferred) together with the set of places D^{n_j} defined in the case of boundary rules. The post-set contains places dissolving_i and a^j (the destination of a transferred token), and the set D^{n_j} .

Finally, we add a transition d_r^i to stop the transfer of tokens and to switch the operating mode back to normal. The pre-set of d_r^i contains the place dissolving_i and its post-set contains the places normal and dissolved_i. Notice that the simulation phase of a dissolution rule for membrane *i* can be activated only if present_i is not empty. This implies that once the dissolving_i flag is reset (i.e. the mode goes back to normal) it cannot be set in successive executions (a membrane can dissolve at most once).

Places, Transitions and Configurations

The Petri net \mathcal{N} is built by taking the union of the places and transitions used in the encoding described before. Let M be a membrane structure with a subset of the nodes in M_0 (initial structure of Π). A configuration $c = (M, \mu)$ is encoded by a marking m_c in which there is one token in *normal*, one token in *present*_i for each membrane *i* in M, and one token in *dissolved*_j for each *j* not in M. Furthermore, for each membrane i in c and $a \in \Gamma$, place a^i has as many tokens as the number occurrences of a in $\mu(i)$. All the remaining places in \mathcal{N} (dissolving_i for $i : 0, \ldots, k$ and present_j for all membranes j not in M) are empty.

Reachability Problem

By construction of \mathcal{N} , it is immediate to see that if there is a sequence of transitions from $c_0 = (M_0, \mu_0)$ to $c = (M, \mu)$ passing through the configurations c_1, \ldots, c_v then there is a firing sequence from m_{c_0} to m_c passing through the markings m_{c_1}, \ldots, m_{c_v} . Such a firing sequence is obtained by completing all transfers of objects required by the simulation of each dissolution rule (i.e. after the simulation of the dissolution of membrane i, the normal mode is reactivate only when the places associated to the objects contained in i are all empty). Vice versa, suppose there exists a firing sequence from m_{c_0} to m_c . We first notice that only the markings in which the place *normal* is not empty correspond to configurations of the original PBD system. Furthermore, suppose that during the simulation of the dissolution of membrane i, the transfer of objects is stopped when some of the places associated to objects in i are not empty. Let m be the resulting marking. Now we notice that the first step of the simulation of dissolution is to set the $present_i$ flag to false. This implies that in the marking m place $present_i$ is empty, while there exists $a \in \Gamma$ such that a^i is not empty (some token has not been transferred). It is easy to check that if m has these two properties, for any marking m' derived from m by applying transitions of \mathcal{N} , the content of the place a^i in m' is the same as in m. Indeed, transitions that simulate internal, boundary and dissolution rules operating directly on i are no more enabled (the condition $present_i$ fails). Furthermore, a dissolution rule on a membrane j nested into i in M_0 cannot transfer tokens to i since $dissolved_i$ is checked when searching for the father of j in the current tree structure. In other words, if the simulation of a dissolution rule is not correctly executed, then there exists at least one non-empty place a^i for a dissolved membrane i. By definition, however, m_c is the marking in which all places associate to dissolved membranes are empty. Thus, if m_c is reachable from m_{c_0} then the corresponding firing sequence corresponds to a real computation in \mathcal{N} . Thus, we have that the reachability of a configuration $c = (M, \mu)$ in Π can be encoded as the reachability of the marking m_c in \mathcal{N} from m_{c_0} . From the decidability of reachability in Petri nets, we obtain the following theorem.

Theorem 1. Reachability is decidable in PBD systems.

This result extends (and is consistent with) the Petri net encoding and the decidability result for reachability in PB systems of [5].

4 PB System with Creation

In this section we consider an extension of PB systems inspired to the membrane creation operation studied in [13]. Let N be a possibly infinite list of membrane

names. A PB system with creation rules (PBC) provides, in addition to internal and boundary rules, a third kind of rules of the following form:

(4) Creation :
$$a \rightarrow [i \ v]_i$$

where $a \in \Gamma$, $v \in \Gamma^{\otimes}$, and $i \in \mathbb{N}$. The intuitive meaning of this rule is that after applying the rule $a \to [{}_iv]_i$ inside a membrane j, object a is replaced by the new membrane i containing the multiset of objects v. To make the semantics formal, we assume that membrane structures are trees whose nodes are labelled with names in N. Furthermore, we make both the set of used names and the membrane structure part of the current configuration, $N_0 \subseteq \mathbb{N}$ being the initial set of used names, and M_0 being the initial tree defined over N_0 . Thus, a configuration is now a triple $c = (N, M, \mu)$ where N is a set of names, M is a tree with nodes labelled in N, and μ is a mapping $M \rightsquigarrow \Gamma^{\otimes}$. Rules of type (1) and (2) operate on a configuration $c = (N, M, \mu)$ without changing N and M. A creation rule like (4) operates on a configuration $c = (N, M, \mu)$ as follows. Suppose that n is a node in M. The rule is enabled if $a \in \mu(n)$. Its application leads to a new configurations $c' = (N', M', \nu')$ such that

- $N' = N \cup \{i\};$
- M' is the tree obtained by adding a new node m labelled by i as a successor of node n;
- ν' is the mapping defined as $\nu'(n) = \nu(n) \ominus a$, $\nu'(m) = v$, and and $\nu'(p) = \nu(p)$ for $p \neq m, n, p \in N$.

Notice that rules of type (4) can be applied in any membrane. Indeed, the only precondition for the application of rule 4 is the existence of object a in a membrane. Furthermore, such an application may create different nodes with the same membrane name.

The reachability problem can naturally be reformulated for the extended semantics of rules. Specifically, it consists in checking whether a given target configuration c is reachable from the initial configuration c_0 .

In presence of creation rules the membrane structure can grow in an arbitrary manner both in width and depth. Notice that in our model we distinguish nodes from membrane names. Thus, different nodes may have the same name. As a simple example, consider the rule $a \rightarrow [_1a]_1$ in a system with a single membrane $[_0a]_0$. The evolution of this PBC system may lead to membrane structures of arbitrary nesting level, e.g.,

 $[0[1a]_1]_0 \quad [0[1[1a]_1]_1]_0 \quad [0[1[1a]_1]_1]_1]_0 \quad \dots$

Now consider the rules $[_0a \rightarrow [_0a \cdot a \text{ and } a \rightarrow [_1b]_1$. Then the membrane $[_0a]_0$ can generate membrane structures of arbitrary width, e.g.,

$$\begin{bmatrix} 0 \\ 1 \end{bmatrix} \begin{bmatrix} 1 \\ 0 \end{bmatrix} \begin{bmatrix} 1 \\ 0 \end{bmatrix} \begin{bmatrix} 1 \\ 0 \end{bmatrix} \begin{bmatrix} 1 \\ 1 \end{bmatrix} \begin{bmatrix} 1 \\ 0 \end{bmatrix} \begin{bmatrix} 1$$

Despite of these powerful features of PBC systems, the reachability problems can still be decided by resorting to an encoding into Petri net reachability as explained in the next section.

4.1 Decidability of Reachability in PBC Systems

Differently from the encoding defined for PBD systems in the previous section, the encoding needed here is function of both the initial and the target configuration. Indeed, since PBC rules can only add new nodes, to decide if a configuration $c = (N, M, \mu)$ is reachable from c_0 we can restrict our attention to membrane structures of size comprised between the size of M_0 and the size of M and with (possibly repeated) labels in N.

Actually, we can make some simplification that allows us to build a Petri net by considering only the target configuration M. Indeed, as shown in [13], with creation rules we can safely consider initial configurations with only the root membrane (we can always add a finite number of creation rules to generate any initial configuration in a preliminary phase of the computation).

So let Π be a PBC system with initial configuration $c_0 = (N_0, M_0, \mu_0)$ where M_0 is a single node labelled n_0 . Consider now a target configuration (N, M, μ) where $N = \{n_0, \ldots, n_k\}$ and M has m nodes with $k \leq m$ and the root node of M is labelled n_0 .

Starting from Π and c we build the Petri net \mathcal{N} described next. For each node n in M, the Petri net \mathcal{N} has places $used_n$ and $notused_n$ (used as one flip-flop), and a^n for each $a \in \Gamma$ (to model the content of membrane n). We assume that $used_n$ is not empty iff the membrane has been created and it is in use, and $notused_n$ is not empty iff the membrane has still to be created. PBC rules are modelled as follows.

Internal Rules

For each node n in M with label i, an internal rule $r = [{}_{i}u \rightarrow [{}_{i}v$ is encoded by a transition t_{r}^{n} that satisfies the following conditions: The pre-set contains place $used_{n}$ together with multiset $\pi_{n}(u)$; The post-set contains $used_{n}$ together with multiset $\pi_{n}(u)$.

The differences with the encoding of PB/PBD systems is the condition on the $used_n$ flag and the fact that we work on nodes of membrane structures and not directly on membrane names (as said before two different nodes may have the same name). The pre-condition on $used_n$ is needed in order to enable rules operating on node n only after the corresponding creation rule has been fired.

Boundary Rules

For each node m in M that has an immediate successor n with label i, a boundary rule $r = u_{i}v \to u'_{i}v'$ is encoded by a transition $b_{r}^{m,n}$ that satisfies the following conditions. The pre-set contains places $used_n$ and $used_m$ together with the multisets $\pi_m(u)$ and $\pi_n(v)$. The post-set contains places $used_n$ and $used_m$ together with the multisets $\pi_m(u')$ and $\pi_n(v')$. Notice that, differently from the encoding used in PBD, in PBC we do not have to consider paths in the membrane structure M.

Creation Rules

For each node m in M that has an immediate successor n with label i, a creation rule $r = a \rightarrow [iv]_i$ is encoded by a transition $c_r^{m,n}$ that satisfies the following conditions. The pre-set contains places $used_m$, a^m and $notused_n$. The post-set contains places $used_m$ and $used_n$, together with the multiset $\pi_n(v)$

Places, Transitions and Configurations

The Petri net \mathcal{N} is built by taking the union of the places associated to each membrane and the union of the set of transitions used to encode internal, boundary and creation rules described before.

A configuration $c' = (N', M', \mu')$ is encoded by a marking $m_{c'}$ in which for each node n in M' there is one token in $used_n$, and for each $a \in \Gamma$, a^n has as many tokens as the number occurrences of a in $\mu(n)$. Furthermore, for each node n' in M that do not occur in M', we put a token in $notused_{n'}$. All the remaining places are empty.

Reachability Problem

By construction of \mathcal{N} , it is immediate to see that there is a sequence of transitions from $c_0 = (N_0, M_0, \mu_0)$ to $c = (N, M, \mu)$ passing through the configurations c_1, \ldots, c_v if and only if there is a firing sequence from m_{c_0} to m_c passing through the markings m_{c_1}, \ldots, m_{c_v} . The creation of a new node *n* corresponds to the activation of the part of the Petri net \mathcal{N} that models node *n*. Since nodes are created in "cascade", a node *m* is created only after all ancestors have been created. This property is ensured by the condition on the *used* flag inserted in the transitions modelling creation rules.

Following from the decidability of reachability in Petri nets, we obtain the following theorem.

Theorem 2. Reachability is decidable in PB systems with creation rules.

This result extends (and is consistent with) the Petri net encoding and the decidability result for reachability in PB systems of [5].

5 PB Systems with Dissolution and Creation

In this section we consider an extension of PB systems with both dissolution and creation rules (PBDC systems). The semantics is obtained in a natural way by adapting the semantics of dissolution rules to membrane structures with labelled nodes. More precisely, a dissolution rule applied to a configuration (N, M, μ) modifies M and μ as specified in Section 3 while it does not modify N, i.e., the set N of used names can only grow monotonically. Notice that the reachability problem for PBDC systems allows to determine if, for a PBDC system Π , it is possible

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from the initial configuration to build a membrane structure M with a mapping μ that associates multisets of objects to nodes, whatever the set of names N is. Indeed, the set of possible names for membranes that appear in executions of Π is determined by the creation rules of the PBDC system (and its initial configuration). Hence, the number of possibilities for N is finite and the problem can be reduced to a finite number of reachability problems.

In presence of both creation and dissolution the membrane structure can change in an arbitrary manner. As a simple example consider the rules $a \to [1a]_1$, and $[1a \to [1a \cdot \delta \text{ in a system with a single membrane } [0a]_0$. The evolution of this PB system may lead to membrane structures that grow and shrink in an arbitrary way as in the sequence:

$$[0[1a]_1]_0 \quad [0[1[1a]_1]_1]_0 \quad [0[1a]_1]_0 \quad \dots$$

This feature gives additional power to PB systems. Indeed, we can reduce the reachability problem for two counter machines (known to be undecidable) to reachability in a PBDC system. For this reduction it is enough to consider dissolution and creation rules working on three membranes with the same name. Specifically, consider a system with initial configuration

$$c_0 = [_0 \ s_0 \ [_0 \ c_1 \]_0 \ [_0 \ c_2 \]_0 \]_0$$

here s_0 represents the initial control state of a two counter machine. Membrane $[_0c_i]_0$ is used to represent counter c_i with value zero for i : 1, 2. Counter c_i with value k is represented by the membrane $[_0c_i \cdot u]_0$ where u is the multiset with k occurrences of a special object a.

The increment of counter c_i in control state s and update to control state s' is encoded by the boundary rule

$$s \ [0 \ c_i \rightarrow s' \ [0 \ c_i \cdot a]$$

for i : 1, 2.

The decrement of counter c_i in control state s and update to control state s' is encoded by the boundary rule

$$s \ [0 \ c_i \cdot a \to s' \ [0 \ c_i$$

for i : 1, 2.

The zero test on counter c_i in control state s and update to control state s' is simulated by three rules. We first move to an auxiliary state aux_s and dissolve the membrane with label i.

$$s \ [0 \ c_i \rightarrow aux_s \ [0 \ \epsilon \cdot \delta]$$

where ϵ is the empty multiset. We then create a new empty instance of the same membrane containing the objects c_i and out'_s via the rule

$$aux_s \rightarrow [0 \ c_i \cdot out_{s'}]_0$$

Finally, we use a boundary rule to move to the next state s'

$$\begin{bmatrix} 0 & c_i \cdot out_{s'} \to s' \end{bmatrix} \begin{bmatrix} 0 & c_i \end{bmatrix}$$

We assume here that no other rule uses aux_s and $out_{s'}$. The effect of the execution of these three rules is that of moving the contents of the counter on which the zerotest is executed to the top level membrane 0. Indeed, the membrane containing c_i is first dissolved and then re-created. If the zero-test is executed when a counter is not zero, some object will remain inside membrane 0 in all successive configurations. This feature can be used to distinguish good simulations from bad ones. Specifically, let us consider the reachability problem for a two counter machine in which the initial and the target configurations both coincide with the configuration with a given control state s and both counters set to zero. This problem can be expressed as the reachability of the configuration c_0 from c_0 . Thus, the following property holds.

Theorem 3. Reachability is undecidable in PBDC systems in which configurations have at most three different membranes with the same name.

5.1 PBDC Systems with Restricted Semantics

As a final analysis, we consider a restricted semantics for PBDC systems in which newly created membranes must be assigned fresh and unused names. In other words we assume that creation rules can be applied at most once for each *type* of membrane. Another possible view is that membrane names are themselves resources that can be used at most once.

Formally, assume a configuration $c = (N, M, \mu)$. Suppose that n is a node in M. In the restricted semantics, the creation rule (4) is enabled if $a \in \mu(n)$ and $i \notin N$, i.e., the name i is *fresh*. Its application leads to a new configurations $c' = (N', M', \nu')$ such that $N' = N \cup \{i\}$, M' is the tree obtained by adding a new node m labelled i as a successor of node n, and ν' is the mapping defined as $\nu'(n) = \nu(n) \ominus a, \nu'(m) = v$, and $\nu'(p) = \nu(p)$ for $p \neq m, n, p \in N$).

Since with creation rules in the style of [13] the set of rules operating on membranes is fixed and known a priori, we can assume that the number of distinct names is finite (it corresponds to the set of names occurring in internal, boundary, dissolution and creation rules and in the initial configuration). This restriction yields the following key observation.

Observation 1.

If the set of possible membrane names N is finite and every name in N can be used only once, then starting from a configuration with a single membrane, the number of distinct membrane structures that we can generate is finite. Every such membrane structure has at most |N| nodes.

This property does not imply that the number of configurations is finite. Indeed, there are no restrictions on creation and deletion of *objects* inside membranes. As

an example, the PBDC system with the internal rule $[_0a \rightarrow [_0a \cdot a \text{ and the initial} membrane <math>[_0a]_0$ generates an infinite set of configurations (membrane 0 with any number of repetitions of object a).

The aforementioned property can be used to show that reachability is decidable in PBDC Systems with restricted semantics. Let Π be a PBDC system defined over a finite set of names Λ . Suppose that Λ has cardinality K. Furthermore, assume that the initial configuration $c_0 = (N_0, M_0, \mu_0)$ is such that M_0 is a single node. We first build the set Θ of all possible membrane structures with at most Knodes labelled with distinct labels taken from Λ . As an example, if $\Lambda = \{0, 1, 2\}$, then we will consider all trees with at most three nodes and such that each node has a distinct label taken from Λ , i.e., $[0 \]_0, [1 \]_1, [1 \ [0 \]_0, [1 \]_1, [2 \ [0 \]_0, [1 \]_1]_2$, and so on. Notice that for a fixed membrane structure T we can always determine the immediate ancestor j of a node i (if it exists) at static time.

Starting from Π and Θ , we now define a Petri net \mathcal{N} that satisfies the following conditions. First of all, we associate a place T to each membrane structure $T \in \Theta$. We assume that only one of such places can be non empty during the simulation of the restricted semantics. A non-empty place $T \in \Theta$ corresponds to the current membrane structure. Furthermore, for each $i \in \Lambda$ we add to \mathcal{N} places $used_i$ and $notused_i$ (to model freshness of name i), and, for each $a \in \Gamma$, place a^i (to model the content of membrane i in the current membrane structure). Notice that since names are used at most once, we can safely confuse nodes of membrane structure with their labels (each node has a different label in Λ). PBDC rules are modelled as the finite set of transitions in \mathcal{N} defined as follows.

Internal Rules

For each membrane structure $T \in \Theta$ with a membrane *i*, an internal rule $r = [_i u \rightarrow [_i v \text{ is encoded by a transition } t_r^T \text{ that satisfies the following conditions. The pre-set contains the places <math>T$ (the current membrane structure) and $used_i$ (*i* is in use) together with multiset $\pi_i(u)$. The post-set contains places T and $used_i$ together with multiset $\pi_i(v)$. Thus, only the internal rules defined on the current membrane structure are enabled.

Boundary Rules

For each membrane structure $T \in \Theta$ with a membrane j with immediate successor i, a boundary rule $r = u_i v \to u'_i v'$ is encoded by a transition $b_r^{T,i,j}$ that satisfies the following conditions. The pre-set contains places T, $used_i$, $used_j$, and the multisets $\pi_i(u)$ and $\pi_i(v)$. The post-set contains places T, $used_i$, $used_j$, and the multisets $\pi_i(u')$ and $\pi_i(v')$. Thus, only boundary rules defined on the current membrane structure are enabled.

Creation Rules

For each $T \in \Theta$ such that *i* does not occur in the set of names in *T* (the side condition that ensures the freshness of generated membrane names), and for each

name j occurring in T, a creation rule $r = a \rightarrow [iv]_i$ is encoded by a transition $c_r^{T,i,j}$ that satisfies the following conditions. The pre-set contains places T, $used_j$, $notused_i$, and a^j . The post-set contains places $used_i$ and $used_j$, the multiset $\pi_i(v)$, and the place $T_{j+i} \in \Theta$ associated to the membrane structure obtained from T by adding a new node labelled i as immediate successor of j.

Dissolution Rules

For each membrane structure $T \in \Theta$ with a membrane j with immediate successor i, a dissolution rule $r = [{}_{i}u \rightarrow [{}_{i}v \cdot \delta$ is encoded by the following set of transitions. We first define a transition $c_r^{T,i,j}$ that starts the dissolution phase of node *i*. The pre-set of $c_r^{T,i,j}$ contains places T, $used_i$, $used_j$, and the multiset $\pi_i(u)$. The postset contains the place $dissolve^{T,i,j}$, and, the multiset $\pi_i(v)$. Notice that, by removing a token from the place T, we automatically disable all transitions not involved in the dissolution phase (i.e. T plays the role of flag normal used for simulating dissolution rules in PBD systems). Now, for each $a \in \Gamma$, we model the transfer of the content of node i to node j via a transition $m_r^{T,i,j,a}$ that satisfies the following conditions: The pre-set contains the places $dissolve^{T,i,j}$ and a^i (the source of a token to be transferred). The post-set contains the places $dissolve^{T,i,j}$ and a^j (the destination of a transferred token). Finally, let T_{j-i} be the membrane structure obtained by T by removing membrane i and moving all of its sub-membranes into membrane j. Then, we add transition $d_r^{T,i,j}$ to non-deterministically stop the transfer of tokens and to update the membrane structure to T_{i-i} , i.e., the pre-set of this transition contains the place $dissolve^{T,i,j}$ and its post-set contains the places T_{i-i} , used_i and used_i. Notice that name i remains marked as used after dissolving the corresponding membrane (i.e. it cannot be used in successive creation rules).

Places, Transitions and Configurations

The Petri net \mathcal{N} is built by taking the union of the places and transitions used to encode internal, boundary, creation and dissolution rules described before. A generic configuration $c = (N, M, \mu)$ is encoded by a marking m_c in which: there is one token in the place associated to the membrane structure M, one token in $used_i$ for each $i \in N$, one token in $notused_i$ for each $i \in \Lambda \setminus N$, and, for each i that occurs in M and for each $a \in \Gamma$, as many tokens in a^i as the number of occurrences of object a in $\mu(i)$. All other places are empty.

Reachability Problem

Notice that after a membrane with name i is introduced by a creation rule (i.e. the place $unused_i$ is emptied while one token is put in $used_i$), no other membranes with the same name can be created (there is no rule that puts a token back to $unused_i$). The membrane i however can dissolve in a successive transition, i.e. in a target configuration $used_i$ can be non-empty (i.e. $i \in N$), even if i does not occur in the current membrane structure. Also notice that in m_c we enforce all places associated to membranes not occurring in M to be empty. The combination of these

two properties allows us to distinguish good simulations (i.e. in which after the application of dissolution rules all tokens are transferred to the father membrane) from bad ones (some tokens are left in a place a^i , $used_i$ is non empty, but i is no more in the current membrane structure). Following from this observation and from the construction of \mathcal{N} , we have that $c = (N, M, \mu)$ is reachable from c_0 if and only if the marking m_c is reachable from m_{c_0} .

Following from the decidability of reachability in Petri nets, we obtain the following theorem.

Theorem 4. Reachability is decidable in PBDC systems with restricted semantics.

6 Boundedness Problem for Extended PB Systems

In [5] Dal Zilio and Formenti exploit the Petri net encoding used for deciding reachability to prove that boundedness is decidable too for PB systems with symbol objects. In this section we investigate the boundedness problem for the different extensions of PB systems proposed in the present paper. In particular, we show that the boundedness problem is decidable for PBDC systems with restricted semantics where a membrane name can be used at most once. This result implies that boundedness is decidable for PB systems with dissolution (and standard semantics). Finally, we show undecidability of the boundedness problem for PB systems with creation and standard semantics.

6.1 Boundedness for PBDC Systems with Restricted Semantics

To prove decidability of boundedness in PBDC systems with restricted semantics, let us first define the following partial order \sqsubseteq over configurations. Assume two configurations $c_1 = (N_1, M_1, \mu_1)$ and $c_2 = (N_2, M_2, \mu_2)$. We define $c_1 \sqsubseteq c_2$ if and only if $N_1 = N_2$ and $M_1 = M_2$ (i.e. c_1 and c_2 have the same tree structure), and $\mu_1(n) \le \mu_2(n)$ (the multiset associated to n in c_1 is contained in that associated to n in c_2) for all node n in M_1 . If we fix an upper bound on the number of possible nodes occurring in a membrane structure along a computation, then \sqsubseteq has the following property.

Proposition 1. Fixed $a \ k \in \mathbb{N}$, for any infinite sequence of configurations $c_1c_2...$ with membrane structure of size at most k, there exist positions i < j such that $c_i \sqsubseteq c_j$ (i.e. \sqsubseteq is a well-quasi ordering).

The proof is a straightforward application of composition properties of well-quasi ordering, see e.g. [1]. Now assume an infinite computation $c_0 = (N_0, M_0, \mu_0)c_1 = (N_1, M_1, \mu_1) \dots$ of a PBDC system with restricted semantics. From Observation 1 it follows that for all $i \ge 0$ the number of nodes in M_i is bounded by the number of possible names. Hence, by Prop. 1 we know that there exist positions i < jsuch that $c_i \sqsubseteq c_j$. Furthermore, if $c_i \sqsubseteq c_j$ and $c_i \ne c_j$, then $N_i = N_j$, $M_i = M_j$, and $\mu_i \prec \mu_j$. Thus, the transition sequence σ from c_i to c_j does not modify the membrane structure but strictly increases the number of objects contained at each of its node. This implies that the application of σ can be iterated starting from c_j , leading to a infinite strictly increasing, w.r.t \sqsubseteq , sequence of configurations.

As a consequence of these properties, the boundedness problem for PBDC systems can be decided by building a computation tree T such that: the root node n_0 of T is labelled by the initial configuration c_0 , if n_0, \ldots, n_k is a path in T such that n_i is labelled with c_i $i : 0, \ldots, k$, and for all $i : 0, \ldots, k - 1$ $c_i \neq c_k$ and $c_i \not\sqsubseteq c_k$ then we add a node n' labelled c' as successor of n_k if and only if $c_k \Rightarrow c'$. Furthermore, the PBDC system is not bounded if and only if there exists a leaf n labelled with c and a predecessor n' of n labelled with c' in T such that $c' \sqsubseteq c$. Since, \sqsubseteq is a decidable relation and, the tree T is finite (by Observation 1 and Prop. 1), the following property then holds.

Theorem 5. The boundedness problem is decidable for PBDC systems with restricted semantics.

From Theorem 5, we know that boundedness is decidable for PB systems (consistently with the result in [5]) and for PB systems with dissolution (they form a subclass of PBDC systems where the restricted and standard semantics coincides).

6.2 Boundedness Problem for PBC Systems

The boundedness problem turns out to be undecidable for PBC systems with standard semantics in which there is no limit on the number of instances of a given type of membranes that can be created during a computation. The proof is based on a reduction of counter machines with increment, decrement and zero-test to PBC systems. The idea is to use nested membranes to model the current value of a counter. For instance,

$$[_1 used \ [_1 used \ [_1 unused \ [_1 end \]_1]_1]_1$$

can be used to encode counter c_1 with value 2 (the number of occurrences of symbol *used*). Hence a configuration of a two counter machine with both counters set to zero is encoded as a configuration of the form

$$[0 \ \ell \ [1 \ unused \ [1 \ \dots \ [1 \ end \]_1 \ \dots \]_1]_1 [2 \ unused \ [2 \ \dots \ [2 \ end \]_2 \ \dots \]_2 \]_2 \]_0$$

where ℓ is a symbol corresponding to the current location of the two counter machine, and membrane with name *i* encodes counter *i* for *i* : 1, 2. Increment of counter *i* in location ℓ with ℓ_1 as successor location is simulated by replacing the first *unused* symbol encountered when descending the tree from the membrane 0 with the symbol *used*. This is implemented by the following set of rules that descend the structure of a membrane of type *i* in search for the first occurrence of symbol *unused*:

```
 \begin{split} \ell & [i \text{ unused } \to \ell_1 \text{ } [i \text{ used } \\ \ell & [i \text{ used } \to \ell'_i \text{ } [i \text{ down } \\ down \text{ } [i \text{ used } \to \text{ down } [i \text{ down } \\ down \text{ } [i \text{ unused } \to \text{ up } [i \text{ used } \\ down \text{ } [i \text{ up } \to \text{ up } [i \text{ used } \\ \ell'_i \text{ } [i \text{ up } \to \ell_1 \text{ } [i \text{ used } \\ \end{split}
```

where ℓ'_i , down, up are auxiliary symbols (ℓ'_i is blocking for the other counter(s), down propagate the search down in the tree, up propagate the success notification up in the tree). Furthermore, if the current tree has no membrane containing the unused symbol (i.e. all membranes contain used) then a new membrane is created by applying the following additional rules:

$$down [_i end \rightarrow down [_i create create \rightarrow [_i exit \cdot end]_i \epsilon [_i exit \rightarrow up [_i \epsilon$$

where ϵ denotes the empty multiset. Decrement is simulated by changing the last occurrence of *used* encountered by descending the membrane tree from the root into symbol *unused*. We can safely assume here that the counter is non-zero, i.e., that there is at least one membrane with *used* object. The rules that implement decrement are defined as follows.

$$\begin{array}{ll} \ell \; [i \; used \; \rightarrow \; \ell_i' \; [i \; used_1 \\ used_1 \; [i \; used \; \rightarrow \; used_1 \; [i \; used_1 \\ used_1 \; [i \; unused \; \rightarrow \; used_2 \; [i \; unused \\ used_1 \; [i \; end \; \rightarrow \; used_2 \; [i \; end \\ used_1 \; [i \; used_2 \; \rightarrow \; used_3 \; [i \; unused \\ used_1 \; [i \; used_3 \; \rightarrow \; used_3 \; [i \; unused \\ used_1 \; [i \; used_3 \; \rightarrow \; used_3 \; [i \; used \\ \ell_i' \; [i \; used_2 \; \rightarrow \; \ell_1 \; [i \; unused \\ \ell_i' \; [i \; used_3 \; \rightarrow \; \ell_1 \; [i \; used \\ \end{array}$$

where ℓ'_i , $used_1$, $used_2$, $used_3$ are auxiliary symbols, $used_1$ is used to mark nodes during the downward search (for unused), $used_2$ is used to mark the used node to be replaced by unused, and $used_3$ is used to replace nodes marked with $used_1$ with used during the return from the search. The zero-test on counter c_i in location ℓ with successor ℓ_1 can be implemented by testing if all objects in membranes i are unused (or end). Note that increment, resp. decrement, of c_i is encoded by a topdown traversal of membranes i until reaching a membrane containing an object unused, resp. used, which is then replaced by used, resp. unused. Furthermore, each membrane contains one object. Hence, no membrane i contain a used symbols if and only if the top level membrane i has object unused or end. This can be checked with the following rules:

$$\ell [i \text{ unused } \rightarrow \ell_1 [i \text{ unused } \\ \ell [i \text{ end } \rightarrow \ell_1 [i \text{ end }]$$

Similarly, testing that c_i is not equal to zero in location ℓ with successor ℓ_1 amounts to check that the top level membrane *i* has object *used*, i.e.,

$$\ell [i used \rightarrow \ell_1 [i used]$$

By construction, we directly have that a two counter machine is bounded (Are its counters bounded by a constant $k \in \mathbb{N}$?) if and only if its encoding into PBC systems is bounded. Since boundedness is undecidable for two-counter machines with zero-test, we obtain the following negative result.

Theorem 6. The boundedness problem is undecidable for PBC systems.

Remark The proof of Theorem 6 shows that PBC systems can simulate two-counter machines by means of membrane structures with unbounded depth. Let us now go back to the reachability problem of a target configuration μ . Since PBC rule never remove membranes, the target configuration μ gives us an upper bound on the size of membrane structures that may occur in a sequence of transition leading to it. Thus, the set of possible membrane structures than we have to consider to solve a reachability problem is always finite (Notice that this not imply that we have to consider a finite number of configurations). In other words, for reachability problems, we do not have to deal with the full computational power of PBC systems. For this reason, the undecidability of boundedness proved in Theorem 6 is not in contradiction with the decidability of reachability proved in Theorem 2. Similar results obtained for fragments of process calculi [3, 4] seem to indicate that, in general, the decidability of reachability cannot be use to give an estimation of the expressive power of a computational model.

7 Conclusions

In this paper we have investigated the decidability of reachability and boundedness in extensions of PB systems with rules that dynamically modify the tree structure of membranes. We conjecture that some of the positive results presented here can be extended to PB systems with some form of movement and with dynamic generation of membrane names. We plan to investigate these problems in future work.

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A Cellular Solution to Subset Sum Using Division of Non-elementary Membranes and Dissolution, with Time and Initial Resources Bounded by $\log k$

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Summary. The aim of our paper is twofold. On one hand we prove the ability of polarizationless P systems with dissolution and with division rules for non-elementary membranes to solve **NP**-complete problems in a polynomial number of steps, and we do this by presenting a solution to the Subset Sum problem. On the other hand, we improve some similar results obtained for different models of P systems by reducing the number of steps and the necessary resources to be of a logarithmic order with respect to k (recall that n and k are the two parameters used to indicate the size of an instance of the Subset Sum problem).

As the model we work with does not allow cooperative rules and does not consider the membranes to have an associated polarization, the strategy that we will follow consists on using objects to represent the weights of the subsets through their multiplicities, and comparing the number of objects against a fixed number of membranes. More precisely, we will generate k membranes in log k steps.

1 Introduction

This paper is the continuation of a series of results on Complexity Classes in Membrane Computing that are trying to establish the relevance, in terms of computing power, of each one of the possible features of a P system (see [3]).

The Subset Sum problem is a well-known **NP**-complete problem which can be formulated as follows: Given a finite set A, a weight function, $w : A \to \mathbb{N}$, and a constant $k \in \mathbb{N}$, determine whether or not there exists a subset $B \subseteq A$ such that w(B) = k. It has been a matter of study in Membrane Computing several times, being mainly used to prove the ability of different P system models in order to solve problems from the **NP** class in a polynomial time.

This speed-up is achieved by trading space for time, in the sense that the considered models allow that an exponential amount of membranes can be produced by a P system in a polynomial number of steps. For example, solutions to the Subset Sum problem working in a number of steps which is linear with respect to the parameters n and k have been designed using P systems with active membranes [9], using tissue P systems with cell division [2], and using P systems with membrane creation [4].

In this paper we work with P systems using division of non-elementary membranes and dissolution rules. Our aim goes beyond adding this P system model to the above mentioned list; we improve previous complexity results by solving the Subset Sum problem in a linear number of steps with respect to n and log k. We also improve the pre-computation process, as the initial resources are also bounded by log k.

The paper is structured as follows: in the next section we present the formal framework, i.e., we recall the definition of recognizing P systems, the P system model used along the paper is settled and the class $\mathbf{PMC}_{\mathcal{AM}^0(+d,+ne)}$ is presented. In Section 3, our design of the solution of the Subset Sum problem is presented and some conclusions are given in the last section.

2 Formal Framework

In this paper we are using cellular systems for attacking the resolution of decision problems. This means that for each instance of a problem that we try to solve, we are only interested in obtaining a Boolean answer (*Yes* or *No*). Therefore, the P system can behave as a *black box* to which the user supplies an input and from which an affirmative or negative answer is received. This is indeed the motivation for defining the concept of *recognizing P systems* (introduced in [13]).

2.1 Recognizing P Systems

Let us recall that a decision problem, X, is a pair (I_X, θ_X) where I_X is a language over an alphabet whose elements are called *instances* and θ_X is a total Boolean function over I_X . If u is an instance of the problem X such that $\theta_X(u) = 1$ (respectively, $\theta_X(u) = 0$), then we say that the answer to the problem for the instance considered is Yes (respectively, No).

Keeping this in mind, recognizing P systems are defined as a special class of membrane systems that will be used to solve decision problems, in the framework of the complexity classes theory. Note that this definition is stated informally, and it can be adapted for any kind of membrane system paradigm.

A recognizing P system is a P system with input and with external output having two distinguished objects **yes** and **no** in its working alphabet such that:

- All computations halt.
- If C is a computation of Π , then either the object yes or the object no (but not both) must have been released into the environment, and only in the last step of the computation.

2.2 The P System Model

The power of membrane division as a tool for efficiently solving **NP** problems in Membrane Computing has been widely proved. Many examples of designs of P systems solving **NP**-complete problems have been proposed in the framework of P systems with active membranes with two polarizations and three polarizations and in the framework of P systems with non-elementary membrane division. The key of such solutions is the creation of an exponential amount of workspace (membranes) in a polynomial time.

In the literature, one can find two quite different rules for performing membrane division. On the one hand, in [7], P systems with active membranes were presented. In this model new membranes were obtained through the process of *mitosis* (membrane division). In these devices membranes have polarizations, one of the "electrical charges" 0, -, +, and several times the problem was formulated whether or not these polarizations are necessary in order to obtain polynomial solutions to **NP**-complete problems. The last result is that from [1], where one proves that two polarizations suffice.

P systems with active membranes have been successfully used to design (uniform) solutions to well-known **NP**-complete problems, such as SAT [13], Subset Sum [9], Knapsack [10], Bin Packing [11], Partition [5], and the Common Algorithmic Problem [12].

The syntactic representation of membrane division rule is

$$[a]_{h}^{e_{1}} \to [b]_{h}^{e_{2}} [c]_{h}^{e_{3}} \tag{1}$$

where h is a label, e_1, e_2 and e_3 are electrical charges and a, b and c are objects. The interpretation is well-known: An elementary membrane can be divided into two membranes with the same label, possibly transforming some objects and changing the electrical charge. All objects present in the membrane except the object triggering the rule are copied into both new membranes.

In [6], a variant of this rule was used in which the polarization was dropped:

$$[a]_h \to [b]_h [c]_h. \tag{2}$$

In both cases (with and without polarizations) the key point is that the membranes are always elementary membranes. In the literature, there also exist rules for the division of non-elementary polarizationless membranes, as

$$[[]_{h_1}[]_{h_2}]_{h_0} \to [[]_{h_1}]_{h_0}[[]_{h_2}]_{h_0} \tag{3}$$

where h_0, h_1 and h_2 are labels. There exists an important difference with respect to elementary membrane division: in the case 3, the rule is not triggered by the occurrence of an object inside a membrane, but by the membrane structure instead. This point has a crucial importance in the design of solutions, since a membrane can be divided by the corresponding rule even if there are no objects inside it. 304 D. Díaz-Pernil et al.

According to the representation (3), the membrane h_2 divides into two new membranes also with label h_2 and all the information (objects and membranes) inside is duplicated.

In this paper we use a type of membrane division which is syntactically equivalent to (2)

$$[a]_h \to [b]_h [c]_h, \tag{4}$$

but we will consider a semantic difference; the dividing membrane can be elementary or non-elementary and after the division, all the objects and membranes inside the dividing membrane are duplicated, except the object a that triggers the rule, which appears in the new membranes possibly modified (represented as objects band c).

In this paper we work with a variant of P systems with active membranes and weak division that does not use polarizations.

Definition 1. A P system with active membranes with weak division is a P system with Γ as working alphabet, with H as the finite set of labels for membranes, and where the rules are of the following forms:

- (a) $[a \to u]_h$ for $h \in H$, $a \in \Gamma$, $u \in \Gamma^*$. This is an object evolution rule, associated with a membrane labeled with h: an object $a \in \Gamma$ belonging to that membrane evolves to a multiset $u \in \Gamma^*$.
- (b) $a[]_h \to [b]_h$ for $h \in H$, $a, b \in \Gamma$. An object from the region immediately outside a membrane labeled with h is introduced in this membrane, possibly transformed into another object.
- (c) $[a]_h \to b []_h$ for $h \in H$, $a, b \in \Gamma$. An object is sent out from membrane labeled with h to the region immediately outside, possibly transformed into another object.
- (d) $[a]_h \to b$ for $h \in H$, $a, b \in \Gamma$: A membrane labeled with h is dissolved in reaction with an object. The skin is never dissolved.
- (e) $[a]_h \to [b]_h [c]_h$ for $h \in H$, $a, b, c \in \Gamma$. A membrane can be divided into two membranes with the same label, possibly transforming some objects. The content of the membrane is duplicated. The membrane can be elementary or not.

These rules are applied according to the following principles:

- All the rules are applied in parallel and in a maximal manner. In one step, one object of a membrane can be used by only one rule (chosen in a non-deterministic way), but any object which can evolve by one rule of any form, must evolve.
- If at the same time a membrane labeled with *h* is divided by a rule of type (e) and there are objects in this membrane which evolve by means of rules of type (a), then we suppose that first the evolution rules of type (a) are used, and then the division is produced. Of course, this process takes only one step.

• The rules associated with membranes labeled with *h* are used for all copies of this membrane. At one step, a membrane can be the subject of *only one* rule of types (b)-(e).

Let us note that in this framework we work without cooperation, without priorities, with weak division, and without changing the labels of membranes.

In this paper we work within the model of polarizationless P systems using weak division of non-elementary membranes and dissolution. Let $\mathcal{AM}^0(+d, +ne)$ be the class of such systems.

2.3 The Class $PMC_{\mathcal{AM}^0(+d,+ne)}$

Definition 2. We say that a decision problem $X = (I_X, \theta_X)$ is solvable in polynomial time by a family $\Pi = \{\Pi(n) : n \in \mathbb{N}\}$ of recognizing P systems from $\mathcal{AM}^0(+d, +ne)$ if the following holds:

- The family Π is polynomially uniform by Turing machines, that is, there exists a deterministic Turing machine working in polynomial time which constructs the system $\Pi(n)$ from $n \in \mathbb{N}$.
- There exists a pair (cod, s) of polynomial-time computable functions over I_X such that:
 - for each instance $u \in I_X$, s(u) is a natural number and cod(u) is an input multiset of the system $\Pi(s(u))$;
 - the family Π is polynomially bounded with regard to (X, cod, s), that is, there exists a polynomial function p, such that for each $u \in I_X$ every computation of $\Pi(s(u))$ with input cod(u) is halting and, moreover, it performs at most p(|u|) steps;
 - the family Π is sound with regard to (X, cod, s), that is, for each $u \in I_X$, if there exists an accepting computation of $\Pi(s(u))$ with input cod(u), then $\theta_X(u) = 1$;
 - the family Π is complete with regard to (X, cod, s), that is, for each $u \in I_X$, if $\theta_X(u) = 1$, then every computation of $\Pi(s(u))$ with input cod(u) is an accepting one.

In the above definition we have imposed to every P system $\Pi(n)$ a confluent condition, in the following sense: every computation of a system with the same input multiset must always give the same answer. The pair of functions (cod, s) is called a *polynomial encoding* of the problem in the family of P systems.

We denote by $\mathbf{PMC}_{\mathcal{AM}^0(+d,+ne)}$ the set of all decision problems which can be solved by means of recognizing polarizationless P systems using division of non-elementary membranes and dissolution in polynomial time.

3 Designing the Solution to Subset Sum

In this section we address the resolution of the problem following a brute force algorithm, implemented in the framework of recognizing P systems from the $\mathcal{AM}^{0}(+d, +ne)$ class. The idea of the design is better understood if we divide the solution to the problem into several stages:

- Generation stage: for every subset of A, a membrane labeled by e is generated via membrane division.
- Calculation stage: in each membrane the weight of the associated subset is calculated (using the auxiliary membranes e_0, \ldots, e_n).
- Checking stage: in each membrane it is checked whether the weight of its associated subset is exactly k (using the auxiliary membranes ch).
- *Output stage*: the system sends out the answer to the environment, according to the result of the checking stage.

Let us now present a family of recognizing P systems from the $\mathcal{AM}^0(+d, +ne)$ class that solves Subset Sum, according to Definition 2.

We shall use a tuple $(n, (w_1, \ldots, w_n), k)$ to represent an instance of the Subset Sum problem, where *n* stands for the size of $A = \{a_1, \ldots, a_n\}, w_i = w(a_i)$, and *k* is the constant given as input for the problem. Let $g : \mathbb{N} \times \mathbb{N} \to \mathbb{N}$ be a function defined by

$$g(n,k) = \frac{(n+k)(n+k+1)}{2} + n$$

This function is primitive recursive and bijective between $\mathbb{N} \times \mathbb{N}$ and \mathbb{N} and computable in polynomial time. We define the polynomially computable function s(u) = g(n, k).

We shall provide a family of P systems where each P system solves all the instances of the Subset Sum problem with the same size. Let us consider the binary decomposition of k, $\Sigma_{i \in I} 2^i = k$, where the indices $i \in I$ indicate the positions of the binary expression of k where a 1 occurs. Let $I' = \{1, \ldots, \lfloor \log k \rfloor\} - I$ be the complementary set, that is, the positions where a 0 occurs. This binary encoding of k, together with the weight function w of the concrete instance, will be provided via an input multiset determined by the function cod as follows:

$$cod(u) = cod_1(u) \cup cod_2(u),$$

where
$$cod_1(u) = \{\{b_i^{w_i} : 1 \le i \le n\}\}$$
 and
 $cod_2(u) = \{\{c_j : j \in I\}\} \cup \{\{c'_j : j \in I'\}\}$

Next, we shall provide a family $\mathbf{\Pi} = \{\Pi(g(n,k)) : n, k \in \mathbb{N}\}$ of recognizing P systems which solve the Subset Sum problem in a number of steps being of $O(n + \log k)$ order. We shall indicate for each system of the family its initial configuration and its set of rules. We shall present the list of rules divided by groups, and we shall provide for each of them some comments about the way their rules work.

Let us consider an arbitrary pair $(n,k) \in \mathbb{N} \times \mathbb{N}$. The system $\Pi(g(n,k))$ is determined by the tuple $(\Gamma, \Sigma, \mu, M, \mathcal{R}, i_{in}, i_0)$, that is described next:

• Alphabet:



Fig. 1. Initial Configuration

$$\begin{split} \Gamma &= \Sigma \cup \{b_i^+, b_i^-, b_i^=, d_i, d_i^+, d_i^-, p_i, q_i : i = 1, \dots, n\} \\ &\cup \{g_0, \dots, g_2\lfloor \log k \rfloor + 2, h_0, \dots, h_2\lfloor \log k \rfloor + 2n + 8, l_0, \dots, l_2\lfloor \log k \rfloor + 2n + 10\} \\ &\cup \{r_0, \dots, r_2\lfloor \log k \rfloor + 2n + 7, v_0, \dots, v_2\lfloor \log k \rfloor + 2n + 12\} \\ &\cup \{w_0, \dots, w_2\lfloor \log k \rfloor + 2n + 18\} \\ &\cup \{x_0, \dots, x_2\lfloor \log k \rfloor + 2n + 15, z_0, \dots, z_{\lfloor \log k \rfloor}\} \\ &\cup \{s, \text{yes}, \text{no}, Trash\} \end{split}$$

• Input alphabet: $\Sigma(n,k) = \{b_1, \ldots, b_n, c_0, \ldots, c_{\lfloor \log k \rfloor}, c'_0, \ldots, c'_{\lfloor \log k \rfloor}\}.$

The initial configuration consists of $n + \lfloor \log k \rfloor + 9$ membranes, arranged as shown in Figure 1. Formally, the membrane structure μ is

$$[[[[[\dots n \quad [[[[[]_{ch} \dots []_{ch}]_{a'}]_a]_{e_0}]_{e_1} \quad \stackrel{n}{\dots}]_{e_n}]_{a_2} \quad []_c]_{a_1}]_e]_f]_{skin}$$

where there are exactly $|\log k| + 1$ copies of membrane $[]_{ch}$.

Roughly speaking (more precise explanations will be given for the rules), we can classify the membranes according to their role as follows:

- n+2 membranes that take care of the generation stage, namely those labeled by $e_0, e_1, \ldots e_n$ and e.
- $\lfloor \log k \rfloor + 3$ membranes that take care of preparing and implementing the checking stage, namely those labeled by ch, a and a'.
- 4 membranes that take care of the answer stage, handling and synchronizing the results of the checking, namely those labeled by a_1, a_2, c and f.
- The initial multisets are:

$$M(f) = \{\{w_0\}\}; M(e) = \{\{g_0\}\}; M(a_1) = \{\{v_0\}\}; M(a_2) = \{\{h_0\}\};$$
$$M(c) = \{\{x_0\}\}; M(a) = \{\{r_0\}\}; M(a') = \{\{z_0\}\}; M(ch) = \{\{l_0\}\}$$
$$M(skin) = M(e_0) = \dots = M(e_n) = \emptyset$$

• The input membrane is $i_{in} = e_0$, and the output region is the environment $(i_0 = env)$.

First task: generate k membranes ch

At the beginning of the computation, k membranes ch will be generated inside the innermost region of the structure.

The strategy works as follows:

- 1. Initially, there are $\lfloor \log k \rfloor$ membranes ch in the region a', and the input multiset is located in region e_0 (recall that $cod_2(u)$ consists of $\lfloor \log k \rfloor$ objects c_i or c'_i representing the binary encoding of k).
- 2. In the first $\lfloor \log k \rfloor$ steps, the objects from $cod_2(u)$ get into membrane a (the objects enter one by one membrane a). Simultaneously, the counter z_i is evolving inside membrane a' and dissolves it at the $\lfloor \log k \rfloor$ step.
- 3. Thus, in the next step each element from $cod_2(u)$ will go inside a membrane ch (all objects go in parallel into different membranes in a one-to-one manner).
- 4. Objects c'_i will dissolve the membranes where they enter, while each object c_i will generate by division 2^i membranes ch.
- 5. After at most $\lfloor \log k \rfloor$ further steps all divisions have been completed, and the number of membranes ch is exactly k.

Membrane a will not be divided until the generation and weight calculation stages have been completed, acting as a separator between objects from $cod_1(u)$ and membranes ch.

$$\begin{array}{ll} \mathbf{Set} \ (\mathbf{A1}). & c_i[]_a \to [c_i]_a \\ & c'_i[]_a \to [c'_i]_a \\ & c_i[]_{ch} \to [c_i]_{ch} \\ & c'_i[]_{ch} \to [c'_i]_{ch} \\ & [c'_i]_{ch} \to Trash \end{array} \right\} \text{ for } i \in \{0, \dots, \lfloor \log k \rfloor\}.$$

Set (A2).

$$\begin{bmatrix} c_0 \to Trash \end{bmatrix}_{ch} \\ \begin{bmatrix} c_i \end{bmatrix}_{ch} \to \begin{bmatrix} c_{i-1} \end{bmatrix}_{ch} \begin{bmatrix} c_{i-1} \end{bmatrix}_{ch} & \text{for } i = 1, \dots, \lfloor \log k \rfloor \\ \begin{bmatrix} z_i \to z_{i+1} \end{bmatrix}_{a'} & \text{for } i = 0, \dots, \lfloor \log k \rfloor - 1 \\ \begin{bmatrix} z_{\lfloor \log k \rfloor} \end{bmatrix}_{a'} \to Trash \\ \begin{bmatrix} g_i \to g_{i+1} \end{bmatrix}_{a'} & \text{for } i = 0, \dots, 2\lfloor \log k \rfloor + 1 \\ \begin{bmatrix} g_{2 \lfloor \log k \rfloor + 2} \to d_1 s \end{bmatrix}_e \end{bmatrix}$$

In the last step of this stage, the counter g_i produces the objects d_1 and s which will trigger the beginning of the next stage.

$$\begin{aligned} \mathbf{Set}(\mathbf{B}). & \begin{bmatrix} w_i \to w_{i+1} \end{bmatrix}_f \\ \begin{bmatrix} v_i \to v_{i+1} \end{bmatrix}_{a_1} \\ \begin{bmatrix} h_i \to h_{i+1} \end{bmatrix}_{a_2} \\ \begin{bmatrix} x_i \to x_{i+1} \end{bmatrix}_c \\ \begin{bmatrix} r_i \to r_{i+1} \end{bmatrix}_a \\ \begin{bmatrix} l_i \to l_{i+1} \end{bmatrix}_{ch} \end{aligned} \right\} \text{ for } i \in \{0, \dots, 2\lfloor \log k \rfloor + 2\}. \end{aligned}$$

The rest of the counters simply increase their indices in this stage. (See Fig. 2).

Second task: generate 2^n membranes e

Objects d_i residing inside membrane(s) e will produce n consecutive divisions, thus yielding 2^n copies of membrane e. To each one of them, a subset of A is associated in the following way: after each division, the membranes where object p_i occurs correspond to subsets of A containing a_i , and conversely, membranes where q_i occurs will be associated with subsets not containing a_i .

$$\begin{aligned} \mathbf{Set}(\mathbf{C}). \quad & [d_i]_e \to [d_i^+]_e [d_i^-]_e \quad \text{ for } i = 1, \dots n \\ & [d_i^+ \to p_i d_{i+1}]_e \quad \text{ for } i = 1, \dots n-1 \\ & [d_i^- \to q_i d_{i+1}]_e \quad \text{ for } i = 1, \dots n-1 \\ & [d_n^+ \to p_n]_e \\ & [d_n^- \to q_n]_e \end{aligned}$$

Membrane divisions take place every two steps, so in the $(2\lfloor \log k \rfloor + 2n + 2)$ -th step there will be 2^n membranes e.

$$\begin{aligned} \mathbf{Set}(\mathbf{D}). & s\,[\,]_{a_i} \to [s]_{a_i} & \text{for } i = 1, 2 \\ & s\,[\,]_{e_i} \to [s]_{e_i} & \text{for } i = 0, \dots, n \\ & [s]_{e_0} \to Trash \\ & p_j\,[\,]_{a_i} \to [p_j]_{a_i} & \text{for } i = 1, 2 \quad j = 1, \dots, n \\ & p_j\,[\,]_{e_i} \to [p_j]_{e_i} & \text{for } j = 1, \dots, n & i = j, \dots, n \\ & [p_i \to q_i]_{e_i} & \text{for } i = 1, \dots, n \\ & q_j\,[\,]_{a_i} \to [q_j]_{a_i} & \text{for } i = 1, 2 \quad j = 1, \dots, n \\ & q_j\,[\,]_{e_i} \to [q_j]_{e_i} & \text{for } j = 1, \dots, n & i = j, \dots, n \\ & [q_i]_{e_i} \to Trash & \text{for } i = 1, \dots, n \end{aligned}$$

While the divisions are being carried out, objects s, p_j and q_j , for j = 1, ..., n, travel into inner membranes (recall that whenever membrane e gets divided, the



Fig. 2. TIME $2\lfloor \log k \rfloor + 3$

internal nested structure of membranes e_i is duplicated). In the $(2\lfloor \log k \rfloor + n + 2)$ -th step, an object s arrives to every membrane e_0 . This object dissolves the membrane in the next step, and therefore in the $(2\lfloor \log k \rfloor + n + 3)$ -th step we find inside every membrane e_1 the multiset $cod_1(u)$, and in this moment the weight calculation stage begins (see rules in **Set** (**E**)).

As we said before, objects p_j and q_j are traveling into inner membranes, until they reach e_j . This is done in such a way that in the $(2\lfloor \log k \rfloor + n + 3)$ -th step there is in each membrane e_1 either an object p_1 or an object q_1 , in addition to the multiset $cod_1(u)$.

Before going on, let us state two points. First, recall that in the input multiset, introduced in e_0 at the beginning of the computation, there are $w(a_i)$ copies of b_i , for i = 1, ..., n. Second, let us note that objects q_i dissolve membrane e_i immediately after arriving to it, while objects p_i take two steps to dissolve membrane e_i (first they are transformed into q_i and in the next step the dissolution takes place). Set(**F**) $[b_i \rightarrow b^+]$

 $\mathbf{Set}(\mathbf{E}).$ [b_1

The basic strategy consists on allowing objects b_i to get transformed into objects b_0 only if the element $a_i \in A$ belongs to the associated multiset.

Let us summarize informally the evolution of objects b_i for all possible cases. Recall that in the $(2\lfloor \log k \rfloor + 2)$ -th step, the counter g_i produces an object s in membrane e:

- At step $t = 2\lfloor \log k \rfloor + 3$ object s enters in e_n and either d_1^+ or d_1^- appear in each one of the two existing copies of membrane e.
- At step $t = 2\lfloor \log k \rfloor + 4$ object s enters in e_{n-1} and either p_1 or q_1 appear in membranes e.
- At step $t = 2\lfloor \log k \rfloor + 5$, after the second division has been carried out, there are 4 membranes labeled by e. Object s enters in e_{n-2} (this happens in all 4 copies) and p_1 or q_1 get into e_n (there are two of each).

• At step $t = 2\lfloor \log k \rfloor + n + 3$ object s arrives into e_0 , and p_1 or q_1 enter in e_2 .

^{• ...}

- At step $t = 2\lfloor \log k \rfloor + n + 4$ object s dissolves e_0 (and hence objects b_i are moved to e_1), and p_1 or q_1 arrive into e_1 .
- At step t = 2⌊log k⌋ + n + 5 objects b₁, b₂ and b₃ have been transformed in b₁⁺, b₂⁻ and b₃⁻, respectively, and they will be located either in e₁ (if the membrane contained an object p₁) or in e₂ (if there was an object q₁ in e₁). Besides, in the same step p₂ or q₂ get into e₂.
- At step $t = 2\lfloor \log k \rfloor + n + 6$
 - Objects b_1^+ evolve to b_0 (if they were in e_1) or to Trash (if they were in e_2).
 - Objects b_2^- evolve to b_2^+ .
 - Objects b_3^{\pm} have been transformed into b_3^{-} (both those that were in e_2 and those in e_1).
 - All the objects b_i^{α} $(i = 1, ..., n \text{ and } \alpha \in \{+, -, =\})$ will be located either in membrane e_2 (if the latter contained an object p_2) or in e_3 (if there was an object q_2 in e_2).
 - Besides, in this moment p_3 or q_3 get into e_3 .

The design has been adjusted in such a way that in the moment when objects p_i and q_i arrive into membranes e_i it happens that the objects b_j^{α} $(j = i, \ldots, n)$ and $\alpha \in \{+, -, =\}$ are located in e_i in half of the membranes or in e_{i+1} in the rest of membranes. In the next step there will be objects b_i^+ in e_i only for those cases where there was an object p_i , and hence the weight of element $a_i \in A$ should be added to the weight of the associated multiset (that is, $w(a_i)$ copies of b_0 will be produced in those membranes).

$$\begin{array}{ll} \mathbf{Set}(\mathbf{F}). & \begin{bmatrix} w_i \to w_{i+1} \end{bmatrix}_f \\ \begin{bmatrix} v_i \to v_{i+1} \end{bmatrix}_{a_1} \\ \begin{bmatrix} h_i \to h_{i+1} \end{bmatrix}_{a_2} \\ \begin{bmatrix} x_i \to x_{i+1} \end{bmatrix}_c \\ \begin{bmatrix} r_i \to r_{i+1} \end{bmatrix}_a \\ \begin{bmatrix} l_i \to l_{i+1} \end{bmatrix}_{ch} \end{array} \right\} \text{ for } i \in \{2\lfloor \log k \rfloor + 3, \dots, 2\lfloor \log k \rfloor + 2n + 6\}.$$

The rest of the counters simply increase their indices during this stage. At the end of the stage, in the $(2\lfloor \log k \rfloor + 2n + 7)$ -th step, r_i will dissolve all membranes a. Therefore, in the next step we have 2^n membranes labeled by e, and inside them (more precisely, inside membranes a_2) we have a multiset of objects encoding the weight of a subset and also exactly k copies of membrane ch, see Fig. 3.

Third task: compare k to the weight of each subset

We shall focus next on the checking stage. That is, the system has to check in all membranes a_2 if the number of objects b_0 (encoding the weight of the associated subset) matches or not the parameter k (represented as the number of membranes ch). This task is performed by the following set of rules (for the sake of simplicity, we denote $\beta = 2\lfloor \log k \rfloor + 2n + 8$):



Fig. 3. TIME $2\lfloor \log k \rfloor + 2n + 8$

 $\begin{array}{lll} \mathbf{Set}(\mathbf{G}). & b_0\,[\,]_{ch} \to [c^*]_{ch} \\ & [b_0 \to u_1]_{a_1} \\ & [c^*]_{ch} \to Trash \\ & [h_\beta]_{a_2} \to Trash \end{array}$

At the step $t = \beta$, objects b_0 get into membranes ch, and simultaneously membrane a_2 is dissolved. There are three possible situations:

- 1. There are exactly k objects b_0 . In this case at step $t = \beta + 1$ there will not be any object b_0 remaining, and all membranes ch have been dissolved.
- 2. The number of objects b_0 is lower than k. In this case at step $t = \beta + 1$ there will not be any object b_0 remaining, but there will be some membranes ch that have not been dissolved (because no object b_0 entered them).
- 3. The number of objects b_0 is greater than k. In this case there are some objects b_0 that could not get inside a membrane ch (recall that the rules are applied in a maximal parallel way, but for each membrane only one object can cross it at a time).

In the second case, inside each membrane ch that has not been dissolved the rules $[l_{\beta+1} \rightarrow l_{\beta+2}]_{ch}$ and $[l_{\beta+2}]_{ch} \rightarrow u_2$ are applied in the steps $t = \beta + 1$ and $t = \beta + 2$, respectively. Hence at step $t = \beta + 3$ there will be an object u_2 in a_1 .

In the third case, the exceeding objects b_0 may, nondeterministically, either get into a membrane ch (avoiding that the dissolution rule is applied to that membrane) or evolve into object u_1 . Irrespectively of the nondeterministic choice, we know that there will be no more objects b_0 in a_1 at step $t = \beta + 2$.

Of course, during this stage the rest of the counters continue evolving:

$$\begin{aligned} \mathbf{Set}(\mathbf{H}). \quad & [l_{\beta+i} \to l_{\beta+i+1}]_{ch} \text{ for } i = 0, 1\\ & [v_{\beta+i} \to v_{\beta+i+1}]_{a_1} \text{ for } i = 0, \dots, 3\\ & [x_{\beta+i} \to x_{\beta+i+1}]_c \text{ for } i = 0, \dots, 6\\ & [w_{\beta+i} \to w_{\beta+i+1}]_f \text{ for } i = 0, \dots, 9 \end{aligned}$$

The next set of rules guarantees that in every membrane where the weight of the associated subset was different from k (and only in such membranes) there will be some objects u_3 .

These objects u_3 , being in membrane a_1 , will go into membranes c and dissolve them. We have here a similar situation as before, as there may be several objects u_3 willing to go into a membrane c. The counter v_i takes care of dissolving membrane a_1 so that any exceeding object u_3 will be moved to membrane e and subsequently transformed into Trash.

Set(I2).
$$u_3 []_c \rightarrow [u_4]_c$$

 $[v_{\beta+4}]_{a_1} \rightarrow Trash$
 $[u_3 \rightarrow Trash]_e$
 $[u_4 \rightarrow u_5]_c$
 $[u_5]_c \rightarrow Trash$

Final task: answer stage

Therefore, only in the branches where the number of objects b_0 were equal to k we have a membrane c inside membrane e at step $\beta + 7$. Besides, we also have a counter w_i evolving in membrane f:

• If the instance of the Subset Sum problem has an affirmative answer, i.e., if there exists a subset of A whose weight is k, then in the step $\beta + 7$ there will be a membrane e with a membrane c inside and an object $x_{\beta+7}$ in it. This object will produce an object **yes** which will dissolve his way out to the environment. On the contrary, if the instance has a negative answer, then there will not exist any membrane c in the system in the step $\beta + 7$ and the object **yes** will not be produced. Hence, the membrane f will not be dissolved by **yes** and when the counter w_i reaches $w_{\beta+10}$, an object **no** will appear and will be sent to the environment.

The set of rules is the following one:

$$\begin{split} \mathbf{Set}(\mathbf{J}). & [x_{\beta+7}]_c \to yes \\ & [yes]_e \to yes \\ & [yes]_f \to yes \\ & [yes]_{skin} \to yes []_{skin} \\ & [w_{\beta+10}]_f \to no \\ & [no]_{skin} \to no []_{skin} \end{split}$$
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Consequently, if the answer is affirmative the P system halts after $\beta + 11$ steps and otherwise after $\beta + 12$ steps.

4 Conclusions

In this paper we have combined different techniques for designing P systems in order to get a uniform family of P systems that solves the Subset Sum problem in the framework of P systems with weak division, with dissolution and without polarization. The main contribution of this paper is related to the Complexity Theory of P systems. The best solution of the **NP**-complete problem Subset Sum in any P system model up to now was linear in both input parameters n and k. In this paper we show that the dependency on k can be significantly reduced, since we show a solution where the resources and the number of steps are of a logarithmic order with respect to k.

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A Formal Framework for P Systems

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Summary. The formalism of P systems is known for many years, yet just recently new derivation modes and halting conditions have been proposed. For developing comparable results, a formal description of their functioning, in particular, of the derivation step is necessary. We introduce a formal general framework for static membrane systems that aims to capture most of the essential features of (tissue) P systems and to define their functioning in a formal way.

1 Introduction

P systems were introduced by Gh. Păun (see [8], [14]) as distributed parallel computing devices, based on inspiration from biochemistry, especially with respect to the structure and the functioning of a living cell. The cell is considered as a set of compartments enclosed by membranes; the membranes are nested one in another and contain objects and evolution rules. The basic model neither specifies the nature of these objects nor the nature of the rules. Specifying these two parameters, a lot of different models of computing have been introduced, see [20] for a comprehensive bibliography. Tissue P systems, first considered by Gh. Păun and T. Yokomori in [18] and [19], also see [11], use the graph topology in contrast to the tree topology used in the basic model of P systems.

In this paper, we design a general class of multiset rewriting systems containing, in particular, P systems and tissue P systems. We recall that any P system may be seen at the most abstract level as a multiset rewriting system with only one compartment, encoding the membrane as part of the object representation. However, this approach completely ignores the inner structure of the system because all structural information is hidden (by an encoding) which makes it difficult do deduce any compartment-related information or to model (processes in) biological systems. At a lower level of abstraction, a P system may be seen as networks of cells (compartments) evolving with multi-cell multiset rewriting rules. At the lowest level, the graph/tree structure appears as well as a specialization of rules which are of a very particular form. This last level is usually used in the area of P systems because it permits to easily specify the system and to incorporate different new types of rules.

It is worth noting that in the definition of membrane systems the application of rules often is defined in a quite informal way. This is related to the fact that for a long time only the maximally parallel derivation mode was considered and a P system was supposed to work only in this mode. Recent developments in P systems area have revealed that other derivation modes as the minimally parallel derivation mode might be considered [5] and allow for many interesting new results, yet depending on specific interpretations of this notion. Moreover, different halting conditions have been investigated (see [10], [1]), too. All these articles have shown that there is a need for a formal definition of part of the semantics of membrane systems as the derivation step and the halting procedure like it was done for splicing test tube systems [6] or networks of language processors [7]. In particular, this is important for a classification of P systems as well as for their implementation. For approaches to find operational and logic based semantics for P systems we refer to [4] and [2]; a Petri net semantics for membrane systems is discussed in [12].

This article is an attempt to fulfill the goal of formally defining a procedural semantics for a quite large number of well-known variants of P systems considered so far in the literature, but, of course, we do not at all claim to have captured all the variants having already appeared in the literature. In order to be quite general we place our reasoning at the abstract level of *networks of cells*, already considered in a slightly different way in [3]. We adapt an implementational point of view and also give a formal definition of the derivation step, the halting condition and the procedure for obtaining the result of a computation. Moreover, we give examples of applying our concepts to some well-known variants of P systems.

2 Preliminaries

We recall some of the notions and the notations we use (for further details see [8] and [17]). Let V be a (finite) alphabet; then V^* is the set of all strings (a language) over V, and $V^+ = V^* - \{\lambda\}$ where λ denotes the empty string. FIN (FIN (V)) denotes the set of finite languages (over the alphabet V), and RE, REG, and MAT^{λ} denote the families of recursively enumerable and regular languages as well as matrix languages, respectively. For any family of string languages F, PsF denotes the family of Parikh sets of languages from F and NF the family of Parikh sets of languages from F we denote the set of all non-negative integers, by \mathbb{N}^k the set of all vectors of non-negative integers.

Let V be a (finite) set, $V = \{a_1, ..., a_n\}$. Then a finite multiset S over V is a mapping $f_S : V \longrightarrow \mathbb{N}$. The mapping f_S specifies the number of occurrences of each $x \in V$ in S. The size of the multiset S is $|S| = \sum_{x \in V} f_S(x)$. A multiset S over V

can also be represented by any string x that contains exactly $f_S(a_i)$ symbols a_i for all $1 \leq i \leq n$, e.g., by $a_1^{f_S(a_1)} \dots a_n^{f_S(a_n)}$, or else by the set $\left\{a_i^{f_S(a_i)} \mid 1 \leq i \leq n\right\}$. The support of S is the set $supp(S) = \{a \in V \mid f(a) \geq 1\}$. For example, the multiset over $\{a, b, c\}$ defined by the mapping $a \to 3, b \to 1, c \to 0$ can be specified by a^3b or $\{a^3, b\}$, its support is $\{a, b\}$.

The set of all finite multisets over the set V is denoted by $\langle V, \mathbb{N} \rangle$. We may also consider mappings f_S of form the $f_S : V \longrightarrow \mathbb{N}_{\infty}$ where $\mathbb{N}_{\infty} = \mathbb{N} \cup \{\infty\}$, i.e., elements of S may have an infinite multiplicity; we shall call such multisets where $f_S(a_i) = \infty$ for at least one $i, 1 \leq i \leq n$, infinite multisets. The set of all such multisets S over V with $f_S : V \longrightarrow \mathbb{N}_{\infty}$ is denoted by $\langle V, \mathbb{N}_{\infty} \rangle$.

Let x and y be two multisets over V, i.e., from $\langle V, \mathbb{N} \rangle$ or $\langle V, \mathbb{N}_{\infty} \rangle$. Then x is called a submultiset of y, written $x \leq y$ or $x \subseteq y$, if and only if $f_x(a) \leq f_y(a)$ for all $a \in V$; if, moreover, $f_x(a) < f_y(a)$ for some $a \in V$, then x is called a strict multiset of y. Observe that for all $n \in \mathbb{N}$, $n + \infty = \infty$, and $\infty - n = \infty$. The sum of x and y, denoted by x+y or $x \cup y$, is a multiset z such that $f_z(a) = f_x(a) + f_y(a)$ for all $a \in V$. The difference of two multisets x and y, denoted by x-y or $x \setminus y$, provided that $y \subseteq x$, is the multiset z with $f_z(a) = f_x(a) - f_y(a)$ for all $a \in V$. Observe that in the following, when taking the sum or the difference of two multisets x and y from $\langle V, \mathbb{N}_{\infty} \rangle$, we shall always assume $\{f_x(a), f_y(a)\} \cap \mathbb{N} \neq \emptyset$.

If $X = (x_1, \ldots, x_n)$ and $Y = (y_1, \ldots, y_n)$ are vectors of multisets over V, then $X \leq Y$ if and only if $x_i \subseteq y_i$ for all $i, 1 \leq i \leq n$; in the same way, sum and difference of vectors of multisets are defined by taking the sum and the difference, respectively, in each component.

3 Network of Cells

In this section we consider a general framework for describing membrane systems with a static membrane structure. We consider membrane systems as a collection of interacting cells containing multisets of objects [3].

Definition 3.1 A network of cells of degree $n \ge 1$ (an NC of degree $n \ge 1$, for short) is a construct

$$\Pi = (V, w_1, w_2, \dots, w_n, R)$$

where

1. V is a finite alphabet;

- 2. $w_i \in \langle V, \mathbb{N}_{\infty} \rangle$, for all $1 \leq i \leq n$, is the multiset initially associated to cell *i*;
- 3. R is a finite set of interaction rules of the form

$$(X \to Y; P, Q)$$

where $X = (x_1, \ldots, x_n)$, $Y = (y_1, \ldots, y_n)$, $x_i, y_i \in \langle V, \mathbb{N} \rangle$, $1 \le i \le n$, are vectors of multisets over V and $P = (p_1, \ldots, p_n)$, $Q = (f_1, \ldots, f_n)$, p_i, f_i , $1 \le i \le n$ are finite sets of multisets over V.

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We remark that in the definition given above w_i might be an infinite multiset. However, in most of the cases, only one cell, called *the environment*, will contain an infinite multiset. Hence we define $Infinite(\Pi)$ as the vector specifying the symbols with infinite multiplicity. More exactly,

Infinite(
$$\Pi$$
) = (inf_1, \ldots, inf_n) where $inf_i = \{a \in V \mid f_{w_i}(a) = \infty\}, 1 \le i \le n$.

Moreover, we define inf'_i , $1 \leq i \leq n$, to be the infinite submultisets of w_i taking into account only the symbols with infinite multiplicity, i.e., $f_{inf'_i}(a) = \infty$ for $f_{w_i}(a) = \infty$ and $f_{inf'_i}(a) = 0$ for $f_{w_i}(a) < \infty$, $a \in V$, as well as w'_i , $1 \leq i \leq n$, to be the finite submultisets of w_i taking into account only the symbols with finite multiplicity, i.e., $f_{w'_i}(a) = 0$ for $f_{w_i}(a) = \infty$ and $f_{w'_i}(a) = f_{w_i}(a)$ for $f_{w_i}(a) < \infty$, $a \in V$.

Remark 3.1 We will also use the notation

$$((x_1,1)\dots(x_n,n)\to (y_1,1)\dots(y_n,n); (p_1,1)\dots(p_n,n), (f_1,1)\dots(f_n,n))$$

for a rule $(X \to Y; P, Q)$. Moreover, if some p_i or f_i is an empty set or some x_i or y_i is equal to the empty multiset, $1 \le i \le n$, then we may omit it from the specification of the rule.

A network of cells consists of n cells, numbered from 1 to n, that contain (possibly infinite) multisets of objects over V; initially cell i contains multiset w_i . Cells can interact with each other by means of the rules in R. An interaction rule

$$((x_1, 1) \dots (x_n, n) \to (y_1, 1) \dots (y_n, n); (p_1, 1) \dots (p_n, n), (f_1, 1) \dots (f_n, n))$$

rewrites objects x_i from cells *i* into objects y_j in cells *j*, $1 \le i, j \le n$ if cells *k*, $1 \le k \le n$, contain all multisets from p_k and do not contain any multiset from f_k . In other words, the first part of the rule specifies the rewriting of symbols, the second part of the rule specifies permitting conditions and the third part of the rule specifies the forbidding conditions. In the next section we give a precise definition for the application of an interaction rule.

For an interaction rule r of the form above, the set

$$\{i \mid x_i \neq \lambda \text{ or } f_i \neq \emptyset \text{ or } p_i \neq \emptyset \text{ or } y_i \neq \lambda\}$$

induces a relation between the interacting cells. However, this relation need not give rise to a *structure* relation like a tree as in P systems or a graph as in tissue P systems (e.g., see [15] for definitions of P systems and tissue P systems), though most models of membrane systems with a static membrane structure can be seen as special variants of NCs, and moreover, a lot of important features of membrane systems, in particular the derivation step and the halting condition, may be described at the level of NCs.

4 Systems with a Static Structure

In this section we consider networks of cells having a static structure, i.e., the number of cells does not change during the evolution of the system. We first define a transition step and then halting conditions.

Definition 4.1 Consider a network of cells $\Pi = (V, w_1, w_2, \ldots, w_n, R)$. A configuration of Π is an n-tuple of finite multisets $C = (u_1, \ldots, u_n)$ satisfying $inf_i \cap u_i = \emptyset$. The initial configuration of Π is defined as $C_0 = (w'_1, \ldots, w'_n)$ where w'_i , $1 \leq i \leq n$, are the finite multisets from $\langle V, \mathbb{N} \rangle$ with $f_{w'_i}(a) = 0$ for $f_{w_i}(a) = \infty$ and $f_{w'_i}(a) = f_{w_i}(a)$ for $f_{w_i}(a) < \infty$, $a \in V$.

Definition 4.2 We say that an interaction rule $r = (X \to Y; P, Q)$ is eligible for the configuration $C = (u_1, \ldots, u_n)$ if and only if for all $i, 1 \le i \le n$, we have

- $p_i \subseteq u_i \cup inf'_i$ (p_i is a submultiset of $u_i \cup inf'_i$),
- $f_i \not\subseteq u_i \cup inf'_i$ (f_i is not a submultiset of $u_i \cup inf'_i$), and
- $x_i \subseteq u_i \cup inf'_i$ (x_i is a submultiset of $u_i \cup inf'_i$).

Moreover, we require that $x_j \cap (V - inf_j) \neq \emptyset$ for at least one $j, 1 \leq j \leq n$. This last condition ensures that at least one symbol appearing only in a finite number of copies is involved in the rule. The set of all rules eligible for C is denoted by Eligible (Π, C) .

The marking algorithm.

Let $C = (w_1, \ldots, w_n)$ be a configuration of a network of cells Π and let R' be a finite multiset over M with M consisting of the (copies of) rules r_1, \ldots, r_k , where each $r_i = (X_i \to Y_i; P_i, Q_i) \in Eligible(\Pi, C), X_i = (x_{i,1}, \ldots, x_{i,n}), Y_i = (y_{i,1}, \ldots, y_{i,n}), 1 \leq i \leq k$. Moreover, set $x'_{i,j}, 1 \leq j \leq n$, to be the finite multisets from $\langle V, \mathbb{N} \rangle$ with $f_{x'_{i,j}}(a) = 0$ for $a \in inf_j$ and $f_{x'_{i,j}}(a) = f_{x_{i,j}}(a)$ for $a \notin inf_j$, $a \in V$, as well as $y'_{i,j}, 1 \leq j \leq n$, to be the finite multisets from $\langle V, \mathbb{N} \rangle$ with $f_{y'_{i,j}}(a) = 0$ for $a \in inf_j$ and $f_{y'_{i,j}}(a) = f_{y_{i,j}}(a)$ for $a \notin inf_j, a \in V$. Then:

- 1. consider a vector of multisets $Marked_0(\Pi, C, r_1, \ldots, r_k) = (\lambda, \ldots, \lambda)$ of size n and let i = 1;
- 2. if $X'_i \leq C Marked_{i-1}(\Pi, C, r_1, \ldots, r_k)$, then set

$$Marked_i (\Pi, C, r_1, \dots, r_k) = C - Marked_{i-1} (\Pi, C, r_1, \dots, r_k) - X'_i,$$

otherwise, end the algorithm and return false;

3. if i = k then end the algorithm and return **true**, otherwise set i to i + 1 and return to step 2.

If the marking algorithm returns **true** for the pair (C, R') then we say that the configuration C may be marked by R', and we define $Marked(\Pi, C, R') = Marked_k(\Pi, C, r_1, \ldots, r_k)$.

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Definition 4.3 Consider a configuration C and $R' \subseteq Eligible(\Pi, C)$ (i.e., a multiset of eligible rules). We say that the multiset of rules R' is applicable to C if the marking algorithm as described above returns **true** and the configuration Marked (Π, C, R') . The set of all multisets of rules applicable to C is denoted by Applicable (Π, C) .

Definition 4.4 Consider a configuration C and a multiset of rules $R' \in Applicable(\Pi, C)$. According to the marking algorithm described above, we define the configuration being the result of applying of R' to C as

 $Apply\left(\Pi, C, R'\right) = C - Marked\left(\Pi, C, R'\right) + \sum_{1 \le i \le k} Y'_{i}.$

We remark that Apply(R', C) is again a configuration.

For the specific *derivation modes* to be defined in the following, the selection of multisets of rules applicable to a configuration C may only be a specific subset of Applicable (Π, C) .

Definition 4.5 For the derivation mode ϑ , the selection of multisets of rules applicable to a configuration C is denoted by Applicable (Π, C, ϑ) .

Definition 4.6 For the asynchronous derivation mode (asyn),

Applicable $(\Pi, C, asyn) = Applicable (\Pi, C)$,

i.e., there are no particular restrictions on the multisets of rules applicable to C.

Definition 4.7 For the sequential derivation mode (sequ),

Applicable $(\Pi, C, sequ) = \{ R' \mid R' \in Applicable (\Pi, C) \text{ and } |R'| = 1 \},$

i.e., any multiset of rules $R' \in Applicable(\Pi, C, sequ)$ has size 1.

The most important derivation mode considered in the area of P systems from the beginning is the *maximally parallel* derivation mode where we only select multisets of rules R' that are not extensible, i.e., there is no other multiset of rules $R'' \supseteq R'$ applicable to C.

Definition 4.8 For the maximally parallel derivation mode (max),

$$Applicable (\Pi, C, max) = \{ R' \mid R' \in Applicable (\Pi, C) \text{ and there is} \\ no \ R'' \in Applicable (\Pi, C) \text{ with } R'' \supseteq R' \}.$$

A derivation mode closely related to the maximally parallel one, yet not considered so far in the literature is the following one, where we not only demand that the chosen multiset R' is not extensible, but also contains the maximal number of rules among all multisets from *Applicable* (Π, C, max): **Definition 4.9** For the maximal in rules maximally parallel derivation mode $(max_{rule}max)$,

$$\begin{aligned} Applicable\left(\Pi, C, max_{rule}max\right) &= \left\{R' \mid R' \in Applicable\left(\Pi, C, max\right) \text{ and } \\ & \text{there is no } R'' \in Applicable\left(\Pi, C, max\right) \\ & \text{with } |R''| > |R'| \right\}. \end{aligned}$$

In the minimally parallel derivation mode, we need an additional feature for the set of rules R, i.e., we consider a partition of R into disjoint subsets R_1 to R_h . For any set of rules $R' \subseteq R$, let ||R'|| denote the number of sets of rules R_j , $1 \leq j \leq h$, with $R_j \cap R' \neq \emptyset$.

There are several possible interpretations of this minimally parallel derivation mode which in an informal way can be described as applying multisets such that from every set R_j , $1 \leq j \leq h$, at least one rule – if possible – has to be used (e.g., see [5]). We start with the basic variant where in each derivation step we only choose a multiset of rules R' from Applicable $(\Pi, C, asyn)$ that cannot be extended to $R'' \in Applicable (\Pi, C, asyn)$ with $R'' \supseteq R'$ as well as $(R'' - R') \cap R_j \neq \emptyset$ and $R' \cap R_j = \emptyset$ for some j, $1 \leq j \leq h$, i.e., extended by a rule from a set of rules R_j from which no rule has been taken into R'.

Definition 4.10 For the minimally parallel derivation mode (min),

$$\begin{aligned} Applicable \left(\Pi, C, min\right) &= \left\{R' \mid R' \in Applicable \left(\Pi, C, asyn\right) \text{ and } \\ & \text{there is no } R'' \in Applicable \left(\Pi, C, asyn\right) \\ & \text{with } \left(R'' - R'\right) \cap R_j \neq \emptyset \\ & \text{and } R' \cap R_j = \emptyset \text{ for some } j, \ 1 \leq j \leq h \right\}. \end{aligned}$$

As in the case of the maximally parallel derivation mode, also for the minimally parallel derivation mode we may choose only multisets of rules with the maximal number of rules thus obtaining the maximal minimally parallel derivation mode:

Definition 4.11 For the maximal in rules minimally parallel derivation mode $(max_{rule}min)$,

$$\begin{aligned} Applicable\left(\Pi, C, max_{rule}min\right) &= \{R' \mid R' \in Applicable\left(\Pi, C, min\right) \text{ and} \\ & \text{there is no } R'' \in Applicable\left(\Pi, C, min\right) \\ & \text{with } |R''| > |R'| \}. \end{aligned}$$

In the case of the minimally parallel derivation mode, we have two more very interesting variants of possible interpretations, the first one maximizing the sets of rules involved in a multiset to be applied $(max_{set}min)$, and the second one demanding that all sets of rules that could contribute should contribute $(all_{aset}min)$:

Definition 4.12 For the maximal in sets minimally parallel derivation mode $(max_{set}min)$,

$$\begin{aligned} Applicable \left(\Pi, C, max_{set}min\right) &= \{R' \mid R' \in Applicable \left(\Pi, C, min\right) \text{ and} \\ & \text{there is no } R'' \in Applicable \left(\Pi, C, min\right) \\ & \text{with } \|R''\| > \|R''\| \} \,. \end{aligned}$$

Definition 4.13 For the using all applicable sets minimally parallel derivation mode $(all_{aset}min)$,

$$Applicable (\Pi, C, all_{aset}min) = \{R' \mid R' \in Applicable (\Pi, C, min) \text{ and } for all j, 1 \leq j \leq h, \\ R_j \cap Applicable (\Pi, C) \neq \emptyset \\ implies R_j \cap R' \neq \emptyset \}.$$

The ideas of taking only multisets of rules involving the maximal number of sets of rules and of taking only multisets of rules involving all sets of rules that can contribute now can also be taken over for the maximally parallel derivation mode:

Definition 4.14 For the maximal in sets maximally parallel derivation mode $(max_{set}max)$,

$$Applicable (\Pi, C, max_{set}max) = \{R' \mid R' \in Applicable (\Pi, C, max) \text{ and} \\ there is no \ R'' \in Applicable (\Pi, C, max) \\ with \ \|R''\| > \|R''\| \}.$$

Definition 4.15 For the using all applicable sets maximally parallel derivation mode $(all_{aset}max)$,

$$\begin{aligned} Applicable\left(\Pi, C, all_{aset}max\right) &= \{R' \mid R' \in Applicable\left(\Pi, C, max\right) \text{ and } \\ & \text{for all } j, \ 1 \leq j \leq h, \\ & R_j \cap Applicable\left(\Pi, C\right) \neq \emptyset \\ & \text{implies } R_j \cap R' \neq \emptyset \}. \end{aligned}$$

Finally, we should like to mention that the derivation modes $max_{set}X$ and $all_{aset}X$ with $X \in \{max, min\}$ could be extended by the constraint that a maximal number of rules has to be used, too, thus yielding derivation modes $max_{set}max_{rule}X$ and $all_{aset}max_{rule}X$ with $X \in \{max, min\}$. Demanding to use at least one rule from every set R_j , $1 \leq j \leq h$, would be another option, yet this case will be covered by the variant of partial halting defined in the succeeding subsection when being combined with derivation modes as $max_{set}X$ and $all_{aset}X$ with $X \in \{max, min\}$.

For all the derivation modes defined above, we now can define how to obtain a next configuration from a given one by applying an applicable multiset of rules according to the constraints of the underlying derivation mode:

Definition 4.16 Given a configuration C of Π and a derivation mode ϑ , we may choose a multiset of rules $R' \in Applicable(\Pi, C, \vartheta)$ in a non-deterministic way and apply it to C. The result of this transition step from the configuration C with applying R' is the configuration $Apply(\Pi, C, R')$, and we also write $C \Longrightarrow_{(\Pi, \vartheta)} C'$. The reflexive and transitive closure of the transition relation $\Longrightarrow_{(\Pi, \vartheta)}$ is denoted by $\Longrightarrow^*_{(\Pi, \vartheta)}$. There are several other derivation modes considered in the literature, e.g., we may apply (at most) k rules in parallel in every derivation step, but we leave the task to define such derivation modes in the general framework elaborated in this paper to the reader.

Definition 4.17 A configuration C is said to be accessible in Π with respect to the derivation mode ϑ if and only if $C_0 \Longrightarrow_{(\Pi,\vartheta)}^* C$ (C_0 is the initial configuration of Π). The set of all accessible configurations in Π is denoted by Accessible (Π).

Definition 4.18 A derivation mode ϑ is said to be deterministic (det- ϑ) if $|Applicable(\Pi, C, \vartheta)| = 1$ for any accessible configuration C.

4.1 Halting conditions

A halting condition is a predicate applied to an accessible configuration. The system halts according to the halting condition if this predicate is true for the current configuration. In such a general way, the notion halting with final state or signal halting can be defined as follows:

Definition 4.19 An accessible configuration C is said to fulfill the signal halting condition or final state halting condition (S) if and only if

 $S(\Pi, \vartheta) = \{ C' \mid C' \in Accessible(\Pi) \text{ and } State(\Pi, C', \vartheta) \}.$

Here $State(\Pi, C', \vartheta)$ means a decidable feature of the underlying configuration C', e.g., the occurrence of a specific symbol (signal) in a specific cell.

The most important halting condition used from the beginning in the P systems area is the *total halting*, usually simply considered as *halting*:

Definition 4.20 An accessible configuration C is said to fulfill the total halting condition (H) if and only if no multiset of rules can be applied to C with respect to the derivation mode anymore, i.e.,

 $H\left(\Pi,\vartheta\right) = \left\{C' \mid C' \in Accessible\left(\Pi\right) \text{ and } Applicable\left(\Pi,C',\vartheta\right) = \emptyset\right\}.$

The adult halting condition guarantees that we still can apply a multiset of rules to the underlying configuration, yet without changing it anymore:

Definition 4.21 An accessible configuration C is said to fulfill the adult halting condition (A) if and only if

$$\begin{split} A\left(\Pi,\vartheta\right) &= \{C' \mid C' \in Accessible\left(\Pi\right), \ Applicable\left(\Pi,C',\vartheta\right) \neq \emptyset \ and \\ Apply\left(\Pi,C',R'\right) &= C' \ for \ every \ R' \in Applicable\left(\Pi,C',\vartheta\right)\}. \end{split}$$

We should like to mention that we could also consider $A(\Pi, \vartheta) \cup H(\Pi, \vartheta)$ instead of $A(\Pi, \vartheta)$.

For introducing the notion of partial halting, we have to consider a partition of R into disjoint subsets R_1 to R_h as for the minimally parallel derivation mode. We then say that we are not halting only if there still is a multiset of rules R' from Applicable (Π, C) with $R' \cap R_j \neq \emptyset$ for all $j, 1 \leq j \leq h$:

Definition 4.22 An accessible configuration C is said to fulfill the partial halting condition (h) if and only if

$$h(\Pi, \vartheta) = \{C' \mid C' \in Accessible(\Pi) \text{ and there is} \\ no \ R' \in Applicable(\Pi, C') \text{ with} \\ R' \cap R_j \neq \emptyset \text{ for all } j, \ 1 \le j \le h\}.$$

4.2 Goal and result of a computation

The computations with a network of cells may have different goals, e.g., to generate (gen) a (vector of) non-negative integers in a specific output cell (membrane) or to accept (acc) a (vector of) non-negative integers placed in a specific input cell at the beginning of a computation. Moreover, the goal can also be to compute (com) an output from a given input or to output yes or no to decide (dec) a specific property of a given input.

The results not only can be taken as the number (N) of objects in a specified output cell, but, for example, also be taken modulo a terminal alphabet (T) or by subtracting a constant from the result (-k).

Such different tasks of a network of cells may require additional parameters when specifying its functioning, e.g., we may have to specify the output/input cell(s) or the terminal alphabet.

We shall not go into the details of such definitions here, we just mention that the goal of the computations $\gamma \in \{gen, acc, com, dec\}$ and the way to extract the results ρ are two other parameters to be specified and clearly defined when defining the functioning of a network of cells or a membrane system.

4.3 Taxonomy of networks of cells and (tissue) P systems

For a particular variant of networks of cells or especially P systems/tissue P systems we have to specify the derivation mode, the halting condition as well as the procedure how to get the result of a computation, but also the specific kind of rules that are used, especially some complexity parameters.

For networks of cells, we shall use the notation

 $O_m C_n \left(\vartheta, \phi, \gamma, \rho\right)$ [parameters for rules]

to denote the family of sets of vectors obtained by networks of cells $\Pi = (V, w_1, w_2, \ldots, w_n, R)$ of degree n with m = |V|, as well as ϑ, ϕ, ρ indicating the

derivation mode, the halting condition, and the way how to get results, respectively; the *parameters for rules* describe the specific features of the rules in R. If any of the parameters m and n is unbounded, we replace it by *.

For P systems, with the interaction between the cells in the rules of the corresponding network of cells allowing for a tree structure as underlying interaction graph, we shall use the notation

$$O_m P_n \left(\vartheta, \phi, \gamma, \rho \right)$$
 [parameters for rules].

Observe that usually the environment is not counted when specifying the number of membranes in P systems, but this usually hides that in many cases the environment takes an important role in the functioning of the system.

For tissue P systems, with the interaction between the cells in the rules of the corresponding network of cells allowing for a graph structure as underlying interaction graph, we shall use the notation

 $O_m t P_n (\vartheta, \phi, \gamma, \rho)$ [parameters for rules].

As a special example, let us now consider *symport/antiport P systems*.

A specific example: P systems with symport/antiport rules

For definitions and results concerning P systems with symport/antiport rules, we refer to the original paper [13] as well as to the overview given in [16]. An *antiport rule* is a rule of the form $((x, i) (u, j) \rightarrow (x, j) (u, i))$ usually written as $(x, out; u, in), xu \neq \lambda$, where j is the region outside the membrane i in the underlying graph structure. A symport rule is of the form $((x, i) \rightarrow (x, j))$ or $((u, j) \rightarrow (u, i))$.

The weight of the antiport rule (x, out; u, in) is defined as $\max \{|x|, |u|\}$. Using only antiport rules with weight k induces the type of rules α usually written as $anti_k$. The weight of a symport rule (x, out) or (u, in) is defined as |x| or |u|, respectively. Using only symport rules with weight k induces the type of rules α usually written as sym_k . If only antiport rules (x, out; u, in) of weight ≤ 2 and with $|x| + |u| \leq 3$ as well as symport rules of weight 1 are used, we shall write $anti_{2'}$.

As is well known,

$$O_*P_2(max, H, gen, N)[anti_{2'}] = NRE.$$

Observe that we only need one membrane separating the environment and the skin region, but this means that two regions corresponding to two cells are involved.

A general result

For any network of cells using rules of type α , with a derivation mode ϑ , $\vartheta \in \{all_{aset}min, asyn, sequ\}$, and partial halting, we only get Parikh sets of matrix languages (regular sets of non-negative integers):

Theorem 1. For every $\vartheta \in \{all_{aset}min, asyn, sequ\}, O_*C_*(\vartheta, h, gen, T)[\alpha] \subseteq PsMAT^{\lambda} and O_*C_*(\vartheta, h, gen, N)[\alpha] \subseteq NREG.$

The proof follows the ideas of a similar result proved for a general variant of P systems with permitting contexts in [1] and therefore is omitted. We should like to mention that this result is still valid if we take the derivation mode $max_{set}min$ instead of $all_{aset}min$ (because when using partial halting we always have to take at least one rule from every set of rules), yet we do not know whether it also holds for the derivation modes min and/or $max_{rule}min$.

5 Conclusions

The main purpose of this paper is to elaborate a general framework for static P systems and tissue P systems, but there are many variants of membrane systems not yet covered by this general framework, especially dynamic changes of the number of cells cannot be handled with the current version. Yet we have already started to extend our approach to such dynamic variants like P systems with active membranes. Moreover, also spiking neural P systems require some efforts for being captured within this framework. Our approach aims at formalizing the main features of membrane systems in such a way that derivation modes and halting conditions can be defined in a clear and unambiguous way to avoid that different interpretations of notions and concepts in the P systems area yield incomparable results (as a special example consider the variants described for the minimally parallel derivation mode). Moreover, specifying the marking algorithm in a procedural way should allow for easier and unambiguous implementations. Considering variants of (tissue) P systems at such a high level of abstraction allows for establishing quite general results.

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Conformon-P Systems with Negative Values

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Summary. Some initial results on the study of conformon-P systems with negative values are reported.

One model of these conformon-P systems is proved to be computationally universal while another is proved to be at least as powerful as partially blind program machines.

1 Introduction

The subdivision of a cell into compartments delimited by membranes inspired G. Păun to define a new class of (distributed and parallel) models of computation called *membrane systems* [8]. The hierarchical structure, the locality of interactions, the inherent parallelism, and also the capacity (in less basic models) for membrane division, represent the distinguishing hallmarks of membrane systems. Research on membrane systems, also called 'P systems' (where 'P' stays for 'Păun'), has really flourished [9].

One of the lines of research within membrane systems deals with the study of the generative power of models of these systems.

Recent results [3, 4] obtained with the use of Petri nets and P/T systems [10] show that the study of the generative variants of computing systems based on symbol objects (membrane systems, program machines, brane calculi, etc.) can be facilitated if someone considers the number of unbounded elements present in these systems. In the present paper we do not introduce the notation of Petri net and P/T systems but only one result obtained with their use. These information can be found in the just mentioned publications.

In particular [Corollary 2] from [4] indicates:

A P/T system with two unbounded elements has computational power equivalent to the one of program machines;

A P/T system with only unbounded number of tokens has computational power equivalent to the one of partially blind program machines;

A P/T system with only unbounded number of places has computational power equivalent to the one of restricted program machines (in this case restrictions in the composition of building blocks are present).

There unbounded elements refers to some components of the P/T systems (as, for instance, number of places and tokens) that are present in unbounded quantity. In [4] it is also proved that maximal parallelism is equivalent to the presence of an unbounded number of places. The results proved in [4] indicate that in the study of a computing system the number and kind of unbounded elements can give an indication (upper bounds and precise characterisation) of the computing power of the system.

The research reported in the present paper does not have the level of generality (i.e., the use of Petri nets) used in [4]. It refers to our initial results on the study of conformon-P systems having one 'extended' unbounded element: the value of the conformons ranges from $-\infty$ to $+\infty$, differently from previous studies in which it was ranging from 0 to $+\infty$.

2 Basic definitions

We assume the reader to have familiarity with basic concepts of formal language theory [6] and program machines [7]. We indicate with \mathbb{N} the set of positive integers, $\mathbb{N}_0 = \{0\} \cup \mathbb{N}$ and $\mathbb{Z} = \mathbb{N}_0 \cup \{-i \mid i \in \mathbb{N}\}$ indicates the set of all integers (positive, negative and zero).

2.1 Program machines

A program machine (also known as (multi)counter machines, multipushdown machines, register machines and counter automata) with n counters $(n \in \mathbb{N})$ is defined as $M = (S, R, s_0, s_d)$, where S is a finite set of states, $s_0, s_d \in S$ are respectively called the *initial* and *final* states, R is the finite set of instructions of the form (s_i, l_-, s_g, s_u) or (s_i, l_+, s_q) , with $s_i, s_g, s_u, s_q \in S$, $s_i \neq s_d, 1 \leq l \leq n$

A configuration of a program machine M with n counters is given by an element in the n + 1-tuples (s_j, \mathbb{N}_0^n) , $s_j \in S$. Given two configurations (s_i, l_1, \ldots, l_n) and $(s'_j, l'_1, \ldots, l'_n)$ we define a computational step as $(s_i, l_1, \ldots, l_n) \vdash (s_j, l'_1, \ldots, l'_n)$:

- if (s_i, l_-, s_g, s_u) , $l = l_p$ and $l_p \neq 0$, then $s_j = s_g$, $l'_p = l_p 1$, $l'_k = l_k$, $k \neq p$, $1 \leq k \leq n$; if $l = l_p$ and $l_p = 0$, then $s_j = s_u$, $l'_k = l_k$, $1 \leq k \leq n$;
- (informally: in state s_i if the content of counter l is greater than 0, then subtract 1 from that counter and change state into s_g, otherwise change state into s_u);
 if (s_i, l₊, s_q), l = l_p, then s_j = s_q, l'_p = l_p + 1, l'_k = l_k, k ≠ p, 1 ≤ k ≤ n;
 - (informally: in state s_i add 1 to counter l and change state into s_q).

The reflexive and transitive closure of \vdash is indicated by \vdash^* .

A computation is a finite sequence of transitions between configurations of a program machine M starting from the initial configuration (s_0, l_1, \ldots, l_n) with

 $l_1 \neq 0, \ l_k = 0, \ 2 \leq k \leq n$. If the last of such configurations has s_d as state, then we say that M accepted the number l_1 . The set of numbers accepted by Mis defined as $L(M) = \{l_1 \mid (s_0, l_1, \dots, l_n) \vdash^* (s_d, l'_1, \dots, l'_n)\}$. For every program machine it is possible to create another one accepting the same set of numbers and having all counters empty in the final state.

Partially blind program machines (also known as partially blind multicounter machines) were introduced in [5] and defined as program machines without test on zero. The only allowed operations are increase and decrease of one unit per time of the counters indicated as (s_i, l_+, s_q) and (s_i, l_-, s_g) respectively. In case the machine tries to subtract from a counter having value zero it stops in a non final state. In [5] it is also proved that such machines are strictly less powerful than non blind ones.

2.2 Conformon-P system with negative values

A conformon-P system with negative values has conformons, a name-value pair, as objects. If V is an alphabet (a finite set of letters), then we can define a conformon as $[\alpha, a]$, where $\alpha \in V$ and $a \in \mathbb{Z}$ (in our previous works on conformons, see for instance [1, 2], we considered $a \in \mathbb{N}_0$). We say that α is the name and a is the value of the conformon $[\alpha, a]$. If, for instance, $V = A, B, C, \ldots$, then [A, 5], [C, 0], [Z, -14] are conformons, while [AB, 21] and [D, 0.5] are not.

Two conformons can interact according to an *interaction rule*. An interaction rule is of the form $r: \alpha \xrightarrow{n} \beta$, where r is the label of the rule (a kind of name, it makes easier to refer to the rule) $\alpha, \beta \in V$ and $n \in \mathbb{N}_0$, and it says that a conformon with name α can give n from its value to the value of a conformon having name β . If, for instance, there are conformons [G, 5] and [R, 9] and the rule $r: G \xrightarrow{3} R$, one application of r leads to [G, 2] and [R, 12], another application of r (to [G, 2] and [R, 12]) leads to [G, -1] and [R, 15].

The compartments (membranes) present in a conformon-P system have a label (again, a kind of name which makes it easier to refer to a compartment), every label being different. Compartments can be unidirectionally connected to each other and for each connection there is a *predicate*. A predicate is an element of the set $\{\geq n, \leq n \mid n \in \mathbb{Z}\}$. Examples of predicates are: $\geq 5, \leq -2$, etc.

If, for instance, there are two compartments (with labels) m_1 and m_2 and there is a connection from m_1 to m_2 having predicate ≥ 4 , then conformons having value greater or equal to 4 can pass from m_1 to m_2 . In a time unit any number of conformons can move between two connected membranes as long as the predicate on the connection is satisfied. Notice that we have *unidirectional connections* that is: m_1 connected to m_2 does not imply that m_2 is connected to m_1 . Moreover, each connection has its own predicate. If, for instance, m_1 is connected to m_2 and m_2 is connected to m_1 , the two connections can have different predicates. It is possible to have multiple connections (with different predicates) between compartments.

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The interaction with another conformon and the passage to another membrane are the only *operations* that can be performed by a conformon.

Formally, a conformon-P system with negative values of degree $m, m \ge 1$, is a construct $\Pi = (V, \mu, \alpha_a, ack, L_1, \ldots, L_m, R_1, \ldots, R_m)$, where V is an alphabet; $\mu = (N, E)$ is a directed labelled graph underlying Π . The set N contains vertices (the membrane compartments), while the set E defines directed labelled edges (the connections) between vertices.

In α_a the value of α can either be *input* or *output*, in the former case Π is an accepting device, in the latter case Π is a generating device, while $a \in \{1, \ldots, m\}$ indicates the input or output membrane, respectively. $ack \in N$ indicates the *acknowledgment membrane*.

The multisets L_i contain conformons associated to region i; R_i are finite sets of rules for conformons interaction associated to region i.

A configuration of Π is an *m*-tuple indicating the multisets of conformons present in each membrane of the system. A *transition* is the passage from one configuration to another as the consequence of the application of operations.

A computation is a finite sequence of transitions between configurations of a system Π starting from (L_1, \ldots, L_m) , the *initial configuration* characterized by the fact that no conformon is present in the acknowledgment membrane. If used as a generating device, then the result of a computation is given by the multisets of conformons associated to membrane a when any conformon is associated to membrane ack. When this happens the computation is halted, that is no other operation is performed even if it could. When a conformon is associated to the acknowledge membrane the number of conformons (counted with their multiplicity) associated to membrane a defines the *number generated* by Π .

If used as an accepting device, then the input is given by the multiset of conformons associated to a in the initial configuration. If Π reaches a configuration with any conformon in ack, then no other operation is performed even if it could and Π accepts the input.

Some of the conformon-P systems considered in this paper work under *maximal parallelism*: in every configuration the maximum number of operations that can be performed is performed. If in one configuration some operations are conflicting (so that they cannot be executed together as they involve the same conformons), then any maximum number of non conflicting operations is performed. The passage of two conformons through the same connection is considered as two different operations (similarly for the interactions of two different pairs of conformons due to one rule).

2.3 Some modules for conformon-P systems

In the following we use the concept of *module*: a group of membranes with conformons and interaction rules in a conformon-P system able to perform a specific task. An example of module is a *splitter* [1]: a module that, when a conformon [X, x] with $x \in \{x_1, \ldots, x_h\}, x_i < x_{i+1}, 1 \le i \le h-1$ is associated with a specific membrane of it, it may pass such a conformon to other specific membranes according to its value x. A detailed splitter is depicted in Figure 1.a. Vertices outgoing a module representation of a splitter, Figure 1.b, have as predicates elements in the set $\{=n \mid \mathbb{N}_0\}$, this is a shorthand indicating the function performed by this module.



Fig. 1. A detailed splitter (a) and its module representation (b)

It should be clear that if a splitter is part of a conformon-P system with maximal parallelism, then the number of steps required to a conformon to pass from the u_{h+1} membrane to any other of the *u* membrane depends on the value of the conformon. If we consider the splitter depicted in Figure 1.a, a conformon present in membrane u_{h+1} requires only two steps to pass to membrane u_1 but it requires h+1 steps to pass to membrane u_h .

In order to have this time constant (equal to h+1 in the example), then delays (i.e., sequences of membranes) have to be introduced. We make this assumption for all the splitters considered in the proof of Theorem 1.

2.4 Figures in this paper

The representation of the conformon-P systems considered in this paper follows some rules aimed to a more concise representation an to an easier understanding of them.

The label of each membrane (a number) is indicated in **bold** on the top right corner of each compartment. Splitters are depicted by a thicker line, their label

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(also in **bold**) starts with **spl**, and their edges have '=' as predicate. The module representation of a splitter is depicted in Figure 1.b.

Oval compartments with a label inside are shortcuts for membranes or modules.

Conformons present in the initial configuration of the system are depicted in **bold**, the remaining conformons are the ones that could be present in the membrane during the computation.

The predicate associated to an edge is indicated close to the edge.

Some predicates and the value of some conformons contain a slash (/). This is a shorthand for multiple predicates or values. For instance, the conformon [A, 3/4]indicates that in a membrane the conformons [A, 3] and [A, 4] can be present. If there is a connection from membrane m_1 to membrane m_2 and the connection has predicate $\leq 0/\geq 5$, then this is equivalent to two connections from m_1 to m_2 , one with predicate ≤ 0 and the other with predicate ≥ 5 .

If the conformon [A, a] is present in m copies in a certain membrane, then this is indicated with ([A, a], m), where an unbounded number of copies is indicated with $+\infty$.

3 Results

Theorem 1. The class of numbers accepted by conformon-P systems with negative values and with maximal parallelism coincides with the one accepted by program machines.

Proof. This proofs follows from the one of [Theorem 2] from [1] (where priorities between interaction rules are present) and from [Theorem 1] from [4].

Figure 1 represents such a conformon-P system used as accepting device simulating a program machine. During this proof we refer to this figure.

For each state s_i of the simulated program machine there is a conformon with name s_i . For each instructions of the kind $(s_i, l_+, s_q) \in R$ there is a conformon with name $s'_{q,l}$; for each instruction of the kind $(s_i, l_-, s_g, s_u) \in R$ there are conformons with name $s''_{g,l}$ and $\bar{s}_{g,l}$. For the final state $s_d \in S$ there is one conformon with name $s''_{d'}$.

The initial configuration of the conformon-P system with priorities has all conformons with name $s'_{q,l}$, $s''_{g,l}$ and s'''_{d} and 0 as value in membrane 1; all the ones with name $\bar{s}_{g,l}$ and 1 as value in membrane 17; all the ones with name s_i and 0 as value in membrane 11 except the one with name of the initial state s_0 that is in membrane 1 with value 9 (in Figure 2 the generic conformon $[s_i, 9]$ is present in membrane 1); conformons [a, 8] and [c, 0] are initially present in membrane 6 and 13 respectively. Moreover for each counter l of the simulated machine there are an unbounded number of occurrences of the conformons [l, 0] in membrane 8, while the input membrane (membrane 14 in the figure) contains as many copies of such conformons as the values k_l of the counters at the initial configuration of the simulated machine.



Fig. 2. The conformons-P system related to Theorem 1

The addition of one unit to one counter l is simulated moving one occurrence of the conformon [l, 0] from membrane 8 to membrane 14; the subtraction of one unit is simulated with the passage of one occurrence of the same conformon from membrane 14 to membrane 8.

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For each instruction of the type $(s_i, l_+, s_q) \in R$ there is in membrane 1 the rule $s_i \xrightarrow{6} s'_{q,l}$; for each instruction of the kind $(s_i, l_-, s_g, s_u) \in R$, there is in membrane 1 the rule $s_i \xrightarrow{7} s''_{a,l}$; for $s_d \in S$ there is in membrane 1 the rule $s_i \xrightarrow{8} s''_d$.

Only one conformons of the kind $[s_i, 9]$ may be associated to membrane 1. When such a conformon is present in membrane 1, then one of the interaction rules indicate above can occur.

Let us consider now the case that the rule $s_i \xrightarrow{8} s_d''$ is applied. As there is only one instance of $[s_d^{\prime\prime\prime}, 0]$ then the newly created $[s_d^{\prime\prime\prime}, 8]$ passes to spl_1 and from here to membrane 6. After the interaction two sets of rules are applicable: one with a second interaction of s_i and s'''_d and another with the passage of $[s'''_d, 8]$ and $[s_i, 1]$ to spl_1 . Maximal parallelism forces this last set to be applied.

In membrane 6 $[s_d^{\prime\prime\prime}, 8]$ interacts with [a, 8] such that $[s_d^{\prime\prime\prime}, 10]$ is created. When this happens this conformon passes first to spl_6 and then to membrane 7, the acknowledgment membrane, halting in this way the computation.

It is important to notice that the presence of [a, 8] in membrane 6 is necessary for the halting of the computation. If in a configuration the conformon $[s''_{d'}, 8]$ passes in membrane 6 but [a, 8] is not there, then the simulation does not halt.

If instead the interaction in membrane 1 involves the conformon with name s_i and either $s'_{q,l}$ or $s''_{g,l}$ (due to either $(s_i, l_+, s_q) \in R$ or $(s_i, l_i, s_g, s_u) \in R$), then the following sets of applicable operations are.

- 1. the conformon with name s_i can interact again with the same instance of either
- $s'_{q,l}$ or $s''_{g,l}$; 2. if there is either $(s_i, l_+, s_{q'}) \in R$ or $(s_i, l_1, s_{g'}, s_{u'}) \in R$, then the conformon with name s_i can interact with an instance of either $s'_{q',l}$ or $s''_{q',l}$ and the conformon created in the previous interaction (either $[s'_{q,l}, 6]$ or $[s''_{g,l}, 7]$) can pass to spl_1 ;
- 3. both s_i and either $[s'_{q,l}, 6]$ or $[s''_{q,l}, 7]$ can pass to spl_1 .

Because of maximal parallelism only the second and third set of operations in the previous list can take place (as they contain two elements while the first set contains only one element).

If the second set occurs, then the system never reaches an halting configuration. This can be seen if, for instance, we consider the conformon $[s'_{a,l}, 6]$. Once in spl_1 this conformons passes to membrane 2, then to spl_2 and from here to membrane 6 where it interacts with [a, 8]. As a consequence of this interaction the conformon with name a passes to spl_6 so that the system does never halt.

The role of [a, 8] in membrane 6 is just this: if in any stage during the computation the system performed an operation that does not follow the simulation of the program machine, then a conformon passes to membrane 6 and interacts with [a, 8] making it unavailable for $[s'''_d, 8]$.

The creation of $[s_i, 3]$ and $[s'_{q,l}, 6]$ in membrane 1 starts the simulation of the instruction $(s_i, l_+, s_q) \in R$. As we said, the simulation of the addition of 1 to the value of the counter is performed with the passage of one instance of [l, 0] from membrane 8 to membrane 14. When in membrane 2 $[s_i, 3]$ and $[s'_{a,l}, 6]$ interact,

 $[s_i, -1]$ and $[s'_{q,l}, 10]$ are created and they pass to spl_2 (in case $[s'_{q,l}, 6]$ passes to spl_2 , then the system never halts). From spl_2 $[s_i, -1]$ and $[s'_{q,l}, 10]$ pass together to membrane 3 where they interact, $[s_i, 0]$ and $[s'_{q,l}, 9]$ are created and pass to spl_3 . From here $[s_i, 0]$ passes to membrane 11 while $[s'_{q,l}, 9]$ passes to membrane 8.

The membrane-splitter-membrane-splitter sequence that we just described (membrane 2 - spl_2 - membrane 3 - spl_3) is present in other parts of the system. This sequence allows to control the interaction of two conformons in a very precise way and to discard the outcome (i.e., conformons) of undesired interactions.

Once in membrane 8 the conformon with name $s'_{q,l}$ goes under another membrane-splitter sequence. In membrane 8 $[s'_{q,l}, 9]$ interacts with an instance of [l, 0]. After this interaction three sets of applicable operations are possible, this situation is similar to the one described before for membrane 1. In this case the undesired computation sees a conformons [l, 5] passing from spl_8 to membrane 6, while a proper simulation sees a conformon [l, 0] passing to membrane 14 and $[s'_{q,l}, 9]$ passing to membrane 11.

In membrane 11 $[s'_{q,l}, 9]$ interacts with $[s_q, 0]$ such that $[s'_{q,l}, 3]$ and $[s_q, 6]$ are created. Again a membrane-splitter sequence allows to create $[s'_{q,l}, 0]$ and $[s_q, 9]$ and let them pass to membrane 1. The instruction $(s_i, l_+, s_q) \in R$ has been performed and the system simulates the program machine being in state q.

The simulation of the instruction $(s_i, l_-, s_g, s_u) \in R$ starts with the interaction of $[s_i, 9]$ and $[s''_{g,l}, 0]$. If when this happens no [l, 0] conformon is present in membrane 14, then the conformon $[s_u, 9]$ passes to membrane 1, otherwise an occurrence of [l, 0] passes from membrane 14 to membrane 8 and the conformon $[s_g, 9]$ passes to membrane 1.

One interaction of $[s_i, 9]$ and $[s''_{g,l}, 0]$ in membrane 1 creates $[s_i, 2]$ and $[s''_{g,l}, 7]$ and, similarly to what described before, they can follow a membrane-splitter sequence at the end of which $[s_i, 2]$ is in membrane 11 and $[s''_{g,l}, 9]$ is in membrane 13.

In this last membrane $[s''_{g,l}, 9]$ interacts with [c, 0] so that $[s''_{g,l}, 7]$ and [c, 2] are created, then these two conformons pass to spl_{11} . From here [c, 2] passes to membrane 14. The conformon $[s''_{g,l}, 7]$ also passes to this membrane but only after two steps (in the meantime it goes in membrane 15 and 16).

If in membrane 14 there is at least an occurrence of [l, 0], then [c, 2] interacts with any of these so that [c, -3] and [l, 5] are created (at the same time $[s''_{g,l}, 7]$ pass to membrane 17). In this configurations a few things can happen. Similarly to the second and third set of operations indicated in the list above [c, -3] can either remain in membrane 14 and interact with another instance of [l, 0] (if present) or it can pass to spl_{12} and from here to membrane 15. In any case [l, 5] passes to spl_{13} and from here to membrane 15. If [c, -3] is not present in this membrane when [l, 5] is present, then this last conformon passes to spl_{14} and from here to membrane 6 (and here it interacts with [a, 8] such that the system never halts).

It should be clear now that if [c, -3] and [l, 5] do not move together out of membrane 14 the system never halts. If they do so, then they pass to membrane

15 at the same time. Here [l, 5] can either pass to spl_{14} (and then to membrane 6) or it can interact with [c, -3] so that [l, 0] and [c, 2] are created and they pass together to spl_{14} . From here [l, 0] passes to membrane 8 and [c, 2] to membrane 16 (where it waits until $[s''_{g,l}, 7]$ arrives).

When $[s_{g,l}', 7]$ passes to membrane 14 it can be that the conformon with name c is there or not. This last conformon can be in membrane 14 for two reasons: either no occurrence of [l, 0] was in that membrane, or one occurrence of [l, 0] was there and [c, -2] did not pass to spl_{12} . We know from the above that in this last case the system does not halt (because an [l, 2] is heading membrane 6), so we are not going to discuss the consequences of the interaction between $[s_{g,l}'', 7]$ and the conformon with name c when this last has a negative value.

If [c, 2] is present in membrane 14 when $[s''_{g,l}, 7]$ arrives there, too, then two things can happen: the two conformons interact or not. In this last case $[s''_{g,l}, 7]$ passes to spl_{13} and from here to membrane 16 and no operation can happen in the system. If instead the two conforms interact, then $[s''_{g,l}, 11]$ and [c, -1] are created and they pass to membrane 18 (through spl_{13} and spl_{12} , respectively). Here either [c, -2] passes to spl_{15} and no further operation is applied, or the two conformons interact so to create $[s''_{g,l}, 9]$ and [c, 0] and then these two conformons pass to membrane 11 and 13, respectively. So when $[s''_{g,l}, 9]$ is present in spl_{15} , then in the simulation the counter l was empty.

What happens to the conformon with name $s''_{g,l}$ in membrane 11 is similar to what discussed for the conformon with name $s'_{q,l}$ earlier on. The result of these operations is that $[s_g, 9]$ and $[s''_{g,l}, 0]$ are created and pass to membrane 1.

We still have to discuss the case in which no conformon with name c is present in membrane 14 when $[s''_{g,l}, 7]$ arrives. Here maximal parallelism forces this conformon to pass to spl_{13} and from here to membrane 16 where [c, 2] is also present. This means that if [c, 2] and $[s''_{g,l}, 7]$ are present in membrane 16, then the simulation of the subtraction of 1 from counter l has been performed.

When in membrane 16 [c, 2] and $[s''_{g,l}, 7]$ interact so that [c, 0] and $[s''_{g,l}, 9]$ are created and pass to membrane 13 and 17, respectively. In this last membrane $[s''_{g,l}, 9]$ interacts with $[\bar{s}_{g,l}, 1]$ so that $[s''_{g,l}, 0]$ and $[\bar{s}_{g,l}, 10]$ are created. Because of maximal parallelism these last two conformons pass to membrane 1 and 11, respectively.

What happens to the conformon with name $\bar{s}_{g,l}$ in membrane 11 is similar to what discussed for the conformon with name $s'_{g,l}$ earlier on. The result of these operations is that $[s_u, 9]$ and $[\bar{s}_{g,l}, 1]$ are created and pass to membrane 1 and 17 respectively.

If on a given input the program machine reaches an halting state, then the simulating conformon-P system can reach a final configuration.

The assumption that a program machine can simulate any such conform on-P system derives from the Turing-Church thesis. $\hfill\square$ In the figure related to the following proof some conformons have a parametric value of the kind a + bn, $a, b \in \mathbb{N}, n \geq 0$. This indicates all the possible values that a conformons can have as a consequence of interactions. If, for instance, the conformons [A, a], [C, c] and the interaction rule $C \xrightarrow{b} A$ are present in the same membrane, then the value of the A conformon can change into a + bn, where n indicates the number of interactions between the A and the C conformon.

Theorem 2. Conformon-P systems with negative values and without maximal parallelism can simulate partially blind program machines.

Proof. This proof follows from the one of [Theorem 1] from [1].

Figure 2 represents such a conformon-P system used as an accepting device simulating a program machine. During this proof we refer to this figure.



Fig. 3. The conformons-P system related to Theorem 2

For each state s_i of the simulated program machine there is a conformon with name s_i . For each instruction of the kind $(s_i, l_+, s_q) \in R$ there is a conformon with name $s'_{q,l}$; for each instruction of the kind $(s_i, l_-, s_g) \in R$ there is a conformon with name $s''_{q,l}$. For the final state $s_d \in S$ there is one conformon with name $s''_{d,l}$.

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The initial configuration of the conformon-P system has all conformons with name $s'_{a,l}$, $s''_{a,l}$ and s'''_{d} and 0 as value in membrane 1; all the ones with name s_i and 0 as value in membrane 8 except the one with name of the initial state s_0 that is in membrane 1 with value 7 (in Figure 3 the generic conformon $[s_i, 7]$ is present in membrane 1).

Moreover, for each counter l of the simulated machine there are an unbounded number of occurrences of the conformons [l, 0] in membrane 5, while the input membrane (membrane 4 in the figure) contains as many copies of such conformons as the values k_l of the counters at the initial configuration of the simulated machine. The addition of one unit to one counter l is simulated moving one occurrence of the conformon [l, 0] from membrane 5 to membrane 4; the subtraction of one unit is simulated with the passage of one occurrence of the same conformon from membrane 4 to membrane 5.

For each instruction of the type $(s_i, l_+, s_q) \in R$ there is in membrane 1 the rule $s_i \xrightarrow{6} s'_{a,l}$; for each instruction of the kind $(s_i, l_-, v) \in R$, there is in membrane 1 the rule $s_i \xrightarrow{5} s''_{g,l}$; for $s_d \in S$ there is in membrane 1 the rule $s_i \xrightarrow{4} s''_d$.

For any configuration of the conformon-P system only one conformon of the kind $[s_i, 7]$ may be associated to membrane 1. As we already said, initially this conformon is the one related to the initial state of the program machine.

When a conformon of the kind $[s_i, 7]$ is present in membrane 1, then one of the interaction rules indicate above can occur.

Let us consider now the case that the rule $s_i \stackrel{8}{\to} s_d''$ can be applied. Of course this rule can be applied more than once, if this happens the value of the s_i conformon goes below 0 and the one of the $s_d^{\prime\prime\prime}$ conformon is 4n. If n is at least 1, then the $s_d^{\prime\prime\prime}$ conformon can pass to spl_1 , but only if n = 1, then $[s_d^{\prime\prime\prime}, 4]$ passes to membrane 2, the acknowledgment membrane, halting in this way the computation.

This process of 'filtering out' (with splitters) conformons with an undesired value is present in many places in this conformon-P system. If two conformons over interacted, then the system never reaches an halting configuration.

If instead a rule of the kind $s_i \xrightarrow{6} s'_{q,r}$ is applied, then $[s'_{q,r}, 6+6n]$ can pass to spl_1 , but only $[s'_{q,r}, 6]$ can pass to membrane 5. If in membrane 1 $[s_i, 1]$ is produced, then it passes to membrane 3 (through spl_1).

In membrane 5 two things can happen:

- 1. $s'_{q,l}$ interacts several times with the same l conformon; 2. $s'_{q,l}$ interacts with different l conformons.

The only case such that $[s'_{a,l}, 1]$ is produced and passes to membrane 12 (through spl_3) is when $[s'_{a,l}, 6]$ interacts only once with one [l, 0] conformons. In all the other cases the value of the $s'_{q,l}$ conformon becomes negative and such a conformon does not pass to membrane 12 (in this way the system never halts).

If $[s'_{q,l}, 1]$ is produced, then also one [l, 5] is produced and, once in membrane 12, these two conformons interact so that $[s'_{q,l}, 6]$ and [l, 0] are recreated (because of spl_4 , an over interaction in membrane 12 leads the resulting conformons to remain in that splitter) and they can pass to membrane 3 and 4, respectively.

The passage of an instance of [l, 0] from membrane 5 to membrane 4 simulates the subtraction of one unit from the *l* counter. If no [l, 0] conformon is present in membrane 5 when a $s'_{q,l}$ conformon gets there, then the system never reaches an halting configuration.

In membrane 3 s_i and $s'_{q,l}$ can interact such that $[s_i, -n]$ and $[s'_{q,l}, 7+n], n \ge 0$ are produced and they pass to membrane 7 and 6 respectively. Only if n = 0(which means that only one interaction took place), then $[s_i, 0]$ and $[s'_{q,l}, 7]$ pass to membrane 8. Here $s'_{q,l}$ interacts with s_j and, in a way similar to what described until now, this can lead to the production of $[s_j, 7]$ and $[s'_{q,l}, 0]$ and these two conformons can pass to membrane 1.

The simulation of an instruction of the kind (s_i, l_-, v) is performed in a similar way.

If on a given input the partially blind program machine reaches an halting state, then the simulating conformon-P system can reach a final configuration. \Box

4 Final Remarks

The results reported in this paper leave us perplex. How shall we interpret these results in terms of unbounded elements [4]? Must the range of values (from $-\infty$ to $+\infty$) be regarded as one or two unbounded elements or as no unbounded elements at all? (because the two infinities 'cancel' each other)

In this last case [Corollary 2] from [4] is confirmed as Theorem 1 has two unbounded elements (number of conformons and maximal parallelism) and Lemma 2 only one (number of conformons). This also implies that it is possible to prove that a partially blind program machine can simulate any conformon-P system with negative values without maximal parallelism.

In case the range of values is regarded as one unbounded element, then it should be possible to prove that the computational power of conformon-P system with negative values is equivalent to the one of program machines. This implies that Theorem 1 is redundant (because it uses three unbounded elements).

In case the range of values is regarded as two unbounded elements, then some of the results reported in [4] should be extended and made more general.

We believe that the range of values should be regarded an one unbounded element.

Another problem on this line of research regards the characterization of blind program machines in terms of unbounded elements. *Blind program machines* (also known as *blind multicounter machines*) [5] are program machines that cannot sense (so neither halt or check) if the value of a counter is zero. It is know that these machines are strictly less computationally powerful then partially blind program machines.

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Optimizing Membrane System Implementation with Multisets and Evolution Rules Compression

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Summary. Nowadays, several research works on implementation of P systems have been focused on the massively parallel character of the model. In particular, it is possible to find out implementation of P system on cluster of processors, networks of processors, FPGA's ad hoc designed, and microcontrollers. Several published time analysis have proved that there is a very strong relationship between communication and evolution rules application time in membranes of the system. This relation determines the number of membranes that can be allocated per processor in order to obtain the minimum evolution time for the P system. This fact presents a problem related to hardware technologies which have a small amount of memory for storing multisets of objects and evolution rules, in particular microcontrollers.

The aim of this work is to present an algorithm for compressing information of multisets and evolution rules stored in membranes. In particular, this algorithm has to compress information, without penalizing evolution rules application and communication times with complex processes for compressing and decompressing such information. At the same time, keeping the same parallelism degree obtained in P systems implementation over distributed architectures presented in previous research works, but storing more membranes per processor.

1 Introduction

Membrane computing is a new computational model based on the membrane structure of living cells [14]. It must be stressed that they are not intended to model the functioning of biological membranes. Rather, they explore the computational character of several membranes features that can be used for modelling new computational paradigms inspired in Nature. This model has become, during last years, a powerful framework to develop new ideas in theoretical computation and to connect Biology with Computer Science.

The membrane structure of a P System is a hierarchical arrangement of membranes, embedded in a skin that separates the system from its environment. A membrane with no other membrane inside is called elementary. Each membrane defines a region that contains a multiset of objects, and a set of evolution rules. The objects are represented by symbols from a given alphabet. The objects can pass through membranes and membranes can change their permeability, they can be dissolved, or they can be divided. These features are used in defining transitions between system configurations, and sequences of transitions are used to define computations. Membrane systems are synchronous, in the sense that a global clock is assumed, i.e., the same clock holds for all regions of the system. At each time unit, a transformation of a system configuration takes place by applying rules in all regions, in a nondeterministic and maximally parallel manner.

Nowadays, membrane systems have been sufficiently characterized from a theoretical point of view. Their computational power has been settled – many variants are computationally complete. Among their most relevant characteristics appears the fact that they can solve non polynomial time problems in polynomial time, but this is achieved by the consumption of an exponential number of resources, in particular, the number of membranes that evolve in parallel.

An overview of membrane computing software can be found in [2] [15]. However, the way in which these models can be implemented is a persistent problem today, because "the next generation of simulators may be oriented to solve (at least partially) the problems of information storage and massive parallelism by using parallel language programming or by using multiprocessor computers" [2]. In this sense, information storage in membrane computation implementation is an example of Parkinson's First Law [13]: "storage and transmission requirements grow double than storage and transmission improvements".

This work focuses upon the problem due to the storage of multisets and rules information for the membranes in P-systems implementation. In particular, we pretend to get the highest compression level for the information without penalizing compression and decompression time with cost-consuming operations. Thus, we intend to reach a compression ratio that complies with the parallelism level reached in previous research works for P-systems implementation.

The structure of this work is as follows: first, related works in P-systems implementation are presented; next two sections present requirements for information compression in membrane systems and the proposed compression schema; then we analyze the obtained results for a set of tests for a well known P-system. Finally, authors present reached conclusions.

2 Related Works

First works over massively parallel implementation for P-systems started with Syropoulos [19] and Ciobanu [3] that in their distributed implementations of P systems use Java Remote Method Invocation (RMI) and the Message Passing Interface (MPI) respectively, on a cluster of PC connected by Ethernet. These authors do not carry out a detailed analysis about the importance of the time used during communication phase in the total time of P system evolution; although Ciobanu stated that "the response time of the program has been acceptable. There are however executions that could take a rather long time due to unexpected network congestion".

Recently, in [20] and [1] they present analysis for distributed architectures that are technology independent, based on: the allocation of several membranes in the same processor; the use of proxies for communication among processors; and, token passing in the communication. These solutions avoid communication collisions, and reduce the number and size of communication among membranes. All this allows to obtain a better step evolution time than in others suggested architectures congested quickly by the network collisions when the number of membranes grows up. Table 1 summarizes minimum times (T_{min}) , optimal amount of processors needed to implement the P system P_{opt} , membranes allocated at each one of them K_{opt} to reach those minimum times, the throughput obtained with corresponding processors Th_{proc} and communications Th_{com} for the architecture. These analysis consider the P-system number of membranes (M) that would evolve, the maximum time used by the slowest membrane in applying its rules (T_{apl}) , and the maximum time used by the slowest membrane for communication (T_{com}) .

Distributed Architecture [20]	Distributed Architecture [1]		
$T_{\rm min} = 2\sqrt{2 \ M \ T_{apl} \ T_{com}} - 2 \ T_{com}$	$T_{\min} = 2 \sqrt{M T_{apl} T_{com}} + T_{com}$		
$P_{opt} = \sqrt{\frac{M T_{apl}}{2 T_{com}}}$	$P_{opt} = \sqrt{\frac{M T_{apl}}{T_{com}}}$		
$K_{opt} = \sqrt{\frac{2 M T_{com}}{T_{apl}}}$	$K_{opt} = \sqrt{\frac{M T_{com}}{T_{apl}}}$		
$Th_{proc} \sim 50\%$	$Th_{proc} \sim 50\%$		
$Th_{com} \sim 50\%$	$Th_{com} \sim 100\%$		

Table 1. Distributed architecture parameters depending on application rules time (T_{apl}) , communication time (T_{com}) and number of membranes (M)

It may be concluded that to reach minimum times over distributed architectures, there should be a balance between the time dedicated to evolution rules application and the time used for communication among membranes. So, depending from the existing relation between both times, and from the number of membranes in the P-system, it is possible to determine the number of processors and the number of membranes that can be allocated at each one of them to obtain the evolution minimum time.

The difference between both architectures lies on the different topology for the processors net and the policy for token passing. Thus, [1] reaches a throughput near to a 100% of the communication line. With a 40% of increment in the parallelism level produced by the increment of the number of processors in the architecture, which permit to obtain a reduction over the 70% in the evolution time. Both works conclude that, for a specific number of membranes M, if it is possible that:

1. T_{apl} be N faster, the number of membranes that would be hosted in a processor would be multiplied by \sqrt{N} , the number of required processors would be

divided by the same factor and the time required to perform an evolution step would improve approximately with the same factor \sqrt{N} .

2. T_{com} be N faster, the number of required processors would be multiplied by \sqrt{N} , the number of membranes that would be hosted in a processor would be divided by the same factor and the time required to perform an evolution step would improve approximately by the same factor \sqrt{N} .

Table 2 summarizes the importance of reducing T_{apl} and T_{com} over the distributed architectures parameters (minimum evolution time, optimum number of processors and optimum number of membranes per processor).

Conditions	T_{\min}	P_{opt}	K_{opt}
T_{apl} be N faster and T_{com} be equal	$\frac{T_{\min}}{\sqrt{N}}$	$\frac{P_{opt}}{\sqrt{N}}$	$K_{opt} \cdot \sqrt{N}$
T_{apl} be equal and T_{com} be N' faster	$\frac{T_{\min}}{\sqrt{N'}}$	$P_{opt} \cdot \sqrt{N'}$	$\frac{K_{opt}}{\sqrt{N'}}$
T_{apl} be N faster and T_{com} be N' faster	$\frac{T_{\min}}{\sqrt{N'} \cdot \sqrt{N}}$	$\frac{P_{opt} \cdot \sqrt{N'}}{\sqrt{N}}$	$\frac{P_{opt} \cdot \sqrt{N}}{\sqrt{N'}}$

Table 2. Repercussion on distributed architecture parameters depending on T_{apl} and T_{com}

All the previous analysis for distributed architectures is independent of specific technology. In this sense, as it usually happens, implementation of these systems has been addressed from two different approaches: software and hardware models. The main research lines are: simulation over local networks using *cluster of microprocessors*, hardware *ad-hoc*, and generic hardware such as *microcontrollers*:

- 1. The hardware specifically designed has the possible advantage of being a massively parallel solution [17] [5] [12]. Their weak point resides in the lack of flexibility that presents, because this type of solutions only allows the simulation of a specific kind of membrane systems (for each membrane system a specific hardware is needed). They are also very enclosed solutions because they may be applied only to a very little range of problems (reduced number of objects in the alphabet and small number of evolution rules).
- 2. The solutions based on cluster of microprocessors and local networks have as main advantage the use of very common and well-known architectures. They are floppy systems also because a change at software level allows the simulation of any kind of membrane systems. Its main problem is that the best simulation times are reached always with few units, so the obtained solutions have a low degree of parallelism.
- 3. The solution, based on the use of microcontrollers [9] [10], seeks to be an intermediate point between hardware specifically designed and simulation using cluster of microprocessors. Microcontroller architecture is as flexible as a cluster of microprocessors and less enclosed and floppier than the hardware specifically designed. Microcontroller architecture has more level of potential parallelism than cluster of microprocessors but does not have intrinsic parallel

nature of the hardware specifically designed. Therefore, it is the cheapest architecture. But, in return, in [10] it is admitted that it is needed "the definition of more and more efficient data structures and algorithms oriented to minimize the quantity of memory required for the multisets and evolution rules storage that define the membrane".

Thus, in hardware solutions and solutions based upon microprocessors nets, the amount of information that has to be stored and transmitted is very important. In the first case, the main problem is due to their low storage capacity. So, reducing this amount of information needed to represent a membrane, means to be able to extend the variety of problems that can be solved with this technology. In the second case, reducing the amount of information to transmit means to minimize the bottleneck in processor communication and so, increase the parallelism level.

3 Compression Requirements

First, unlike other environments, where it is admissible a lossy information system (i.e. multimedia contents transmission), in our environment it is essential to have a lossless information compression system.

Almost all the compression methods require two phases: the first for analysis followed by a second one for conversion. First, an initial analysis of the information is done to identify repeated strings. From this analysis, an equivalences table is created to assign short codes to those strings. In a second phase, information is transformed using equivalent codes for repeated strings. Besides, this table is required with the information for its future compression/decompression. On the other hand, we must realize that a higher compression without any information loss will take more processing time. *Bitrate* is always variable and it is used mainly in text compression [18]. Because all of this, in spite of the fact that there are compression systems that are able to reach entropy limit - highest limit for data compression (i.e. Huffman codes) - they are not the ideal candidates for our system because of the following reasons:

- 1. Table storage will increase the needs for memory resources and would decrease compression goal.
- 2. Time for the phase of evolution rules application is penalized with compression/decompression processes when accessing compressed information on the P-system. This reduces parallelism level from distributed systems and increases evolution time.
- 3. And also, despite of the fact that communication phase time will be reduced because a lowest amount of information is transmitted, this will be counteracted by the time needed for decompression in the destination.

This way, as it is said in the goal of this work, compression schema for information from P-system membranes should accomplish following requirements:
- 1. there should be no information loss;
- 2. it should use the lowest amount of space for storage and transmission;
- 3. it should not penalize time for rules application phase and communication among membranes while processing compressed information. Thus, this means that the system should:
 - a) encode information for a direct manipulation in both phases without having to use coding/decoding processes.
 - b) do the compression in a previous stage to the P-system evolution
 - c) therefore, abandon entropy limit to be able to maintain parallelism level and evolution time reached in previous research works.

4 Compression Schema

This work pretends to compress the information from multisets that are present in regions and rules antecedents and consequents from each one of the P-system membranes. But it does not address the compression of another kind of information, such as priorities, membrane targets in rule consequents nor dissolving rule capability.

Representation for multisets information in related literature is Parikh's vector [4]. Data compression is very associated with its representation. Proposed compression schema is presented here in three consecutive steps beginning with Parikh's vector codification over the P-system alphabet. To show it, each of the successive steps will be applied over the following P-System [14]. See figure 1.



Fig. 1. A P System generating $n^2, n \ge 1$

4.1 Parikh's Vector over P System Alphabet

Each region of a membrane can potentially host an unlimited number of objects, represented by the symbols from a given alphabet V. We use V^* to denote the

set of all strings over the alphabet V (we consider only finite alphabets). For $a \in V$ and $x \in V^*$ we denote by $|x|_a$ the number of occurrences of a in x. Then, for $V = \{a_1, ..., a_n\}$, the Parikh vector associated with V is the mapping on V^* denoted by $\psi_V(x) = (|x|_{a_i}, \cdots, |x|_{a_n})$ for each $x \in V^*$. Our representation considers an order in the set of objects, hence the byte's order reflects the order of the objects within the alphabet and consequently, the position directly indicates which symbol's multiplicity is being stored.

Figure 2 shows an example for Parikh's vector associated to a multiset over the alphabet $V = \{a, b, c, d, e, f\}$.



Fig. 2. Example for Parikh's vector over the alphabet $V = \{a, b, c, d, e, f\}$

Figure 3 shows previous P-system information by the use of Parikh's vector over the alphabet $V = \{a, b, b', c, f\}$ for all the multisets that are present in each region and evolution rules for each membrane.

As we can see, codification using Parikh's vector over the P system alphabet requires 90 storage units for the multiplicities present in the multisets

4.2 Parikh's Vector for each Membrane's Alphabet

First step in compression considers only the alphabet subset for the P system that may exist in each of the regions for the membrane system, whatever are the possible configurations for the P system evolution. This subset may be calculated by a static analysis, previous to P system evolution time. In order to determine membranes alphabets in a given P system it is needed to consider the following facts:

- 1. Every object present at the region in the P system initial configuration belongs to its membrane's alphabet.
- 2. Every object present at the consequent in a membrane's evolution rule with target "here" belongs to its membrane's alphabet.
- 3. Every object present at the consequent in a membrane's evolution rule with target "in" to another membrane, belongs to the target membrane's alphabet.
- 4. Every object present at the consequent in a membrane's evolution rule with target "out", belongs to it's father alphabet.

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Fig. 3. A P System generating $n^2, n \ge 1$ representing multiset and evolution rules with Parikh's vector associated with V alphabet

5. Every object present at any membrane's alphabet with a evolution rule with dissolution capability belongs to it's father alphabet.

We will designate alphabet for a membrane $i, V_i \subseteq V$, where $\forall V_i, |V_i| \leq |V|$. For the example in figure 1, alphabets for the four membranes are: $V_1 = \{a, b, b', f\}$; $V_2 = \{a, b, b', f\}$; $V_3 = \{a, b, f\}$; and $V_4 = \{c\}$. According to these alphabets, each of the multisets in the region and the antecedents for a membrane evolution rules are codified by the Parikh's vector over its membrane alphabet. On the other hand, consequent multisets in each evolution rule are codified by the Parikh's vector over the target membrane alphabet.

Figure 4 shows previous P System information using Parikh's vector for each membrane alphabet.



Fig. 4. A P System generating $n^2, n \ge 1$ representing multiset and evolution rules with Parikh's vector associated with the membranes' alphabet

Now, P system codification by the use of Parikh's vector over each membrane specific alphabet, requires 63 storage units for present multiplicities in multisets. This codification reduces information size over a 70,0% concerning previous codification.

4.3 Parikh's vector without null values

Next compression step is an alteration over the *Run Length Encoding (RLE)* algorithm [11], used mainly to compress FAX transmissions. In this lossless codification, data sequences with same value (usually zeros) are stored as a unique value plus its count. RLE compression factor is, approximately:

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$$\frac{E\left(X\right)}{E\left\{\log_2 x\right\}}$$

where X is a discrete random variable that represents the number of successive zeros between two ones and E(X) is its expected value (average). Compression value stands between 20% and 30%.

In our case, what we pretend is to eliminate all the null values in Parikh's vector, that is, to eliminate all the references to the alphabet elements in a membrane that do not appear in its multiset. This information may be considered as redundant cause it may be obtained from the new coded information. In a formal way, let $V = \{a_1, a_2, \ldots, a_n\}$ be an ordered finite alphabet, $\forall x \in V *$, the encoded Parikh vector associated with V is defined by $\Psi_V^E(x) = \{(|x|_{a_i}, i) \mid |x|_{a_i} \neq 0\}$. Figure 5 shows an example for Parikh's vector without null values associated to a multiset over the ordered alphabet $V = \{a, b, c, d, e, f\}$.



Instance vector Parikh without null values

Fig. 5. Example for Parikh's vector without null values over the alphabet $V = \{a, b, c, d, e, f\}$

Concerning to multiplicities of objects present in multisets, there is two different situations: for the cases with multisets present at a membrane region, independently from the initial configuration, its multiplicities values are variable depending on the evolution that takes the membrane system in a non deterministic way; on the other hand, for the cases with multisets present at the evolution rules antecedents and consequents, its multiplicities values are constant and known previously to the P system evolution.

According to this situation, the evolution second step encodes without null values just the information that belongs to constant multisets present at evolution rules. Thus, we get a more compressed and lossless representation. The reason that does this representation possible is the fact that the absence of these null values multiplicities does not affect none of the multisets operations (addition, subtraction, applicability, scalar product, ...)

Figure 6 shows previous P system information using Parikh's vector without null values over each membrane's ordered alphabet.

Now, the P system codification using Parikh's vector without null values over each membrane's ordered alphabet requires 46 storage units for the multiplicities



Fig. 6. A P System generating $n^2, n \ge 1$ representing multiset and evolution rules with Parikh's vector associated with membranes alphabet and without null values

present at the multisets. This codification, in figure's 1 example, reduces information size until a 51,1% from the initial codification.

4.4 Storage Unit Compression

Last compression step concerns storage unit size for each of the P system information values. Depending on the storage unit size (measured in bits), we will be able to codify a greater or smaller range of values. In membrane computing, that does not allow negative values, given a size t bits for the storage unit, the range for possible values will vary from 0 to $2^t - 1$.

In this section, we will have to take into account separately multisets present at the membrane's regions in front of the ones present at evolution rules. For the first, storage unit size depends on the value range we want to reach during

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evolution while not having an overflow. Instead of it, for the second ones, we have to take into account, as it was shown in previous sections, that each membrane's ordered alphabet and their multiplicities are constant. Thus, an analysis previous to the P system evolution allows calculating the value ranges that are present in constant multisets for evolution rules and, so, the size that is needed to get their codification:

- 1. value range for multiplicities present at the antecedents and consecuent for each membrane (from 0 to 2 in example in figure 1; 2 bits needed for its codification).
- 2. value range for Parikh's vector positions over the ordered alphabet for each membrane (from 1 to 4 in example in figure 1; 2 bits needed for its codification).

Table 3 presents the size, in bits, needed to represent P system information from figure 1, with different storage unit sizes, using two different representations: the original representation (Parikh's vector over P system alphabet) and our compression schema (Parikh's vector without null values over membrane's alphabet). Last row shows compression rates obtained for different storage units sizes. In particular, it has been considered that the minimum storage unit size should be 8 bits according to actual technologies. In this work, we do not address the possibility of encoding several values at the same byte, which would increase more compression rates.

	Storage unit size			
Representational Schema	8 bits	16 bits	32 bits	64 bits
Parikh's vector for the P sys-	720 bits	1440 bits	2880 bits	5760 bits
tem alphabet				
Parikh's vector whithout null	368 bits	464 bits	656 bits	1040 bits
values associated with mem-				
brane alphabets				
Compression degree	51.1%	32.2%	22.8%	18.1%

Table 3. Size in bits for representing P system information with different size for storageunits and different representations

5 Analysis of Results

At this section we present the analysis of results obtained from the compression schema. First we analyze the schema compression itself. Afterwards we analyze the impact that compression has over the times for evolution rules application and communication among membranes. Finally, we analyze the global impact over distributed architectures parameters: evolution minimum time, optimum number of processors and membranes in each processor. For the analysis of the following sections, we will do a review over a test set composed by the P System published in [14] and [16]. Table 4 describes the considered P System set.

P System	Task	Reference
A.	First example	[14]
В.	Decidability: $n \mod k = 0$	[14]
С.	Generating: $n^2, n \ge 1(1^{st} \text{ version})$	[14]
D.	Generating: $n^2, n \ge 1$ (2 nd version)	[16]

Table 4. P System used for testing

5.1 Compression Schema Analysis

Table 5 shows compression rates reached for each P system from table 4, considering different storage unit sizes. Last row presents average compression rates for each storage size.

	Storage unit size			
P System	8 bits	16 bits	$32 \ bits$	64 bits
А.	59.8%	37.8%	26.8%	21.3%
В.	75.0%	47.7%	34.1%	27.3%
С.	51.1%	32.1%	22.8%	18.1%
D.	52.2%	33.3%	23.9%	19.2%
Average compression degree	59.5%	37.7%	26.9%	21.5%

Table 5. Compression degree for P System from Table 4

Considering the worst case for this compression schema (8 bits for all the storage units), at least, we reach a compression rate of 75,0%, which implies a increase of a 33,3% for memory availability to store information. For average compression rate (59,5%), it is reached an increase of 68,0% of memory availability. So we attenuate the storage problem for information in distributed architectures implemented with low storage capacity microcontrollers based technologies. Using this compression schema, it will be possible allocate more membranes in each microcontroller and so, it will be possible to reach minimum times at the same time that we are maximizing resources.

On the other hand, it has to be underlined that the compression process is done by an previous analysis to the P system evolution. Thus, evolution rules application and communication among membranes phases are not penalized with compression/decompression processes.

5.2 Impact Analysis for Evolution Rules Application Time

Published parallel and sequential algorithms [6] [7] [21] [22] [8] for evolution rules application are based upon a limited set of multisets primitive operations. These are calculation of: applicability, maximum applicability, antecedent/consequent addition and subtraction over its region multiset and the scalar product of a antecedent/consequent.

Algorithmic complexity of any of these operations is determined by the representation of multiset information present at the evolution rules. In worst case, using representation trough Parikh's vector over the P system alphabet, complexity will be equal to its alphabet cardinal. On the other hand, using representation through the proposed compression schema, complexity in worst case will be equal to the multiset support that is present at the evolution rule antecedent/consequent. Table 6 presents, for each of the P system in table 4, its alphabet support, the average support for multisets present in its evolution rules and a percentage based relation among both cardinals. Last row presents these cardinals average values and their relation.

P System	$\mid V \mid$	support(w)	%
А.	4	1.05	26.3%
В.	4	1.50	37.5%
С.	5	1.13	22.6%
D.	5	1.13	22.6%
Average	4.5	1.20	26.7%

Table 6. Alphabet Cardinality and support average from P systems of Table 4

According to these empirical values, each of the primitive operations previously mentioned will decrease its execution time approximately until a 26,7%. And consequently, evolution rules application time will be approximately 3,75 times faster.

5.3 Impact Analysis for Communication among Membranes Time

Communication among membranes addresses submission of multisets present at the applied application rules consequents and, in case of dissolution, the region multiset itself. Depending on information representation, the data packet size to transmit will be smaller or bigger. Table 7 shows, for each of the P systems shown in table 4, information compression rate for its communications for different storage units sizes. Last row presents compression rates average.

According to these empirical values, a reduction until a 55,6% of the information to transmit among membranes may be reached in the worst case. Considering that communication is a linear process that depends upon the amount of information to transmit, communication time among membranes will be approximately 1,80 times faster.

	Storage unit size			
P System	8 bits	16 bits	$32 \ bits$	64 bits
А.	55.0%	45.0%	40.0%	37.5%
В.	60.0%	50.0%	45.0%	42.5%
С.	54.0%	44.0%	39.0%	36.5%
D.	53.3%	44.4%	40.0%	37.8%
Average	55.6%	45.9%	41.0%	$\mathbf{38.6\%}$

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Table 7. Compression degree for communication units from P systems of Table 4

5.4 Global Impact Analysis

Finally, we present a impact analysis of this compression schema over distributed architecture parameters. In particular, we examine, following criteria shown in table 2, implication in optimum number of processors and membranes per processor and minimum evolution time.

On one side, time reduction for evolution rules application increases the number of membranes per processor. It also decreases the number of processors and evolution time. According to the previous empirical data, from a rule application time 3,75 times faster, we get an increment of a 93,5% for membranes per processor, a reduction until a 51,6% for number of processors and for evolution time.

On the other hand, time reduction for communication among membranes increases the number of processors. It also decreases the number of membranes per processor and evolution time. According to the previous empirical data, from a communication time 1,80 times faster, we get, for the worst case, a 34,2% increment for number of processors and a reduction until a 74,5% for the number of membranes per processor and for evolution time.

Taking in account both factors, reduction for application and communication time, counteract their effects over the number of processors and the number of membranes per processor. According to the previous empirical data, we get a reduction of a 69,3% for the number of processors, an increment of a 44,3% for the number of membranes per processor and a reduction until a 38,5% for evolution time.

6 Conclusions

In this work has been presented a schema for compressing multisets and evolution rules for P system membranes. This compressing schema is a variant from Run Length Encoding starting form the Parikh's vector considering the specific membrane alphabet. Empirical results over a reduced set of classical P systems show degrees of compression varying from 51.1 % to 18.1% depending on the size in bits needed for storing objects multiplicities for multisisets of objects in membranes of the P systems. This is a way for allocating more membranes per processor in

the implementation of P system on distributed architectures having a low memory capability.

On the other hand, the compression schema does not penalize evolution rule application nor communication times during P system evolution. The schema does not required compression/decompression process during P system evolution. The whole compression process is performed by mean of a static analysis previous to the P system evolution. These facts, thanks to the representation of information established, improve the system performance reducing evolution rule application and communication times, what is very important because it implies a direct implication on reducing the evolution time of the membrane systems.

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Hill Kinetics Meets P Systems: A Case Study on Gene Regulatory Networks as Computing Agents *in silico* and *in vivo*

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Summary. Modelling and simulation of biological reaction networks is an essential task in systems biology aiming at formalisation, understanding, and prediction of processes in living organisms. Currently, a variety of modelling approaches for specific purposes coexists. P systems form such an approach which owing to its algebraic nature opens growing fields of application. Here, emulating the dynamical system behaviour based on reaction kinetics is of particular interest to explore network functions. We demonstrate a transformation of Hill kinetics for gene regulatory networks (GRNs) into the P systems framework. Examples address the switching dynamics of GRNs acting as inverter, NAND gate, and RS flip-flop. An adapted study *in vivo* experimentally verifies both practicability for computational units and validity of the system model.

1 Introduction

Along with the development of systems biology, a variety of modelling techniques for biological reaction networks have been established during the last years [1]. Inspired by different methodologies, three fundamental concepts emerged mostly independent of each other: *analytic, stochastic,* and *algebraic* approaches. Each paradigm specifically emphasises certain modelling aspects. Analytic approaches, primarily adopted from chemical reaction kinetics, enable a macroscopic view on species concentrations in many-body systems. Based on differential equations considering generation and consumption rates of species, deterministic monitoring and prediction of temporal or spatial system behaviour is efficiently expressed by continuous average concentration gradients. In contrast, stochastic approaches reflect aspects of uncertainty in biological reaction networks by incorporating randomness and probabilities. So, ranges of possible scenarios and their statistical distri-



Fig. 1. Modelling approaches for biological reaction networks and bridges between them. Algorithmic strategies behind these bridges allow model transformations. Stochastic, analytic, and algebraic approaches form fundamental paradigms, categorisable into subclasses (white highlighted) and transformational concepts (black highlighted)

bution can be studied facilitating a direct comparison with wetlab experimental data. Statistical tools help in discovering correlations between network components. Furthermore, algebraic approaches appear as flexible instruments regarding the level of abstraction for system description. Due to their discrete principle of operation, they work by embedding as well as evaluating structural information, modularisation, molecular tracing, and hierarchical graduation of provided system information.

Combining advantages of several paradigms comes more and more into the focus of research. On the one hand, heterogeneous models subsume elements from different approaches into an extended framework. On the other hand, transformation strategies aim to model shifting between approaches, see Figure 1. Thus, specific analysis tools as well as advanced techniques for classification, simplification, comparison, and unification can become applicable more easily. This is additionally motivated by the fact that all three paradigms are independently known to be capable of constructing Turing complete models for computation [14].

In general, P systems represent term rewriting mechanisms, hence algebraic constructs [19, 20]. Substantiated by the progress in proteomics, investigating the

dynamical behaviour of biological reaction networks is essential to understand their function. Although P systems containing appropriate kinetics are useful, reaction kinetics is mostly defined for analytic models. In this paper, we contribute to bridging this gap for GRNs.

Related work addresses corresponding P systems for phenotypic representations of some biological network classes. While metabolic P systems [15] and P systems for cell signalling [11, 18] have already been equipped with mass-action kinetics derived from underlying reaction mechanism [7], P systems for GRNs [2, 5] and for quorum sensing [3] are restricted to formulate inhibiting or activating effects qualitatively. In order to introduce a homogeneous quantitative model, we decided to incorporate Hill kinetics [16] to the P systems framework by describing the cooperativity in GRNs dynamically using sigmoid-shaped transfer functions that are more precise than two-stage on/off switching.

The paper is organised as follows: Based on the definition of Hill kinetics, we present a method for discretisation that leads to P systems Π_{Hill} whose properties are discussed briefly. A case study includes GRNs acting as inverter, NAND gate, and RS flip-flop. For each logic gate, its GRN in concert with ODEs derived from Hill kinetics, corresponding P system, and simulation results are shown. Finally, we verify that a reporter gene encoding the green fluorescent protein (gfp) with transcription factors N-acyl homoserine lactone (AHL) and isopropyl- β D-thiogalactopyranoside (IPTG) can mimic the aforementioned RS flip-flop *in vivo*. Here, gfp expression is quantified using flow cytometry.

2 Transforming Hill Kinetics to P System

Hill Kinetics

Hill kinetics [16] represents a homogeneous analytic approach to model cooperative and competitive aspects of interacting biochemical reaction networks dynamically. It formulates the relative intensity of gene regulations by sigmoid-shaped threshold functions h of degree $m \in \mathbb{N}_+$ and threshold $\Theta > 0$ such that $x \ge 0$ specifies the concentration level of a transcription factor that activates resp. inhibits gene expression. Function value h then returns the normalised change in concentration level of the corresponding gene product:

activation (upregulation) \rightarrow : $h^+(x, \Theta, m) = \frac{x^m}{x^m + \Theta^m}$

inhibition (downregulation) \perp : $h^{-}(x, \Theta, m) = 1 - h^{+}(x, \Theta, m)$

Functions h^+ and h^- together with a proportional factor c_1 quantify the production rate of a certain gene product *GeneProduct*. Here we assume a linear spontaneous decay with rate $c_2[GeneProduct]$ such that the differential equation takes the form $\frac{d[GeneProduct]}{dt} = ProductionRate - c_2[GeneProduct]$. Different activation and inhibition rates are simply multiplied as in the following example illustrated in Figure 2 ($c_1, c_2 \in \mathbb{R}_+$):



Fig. 2. Gene regulatory unit. Repetitive expression of a *Gene* leads to generation of a specific *GeneProduct*, a protein whose amino acid sequence is encoded by the DNA sequence of the *Gene*. Transcription factors (specific single proteins or complexes) quantitatively control the expression rate by their present concentration. Two types of transcription factors can be distinguished: Inhibitors, here symbolised by I_1, \ldots, I_p , repress *Gene* expression by downregulation while activators A_1, \ldots, A_n cause the opposite amplifying effect by upregulation.

$$\frac{\mathrm{d}\left[GeneProduct\right]}{\mathrm{d}t} = c_1 \cdot \mathrm{h}^+(A_1, \Theta_{A_1}, m) \cdot \ldots \cdot \mathrm{h}^+(A_n, \Theta_{A_n}, m) \cdot \left(1 - \mathrm{h}^+(I_1, \Theta_{I_1}, m) \cdot \ldots \cdot \mathrm{h}^+(I_p, \Theta_{I_p}, m)\right) - c_2[GeneProduct]$$

For simplicity, each differential coefficient $\frac{d y}{d t}$ is subsequently denoted as \dot{y} .

By coupling gene regulatory units we obtain GRNs. Here, gene products can act as transcription factors for other genes within the network. Additional complex formation among gene products allows conjunctive composition of transcription factors and the introduction of further nonlinearities. Thus, an effective signal transduction and combination between different network elements becomes feasible.

Discretisation

The analytic nature of Hill kinetics based on continuous concentrations requires a discretisation with respect to value and time in order to derive a homologous term rewriting mechanism. Following the intention to approximate continuous concentrations by absolute particle numbers, we assume a large but finite pool of molecules. The application of a reaction rule in terms of a rewriting process removes a number of reactant particles from this pool and simultaneously adds all products. Therefore, selection and priorisation of reaction rules to apply are controlled by an underlying iteration scheme with temporally stepwise operation.

Since Hill kinetics is characterised by variable reaction rates due to the sigmoidshaped functions h, this variability should also be reflected in the term rewriting mechanism. For this reason, we introduce dynamic stoichiometric factors resulting in time dependent reaction rules. Let $\Delta \tau > 0$ be the constant time discretisation interval (step length), the gene regulatory unit depicted in Figure 2 consists of two reaction rules with variable stoichiometric factors s and u:

$$s \ Gene \longrightarrow s \ GeneProduct + s \ Gene \Big|_{A_1, \dots, A_n, \neg I_1, \dots, \neg I_p} \quad \text{where} \\ s = \lfloor \Delta \tau \cdot c_1 \cdot [Gene] \cdot \\ h^+(A_1, \Theta_{A_1}, m) \cdot \dots \cdot h^+(A_n, \Theta_{A_n}, m) \cdot \\ (1 - h^+(I_1, \Theta_{I_1}, m) \cdot \dots \cdot h^+(I_p, \Theta_{I_p}, m)) \rfloor \\ u \ GeneProduct \longrightarrow \emptyset \quad \text{where} \quad u = \lfloor \Delta \tau \cdot c_2 \cdot [GeneProduct] \rfloor$$

Here, the upper reaction formulates the generation of *GeneProduct* particles with regard to the limiting resource of available *Gene* objects. Reaction conditions coming from the presence of activators A_1, \ldots, A_n and absence (\neg) of inhibitors I_1, \ldots, I_p affect the stoichiometric factor s. The notation of indexes after the vertical bar declares the elements which occur in the h-components (h^+, h^-) of the function regulating the rule. In order to map normalised concentrations from Hill kinetics into absolute particle numbers, we introduce the factor term [*Gene*] which represents the total number of *Gene* objects present in the reaction system. Accordingly, the decay (consumption) of *GeneProduct* is expressed by the lower

The rounding regulation $(\lfloor \rfloor)$ provides for integer numbers as stoichiometric factors. This is necessary for handling the discrete manner of term rewriting. Nevertheless, a discretisation error can occur and propagate over the time course. The higher the total number of particles in the reaction system is initially set, the more this inaccuracy can be reduced.

Now, we incorporate the reaction system obtained by discretisation into the P systems framework. Therefore, we firstly define some syntactical conventions with respect to formal languages and multisets.

Formal Language and Multiset Prerequisites

transition rule.

We denote the empty word by ε . Let A be an arbitrary set and \mathbb{N} the set of natural numbers including zero. A multiset over A is a mapping $F: A \longrightarrow \mathbb{N} \cup \{\infty\}$. F(a), also denoted as $[a]_F$, specifies the multiplicity of $a \in A$ in F. Multisets can be written as an elementwise enumeration of the form $\{(a_1, F(a_1)), (a_2, F(a_2)), \ldots\}$ since $\forall (a, b_1), (a, b_2) \in F : b_1 = b_2$. The support $\operatorname{supp}(F) \subseteq A$ of F is defined by $\operatorname{supp}(F) = \{a \in A \mid F(a) > 0\}$. A multiset F over A is said to be empty iff $\forall a \in A : F(a) = 0$. The cardinality |F| of F over A is said to be empty iff $\forall a \in A : F(a) = 0$. The cardinality |F| of F over A is $|F| = \sum_{a \in A} F(a)$. Let F_1 and F_2 be multisets over A. F_1 is a subset of F_2 , denoted as $F_1 \subseteq F_2$, iff $\forall a \in A : (F_1(a) \leq F_2(a))$. Multisets F_1 and F_2 are equal iff $F_1 \subseteq F_2 \wedge F_2 \subseteq F_1$. The intersection $F_1 \cap F_2 = \{(a, F(a)) \mid a \in A \wedge F(a) = \min(F_1(a), F_2(a))\}$, the multiset sum $F_1 \uplus F_2 = \{(a, F(a)) \mid a \in A \wedge F(a) = \max(F_1(a) - F_2(a), 0)\}$ form multiset operations. The term $\langle A \rangle = \{F: A \longrightarrow \mathbb{N} \cup \{\infty\}\}$ describes the set of all multisets over A while $\mathcal{P}(A)$ denotes the power set of A.

Transformation: Definition of the Corresponding P System

The general form of a P system $\varPi_{\rm Hill}$ emulating the dynamical behaviour of GRNs using Hill kinetics is a construct

$$\Pi_{\text{Hill}} = (V_{\text{Genes}}, V_{\text{GeneProducts}}, \Sigma, [1]_1, L_0, r_1, \dots, r_k, f_1, \dots, f_k, \Delta\tau, m)$$

where V_{Genes} denotes the alphabet of genes, $V_{\text{GeneProducts}}$ the alphabet of gene products (without loss of generality $V_{\text{Genes}} \cap V_{\text{GeneProducts}} = \emptyset$), and $\Sigma \subseteq V_{\text{GeneProducts}}$ represents the output alphabet. Π_{Hill} does not incorporate inner membranes, so the only membrane is the skin membrane $[_1]_1$. The single membrane property results from the spatial globality of GRNs within an organism: Gene expression is located in the cell nuclei flanked by a receptor-controlled intercellular transduction and combination of transcription factors. Resulting GRNs form independent network structures of high stability within living organisms.

Let $V = V_{\text{Genes}} \cup V_{\text{GeneProducts}}$. The multiset $L_0 \in \langle V \rangle$ over V holds the initial configuration of the system.

Initial reaction rules³ $r_i \in \langle E_{i,0} \rangle \times \langle P_{i,0} \rangle \times \mathcal{P}(TF_i)$ with multiset of reactants $E_{i,0} \subseteq V \times \mathbb{N}$, multiset of products $P_{i,0} \subseteq V \times \mathbb{N}$ and set of involved transcription factors $TF_i \in V_{\text{GeneProducts}}$, $i = 1, \ldots, k$, define the potential system activity at time point 0. A function $f_i : \mathbb{R}_+ \times \langle V \rangle \times \mathbb{N}_+ \to \mathbb{N}$ is associated with each initial reaction rule r_i . This function adapts the stoichiometric factors according to the discretised Hill kinetics as described above.

Furthermore, we introduce two global parameters. The time discretisation interval $\Delta \tau \in \mathbb{R}_+$ corresponds to the length of a time step between discrete time points t and t + 1. The degree $m \in \mathbb{N}_+$ is used for all embedded sigmoid-shaped functions.

Finally, the dynamical behaviour of the P system is specified by an iteration scheme updating both the system configuration L_t and the stoichiometric factors of reaction rules r_i starting from L_0 where $i = 1, \ldots, k$:

$$L_{t+1} = L_t \ominus Reactants_t \uplus Products_t \quad \text{with}$$

$$Reactants_t = \biguplus_{i=1}^k (E_{i,t+1} \cap L_t)$$

$$Products_t = \biguplus_{i=1}^k (P_{i,t+1} \cap L_t)$$

$$E_{i,t+1} = \{(e, a') \mid (e, a) \in E_{i,t} \land a' = f_i(\Delta \tau, L_t, m)\}$$

$$P_{i,t+1} = \{(q, b') \mid (q, b) \in P_{i,t} \land b' = f_i(\Delta \tau, L_t, m)\}$$
(1)

Informally, the specification of $E_{i,t+1}$ and $P_{i,t+1}$ means that all reactants e and products q remain unchanged over the time course. Just their stoichiometric factors are updated from value a to a' (reactants) and from b to b' (products) according

³ Note that in our case the stoichiometry of reaction rules changes over time which is used to implement time-varying reaction rates.

to functions f_i . These functions may utilise the numbers of copies for all |V| types of particles recently present in the system. The cardinality $|L_t \cap \{(w_j, \infty)\}|$ then identifies this amount for any $w_j \in V$.

In terms of computational devices, P systems Π_{Hill} carry an output providing the outcome of a calculation. For this purpose, the multiplicity of those gene products listed in the output alphabet is suitable. We define an output function output : $\mathbb{N} \to \mathbb{N}$ by

$$\operatorname{output}(t) = |L_t \cap \{(w, \infty) \mid w \in \Sigma\}|.$$

For better readability, we subsequently write a reaction rule $r_i = (\{(e_1, a_1), \ldots, (e_h, a_h)\}, \{(q_1, b_1), \ldots, (q_v, b_v)\}, \{tf_1, \ldots, tf_c\})$ with $\operatorname{supp}(E_{i,t}) = \{e_1, \ldots, e_h\}$ and $\operatorname{supp}(P_{i,t}) = \{q_1, \ldots, q_v\}$ as well as $TF_i = \{tf_1, \ldots, tf_c\}$ by using the chemical denotation $r_i : a_1 e_1 + \ldots + a_h e_h \longrightarrow b_1 q_1 + \ldots + b_v q_v|_{tf_1, \ldots, tf_c}$. As a first example, Π_{Hill} of the gene regulatory unit shown in Figure 2 reads:

Note that s_1 at time point t + 1 is equal to $f_1(\Delta \tau, L_t, m)$ at time point t or holds its initialisation value at time point 0. Respectively, s_2 at time point t + 1 is equal to $f_2(\Delta \tau, L_t, m)$ at time point t or holds its initialisation value at time point 0, see equations (1) and (2).

At low molecular concentrations, deterministic application of Hill functions can conflict between different functions which want to update the system configuration. This is the case if the amount of reactants is too small to satisfy the needs of all functions. Since the number of multiset elements always remains nonnegative (see definition of \ominus), the system can violate mass conservation by satisfying these needs.

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A system extension based on stochastic rewriting mechanisms can overcome this insufficiency.

System Classification, Properties and Universality

 $\Pi_{\rm Hill}$ belongs to P systems with symbol objects and time varying transition rules whose evolution is based on conditional rewriting by quantitative usage of promoters and inhibitors. Thus, the dynamical behaviour formulated in Hill kinetics is time- and value-discretely approximated by a stepwise adaptation. This leads to a deterministic principle of operation.

From the view on computational completeness, there are several indicators for Turing universality. On the one hand, we will demonstrate within the next section how NAND gates and compositions of NAND gates can be emulated by P systems of the form Π_{Hill} . Arbitrarily extendable circuits consisting of coupled NAND gates can be seen as computational complete [14]. On the other hand, the multiplicity of each symbol object within the system may range through the whole recursively enumerable set of natural numbers. So, copies of a gene product expressed by a dedicated gene are able to represent the register value of a random access machine. Autoactivation loops keep a register at a certain value while external activation increases the amount of gene product (increment operation) and external inhibition decreases respectively (decrement operation). Incrementing and decrementing transcription factors always form complexes with program counter objects. The interplay of those specific transcription factors manages the program control.

3 Case Study: Computational Units and Circuits

Artificial GRNs have been instrumental in elucidating basic principles that govern the dynamics and consequences of stochasticity in the gene expression of naturally occurring GRNs. The realisation as computational circuits infers inherent evolutionary fault tolerance and robustness to these modular units.

In a case study, we introduce three artificial GRNs for logic gates (inverter, NAND gate, RS flip-flop) and describe their dynamical behaviour quantitatively by an ordinary differential equation model using Hill kinetics and by corresponding P systems Π_{Hill} .

A variety of distinguishable transcription factors given by their concentration over the time course enables communication between as well as coupling of computational units. Thus, circuit engineering becomes feasible.

Inverter

input: concentration level of transcription factor xoutput: concentration level of gene product y



Simulation Result (MATLAB, P system iteration scheme)

dynamical behaviour depicted for m = 2, $\Delta \tau = 0.1$, $\Theta_j = 500$, $j \in \{x, a\}$ rg = 10,000, eg = 10,000, $x_0 = 0$, $y_0 = 0$, $a_0 = 0$



NAND gate

input: concentration levels of transcription factors x (input1), y (input2) output: concentration level of gene product z



Ordinary Differential Equations

 $\begin{aligned} \dot{a} &= h^+(x, \Theta_x, m) - a \\ \dot{b} &= h^+(y, \Theta_y, m) - b \\ \dot{z} &= 1 - h^+(a, \Theta_a, m) \cdot h^+(b, \Theta_b, m) - z \\ \text{Simulation Result (Copasi, ODE solver)} \end{aligned}$

dynamical behaviour depicted for $m = 2, \Theta_j = 0.1, j \in \{x, y, a, b\}$



Corresponding P System

$$\begin{split} &H_{\mathrm{Hill},\mathrm{GRNnand}} = (V_{\mathrm{Genes}}, V_{\mathrm{GeneProducts}}, \Sigma, [1]_{1}, L_{0}, r_{1}, \ldots, r_{6}, f_{1}, \ldots, f_{6}, \Delta\tau, m) \\ &V_{\mathrm{Genes}} = \{RegGeneX, RegGeneY, EffGene\} \\ &V_{\mathrm{GeneProducts}} = \{x, y, z, \neg a, \neg b\} \\ &\Sigma = \{z\} \\ &L_{0} = \{(RegGeneX, rgx), (RegGeneY, rgy), (EffGene, eg), \\ & (x, x_{0}), (y, y_{0}), (z, z_{0}), (\neg a, a_{0}), (\neg b, b_{0})\} \\ &r_{1} : s_{1} RegGeneX \longrightarrow s_{1} \neg a + s_{1} RegGeneX \mid_{x} \\ &r_{2} : s_{2} \neg a \longrightarrow \emptyset \\ &r_{3} : s_{3} RegGeneY \longrightarrow s_{3} \neg b + s_{3} RegGeneY \mid_{y} \\ &r_{4} : s_{4} \neg b \longrightarrow \emptyset \\ &r_{5} : s_{5} EffGene \longrightarrow s_{5} z + s_{5} EffGene \mid_{\neg a, \neg b} \\ &r_{6} : s_{6} z \longrightarrow \emptyset \\ f_{1}(\Delta\tau, L_{t}, m) = \left[\Delta\tau \cdot |L_{t} \cap \{(RegGeneX, \infty)\}| \cdot \frac{|L_{t} \cap \{(x, \infty)\}|^{m}}{|L_{t} \cap \{(x, \infty)\}|^{m}} \right] \\ f_{2}(\Delta\tau, L_{t}, m) = \left[\Delta\tau \cdot |L_{t} \cap \{(ra, \infty)\}|\right] \\ f_{3}(\Delta\tau, L_{t}, m) = \left[\Delta\tau \cdot |L_{t} \cap \{(RegGeneY, \infty)\}| \cdot \frac{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{x}^{m}}{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{y}^{m}} \right] \\ f_{4}(\Delta\tau, L_{t}, m) = \left[\Delta\tau \cdot |L_{t} \cap \{(RegGeneY, \infty)\}| \cdot \frac{|L_{t} \cap \{(y, \infty)\}|^{m}}{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{y}^{m}} \right] \\ f_{5}(\Delta\tau, L_{t}, m) = \left[\Delta\tau \cdot |L_{t} \cap \{(rd, \infty)\}|^{m} + \Theta_{a}^{m}} \cdot \frac{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{b}^{m}}}{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{b}^{m}}} \right] \\ f_{6}(\Delta\tau, L_{t}, m) = \left[\Delta\tau \cdot |L_{t} \cap \{(z, \infty)\}|^{m} + \Theta_{a}^{m}} \cdot \frac{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{b}^{m}}}{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{b}^{m}}} \right] \\ f_{6}(\Delta\tau, L_{t}, m) = \left[\Delta\tau \cdot |L_{t} \cap \{(z, \infty)\}|^{m} + \Theta_{a}^{m}} \cdot \frac{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{b}^{m}}}{|L_{t} \cap \{(z, \infty)\}|^{m} + \Theta_{b}^{m}}} \right] \\ f_{6}(\Delta\tau, L_{t}, m) = \left[\Delta\tau \cdot |L_{t} \cap \{(z, \infty)\}|^{m} + \Theta_{a}^{m}} \cdot \frac{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{b}^{m}}}{|L_{t} \cap \{(z, \infty)\}|^{m} + \Theta_{b}^{m}}} \right] \\ f_{6}(\Delta\tau, L_{t}, m) = \left[\Delta\tau \cdot |L_{t} \cap \{(z, \infty)\}|^{m} + \Theta_{a}^{m}} \cdot \frac{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{b}^{m}}}{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{b}^{m}}} \right] \\ f_{6}(\Delta\tau, L_{t}, m) = \left[\Delta\tau \cdot |L_{t} \cap \{(z, \infty)\}|^{m} + \Theta_{a}^{m}} \cdot \frac{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{b}^{m}}}{|L_{t} \cap \{(y, \infty)\}|^{m} + \Theta_{b}^{m}}} \right] \\ \\ \int d\tau \in \mathbb{R}_{+} \\ m \in \mathbb{N}_{+} \end{aligned}$$

Simulation Result (MATLAB, P system iteration scheme)

dynamical behaviour depicted for m = 2, $\Delta \tau = 0.1$, $\Theta_j = 500$, $j \in \{x, y, a, b\}$ $rgx = 10,000, rgy = 10,000, eg = 10,000, x_0 = 0, y_0 = 0, z_0 = 0, a_0 = 0, b_0 = 0$



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RS Flip-Flop

input: concentration levels of transcription factors $\overline{S}, \overline{R}$ output: concentration level of gene product Q



Ordinary Differential Equations

$$\begin{split} \dot{a} &= 1 - \mathbf{h}^{+}(b, \Theta_{b}, m) \cdot \mathbf{h}^{-}(\overline{S}, \Theta_{\overline{S}}, m) - a \\ \dot{b} &= 1 - \mathbf{h}^{+}(a, \Theta_{a}, m) \cdot \mathbf{h}^{-}(\overline{R}, \Theta_{\overline{R}}, m) - b \\ \dot{Q} &= \mathbf{h}^{+}(b, \Theta_{b}, m) \cdot \mathbf{h}^{-}(\overline{S}, \Theta_{\overline{S}}, m) - Q \end{split}$$

Simulation Result (Copasi, ODE solver)

dynamical behaviour depicted for $m = 2, \, \Theta_j = 0.1, \, j \in \{a, b, \overline{R}, \overline{S}\}$



Corresponding P System

$$\begin{split} \Pi_{\mathrm{Hill},\mathrm{GRNrsff}} &= (V_{\mathrm{Genes}}, V_{\mathrm{GeneProducts}}, \Sigma, [_1]_1, L_0, r_1, \dots, r_6, f_1, \dots, f_6, \Delta\tau, m) \\ V_{\mathrm{Genes}} &= \{ RegGeneResetState, RegGeneSetState, EffGene \} \\ V_{\mathrm{GeneProducts}} &= \{ Q, \neg \overline{S}, \neg \overline{R}, \neg a, \neg b \} \\ \Sigma &= \{ Q \} \\ L_0 &= \{ (RegGeneResetState, rgr), (RegGeneSetState, rgs), \\ (EffGene, eg), (Q, q_0), (\overline{S}, ss_0), (\overline{R}, rs_0), (\neg a, a_0), (\neg b, b_0) \} \\ r_1 &: s_1 \ RegGeneResetState \longrightarrow s_1 \ \neg a + s_1 \ RegGeneResetState |_{\neg \overline{S}, \neg b} \\ r_2 &: s_2 \ \neg a \longrightarrow \emptyset \\ r_3 &: s_3 \ RegGeneSetState \longrightarrow s_3 \ \neg b + s_3 \ RegGeneSetState |_{\neg \overline{R}, \neg a} \\ r_4 &: s_4 \ \neg b \longrightarrow \emptyset \\ r_5 &: s_5 \ EffGene \longrightarrow s_5 \ Q + s_5 \ EffGene \ |_{\neg \overline{S}, \neg b} \end{split}$$

$$\begin{split} r_{6} &: s_{6} \ Q \longrightarrow \emptyset \\ f_{1}(\Delta\tau, L_{t}, m) &= \left[\Delta\tau \cdot |L_{t} \cap \{(\operatorname{Reg}Gene\operatorname{ResetState}, \infty)\}| \cdot \\ & \left(1 - \frac{|L_{t} \cap \{(\neg b, \infty)\}|^{m}}{|L_{t} \cap \{(\neg b, \infty)\}|^{m} + \Theta_{\neg b}^{m}} \cdot \left(1 - \frac{|L_{t} \cap \{(\neg \overline{S}, \infty)\}|^{m}}{|L_{t} \cap \{(\neg \overline{S}, \infty)\}|^{m} + \Theta_{\neg \overline{S}}^{m}} \right) \right) \right] \\ f_{2}(\Delta\tau, L_{t}, m) &= \left[\Delta\tau \cdot |L_{t} \cap \{(\neg a, \infty)\}| \right] \\ f_{3}(\Delta\tau, L_{t}, m) &= \left[\Delta\tau \cdot |L_{t} \cap \{(\operatorname{Reg}GeneSetState, \infty)\}| \cdot \\ & \left(1 - \frac{|L_{t} \cap \{(\neg a, \infty)\}|^{m}}{|L_{t} \cap \{(\neg a, \infty)\}|^{m} + \Theta_{\neg a}^{m}} \cdot \left(1 - \frac{|L_{t} \cap \{(\neg \overline{R}, \infty)\}|^{m}}{|L_{t} \cap \{(\neg \overline{R}, \infty)\}|^{m} + \Theta_{\neg \overline{R}}^{m}} \right) \right) \right] \\ f_{4}(\Delta\tau, L_{t}, m) &= \left[\Delta\tau \cdot |L_{t} \cap \{(\neg b, \infty)\}| \right] \\ f_{5}(\Delta\tau, L_{t}, m) &= \left[\Delta\tau \cdot |L_{t} \cap \{(\operatorname{Cff}Gene, \infty)\}| \cdot \\ & \frac{|L_{t} \cap \{(\neg b, \infty)\}|^{m}}{|L_{t} \cap \{(\neg b, \infty)\}|^{m} + \Theta_{\neg b}^{m}} \cdot \left(1 - \frac{|L_{t} \cap \{(\neg \overline{S}, \infty)\}|^{m}}{|L_{t} \cap \{(\neg \overline{S}, \infty)\}|^{m} + \Theta_{\neg \overline{S}}^{m}} \right) \right] \\ f_{6}(\Delta\tau, L_{t}, m) &= \left[\Delta\tau \cdot |L_{t} \cap \{(Q, \infty)\}| \right] \\ \Delta\tau \in \mathbb{R}_{+} \\ & m \in \mathbb{N}_{+} \end{split}$$

Simulation Result (MATLAB, P system iteration scheme)

dynamical behaviour depicted for m = 2, $\Delta \tau = 0.1$, $\Theta_j = 500$, $j \in \{a, b, \overline{R}, \overline{S}\}$ rgr = 10,000, rgs = 10,000, eg = 10,000, $q_0 = 0$, $ss_0 = 0$, $rs_0 = 0$, $a_0 = 0$, $b_0 = 0$



A homologous analytic model of a bistable toggle switch was introduced in [8]. In case of the forbidden input signalling $\overline{S} = 1$, $\overline{R} = 1$, the normalised concentrations of both inhibitors $\neg a$ and $\neg b$ converge to 0.5. By setting or resetting input signalling, the flip-flop restores.

4 RS Flip-Flop Validation in vivo

In addition to prediction and simulation of GRNs acting as logic gates, we demonstrate the practicability of the RS flip-flop by an experimental study *in vivo*. Resulting output protein data measured over the time course can validate the system model. Following the pioneering implementation of a bistable toggle switch [8], we

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could confirm its function in a previous study [10]. Two extensions were investigated: Firstly, the effects of IPTG and AHL as appropriate intercellular inducers for flip-flop setting were shown. Secondly, flow cytometry was used to quantitatively measure protein concentrations within the flip-flop implementation. In the following, we give a brief overview of the experimental setup and compare obtained experimental results with our simulations based on the models.

Biological Principles and Prerequisites

Quorum Sensing and Autoinduction via AHL

In quorum sensing, bacterial species regulate gene expression based on cellpopulation density [17]. An alteration in gene expression occurs when an intercellular signalling molecule termed autoinducer, produced and released by the bacterial cells reaches a critical concentration. Termed as quorum sensing or auto induction, this fluctuation in autoinducer concentration is a function of bacterial cell-population density. Vibrio fischeri, a well studied bacterium, colonises the light organs of a variety of marine fishes and squids, where it occurs at very high densities $(10^{10} \frac{\text{cells}}{\text{ml}})$ and produces light. The two genes essential for cell density regulation of luminescence are: luxI, which codes for an autoinducer synthase [22]; and luxR, which codes for an autoinducer-dependent activator of the luminescence genes. The luxR and luxI genes are adjacent and divergently transcribed, and luxI is the first of seven genes in the luminescence or lux operon. LuxI-type proteins direct AHL synthesis while LuxR-type proteins function as transcriptional regulators that are capable of binding AHL signal molecules. Once formed, LuxR-AHL complexes bind to target promoters of quorum-regulated genes. Quorum sensing is now known to be widespread among both Gram-positive and Gram-negative bacteria.

Bioluminescence in Vibrio fischeri

Bioluminescence in general is defined as an enzyme catalysed chemical reaction in which the energy released is used to produce an intermediate or product in an electronically excited state, which then emits a photon. It differs from fluorescence or phosphorescence as it is not depended on light absorbed. The mechanism for gene expression and the structure of the polycistronic message of the lux structural genes in *Vibrio fischeri* have been thoroughly characterised [9]. Briefly, there are two substrates, luciferin, which is a reduced flavin mononucleotide (FMNH₂), and a long chain (7 – 16 carbons) fatty aldehyde (RCHO). An external reductant acts via flavin mono-oxygenase oxidoreductase to catalyse the reduction of FMN to FMNH₂, which binds to the enzyme and reacts with O_2 to form a 4a-peroxyflavin intermediate. This complex oxidises the aldehyde to form the corresponding acid (RCOOH) and a highly stable luciferase-hydroxyflavin intermediate in its excited state, which decays slowly to its ground state emitting blue-green light $h\nu$ with a maximum intensity at about 490nm:

 $\mathrm{FMNH}_2 + \mathrm{RCHO} + \mathrm{O}_2 \stackrel{\mathrm{lucif.}}{\longrightarrow} \mathrm{FMN} + \mathrm{H}_2\mathrm{O} + \mathrm{RCOOH} + h\nu$

Transcription Control by LacR and λ CI Repressor Proteins

Escherichia coli cells repress the expression of the lac operon when glucose is abundant in the growth medium. Only when the glucose level is low and the lactose level is high, the operon is fully expressed. The Lac repressor LacR is a 360 residue long protein that associates into a homotetramer. It contains a helix-turnhelix (HTH) motif through which it interacts with DNA. This interaction represses transcription by hindering association with RNA polymerase and represents an example of "combinatorial control" widely seen in prokaryotes and eukaryotes [4]. The CI repressor of bacteriophage lambda is the key regulator in lambda's genetic switch, a bistable switch that underlies the phage's ability to efficiently use its two modes of development [21].

Flow Cytometry

Flow cytometry refers to the technique where microscopic particles are counted and examined as they pass in a hydro-dynamically focused fluid stream through a measuring point surrounded by an array of detectors. Previously, flow cytometry analyses were performed by us using a BD LSRII flow cytometer equipped with 405nm, 488nm and 633nm lasers. 488nm laser was used for gfp and yellow fluorescent protein (yfp) quantification.

Experimental Setup and Implementation

We have shown that an *in vivo* system [10] can potentially be used to mimic a RS flip-flop [13] and have quantified its performance using flow cytometry. The presence or absence of the inducers IPTG or AHL in combination with temperature shift acts as an input signal. The toggle switch comprising of structural genes for reporter/output proteins fused to promoter regions that are regulated by input signals is visualised as a RS flip-flop. The functional modularity of input and output circuits is maintained so that the artificial GRN used can be easily extended for future studies.

This design endows cells with two distinct phenotypic states: where the λ CI activity is high and the expression of lacI is low (referred to as high or 1 state), or where the activity of LacR is high and the expression of λ CI is low (referred to as low or 0 state). *gfp* is expressed only in the high λ CI/low LacR state.



Fig. 4. A schematic diagram of an AHL biosensor module interfaced with the genetic toggle switch adapted from [10]. The transgenic artificial GRN consists of a bistable genetic toggle switch [8] which is interfaced with genes from the lux operon [6] of the quorum sensing signalling pathway of *Vibrio fischeri* [22].

Results and Discussion

For co-relational purposes, all experiments were conducted with both BL21 and Top10 strains of *Escherichia coli*. The concentration of IPTG used in all the experiments was 2mM and that of AHL was 1μ M. Experiments conducted without the use of inducers, lead to an unreliable shifting of the states, signifying the use if inducers in a tightly, mutually regulated circuit. Further experiments conducted to understand the switching dynamics of the circuit revealed that in the current scenario, it was easier to switch from a high to a low state than vice versa. This discrepancy in switching behaviour is attributed to the differing modes of elimination of LacR and λ CI repressor proteins. While switching from low to high state, the repression due to IPTG-bound Lac repressor needs to be overcome by cell growth. Switching from high to low state is effected by immediate thermal degradation of the temperature-sensitive λ CI. Experiments were also conducted to test the sustainability of states. The plug and play property of the circuit was examined by employing yfp as the reporter gene instead of gfp. As shown in Figure 5, the circuit could reliably mimic a RS flip-flop. The massive parallelism permissible by the use of large quantities of cells can compensate for the slow speed of switching. Further tests are to be performed to confirm this hypothesis.

5 Conclusions

The dynamical behaviour of GRNs is able to emulate information processing in terms of performing computations. In order to formalise this capability, we have introduced P systems of the form Π_{Hill} incorporating cooperativity and competitivity between transcription factors based on Hill kinetics. Its transformation to a dedicated iteration scheme for a discrete term rewriting mechanism with variable stoichiometric factors in Π_{Hill} provides a homogeneous approach that allows to compose GRNs towards functional units like computing agents. Examples address computational units (inverter, NAND gate, RS flip-flop), each defined by GRN, its ODE model, and the corresponding P system. Simulations of the dynamical



Fig. 5. Inducer-dependent switching. Repeated activation and deactivation of the toggle switch based on inducers and temperature. Temperature was switched every 24 hours. Cells were incubated with inducers for 12 hours, followed by growth for 12 hours without inducers, initially kept at 30°C (**A**) and 42°C (**B**). The cells successfully switched states thrice.

behaviour quantitatively show the switching characteristics as well as the expected quality of binary output signals. Along with the prediction of GRNs acting as computational units, an experimental study *in vivo* demonstrates their practicability. Although the measurement of the dynamic switching behaviour was condensed to 12 points in time, they approximate the expected course. At the crossroad of modelling, simulation, and verification of biological reaction networks, the potential of amalgamating analytic, stochastic, and algebraic approaches into the P systems framework seems promising for applications in systems biology to explore network functions.

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Some Applications of Spiking Neural P Systems

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Summary. In this paper we investigate some applications of spiking neural P systems regarding their capability to solve some classical computer science problems. In this respect it is studied the versatility of such systems to simulate a well known parallel computational model, namely the Boolean circuits. In addition, another notorious application the sorting - is considered within this framework.

1 Introduction

Spiking neural P systems (shortly called SN P systems) are a class of computing models introduced in [9]. They are using ideas from neural computing, area currently under high investigation, with a focus on spiking neurons (see, e.g., [4], [12], [13]).

The new models are based on the tissue-like and neural-like P systems structure to which various features were added. Details can be found at the website of membrane computing ([21]). For an introduction in the area we refer to [16].

In short, an SN P system consists of a set of neurons placed in the nodes of a graph and sending signals (spikes) along synapses (edges of the graph), under the control of firing rules. One also uses forgetting rules, which remove spikes from neurons. Hence, the spikes are moved and created, destroyed, but never modified (there is only one type of objects in the system).

A generalization of the original model was considered in [15], [3] where rules of the form $E/a^c \to a^p$; d where introduced. The meaning is that when using the rule, c spikes are consumed and p spikes are produced. Because p can be 0 or greater than 0, we obtain at the same time a generalization of both spiking and forgetting rules. Different from the original model of SN P systems, in [10], parallelism inside a neuron was introduced. By that we mean that when a rule $E/a^c \to a; d$ can be applied (the contents of a neuron is described by the regular expression E), then we apply it as many times as possible in that neuron.

Based on the above features, we investigate their power to simulate Boolean gates and circuits. We also introduce here a modality to sort natural numbers (given as number of spikes) with SN P systems in the initial version.

2 Prerequisites

In this section we first introduce the definition of SN P system which we will use during our endeavor, altogether with some explanations on the exhaustive use of the rules. Then, we recall (some) basic notions on Boolean functions and circuits.

2.1 SN P systems

A spiking neural P system (in short, an SN P system), of degree $m \ge 1$, is a construct of the form

$$\Pi = (O, \sigma_1, \ldots, \sigma_m, syn, out),$$

where:

- 1. $O = \{a\}$ is the singleton alphabet (a is called *spike*);
- 2. $\sigma_1, \ldots, \sigma_m$ are *neurons*, of the form $\sigma_i = (n_i, R_i), 1 \leq i \leq m$, where:
 - a) $n_i \ge 0$ is the *initial number of spikes* contained by the neuron;
 - b) R_i is a finite set of *rules* of the following two forms:
 - (1) $E/a^c \to a; d$, where E is a regular expression over $O, c \ge 1$, and $d \ge 0;$ (2) $a^s \to \lambda$, for some $s \ge 1$, with the restriction that $a^s \in L(E)$ for no rule $E/a^c \to a; d$ of type (1) from $R_i;$
- 3. $syn \subseteq \{1, 2, \ldots, m\} \times \{1, 2, \ldots, m\}$ with $(i, i) \notin syn$, for $1 \le i \le m$ (synapses);
- 4. $out \in \{1, 2, ..., m\}$ indicates the *output neuron*.

The rules of type (1) are firing (also called spiking) rules, and the rules of type (2) are called forgetting rules. The first ones are applied as follows: if the neuron contains k spikes, $a^k \in L(E)$ and $k \geq c$, then the rule $E/a^c \to a; d$ can be applied, and this means that c spikes are consumed, only k - c remain in the neuron, the neuron is fired, and it produces one spike after d time units (a global clock is assumed, marking the time for the whole system, hence the functioning of the system is synchronized). If d = 0, then the spike is emitted immediately, if d = 1, then the spike is emitted in the next step, and so on. In the case $d \geq 1$, if the rule is used in step t, then in steps $t, t + 1, t + 2, \ldots, t + d - 1$ the neuron is closed, and it cannot receive new spikes (if a neuron has a synapse to a closed neuron and sends a spike along it, then the spike is lost). In step t + d, the neuron spikes and becomes again open, hence can receive spikes (which can be used in step t + d + 1). A spike emitted by a neuron σ_i is replicated and goes to all neurons σ_j such that $(i, j) \in syn$. The forgetting rules, are applied as follows: if the neuron contains exactly s spikes, then the rule $a^s \to \lambda$ can be used, and this means that all s spikes are removed from the neuron.

In each time unit, in each neuron which can use a rule we have to use a rule, either a firing or a forgetting one. Because two firing rules $E_1/a^{c_1} \rightarrow a; d_1$ and $E_2/a^{c_2} \rightarrow a; d_2$ can have $L(E_1) \cap L(E_2) \neq \emptyset$, it is possible that two or more rules can be applied in a neuron, and then one of them is chosen non-deterministically. Note however that we cannot interchange a firing rule with a forgetting rule, as all pairs of rules $E/a^c \rightarrow a; d$ and $a^s \rightarrow \lambda$ have disjoint domains, in the sense that $a^s \notin L(E)$.

The initial configuration of the system is described by the numbers n_1, n_2, \ldots, n_m of spikes present in each neuron. Starting from the initial configuration and applying the rules, we can define transitions among configurations. A transition between two configurations C_1, C_2 is denoted by $C_1 \implies C_2$. Any sequence of transitions starting in the initial configuration is called a *computation*. A computation halts if it reaches a configuration where all neurons are open and no rule can be used.

With any computation, halting or not, we associate a spike train, a sequence of digits 0 and 1, with 1 appearing in positions $1 \leq t_1 < t_2 < \ldots$, indicating the steps when the output neuron sends a spike out of the system (we also say that the system itself spikes at that time). With any spike train containing at least two spikes we associate a result, in the form of the number $t_2 - t_1$; we say that this number is computed by Π . By definition, if the spike train contains only one occurrence of 1, then we say that we have computed the number zero. The set of all numbers computed in this way by Π is denoted by $N_2(\Pi)$ (the subscript indicates that we only consider the distance between the first two spikes of any computation). Then, by $Spik_2P_m(rule_k, cons_q, forg_r)$ we denote the family of all sets $N_2(\Pi)$ computed as above by spiking neural P systems with at most $m \geq 1$ neurons, using at most $k \geq 1$ rules in each neuron, with all spiking rules $E/a^c \to a; t$ having $c \leq q$, and all forgetting rules $a^s \to \lambda$ having $s \leq r$. When one of the parameters m, k, q, r is not bounded, it is replaced with *.

A rule of the type $E/a^c \to a^p$ is called an *extended rule*, and is applied as follows: if neuron σ_i contains k spikes, and $a^k \in L(E), k \geq c$, then the rule can fire, and its application means consuming (removing) c spikes (thus only k - cremain in σ_i) and producing p spikes, which will exit immediately the neuron.

In this paper, we use SN P systems of the form introduced above, but using the rules in the exhaustive way. Namely if a rule $E/a^c \to a^p; d$ is associated with a neuron σ_i which contains k spikes, then the rule is enabled (we also say *fired*) if and only if $a^k \in L(E)$. Using the rule means the following. Assume that k = sc+r, for some $s \ge 1$ (this means that we must have $k \ge c$) and $0 \le r < c$ (the remainder of dividing k by c). Then sc spikes are consumed, r spikes remain in the neuron σ_i , and sp spikes are produced and sent to the neurons σ_j such that $(i, j) \in syn$ (as usual, this means that the sp spikes are replicated and exactly sp spikes are sent to each of the neurons σ_j). In the case of the output neuron, sp spikes are
also sent to the environment. Of course, if neuron σ_i has no synapse leaving from it, then the produced spikes are lost.

We stress two important features of this model. First, it is important to note that only one rule is chosen and applied, the remaining spikes cannot evolve by another rule. For instance, even if a rule $a(aa)^*/a \rightarrow a; 0$ exists, it cannot be used for the spike remaining unused after applying the rule $a(aa)^*/a^2 \rightarrow a; 0$. Second, is that the covering of the neuron is checked only for enabling the rule, not step by step during its application. For instance, the rule $a^5/a^2 \rightarrow a; 0$ has the same effect as $a(aa)^*/a^2 \rightarrow a; 0$ in the case of a neuron containing exactly 5 spikes: the rule is enabled, 4 spikes are consumed, 2 are produced; both applications of the rule are concomitant, not one after the other, hence all of them have the same enabling circumstances.

If several rules of a neuron are enabled at the same time, one of them is nondeterministically chosen and applied. The computations proceed as in the SN P systems with usual rules, and a spike train is associated with each computation by writing 0 for a step when no spike exits the system and 1 within a step when one or more spikes exit the system. Then, a number is associated – and said to be generated/computed by the respective computation – with a spike train containing at least two occurrences of the digit 1, in the form of the steps elapsed between the first two occurrences of 1 in the spike train. Number 0 is computed by computations whose spike trains contain only one occurrence of 1.

2.2 Boolean Functions and Circuits

An *n*-ary Boolean function is a function $f\{true, false\}^n \mapsto \{true, false\}$. \neg (negation) is a unary Boolean function (the other unary functions are: constant functions and identity function). We say that Boolean expression φ with variables x_1, \ldots, x_n expresses the *n*-ary Boolean function *f* if, for any *n*-tuple of truth values $t = (t_1, \cdots, t_n), f(t)$ is true if $T \vDash \varphi$, and f(t) is false if $T \nvDash \varphi$, where $T(x) = t_i$ for $i = 1, \ldots, n$.

There are three primary Boolean functions that are widely used: The NOT function - this is a just a negation; the output is the opposite of the input. The NOT function takes only one input, so it is called a unary function or operator. The output is true when the input is false, and vice-versa. The AND function - AND function returns true only if all inputs are true; if there is an input which is false the function returns false. The OR function - the output of an OR function is true if the first input is true or the second input is true or the third input is true, etc. (hence, to return true is enough for one input to be true). Both AND and OR can have any number of inputs, with a minimum of two.

Any *n*-ary Boolean function f can be expressed as a Boolean expression φ_f involving variables x_1, \ldots, x_n .

There is a potentially more economical way than expressions for representing Boolean functions, namely *Boolean circuits*. A Boolean circuit is a graph C = (V, E), where the nodes in $V = \{1, \ldots, n\}$ are called the *gates* of C. Graph C has a rather special structure. First, there are no cycles in the graph, so we can assume that all edges are of the form (i, j), where i < j. All nodes in the graph have the "in-degree" (number of incoming edges) equal to 0, 1, or 2. Also, each gate $i \in V$ has a *sort* s(i) associated with it, where $s(i) \in \{true, false, \lor, \land, \neg\} \cup \{x_1, x_2, \ldots\}$. If $s(i) \in \{true, false\} \cup \{x_1, x_2, \ldots\}$, then the in-degree of i is 0, that is, i must have no incoming edges. Gates with no incoming edges are called the *inputs* of C. If $s(i) = \neg$, then i has "in-degree" one. If $s(i) \in \{\lor, \land\}$, then the in-degree of i must be two. Finally, node n (the largest numbered gate in the circuit, which necessarily has no outgoing edges) is called the *output gate* of the circuit.

This concludes our definition of the syntax of circuits. The semantics of circuits specifies a truth value for each appropriate truth assignment. We let X(C) be the set of all Boolean variables that appear in the circuit C (that is, $X(C) = \{x \in X \mid s(i) = x \text{ for some gate } i \text{ of } C\}$). We say that a truth assignment T is appropriate for C if it is defined for all variables in X(C). Given such a T, the truth value of gate $i \in V$, T(i), is defined, by induction on i, as follows: If s(i) = true then T(i) = true, and similarly if s(i) = false. If $s(i) \in X$, then T(i) = T(s(i)). If now $s(i) = \neg$, there is a unique gate j < i such that $(j, i) \in E$. By induction, we know T(j), and then T(i) is true if T(j) = false, and vice-versa. If $s(i) = \lor$, then there are two edges (j, i) and (j', i) entering i. T(i) is true if only if at least one of T(j), T(j') is true. If $s(i) = \land$, then T(i) is true if only if both T(j) and T(j') are true, where (j, i) and (j', i) are the incoming edges. Finally, the value of the circuit, T(C), is T(n), where n is the output gate.

3 Simulating Logical Gates and Circuits

In this section we show how SNP systems can simulate logical gates. We consider that input is given in one neuron while the output will be collected from the output neuron of the system. Boolean value 1 is encoded in the spiking system by two spikes, hence a^2 , while 0 is encoded as one spike.

We collect the result as follows. If the output neuron fires two spikes in the second step of the computation, then the Boolean value computed by the system is 1 (hence true). If it fires only one spike, then the result is 0 (false).

3.1 Simulating Logical Gates

Lemma 1. Boolean AND gate can be simulated by SN P systems using two neurons and no delay on the rules, in two steps.

Proof. We construct the SNP system (formed by only one neuron):

$$\Pi_{AND} = (\{a\}, \sigma_1 = (0, \{a^2 \to a; 0, a^3 \to a; 0, a^4/a^2 \to a; 0\}), \emptyset, 1).$$

The functioning of the system is rather simple (remember that the rules are used in an exhaustive way). Suppose in neuron 1 we introduce three spikes. This

means we compute the logical AND between 1 and 0 (or 0 and 1). The only rule the system can use is $a^3 \rightarrow a; 0$ and one spike (hence the correct result - 0 in this case) is sent to the environment.

If 4 spikes are introduced in neuron 1 (the case 11), the output neuron will fire using the rule $a^4/a^2 \rightarrow a; 0$, and will send two spikes in the environment. The system with the input 00 behaves similarly to the 01 or 10 cases. We have shown how the system we have constructed gives the right answer in one computational step and gets back to its initial configuration for a further use, if necessary.

We want to emphasize here that no "extended" rule was used. Of course, a rule $a^4 \rightarrow a^2$ can substitute, with the same effect, the rule we have preferred above (namely $a^4/a^2 \rightarrow a; 0$) but, in simulating Boolean gates, we have tried to minimize the use of such rules. An extended rule is used only once in simulating Boolean gates, more precisely in the simulation of OR gate.

If in the system above, in the output neuron, we change only the rule $a^3 \rightarrow a; 0$ (with the rule $a^3 \rightarrow a^2; 0$) we obtain the OR gate.

Lemma 2. Boolean OR gate can be simulated by SN P systems using two neurons and no delay on the rules, in two steps.

We now pass to the simulation of logical gate NOT.

Lemma 3. Boolean NOT gate can be simulated by SNP systems using two neurons, no delay on the rules, in two steps.

Proof. We first want to stress that in simulating this gate we did not use any extended rules. The case when such rules are used is left to the reader.

Let us construct the following SN P system:

$$\Pi_{NOT} = (\{a\}, \sigma_1, \sigma_2, \{(1, 2), (2, 1)\}, 1),$$

and:

- $\sigma_1 = (a, \{a^2/a \to a; 0, a^3 \to a; 0\}),$ • $\sigma_2 = (0, \{a/a \to a; 0, a^2/a^2 \to a; 0\}).$
- Let us emphasize that in order to simulate Boolean gate NOT, in the initial configuration, neuron 1 contains 1 spike, which, once used to correctly simulate the gate, has to be present again in the neuron such that the system returns to its initial configuration. This is done with the help of neuron 2 which in step 2 of the computation refills neuron 1 with one spike.

The system is given in its initial configuration in Figure 1. This gives us the opportunity to introduce the way we graphically represent a SN P system: as a directed graph, with the neurons as nodes and the synapses indicated by arrows. Each neuron has inside its specific rules and the spikes present in the initial configuration.

If the input in the Boolean gate is 1, then two spikes are placed in neuron 1. Having three spikes inside (two from the input, and one initially present inside) neuron 1 can use only rule $a^3 \to a; 0$, thus sending one spike to the environment (hence Boolean 0 – the correct result – is obtained), and one spike to neuron 2. The latter one will send the spike back, in the second step of the computation by using rule $a/a \to a; 0$, and the system regains its initial configuration.



Figure 1. SN P systems simulating **NOT** gate

If the input in the Boolean gate is 0, hence one spike is introduced in neuron 1, it uses the rule $a^2/a \rightarrow a; 0$, two spikes are sent to the environment (and the result of the computation is 1), and to neuron 2 in the same time. In the second step of the computation neuron 2 uses the rule $a^2/a^2 \rightarrow a; 0$, consumes the two spikes present inside, and sends one back to neuron 1. The system recovers its initial configuration.

After showing how SN P systems can simulate logical gates, we pass to the simulation of circuits.



Figure 2. Boolean Circuit and the Spiking System

3.2 Simulating Circuits

Next, we are presenting an example of how to construct a SN P system to simulate a Boolean circuit designed to evaluate a Boolean function. Of course, in our goal we are using the systems Π_{AND} , Π_{OR} , and Π_{NOT} constructed before, to which we add extra neurons to synchronize the system for a correct output.

We start with the same **example** considered in [1] and [11], namely the function $f: \{0,1\}^4 \to \{0,1\}$ given by the formula

$$f(x_1, x_2, x_3, x_4) = (x_1 \land x_2) \lor \neg (x_3 \land x_4).$$

The circuit corresponding to the above formula as well as the spiking system assign to it are depicted in Figure 2.

In order for the system that simulates the circuit to output the correct result it is necessary for each sub-system (that simulates the gates AND, OR, and NOT) to receive the input from the above gate(s) at the same time. To this aim, we have to add synchronization neurons, initially empty with a single rule inside $(a \rightarrow a; 0)$. Note that in Figure 2. we have added such a neuron in order for the output of the first AND gate to enter gate OR at the same time with the output of NOT gate (at the end of the second step of the computation).

Having the overall image of the functioning of the system, let us give some more details on the simulation of the above formula. For that we construct the SN P system

$$\Pi_C = (\Pi_{AND}^{(1)}, \Pi_{AND}^{(2)}, \Pi_{NOT}^{(3)}, \Pi_{OR}^{(4)})$$

formed by the sub-SN P systems for each gate, and we obtain the unique result as follows:

- 1. for every gate of the circuit with inputs from the input gates we have a SN P system to simulate it. The input is given in neuron labeled 1 of each gate;
- 2. for each gate which has at least one input coming as an output of a previous gate we construct a SN P system to simulate it by "constructing" a synapse between the output neuron of the gate from which the signal (spike) comes and the input neuron of the system that simulates the new gate.

Note that if synchronization is needed the new synapse is constructed from the output neuron of the output gate to the synchronization neuron and from here another synapse is constructed to the input of the new gate in the circuit.

For the above formula and the circuit depicted in Figure 2 we will have:

- $\Pi_{AND}^{(1)}$ computes the first AND₁ gate $(x_1 \wedge x_2)$ with inputs x_1 and x_2 . $\Pi_{AND}^{(2)}$ computes the second AND₂ gate $(x_3 \wedge x_4)$ with inputs x_3 and x_4 ; these two P systems, $\Pi_{AND}^{(1)}$ and $\Pi_{AND}^{(2)}$, act in parallel.
- $\Pi_{NOT}^{(3)}$ computes NOT gate $\neg(x_3 \wedge x_4)$ with input $(x_3 \wedge x_4)$. While $\Pi_{NOT}^{(3)}$ is working, the output value of the first AND_1 gate passes through the synchronization neuron.
- The input enters in the first neuron of OR gate, and SN P system $\Pi_{OR}^{(4)}$ completes its task. The result of the computation for OR gate (which is the result of the global P system), is sent into the environment of the whole system.

Generalizing the previous observations the following result holds:

Theorem 1. Every Boolean circuit α , whose underlying graph structure is a rooted tree, can be simulated by a SN P system, Π_{α} , in linear time. Π_{α} is constructed from SN P systems of type Π_{AND} , Π_{OR} and Π_{NOT} , by reproducing in the architecture of the neural structure, the structure of the tree associated to the circuit.

4 A Sorting Algorithm

We pass now to a different problem SN P systems can solve, namely to sort n natural numbers, this time not using the rules in the exhaustive way, but as in the original definition of such systems.

We first exemplify our sorting procedure through an example. Let us presume we want to sort the natural numbers 1, 3, and 2, given in this order. For that we construct the following system given only in its pictorial format below:



Figure 3. Sorting three natural numbers

We encode natural numbers in the number of spikes (1 - one spike, 3 - threespikes, 2 - two spikes) which we input in the first line of the system (hence in the neurons labeled i_1 , i_2 , an i_3). It can be noticed that the neurons in the first layer of the structure are having the same rule inside $(a^*/a \to a; 0)$ and outgoing synapses to all the neurons in the second layer of the structure (the ones denoted s_1 , s_2 , and s_3). Neuron labeled s_1 has outgoing synapses with all neurons in the third layer of the system, only one spiking rule inside $(a^3 \to a; 0, \text{ where } 3 \text{ is the}$ number of numbers that have to be sorted), and two deletion rules $(a^2 \to \lambda, \text{ and}$ $a \to \lambda$). For the other neurons in the second layer, the exponent of the firing rule decreases one by one as well as the synapses with the neurons from the third layer of the system.

In the initial configuration of the system we have one spike in neuron i_1 , three spikes in neuron i_2 and 2 spikes in neuron i_3 . In the *first step* of the computation, one spike from each neuron is consumed and sent to neurons from the second layer of the system. Each of them receives the same number of spikes, namely 3.

In the second step of the computation, neuron labeled s_1 consumes all three spikes previously received and fires to neurons o_1 , o_2 and o_3 . Hence, each neuron from the output layer has one spike inside. The other neurons from the second layer delete the three spikes they have received. In the same time neurons i_2 and i_3 fire again sending 2 spikes (one each) to all neurons from the second layer.

In the *third step* of the computation, neuron s_2 fires only to neurons o_2 and o_3 (so, they will have one more spike inside, hence 2, while o_1 remains with only one spike), the other spikes from neurons s_1 and s_3 being deleted. In the same time neuron i_2 refills the neurons in the second layer of the system with one spike, which will be consumed in the *forth step* of the computation by neuron s_3 and sent to the output neuron o_3 .

So, in the last step of the computation there are: 1 spike in the neuron o_1 , 2 spikes in the neuron o_2 , and 3 spikes in the neuron o_3 .

We pass now to the general case, constructing the system in the pictorial form:



Figure 4. Sorting *n* natural numbers

The functioning of the system is similar to the one described in the example above. We introduce n natural numbers encoded as spikes, one in each neuron from the first layer of the structure (denoted by i_j , with $1 \le j \le n$). As long as they are not empty they consume at each step a spike, and send n spikes, one to each neuron from the second layer of the structure (denoted by s_i , with $1 \le i \le n$). The latter neurons have n different thresholds (decreasing one by one from n – neuron labeled s_1 , to 1 – neuron labeled s_n), and have n different number of synapses with the neurons from the third layer of the structure. The latter ones contain the result of the computation.

Theorem 2. SN P systems can sort a vector of natural numbers where each number is given as number of spikes introduced in the neural structure.

Based on the above construction, the time complexity (measured as usually as the number of configurations reached during the computation) is O(T), where Tis the magnitude of the numbers to be sorted. Although the time complexity is better than the "classical", sequential algorithm, in this case one can notice that the construction presented depends on the number of numbers to be sorted.

5 Final Remarks

Spiking neural P systems are a versatile formal model of computation that can be used for designing efficient parallel algorithms for solving known computer science problems. Here we firstly studied the ability of SN P systems to simulate Boolean circuits since, apart for being a well known computational model, there exists many "fast" algorithms solving various problems. In addition, this simulation, enriched with some "memory modules" (given in the form of some SN P sub-systems), may constitute an alternative proof of the computational completeness of the model.

Another issue studied here regards the sorting of a vector of natural numbers using SN P systems. In this case, due to its parallel features, the obtained time complexity for the proposed algorithm overcome the classical sequential ones.

Several open problems arose during our research. For instance, in case of Boolean circuits the simulation is done for such circuits whose underlying graphs have rooted tree structures, therefore a constraint that need further investigations.

In what regards the sorting algorithm, the presented construction depends on the magnitude of the numbers to be sorted. We conjecture that this inconvenient might be eliminated. Also, we conjecture that further improvements concerning time complexity can be made.

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Plain Talk About the Language-Theoretic Models of Multi-Agent Systems

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Summary. The contribution argues for the proposition that formal models based on the theory of formal grammars and languages are adequate for the study of some computationally relevant properties of agents and multi-agent systems. Some questions are formulated concerning the possibilities to enlarge the universality and realism of such models by considering the possibilities to go with their computing abilities beyond the traditional Turing-computability, and by considering very natural properties of any real (multi-)agent system such as not fully predictable functioning (behavior) of agents, their unreliability, dysfunctions, etc.

1 Introduction

Checking the hypothesized list of different areas of the use of different types of formal language-theoretic framework we easily realize that the language-theoretic paradigm works well. This impression seems to be true. However, on the other side of the coin, there is the question of limitations. From the methodology of science we know very well that each modeling framework is in certain sense limited by its own descriptive and predictive boundaries.

A highly challenging field for constructing predictively productive formal models is the field of agents and multi-agent systems (agencies). The reason of the increased interest consists in the fact, that the agent perspective seems to be very effective position for study of a very large spectrum of systems. The second reason consists perhaps in the fact that good, really applicable formal models of agents and agencies are up to now very rear. On the other hand, we must consider very seriously the appeals from the practice, like the one formulated by a recognized specialist in advanced robotics and artificial intelligence, Rodney Brooks: We have become very good at modeling fluids, materials, planetary dynamics, nuclear explosions and all manner of physical systems. Put some parameters into the program, let it crank, and out come accurate predictions of the physical character of modeled system. But we are not good at modeling living systems, at small or large scales. Something is wrong. What is wrong? There are a number of possibilities: (1) we might just be getting a few parameters wrong; (2) we might be building models that are below some complexity threshold; (3) perhaps it is still a lack of computing power; and (4) we might be missing something fundamental and currently unimaginable in our models (Brooks, 2001). OK, but what fundamental we have missing? And why? And how to build the better models?

We must be also sensitive to the skepticism of some of our respected colleagues. It is necessary to take seriously the opinion of Marvin Minsky one among the founders of theoretical computer science and artificial intelligence research, who said: If a theory is very simple, you can use mathematics to predict what it'll do. If it's very complicated, you have to do a simulation (Kruglinski, 2007). We must pose the question: OK, but what to do when the problems are somewhere in between?

This contribution focuses on two among the number of different languagetheoretic models developed during the decades of efforts. Namely, it focuses to the bio-chemically inspired models built up on the idea of membrane activities in living structures, and to the sort of models inspired by distributed and multiagent systems, which have the form of systems of traditional formal grammars which work together (cooperate, compete,) during the process of derivations. We will sketch the close relation of the membrane and multi-grammar models to the field of agents and agencies first, and then we will sketch some questions on the universality and realism of the models. (Unfortunately, without answers.)

2 Agents and Agencies

In the broadest meaning *agent* is any active entity, which is able to sense its environment, and to act in it according to the sensed pattern; cf. e.g. (Ferber, 1999). In (Kelemen, 2006) we provide a taxonomy of different types of agents created according the levels of complicatedness of the generation of appropriate actions on the base of patterns sensed by them in the moment of action (this is the case of the so-called purely reactive agents) or sequences of patterns sensed during the history of their activities and processed by specific inference procedures (deliberations) inside the agents' structures (this is the case of the so-called deliberative agents).

Consideration of any active thing as an agent, it is a very general perspective. From this perspective, agents at different levels of complexity are human beings, computer programs, living cells, social or economic organizations, etc., for instance. So, to find a universal theoretical framework for dealing with such a large spectrum of active entities is really a hard problem. However, we have at hand some appealing approaches. Let us to mention at least two of them.

Systems consisting of biochemical membranes and active entities inside the regions bounded by these membranes can be considered as agencies. The active entities inside are from such perspective the agents belonging to the agency. The agents, (bio-)chemical structures in their substance, act in their unstructured environments (acting means (bio)-chemical reactions in real situations, and the unstructured nature of the environment is modeled by the multi-set of symbols instead of the strings of symbols), and through the membranes the results of their activities changes the environment in neighboring regions (it can defines structure in the environment in certain sense and certain extent because of generating strings of symbols in the environment from previously isolated symbols), where other agents react to the chemical conditions in their environment, etc. Note two important points in this context. First, that in this case, the environment in which the agents act is unstructured and the activities of agents contribute to the emergence of some (local) structures of the environment. Second, that environments can be organized like the (semi-)Chinese box, or (semi-)Russian dolls (the environment of the agent B on the Fig. 1 is the part of the hierarchically higher environment of the agent A, while the environments of A and C are in the same level of the hierarchy in the environment of E.

These changes of the environment(s) caused by the agents' activities might be interpreted in the computationalistic framework as computation, and, in the consequence, the membrane structures might be considered as specific type of computing devices. The systematic study of this type of computation using the language-theoretic framework, is presented in the form of a monograph in (Păun, 2002).

In the model, there exist two types of communication between regions isolated by membranes: The firs type is the communication between the "equal" regions, for instance between the regions A and C through the shared environment E in Fig. 1. The second type of the communication is the communication like between the regions A and B. where the environment A is in fact the environment in which the region B is included and isolated by the corresponding membrane.



Fig. 1. A schematic view of a membrane system consisting of four regions, A, B, C and E isolated by membranes.

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Another example of the use of the agent paradigm is the view of the mind from the perspective of Marvin Minsky's society theory of mind (Minsky, 1986). In this case, the components of the brain/mind machinery are considered as agents and agencies. At the lowest level of the hierarchy are perhaps the neurons. Then as agents are considered the agencies formed from neurons (as some anatomical parts of the brain structure), then other agents are the agencies appearing as psycho-physiological regions of the brain, playing roles in the formation of psychic activities, etc. Note that the neurons might be considered as membrane structures, and in such a simple way, the two models are in fact interconnected and the agent paradigm works as a unifying framework for the whole spectra of activities starting somewhere down by the (bio-)chemical ones up to the psychic on the top. To provide a unifying formal framework for theoretical study with respect to the needs of practice, seems to be a great appeal, and the multi-grammar models (as the membrane system or (eco-)grammar system approaches) are promising candidates for such framework. From the architectural point of view (see Fig. 2), the agents are in the case of the society theory of mind organized in pseudopyramidal hierarchies. Some parts o such hierarchies might be considered - from an outside observers point of view, or from the point of view of their functioning - as agents. The interactions between agents are changeable, what enables to consider in the framework of the society of mind phenomena like learning, remembering, evolution, and similar cognitive processes. Moreover, the theory connects, as it is presented in (Minsky, 2006), in an interesting and inspiring way the cognitive and the emotive parts of the functioning of mind.



Fig. 2. A schematic view of the organization of agents in the society theory of mind. I states for input agents (sensors), O for the output agents (actuators).

The above sketched concepts of agents might be described in different formalisms and then studied in different formalizations. Each formalization emphasizes some of the aspects af the real systems, and suppress others.

In our context, the most interesting are the formal models of agents and agencies constructed in the framework of the theory of formal languages and formal grammars, as presented e.g. in (Csuhaj-Varju et al., 1994) or (Păun, 2002). In the case of grammar-systems, comparing them with membrane systems, the model of environment is supposed to be strictly structured, it is modeled by a string of symbols instead of the model of environment as a multi-set of symbols, as it is in the case of membrane systems. As documented e.g. in (Păun, Salomaa, 1999), there are many results concerning the formal language or the grammar-systems-like models. And the situation is similar in relation to other models, too.

In the prevailing majority of formal models, however, the answers to the questions appearing very naturally inside the framework used for modeling, have values first of all inside the conceptual framework of the model formalism. In the case of grammar systems there are questions inherent in the theory of formal grammars and languages concerning the relation of language hierarchies to the traditional Chomsky hierarchy, and questions generated then with respect the number of components sufficient or necessary to generate some given (and theoretically important) families of languages, etc. In such a way, the models built originally for studying multi-agent systems contribute to the enlargement and further development of the traditional formal language theory. It is important for the development of this theory, first of all. But *how to proceed in order to help with result of such theoretical model to the better understanding of the field of real multi-agent systems?*

3 Universality

Before starting with searching the related questions to that, formulated at the end of the previous section, and looking for the appropriate from of answers, we will deal in short with the question of the universality of multi-agent approaches to describing the different parts of the reality of some processes and phenomena. According to (D'Inverno, Luck, 2004), agents offer an abstraction tool, or a metaphor, for the design and construction of complex systems with multiple distinct and independent components. These abstractions can be used in the design and development of large systems, of individual agents, of ways in which agents may interact to support these concepts, and in the consideration of societal or macro-level issues such as organizations and their computational counterparts. They also enable the aggregation of different functionalities that have previously been distinct in a conceptually embodied and situated whole. So, we may ask why the multi-agent approach is an adequate point of view, and where are the limits of its successful use.

As we have mentioned in the previous section, the agent approach is interesting and attractive, because it provides a unifying view to the processes running in the (bio-)chemical level through many interesting branches of the human scientific and engineering interests, e.g. up to the reflection (some parts of) processes running in the level of (human) consciousness. Moreover, this view unifies in an appropriate way the biological, psychic, social, and technical systems. All the mentioned types (and many other types) of systems might be considered as systems composed from (smaller or larger) number of active, effectively isolable components with their own specific behaviors. The behaviors of the whole systems then emerge (in a more or less predictable ways) from the behaviors (often not coordinated, not centrally governed or managed) ways from the behaviors of the component agents; cf. (Kelemen, 2001).

In the cases of above mentioned formal models of membrane computing, of grammar systems, and of eco-grammar systems, and more generally, in all the cases when the models are built on the conceptual base of formal grammars and languages, the rules governing the dynamics of the behavior of agent-like entities are described in the form of rewriting rules. This is an advantage, because this formulation of the rules is in fact a formulation which defines a (trivially simple, but it is not a disadvantage!) agents (in the meaning we have accepted at the beginning): Each rule has its own sensor capacity (to sense the appearance of its left-hand side string), and an action capacity to make a change in its environment (to rewrite the sensed pattern to the for defined by the rule's right hand side). The ways of rules interactions are specified by different derivation modes and rewriting regulations.

This is, at least from the methodological point of view, a fundamental advantage. We know very well, that some specific multi-agent systems (formal grammars) define very well-specified behaviors (formal languages) with very interesting relation to different models of computation (to different types of automat) which have very important relations to real engineered (computing) machines. What we do not know it is the question of the universality of the approach accepted for describing languages (behaviors). What kind of behaviors are we able to describe using the just described framework behind the Turing-computable ones?

The second question follows from inclusion of the dynamics of the environments in which our trivial agents act. In the traditional formal language theory we do not consider any dynamics of the strings under rewriting. The only changes are those executed as rewriting activities of (some of) the rules. In the case of eco-grammar systems, however, the situation is slightly modified, because of providing an "independent" dynamics of the environment changes using a specific parallel rewriting mechanism (modeled by L-systems) working independently on the agents activities. What will happen when more complicated mechanisms of changes will be included into the models? What we know on the situation, for instance, when some finite subsequences (belonging to a language with specific Turing-computability properties) will be randomly replaced by words from another set of words (of known Turing-computability property)? How to proceed in the case of multi-sets used as models of the environments in the case of membrane systems? Change the number of (some) symbols as the result of applying a non-computable function to generate the changes, for instance?

Of course, there are many more similar questions which can be formulated in a more or less formal ways. We provide some examples only, but related to very actual themes in present days theoretical computer science, too, as documented in (Burgin, Klinger, 2004), for instance.

The most fundamental question, according our meaning, is the following one: Is it possible, and if yes, in what cases, and under what condition, to receive (define) some stabilized (well defined in the framework of Turing-computability, for instance) behavior in the hardly-predictable behavior of the environment in which the agents act? From the standpoint of the practice, this question is very important! To design such stabilizing multi-agent systems working in the unstable environment is often the main goal of many engineering activities. What we are able to say about the possibility of such design in our theoretical framework?

The last question leads us from the speculations about the universality of our models to the question of their realism.

4 Realism

In this Section we will continue the provocations contained in (Di Nola et al., 2004). However, our intention is not to continue in the development of inspiring ideas contained in it. We are much more modest - we try to enlarge, if we will be successful, the number of motivations for some questions connected with the study of realism of grammar-theoretic models of real systems. We note, that systems are in our context the systems composed from agents. We construct conceptual models in order to better understand the studied real systems. We formalize our conceptual models in order to receive rigorously predictive power of our conceptual models, and, as a consequence of that, to have models with rigorous predictive power. The answers derived logically and in formally (logically) correct ways with mathematical rigor are truthful. OK, but are we able to formulate practically interesting questions in our theoretical formalized models? Are our models realistic in this sense? If yes, what are the inherently interesting questions for any multi-agent systems theory?

In this section we will concentrate on the problem of the broad-sense reliability of multi-agent systems. Real (embodied or software) multi-agent systems are naturally not perfectly reliable. To be more particular, let us mention some of really often appearing in multi-agent systems, and because of that practically interesting, phenomena related with reliability of (multi-agent) systems.

One among the most often is the phenomenon of disfunction of (some of the) agents which form parts of the whole system. Suppose that some of the components of a complicated machine go down. Will the whole machine work after this reduction of its components? What kind of changes will appear in its behavior after this change? How to preserve some appropriate level of the functionality of the machine (its resistance with respect of the "small" changes in its architecture)?

We see, that the parts and the *reliability* (in weaker sense) of the parts are in very close relation to the whole systems functioning.

In other type of systems, despite of the reliability of the agents, their involvement into the work of the whole system is important. In the society ants, for instance, it is practically impossible to organize the work of any particular agent. Some ants work in some time period, some of them not, and, moreover, we have absolutely no predictive power to know exactly what ant will or will not do in the next time period. This problem I will call the problem of *randomness* in multiagent systems.

Mention, that both of the sketched types of problems are imaginable in the context of membrane computing and in the case of (eco-)grammar systems as well, and that there are perhaps expressible also in some not very complicated ways mathematically. *How to do that?*

Concerning the problem of reliability, first. An approach to incorporating reliability into the multi-grammar models (like the membrane computing and the (eco-)grammar frameworks) may be inspired by the incorporation of the fuzzyapproaches into the traditional grammar-theoretic models. Is seems to be possible to fuzzyfy the rewriting rules, and in the consequence of the derived strings, and to receive formal languages az fuzzy sets, in such a way. It is possible to fuzzyfy also the components of grammar systems, or of the regions of membrane systems, and then to propagate the fuzzyness toward the generated sets of behaviors, etc. It is then possible to compare the behaviors of such models with the behavior (generative capacity) of the unfuzzyfied models. *How to define the necessary notions, and what will be the results derived from the resulting model?*

Concerning the randomness of the impact of particular components of multigrammar models to the derivative capacity of the whole systems, we mention (Wätjen, 2003) as an example of an interesting approach. The participation of the components in each derivation step is defined by a function defined on the number of derivation step with values in the superset of the components. In such a form, for each derivation step, a team - similar to that introduced in (Csuhaj-Varjú, Kelemenová, 1998) – is created from all of the components, which execute the derivation. The relation of particular forms of this team-forming function and their computational properties considerably influence the behavior of the multigrammar models. *Exactly in what ways, and in what extent?*

Another way of incorporation of dynamics of components behaviors in the models may consist in timing, as defined in (Kelemenová, 1999) for colonies, but defined also for other models, e.g. by functions defined of the length of the derivation chains. Note that the similar approaches are incorporable also into the fuzzyfied models, so it seems to be realistic also the ability to combine different models of reliability and randomness into one theoretical model. What is the most perspective way of doing that?

5 Conclusions

The language-theoretic models seem to be well inspired by large spectrum of multiagent systems, and the agent and multi-agent paradigm seems to be promising for better understanding of events and processes appearing in the real word in which different individually more or less autonomous entities (agents, in our terminology) cooperate, compete, or simply cohabitate, but inevitably participate in generation of the dynamics of the whole. We have argue that the language-theoretic models form a suitable formal framework form study of systems of the above mentioned type, and we have posed maybe too many questions the answers to which will contribute to the better understanding of at least some multi-agent phenomena. But almost no answers yet!. However, let us hope that, as Marcel Proust wrote, *each reader reads only what is inside himself. A text (book,* in the original) *is only a sort of optical instrument which the writer offers to let the reader discover in himself* So, be successful!

And remember: If you "understand" something in only one way, then you scarcely understand it at all – because when you get stuck, you'll have nowhere to go. But if you represent something in several ways, then when you get frustrated enough, you can switch among different points of view, until you find one that works for you! (Minsky, 2006).

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Solving Numerical NP-Complete Problems with Spiking Neural P Systems

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Summary. Starting from an extended nondeterministic spiking neural P system that solves the SUBSET SUM problem in a constant number of steps, recently proposed in a previous paper, we investigate how different properties of spiking neural P systems affect the capability to solve numerical **NP**-complete problems. In particular, we show that by using maximal parallelism we can convert any given integer number from the usual binary notation to the unary form, and thus we can initialize the above P system with the required (exponential) number of spikes in polynomial time. On the other hand, we show that this conversion cannot be performed in polynomial time if the use of maximal parallelism is forbidden. Finally, we show that by *selectively* using nondeterminism and maximal parallelism (that is, for each neuron in the system we can specify whether it works in deterministic or nondeterministic way, as well as in sequential or maximally parallel way) there exists a *uniform* spiking neural P system that solves all the instances of SUBSET SUM of a given size.

1 Introduction

Membrane systems (also called *P systems*) were introduced in [16] as a new class of distributed and parallel computing devices, inspired by the structure and functioning of living cells. The basic model consists of a hierarchical structure composed by several membranes, embedded into a main membrane called the *skin*. Membranes divide the Euclidean space into *regions*, that contain some *objects* (represented by symbols of an alphabet) and *evolution rules*. Using these rules, the objects may evolve and/or move from a region to a neighboring one. Usually, the rules are applied in a nondeterministic and maximally parallel way; moreover, all the objects that may evolve are forced to evolve. A *computation* starts from an initial configuration of the system and terminates when no evolution rule can be applied. The result of a computation is the multiset of objects contained into an *output* *membrane*, or emitted to the environment from the skin of the system. For a systematic introduction to P systems we refer the reader to [18], whereas the latest information can be found in [23].

In an attempt to pass from cell-like to tissue-like architectures, in [14] tissue P systems were defined, in which cells are placed in the nodes of a (directed) graph. Since then, this model has been further elaborated, for example, in [4] and [21], with recent results about both theoretical properties [1] and applications [15]. This evolution has led to explore also neural-like architectures, yielding to the introduction of spiking neural P systems (SN P systems, for short) [8], based on the neurophysiological behavior of neurons sending electrical impulses (spikes) along axons to other neurons. We recall that this biological background has already led to several models in the area of neural computation, e.g., see [12, 13, 6].

Similarly to tissue P systems, in SN P systems the cells (neurons) are placed in the nodes of a directed graph, called the synapse graph. The contents of each neuron consist of a number of copies of a single object type, called the *spike*. The firing rules assigned to a cell allow a neuron to send information to other neurons in the form of electrical impulses (also called spikes) which are accumulated at the target cell. The application of the rules depends on the contents of the neuron; in the general case, applicability is determined by checking the contents of the neuron against a regular set associated with the rule. As inspired from biology, when a cell sends out spikes it becomes "closed" (inactive) for a specified period of time, that reflects the refractory period of biological neurons. During this period, the neuron does not accept new inputs and cannot "fire" (that is, emit spikes). Another important feature of biological neurons is that the length of the axon may cause a time delay before a spike arrives at the target. In SN P systems this delay is modeled by associating a delay parameter to each rule which occurs in the system. If no firing rule can be applied in a neuron, there may be the possibility to apply a *forgetting rule*, that removes from the neuron a predefined number of spikes.

In the original model of SN P systems defined in [8], computations occur as follows. A configuration specifies, for each neuron of the system, the number of spikes it contains and the number of computation steps after which the neuron will become "open" (that is, not closed). Starting from an initial configuration, a positive integer number is given in input to a specified *input neuron*. The number is encoded as the interval of time steps elapsed between the insertion of two spikes into the neuron (note that this is a unary encoding). To pass from a configuration to another one, for each neuron a rule is chosen among the set of applicable rules, and is executed. The computation proceeds in a sequential way into each neuron, and in parallel among different neurons. Generally, a computation may not halt. However, in any case the output of the system is considered to be the time elapsed between the arrival of two spikes in a designated *output cell*. Defined in this way, SN P systems compute functions of the kind $f : \mathbb{N} \to \mathbb{N}$ (they can also indirectly compute functions of the kind $f : \mathbb{N}^k \to \mathbb{N}$ by using a bijection from \mathbb{N}^k to \mathbb{N}). By neglecting the output neuron we can define *accepting* SN P systems, in which the natural number given in input is accepted if the computation halts. On the other hand, by ignoring the input neuron (and thus starting from a predefined input configuration) we can define *generative* SN P systems.

In [8] it was shown that generative SN P systems are universal, that is, can generate any recursively enumerable set of natural numbers. Moreover, a characterization of semilinear sets was obtained by spiking neural P systems with a bounded number of spikes in the neurons. These results can also be obtained with even more restricted forms of spiking P systems; for example, [7] shows that at least one of these features can be avoided while keeping universality: time delay (refractory period) greater than 0, forgetting rules, outdegree of the synapse graph greater than 2, and regular expressions of complex form. Finally, in [19] the behavior of spiking neural P systems on infinite strings and the generation of infinite sequences of 0 and 1 was investigated, whereas in [2] spiking neural P systems were studied as language generators (over the binary alphabet $\{0, 1\}$).

In [10] we have shown that by slightly extending the original definition of SN P system given in [8] and [9], it is possible to solve any given instance of SUBSET SUM by using a nondeterministic (extended) SN P system. The solution is given in the so called *semi-uniform* setting, that is, for every fixed instance of SUBSET SUM a specific SN P system that solves it is built. In particular, the rules of the system and the number of spikes which occur in the initial configuration depend upon the instance to be solved. A drawback of this solution is that in general the number of spikes needed to initialize the system is exponential with respect to the usually agreed instance size of SUBSET SUM. However, in this paper we show that this preparation can be performed in polynomial time by traditional SN P systems that, endowed with the power of maximal parallelism, read from the environment the k-bit integer numbers v_1, v_2, \ldots, v_n encoded in binary and produce v_1, v_2, \ldots, v_n spikes, respectively, in n specified neurons. We also prove that this operation cannot be performed in polynomial time if the use of maximal parallelism is forbidden. Then we design an SN P system that performs the opposite conversion: it takes a given (k-bit) number N of spikes occurring in a certain neuron, and produces the coefficients of the binary encoding of N in k predefined neurons. Thanks to these two modules, that allow us to move from binary to unary encoding and back, we finally design a uniform family $\{\Pi(\langle n,k \rangle\}_{n,k \in \mathbb{N}})$ of SN P systems, where $\Pi(\langle n, k \rangle)$ solves all possible instances $(\{v_1, v_2, \ldots, v_n\}, S)$ of SUBSET SUM such that all v_i and S are k-bit positive integer numbers. As we will see, the construction of $\Pi(\langle n, k \rangle)$ relies upon the assumption that different subsystems can work under different regimes: deterministic vs. nondeterministic, and sequential vs. maximally parallel.

The rest of this paper is organized as follows. In section 2 we give some mathematical preliminaries, and we define the standard version of SN P systems (as found in [9]) as well as a slightly extended version. In section 3 we recall from [10] how the **NP**-complete problem SUBSET SUM can be solved in constant time by exploiting nondeterminism in our extended SN P systems. In section 4 we convert positive integer numbers from binary notation to the unary form through maximally parallel SN P systems, and we use such a convertion as an initialization stage to solve SUBSET SUM. In section 5 we perform also the opposite conversion, and we design a family of SN P systems that solves SUBSET SUM in a uniform way (according to the above definition). Section 6 concludes the paper and gives some directions for future research.

2 Preliminaries

Let us start by recalling the standard definition of a spiking neural P system, taken from [9]. A spiking neural membrane system (SN P system, for short), of degree $m \ge 1$, is a construct of the form

$$\Pi = (O, \sigma_1, \sigma_2, \dots, \sigma_m, syn, in, out),$$

where:

- 1. $O = \{a\}$ is the singleton alphabet (a is called *spike*);
- 2. $\sigma_1, \sigma_2, \ldots, \sigma_m$ are *neurons*, of the form $\sigma_i = (n_i, R_i)$, with $1 \le i \le m$, where: a) $n_i \ge 0$ is the *initial number of spikes* contained in σ_i ;
 - b) R_i is a finite set of *rules* of the following two forms:
 - (1) $E/a^c \to a; d$, where E is a regular expression over a, and $c \ge 1, d \ge 0$ are integer numbers; if $E = a^c$, then it is usually written in the following simplified form: $a^c \to a; d;$
 - (2) $a^s \to \lambda$, for $s \ge 1$, with the restriction that for each rule $E/a^c \to a; d$ of type (1) from R_i , we have $a^s \notin L(E)$ (where L(E) denotes the regular language defined by E);
- 3. $syn \subseteq \{1, 2, ..., m\} \times \{1, 2, ..., m\}$, with $(i, i) \notin syn$ for $1 \leq i \leq m$, is the directed graph of synapses between neurons;
- 4. $in, out \in \{1, 2, ..., m\}$ indicate the *input* and the *output* neurons of Π .

The rules of type (1) are called *firing* (also *spiking*) rules, and they are applied as follows. If the neuron σ_i contains $k \geq c$ spikes, and $a^k \in L(E)$, then the rule $E/a^c \to a; d \in R_i$ can be applied. The execution of this rule removes c spikes from σ_i (thus leaving k - c spikes), and prepares one spike to be delivered to all the neurons σ_j such that $(i, j) \in syn$. If d = 0, then the spike is immediately emitted, otherwise it is emitted after d computation steps of the system. (Observe that, as usually happens in membrane computing, a global clock is assumed, marking the time for the whole system, hence the functioning of the system is synchronized.) If the rule is used in step t and $d \geq 1$, then in steps $t, t + 1, t + 2, \ldots, t + d - 1$ the neuron is *closed*, so that it cannot receive new spikes (if a neuron has a synapse to a closed neuron and tries to send a spike along it, then that particular spike is lost), and cannot fire new rules. In the step t + d, the neuron spikes and becomes open again, so that it can receive spikes (which can be used starting with the step t + d + 1) and select rules to be fired. Rules of type (2) are called *forgetting* rules, and are applied as follows: if the neuron σ_i contains *exactly* s spikes, then the rule $a^s \to \lambda$ from R_i can be used, meaning that all s spikes are removed from σ_i . Note that, by definition, if a firing rule is applicable then no forgetting rule is applicable, and vice versa.

In each time unit, if a neuron σ_i can use one of its rules, then a rule from R_i must be used. Since two firing rules, $E_1 : a^{c_1} \to a; d_1$ and $E_2 : a^{c_1} \to a; d_2$, can have $L(E_1) \cap L(E_2) \neq \emptyset$, it is possible that two or more rules can be applied in a neuron. In such a case, only one of them is chosen nondeterministically. Thus, the rules are used in the sequential manner in each neuron, but neurons function in parallel with each other.

The *initial configuration* of the system is described by the numbers n_1, n_2, \ldots , n_m of spikes present in each neuron, with all neurons being open. During the computation, a configuration is described by both the number of spikes present in each neuron and by the state of each neuron, which can be expressed as the number of steps to count down until it becomes open (this number is zero if the neuron is already open). A *computation* in a system as above starts in the initial configuration. In order to compute a function $f: \mathbb{N}^k \to \mathbb{N}$, one possibility is to introduce k natural numbers n_1, n_2, \ldots, n_k in the system by "reading" from the environment a binary sequence $z = 0^b 10^{n_1} 10^{n_2} 1 \dots 10^{n_k} 10^g$, for some b, q > 0; this means that the input neuron of Π receives a spike in each step corresponding to a digit 1 from the string z. Note that we input exactly k + 1 spikes. The result of the computation is also encoded in the distance between two spikes: we impose to the system to output exactly two spikes and halt (sometimes after the second spike) hence producing a spike train of the form $0^{b'}10^r 10^{g'}$, for some $b', g' \ge 0$ and with $r = f(n_1, n_2, \ldots, n_k)$. As discussed in [9], there are other possibilities to encode natural numbers read from and/or emitted to the environment by SN P systems; for example, we can consider the number of spikes arriving to the input neuron and leaving from the output neuron, respectively, or the number of spikes read/produced in a given interval of time.

If we do not specify an input neuron (hence no input is taken from the environment) then we use SN P systems in the *generative* mode; we start from the initial configuration, and the distance between the first two spikes of the output neuron (or the number of spikes, etc.) is the result of the computation. Note that generative SN P systems are inherently nondeterministic, otherwise they would always reproduce the same sequence of computation steps, and hence the same output. Dually, we can neglect the output neuron and use SN P systems in the *accepting* mode; for $k \geq 1$, the natural number n_1, n_2, \ldots, n_k are read in input and, if the computation halts, then the numbers are accepted.

We define the *description size* of an SN P system Π as the number of bits which are necessary to describe it. Since the alphabet O is fixed, no bits are necessary to define it. In order to represent syn we need at most m^2 bits, whereas we can represent the values of *in* and *out* by using log *m* bits each. For every neuron σ_i we have to specify a natural number n_i and a set R_i of rules. For each rule we need to specify its type (firing or forgetting), which can be done with 1 bit, and in the worst case we have to specify a regular expression and two natural numbers. If we denote by N the maximum natural number that appears in the definition of Π , R the maximum number of rules which occur in its neurons, and S the maximum size required by the regular expressions that occur in Π (more on this later), then we need a maximum of $\log N + R(1 + S + 2\log N)$ bits to describe every neuron of Π . Hence, to describe Π we need a total of $m^2 +$ $2\log m + m(\log N + R(1 + S + 2\log N))$ bits. Note that this quantity is polynomial with respect to m, R, S and $\log N$. Since the regular languages determined by the regular expressions that occur in the system are *unary* languages, the strings of such languages can be bijectively identified with their lengths. Hence, when writing the regular expression E, instead of writing unions, concatenations and Kleene closures among strings we can do the same by using the lengths of such strings. (Note that, when concatenating two languages L_1 and L_2 represented in this way, the lengths in L_1 are summed with the lengths of L_2 by combining them in all possible ways). In this way we obtain a representation of E which is *succint*, that is, exponentially more compact than the usual representation of regular expressions. As we have seen in [10], this succint representation yields some difficulties when we try to simulate a deterministic accepting SN P system that contains general regular expressions, by a deterministic Turing machine. However, as shown in [7], it is possible to restrict our attention to particularly simple regular expressions, without loosing computational completeness. For these expressions, the membership problem (is a given string into the language generated by the regular expression?) is polynomial also when representing the instances in succint form, and thus they do not yield problems when simulating the system with a deterministic Turing machine.

In what follows it will be convenient to consider also the following slightly extended version of SN P systems. Precisely, we will allow rules of the type $E/a^c \rightarrow a^p; d$, where $c \ge 1, p \ge 0$ and $d \ge 0$ are integer numbers. The semantics of this kind of rules is as follows: if the contents of the neuron matches the regular expression E, then the rule can be applied. When the rule is applied, c spikes are removed from the contents of the neuron and p spikes are prepared to be delivered to all the neurons which are directly connected (through an arc of syn) with the current neuron. If d = 0, then these p spikes are immediately sent, otherwise the neuron becomes closed for the neuron does not receive spikes from other neurons, and does not apply any rule. If p = 0, then we obtain a forgetting rule as a particular case of our general rules.

Also in the extended SN P systems it may happen that, given two rules $E_1/a^{c_1} \rightarrow a^{p_1}$; d_1 and $E_2/a^{c_2} \rightarrow a^{p_2}$; d_2 , if $L(E_1) \cap L(E_2) \neq \emptyset$ then for some contents of the neuron both the rules can be applied. In such a case, one of them is nondeterministically chosen. Note that we do not require that forgetting rules are applied only when no firing rule can be applied. We say that the system is *deterministic* if, for every neuron that occurs in the system, any two rules $E_1/a^{c_1} \rightarrow a^{p_1}$; d_1 and $E_2/a^{c_2} \rightarrow a^{p_2}$; d_2 in the neuron are such that $L(E_1) \cap L(E_2) = \emptyset$. This means

that, for any possible contents of the neuron, at most one of the rules that occur in the neuron may be applied.

By using an *input neuron* and an *output neuron*, we have SN P systems that compute functions of the kind $f : \mathbb{N} \to \mathbb{N}$ (as well as functions of the kind $f : \mathbb{N}^k \to \mathbb{N}$, by appropriate bijections between \mathbb{N}^k and \mathbb{N}), and hence we cover both the generative and the accepting cases. If out = 0, then it is understood that the output is sent to the environment (as the number of spikes produced by the system, as the distance between the first two spikes, etc.). As usual, to use an SN P system in the generative mode we do not consider the input neuron, whereas by ignoring the output neuron we obtain an accepting SN P system.

The *description size* of an extended SN P system is defined exactly as we have done for standard systems, the only difference being that now we require (at most) three natural numbers to describe a rule.

3 Solving Numerical NP–complete Problems with Extended Spiking Neural P Systems

Let us start by recalling the nondeterministic extended SN P system introduced in [10] to solve the **NP**-complete problem SUBSET SUM in a *constant* number of computation steps. The SUBSET SUM problem can be defined as follows.

Problem 1. NAME: SUBSET SUM.

- INSTANCE: a (multi)set $V = \{v_1, v_2, \dots, v_n\}$ of positive integer numbers, and a positive integer number S.
- QUESTION: is there a sub(multi)set $B \subseteq V$ such that $\sum_{b \in B} b = S$?

If we allow to nondeterministically choose among the rules which occur in the neurons, then the extended SN P system depicted in Figure 1 solves any given instance of SUBSET SUM in a constant number of steps. We emphasize the fact that such a solution occurs in the *semi-uniform* setting, that is, for every instance of SUBSET SUM we build an SN P system that specifically solves that instance.

Let $(V = \{v_1, v_2, \ldots, v_n\}, S)$ be the instance of SUBSET SUM to be solved. In the initial configuration of the system, the leftmost neurons contain (from top to bottom) v_1, v_2, \ldots, v_n spikes, respectively, whereas the rightmost neurons contain zero spikes each. In the first step of computation, in each of the leftmost neurons of the SN P system depicted in Figure 1 it is nondeterministically chosen whether to include or not the element v_i in the (candidate) solution $B \subseteq V$; this is accomplished by nondeterministically choosing among one rule that forgets v_i spikes (in such a case, $v_i \notin B$) and one rule that propagates v_i spikes to the rightmost neurons. At the beginning of the second step of computation a certain number N = |B| of spikes, that corresponds to the sum of the v_i which have been chosen, occurs in the rightmost neurons. We have three possible cases:



Fig. 1. A nondeterministic extended SN P system that solves the SUBSET SUM problem in constant time

- N < S: in this case neither the rule a*/a^S → a; 0 nor the rule a*/a^{S+1} → a; 1 (which occur in the neuron at the top and at the bottom of the second layer, respectively) fire, and thus no spike is emitted to the environment;
- N = S: only the rule $a^*/a^S \rightarrow a$; 0 fires, and emits a single spike to the environment. No further spikes are emitted;
- N > S: both the rules $a^*/a^S \to a; 0$ and $a^*/a^{S+1} \to a; 1$ fire. The first rule immediately sends one spike to the environment, whereas the second rule sends another spike at the next computation step (due to the delay associated with the rule).

Hence, by counting the number of spikes emitted to the environment at the second and third computation steps we are able to read the solution of the given instance of SUBSET SUM: the instance is positive if and only if a single spike is emitted.

The proposed system is generative; its input (the instance of SUBSET SUM to be solved) is encoded in the initial configuration. We stress once again that the ability to solve SUBSET SUM in constant time derives from the fact that the system is nondeterministic. As it happens with Turing machines, nondeterminism can be interpreted in two ways: (1) the system "magically" chooses the correct values v_i (if they exist) that allow to produce a single spike in output, or (2) at least one of the possible computations produces a single spike in output.

The formal definition of the extended (generative) SN P system depicted in Figure 1 is as follows:

$$\Pi = (\{a\}, \sigma_1, \ldots, \sigma_{n+2}, syn, out),$$

where:

- $\sigma_i = (v_i, \{a^{v_i} \to \lambda, a^{v_i} \to a^{v_i}; 0\})$ for all $i \in \{1, 2, \dots, n\};$
- $\sigma_{n+1} = (0, \{a^*/a^S \to a; 0\});$
- $\sigma_{n+2} = (0, \{a^*/a^{S+1} \to a; 1);$
- $syn = \bigcup_{i=1}^{n} \{(i, n+1), (i, n+2)\};$
- out = 0 indicates that the output is sent to the environment.

However, here we are faced with a problem that we have already met in [11], and that we will meet again in the rest of the paper. In order to clearly expose the problem, let us consider the following algorithm that solves SUBSET SUM using the well known Dynamic Programming technique [3]. In particular, the algorithm returns 1 on positive instances, and 0 on negative instances.

 $\begin{array}{l} \underline{\text{SUBSET SUM}}(\{v_1,v_2,\ldots,v_n\},S)\\ \text{for }j\leftarrow 0 \text{ to }S\\ \text{ do }M[1,j]\leftarrow 0\\ M[1,0]\leftarrow M[1,v_1]\leftarrow 1\\ \text{for }i\leftarrow 2 \text{ to }n\\ \text{ do for }j\leftarrow 0 \text{ to }S\\ \text{ do }M[i,j]\leftarrow M[i-1,j]\\ \text{ if }j\geq v_i \text{ and }M[i-1,j-v_i]>M[i,j]\\ \text{ then }M[i,j]\leftarrow M[i-1,j-v_i]\\ \end{array}$

In order to look for a subset $B \subseteq V$ such that $\sum_{b \in B} b = S$, the algorithm uses an $n \times (S+1)$ matrix M whose entries are from $\{0,1\}$. It fills the matrix by rows, starting from the first row. Each row is filled from left to right. The entry M[i,j] is filled with 1 if and only if there exists a subset of $\{v_1, v_2, \ldots, v_i\}$ whose elements sum up to j. The given instance of SUBSET SUM is thus a positive instance if and only if M[n, S] = 1 at the end of the execution.

Since each entry is considered exactly once to determine its value, the time complexity of the algorithm is proportional to $n(S+1) = \Theta(nS)$. This means that the difficulty of the problem depends on the value of S, as well as on the magnitude of the values in V. In fact, let $K = max\{v_1, v_2, \ldots, v_n, S\}$. If K is polynomially bounded with respect to n, then the above algorithm works in polynomial time. On the other hand, if K is exponential with respect to n, say $K = 2^n$, then the above algorithm may work in exponential time and space. This behavior is usually referred to in the literature by telling that SUBSET SUM is a *pseudo-polynomial* **NP**-complete problem.

The fact that in general the running time of the above algorithm is not polynomial can be immediately understood by comparing its time complexity with the instance size. The usual size for the instances of SUBSET SUM is $\Theta(n \log K)$, since for conciseness every "reasonable" encoding is assumed to represent each element of V (as well as S) using a string whose length is $O(\log K)$. Here all logarithms are taken with base 2. Stated differently, the size of the instance is usually considered to be the number of bits which must be used to represent in binary S and all the

integer numbers which occur in V. If we would represent such numbers using the unary notation, then the size of the instance would be $\Theta(nK)$. But in this case we could write a program which first converts the instance in binary form and then uses the above algorithm to solve the problem in polynomial time with respect to the new instance size. We can thus conclude that the difficulty of a numerical **NP**-complete problem depends also on the measure of the instance size we adopt.

The problem we mentioned above about the SN P system depicted in Figure 1 is that the rules $a^{v_i} \rightarrow \lambda$ and $a^{v_i} \rightarrow a^{v_i}$; 0 which occur in the leftmost neurons, as well as those that occur in the rightmost neurons, check for the existence of a number of spikes which may be exponential with respect to the usually agreed instance size of SUBSET SUM. Moreover, to initialize the system the user has to place a number of objects which may also be exponential. This is not fair, because it means that the SN P system that solves the **NP**-complete problem has in general an exponential effort is thus needed to build and initialize the system, that easily solves the problem by working in unary notation (hence in polynomial time with respect to the size of the system, but not with respect to its *description size*). This problem is in some aspects similar to what has been described in [11], concerning traditional P systems that solve **NP**-complete problems.

4 Solving SUBSET SUM with Inputs Encoded in Binary

Similarly to what we have done in [11], in this section we show that the ability of the SN P system depicted in Figure 1 to solve SUBSET SUM does not derive from the fact that the system is initialized with an exponential number of spikes, at least if we allow the application of rules in the maximal parallel way.

In this paper, maximal parallelism is intended exactly as in traditional P systems. Since in SN P systems we have only one kind of objects (the spike), this means that at every computation step the (multi)set of rules to be applied in a neuron is determined as follows. Let k denote the number of spikes contained in the neuron. First, one rule is nondeterministically chosen among those which can be applied. If such a rule consumes c spikes, then the selection process is repeated to the remaining k - c spikes, until no rule can be applied. Note that a rule may eventually be chosen many times, and thus at the end of the process we obtain a multiset of rules. However let us note that, for our purposes, it will suffice to define maximally parallel neurons that contain just one rule. Hence, the process with which the neuron chooses the rules to be applied is uninfluent: at every computation step the only existing rule is chosen, and is applied as many times as possible (i.e., maximizing the number of spikes which are consumed).

Consider the SN P system depicted in Figure 2, in which all the neurons work in the maximal parallel way. Assume that a sequence of spikes comes from the environment, during k consecutive time steps. Such spikes can be considered as the binary encoding of a k-bit natural number N, by simply interpreting as 1 (resp.,



Fig. 2. A maximally parallel SN P system that converts a binary encoded positive integer number to unary form

0) the presence (resp., the absence) of a spike in each time step. The system works as follows. In the first step, the most significant bit of N enters into the neuron labelled with 0. Simultaneously, neuron st fires and sends a spike to neuron out, that will contain the resulting unary encoding of N. This is done in order to close such a neuron, so that it does not receive the intermediate results produced by neurons $0, 1, \ldots, k-1$ during the conversion. During the next k-1 steps, all subsequent bits of N enter into the system. Neurons $0, 1, \ldots, k-1$ act as a shift register, and they duplicate every spike before sending both copies to the neighbouring neuron. In this way, since rules are applied in the maximally parallel way, at the end of the k-th step each neuron j, with $j \in \{0, 1, \dots, k-1\}$, will contain 2^{j} spikes if the j-th bit of N is 1, otherwise it will contain 0 spikes. At the (k+1)-th step, neuron out becomes open again, and receives exactly N spikes. Two little annoying details are that this neuron emits a "spurious" spike at the (k+1)-th computation step, and that it becomes again closed for further k-1time steps. The first spike emitted from the subsystem has obviously to be ignored, whereas during the (2k)-th step neuron out emits the N spikes we are interested in. Note that this module can be used only once, since neuron st initially contains just one spike. By making neuron st work in the sequential mode (instead of the maximally parallel mode), and slightly complicating the structure of the system, we can also convert a sequence of n numbers arriving from the environment in $n \cdot k$ consecutive time steps.

By looking at Figure 3, we can see that for any instance $(\{v_1, v_2, \ldots, v_n\}, S)$ of SUBSET SUM it is possible to build a maximally parallel nondeterministic SN P system that solves it as follows. During the first k computation steps, the system reads n sequences of spikes, each one encoding in binary the natural number v_i . Each sequence goes to an SN subsystem which performs the conversion from binary



Fig. 3. A nondeterministic SN P system that solves the SUBSET SUM problem by working in the maximal parallel way (but for the neuron Sum)

to unary, as illustrated in Figure 2. Thus in the (2k)-th step, for all $i \in \{1, 2, \ldots, n\}$, v_i spikes reach the neuron labelled with v_i . At the next step, each of these neurons *nondeterministically* decides whether to propagate the spikes it has received, or to delete them. Hence, the rules of neurons v_i are applied not only in the maximal parallel way, but also in a nondeterministic way (in the sense that one of the two rules is nondeterministically chosen, and then is applied in the maximal parallel way). In step 2k + 2, the neuron labelled with Sum checks whether the number of spikes it has gathered is equal to S; if so, it fires one spike to the environment, thus signalling that the given instance of SUBSET SUM is positive. Conversely, the instance is negative if and only if no spike is emitted from the system during the (2k+2)-nd computation step. The forgetting rules which occur in neuron Sum are needed so that at step k+2 all the spurious spikes that (eventually) reach the neuron (coming from the modules that have performed the conversions from binary to unary) are removed from the system, and are not added to the spikes that arrive at step 2k+1. Of course, here we are assuming that S > n; if this is not the case, then the rules must be modified accordingly. Note that neuron Sum is deterministic, and works in the sequential way. We also observe that, if desired, we can use two neurons instead of one in the last layer of the system, as we have done in Figure 1. The first neuron would be just like Sum, the only difference being that the rule $a^S \to a; 0$ becomes $a^*/a^S \to a; 0$. The second neuron would contain the same forgetting rules as Sum, and the firing rule $a^*/a^{S+1} \rightarrow a; 1$ instead of $a^S \rightarrow a; 0$. In this way, the instance would be signalled as positive if and only if a single spike is emitted during the steps 2k + 2 and 2k + 3.

This solution to the SUBSET SUM problem is still semi-uniform: a single system is able to solve all the instances that have the same value of S, and in which all v_i are k-bit numbers. A way to make the system uniform would be to read from the environment also the value of S, encoded in binary form, and send a corresponding number of spikes to a predefined neuron. The problem would thus reduce to comparing with S the number of spikes obtained by nondeterministically choosing some of the v_i . In the next section we will operate in a similar way; however, instead of comparing the contents of two neurons, expressed in unary form, we will operate as follows: we will keep S in binary form, and we will convert the sum of v_i from unary to binary. In this way, the problem to compare S with the sum of v_i is reduced to a bit-by-bit comparison.



Fig. 4. A maximally parallel SN P system that converts a unary encoded positive integer number to binary form

Before doing all this, let us show that the conversion from binary to unary of a given natural number cannot be performed in polynomial time without using maximal parallelism. Let Π be a deterministic SN P system that works in the sequential way: all the neurons compute in parallel with respect to each other, but in each neuron only one rule is chosen and applied at every computation step. To be precise, even if the contents of the neuron would allow to apply the chosen rule many times (such as it happens, for example, with the rule $a \to a^2$; 0 and five spikes occurring in the neuron), only one instance of the rule is applied (in the example, one spike is consumed and two spikes are produced). Without loss of generality, we can assume that the regular expressions that occur in \varPi have the form a^i with $i \leq 3$ or $a(aa)^+$, which suffice to obtain computationally complete SN P systems [7]. Let m be the number of neurons in Π , and let t(k) be the polynomial number of steps needed by Π to convert the k-bit natural number N given in input from the binary to the unary form. Moreover, let Q be the maximum number of spikes produced by any rule of Π . Since in the worst case every neuron is connected with every other neuron, the total number of spikes occurring in the system is incremented by at most mQ units during each computation step. If we denote by M the number of spikes occurring in the initial configuration, then after t(k) computation steps the number of spikes in the system will be at most M + mQt(k). This quantity is polynomial with respect to both the number of steps and the description size of Π , and thus it cannot cover the exponential gap that exists between the number of objects needed to represent N in binary and in unary form.

5 A Uniform Family of SN P Systems for SUBSET SUM

Let us present now a uniform family $\{\Pi(\langle n, k \rangle)\}_{n,k \in \mathbb{N}}$ of SN P systems such that for every n and k in \mathbb{N} , the system $\Pi(\langle n, k \rangle)$ solves all possible instances $(\{v_1, v_2, \ldots, v_n\}, S)$ of SUBSET SUM in which v_1, v_2, \ldots, v_n and S are all k-bit natural numbers.



(To the environment)

Fig. 5. The uniform SN P system $\Pi(\langle n, k \rangle)$ that solves all instances of SUBSET SUM composed by k-bit natural numbers

As told in the previous section, we first need a subsystem that allows to convert natural numbers from the unary to the binary form. Consider the system depicted in Figure 4. All the neurons work in the maximally parallel way. Initially, neuron



Fig. 6. An SN P system that delays of k steps the sequence of spikes given in input

in contains N spikes, where N is the k-bit integer number we want to convert. In the first computation step, all the spikes contained in neuron *in* are sent to neuron 0 (thus entering into the subsystem), thanks to the rule $a \rightarrow a$ applied in the maximally parallel way. In the second step, rule $a^2 \rightarrow a$ in neuron 0 halves the number of spikes (indeed, computing an integer division by 2) and sends the result to neuron 1. If the initial number of spikes was even, then in neuron 0 no spikes are left; instead, if the initial number of spikes was odd, then exactly one spike will remain in neuron 0. Hence, the number of spikes remaining in neuron 0 is equal to the value of the least significant bit of the binary encoding of N. The computation proceeds in a similar way during the next k-1 steps; in each step, the next bit (from the least significant to the most significant) of the binary encoding of N is computed. Note that the bits that have already been computed are unaffected by subsequent computation steps. After k computation steps, the neurons labelled with $0, 1, \ldots, k-1$ contain all the bits of the binary encoding of N. In order to use such bits, we can connect these neurons to other k neurons, which should be kept closed during the conversion by means of a trick similar to that used in Figure 2.

The SN P system $\Pi(\langle n, k \rangle)$ that solves all the instances $(\{v_1, v_2, \ldots, v_n\}, S)$ of SUBSET SUM which are composed by k-bit natural numbers is depicted (in a schematic way) in Figure 5. The sequences of spikes that encode v_1, v_2, \ldots, v_n and S in binary form arrive simultaneously from the environment, and enter into the system from the top. The values v_1, \ldots, v_n are first converted to unary and then some of them are summed, as before; the sequence of bits in S, instead, is just delayed (using the subsystem depicted in Figure 6) so that it arrives in the "Bit by bit comparison" subsystem simultaneously with the binary representation of the sum of the v_i . Such a binary representation is obtained through the subsystem depicted in Figure 7) emits a spike if and only if all the bits of the two integer numbers given in input match, that is, if and only if the two numbers are equal. If we denote by $x = \sum_{i=0}^{k-1} x_i 2^i$ and $y = \sum_{i=0}^{k-1} y_i 2^i$ the numbers to be compared, the subsystem computes the following boolean function:

$$\operatorname{COMPARE}(x_0,\ldots,x_{k-1},y_0,\ldots,y_{k-1}) = \bigwedge_{i=0}^{k-1} \left(\neg (x_i \oplus y_i) \right) = \neg \left(\bigvee_{i=0}^{k-1} (x_i \oplus y_i) \right)$$

where \oplus denotes the logical XOR operation. The subsystem works as follows. Bits x_i and y_i are XORed by the neurons depicted on the top of Figure 7. The neuron



Fig. 7. A standard SN P system that compares two k-bit natural numbers

labelled with \lor computes the logical OR of its inputs: precisely, it emits one spike if and only if at least one spike enters into the neuron. Neuron *out* receives the output produced by \lor and computes its logical negation (NOT). In order to be able to produce one spike if no spikes come from *out*, we use two auxiliary neurons that send to *out* one spike at every computation step. The number of neurons, as well as the total number of rules, used by $\Pi(\langle n, k \rangle)$ is polynomial with respect to n and k.

We conclude by observing that the output of the SN P system depicted in Figure 5 has to be observed exactly after 3k + 6 computation steps. One spike will eventually be emitted by the system before this time, since the conversion from binary to unary of v_1, v_2, \ldots, v_n produces some spurious spikes before emitting the result. These spurious spikes are added in neuron *sum*, and the result of this addition is first converted to binary and then sent to the comparison subcircuit. However, by carefully calibrating the delay subsystem this value does not interfere with the bits of S, that will arrive to the comparison subsystem only later. From a direct inspection of the system in Figure 5, it is easily seen that the correct delay to be applied is of 3k + 2 steps.

6 Conclusions and Directions for Future Research

In this paper we have continued the study concerning the computational power of SN P systems, started in [10]. In particular, by slightly extending the original definition of SN P systems given in [8] and [9] we have shown that by exploiting nondeterminism it is possible to solve numerical **NP**-complete problems such as SUBSET SUM and PARTITION (which can be considered as a particular case of SUBSET SUM).

However, a drawback of this solution is that the system may require to specify an exponential number of spikes both when defining the rules and when describing the contents of the neurons in the initial configuration. Hence, we have shown that the numbers v_1, v_2, \ldots, v_n occurring in the instance of SUBSET SUM can be given to the system in binary form, and subsequently converted to the unary form in polynomial time. In this way we have proved that the capability of the above system to solve SUBSET SUM does not derive from the fact that it requires an exponential effort to be initialized.

The new SN P system thus obtained still provides a *semi-uniform* solution, since for each instance of the problem we need to build a specifically designed SN P system to solve it. Thus, we have finally proposed a family $\{\Pi(\langle n, k \rangle)\}_{n,k \in \mathbb{N}}$ of SN P systems such that for all $n, k \in \mathbb{N}$, $\Pi(\langle n, k \rangle)$ solves all the instances $(\{v_1, v_2, \ldots, v_n\}, S)$ of SUBSET SUM such that v_1, v_2, \ldots, v_n and S are all k-bit natural numbers. This solution assumes that for each neuron (or, at least, for each subsystem) it is possible to choose whether such a neuron (resp, subsystem) works in a deterministic vs. nondeterministic way, and in the sequential vs. the maximally parallel way.

In [10] we have also studied the computational power of *deterministic* accepting SN P systems working in the sequential way. In particular, we have shown that they can be simulated by deterministic Turing machines with a polynomial slowdown. This means that they are not able to solve **NP**–complete problems in polynomial time unless $\mathbf{P} = \mathbf{NP}$, a very unlikely situation. In future work, we will address the study of the computational power of deterministic accepting SN P systems working in the maximally parallel way.

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Towards a Complete Covering of SBML Functionalities

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Summary. The complexity of biological systems is at times made worse by the diversity of ways in which they are described: the organic evolution of the science over many years has led to a myriad of conventions. This confusion is reflected by the in-silico representation of biological models, where many different computational paradigms and formalisms are used in a variety of software tools.

The Systems Biology Markup Language (SBML) is an attempt to overcome this issue and aims to simplify the exchange of information by imposing a standardized way of representing models. The success of the idea is attested to by the fact that more than 110 software tools currently support SBML in one form or another.

This work focuses on the translation of the Cyto-Sim simulation language (based on a discrete stochastic implementation of P systems) to SBML. We consider the issues both from the point of view of the employed software architecture and from that of the mapping between the features of the Cyto-Sim language and those of SBML.

1 Introduction

Nowadays, very few common exchange formats exist. We face difficulties to exchange models among different analysis and simulation tools. Therefore, taking advantage of the different tools power and capabilities is the main issue among scientists.

To overcome this issue, in March 2001, a first step was taken. During the *First International Symposium on Computational Cell Biology*, (Massachussetts, USA), Michael Hucka presented a new simple, well-supported and with textual substrate (XML) language adding components that reflect the natural conceptual constructs used by modellers in the domain, **SBML: Systems Biology Markup Language**.

In the following November 2002, M. Hucka talked again about *The Systems Biology Markup Language* at the *I3C 4th Quarter Technical Meeting* (San Diego, CA). On that occasion, starting from the observation of the enormous proliferation of software tools in this domain, he observed that a single package able to

cover all needs does not exist. Different packages can have different *niche strengths* and their strengths are often complementary. Moreover, he highlighted that much likely, no single tool is able perform likewise in the near future because the range of capabilities needed is large and new techniques and new tools evolve all the time making simulations and results often not shareable or reusable. Finally, he remarked that SBML is intended to be a common exchange format for transferring network models among tools, even if it may not capture everything represented by every tool (lossy transformation). At the same time, SBML is not suited to represent experimental data and numerical results, even if with the addition of the metadata support it may be suitable as a storage format for models too.

In May 2003, during the I3C May 2003 Meeting (Cambridge, MA) with a presentation entitled Update on the Status of the Systems Biology Markup Language (SBML) [17], M. Hucka dealt with the Related Efforts, the current status of the features and the *software libraries* for SBML. In particular he talked about similarities with *CellML* and the current joint work with the aim to bring them together. He also discussed the adoption in SBML of some features from CellML, like the MathML Subset and the Metadata specification. Moreover, he presented *BioPAX*, specifying that it is oriented towards being a common exchange format for databases of pathways Complementary efforts and then not a competing tool. However, SBML and BioPAX teams will work together to define linkages between SBML and BioPAX representations. Finally, he presented the version 1.0.1 of *libsbml*, a library designed to help modellers to read, write, manipulate, translate, and validate SBML files and data streams. In the same year many other presentations have been given by M. Hucka and others. In fact, in June 2003 MIPNETS Meeting (Liverpool, UK), Finney presented The Systems Biology Workbench and Systems Biology Markup Language [9], a project funded by Japan Science and Technology Corporation ERATO program and started in the summer 2000. The project goal was to provide software infrastructure which (i) enables sharing of simulation/analysis software and models and (ii) enables collaboration between software developers. Focused on biochemical modelling, it is an environment that enables tools to interact using SBML to transfer models between tools and supporting resource sharing.

All 2004 long was spent to delineate the limitations of the current SBML specification. In the same year Shapiro at al. talked about *MathSBML* [39], a *Mathematica* package designed for manipulating SBML models. It converts SBML models into Mathematica data structures and provides a platform for manipulating and evaluating these models. In [16], Hucka et al. summarise the current and upcoming versions of SBML and their efforts at developing software infrastructure for supporting and broadening its use. They also provided a brief overview of the many available SBML-compatible software tools.

In the following year other interesting publications appeared. The one about MIRIAM [26] is related to the way to define a minimum quality standard for the encoding of biochemical models by means of a set of rules about quantitative models of biological systems. These rules define procedures for encoding and an-

notating models represented in machine-readable form. In May 2005 as well, an ideal test bed in the understanding of the cellular systems by means of computational modelling appeared in [42]. In this chapter one goes over (i) computing for modelling cells, (ii) SBML and (iii) developing the International E. coli Alliance, which has been created to tackle the whole cell problem. Again in [15] and [8], one presents SBML as an XML-based exchange format for computational models of biochemical networks, including overview, enhancements, several proposals for the language extension, model composition and multi-component chemical species. In [32], the first step forward is taken by the *P System* community for representing P systems which model biological reaction networks as SBML models.

In 2006 a very good paper about the *BioModels database* was published, an annotated resource of quantitative models of biomedical interest. Models are carefully curated to verify their correspondence to their source articles. They are also extensively annotated, with (i) terms from controlled vocabularies, such as disease codes and Gene Ontology terms and (ii) links to other data resources, such as sequence or pathway databases. Researchers in the biomedical and life science communities can then search and retrieve models related to a particular disease, biological process or molecular complex [25]. In February of the same year, the joint work between CellML and SBML team groups had a big result, namely the CellML2SBML tool [38] implemented as a suite of XSLT stylesheets that, when applied consecutively, convert models expressed in CellML into SBML without significant loss of information. One month later, Sarah Keating et al. presented another similar conversion tool: SBMLToolbox [22] a toolbox that facilitates importing and exporting models represented in SBML in and out of the MATLAB environment and provides functionality that enables an experienced user of either SBML or MATLAB to combine the computing power of MATLAB with the portability and exchangeability of an SBML model. Other two useful tools appeared in October of the same year, they were *SBMLSupportLayout* and *SBWAutoLayout*, supporting reading, creating, manipulating and writing layout information for biochemical models. SBMLSupportLayout can read, update, add and render model layout information. SBWAutoLayout can automatically layout models, graphically manipulate model layouts and generate layout information for models without layout information. Using them, researchers can study large or complex biochemical networks and benefit from the ability to automatically create lucid visualizations and store them in a portable and widely accepted format [5]. In the winter of the same year, Bergmann and Sauro described the current state of the Systems Biology Workbench talking about how users and developers can perceive SBW and then focusing on currently available SBW modules [2]. Yet other four smart tools have been presented in the current year: SBML ODE Solver Library [28] (SOSlib), a programming library for symbolic and numerical analysis of chemical reaction network models encoded in SBML; SBML-PET [43], a tool designed to enable parameter estimation for biological models including signalling pathways, gene regulation networks and metabolic pathways. It can estimate the parameters by fitting a variety of experimental data from different experimental conditions.

In the same year, Eccher and Priami presented a tool to translate SBML into pi-calculus [6] while Gheorghe presented the *P System Modelling Framework* [10], a framework able to simulate the evolution of multi-compartmental Gillespie algorithm over a hierarchy of compartment structures.

In 2007, other tools have appeared. The first one has been $SBMLR^1$, a tool able to link R to libsbml for SBML parsing and output converting SBML to R graph objects, and more. Another emerging project is $SemanticSBML^2$: a suite of tools to facilitate merging of SBML models for systems biology starting from all elements in the SBML files described by MIRIAM-type annotations. SemanticSBML will help to insert and check such annotations. The need to build a tool to facilitate the quick creation and editing of models encoded in SBML has been growing with the number of users and the increased complexity of the language. SBMLeditor[36] tries to answer this need by providing a very simple, low level editor of SBML files. Users can create and remove all the necessary bits and pieces of SBML in a controlled way, that maintains the validity of the final SBML file.

As many tools have been implemented all around SBML just to highlight the trust of developers on the standardizing initiatives related to the software biological infrastructures towards commons exchange formats. In particular, it is an undeniable fact the increasing and unison consensus among developers in favour of SBML. In fact, several languages have been recently developed to overcome these kind of problems (integrations, standardizing, reuse of biological models) [27], [14], [7], [40], [41], [12], [29], [1], [19], [18]. However, only two XML-based formats are suitable for representing compartmental reaction network models with sufficient mathematical depth that the descriptions can be used as direct input to simulation software. The two are CellML [4], [13] and SBML[17]. The latter is becoming a defacto standard for a common representation supporting basic biochemical models. In fact, today, SBML is supported by over 110 software systems. As a consequence, many SBML models of gene regulatory networks and metabolic pathways that code a considerably body of biological knowledge have been accumulated in repositories. Among all databases, I recall (i) the PANTHER Classification System, [31], an unique resource that classifies genes by their functions, using published scientific experimental evidence and evolutionary relationships to predict function even in the absence of direct experimental evidence; (ii) KEGG [21], a knowledge base for systematic analysis of gene functions, linking genomic information with higher order functional information; (iii) JWS Online [34], a Systems Biology tool for simulation of kinetic models from a curated model database and (iv) Reactome |20|, a curated resource of core pathways and reactions in human biology. The information in this database is cross-referenced with the sequence databases at NCBI, Ensembl and UniProt, the UCSC Genome Browser, HapMap, KEGG (Gene and Compound), ChEBI, PubMed and GO. In addition to curated human events, inferred orthologous events in 22 non-human species including mouse, rat, chicken, zebra fish, worm, fly, yeast, two plants and E.coli are also available.

¹ Web Site of SBMLR: http://cran.r-project.org/src/contrib/Descriptions/rsbml.html

² Web Site of SemanticSBML: http://sysbio.molgen.mpg.de/semanticsbml/

Therefore, with the constant focus on SBML, in this paper I am going to inspect in section 2 all the features and the internal structure of SBML, in the section 4 the software facilities employed and the software packages implemented to build a pure Java library to handle SBML documents, in the section 3 I am going to show how to use the library to encode and decode information from a SBML file to Cyto-Sim model and vice-versa. In the section 5 I am going to test the software package implemented on real SBML files taken from different data sources and in the last section I am going to delineate the future works.

2 SBML Structure

In this paper I am going to take into consideration the latest stable release of SBML highlighting differences with the previous versions. SBML is primarily oriented to allow models to be encoded using XML. Major release of SBML are termed *levels* and represent substantial changes to the composition and structure of the language. The latest release³ is the SBML level 2. It represents an incremental evolution of the level 1, therefore a valid SBML level 1 can be mapped in a valid SBML level 2 file while only a subset of level 2 can be mapped in a level 1 file. However, both levels remain separated and distinct. Instead, minor revisions of whatever level of a SBML file will bring a new *version*. A new version carries new corrections, adjusts and refinings of the language. Some languages features of previous levels or versions can be deprecated or entirely removed in the time. A feature can be directly removed, but not recommended, although the usual path to completely remove a feature is to deprecate it before and then remove to maintain backward compatibility as much as possible.

SBML allows models of arbitrary complexity to be represented. Each type in a model is described using a specific type of data structure that organizes the relevant information and derive directly or indirectly from a single abstract type called *Sbase*. In addition to serving as the parent class for most other classes of objects in SBML, this base type is designed to allow modeller or a software package to attach arbitrary information to each major structure or list in an SBML model. Sbase contains two default elements: (i) *notes*, intended to serve as a place for storing optional information intended to be seen by humans; (ii) *annotations*, it is a container for optional software-generated content not meant to be shown to humans. All other types have at least other three parameters: (i) *id*, a mandatory field on most SBML structures used to identify a component within the model definition; (ii) *name*, an optional field, not intended to be used for cross-referencing purposes within a model but just to provide a human-readable label for the component; (iii) *sboTerm*, an optional field used to identify a term from an ontology where vocabulary describes entities and processes in computational models.

 $^{^{3}}$ at the writing time

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2.1 SBML in more depth ...

The top level of an SBML model definition consists of a list of all data type (figure 1), all being optional. These elements are contained in the *model* element directly enveloped in the most outer element properly called *sbml*.



Fig. 1. List of all data types in a SBML model

- SBML envelope: The root of a SBML file is an *SBML element* with three required fields: two attributes (*level* and *version*) and exactly one child node (*model*).
- Model: It is the highest level construct in a SBML document. It has got three optional parameters: *id*, *name*, *sboTerm* and the hierarchy of data types just before shown and in the exact order of the listed terms.
- Function definitions: A *function* (or often called *user-defined function*) is a mere textual macro containing a MathML lambda element, then its call can be implemented as a textual substitutions. It has limited capabilities because none external parameter can be referenced by a function.
- Unit definitions: SBML optionally allows for the employment of units of measurements for some mathematical entities (e.g. compartment size, reaction rates, mathematical formulas, etc...). By means of two operative classes (UnitDefinition and Unit), a new unit of measurement (UnitDefinition(newid, newname)) can be obtained by composition of the elementary ones already defined in SBML (Unit(id, name)).

- Compartment types: If compartments share either a common biological function or similar reactions or other underlying conceptual features, they can be grouped assigning to them the same compartment type value. After defining a compartment type with its univocal identifier, it can be referred by each compartment belonging to the category specified by that compartment type.
- Species types: As compartments, if species share common characteristics, they can be grouped in a same logical set specified by a new species type. The existence of SpeciesType structures in a model has not effect on the model's numerical interpretation.
- Compartments: A compartment represents a bounded space in which species are located. Compartments can be arranged in hierarchical structures and can have a static or dynamical size.
- Species: A species refers to a multiset of entities of a specific species type that take part in reactions and are located in a specific compartment. With a species it is possible to specify a number of parameters to better explaining its characteristic. For example for a species one can specify the substance quantity or the concentrations, the charge, the substance unit, etc...
- Parameters: As usual programming languages, they are variables that can be used in mathematical formulas. They are defined as constant values and can be global, if defined at the beginning of the model or local if defined within the scope of a reaction. Local parameters overwrite the global ones with the same name.
- Initial Assignments: An initial assignment is an alternative way to set quantities of species, size of compartments and value of parameters with complex mathematical expressions at the beginning of a simulation. It overwrites eventual already existing numerical values for quantities, sizes and parameter with the mathematical expression it carries.
- Rules: A rule furnishes a supplementary way to set a variable of the system that cannot be set otherwise by any reaction rules or initial assignments. There are three kind of rules: (i) assignment, used to express equations that set the values of the variables. As an initial assignment, it overrides eventual existing numerical values of species quantity, compartments size and parameters value with its mathematical expression. It is forbidden to have both an initial assignment and a assignment rule related to the same object; (ii) rate, it is used to express equations that determine the rates of change of variables. It can refer to species, compartments and parameters as before explained. In the context of a simulation, it can have effect either at t = 0 to obtain consistent initial conditions or at t > 0 during the simulation run. To avoid indetermination problems, for each object in a system, only one between assignment rules and rate rules can exist; (iii) algebraic rule, used to express equations that are neither assignments of model variables nor rates of changes. The only one role is to distinguish this case from the other cases.
- Constraints: Constraints are mathematical expressions used to check permissible values of different quantities in a model. A constraint should be checked against

a variable at all time and triggers a message warning if the conditions expressed by the formula is not verified.

- Reactions: Reactions are processes that result in the interconversion of substances that can change their quantities. The participants of a reaction are called *reactants* and *products*. They change their quantities according to their own stoichiometric coefficient and the kinetic law which rules the reaction. All participants must belong to the list of species. A reaction can be optionally set as *reversible*, *fast* and can contains modifiers, namely species acting as catalysts or inhibitors.
- Events: Events defines changes of variables of a system when a triggering condition fires. In particular an event specify (i) under which mathematical condition, (ii) how and (iii) when a variable changes. The triggered events can be delayed and applied to more than one variables at t > 0. As usual, events have effect on species quantity, compartments size and parameters value.

2.2 Differences between SBML Level 1 and Level 2

Many significant changes characterize the new level 2. They cover the (i) identification of the objects, (ii) annotation within the model, (iii) language for kinetic expressions, (iv) dimensionality of compartments and (v) rule specification.

In particular SBML Level 2 supports the inclusion of metadata. In fact, all structures in SBML can be annotated with optional content in RDF. The top-level *Model* structure can contain an optional list of global user-defined functions expressed in *MathML* and organized in new structures of type *FunctionDefinition* and can contain an optional list of event definitions organized in structures of type *Event*. All data structures, including SBML and listOf elements, are now derived from the type SBase. This means all major structures in SBML can have separate annotations and metadata associated with them.

A new field, *id*, replaces the name field previously defined for most SBML structures to identify each part of a model. Formulas in Level 2 are expressed using MathML 2.0. The field named formula previously available on the KineticLaw and Rule structures has been replaced by a MathML element named math containing MathML content. In addition, stoichiometry numbers may now be expressed using MathML, allowing for more flexibility in defining reactions. Unlike in SBML Level 1, unit identifiers in Level 2 are in a separate namespace from the namespace used for models, functions, species, compartments, reactions and parameters and have the additional fields *multiplier* and *offset* to enable the definition of non-SI units.

The Compartment structure has a new field, *spatialDimensions*, whose value is a positive integer specifying the number of dimensions in space the compartment possesses. This enables the definition of such things as two-dimensional membranes. All fields representing initial conditions or parameter values, including compartment sizes and species concentrations, are optional in Level 2. A missing value for one of these fields implies that the value is either unknown, not required for analysis, or should be obtained from an external source. The Compartment, Species and Parameter structures each have a new boolean field named constant. This field specifies whether the variables represented by these structures can be changed by rules and reactions. In particular, the Species structure has a new field, *initialConcentration*, for setting the initial value of a species in terms of its concentration. This is in addition to the ability, carried over from Level 1, to set the values in terms of amounts.

There is no longer a type field on Rule, and new structures AssignmentRule and RateRule replace SBML Level 1's ParameterRule, SpeciesConcentrationRule and CompartmentVolumeRule. The Reaction structure has a new list of modifiers in addition to the list of reactants and products. The listOfModifiers enumerates species that affect a reaction but are neither created nor destroyed by the reaction.

$3 \text{ SBML} \Leftrightarrow \text{Cyto-Sim}$

As deeply shown, SBML is a powerful and well defined language for modelling biological interactive systems in standard way. The aim of my work has been to make Cyto-Sim able to *speak* and *understand* SBML.

Cyto-Sim [3] is a stochastic simulator of biochemical processes in hierarchical compartments which may be isolated or may communicate via peripheral and integral membrane proteins. It is available online as a Java applet [33] and as standalone application. For security issue, although the functionalities of the applet has been reduced, it fully and correctly works. By means of it, it is possible to model: (i) interacting species; (ii) compartmental hierarchies; (iii) species localizations inside compartments and membranes and (iv) rules and their and correlated velocity formulas which govern the dynamics of the system to be simulated, as chemical equations.

Some real biological systems have already been successfully simulated in the past by means of Cyto-Sim. Now I am going to try to explain at first how to translate a Cyto-Sim model into SBML (and vice-versa) and later I will test the quality of the translation comparing the simulations available in literature against those obtained by Cyto-Sim about the same models.

3.1 Speaking SBML

The conversion process from the Cyto-Sim syntax to the SBML one is quite straightforward. In Cyto-Sim, users must declare the species present into the system writing something like this:

```
/* Object Declaration */
object speciesA, speciesB, speciesC
```

This line of code corresponds to the following SBML chunk of code:

```
<listOfSpecies>
<species id="compartmentA_0_speciesA" name="speciesA"</li>
```

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```
compartment="compartmentA" initialAmount="0.0"/>
<species id="compartmentB_0_speciesB" name="speciesB"
    compartment="compartmentB" initialAmount="1.0"/>
<species id="compartmentC_2_speciesC" name="speciesC"
    compartment="compartmentB" initialAmount="2.0"/>
</listOfSpecies>
```

Not all the information in this XML code can be retrieved by the previous objects specification⁴. In fact, the compartment, membrane and initial amount related to one species are reached both from the following code:

```
/* Compartments Declarations */
compartment compartmentA [ruleA]
compartment compartmentB [compartmentA, ruleB, ruleC,
    speciesB, 100 speciesB@7000 : |2 speciesC|]
system compartmentB
```

and from this:

```
/* Rules Declarations */
rule ruleA {
   speciesA k1-> *
    || + speciesA k2-> speciesA + ||
}
rule ruleB speciesB k3-> speciesC
rule ruleC |speciesC| k4-> || + speciesC
```

From the code related to the compartments we take information about (i) the compartment hierarchy, (ii) which rule happens and in what compartment, (iii) the declared initial quantities (species not declared in this context will not still exist as default at the beginning of the simulation) and (iv) eventual re-feeding events at specified evolution times. Considering now that a reaction happening in a compartment acts only on the species within it, looking the localization of a reaction we can infer the localization of its reactant species. Moreover it is possible to notice that the rule *ruleC* acts on the species *speciesC* inside the membrane⁵ (membrane number 2) of the compartment *compartmentB*.

In SBML each compartment is quadruplicated to easily handle membranes.

```
<listOfCompartments>
<compartment id="compartmentA_0" compartmentType="compartmentA"
outside="compartmentA_1"/>
```

⁴ The figure between the compartment and the species names within the string assigned to each species id corresponds to the membrane in which a species sits. For more information about the syntax, look at [3]

⁵ Recall that in this context a compartment is surrounded by a membrane with a not negligible thickness, therefore a compartment is logically divided into the internal (membrane 0), internal and superficial (membrane 1), intra (membrane 2), external and superficial (membrane 3) and external (membrane 4) membranes.

```
<compartment id="compartmentA_1" compartmentType="compartmentA"
        outside="compartmentA_2"/>
    <compartment id="compartmentA_2" compartmentType="compartmentA"
        outside="compartmentA_3"/>
    <compartment id="compartmentA_3" compartmentType="compartmentA"
        outside="compartmentB_0"/>
    <compartment id="compartmentB_0" compartmentType="compartmentB"
        outside="compartmentB_1"/>
    <compartment id="compartmentB_1" compartmentType="compartmentB"
        outside="compartmentB_2"/>
    <compartment id="compartmentB_2" compartmentType="compartmentB"
        outside="compartmentB_3"/>
    <compartment id="compartmentB_3" compartmentType="compartmentB"
        outside="system_0"/>
    <compartment id="system_0" compartmentType="system"
        outside="system_1"/>
    <compartment id="system_1" compartmentType="system"
        outside="system_2"/>
    <compartment id="system_2" compartmentType="system"
        outside="system_3"/>
    <compartment id="system_3" compartmentType="system"/>
</listOfCompartments>
```

Then a single compartment generates four independent concentric compartments, as a matrioska doll toy, related to the same compartment but enclosing different spatial areas and then species. To keep conceptually linked these compartments, a compartment type specification is provided.

```
<listOfCompartmentTypes>
<compartmentType id="compartmentA"/>
<compartmentType id="compartmentB"/>
<compartmentType id="system"/>
</listOfCompartmentTypes>
```

The previously seen reactions are easily translated into SBML differenting the names of the grouped rules (e.g. the ruleA group contains two reactions. Their names will become: ruleA.0 and ruleA.1). Moreover, the kinetic formulas just touched (k1, k2, etc) before are expressed by MathML expressions inside <kineticLaw> tags.

```
<listOfReactions>
[...]
<reaction id="ruleA.1" name="compartmentA_0_ruleA.1">
<listOfReactants>
<speciesReference species="compartmentA_0_speciesA"
stoichiometry="1.0"/>
</listOfReactants>
<!--listOfProducts>No Products</listOfProducts-->
<kineticLaw>
```

```
<math xmlns="http://www.w3.org/1998/Math/MathML">
        <apply>
        <times/>
        <cn>k1_value</cn>
        <ci>ccompartmentA_0_speciesA</ci>
        </apply>
        </math>
        </kineticLaw>
        </reaction>
        </listOfReactions>
```

Cyto-Sim also requires the specification of a range of evolution times and of the species whose quantities have to be plotted on the screen.

```
evolve 0 - 1000
plot compartmentA[speciesA], compartmentB[speciesB:|speciesC|]
```

This information can be encoded in SBML by the use of an annotation which is auto-explicative.

3.2 Understanding SBML

The process to make an existing SBML file comprehensible to Cyto-Sim is more complex than the opposite step. Keeping in mind the correspondences among structures before shown, during this kind of translation we have to check some restrictions and to guarantee some constraints which are now explained.

Parameters: SBML optionally carries global parameters, visible everywhere in the file and local ones with more restricted scope. During the parsing time of an SBML file, Cyto-Sim loads all global parameters putting them into a global HashMap. In the case of local parameters inside kineticLaw of reactions, Cyto-Sim considers local and global parameters together taking care to overwrite eventual global parameters with the same name of local ones.

- Species Quantities: SBML provides optional size for compartments. Cyto-Sim handles quantities and not concentration for species, then each concentration (if any) has to be converted into quantity. To do that, Cyto-Sim requires the size specification for each compartment if there are any specification of the species concentrations inside it.
- Assignments: Cyto-Sim handles assignment rules at the moment of parsing and use them to replace eventual existing fixed values specified for species quantity, compartment size or parameters value. Up to now, it does not understand initial assignments, rate rules and algebraic rules. These features will be made available soon.

Functions: Cyto-Sim does not still handle λ -functions.

Units & Constraints: Cyto-Sim does not still make use of units of measurements and costraints.

4 Binding to the SBML Schema

After having conceptually explained how Cyto-Sim converts SBML in its own language and vice-versa, now I am going to show which software architecture gives it the possibility to do that. I used two well known tools for this aim: the Java Architecture for XML Binding (JAXB) package and the XML DOM parser, both build-in the latest release of Java (Java Mustang).

JAXB [11] simplifies access to an XML document from a Java program by presenting the XML document to the program in a Java format. The first step in this process is to bind the schema for the XML document into a set of Java classes that represents the schema. Binding a schema means generating a set of Java classes that represents the schema. All JAXB implementations provide a tool called *binding compiler* in order to bind a schema. In response, the binding compiler generates a set of interfaces and a set of classes that implement the interface. I obtained Java classes for each available XML levels and versions, I mean SBML level 1 version 1, level 1 version 2, level 2 version 1 and level 2 version 2. Later, I compiled and packaged them into just one package. The second step is to unmarshal an SBML document. Unmurshalling means creating a tree of content objects that represents the content and the organization of the document. The content tree is not a DOM-based tree. In fact, content trees produced through JAXB can be more efficient in terms of memory use than DOM-based trees. The content objects are instances of the classes produced by the binding compiler. In addition to providing a binding compiler, JAXB provides runtime APIs for JAXBrelated operations such as marshalling. It is possible to validate source data against an associated schema as part of the unmarshalling operation. If the data is found to be invalid (that is, it doesn't conform to the schema) the JAXB implementation can report it and might take further action. JAXB providers have a lot of flexibility here. The JAXB specification mandates that all provider implementations report validation errors when the errors are encountered, but the implementation does

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not have to stop processing the data. Some provider implementations might stop processing when the first error is found, others might stop even if many errors are found. In other words, it is possible for a JAXB implementation to successfully unmarshal an invalid XML document, and build a Java content tree. However, the result will not be valid. The main requirement is that all JAXB implementations must be able to unmarshal valid documents. I unmarshal and validate each SBML file at runtime.



Fig. 2. Software Architecture for SBML Binding

The W3C Document Object Model (DOM) is a platform and language-neutral interface that allows programs and scripts to dynamically access and update the content, structure, and style of a document. The XML DOM is the tool to define (i) a standard set of objects for XML, (ii) a standard way to access XML documents; (iii) a standard way to manipulate XML documents. Cyto-Sim uses the DOM parser contained into xerces2-j [35] built into the Java Mustang release. The DOM parser is used to:

Check Levels and Versions: Cyto-Sim preliminary opens SBML files and checks levels and versions (delegating validation and comprehension to JAXB). It acquires knowledge about which JAXB context instantiating or, more clearly, which SBML schema considering for binding, unmarshalling and validation; Parse MathML expression: Due to intrinsic limitations of JAXB to handle recursively nested xml tags, Cyto-Sim makes use of DOM to explore MathML expressions and parse their components.

5 Experimental Tests

The capability of Cyto-Sim to understand all currently existing SBML levels and versions has been tested on almost all official existing SBML files available on the web. I successfully imported all SBML files generated by Gepasi [30], a software package for modelling biochemical systems and the most part of the models stored into the BioModels database [25]. Gepasi makes available 9 SBML level 1 version 1 files 6 while BioModels has 70 curated and 43 not curated models exported as SBML level 2 version 1 files. I have also tested models from the PANTHER (130 SBML level 1 version 2 files) Classification System, and from KEGG (77 SBML level 2 version 1 files). All SBML files were converted from KEGG by using a conversion script keqg2sbml. Moreover, I retrieved some interesting models among all 238 CellML models and tested them. To do that, I had to manually convert from the CellML format to SBML by means of CellML2SBML [38] and later import and simulate them with Cyto-Sim. All imported files have been successfully parsed by Cyto-Sim. This testifies the quality of the conversion routines and of the architecture employed. Summarising, I retrieved 567 models from the most known and famous biological model containers available in SBML (or in formats having reference to SBML), and tested them. I obtained a successful test, when Cyto-Sim had been able to correctly parse the inferred model. In particular, now I am going to show a couple of examples which Cyto-Sim has been able not only to correctly parse, but also to simulate and get the same results shown in literature.

The first test is related to the model *BIOMD000000010* picked up from the BioModels database. It concerns the functional organization of signal transduction into protein phosphorylation cascades and in particular the mitogen-activated protein kinase (MAPK) cascades. It greatly enhances the sensitivity of cellular targets to external stimuli [23]. In this paper it is demonstrated that a negative feedback loop combined with intrinsic ultrasensitivity of the MAPK cascade can bring about sustained oscillations in MAPK phosphorylation. The conversion of the SBML file produces the following model with 1 compartment, 8 species and 10 reactions.

```
object MKKK, MKKK_P, MKK, MKK_P, MKK_PP, MAPK, MAPK_P, MAPK_PP
```

```
rule J0 MKKK ((1.0*2.5*MKKK)/((1+((MAPK_PP/9.0)^1.0))*(10.0+MKKK)))-> MKKK_P
rule J1 MKKK_P ((1.0*0.25*MKKK_P)/(8.0+MKKK_P))-> MKKK
rule J2 MKK ((1.0*0.025*MKKK_P*MKK)/(15.0+MKK))-> MKK_P
rule J3 MKK_P ((1.0*0.025*MKKK_P*MKK_P)/(15.0+MKK_P))-> MKK_PP
rule J4 MKK_PP ((1.0*0.75*MKK_PP)/(15.0+MKK_PP))-> MKK_P
```

⁶ among all, a very large model representing a set of 100 yeast cells in a liquid culture whose dynamics is represented by means of 2000 reactions

```
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```

```
rule J5 MKK_P ((1.0*0.75*MKK_P)/(15.0+MKK_P))-> MKK
rule J6 MAPK ((1.0*0.025*MKK_PP*MAPK)/(15.0+MAPK))-> MAPK_P
rule J7 MAPK_P ((1.0*0.025*MKK_PP*MAPK_P)/(15.0+MAPK_P))-> MAPK_PP
rule J8 MAPK_PP ((1.0*0.5*MAPK_PP)/(15.0+MAPK_PP))-> MAPK_P
rule J9 MAPK_P ((1.0*0.5*MAPK_P)/(15.0+MAPK_P))-> MAPK
compartment uVol[J0, J1, J2, J3, J4, J5, J6, J7, J8, J9, 280.0 MAPK,
10.0 MKK_P, 10.0 MKK_PP, 10.0 MKKK_P, 10.0 MAPK_PP, 280.0 MKK,
10.0 MAPK_P, 90.0 MKKK]
system uVol
evolve 0-33000
plot uVol[MAPK,MAPK_PP]
```

In the figure 3, on the left is shown the simulation result coming from the literature and on the right that one obtained with Cyto-Sim. The graphs are identical.



Fig. 3. Sustained oscillations in MAPK cascade

The second test is related to the glucose transport by the Bacterial Phosphoenolpyruvate [37] whose model has been found in JWS Online. The resulting model has 1 compartment, 17 species and 10 reactions.

```
object EI, PyrPI, EIP, HPr, EIPHPr, HPrP, EIIA, HPrPIIA, EIIAP, EIICB,
EIIAPIICB, EIICBP, EIICBPGlc, PEP, Pyr, GlcP, Glc
rule v1 PEP + EI ((1960.0*PEP*EI)-(480000.0*PyrPI))-> PyrPI
rule v2 PyrPI ((108000.0*PyrPI)-(294.0*Pyr*EIP))-> EIP + Pyr
rule v3 HPr + EIP ((14000.0*EIP*HPr)-(14000.0*EIPHPr))-> EIPHPr
rule v4 EIPHPr ((84000.0*EIP*HPr)-(14000.0*EIPHPr))-> HPrP + EI
rule v5 HPrP + EIIA ((21960.0*HPrP*EIIA)-(21960.0*HPrPIIA))-> HPrPIIA
rule v6 HPrPIIA ((4392.0*HPrPIA)-(3384.0*HPr*EIIAP))-> EIIAP + HPr
rule v7 EIICB + EIIAP ((880.0*EIIAP*EIICB)-(880.0*EIIAPIICB))-> EIIAPIICB
rule v8 EIIAPIICB ((2640.0*EIIAP*EIICB)-(960.0*EIIA*EIICBP))-> EIICBP + EIIA
rule v9 EIICBP + Glc ((260.0*EIICBP*Glc)-(389.0*EIICBPGlc))-> EIICBPGlc
rule v10 EIICBPGlc ((4800.0*EIICBPGlc)-(0.0054*EIICB*GlcP))-> EIICB + GlcP
```

```
compartment compartment_cyto_sim[v1, v2, v3, v4, v5, v6, v7, v8, v9, v10,
 0.0 EIICBPGlc, 5.0 EIICBP, 25.0 HPrP, 2.0 EIP, 20.0 EIIA, 5.0 EIICB,
 25.0 HPr, 2800.0 PEP, 0.0 PyrPI, 0.0 EIPHPr, 50.0 GlcP, 900.0 Pyr,
 0.0 HPrPIIA, 20.0 EIIAP, 500.0 Glc, 0.0 EIIAPIICB, 3.0 EI]
system compartment_cyto_sim
evolve 0-10000
plot compartment_cyto_sim[HPrP,EIIAPIICB,HPrPIIA]
```

In the figure 4 it is possible to notice that both graphs represent the same behaviour. The differences are due to the deterministic (on the left) or stochastic (on the right) nature of the simulations.



Fig. 4. Glucose Transport by the Bacterial Phosphoenolpyruvate

At the end, I tested the whole human reactome derived from Reactome. The actual release of the human reactome I used is an SBML file containing 28 compartments (even including internal membranes of the same compartment), 3054 species (in all their forms) and 1979 interactions represented by means of reactions. Cyto-Sim is able to parse and even to simulate it, although at the moment it cannot have meaning because of the lack of quantitative parameters (reaction rates and initial species quantities).

6 Conclusion

kosmopolitês (citizen of the world), has been used to describe a wide variety of important views in moral and socio-political philosophy. The nebulous core shared by all cosmopolitan views is the idea that all human beings, regardless of their political affiliation, do (or at least can) belong to a single community, and that this community should be cultivated. Different versions of cosmopolitanism envision this community in different ways, some focusing on political institutions, others on moral norms or relationships, and still others focusing on shared markets or forms of cultural expression [24].

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In the context of the present work, a citizen of the world is anyone who speaks and understands a common language, who can travel to the ends of the earth without worrying about misunderstanding or being misunderstood. Limited comprehension of language is the greatest barrier for people who need to spread information and ideas. This is exactly the case for scientists who wish to share their results and models with the widest possible audience.

In this paper I have presented an extended overview of the SBML story, continually remarking on the increasing interest of scientists both to support and write their biological models in SBML. I talked about the internal structure of SBML, in order to focus on its expressive potentialities, and later I presented a possible software architectural arrangement to allow simple binding to SBML schemas and correct unmarshalling of SBML files. Finally, I presented tests performed on two models coming from separate databases. I demonstrated the correctness of the translation routines and highlighted the similarities of the obtained simulation results.

Today there are more than 600 models written in SBML, ready to be more accurately studied, confirmed or refuted. Challenging existing knowledge is the means to increase understanding and therefore to *grow* knowledge. The best way to achieve this is to maximize the number of people that speak the same language, in this case SBML. My work sits perfectly in this context and my hope is that it has wide application.

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Frequency Membrane Systems

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Summary. We define a model of membrane system where each membrane is clocked independently from the others, in the sense that every derivation step is applied without a global synchronization. The computation is obtained by the execution of a limited amount of rules in each membrane, and only when they are allowed to execute a derivation step. Indeed, each membrane operates with a certain work frequency that could change across the system and during the execution. Simple results show that this model is at least as powerful as the usual one, and a few examples demonstrate that it gives rise to interesting dynamic behaviors.

1 Introduction: Biological Motivation

This work introduces a new class of P systems, where we want to come closer to biological cell's behavior by adopting some new features of membrane computing that could have a higher correspondence in biology. As we known, in general, different chemical reactions can take different time to be executed and moreover the same reaction could take different time depending on some environment conditions (for example concentration, temperature, etc.).

Some models have already adopted different approaches to the timing in P systems. For instance, in [5] maximal parallelism is not enforced, while in [1] a single rule is probabilistically chosen inside each membrane, but all selected rules work in parallel. Interestingly, paper [1] specifically aims at modeling some biological phenomena. In [2,6,4] authors study a model where each rule can take a different time to perform its action, and look for systems where even if timing changes results stay the same. Further results along this line are in [3].

We want to explore other changes to timing inside P systems but, above all, we want to focus the attention on the real asynchronous nature of biological cells. We want to consider every single membrane as a separate domain with a specific clock, where the rules can be applied as often as specified by their membranes' clock frequency. In this behavior, a single derivation step can occur only when the membrane has the right, specified by its own clock and by other constraints, to carry out one or more rules, so each component of the system, regarding the overall computation, is totally independent from the others, hence the system is partially asynchronous.

As we said before we want, also, to adopt a different approach to the maximal parallel way to execute the rules. We think maximal parallelism is a very strong assumption, so we want to bound the amount of rules that can be applied in each membrane's derivation step. The motivations for this feature come from the fact that, in a living cell, there is a limited amount of energy, hence a limited amount of reactions can take place in a given time.

In this paper we introduce also a decay time for some symbol objects; we have made this assumption thinking that other reactions (here not described) could be modeled by such feature. For instance, symbols could decay because the cell itself uses them to get energy.

One more detail related to clocking is the offset on the starting time of operations in each membrane. This, and the other modeling features are discussed by means of examples and simple formal results in the following sections.

2 Frequency P Systems: Definition

A frequency P system is a P system where each rule has a time of execution. This execution's time is expressed in clock steps. Each membrane has its own clock, and a limited amount of rules that can be applied in each clock step (or a fixed amount of energy, if we want to associate different levels of energy to the rules). The clock period is a multiple of the unit time of an external observer.

The system will use symbol objects with evolution rules. We denote by N the set of natural numbers.

A Frequency P system with symbol objects of degree $m \ge 1$, is a construct

$$\Pi = (O, D, T, \mu, \omega_1, \dots, \omega_m, E, t_D, C, R_1, \dots, R_m, i_O)$$

where:

- *O* is the alphabet of the objects;
- $D \subseteq O$ is the alphabet of decaying symbols;
- $T \subseteq O$ is the alphabet of non-decaying symbols;
- μ is a membrane structure consisting of *m* membranes labeled with $1, 2, \ldots, m$;
- $\omega_i, 1 \leq i \leq m$, specifies the multiset of objects present in the corresponding region *i* at the beginning of a computation;
- E ⊆ N^m is a set of m numbers indicating the energy value assigned to each membrane at every membrane's clock step, overriding any previous energy level associated to them;
- t_D ⊆ Nⁿ is a set of n numbers indicating the decay time of the n decay symbols in D;

- $C \subseteq N^m$ is a set of *m* numbers indicating the clock value (referred to an external observer) assigned on each membrane;
- $R_i, 1 \leq i \leq m$, are finite sets of evolutionary rules over O associated with regions $1, 2, \ldots, m$ of μ ; the rules can be either cooperative or non-cooperative rules of the form $A \longrightarrow_s^k v$, where A is an object from O and v is a string over $\{a_{here}, a_{out}, a_{in} \mid a \text{ in } O, 1 \leq j \leq m\}$, if the target is not specified, then it is intended to be here; k is an integer representing the energy to consume to apply the rule. Note that k could be a negative number, in this case we assume that the reaction modeled by the rule produces energy for the cell, when k is not specified we assume that k=1; s in N is the number of clock's steps necessary to the rules to act (and produce the objects in the right hand side), when s is not specified we assume that s=0;
- i_O in $\{0, 1, \dots, m\}$ is the output region (0 for the environment).

A configuration of \prod at a given time t is represented by a string of parentheses (the structure μ) and strings over O (contents of the regions). For instance, a possible configuration of the system at time 0 (the starting time) with an alphabet $O = \{A, B\}$ and a structure $\mu = [1[2]2[3]3]1$ could be:

 $\varsigma(\Pi(0)) = [{}_{1}[{}_{2}AA]_{2}[{}_{3}BB]_{3}]_{1}$

Given a string ω representing a configuration at time t, then all the strings obtained from ω by taking any permutation of the strings, representing the contents of the regions, represent the same configuration at time t.

We suppose the existence of an external global clock that ticks at uniform intervals, taken as time unit, starting at time 0; we also suppose that each membrane has its own clock that ticks at uniform intervals, taken as multiple of the observer's time unit. In the former example we assume that each membrane starts at same time (time 0 of the observer).

In each membrane of the system we have a finite number of objects from alphabet O, a finite set of evolution rules each one with its own costs (in time and energy) and a finite amount of energy. At each time step of the observer's clock, we have membranes in the system that are allowed to execute their rules according to their own clock ticks and membranes that are not allow to execute their rules (until their next clock ticks). These membranes could receive object symbols from other membranes if they are target of some rules but they are not allowed to deal with those objects until their next clock ticks. We identify former membranes as active membranes and the latter as passive membranes (in a given observer's time).

When a membrane becomes active we apply as many rules as we can, according to the left hand side of the rules and the energy necessary to carry out the rules. It's important to understand that the rules are NOT applied in maximum parallel manner as in standard P systems, since only one instance of objects on the left hand side of the rule will be consumed and only one instance of the objects on the right hand side of the rule will be produced after a fixed number of time step specified by the rule. During the execution of a rule, the occurrences of symbol objects in the right hand side are no longer available for other rules. Each time a rule is applied we decrease the energy value of the membrane by the value specified by the rule, in this way a membrane could execute a fixed number of rules for each clock ticks. The energy level of a membrane will be reset to the initial value at the beginning of the next membrane clock ticks.

If two or more rules in an active membrane are allowed to be executed, then possible conflicts for using the occurrences of symbol objects are solved by assigning the objects in a non-deterministic way. The computation halts when at a certain observer's clock step no rule can be applied in any region and there are no rules in execution. The output of a halting computation is the vector of numbers representing the multiplicities of objects present in the output region in the halting configuration.

3 Frequency P Systems: Examples

The following example shows how a frequency P system works:

$$\begin{aligned} \Pi_0 &= (O, D, T, \mu = [{}_1[2]_2[_3]_3[_4]_4]_1, \omega_1 = \lambda, \omega_2 = A^5, \omega_3 = B^7, \omega_4 = C^7, \\ E, t_D, C, R_1, R_2, R_3, R_4, i_O = 1), \end{aligned}$$

where:

$$\begin{split} & O = \{A, B, C, a, b, c, e\} \; ; \; D = \emptyset \; ; \; T = O; \\ & E = \{2, 1, 1, 1\} \\ & t_D(i) = 1, \forall i \in D \\ & C = \{C_1 = 1, C_2 = 3, C_3 = 2, C_4 = 2\} \\ & R_1 = \{r_1 : abc \to e, r_2 : bc \to a, r_3 : aa \to e\} \\ & R_2 = \{r_4 : A \to a_{out}\} \\ & R_3 = \{r_5 : B \to b_{out}\} \\ & R_4 = \{r_6 : C \to c_{out}\} \end{split}$$

The starting configuration of this frequency P system represented by Figure 1. The configuration of the system at different times is shown in Figure 2.

In this and in the following figures, we focus the attention only to the objects in skin membrane $\mu 1$, in particular by displaying the availability in there of symbols a, b, c at different times, by marking in the picture the axis labeled as "Symbol object...".

Note that if we set to 0 the time of execution of each rule we get the same result of a computation carried out with a standard P system; the only difference is that we introduce a sort of priority in rules execution. For example, if we introduce a new rule in membrane $\mu 1$ (r7: $a \rightarrow f$) we force the system to apply this rule before the others at least in t5 (because in each clock step the membrane could execute at most 2 rules if it can, and in t5 no other rule can be applied except r7). In this case we reach the same final configuration of a standard P system also because







Fig. 2. The evolution of skin membrane in Π_0 .

we set an energy value for the membrane $\mu 1$ to E=2. This parameter in the given example enforce maximality because in case of presence of the symbol objects a, b, cin the membrane we could apply either the rule r1 or both the rules r2 and r3 in the same clock step. This behavior produces the same result (the symbol object e) but in the former case it consumes only a single unit of energy indeed, while in the latter, it consumes two energy units.

Note that this sort of "chain reaction" is allowed by the definitions of the system, even though this happens only because we set to 0 the time of execution

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of the rules; but, despite of that, this feature will become useful in some further variants of the system (for example we could apply a rule even if the membrane is not active if we have enough energy units left, or we could increase or decrease the clock frequency of the membrane in function of its own energy level).

Note also that if we do not associate decay time to some object symbols we get the same result even if we starts the membranes not at the same time.

In the following example we run the same P system as above, but with the difference that we introduce a delay of one observer's clock step for the membrane $\mu 4$. The resulting dynamics is shown in Figure 3.



Fig. 3. Dynamics in skin membrane of a system derived from Π_0 by introducing a delay for μ_4 .

4 Frequency P Systems with Decay Time: Examples

Now we shows the same example with a delay time 1 for the symbol objects $\{a, b, c\}$.

The following is the definition of the system Π_1 .

$$\begin{split} \Pi_1 &= (O, D, T, \mu = [_1[_2]_2[_3]_3[_4]_4]_1, \omega_1 = \lambda, \omega_2 = A^5, \omega_3 = B^7, \omega_4 = C^7, \\ &E, t_D, C, R_1, R_2, R_3, R_4, i_O = 1), \end{split}$$

where:

$$\begin{array}{l} O = \{A,B,C,a,b,c,e\} \; ; \; D = \{a,\mathbf{b},\mathbf{c}\} \; ; \; T = O\text{-}D = \{A,B,C,e\}; \\ E = \{2,1,1,1\} \\ t_D(i) = 1, \forall i \in D \\ C = \{C_1 = 1,C_2 = 3,C_3 = 2,C_4 = 2\} \end{array}$$

$$R_1 = \{r_1 : abc \to e, r_2 : bc \to a, r_3 : aa \to e\}$$

$$R_2 = \{r_4 : A \to a_{out}\}$$

$$R_3 = \{r_5 : B \to b_{out}\}$$

$$R_4 = \{r_6 : C \to c_{out}\}$$

As we have done before, the configuration of the system will be showed below. However, before, we want to justify this feature by comparing it with its biological counterpart. We could think of an object produced by a membrane as a sort of bio-chemical stimulus that is a part of a reaction. If this stimulus does not reach a certain concentration the reaction could not take place. We could think this stimulus will be suppressed if it does not match with a proper receptor in a limited period of time.

When we set a decay time for some symbol objects, the system could have a different behavior (and a different final configuration) depending of the starting time of activity of certain membranes.

For the following examples we have set the decay time $t_D()$ equal to 1 for the symbols objects $\{a,b,c\}$; this means that a symbol object *i* in *D* produced at time *j*, could be used by a rule only from time *j* (included) to time j+1 (excluded). The dynamics can be seen in Figure 4.



Fig. 4. The dynamics in skin membrane of Π_1 .

As you can see, in this case the same system with decay times produces a different output with respect to a standard P system and moreover, the final configuration could be different if we introduce a random delay in membranes starting.

In the previous example we see that the rule r1 could be applied only where are present all the three symbol objects $\{a, b, c\}$, and this situation occurs three times: at time t0, t6 and t12.

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Also the rule r2 is applied but the symbol object a decays before another instance of the symbol could reach the membrane, hence the rule r3 is never carried out.

The following example, described by Figure 5, shows the configuration of the same system, but with a delay of one observer's clock step for the membrane $\mu 4$.



Fig. 5. Dynamics in skin membrane of a system derived from Π_1 by introducing a delay for μ_4 .

Note that no rules of type r1 or r3 could be applied during the computation because, due to the time shifting on membrane $\mu 4$, the symbol objects $\{a, b, c\}$ are never present, at the same time, in the membrane. Hence in this case the halting configuration of the system is \emptyset .

In the last example we want to force the production of the symbol object e by increasing the reactivity frequency of membrane $\mu 4$. The following is the definition of the system Π_2 .

$$\begin{split} \Pi_2 &= (O, D, T, \mu = [_1[_2]_2[_3]_3[_4]_4]_1, \omega_1 = \lambda, \omega_2 = A^5, \omega_3 = B^7, \omega_4 = C^7, \\ &E, t_D, C, R_1, R_2, R_3, R_4, i_O = 1), \end{split}$$

where:

 $O = \{A, B, C, a, b, c, e\} ; D = \{a, b, c\} ; T = O - D = \{A, B, C, e\};$ $E = \{2, 1, 1, 1\}$ $t_D(i) = 1, \forall i \in D$ $C = \{C_1 = 1, C_2 = 3, C_3 = 2, C_4 = 1\}$ $R_1 = \{r_1 : abc \to e, r_2 : bc \to a, r_3 : aa \to e\}$ $\begin{aligned} R_2 &= \{r_4 : A \rightarrow a_{out}\}\\ R_3 &= \{r_5 : B \rightarrow b_{out}\}\\ R_4 &= \{r_6 : C \rightarrow c_{out}\} \end{aligned}$

The resulting dynamics is shown in Figure 6.



Fig. 6. The evolution of symbols in skin membrane of Π_2 .

This behavior is of interest with respect to the modeling of biological mechanisms. We could think that the presence of symbol object e in the halting configuration models certain gene activation. If we simulate several cells with several P systems, derived from Π_1 but with different shifting of their membranes' starting time, we obtain only a little gene activation depending of the starting configuration of the membranes of each system, but if we increase the concentration of our stimulus (represented by the frequency increasing of membrane $\mu 4$) we force the activation of the gene in every system for every permutation of their membranes' starting time.

5 Some Properties of Frequency P Systems

It is easy to state that the class of frequency P systems includes all usual P systems, since we can: assign 0 to the value of energy consumed by each rule, set to 0 the time required by rules to produce their output, have no decaying symbols, have all membranes' clocks with period equal to the unit time of the observer.

But this remark also leads us to a key aspect: a global clock does exist. It is the clock of the observer, which has a frequency which is a multiple of each

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membranes' frequency, and above all, all clocks, in membranes and for the observer, *are* synchronized among themselves, along the grid of time points defined by the observer's clock. Even the eventual offset assigned to membranes is of length equal to an exact multiple of the period of the observer's clock.

This further allows us to state the following:

Proposition: Frequency P systems not consuming energy and without decaying symbols can be simulated by usual P systems with maximal parallelism.

Proof sketch:

We consider observer's clock of simulated frequency P system as the global clock applied in the simulating (usual) P system to each membrane. Membranes of the frequency P system apply rules exactly on (some) ticks of observer's clock. Thus, we could introduce, in the simulating P systems, rules derived from those of the simulated frequency P system, with a left side modified by also requiring the presence a new symbol, associated to the correct (cyclic) counting of global clock's ticks: those new symbols are evolved by specific rules so to represent the waiting of simulated frequency membranes.

(end of Proof sketch)

Nonetheless, our examples show that the dynamics emerging from the behavior of frequency P systems make them an interesting alternative to usual ones as a way to model some basic mechanisms of biological inspiration.

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Active Membrane Systems Without Charges and Using Only Symmetric Elementary Division Characterise P

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Summary. In this paper we introduce a variant of membrane systems with elementary division and without charges. We allow only elementary division where the resulting membranes are identical; we refer to this using the biological term symmetric division. We prove that this model characterises P. This result characterises the power of a class of membrane systems that fall under the so called P conjecture for membrane systems.

1 Introduction

The **P**-conjecture states that recogniser membranes systems with division rules (active membranes [5]), but without charges, characterise **P**. This was shown for a restriction of the model: without dissolution rules [4]. However, it has been shown that systems with dissolution rules and non-elementary division characterise **PSPACE** [2, 8]. In this setting, using dissolution rules allows us to jump from **P** to **PSPACE**. As a step towards finding a bound (upper or lower) on systems with only elementary division rules, we propose a new restriction, and show that it has an upper bound of **P**.

Using our restriction insists that the two membranes that result from an elementary division rule must be identical. This models the usual biological process of cell division [1] and we refer to it using the biological term "symmetric division". We refer to division where the two resulting daughter cells are different by the biological term "asymmetric division". In nature asymmetric division occurs, for example, in stem cells as a way to achieve cell differentiation.

Since our model is uniform via polynomial time deterministic Turing machines it trivially has a lower bound of **P**. All recogniser membrane systems with division rules are upper bounded by **PSPACE** [8]. In this paper we show that systems with symmetric elementary division and without charges are upper bounded by **P**. From an algorithmic point of view, this result result allows one to write a polynomial time

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algorithm that models certain membrane systems which use exponential numbers of membranes and objects.

2 Preliminaries

In this section we define membrane systems and complexity classes. These definitions are from Păun [5, 6], and Sosík and Rodríguez-Patón [8].

2.1 Recogniser membrane systems

Active membranes systems are membrane systems with membrane division rules. Division rules can either only act on elementary membranes, or else on both elementary and non-elementary membranes. An elementary membrane is one which does not contain other membranes (a leaf node, in tree terminology). In Definition 1 we make a new distinction between two types of elementary division rules. When we refer to symmetric division (e_s) we mean division where the resulting two child membranes are identical. When the two child membranes are not identical we refer to the rule as being asymmetric (e).

Definition 1. An active membrane system without charges using elementary division is a tuple $\Pi = (V, H, \mu, w_1, \dots, w_m, R)$ where,

- 1. m > 1 the initial number of membranes;
- 2. V is the alphabet of objects;
- 3. H is the finite set of labels for the membranes;
- 4. μ is a membrane structure, consisting of m membranes, labelled with elements of H;
- 5. w_1, \ldots, w_m are strings over V, describing the multisets of objects placed in the m regions of μ .
- 6. R is a finite set of developmental rules, of the following forms:

a)
$$[a \rightarrow v]_{h},$$

for $h \in H, a \in V, v \in V^{*}$
b) $a[_{h}]_{h} \rightarrow [_{h} b]_{h},$
for $h \in H, a, b \in V$
c) $[_{h} a]_{h} \rightarrow [_{h}]_{h} b,$
for $h \in H, a, b \in V$
d) $[_{h} a]_{h} \rightarrow b,$
for $h \in H, a, b \in V$
(e_{s}) $[_{h} a]_{h} \rightarrow [_{h} b]_{h} [_{h} b]_{h},$
for $h \in H, a, b \in V$
(e_{s}) $[_{h} a]_{h} \rightarrow [_{h} b]_{h} [_{h} b]_{h},$
for $h \in H, a, b, c \in V$.

These rules are applied according to the following principles:

- All the rules are applied in maximally parallel manner. That is, in one step, one object of a membrane is used by at most one rule (chosen in a non-deterministic way), but any object which can evolve by one rule of any form, must evolve.
- If at the same time a membrane labelled with h is divided by a rule of type (e) or (e_s) and there are objects in this membrane which evolve by means of rules of type (a), then we suppose that first the evolution rules of type (a) are used, and then the division is produced. This process takes only one step.
- The rules associated with membranes labelled with h are used for membranes with that label. At one step, a membrane can be the subject of only one rule of types (b)-(e).

In this paper we study the language recognising variant of membrane systems which solves decision problems. A distinguished region contains, at the beginning of the computation, an input – a description of an instance of a problem. The result of the computation (a solution to the instance) is "yes" if a distinguished object **yes** is expelled during the computation, otherwise the result is "no". Such a membrane system is called *deterministic* if for each input a unique sequence of configurations exists. A membrane system is called *confluent* if it always halts and, starting form the same initial configuration, it always gives the same result, either always "yes" or always "no". Therefore, given a fixed initial configuration, a confluent membrane system can non-deterministically choose from various sequences of configurations, but all of them must lead to the same result.

2.2 Complexity classes

Complexity classes have been defined for membrane systems [7]. Consider a decision problem X, i.e. a set of instances $\{x_1, x_2, \ldots\}$ over some finite alphabet such that to each x_1 there is an unique answer "yes" or "no". We consider a *family* of membrane systems to solve each decision problem so that each instance of the problem is solved by some class member.

We denote by $|x_i|$ the size of any instance $x_i \in X$.

Definition 2 (Uniform families of membrane systems). Let \mathcal{D} be a class of membrane systems and let $f : \mathbb{N} \to \mathbb{N}$ be a total function. The class of problems solved by uniform families of membrane systems of type \mathcal{D} in time f, denoted by $\mathbf{MC}_{\mathcal{D}}(f)$, contains all problems X such that:

- There exists a uniform family of membrane systems, $\Pi_X = (\Pi_X(1); \Pi_X(2); \ldots)$ of type \mathcal{D} : each $\Pi_X(n)$ is constructable by a deterministic Turing machine with input n and in time that is polynomial of n.
- Each Π_X(n) is sound: Π_X(n) starting with an input (encoded by a deterministic Turing machine in polynomial time) x ∈ X of size n expels out a distinguished object yes if an only if the answer to x is "yes".
- Each Π_X(n) is confluent: all computations of Π_X(n) with the same input x of size n give the same result; either always "yes" or else always "no".
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• Π_X is f-efficient: $\Pi_X(n)$ always halts in at most f(n) steps.

Semi-uniform families of membrane systems $\Pi_X = (\Pi_X(x_1); \Pi_X(x_2); \ldots)$ whose members $\Pi_X(x_i)$ are constructable by a deterministic Turing machine with input x_i in a polynomial time with respect to $|x_i|$. In this case, for each instance of X we have a special membrane system which therefore does not need an input. The resulting class of problems is denoted by $\mathbf{MC}_{\mathcal{D}}^S(f)$. Obviously, $\mathbf{MC}_{\mathcal{D}}(f) \subseteq$ $\mathbf{MC}_{\mathcal{D}}^S(f)$ for a given class \mathcal{D} and a complexity [3] function f.

We denote by

$$\mathbf{PMC}_{\mathcal{D}} = \bigcup_{k \in \mathbb{N}} \mathbf{MC}_{\mathcal{D}}(O(n^k)), \ \mathbf{PMC}_{\mathcal{D}}^S = \bigcup_{k \in \mathbb{N}} \mathbf{MC}_{\mathcal{D}}^S(O(n^k))$$

the class of problems solvable by uniform (respectively semi-uniform) families of membrane systems in polynomial time. We denote by \mathcal{AM} the classes of membrane systems with active membranes. We denote by \mathcal{EAM} the classes of membrane systems with active membranes and only elementary membrane division. We denote by \mathcal{AM}_{-a}^{0} (respectively, \mathcal{AM}_{+a}^{0}) the class of all recogniser membrane systems with active membranes without charges and without asymmetric division (respectively, with asymmetric division). We denote by $\mathbf{PMC}_{\mathcal{EAM}_{-a}}^{S}$ the classes of problems solvable by semi-uniform families of membrane systems in polynomial time with no charges and only symmetric elementary division.

We let poly(n) be the set of polynomial (complexity) functions of n.

3 An upper bound on $PMC^{S}_{\mathcal{EAM}^{0}}$

In this section we give an upper bound of \mathbf{P} on the membrane class $\mathbf{PMC}_{\mathcal{EAM}_{-a}}^{S}$. We provide a random access machine (RAM) algorithm that simulates this class using a polynomial number of registers of polynomial length, in polynomial time. We begin with an important definition and an informal description of our contribution.

Definition 3 (Equivilance class of membranes). An equivalence class of membranes is a multiset of membranes where: each membrane shares a single parent, each has the same label, and each has identical contents. Further, only membranes without children can be part of an equivalence class of size greater than one; each membrane with one or more children has its own equivalence class of size one.

Throughout the paper, when we say that a membrane system has |E| equivalence classes, we mean that |E| is the minimum number of equivalence classes that includes all membranes of the system.

While it is possible for a computation path of $\mathbf{PMC}^{S}_{\mathcal{EAM}^{0}_{-a}}$ to use an exponential number of equivalence classes, our analysis guarantees that there is another, equally valid, computation path that uses at most a polynomial number of equivalence classes. Our algorithm finds this path in polynomial time. Moreover, via our algorithm, after a single timestep the increase in the number of equivalence classes is never greater than $|E_0||V|$, the product of the number of initial equivalence classes and the number of object types in the system. Since the system is confluent, our chosen computation path is just as valid to follow as any alternative path.

In Section 3.2 we prove that by using our algorithm:

- Type (a) rules do not increase the number of equivalence classes since the rule has the same effect on each membrane of a given equivalence class.
- Type (c) rules do not increase the number of equivalence classes since objects exit all child membranes for the parent membrane (which is already an equivalence class with one membrane).
- Type (d) rules do not increase the number of equivalence classes since the rule is applied to all membranes in the equivalence class. The contents and child membranes are transferred to the parent (already an equivalence class).
- Type (e_s) rules do not increase the number of equivalence classes, the number of membranes in the existing equivalence classes simply increases.

Type (b) rules require a more detailed explanation. In Section 3.3 we show that there is a deterministic polynomial sequential time algorithm that finds a computation path that uses only a polynomial number of equivalence classes.

Our RAM algorithm operates on a number of registers that can be thought of as a data structure (see Section 3.1). The data structure stores the state of the membrane system at each timestep. It compresses the amount of information to be stored by storing equivalence classes instead of explicitly storing all membranes. Each equivalence class contains the number of membranes in the class, a reference to each of the distinct objects in one of those membranes, and the number of copies (in binary) of that object. Type (a) rules could therefore provide a way to create exponential space. However, we store the number of objects in binary thus we store it using space that is the logarithm of the number of objects.

Our RAM algorithm operates in a deterministic way. To introduce determinism we sort all lists of object multisets by object multiplicity, then lexicographically. We sort all equivalence classes by membrane multiplicity, then by label, and then by object. We sort all rules by rules type, matching label, matching object, and then by output object(s). The algorithm iterates through the equivalence classes and applies all rules of type (a), (c), (d), and (e_s) . It then checks to see if any rules of type (b) are applicable. If so, it takes each object in its sorted order and applies it to the relevant membranes in their sorted order.

Theorem 1. $PMC^{S}_{\mathcal{EAM}^{0}} \subseteq P$

The proof is in the remainder of this section.

3.1 Structure of RAM Registers

Our RAM uses a number of binary registers that is a polynomial (poly(n)) of the length n of the input. The length of each register is bounded by a polynomial of n. For convenience our registers are grouped together in a data structure (as illustrated in Figure 1).



Fig. 1. A representation of our polynomial sized registers as a data structure.

Object registers

For each distinct object type v_i , the following registers are used to encode the object in an equivalence class $e_k \in E$.

The register \mathbf{v} represents the type of the object, $v_i \in V$ (see Definition 1). Throughout the computation, the size of the set V is fixed so this register does not grow beyond its initial size.

The copies register is the multiplicity of the distinct object v_i encoded in binary. At time 0 we have $|v_i|$ objects. At time 1 the worst case is that each object evolves via a type (a) rule to give a number of objects that is poly(n). This is an exponential growth function, however, since we store it using binary, the register length does not grow beyond space that is poly(n).

The register used represents the multiplicity v_i objects that have been used already in this computation step. It is always the case that used \leq copies for each object type v_i .

Equivalence class registers

The following registers are used to store information about each equivalence class. To conserve space complexity we only explicitly store equivalence classes (rather than explicitly storing membranes); the number of equivalence classes is denoted |E|.

The register **h** stores the label of equivalence class e_k and is an element of the set H (see Definition 1). The size of register **h** is fixed and is bounded by poly(n).

The register **parent** stores a reference to the equivalence class (a single membrane in this case) that contains this membrane. This value is bounded by the polynomial depth of the membrane structure. Since the depth of the membrane structure is fixed throughout a computation, the space required to store a parent reference is never greater than a logarithm of the depth.

The children register references all of the child equivalence classes of e_k at depth one. Its size is bounded by poly(n) via Theorem 2.

The register **copies** stores the number, denoted $|e_k|$, of membranes in the equivalence class. We store this number in binary. In the worst case, the number that is stored in copies doubles at each timestep (due to type (e_s) rules). Since we store this number in binary we use space that is poly(n).

The register used stores the number of membranes in the equivalence class that have been used by some rule in the current timestep and so this value is $\leq |e_k|$.

Rules registers

The **rules** registers store the rules of the membrane system; their number is bounded by the polynomial |R| and is fixed for all time t. The **rules** registers can not change or grow during a computation. The **type** register stores if the rule is of type (a), (b), (c), (d) or (e_s) . The **lhsObject** register stores the object on the left hand side of the rule. The **lhsLabel** register stores the label on the left hand side of the rule. The **rhsObject** register stores the object on the right hand side of the rule. The **rhsObject** register stores the object on the rule.

3.2 There is a computation path using polynomially many equivalence classes.

In Section 3.2 we prove Theorem 2. Before proceeding to this theorem we make an important observation. Suppose we begin at an initial configuration of a recogniser membrane system. Due to non-determinism in the choice of rules and objects, after t timesteps we could be in any one of a large number of possible configurations. However all computations are *confluent*. So if we are only interested in whether the computation accepts or rejects, then it does not matter which computation path we follow.

Theorem 2 asserts that after a polynomial number of timesteps, there is at least one computation path where the number of equivalence classes of a $\mathbf{PMC}_{\mathcal{EAM}_{-a}}^{S}$ system is polynomially bounded. This is shown by proving that there is a computation path where the application of each rule type (a) to (e_s) , in a single timestep, leads to at most an additive polynomial increase in the number of equivalence classes. **Theorem 2.** Given an initial configuration of a $\text{PMC}_{\mathcal{EAM}_{-a}}^{S}$ system Π with $|E_0|$ equivalence classes and |V| distinct object types, then there is a computation path such that at time $t \in poly(n)$ the number of equivalence classes is $|E_t| = O(|E_0| + t|E_0||V|)$ which is poly(n).

Proof. Base case: From Definition 3, $|E_0|$ is bounded above by the (polynomial) number of membranes at time 0. Thus $|E_0| \in \text{poly}(n)$. Each of lemmata 1 to 5 gives an upper bound on the increase in the number of equivalence classes after one timestep for rule types (a) to (e_s) , respectively. Lemma 2 has an additive increase of $|E_0||V|$ and the other four lemmata have an increase of 0. Thus at time 1 there is a computation path where the number of equivalence classes is $|E_1| \leq |E_0| + |E_0||V|$. (From Definitions 1 and 2, $|V| \in \text{poly}(n)$ and |V| is fixed for all t.)

Inductive step: Assume that $|E_i|$, the number of equivalence classes at time *i*, is polynomial in *n*. Then, while Lemmata 1 to 5, there exists a computation path where $|E_{i+1}| \leq |E_i| + |E_0||V|$.

After t timesteps we have $|E_t| = O(|E_0| + t|E_0||V|)$, which is polynomial in n if t is. \Box

The proofs of the following five lemmata assume some ordering on the set of object types V and on the rules R. For the proof of Lemma 2, we give a specific ordering, however for the other proofs any ordering is valid.

Lemma 1. Given a configuration C_i of a $\mathbf{PMC}^S_{\mathcal{EAM}^0_{-a}}$ system with |E| equivalence classes. After a single timestep, where only rules of type (a) (object evolution) are applied, there exists a configuration C_{i+1} such that $C_i \vdash C_{i+1}$ and C_{i+1} has $\leq |E|$ equivalence classes.

Proof. If a type (a) rule is applicable to an object in a membrane in equivalence class e_k , then the rule is also applicable in exactly the same way to all membranes in e_k . Due to non-determinism in the choice of rules and objects, it could be the case that the membranes in e_k evolve differently. However let us assume an ordering on the object types V and on the rules R. We apply the type (a) rules to objects using this ordering. Then all membranes in an equivalence class evolve identically in a timestep, and no new equivalence classes are created. Thus there is a computation path $C_i \vdash C_{i+1}$ where there is no increase in the number of equivalence classes. □

Observe that type (b) rules have the potential to increase the number of equivalence classes in one timestep by sending different object types into different membranes from the same class. For example, if objects of type v_1 are sent into some of the membranes in an equivalence class, and v_2 objects are sent into the remainder, then we get an increase of 1 in the number of equivalence classes. The following lemma gives an additive polynomial upper bound on this increase.

Lemma 2. Given a configuration C_i of a $\mathbf{PMC}^S_{\mathcal{EAM}^{0}_{-a}}$ system Π with |E| equivalence classes. Let $|E_0|$ be the number of equivalence classes in the initial configuration of Π . Let |V| be the number of distinct object types in Π . After a single timestep, where only rules of type (b) (incoming objects) are applied, there exists a configuration C_{i+1} such that $C_i \vdash C_{i+1}$ and C_{i+1} has $\leq |E| + |E_0||V|$ equivalence classes.

Proof. Let e_j be a parent equivalence class, thus e_j represents one membrane (by Definition 3). If the child membranes of e_j are all parent membranes themselves, then the type (b) communication rule occurs without any increase to the number of equivalence classes. The remainder of the proof is concerned with the other case, where e_j contains a non-zero number of equivalence classes of elementary membranes; by the lemma statement this number is $\leq |E|$.

For the remainder of this proof let $V' \subseteq V$ be the set of distinct object types in the membrane defined by e_j , let $E' \subseteq E$ be the total number of objects in the membrane defined by e_j , let $E' \subseteq E$ be the set of equivalence classes that describe the children of the membrane defined by e_j , and let \mathbb{M} be the total number of membranes that are children of the membrane defined by e_j (therefore \mathbb{M} is the number of membranes in E'). Furthermore we assume that E' is ordered by number of membranes, i.e. we let $E' = (e_1, e_2, \ldots, e_{|E'|})$ where $|e_k|$ is the number of membranes in equivalence class e_k and $\forall k$, $|e_k| \leq |e_{k+1}|$. Similarly we assume that V' is ordered by the number of each object type, i.e. we let $V' = (v_1, v_2, \ldots, v_{|V'|})$ where $|v_k|$ is the multiplicity of objects of type v_k and $\forall k$, $|v_k| \leq |v_{k+1}|$. We divide the proof into two cases.

Case 1: $\mathbb{V} < \mathbb{M}$. Table 1 gives the proof for this case. The "Range" column gives a series of ranges for \mathbb{V} with respect to the numbers of membranes in the elements of E'. The "Total EC" gives the total number of equivalence classes if we place symbols into membranes according to the orderings given on E' and V'. The "Increase EC" column gives the increase in the equivalence classes after one timestep (i.e. |E'| subtracted from the "Total EC" value). Although we omit the tedious details, it is not difficult to show that by using the above ordering and going through the various sub-cases, the worst case for the total number of equivalence classes after one timestep is |E'| + |V'|.

Case 2: $\mathbb{V} \geq \mathbb{M}$. Case 2 is proved using Table 2. Again we omit the details, however it is not difficult to show that by using the above ordering and going through the various sub cases, the worst case for the total number of equivalence classes after one timestep is |E'| + |V'|.

This procedure is iterated over all parent membranes e_j where type (b) rules are applicable, by Definition 3 the number of such parent membranes $\leq |E_0|$. For each parent it is the case that $|V'| \leq |V|$. Thus there is a computation path $C_i \vdash C_{i+1}$ where the increase in the number of equivalence classes is $\leq |E_0||V'| \leq |E_0||V|$. \Box

Lemma 3. Given a configuration C_i of a $\mathbf{PMC}^{S}_{\mathcal{EAM}^{0}_{-a}}$ system with |E| equivalence classes. After a single timestep, where only rules of type (c) (outgoing objects) are applied, there exists a configuration C_{i+1} such that $C_i \vdash C_{i+1}$ and C_{i+1} has $\leq |E|$ equivalence classes.

Proof. If a type (c) rule is applicable to an object in a membrane in equivalence class e_k , then the rule is also applicable in exactly the same way to all membranes in e_k . Due to non-determinism in the choice of rules and objects it could be the case that membranes in e_k eject different symbols. However lets assume an ordering on the object types V and on the rules R. We apply the type (c) rules to objects using this ordering. Then all membranes in an equivalence class evolve identically in one (each membrane ejects the same symbol), and so no new equivalence classes are created from e_k . The single parent of all the membranes in e_k is in an equivalence class e_j which, by Definition 3, contains exactly one membrane and so no new equivalence classes are created from e_j .

Thus there is a computation path $C_i \vdash C_{i+1}$ where there is no increase in the number of equivalence classes. \Box

Interestingly, dissolution is the easiest rule to handle using our approach. The following lemma actually proves something stronger than the other lemmata: dissolution *never* leads to an increase in the number of equivalence classes.

Lemma 4. Given a configuration C_i of a $\mathbf{PMC}_{\mathcal{EAM}_{-a}}^S$ system with |E| equivalence classes. After a single timestep, where only rules of type (d) (membrane dissolution) are applied then for all C_{i+1} , such that $C_i \vdash C_{i+1}$, C_{i+1} has $\leq |E|$ equivalence classes.

Proof. If there is at least one type (d) rule that is applicable to an object and a membrane in equivalence class e_k , then there is at least one rule that is also applicable to all membranes in e_k . Unlike previous proofs, we do not require an ordering on the objects and rules: all membranes in e_k dissolve and equivalence class e_k no longer exists. The single parent of all the membranes in e_k is in an equivalence class e_j which, by Definition 3, contains exactly one membrane and so no new equivalence classes are created from e_j .

Thus for all C_{i+1} , where $C_i \vdash C_{i+1}$, there is no increase in the number of equivalence classes. \Box

Lemma 5. Given a configuration C_i of a $\mathbf{PMC}^S_{\mathcal{EAM}^{0}_{-a}}$ system with |E| equivalence classes. After a single timestep, where only rules of type (e_s) (symmetric membrane division) are applied, there exists a configuration C_{i+1} such that $C_i \vdash C_{i+1}$ and C_{i+1} has $\leq |E|$ equivalence classes.

Proof. If a type (e_s) rule is applicable to an object and membrane in equivalence class e_k , then the rule is also applicable in exactly the same way to all membranes in e_k . Due to non-determinism in the choice of rules and objects it could be the case that membranes in e_k divide using and/or creating different symbols. However lets assume an ordering on the object types V and on the rules R. We apply the type (e_s) rules to objects (and membranes) using this ordering. Then all membranes in

an equivalence class evolve identically in a timestep (each membrane in e_k divides using the same rule). The number of membranes in e_k doubles, but since each new membrane is identical, no new equivalence classes are created from e_k .

Thus there is a computation path $C_i \vdash C_{i+1}$ where there is no increase in the number of equivalence classes. \Box

Range	Total EC	Increase EC
$1 \le \mathbb{V} < e_1 $	E' + V'	V'
$ e_1 \leq \mathbb{V} < e_1 + e_2 $	$\frac{ E' + V' - 1}{ E' + V' }$	$\frac{ V' -1}{ V' }$
$ e_1 + e_2 \le \mathbb{V} < e_1 + e_2 + e_3 $	E' + V' - 2 E' + V' - 1 E' + V'	V' - 2 V' - 1 V'
	:	:
$\sum_{\ell=1}^{ E' -1} e_{\ell} \le \mathbb{V} < \sum_{\ell=1}^{ E' } e_{\ell} $	E' + V' - (E' - 1) + 1 $ E' + V' $	V' - (E' - 1) + 1 V'

Table 1. Increase in the number of equivalence classes (EC) when $\mathbb{V} < \mathbb{M}$. This table is used in the proof of Lemma 2.

	~ .		-
Range	Sub-case	Total EC	Increase
			EC
$0 < \mathbb{M} \le v_1 $	-	E'	0
$ v_1 < \mathbb{M} \le v_1 + v_2 $	$\exists m \text{ s.t. } 1 \le m < E' ,$	E'	0
	$\sum_{\ell} e_{\ell} = v_1 ,$		
	$ v_2 \ge \mathbb{M} - v_1 $		
	$ \nexists m \text{ s.t. } 1 \le m < E' ,$	E' + 1	1
	$\sum^m e_\ell = v_1 ,$		
	$ \overset{\ell_1}{v_2} \ge \mathbb{M} - v_1 $		
$ v_1 + v_2 < \mathbb{M} \le v_1 + v_2 + v_3 $:	:	÷
:	:	•	:
$\sum_{i=1}^{ V' -1} v_\ell < \mathbb{M} \le \sum_{i=1}^{ V' } v_\ell $:	E' + V' - 1	V' - 1
$\ell=1$ $\ell=1$			

Table 2. Increase in the number of equivalence classes (EC) when $\mathbb{V} \geq \mathbb{M}$. This table is used in the proof of Lemma 2.

3.3 RAM Algorithm

Here we outline a RAM algorithm that simulates the computation of any membrane system of the class $\mathbf{PMC}_{\mathcal{EAM}_{-a}^{0}}^{S}$ in polynomial time (in input length *n*). The algorithm operates on any initial configuration and successively applies the evolution rules of the membrane system. It orders all lists of objects, rules and equivalence classes which results in a deterministic computation path that never uses more than polynomial space. The algorithm makes explicit use of the polynomial size bounded registers described in Section 3.1. It also relies on the confluent nature of recogniser membrane systems and simulates only one of the set of valid computation paths. In particular, using the results from Section 3.2, the algorithm chooses a computation path that uses at most a polynomial number of equivalence classes.

Our sort function runs in polynomial time (in input length n) and sorts lists of

- object multisets by object multiplicity, then lexicographically.
- equivalence classes by membrane multiplicity, then by label, and then by objects.
- rules by rules type, matching label, matching object, and then by output object(s).

Since instances of $\mathbf{PMC}^{S}_{\mathcal{EAM}^{0}_{-a}}$ are constructed by polynomial time deterministic Turing machines they are of polynomial size. Also, since all instances of $\mathbf{PMC}^{S}_{\mathcal{EAM}^{0}_{-a}}$ run in polynomial time, if our algorithm simulates it with a polynomial time overhead we obtain a polynomial time upper bound.

Our algorithm begins with a configuration of $\mathbf{PMC}_{\mathcal{EAM}_{-a}}^{S}$ (see Algorithm 1). The input configuration is encoded into the registers of the RAM in polynomial time. The rules of the system are sorted and the algorithm then enters a loop. At each iteration all available rules are applied which simulates a single timestep of the membrane systems computation. The loop terminates when the system ejects a **yes** or **no** object indicating that the computation has halted. Since all instances of $\mathbf{PMC}_{\mathcal{EAM}_{-a}}^{S}$ run in polynomial time, this loop iterates a polynomial number of times.

At each iteration the algorithm iterates through all equivalence classes and applies all applicable rules to it. The worst case time complexity for each function is given. The total time complexity for running the simulation for time t is $O(t|R||E|^2|V|)$.

Algorithm 1: The is the main body of the membrane simulation algorithm. The systems rules are sorted and then applied to the membrane system at each timestep until the system accepts or rejects its input.

	Input: a configuration of $\mathbf{PMC}^{S}_{\mathcal{EAM}^{0}}$	
	Output : The deciding configuration of the system	
	Initialise registers with input system;	
	sortedRules \leftarrow sort(<i>rules</i>);	
O(t)	repeat	
	/* evolve the membrane system one step	*/
O(E)	forall equivalence_class in membraneSystem do	
O(R E V)	ApplyRules(equivalence_class);	
	end	
	until yes or no object is in skin membrane ;	

Function ApplyRules (*equivalence_class*) Applies all applicable rules for an equivalence class for one timestep

	Input: equivalence_class
	Output: equivalence_class after one timestep of computation
	$b_{rules} \leftarrow \emptyset;$
	$b_{ecs} \leftarrow \emptyset;$
	$b_{objs} \leftarrow \emptyset;$
O(R)	forall rule in sortedRules do
	if rule.label matches equivalence_class.label and rule is not type (b) then
O(V)	forall object in sortedObjects do
	if not all copies of object have been used then
	if rule is type (a) then
O(V)	<pre>Apply_a_rule(equivalence_class, object, rule);</pre>
	else if rule is type (c) then
O(1)	<pre>Apply_c_rule(equivalence_class, object, rule);</pre>
	else if rule is type (d) then
O(V)	Apply_d_rule(equivalence_class, object, rule);
	else if rule is type (e_s) then
O(1)	Apply_e_rule(equivalence_class, object, rule);
	end
	end
	end
	end
	if rule is type (b) then
O(E)	forall child_ec in equivalence_class do
	if child_ec.label = rule.lhsLabel and object.used ≥ 1 then
	append child_ec to b_ecs ;
	append object to b_objs ;
O(V E)	Apply_b_rule(b_ecs, b_objs, rule)
	end
	end
	end
	end
$O(V \times E)$	reset all used counters to 0;

Function Apply_a_rule(*equivalence_class, object, rule*) applies a single type (a) rule to instances of an object in an equivalence class. Total time complexity O(|V|).

Input: equivalence_class, object, rule

Output: equivalence_class after a type (a) rule on an object has been applied O(|V|) forall resultingObject *in* rule.outAobjects do

multiplicity of resultingObject in equivalence_class + = the multiplicity of matching object - the number of object.used × the resultingObject.multiplicity ; used number of resultingObject in the equivalence_class + = the multiplicity of resultingObject × object.multiplicity - object.used ;

 \mathbf{end}

decrement object.multiplicity ;
set object.used = object.multiplicity ;

Function Apply_c_rule(*equivalence_class, object, rule*) applies a single rule of type (c) to a membrane. Total time complexity O(1).

 $\mathbf{Input:} \ \mathsf{equivalence_class}$

Output: equivalence_class after a (c) rule have been applied decrement object.multiplicity ; increment object.multiplicity in equivalence_class.parent of the generated object; increment object.used in equivalence_class.parent of the generated object;

increment equivalence_class.used ;

Function Apply_d_rule(equivalence_class, object, rule). This function applies dissolution rules to an equivalence class. It calculates the total number of each object in the equivalence class and adds it to the parent. It also copies the child membranes from the dissolving membrane and adds them to the parents child list. The total time complexity is O(|V|).

	Input: equivalence_class	
	Output : equivalence_class after (d) rule has been applied	
	decrement object.multiplicity;	
	increment object.multiplicity in equivalence_class.parent from the rule;	
	increment object.used in equivalence_class.parent from the rule;	
	<pre>/* move contents of the dissolved membrane to its parent</pre>	*/
O(V)	${f forall}$ move_object in equivalence_class $objects$ ${f do}$	
	add move_object.multiplicity $ imes$ equivalence_class.multiplicity to	
	move_object.multiplicity in equivalence_class.parent ;	
	add move_object.used \times equivalence_class.multiplicity to move_object.used in	
	equivalence_class.parent ;	
	$move_object.multiplicity \leftarrow 0;$	
	$move_object.used \leftarrow 0;$	
	end	
	$equivalence_class.parent.children \gets equivalence_class.parent.children \cup$	
	equivalence_class.children ;	
	equivalence_class.multiplicity $\leftarrow 0$;	
	$equivalence_class \leftarrow \emptyset;$	

Function Apply_es_rule (*equivalence_class, object, rule*). Applies a single rule of type (e_s) to a membrane. Total time complexity O(1).

Input: equivalence_class
Output : equivalence_class after (e_s) rule has been applied
decrement object.multiplicity;
increment object.multiplicity from the rule;
increment object.used from the rule;
increment equivalence_class.used ;
$\texttt{equivalence_class.multiplicity} \gets \texttt{equivalence_class.multiplicity} \times 2;$

Function Apply_b_rules($b_{equivalence_classes}$, $b_{objects}$, b_{rules}). Total time complexity O(|V||E|).

Input: membrane		
Output : membrane after (b) rules have been applied		
$b_objects_sorted \leftarrow sort(b_objects);$		
$b_equivalence_classes_sorted \leftarrow sort(b_equivalence_classes);$		
$O(V)$ forall object <i>in</i> b_objects_sorted do		
$O(E)$ forall equivalence_class in b_equivalence_classes_sorted do		
<pre>if object.multiplicity < equivalence_class.multiplicity then</pre>		
increment equivalence_class.object.used;		
end		
$\begin{array}{c c c c c c c c c c c c c c c c c c c $		
increment equivalence_class.object.used ;		
$equivalence_class.used \leftarrow equivalence_class.multiplicity \ ;$		
subtract equivalence_class.multiplicity from object.multiplicity ;		
end		
end		
end		

4 Conclusion

We have given a \mathbf{P} upper bound on the computational power of one of a number of membrane systems that fall under the so-called \mathbf{P} -conjecture. In particular we consider a variant of membrane systems that allows only symmetric division. This variant can easily generate an exponential number of membranes and objects in polynomial time. Our technique relies on being able to find computation paths that use only polynomial space in polynomial time. It seems that this technique is not naïvely applicable to the case of asymmetric division: it is possible to find examples where all computation paths are forced to use an exponential number of equivalence classes.

Furthermore the result seems interesting since before before now, all models without dissolution rules were upper bounded by \mathbf{P} and all those with dissolution rules characterised **PSPACE**. This result shows that despite having dissolution rules, by using only symmetric elementary division we restrict the system so that it does not create exponential space on all computation paths in polynomial time.

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Balancing Performance, Flexibility and Scalability in a Parallel Computing Platform for Membrane Computing Applications

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Summary. Membrane computing investigates models of computation inspired by certain features of biological cells. To exploit the performance advantage of the large-scale parallelism of membrane computing models, it is necessary to execute them on a parallel computing platform. However, it is an open question whether it is feasible to develop a parallel computing platform for membrane computing applications that significantly outperforms equivalent sequential computing platforms while still achieving acceptable flexibility and scalability. To move closer to an answer to this question, we have investigated a novel approach to the development of a parallel computing platform for membrane computing applications that has the potential to deliver a good balance between performance, flexibility and scalability. This approach involves the use of reconfigurable hardware and an intelligent software component that is able to configure the hardware to suit the specific properties of the membrane computing model to be executed. We have developed a prototype computing platform called Reconfig-P based on the approach. Reconfig-P is the first computing platform of its type to implement parallelism at both the system and region levels. In this paper, we describe Reconfig-P and evaluate its performance, flexibility and scalability. Theoretical and empirical results suggest that the implementation approach on which Reconfig-P is based is a viable means of attaining a good balance between performance, flexibility and scalability in a parallel computing platform for membrane computing applications.

1 Introduction

Membrane computing investigates models of computation inspired by the structural and functional properties of biological cells. Such models have been applied in a variety of domains. To exploit the performance advantage of the large-scale parallelism of membrane computing models, it is necessary to execute them on a parallel computing platform. However, the use of a parallel computing platform instead of a sequential computing platform often comes at the cost of reduced flexibility and scalability.

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The first parallel computing platforms for membrane computing applications to be published [15, 27, 30] do not exhibit sufficient flexibility or scalability. Even so, because research in this area is in its early stages, it is still an open question whether it is feasible to develop a parallel computing platform for membrane computing applications that significantly outperforms equivalent sequential computing platforms while still achieving acceptable flexibility and scalability. To move closer to an answer to this question, it is important to investigate the viability of implementation approaches that have the potential to deliver a good balance between performance, flexibility and scalability.

The research presented in this paper involves an investigation of a novel approach to the development of a parallel computing platform for membrane computing applications. This approach involves the use of reconfigurable hardware and an intelligent software component that is able to configure the hardware to suit the specific properties of the membrane computing model to be executed. We have developed a prototype computing platform called Reconfig-P based on the approach. In this paper, we describe Reconfig-P and evaluate its performance, flexibility and scalability.

The paper is organised as follows. In Section 2, we introduce key concepts and previous research associated with parallel computing platforms for membrane computing applications. In Section 3, we describe Reconfig-P. In Section 4, we evaluate the performance, flexibility and scalability of Reconfig-P. And in Section 5 we draw a conclusion regarding the viability of the implementation approach on which Reconfig-P is based.

2 Background

In this section, we introduce key concepts and previous research associated with parallel computing platforms for membrane computing applications. First, we introduce membrane computing and its applications. Second, we define the attributes of performance, flexibility and scalability in the context of a computing platform for membrane computing applications, explain the significance of these attributes, and indicate the connections that exist between them. Third, we describe the general characteristics of sequential computing platforms, software-based parallel computing platforms and hardware-based parallel computing platforms, and discuss the implications of these characteristics for performance, flexibility and scalability. Finally, we briefly describe existing computing platforms for membrane computing applications and evaluate their performance, flexibility and scalability.

2.1 Membrane computing and its applications

Membrane computing [24, 25] investigates models of computation inspired by certain structural and functional features of biological cells, especially features that arise because of the presence and activity of biological membranes. Biological membranes define compartments inside a cell or separate a cell from its environment. The compartments of a cell contain chemical substances. The substances within a compartment may react with each other or be selectively transported through the membrane surrounding the compartment (e.g., through protein channels) to another compartment as part of the cell's operations.

In a membrane computing model, called a *P* system, multisets of objects (chemical substances) are placed in the regions defined by a hierarchical membrane structure, and the objects evolve by means of reaction rules (chemical reactions) also associated with the regions. The reaction rules are applied in a maximally parallel, nondeterministic manner. The objects can interact with other objects inside the same region or pass through the membrane surrounding the region to neighbouring regions or the cell's environment. These characteristics are used to define transitions between configurations of the system, and sequences of transitions are used to define computations. A computation halts when for every region it is not possible to apply any reaction rule. The input of the computation is defined by the multisets of objects in the initial configuration of the system. The output of the computation may be defined in various ways. For example, the output might be defined as the number of objects located in a particular region in the halting configuration of the system, or as the number of objects emitted to the system's environment during the course of the computation.

Following is a definition of an example P system model. All P systems Π that instantiate the model have the features specified in the definition. The model, which we call the *core P system model*, defines all the essential components of a P system plus two simple and commonly used additional features (namely, catalysts and reaction rule priorities).

 $\Pi = (V, T, C, \mu, w_1, ..., w_m, (R_1, \rho_1), ..., (R_m, \rho_m)),$ where

- V is an alphabet that contains labels for all the *types of objects* in the system;
- $T \subseteq V$ is the *output alphabet*, which contains labels for all the types of objects that are relevant to the determination of the system output;
- $C \subseteq V T$ is the alphabet that contains labels for all the *types of catalysts*, which are the types of objects whose multiplicities cannot change through the application of a reaction rule;
- μ is a hierarchical membrane structure consisting of m membranes, with the membranes (and hence the regions defined by the membranes) injectively labelled by the elements of a given set H of m labels (in this paper, $H = \{1, 2, ..., m\}$);
- each $w_i, 1 \leq i \leq m$, is a string over V that represents the multiset of objects contained in region i of μ in the initial configuration of the system;
- each $R_i, 1 \leq i \leq m$, is a finite set of reaction rules over V associated with the region i of μ ;
- a reaction rule is a pair (r, p), written in the form $r \to p$, where r is a string over V representing a multiset of reactant objects and p is a string over $\{a_{\text{here}}, a_{\text{out}}, a_{\text{in}} \mid a \in V\}$ representing a multiset of product objects, each

of which either (a) stays in the region to which the rule is associated (the subscript 'here' is usually omitted), (b) travels 'out' into the region that immediately contains the region to which the rule is associated, or (c) travels 'in' to one of the regions that is immediately contained by the region to which the rule is associated; and

• each ρ_i is a partial-order relation over R_i which defines the relative priorities of the reaction rules in R_i .

Several P system models have been developed that extend in various ways the core P system model. Examples of additional features found in these extended models include: structured (i.e., non-atomic) objects, membrane creation and dissolution, special inter-region communication rules (e.g., symport and antiport rules), membrane permeability, and electronic charge for objects and membranes. See [24] for a discussion of these features.

P system models have been applied in a variety of domains, including algorithm solving and analysis [1, 19, 20, 23], linguistics [6] and biology [2, 8, 10, 11, 12, 22, 29]. Most existing applications of membrane computing are targeted at the modelling and simulation of biological systems. For example, researchers have modelled the following biological systems as P systems: respiration [10], photosynthesis [22], cell-mediated immunity [12], mechanosensitive channels [2] and protein signalling pathways [29]. In general, biologists are interested in using P systems to perform simulations, rather than to produce computational outputs as in classical computing. That is, they are interested in viewing the configuration-by-configuration evolution of a P system and not only in the final output produced by the P system. Once a biological system has been modelled as a P system, it is possible to simulate the biological system by executing the P system. In many cases, the P system will be executed many times so that the effect of different initial conditions on the evolution of the biological system can be studied.

2.2 Quality attributes of computing platforms for membrane computing applications

The overall quality of a computing platform depends on the extent to which it possesses certain positive attributes, including usability, performance, flexibility, maintainability and scalability. Performance, flexibility and scalability are three of the most important quality attributes for a computing platform for membrane computing applications. Ensuring that a computing platform for membrane computing applications has all three of these attributes to an acceptable degree is a challenge, because a factor that promotes one of the attributes can sometimes demote another one of the attributes. In this section, we define the attributes of performance, flexibility and scalability in the context of a computing platform for membrane computing applications, explain the significance of these attributes, and indicate the connections that exist between them.

Performance

By the performance of a computing platform for membrane computing applications we mean the speed at which it executes P systems; that is, the amount of useful processing it performs per unit time. A suitable measure of the amount of useful processing performed is the number of reaction rule applications performed. Thus the performance of a computing platform for membrane computing applications can be measured in reaction rule applications per unit time.

Flexibility

By the flexibility of a computing platform for membrane computing applications we mean the extent to which it can support the execution of a wide range of P systems. Thus a flexible computing platform for membrane computing applications must be able to adapt to the specific properties of the P system to be executed. The greater the flexibility of the computing platform, the greater the diversity among the P systems in the class of P systems that the computing platform is able to execute.

Scalability

By the scalability of a computing platform for membrane computing applications we mean the extent to which increases in the size of the P system to be executed do not lead to a reduction in the ability of the computing platform to perform its functions or a reduction in the performance of the computing platform. We take the size of a P system as being largely determined by the number of regions and the number of reaction rules it contains.

Connections between performance, flexibility and scalability

The performance of a computing platform for membrane computing applications can be increased by tailoring its implementation to the specific properties of the P systems it is intended to execute. However, the greater the diversity of these P systems, the more difficult it is to efficiently tailor the implementation to their specific properties. Therefore, increasing the performance of the computing platform is likely to come at the cost of reduced flexibility, while increasing the flexibility of the computing platform is likely to come at the cost of reduced performance.

Increasing the flexibility of a computing platform for membrane computing applications involves supporting additional P system features. Naturally, this usually requires the implementation of additional data structures and algorithms. In software-based computing platforms, the implementation of additional data structures is likely to come at the cost of increased memory consumption. In hardware-based computing platforms, the implementation of additional data structures comes at the cost of increased hardware resource consumption, as does the implementation of additional algorithms. Therefore, given that memory resources and hardware resources are limited, implementing additional P system features reduces the maximum size of the P systems that a computing platform for membrane computing applications can execute. Thus increasing the flexibility of a computing platform for membrane computing applications is likely to come at the cost of reduced scalability, while increasing the scalability of such a computing platform is likely to come at the cost of reduced flexibility.

2.3 Types of computing platforms

We identify three major types of computing platforms: sequential computing platforms, software-based parallel computing platforms and hardware-based parallel computing platforms.

Sequential computing platforms are typically based on a software-programmed microprocessor. When such a microprocessor is used, the execution hardware is abstracted by the instruction set architecture, which provides a set of specific instructions that the microprocessor can process to perform computations. This is a very flexible computing solution since it is possible to change the functionality of the computing platform simply by modifying its software — there is no need to modify the hardware configuration. As a result of this flexibility, the same fixed hardware can be used for many applications. However, the flexibility comes at the cost of lower performance. As each instruction needs to be sequentially fetched from memory and decoded before being executed, there is a high execution overhead associated with each individual operation. Furthermore, only one instruction can be executed at a time.

Software-based parallel computing platforms are typically based on a cluster of software-programmed microprocessors. Because the microprocessors execute in parallel, software-based parallel computing platforms can significantly outperform sequential computing platforms for many applications. The microprocessors synchronise their activities by using shared memory or by sending messages to each other (often over a network). Such synchronisation can be very time consuming, and therefore can hinder performance significantly. Increasing the performance of a software-based parallel computing platform involves increasing the amount of parallelism and therefore requires the inclusion of additional microprocessors. However, as the number of microprocessors increases, the overheads associated with synchronisation increase substantially (unless the overall algorithm executed by the computing platform can be neatly partitioned into separate procedures that are largely independent of each other). This fact limits the scalability of softwarebased parallel computing platforms.

Hardware-based parallel computing platforms execute algorithms that have been directly implemented in hardware. In one approach, an application-specific integrated circuit (ASIC) is used. The design of an ASIC is tailored to a specific algorithm. As a consequence, ASICs usually achieve a higher performance than software-programmed microprocessors when executing the algorithm for which they were designed. However, with this higher performance comes reduced flexibility: as the implemented algorithm is fabricated on a silicon chip, it cannot be altered without creating another chip. In another approach, reconfigurable hardware is used. Unlike ASICs, reconfigurable hardware can be modified. Therefore, by using reconfigurable hardware, it is possible to improve on the performance of software-based computing platforms while retaining some of their flexibility. A field-programmable gate array (FPGA) is a type of reconfigurable hardware device. As shown in Figure 1, an FPGA consists of a matrix of logic blocks which are connected by means of a network of wires. The logic blocks at the periphery of the matrix can perform I/O operations. The functionality of the logic blocks and the connections between them can be modified by loading configuration data from a host computer. In this way, any custom digital circuit can be mapped onto the FPGA, thereby enabling it to execute a variety of applications. The digital circuits used in hardware-based parallel computing platforms are specified in hardware description languages. A very popular hardware description language is VHDL. VHDL allows circuits to be specified either in terms of a structural description of the circuit or in terms of low-level algorithmic behaviours of the circuit. Another popular hardware description language is Handel-C. Unlike VHDL, Handel-C does not support the specification of the structural features of a hardware circuit. However, sharing a syntax similar to that of the C programming language, Handel-C allows algorithms to be specified at a very abstract level, and therefore eases the process of designing a circuit for an application.



Fig. 1. The basic architecture of an FPGA.

2.4 Existing computing platforms for membrane computing applications

In this section, we provide a brief survey of existing computing platforms for membrane computing applications.

Sequential computing platforms

As P systems are inherently parallel devices, it is not possible to truly implement them on sequential computing platforms. Nevertheless, sequential computing platforms exist that enable one to simulate in a sequential manner the execution of P systems [4, 7, 9, 13, 16, 21, 26, 28]. For example, Nepomuceno-Chamarro's software tool SimCM [21] is able to simulate the execution of P systems that have the features specified in the definition of the core P system model as well as the feature of membrane dissolution. SimCM is written in Java. It provides a graphical user interface that enables the user to specify and view the evolution of a P system in a visual manner.

Software-based parallel computing platforms

Two research groups have created prototypes of software-based parallel computing platforms for membrane computing applications. Ciobanu and Guo [15] have implemented a simulation of P systems on a Linux cluster using C++ and a library of functions for message-passing parallel computation called the Message Passing Interface (MPI), while Syropoulos and colleagues [30] have implemented a distributed simulation of P systems using Java Remote Method Invocation (RMI). We discuss Ciobanu and Guo's computing platform below.

Ciobanu and Guo's computing platform

Ciobanu and Guo's computing platform is a software program written in C++ that is designed to run on a cluster of computers. The communication mechanism for the computing platform is implemented using MPI. In its prototype form, the computing platform consists of a Linux cluster, in which each node has two 1.4GHz Intel Pentium III CPUs and 1GB of memory, and the nodes are connected by gigabit Ethernet.

Ciobanu and Guo's computing platform supports the execution of a class of P systems that is very similar to the class of P systems that instantiate the core P system model. That is, the computing platform implements most of the basic features of P systems, but does not implement additional features such as membrane creation and dissolution. In the computing platform, each region of a P system is modelled as a separate computational process. Such a process implements the application of the reaction rules in its corresponding region. The processes for the regions in the P system execute in parallel. Communication and synchronisation

between regions is implemented using MPI. The application of reaction rules is performed in rounds. At the end of each round, each region exchanges messages with its parent region and child regions. The reaction rules associated with a region are implemented as threads. If multiple reaction rules require objects of the same type in a round, then only one of them is allowed to consume objects of that type in that round. If two reaction rules do not have relative priorities, which of the two reaction rules is allowed to consume objects first is determined at random.

The process associated with the outermost region of the P system executes a halting detection algorithm. If it detects that the P system has halted, it broadcasts this information to the processes associated with the other regions in the P system.

As the threads for the reaction rules in a region execute on the same node in the cluster, and there are only two processors per node, it would seem that it is impossible for the computing platform to achieve region-level parallelism for anything other than small P systems. To achieve region-level parallelism for larger P systems, it would be necessary to increase the number of processors in a node from two to a number at least equal to the number of reaction rules in the region corresponding to that node. Thus without the inclusion of additional nodes, the computing platform cannot be said to implement region-level parallelism, although it does implement system-level parallelism. Nevertheless, Ciobanu and Guo's use of multithreading as a means of representing the concurrent application of reaction rules within a region is promising, and could be used to implement region-level parallelism if sufficient hardware resources were available.

Ciobanu and Guo do not provide a detailed evaluation of the performance of their computing platform. However, they do report that the performance of the computing platform is somewhat unpredictable. While the execution times exhibited by the computing platform are often acceptable, some execution times are unacceptably long owing to unexpected network congestion. Ciobanu and Guo indicate that the major problem with their computing platform from the point of view of performance is the overhead associated with communication and cooperation between regions. Such communication and cooperation consumes most of the total execution time.

Ciobanu and Guo do not evaluate the scalability of their computing platform. However, it is clear that the scalability of the computing platform is limited to a large extent by the nature of a cluster-based implementation approach. For example, to execute P systems with a large number of regions, the computing platform would have to include a large number of nodes, since there is a one-toone correspondence between regions and nodes. As a consequence, there would be very significant overheads associated with communication and synchronisation between regions, and this would have an adverse impact on the performance of the computing platform.

As it implements only a basic P system model, Ciobanu and Guo's computing platform is not capable of executing P systems that have additional features such as symport and antiport rules. This detracts from its flexibility. Nevertheless, since the existing implementation is expressed at a level of abstraction at which the high-

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level features of a P system are apparent, it seems very feasible that the computing platform could be extended to support additional P system features.

Hardware-based parallel computing platforms

A few researchers have designed digital circuits for particular aspects of P systems (e.g., see [17, 18]). However, to the best of our knowledge, only Petreska and Teuscher [27] have implemented a hardware-based computing platform for membrane computing applications. We discuss Petreska and Teuscher's computing platform below.

Petreska and Teuscher's computing platform

Petreska and Teuscher [27] have developed a full implementation of a particular P system model on reconfigurable hardware. This P system model is similar to the core P system model, except that it also includes the feature of membrane creation and dissolution. The hardware architecture for the specific P system to be executed, which is specified in structural VHDL, is elegant in that it contains only one type of high-level hardware component (a universal component) and interconnections between components of this type.

Petreska and Teuscher have demonstrated the feasibility of implementing some of the important features of membrane computing on reconfigurable hardware. Nevertheless, their computing platform has four main limitations.

First, the computing platform does not exploit the performance advantages of the membrane computing paradigm. This is primarily because it does not implement parallelism at the region level (i.e., the reaction rules in a region are applied sequentially). Achieving region-level parallelism requires the implementation of a scheme for the resolution of conflicts that arise when different reaction rules compete for or produce the same types of objects in the same region at the same time. It is difficult to implement such a scheme efficiently in hardware, especially when a low-level hardware description language is used, and this is perhaps a major reason why Petreska and Teuscher did not attempt to do so. Conflicts do not arise when reaction rules do not compete for or produce the same types of objects in the same region. Nevertheless, in Petreska and Teuscher's computing platform, even such non-conflicting reaction rules must be applied sequentially. Furthermore, if reaction rules from different regions need to update the same multiset, their respective update operations must occur sequentially.

Second, the computing platform is inflexible. As the computing platform uses only one type of high-level hardware component and connects components of this type to build hardware architectures in a fixed manner, the extent to which the hardware architecture for a P system can be tailored to the specific characteristics of the P system is limited.

Third, the computing platform is not extensible. As it is specified at the hardware level in a low-level hardware description language, adding support for additional P system features would require redesigning the hardware for the computing platform directly. This is likely in most cases to be a difficult and time-consuming task, given the dependence of the computing platform on the design of a single universal hardware component. Thus there is limited opportunity to improve the flexibility of the computing platform.

Fourth, the computing platform has limited scalabilty. As there is only a limited ability to tailor the hardware architecture to the specific characteristics of the P system to be executed, the hardware architecture often includes many redundant hardware components. These redundant components unnecessarily consume hardware resources.

As it implements membrane creation and dissolution in addition to the basic P system features included in the core P system model, Petreska and Teuscher's computing platform can execute a wider range of P systems than Ciobanu and Guo's computing platform. So, in this respect, it is more flexible than Ciobanu and Guo's computing platform. However, being specified in a low-level hardware description language, the implementation of Petreska and Teuscher's computing platform is more brittle, and therefore less extensible, than the implementation of Ciobanu and Guo's computing platform. Therefore, unlike in the case of Ciobanu and Guo's computing platform, it seems that it would be very difficult to increase the flexibility of Petreska and Teuscher's computing platform without significantly changing its existing implementation.

3 Description of Reconfig-P

In this section, we describe Reconfig-P, our prototype hardware-based parallel computing platform for membrane computing applications. Being the first computing platform based on reconfigurable hardware to implement parallelism at both the system and region levels, Reconfig-P advances the state-of-the-art in hardware implementations of membrane computing. First, we specify the key features of the novel implementation approach on which Reconfig-P is based, and explain why this implementation approach has the potential to deliver a good balance between performance, flexibility and scalability. Second, we specify the functional requirements of Reconfig-P. Third, we provide an overview of the major components of Reconfig-P and the role of these components in the execution of membrane computing applications. Fourth, we provide an overview of the functionality of P Builder, a software component of Reconfig-P that is responsible for generating customised hardware representations for P systems. Finally, we describe how P Builder represents the fundamental structural and behavioural features of P systems in hardware.

3.1 Implementation approach

Key features of the implementation approach

The implementation approach on which Reconfig-P is based involves

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- use of a reconfigurable hardware platform,
- generation of a customised digital circuit for each P system to be executed, and
- use of a hardware description language that allows digital circuits to be specified at a level of abstraction similar to the level of abstraction at which a general-purpose procedural software programming language (such as C) allows algorithms to be specified.

In the approach, a software component of the computing platform is responsible for analysing the structural and behavioural features of the P system to be executed and producing a hardware description for the P system that is tailored to these features. When determining the hardware description for the P system, the software component aims to maximise performance and minimise hardware resource consumption.

Potential of the implementation approach

The use of reconfigurable hardware opens up the possibility of generating custom digital circuits for P systems. The ability to generate a custom circuit for the P system to be executed makes it possible to design this circuit according to the specific structural and behavioural features of the P system, and therefore facilitates the design of circuits that exhibit good performance and economical hardware usage. Therefore the implementation approach facilitates the development of a computing platform that exhibits good performance and economical hardware usage. For example, because the number of reaction rules in the P system to be executed is known before it is executed, the circuit for the P system can be designed in such a way that it includes exactly that number of processing units to implement the reaction rules. Without the possibility of generating a custom circuit, the circuit for the P system would have to include a fixed number of processing units for reaction rules, and therefore would often include redundant hardware components. Also, because it is possible by inspection of the definitions of the reaction rules in a P system to determine for any two regions in the P system whether it is possible for objects to traverse between these regions, the circuit for a P system can be designed in such a way that the logic that implements object traversal is included only for those inter-region connections over which object traversal is possible.

The fact that digital circuits are specified at a level of abstraction similar to that at which a general-purpose procedural software programming language specifies algorithms, rather than at a level of abstraction that reveals the structure or low-level algorithmic behaviour of the circuits, makes it more feasible to develop a software component that is able to flexibly adapt to the specific features of the P system to be executed when generating a circuit for that P system. The greater the ability of the software component to flexibly adapt to the specific features of the P system to be executed, the greater the range of P systems for which it is capable of generating circuits that exhibit good performance and economical hardware usage. Therefore the implementation approach facilitates the development of a computing platform that exhibits good flexibility. For example, as mentioned in Section 2.4, implementing parallelism at the region level of a P system requires resolving conflicts that may occur when different reaction rules update the same multiplicity values. If a low-level hardware description language were used, it would be very difficult to resolve such conflicts in an efficient manner. The use of a high-level hardware description language makes it more feasible that a solution to the conflict resolution problem can be found.

Because it involves the use of a hardware description language that is incapable of expressing the low-level structure and behaviour of digital circuits, the implementation approach limits the extent to which low-level optimisations of circuits can be carried out. However, it is unlikely that the benefits of customisation and flexibility mentioned above could be achieved if a low-level hardware description language were used.

The above considerations suggest that the implementation approach has the potential to deliver a good balance between performance, flexibility and scalability in a parallel computing platform for membrane computing applications.

3.2 Functional requirements

Reconfig-P is required to execute P systems that instantiate the core P system model on reconfigurable hardware. In addition, to facilitate testing of P system designs, Reconfig-P is required to enable the user to execute a P system in software, and view the configuration-by-configuration evolution of the P system, before generating a hardware circuit for the P system.

It is not a strict requirement that Reconfig-P implement the nondeterminism of P systems. In particular, Reconfig-P is not required to implement the assignment of objects to reaction rules in a nondeterministic manner. Although nondeterminism is very important from a theoretical perspective and can be useful in some applications, many applications of membrane computing do not depend on the nondeterministic aspects of P systems. Therefore we do not regard it as crucial that a computing platform for membrane computing applications implement the nondeterminism of P systems. Even so, it is certainly desirable that a computing platform for membrane computing implement the nondeterminism of P systems. We intend to investigate the feasibility of implementing the nondeterminism of P systems in a future version of Reconfig-P.

3.3 System overview

Figure 2 shows the major components of Reconfig-P and the roles of these components in the execution of a P system.

(1) The user begins using Reconfig-P by writing a P system specification. This specification defines a P system that is described in terms of the core P system



Fig. 2. An overview of Reconfig-P. The shaded region covers the components of Reconfig-P that are transparent to the user.

model. (2) The hardware source code generator called P Builder (which is hidden from the user) processes the input information. (3) P Builder analyses the P system specification, determines a customised hardware representation for the P system, and then generates Handel-C source code that implements the hardware representation. (4) The user can choose to (a) execute the source code in hardware, or (b) simulate the execution of the source code in software. (5) The ability to generate simulation source code enables users to examine their P system design before building a corresponding hardware circuit. (6) The simulation instance (specified by a DLL file) is executed on a *host* computer. The host computer invokes the simulation feature provided by the Celoxica DK Design Suite to allow users to (a) view the evolution of their P system one configuration at a time, or (b) return the output of the simulation in an *output file*. (7) The generation of *hardware execution* source code allows the user, once they have finalised the design of their P system, to build a hardware circuit for the P system. (8) The hardware execution source code is then synthesised into a hardware circuit. A hardware execution instance (specified by a bitstream) can then be executed on a reconfigurable hardware platform (an FPGA). The FPGA communicates with the host computer via a PCI bus. The output of the execution instance is stored in an *output file*, which can then be analysed by the user.

Much of the process of executing a P system is transparent to the user. The shaded region in Figure 2 covers the components of Reconfig-P that are transparent to the user.

3.4 P Builder

P Builder is responsible for implementing the hardware reconfiguration capability of Reconfig-P. It generates customised Handel-C source code for a P system based on the specific characteristics of the P system.

P Builder interprets a simple declarative language which is used by the user to specify P systems. More specifically, P Builder supports the execution of a P system by

- 1. converting a text representation of the P system (the P system specification) into software objects (written in Java);
- 2. converting the object representation into an abstract hardware representation and then into Handel-C source code that implements the abstract hardware representation; and then
- 3. converting the Handel-C source code that implements the abstract hardware representation into a hardware circuit (by invoking Xilinx tools) or initiating a simulation of the abstract hardware representation in software (by invoking the DK Design Suite).

Since P Builder is hidden from the user, the mechanics of the conversion process it performs are transparent to the user. The conversion process is illustrated in Figure 3. The innovation of P Builder lies in the way it converts a P system specification into an abstract hardware representation. In the next section, we describe how P Builder represents the core structural and behavioural features of P systems in hardware.

3.5 Hardware implementation of core P system features

P systems can differ significantly with respect to size, structure and information content. Reconfig-P takes advantage of this fact by configuring the hardware according to the specific requirements of the P system to be executed.

Although P systems can differ significantly, there are certain core features common to all P systems. These include (a) regions and their containment relationships, (b) the mutiset of objects in every region, (c) application of reaction rules, and (d) synchronisation of the application of reaction rules. This section describes how these core features are implemented in hardware.

Regions and their containment relationships

As the evolution of a P system is essentially a matter of the modification of the contents of regions according to certain rules, regions do not need to be explicitly



Fig. 3. P Builder converts the text specification of a P system into an executable hardware circuit or a software simulation of such a hardware circuit. In the intermediate stages of the conversion process, the P system is represented as software objects and then as abstract hardware components.

represented in hardware. Instead, a region is represented in hardware implicitly via its contents. The only inter-region containment relationships that it is important to represent are those between regions between which it is possible for objects to traverse through the application of a reaction rule. These containment relationships are represented implicitly by ensuring that each reaction rule with an 'in' or 'out' target directive has, for each region to/from which it sends/receives objects, access to the multiset of objects in that region.

Multisets of objects in regions

Because the multiplicity values of objects in a region can be accessed by multiple reaction rules simultaneously, the hardware elements that store them should support concurrent accesses. Therefore a multiset is implemented as an array of registers (see Figure 4). Because it is infeasible to predict which types of objects may become available in which regions during the evolution of a P system, the array of registers that represents the multiset of objects in a region contains one register for every type of object in the alphabet of the P system. A common bitwidth is used for all object types (the default width is 8 bits).

Using registers can be expensive if a large amount of data needs to be stored. However, because in the hardware design each register corresponds to the multiplicity of a *type* of object in a region (rather than an individual object), for most P systems only a relatively small amount of data needs to be stored.



Fig. 4. A multiset of objects in a region is implemented as an array of registers.

Reaction rules

A reaction rule is implemented as a processing unit. This processing unit is represented in Handel-C as a potentially infinite while loop that contains code that specifies the processing associated with the application of the reaction rule. If a reaction rule operates on the multiplicity value for a particular object type in a particular region, then the section of the code for its corresponding processing unit that accomplishes this operation contains a reference to the array element representing that multiplicity value.

In a transition of a P system, all the reaction rules in the system complete one instance of execution, which consists of two phases. In the first phase, called the *preparation phase*, objects are assigned to the reaction rules that require them as reactants or catalysts. That is, for each reaction rule in each region, the number of instances of the reaction rule that can be applied is determined. In the second phase, called the *updating phase*, each applicable reaction rule updates one or more multisets of objects according to its definition and the number of instances of the reaction rule that can be applied.

The rest of this section describes the processing performed by the processing units for reaction rules during the preparation and updating phases.



Fig. 5. An illustration of how the structural aspects of a P system are represented as high-level hardware components. (The multiset replication coordinator and extra array elements for object type b in region 3 are included only if the space-oriented conflict resolution strategy is used.)

Preparation phase

In the preparation phase, each reaction rule attempts to obtain as many of each of its required types of object as possible so as to maximise the number of instances of the reaction rule that can be applied in the updating phase. Therefore implementing the preparation phase involves calculating for each reaction rule r the value max-instances_r, which is the maximum number of instances of r that can be applied in the current transition of the P system given (a) the current state of the multiset of objects in its region and (b) the relative priorities and requirements of the other reaction rules in its region. The processing unit corresponding to r performs the calculation.

To calculate max-instances_r, the processing unit for a reaction rule r first calculates for each of its required object types (using integer division) the ratio of the number of available objects of that type in the region of r to the number of objects of that type needed to apply one instance of r. This is done in one clock cycle. It then calculates max-instances_r, which is equal to the minimum ratio calculated in the previous step. The operation of determining the minimum ratio can be represented as a binary tree in which each node corresponds to the execution of a binary MIN operation and executing the MIN operation at the root node gives the value of the minimum ratio. This tree has $\log_2 n$ levels, where n is the sum of the number of reactants and the number of catalysts in the definition of r. The processing unit for r evaluates max-instances_r by first executing in parallel all the MIN operations at the bottom (leaf) level of the tree, then executing in parallel all the MIN operation at the root node to obtain the value of max-instances_r. Therefore calculating max-instances_r takes $\log_2 n$ clock cycles.

If two reaction rules attempt to obtain objects of the same type, then their corresponding processing units execute the relevant operation one after the other according to their relative priorities. (It is assumed that reaction rules that attempt to obtain objects of the same type have been assigned relative priorities.) Otherwise, the processing units for different reaction rules execute in parallel. Therefore the number of clock cycles taken to complete the preparation phase for the entire P system is the maximum number of clock cycles taken by an individual reaction rule, out of all the reaction rules in the P system, to complete its preparation phase.

Updating phase

At the start of the updating phase, the processing unit for a reaction rule r inspects the value of max-instances_r to determine whether r is applicable in the current transition. If max-instances_r = 0, r is inapplicable; otherwise r is applicable. As it takes zero clock cycles to evaluate a conditional expression in Handel-C, determining the applicability of r takes zero clock cycles. The applicability status of r is recorded in the isApplicableFlag of r (see Figure 6). Once the applicability status of each reaction rule has been determined and recorded, the processing unit that coordinates the execution of reaction rules is able to determine whether the P system should halt or continue the updating phase. Assume that the P system should continue the updating phase. If r is inapplicable, the processing unit for rsimply waits for the next transition. If r is applicable, it moves on to the next step of the updating phase.

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In the next step of the updating phase, every instance of every applicable reaction rule is applied. This is implemented by having the processing unit for each applicable reaction rule r bring about the combined effect of the execution of the instances of r. That is, the processing unit decreases/increases certain multiplicity values in certain multiset data structures according to the type, amount and source/destination of the objects consumed/produced by the instances of the reaction rule. For example, in Figure 5, in the next transition of the P system represented at the top of the figure, the processing unit R_2 would decrease by 2 the value stored in the register corresponding to object type a in the multiset data structure for region 2, and increase by 2 the value stored in the register corresponding to object type b in the multiset data structure for region 1.

If a reaction rule includes 'in' target directives, the definition of a P system calls for nondeterministic targeting of objects if there are multiple child regions. Such nondeterministic targeting can be approximated through the use of pseudorandom numbers. Therefore, the hardware design associates a random number generator to each processing unit for a reaction rule that might produce objects in multiple child regions of its own region. When such a processing unit needs to select a destination child region, it invokes its random number generator to obtain a number which identifies the child region to be selected. For example, the processing unit R_1 in Figure 5 invokes its random number generator to determine whether to produce b objects in region 2 or in region 3.

Processing units for reaction rules that do not manipulate any multiplicity values in common execute in parallel during the updating phase. This is not necessarily the case for processing units for reaction rules that do manipulate at least one multiplicity value in common, since without further measures being taken, the parallel execution of such processing units would lead to situations where multiple processing units write to the same register at the same time. Section 3.6 describes two alternative techniques Reconfig-P makes available for the prevention of such situations, and shows the extent to which each technique allows conflicting processing units to execute in parallel during the updating phase. The number of clock cycles taken to complete the updating phase depends on the conflict resolution strategy that is adopted.

Synchronisation of reaction rules

Figure 6 illustrates the synchronisation of reaction rules involved in the execution of a transition of a P system.

The synchronisation of reaction rules is controlled by three sentinels — preparationSentinel, applicableSentinel and updatingSentinel — and corresponding flags associated with each reaction rule — preparationCompleteFlag, updatingCompleteFlag and isApplicableFlag. The sentinels are implemented as 1-bit registers. Each type of flag is implemented as an array of 1-bit registers, each element of which being associated with one reaction rule in the P system. The flags preparationCompleteFlag, isApplicableFlag and updatingCompleteFlag for a reaction rule are used to indicate whether the reaction rule has completed its preparation phase, is applicable, and has completed its updating phase, respectively. The value of each sentinel is the result of performing the AND or OR function to the values of all its corresponding flags. The value of preparationSentinel indicates whether all reaction rules in the P system have completed their preparation phase. The value of applicableSentinel indicates whether at least one reaction rule is applicable (i.e., whether the P system should continue execution). And the value of updatingSentinel indicates whether all applicable reaction rules in the P system have completed their updating phase (and hence whether the P system is ready to proceed to the next transition).

The management of synchronisation is the responsibility of the *rule application* coordinator, a processing unit that executes in parallel with the processing units for the reaction rules (see Figure 5). The rule application coordinator monitors the conditions relevant to synchronisation at each clock cycle.

Let n be the number of reaction rules in the P system. The most natural way to implement the updating of a sentinel value in Handel-C is to write an assignment statement of the form $s = f_1 # f_2 # \dots # f_n$, where s is the variable that stores the sentinel value, each $f_i(1 \le i \le n)$ is a flag for a reaction rule, and # is either the AND operator or the OR operator. Clearly, the greater the number of reaction rules in the P system, the greater the number of AND or OR operations that need to be performed, and therefore the greater the depth of the logic that implements the assignment statement. Since in Handel-C an assignment statement always takes one clock cycle to execute, increasing the depth of the logic that implements an assignment statement can result in a reduction in the clock rate of the FPGA. Therefore, in some situations, decomposing an assignment statement into multiple assignment statements of reduced logical depth can prevent or mitigate a reduction in the clock rate of the FPGA. Whether or not performing such a decomposition is advantageous depends on whether the beneficial effect of reducing the clock cycle length outweighs the detrimental effect of introducing extra clock cycles. For P systems with a large number of reaction rules it might be advantageous to decompose each assignment statement that implements the updating of a sentinel value into multiple assignment statements of reduced logical depth. Therefore Reconfig-P incorporates a logic depth reduction feature. It decomposes an assignment statement with n operands into multiple assignment statements, each of with has at most $x \leq n$ operands. If as many of these assignment statements as possible contain x operands, then the original assignment statement is replaced by $\lfloor \log_r n \rfloor$ assignment statements. The user sets the value of x in order to obtain the best results. By default, Reconfig-P does not perform logic depth reduction (i.e., x = n by default).

3.6 Conflict resolution in the updating phase

As discussed in Section 3.5, a conflict occurs in the updating phase when multiple processing units for reaction rules write to the same register at the same time.



Fig. 6. An illustration of the synchronisation performed to accomplish a transition of a P system.

This occurs if the reaction rules consume or produce the same type of object in the same region in the same transition. As mentioned in Section 2.4, Petreska and Teuscher's hardware implementation avoids the conflict problem by totally sacrificing the parallelism that gives rise to the problem. This is an undesirable strategy, because it hinders performance significantly. Reconfig-P implements two alternative conflict resolution strategies: the *time-oriented strategy* and the *space-oriented strategy*. The time-oriented strategy consumes time, whereas the space-oriented strategy consumes space. Therefore, the best strategy to use depends on whether it is more important to optimise space or time usage. The user selects the strategy to be used.

Both strategies involve determining in software before run-time all of the potential conflicts that might occur between reaction rules, and then generating the hardware circuit for the P system in such a way that all processing units can execute independently without any possibility of writing to the same register at the same time. The task of determining the resource conflicts has a time complexity of $\Theta(n_r n_o)$, where n_r is the number of reaction rules in the P system, and n_o is the number of object types in the alphabet of the P system. Therefore it has a negligible impact on performance. Note that, since the circuit for the P system need be generated only once, the task is performed only once.

In both strategies, potential conflicts are determined through the construction of a *conflict matrix*. Each row of a conflict matrix for a P system is a quadruple (p, q, r, s), where p is an object type in the alphabet of the P system, q is a region in the P system, r is the set of reaction rules whose application results in the consumption and/or production of objects of type p in q, and s — called the *conflict degree* of (p, q) — is the size of r. There is a row for every pair (p, q).

We now describe how the updating phase occurs when (a) the time-oriented strategy is used, and (b) the space-oriented strategy is used.

Time-oriented conflict resolution

In the time-oriented conflict resolution strategy, if two reaction rules need to update the multiplicity value for the same type of object in the same region, then they do so one after the other (the order in which they do so is not important and so is chosen arbitrarily).

Table 1 illustrates the time-oriented strategy. In the table, 'u(p,q)' denotes the operation of updating the multiplicity value of object type p in region q.

The correct interleaving of the various conflicting operations of the processing units is determined by means of analysis of the conflict matrix for the P system before run-time. That is, the Handel-C source code that is generated for the P system specifies the interleaving directly. This is achieved by inserting the appropriate number of single-clock-cycle delay statements in the appropriate places in the source code for the processing units. For example, the code in the processing unit for r_1^3 that updates the multiplicity value of object type a in region 2 is preceded by two delay statements, whereas the corresponding code for object type b in region 3 is not preceded by any delay statements. For the general case, take a quadruple (p, q, r, s) from the conflict matrix for a P system. Assume that the reaction rules $r_1, r_2, \ldots, r_n \in r$ are ordered (for the purpose of conflict resolution) according to the natural ordering of their subscripts. Then the number of delay statements to be inserted immediately before the code in the processing unit for
the reaction rule $r_i \in r$ that updates the multiplicity value of object type p in region q is equal to i - 1. As Table 1 illustrates, the number of clock cycles taken



Fig. 7. An example P system configuration. An arrow labelled by reaction rule r from object type p_1 in region q_1 to object type p_2 in region q_2 means that p_1 is a reactant of r (taken from q_1) and p_2 is a product of r (produced in q_2).

to update the multiplicity value for object type p in region q of a P system is equivalent to the conflict degree of (p,q), which is recorded in the conflict matrix for the P system.

Let k be the highest conflict degree in the conflict matrix for a P system. Then the updating phase for the P system takes k clock cycles to complete when the time-oriented conflict resolution strategy is used.

Table	1.	How	$_{\mathrm{the}}$	processin	ig units	s for	$_{\mathrm{the}}$	react	ion	rules	in	$_{\mathrm{the}}$	Ρ	system	in	Figure	7
execute	e di	ıring	the	updating	phase	of th	e cu	rrent	tra	nsitior	n if	the	tir	ne-orier	nted	l confli	ct
resolut	ion	strat	egy	is used.													

Clock cycle	r_{1}^{1}	<i>r</i> ₁ ²	<i>r</i> ² ₂	r_{1}^{3}	r_{2}^{3}	<i>r</i> ₁ ⁴
1	u(<i>a</i> ,2), u(<i>b</i> ,1), u(<i>c</i> ,1) end	u(<i>f</i> ,2)	u(<i>b</i> ,2), u(<i>e</i> ,2)	u(<i>b</i> ,3)	u(<i>c</i> ,3), u(<i>e</i> ,3) end	u(<i>d</i> ,4), u(<i>e</i> ,2)
2		u(<i>a</i> ,2) end	u(<i>f</i> ,2) end			u(<i>b</i> ,2) end
3				u(<i>a</i> ,2) end		

Space-oriented conflict resolution

In the space-oriented conflict resolution strategy, if n reaction rules need to update the multiplicity value for the same type of object in the same region, then ncopies are made of the register that stores that multiplicity value. The processing units for the conflicting reaction rules are assigned one copy register each, and in the updating phase write to their respective copy registers (see Figure 5 for an example). Once all of the processing units for reaction rules have completed writing to their registers, processing units called *multiset replication coordinators* (each of which is associated with one object type in one region and runs in parallel with the other processing units in Reconfig-P) read the values that have been stored in the copy registers, and set the original registers in the relevant multiset data structures accordingly (again see Figure 5). This step takes one clock cycle to complete. However, for P systems with a large number of object copies, it may be beneficial to perform logic depth reduction (see Section 3.5).

Table 2 illustrates the space-oriented strategy.

Table 2. How the processing units for the reaction rules in the P system in Figure 7 execute during the updating phase of the current transition if the space-oriented conflict resolution strategy is used.

Clock cycle	r_1^1	<i>r</i> ₁ ²	r_{2}^{2}	<i>r</i> ₁ ³	r_{2}^{3}	<i>r</i> ⁴
1	u(<i>a</i> ,2), u(<i>b</i> ,1), u(<i>c</i> ,1) end	u(<i>f</i> ,2) u(<i>a</i> ,2) end	u(<i>b</i> ,2), u(<i>e</i> ,2) u(<i>f</i> ,2) end	u(<i>b</i> ,3) u(<i>a</i> ,2) end	u(<i>c</i> ,3), u(<i>e</i> ,3) end	u(<i>d</i> ,4), u(<i>e</i> ,2) u(<i>b</i> ,2) end
2	Multiset replication coordinators update original registers in relevant multiset data structures					

4 Evaluation of Reconfig-P

In this section, we evaluate the performance, flexibility and scalability of Reconfig-P. First, we present a theoretical analysis of the performance of Reconfig-P. Then we present and discuss empirical results that give insight into the performance and hardware resource usage of Reconfig-P. Finally we comment on the flexibility of Reconfig-P.

4.1 Theoretical evaluation of the performance of Reconfig-P

Figure 8 presents the time complexity of the parallel algorithm executed by Reconfig-P (in both the time-oriented and space-oriented modes) as well as the time complexity of the sequential algorithm used in sequential implementations of

membrane computing. Table 3 illustrates the relative theoretical performances of (a) the sequential algorithm, (b) an algorithm that implements parallelism only at the system level (as in Petreska and Teuscher's computing platform), (c) the timeoriented parallel algorithm executed by Reconfig-P, and (d) the space-oriented parallel algorithm executed by Reconfig-P. It does so by evaluating their time complexities for example P systems.

In Table 3, larger and larger P systems are derived from one initial basic P system using either horizontal cascading or vertical cascading. In horizontal cascading, more and more regions are added, but the number of reaction rules per region is held constant. In vertical cascading, the number of reaction rules per region increases, but the number of regions is held constant. The following assumptions, deemed to represent the average case, are made: (a) there are 20 object types; (b) each reaction rule has four reactant object types, four catalyst object types and four product object types; (c) there are conflicts on 20% of the object types in the preparation phase; and (d) there are conflicts on 60% of the object types in the updating phase. Assumption (d) gives rise to a k value for each P system, which is the highest conflict degree in the conflict matrix for the P system. P systems are also assigned an arbitrary a value, which is the percentage of reaction rules that are applicable in a transition on average.

Table 3 clearly demonstrates the superior speed of the algorithm executed by Reconfig-P over both the sequential algorithm and the algorithm with one level of parallelism. When horizontal cascading is applied, the time-oriented and space-oriented algorithms both show exceptional scalability. When the k value is small and reaction rules are evenly distributed across regions, the time-oriented algorithm is more effective than the space-oriented algorithm because it uses less space while achieving similar speeds. When vertical cascading is applied, the timeoriented and space-oriented algorithms are significantly faster than the algorithm with one level of parallelism. The space-oriented algorithm is faster than the timeoriented algorithm. The main reason is that, whereas increasing the k value reduces the degree of parallelism in the updating phase when the time-oriented algorithm is used, this is not the case when the space-oriented algorithm is used.

4.2 Empirical evaluation of the performance and scalability of Reconfig-P

We have conducted a series of experiments to investigate the performance and hardware resource usage of Reconfig-P. In this section, we present and discuss the results of these experiments, and evaluate the performance and scalability of Reconfig-P in light of these results.

Details of the experiments

Table 4 shows the P systems that were executed in the experiments. Each P system was constructed by first taking n copies of the basic P subsystem shown

R	Б				Number of clock	cycles per transitior	١		
e g i o n s	g u i l o e n s s		a (%)	Sequential	1-level parallelism	2-level parallelism (time-oriented)	2-level parallelism (space-oriented)		
Horizontal cascading									
10	50	3	30	470	25	10	10		
10	50	3	70	630	28	10	10		
50	250	3	30	2350	27	12	13		
50	250	3	70	3150	30	12	13		
100	500	3	30	4700	27	12	13		
100	500	3	70	6300	30	12	13		
				Vert	ical cascading				
10	50	3	30	470	25	10	10		
10	50	3	70	630	28	10	10		
10	250	15	30	2350	186	36	25		
10	250	15	70	3150	351	36	25		
10	500	30	30	4700	606	66	40		
10	500	30	70	6300	1206	66	40		

Table 3. An illustration of the time complexity results presented in Figure 8.

at the top-right of Figure 9, then cascading these copies in a horizontal, vertical or horizontal and vertical manner (as shown in Figure 9), and finally placing the copies into the region shown at the top-left of Figure 9. The value of n is a measure of the size of the constructed P system; the larger the value of n, the larger the P system. Thus in the experiments a series of P systems of different sizes and different structures were executed.

Table 4 lists a C value for each P system. The C value for a P system is a measure of the amount of conflict that exists between reaction rules in the P system. More specifically, C is the sum of the conflict degrees of all pairs (p,q) for the P system, where p is an object type, q is a region and the conflict degree of the pair is greater than 1.

The target circuit for the experiment was the Xilinx Virtex-II XC2V6000FF11 52-4, and the Handel-C code for the P systems was synthesised, placed and routed using Xilinx tools.

P	n	Regions	Horizontal cascading			Vertical cascading			Horizontal and vertical cascading		
oyotom			Rules	С	k	Rules	С	k	Rules	С	k
1	1	4	11	27	7	11	27	7	11	27	7
2	2	7	21	53	7	22	54	7	22	54	7
3	4	13	41	97	7	44	108	7	42	106	7
4	8	25	81	193	9	88	216	7	83	211	7
5	16	49	161	377	17	176	432	7	165	415	7

Table 4. Details of the P systems used in the experiments.

Evaluation of the hardware resource usage of Reconfig-P

Table 5 shows experimental data related to the hardware resource usage of Reconfig-P, both when it executes in time-oriented mode and when it executes in space-oriented mode. We use the number of LUTs (lookup tables) on the circuit generated for a P system as the measure of the hardware resource usage of Reconfig-P for that P system. We also record the percentage of the LUTs available on the FPGA that is used by the circuit, because this percentage provides an indication of the extent to which current FPGA technology meets the hardware resource requirements of Reconfig-P.

		Time-orier	nted mode	Space-oriented mode					
P system	n	Number of LUTs	% of LUTs	Number of LUTs	% of LUTs				
		Horizo	ontal cascading	g					
1	1	1046	1.55%	1102	1.63%				
2	2	1667	2.47%	1801	2.66%				
3	4	3058	4.52%	3248	4.81%				
4	8	5752	8.51%	6060	8.97%				
5	16	11106	16.43%	11719	17.34%				
	Vertical cascading								
1	1	1046	1.55%	1102	1.63%				
2	2	1959	2.9%	2075	3.07%				
3	4	3570	5.32%	3790	5.65%				
4	8	6771	10.09%	7344	10.87%				
5	16	13207	19.79%	14486	21.43%				
		Horizontal a	nd vertical cas	cading					
1	1	1046	1.55%	1102	1.63%				
2	2	1934	2.94%	1934	2.94%				
3	4	3479	5.16%	3689	5.46%				
4	8	6597	9.76%	7293	10.79%				
5	16	12780	19.41%	13360	20.1%				

Table 5. Experimental results related to the hardware resource usage of Reconfig-P inboth the time-oriented and space-oriented modes.

Figures 10 and 11 illustrate the experimental data in graphical form. We make the following observations:

• The hardware resource usage of Reconfig-P scales linearly with respect to the size of the P system executed (i.e., with respect to n). This is as good as can reasonably be expected, and indicates that Reconfig-P is scalable with respect to hardware resource usage.

- The type of cascading employed in the construction of the P system that is executed has little effect on hardware resource usage.
- For all P systems, Reconfig-P uses less than 22% of the LUTs available on the FPGA. Given that the largest P system has 49 regions and 176 reaction rules, this is an impressive result. Not only does it strongly suggest that current FPGA technology meets the hardware resource requirements of Reconfig-P, it also indicates that it would be feasible to extend Reconfig-P to support P system features not covered by the core P system model.
- Reconfig-P uses only slightly more hardware resources in space-oriented mode than in time-oriented mode. This suggests that, at least for the P systems executed in the experiments, multiset replication has only a relatively small effect on hardware usage. Indeed, even for P systems with C > 400 and therefore with more than 400 copies of multiplicity values, the hardware resources consumed by Reconfig-P to store, access and coordinate these copies is relatively small.

In summary, the experimental results indicate that Reconfig-P makes efficient use of hardware resources, and therefore is scalable with respect to hardware resource usage. The fact that less than 22% of the available hardware resources is used for even relatively large P systems augurs well for the flexibility of Reconfig-P (see Section 4.3 for more on this point).

Evaluation of the performance of Reconfig-P

Table 6 shows the number of clock cycles that Reconfig-P executes per transition for each of the P systems used in the experiments, both when it executes in time-oriented mode and when it executes in space-oriented mode. The values shown in the table were determined empirically.

We make the following observations:

- Nearly all of the P systems used in the experiments have k = 7. When it executes in time-oriented mode, Reconfig-P takes 14 clock cycles to execute a transition if k = 7. When k = 9, it takes 16 clock cycles, and when k = 17, it takes 24 clock cycles. This is exactly as expected, given that the number of clock cycles taken to execute the updating phase across all regions in a P system is equal to k when the time-oriented conflict resolution strategy is used (see Section 3.6).
- When it executes in space-oriented mode, Reconfig-P takes 7 clock cycles to execute a transition for all the P systems used in the experiments. This suggests that when the space-oriented conflict resolution strategy is used, Reconfig-P shows exceptional scalability with respect to the number of clock cycles it takes per transition.
- Overall, Reconfig-P shows excellent scalability with respect to the number of clock cycles it takes per transition. However, when it executes in time-oriented

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P system	n	k	Number of clock cycles per transition					
1 System			Time-oriented mode	Space-oriented mode				
		Horizor	ntal cascading					
1	1	7	14	7				
2	2	7	14	7				
3	4	7	14	7				
4	8	9	16	7				
5	16	17	24	7				
	Vertical cascading							
1	1	7	14	7				
2	2	7	14	7				
3	4	7	14	7				
4	8	7	14	7				
5	16	7	14	7				
	Horiz	ontal an	d vertical cascadir	Ig				
1	1	7	14	7				
2	2	7	14	7				
3	4	7	14	7				
4	8	7	14	7				
5	16	7	14	7				

Table 6. Experimental data related to the number of clock cycles Reconfig-P takes toexecute one P system transition.

mode and horizontal cascading is applied, increases in the size of the P system to be executed can lead to increases in the value of k, and hence increases in the number of clock cycles per transition.

• Reconfig-P takes considerably less clock cycles per transition when it executes in space-oriented mode than when it executes in time-oriented mode. This is as expected, given that the updating phase executes in a maximally parallel manner when the space-oriented conflict resolution strategy is used, but only in a partially parallel manner when the time-oriented conflict resolution strategy is used.

Table 7 shows the performance of Reconfig-P for each of the P systems used in the experiments, both when it executes in time-oriented mode and when it executes in space-oriented mode. It also shows, for the sake of comparison, the corresponding results for a software-based sequential computing platform (i.e., a Java simulator for the core P system model).

Figures 12, 13 and 14 illustrate the experimental data in graphical form. We make the following observations:

	n	Reaction rule applications per second								
P system		Software-based sequential computing platform	Reconfig-P (space- oriented mode)	Reconfig-P (time- oriented mode)						
	Horizontal cascading									
1	1	3.7×10^{5}	57×10^5	$53 imes 10^5$						
2	2	4.5×10^{5}	120×10^5	110×10^5						
3	4	$5.3 imes 10^5$	230×10^5	210×10^5						
4	8	$4.7 imes 10^5$	460×10^5	410×10^{5}						
5	16	$3.5 imes 10^5$	$910 imes 10^5$	$710 imes 10^5$						
Vertical cascading										
1	1	$3.7 imes 10^5$	57×10^5	$53 imes 10^5$						
2	2	$5.2 imes 10^5$	126×10^5	116×10^5						
3	4	4.3×10^{5}	250×10^5	230×10^5						
4	8	$3.2 imes 10^5$	$500 imes 10^5$	460×10^5						
5	16	2×10^5	$1000 imes 10^5$	$890 imes 10^5$						
		Horizontal an	d vertical cascading							
1	1	$3.7 imes 10^5$	57×10^5	$53 imes 10^5$						
2	2	$4.1 imes 10^5$	126×10^5	115×10^5						
3	4	$4.5 imes 10^5$	240×10^5	220×10^5						
4	8	$3.7 imes 10^5$	470×10^5	430×10^5						
5	16	$2.7 imes 10^5$	930×10^5	820×10^5						

Table 7. Experimental data related to the performance of Reconfig-P (in both the spaceoriented and time-oriented modes) and a software-based sequential computing platform.

- Reconfig-P executes P systems significantly faster than the software-based sequential computing platform (from 14 to 500 times faster). The larger the P system that is executed, the greater the extent to which Reconfig-P outperforms the sequential computing platform. This is as expected, because larger P systems have more regions and more reaction rules and therefore more opportunity for parallelism at both the system and region levels.
- In general, the performance of Reconfig-P in both the space-oriented and timeoriented modes increases linearly with respect to the size of the P system that is executed. This is a good result, because it indicates that as the size of the P system to be executed increases, Reconfig-P is able to take advantage of the increased opportunities for parallelism. That is, there does not appear to be any significant problems of scale in the hardware design (e.g., nonlinearly growing logic depths in certain parts of the hardware circuit that would reduce the clock rate of the FPGA).
- Reconfig-P performs better in space-oriented mode than in time-oriented mode, although only by approximately 10%. This relatively small difference is a con-

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sequence of the fact that the various P systems used in the experiments have small k values. If the k values were larger, we would expect to observe a more pronounced difference in performance between the space-oriented and time-oriented modes.

In summary, the experimental results indicate that Reconfig-P achieves very good performance.

4.3 Evaluation of the flexibility of Reconfig-P

In its current prototype form, Reconfig-P supports the basic P system features covered by the core P system model. Therefore it is not able to execute P systems that include additional features such as structured objects and membrane permeability. This counts against its flexibility. However, there is good reason to believe that Reconfig-P can be extended to support additional P system features. As we have observed, Reconfig-P exhibits exceptionally economic hardware resource usage: for the P systems used in the experiments, approximately 75% of the available hardware resources are left unused. Thus there is ample space on the FPGA for the inclusion of additional data structures and logic required for the implementation of additional features. Furthermore, the fact that Reconfig-P is implemented in a high-level hardware description language should ease the process of incorporating additional features into the existing implementation.

5 Conclusion

By developing Reconfig-P, we have demonstrated that it is possible to efficiently implement both the system-level and region-level parallelism of P systems on reconfigurable hardware and thereby achieve significant performance gains.

Theoretical results demonstrate that the algorithm executed by Reconfig-P is significantly faster than the sequential algorithm used in sequential implementations of membrane computing. Empirical results show that for a variety of P systems Reconfig-P achieves very good performance while making economical use of hardware resources. And there is good reason to believe that Reconfig-P can be extended in the future to support additional P system features. Therefore, there is strong evidence that the implementation approach on which Reconfig-P is based is a viable means of attaining a good balance between performance, flexibility and scalability in a parallel computing platform for membrane computing applications.

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Definitions

$$\begin{split} M &= \{m_1, m_2, ..., m_n\} \text{ is the set of membranes in the P system. } V &= \{o_1, o_2, ..., o_v\} \text{ is the alphabet of the P system. } R_{m_x} &= \{r_{1,m_x}, r_{2,m_x}, ..., r_{k_{m_x},m_x}\} \text{ is the set of reaction rules in the region defined by membrane } m_x. r_{y,m_x} \text{ is the } y^{\text{th}} \text{ reaction rule in the region defined by membrane } m_x. r_{y,m_x} \text{ is applicable. } n^{\text{R}}(r_{y,m_x}), n^{\text{C}}(r_{y,m_x}) \text{ and } n^{\text{P}}(r_{y,m_x}) \text{ denote the number of reactant, catalyst and product object types in reaction rule } r_{y,m_x}, \text{ respectively. } M_{m_x}^{\text{UPDATE}} : V \longrightarrow R_{m_x} \text{ maps each object type in the region defined by membrane } m_x \text{ to the set of reaction rules that might update its multiplicity. } Max_{i=1}^n x_i \text{ is the function that returns the maximum value in the set } \{x_1, x_2, ..., x_n\}. \text{ In the following performance analysis, } e \text{ denotes the time taken to execute the preparation phase } (p), the updating phase <math>(u)$$
 and (in the parallel algorithm) synchronisation operations (s). Synchronisation operations include updates of the sentinels preparation. Times are measured in clock cycles. \end{split}

Sequential algorithm

$$e^{\text{SEQ}} = \sum_{i=1}^{n} \sum_{j=1}^{k_{m_i}} = \begin{cases} p^{\text{SEQ}}(r_{j,m_i}) &, \text{ if } r_{j,m_i} \text{ is not applicable} \\ p^{\text{SEQ}}(r_{j,m_i}) + u^{\text{SEQ}}(r_{j,m_i}) &, \text{ if } r_{j,m_i} \text{ is applicable} \end{cases}$$

where

$$p^{\text{SEQ}}(r_{y,m_x}) = n^{\text{R}}(r_{y,m_x}) + n^{\text{C}}(r_{y,m_x}) - 1$$

and

$$u^{\text{SEQ}}(r^a_{y,m_x}) = n^{\text{R}}(r^a_{y,m_x}) + n^{\text{P}}(r^a_{y,m_x})$$

Parallel algorithm

$$e^{\text{PAR}} = Max_{i=1}^{n} Max_{j=1}^{k_{m_{i}}} p^{\text{PAR}}(r_{j,m_{i}}) + Max_{i=1}^{n} Max_{j=1}^{k_{m_{i}}} u^{\text{PAR}}(r_{j,m_{i}}^{a}) + s^{\text{PAR}},$$

where

$$p^{\text{PAR}}(r_{y,m_x}) = \begin{cases} \sum_{s=1}^{y} \log_2(n^{\text{R}}(r_{s,m_x}) + n^{\text{C}}(r_{s,m_x})), \\ \text{if } r_{s,m_x} \text{ has an assigned priority} \\ \log_2(n^{\text{R}}(r_{y,m_x}) + n^{\text{C}}(r_{y,m_x})), \\ \text{if } r_{y,m_x} \text{ does not have an assigned priority} \end{cases}$$

and

$$Max_{i=1}^{n}Max_{j=1}^{k_{m_{i}}}u^{\text{PAR}}(r_{j,m_{i}}^{a}) = \begin{cases} Max_{i=1}^{n}Max_{j=1}^{v} \left| M_{m_{i}}^{\text{UPDATE}}(o_{j}) \right|, \\ \text{if the time-oriented strategy is used} \\ 1, \\ \text{if the space-oriented strategy is used} \end{cases}$$

and

$$s^{\text{PAR}} = \begin{cases} 2(\log_x \left[\sum_{i=1}^n |R_{m_i}|\right] - 1) &, \text{ if the time-oriented strategy is used.} \\ 2(\log_x \left[\sum_{i=1}^n |R_{m_i}|\right] - 1) + 1 &, \text{ if the space-oriented strategy is used.} \end{cases}$$

Fig. 8. The time complexities of the parallel algorithm executed by Reconfig-P and the sequential algorithm executed by sequential implementations of membrane computing.

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Fig. 9. Each P system used in the experiments was constructed by first cascading n copies of a basic P subsystem in a horizontal, vertical or horizontal and vertical manner, and then placing these copies into an outermost region.



Fig. 10. An illustration of the experimental results related to the hardware resource usage of Reconfig-P in space-oriented mode.



Fig. 11. An illustration of the experimental results related to the hardware resource usage of Reconfig-P in time-oriented mode.





Fig. 12. An illustration of the experimental data related to the performance of Reconfig-P (in both the space-oriented and time-oriented modes) and a software-based sequential computing platform when horizontal cascading is applied.



Fig. 13. An illustration of the experimental data related to the performance of Reconfig-P (in both the space-oriented and time-oriented modes) and a software-based sequential computing platform when vertical cascading is applied.



Fig. 14. An illustration of the experimental data related to the performance of Reconfig-P (in both the space-oriented and time-oriented modes) and a software-based sequential computing platform when horizontal and vertical cascading is applied.

Multigraphical Membrane Systems: a Visual Formalism for Modeling Complex Systems in Biology and Evolving Neural Networks

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Summary. A concept of a (directed) multigraphical membrane system, akin to membrane systems in [9] and [10], for modeling complex systems in biology, evolving neural networks, perception, and brain function is introduced.

1 Introduction

Statecharts described in [7] and their wide applications, including applications in system biology, cf. [6], and the formal foundations for natural reasoning in a visual mode presented in [11] challenge a prejudice against visualizations in exact sciences that they are heuristic tools and not valid elements of mathematical proofs.

We introduce a concept of a (directed) multigraphical membrane system to be applied for modeling complex systems in biology, evolving neural networks, perception, and brain function. A precise mathematical definition of this concept and its topological representation by Venn diagrams and the usual graph drawings constitute a kind of visual formalism related to that discussed in [7]. The concept of a multigraphical membrane system is some new variant of the notion of a membrane system in [9] and [10].

2 Multigraphical Membrane Systems

Membrane system in [9] and [10] are simply finite trees with nodes labeled by multisets, where the finite trees have a natural visual presentation by Venn diagrams.

We introduce (*directed*) multigraphical membrane systems to be finite trees with nodes labeled by (directed) multigraphs.

We consider directed multigraphical membrane systems of a special feature described formally in the following way.

A sketch-like membrane system S is given by:

- its underlying tree $\mathbb{T}_{\mathcal{S}}$ which is a finite graph given by the set $V(\mathbb{T}_{\mathcal{S}})$ of vertices, the set $E(\mathbb{T}_{\mathcal{S}}) \subseteq V(\mathbb{T}_{\mathcal{S}}) \times V(\mathbb{T}_{\mathcal{S}})$ of edges, and the root r which is a distinguished vertex such that for every vertex v different from r there exists a unique path from v into r in $\mathbb{T}_{\mathcal{S}}$, where for every vertex v we define $\operatorname{rel}(v) = \{v' \mid (v', v) \in E(\mathbb{T}_{\mathcal{S}})\}$ which is the set of vertices immediately related to v;
- its family $(G_v | v \in V(\mathbb{T}_S))$ of finite directed multigraphs for G_v given by the set $V(G_v)$ of vertices, the set $E(G_v)$ of edges, the source function $s_v : E(G_v) \to V(G_v)$, and the target function $t_v : E(G_v) \to V(G_v)$ such that the following conditions hold:
 - 1) $V(G_v) = \{v\} \cup \operatorname{rel}(v),$
 - 2) $E(G_v)$ is empty for every *elementary* vertex v, i.e. such that rel(v) is empty,
 - 3) for every *non-elementary* vertex v, i.e. such that rel(v) is a non-empty set, we have
 - (i) $G_v(v, v')$ is empty for every $v' \in V(G_v)$,
 - (ii) $G_v(v', v)$ is a one-element set for every $v' \in \operatorname{rel}(v)$, where $G_v(v_1, v_2) = \{e \in E(G_v) \mid s_v(e) = v_1 \text{ and } t_v(e) = v_2\}.$

For every non-elementary vertex v of $\mathbb{T}_{\mathcal{S}}$ we define:

• the *v*-diagram Dg(v) to be that directed multigraph which is the restriction of G_v to rel(v), i.e. V(Dg(v)) = rel(v),

$$E(\mathrm{Dg}(v)) = \big\{ e \in E(G_v) \, | \, \{s_v(e), t_v(e)\} \subseteq \mathrm{rel}(v) \big\},\$$

the source and target functions of Dg(v) are the obvious restrictions of s_v, t_v to E(Dg(v)), respectively,

• the *v*-cocone to be a family $(e_{v'} | v' \in \operatorname{rel}(v))$ of edges of G_v such that $s_v(e_{v'}) = v'$ and $t_v(e_{v'}) = v$ for every $v' \in \operatorname{rel}(v)$.

By a model of a sketch-like membrane system S in a category \mathbb{C} with finite colimits we mean a family of graph homomorphisms $h_v : G_v \to \mathbb{C}$ (v is a nonelementary vertex of \mathbb{T}_S) such that $h_v(v)$ is a colimit of the diagram $h_v \upharpoonright \mathrm{Dg}(v) :$ $\mathrm{Dg}(v) \to \mathbb{C}$ and $(h_v(e_{v'}) | v' \in \mathrm{rel}(v))$ is a colimiting cocone for the v-cocone $(e_{v'} | v' \in \mathrm{rel}(v))$, where $h_v \upharpoonright \mathrm{Dg}(v)$ is the restriction of h_v to $\mathrm{Dg}(v)$.

For all categorical and sketch theoretical notions like graph homomorphism, colimit of the diagram, and colimiting cocone we refer the reader to [3].

The idea of a sketch-like membrane system and its categorical model is a special case of the concept of a sketch and its model described in [3] and [8], where one finds that sketches can serve as a visual presentation of some data structure and data type algebraic specifications. On the other hand the idea of a sketch-like membrane system is a generalization of the notion of ramification used in [4] and [5] to investigate hierarchical categories with hierarchies determined by iterated colimits understood as in [4]. Hierarchical categories with hierarchies determined by iterated colimits are applied in [1] and [5] to describe various emergence phenomena in biology and general system theory. The iterated colimits identified with binding of patterns in neural net systems are expected in [5] to be applied in the investigations of binding problems in vision systems (associated with perception and brain function) in [12] and [13], hence the notion of sketch-like membrane system is aimed to be a tool for these investigations.

More precisely, sketch-like membrane systems are aimed to be presentations of objects of state categories of Memory Evolutive Systems in [4] and [5] similarly like strings of digits serve for presentation of numbers, where these state categories are hierarchical categories with hierarchies determined by iterated colimits. Hierarchical shape of sketch-like membrane systems and their categorical semantics reflect iterated colimit feature of objects of state categories of Memory Evolutive Systems.

If we drop condition 3) in the definition of a sketch-like membrane system, we obtain these directed multigraphical membrane systems which appear useful to describe alternating organization of living systems discussed in [2] with a regard to nesting (represented by the underlying tree \mathbb{T}_{S}) and interaction of levels of organization (represented by family of directed multigraphs G_v ($v \in V(\mathbb{T}_{S})$)). According to [2] the edges in $G_v(v', v)$ describe integration, the edges in $G_v(v, v')$ describe regulation, and the edges of v-diagram Dg(v) describe interaction.

A directed multigraphical (a sketch-like) membrane system is illustrated in Fig. 1, whose semantics (model) in a hierarchical category is illustrated in Fig. 2.



Multigraphical membrane system corresponding to 2-ramification:

nodes—membranes, edges—objects, neurons—membranes, synapses—objects.



the fat arrows are colimiting injections, i.e. the elements of colimiting cocones, respectively

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On Flip-Flop Membrane Systems with Proteins

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Summary. We consider once more the membrane systems with proteins on membranes. This model is bridging the membrane systems and brane calculi areas together, thus it is interesting to study it in more depth. We have improved previous results in the area and also defined a new variant of these systems based on time as the output of the computation. The new model allows (due to its flexibility) even stronger improvements with respect to the number of proteins needed to perform the computation.

1 Introduction

We continue the work on a membrane systems model combining membrane systems and brane calculi as introduced in [14]. In brane calculi introduced in [5], one works only with objects - called proteins - placed on membranes, while the evolution is based on membrane handling operations, such as exocytosis, phagocytosis, etc. In the membrane computing area we have rules associated with each region defined by a membrane, and in the recent years the rules in membrane computing have been considered mainly to work on symbol objects rather than other structures such as strings. The extension considered in [14] and in [15] was to have both types of rules (both at the level of the region delimited by membranes and also at the level of membrane controlled by a protein). The reason for considering both extensions was that in biology, many reactions taking place in the compartments of living cells are controlled/catalysed by the proteins embedded in the membranes bilayer. For instance, it is estimated that in the animal cells, the proteins constitute about 50% of the mass of the membranes, the rest being lipids and small amounts of carbohydrates. There are several types of such proteins embedded in the membrane of the cell; one simple classification places these proteins into two classes, that of integral proteins (these molecules can "work" in both inside the membrane as

well as also in the region outside the membrane), and that of peripheral proteins (macromolecules that can only work in one region of the cell) – see [1].

In the present paper we continue the discussion in the direction of membrane systems with proteins, but we extend the model to have also a "more natural" output of the computation with ideas from [8].

Briefly, the systems that we consider in this paper extend the original definition by using the paradigm of time as the output of a computation as previously introduced in [6] and [8]. The idea originates in [17] as Problem W; the novelty is that instead of the "standard" way to output, like the multiplicities of objects found at the end of the computation in a distinguished membrane as it was defined in the model from [14] and in [15], it seems more "natural" to consider certain *events* (i.e., configurations) that may occur during a computation and to relate the output of such a computation with the time interval between such distinguished configurations. Our system will compute a set of numbers similarly with the case of "normal" symport/antiport systems as defined in [14], but the benefit of the current setting is that the computation and the observance of the output are now close to the biology and to the tools used for cell biology (fluorescence microscopy, FACS).

2 The Types of Rules in the System

In what follows we assume that the reader is familiar with membrane computing basic elements, e.g., from [16] and from [19], as well as with basic elements of computability, so that we only mention here a few notations we use. The rules based on proteins on membranes were described in detail in [14], and we refer the interested reader to that publication and to [15] for further details.

As usual, we represent multisets of objects from a given alphabet V by strings from V^* , and the membrane structures by expressions of correctly matching labeled parentheses. The family of recursively enumerable sets of natural numbers is denoted by NRE.

In the P systems which we consider below, we use two types of objects, proteins and usual *objects*; the former are placed **on** the membranes, the latter are placed **in** the regions delimited by membranes. The fact that a protein p is on a membrane (with label) i is written in the form $[_ip]$. Both the regions of a membrane structure and the membranes can contain multisets of objects and of proteins, respectively.

We consider the following types of rules for handling the objects and the proteins; in all of them, a, b, c, d are objects, p is a protein, and i is a label ("cp" stands for "change protein"), where p, p' are two proteins (possibly equal; if p = p', then the rules of the type cp become rules of the type res; i.e. restricted):

Type	Rule	Effect (besides changing also the protein)
1cp	$[_i p a \rightarrow [_i p' b$	
	$a[_ip] \rightarrow b[_ip']$	modify an object, but not move
2 cp	$[_ip a \to a[_ip' $	
	$a[_ip] \rightarrow [_ip' a$	move one object unmodified
3cp	$[_i p a \rightarrow b [_i p' $	
	$a[_ip] \rightarrow [_ip' b$	modify and move one object
4cp	$a[_ip b \rightarrow b[_ip' a]$	interchange two objects
5cp	$a[_ip b \to c[_ip' d$	interchange and modify two objects

An intermediate case between *res* and *cp* can be that of changing proteins in a restricted manner, by allowing at most two states for each protein, p, \bar{p} , and the rules working either in a *res* manner (without changing the protein), or changing it from p to \bar{p} and back (like in the case of bistable catalysts). Rules with such flip-flop proteins are denoted by nff, n = 1, 2, 3, 4, 5 (note that in this case we allow both rules which do not change the protein and rules which switch from pto \bar{p} and back).

Both in the case of rules of type ff and of type cp we can ask that the proteins are always moved in another state (from p into \bar{p} and vice versa for ff). Such rules are said to be of *pure* ff or cp type, and we indicate the use of pure ff or cp rules by writing ffp and cpp, respectively.

We can use these rules in devices defined in the same way as the symport/antiport P systems (hence with the environment containing objects, in arbitrarily many copies each – we need such a supply of objects, because we cannot create objects in the system), where also the proteins present on each membrane are mentioned.

That is, a P system with proteins on membranes is a device of the form

$$\Pi = (O, P, \mu, w_1/z_1, \dots, w_m/z_m, E, R_1, \dots, R_m, i_o),$$

where:

- 1. m is the degree of the system (the number of membranes);
- 2. *O* is the set of objects;
- 3. *P* is the set of proteins (with $O \cap P = \emptyset$);
- 4. μ is the membrane structure;
- 5. w_1, \ldots, w_m are the (strings representing the) multisets of objects present in the *m* regions of the membrane structure μ ;
- 6. z_1, \ldots, z_m are the multisets of proteins present on the *m* membranes of μ ;
- 7. $E \subseteq O$ is the set of objects present in the environment (in an arbitrarily large number of copies each);
- 8. R_1, \ldots, R_m are finite sets of rules associated with the *m* membranes of μ ;
- 9. i_o is the output membrane, an elementary membrane from μ .

The rules can be of the forms specified above, and they are used in a nondeterministic maximally parallel way: in each step, a maximal multiset of rules is used, that is, no rule can be applied to the objects and the proteins which remain unused by the chosen multiset. As usual, each object and each protein can be involved in the application of only one rule, but the membranes are not considered as involved in the rule applications, hence the same membrane can appear in any number of rules at the same time.

If, at one step, two or more rules can be applied to the same objects and proteins, then only one rule will be non-deterministically chosen. At each step, a P system is characterized by a configuration consisting of all multisets of objects and proteins present in the corresponding membranes (we ignore the structure μ , which will not be changed, and the objects from the environment). For example, $C = w_1/z_1, \ldots, w_m/z_m$ is the initial configuration, given by the definition of the P system. By applying the rules in a non-deterministic maximally parallel manner, we obtain transitions between the configurations of the system. A finite sequence of configurations is called computation. A computation halts if it reaches a configuration where no rule can be applied to the existing objects and proteins.

Only halting computations are considered successful, thus a non-halting computation will yield no result. With a halting computation we associate a result, in the form of the multiplicity of objects present in region i_o in the halting configuration. We denote by $N(\Pi)$ the set of numbers computed in this way by a given system Π . (A generalization would be to distinguish the objects and to consider vectors of natural numbers as the result of a computation, but we do not examine this case here.)

We denote, in the usual way, by $NOP_m(pro_r; list-of-types-of-rules)$ the family of sets of numbers $N(\Pi)$ generated by systems Π with at most m membranes, using rules as specified in the list-of-types-of-rules, and with at most r proteins present on a membrane. When parameters m or r are not bounded, we use * as a subscript.

The new definition introduced by the current paper is the addition of time to the above model, in brief, P system with proteins on membranes and time is a device of the form

$$\Pi = (O, P, \mu, w_1/z_1, \dots, w_m/z_m, E, R_1, \dots, R_m, C_{start}, C_{stop}),$$

where:

- 1. $m, O, P, \mu, w_1, \ldots, w_m, z_1, \ldots, z_m, E, R_1, \ldots, R_m$ are as defined above;
- 2. C_{start} , C_{stop} are regular subsets of $(O^*)^m$, describing configurations of Π . We will use a regular language over $O \cup \{\$\}$ to describe them, the special symbol $\$ \notin O$ being used as a marker between the configurations³ in the different regions of the system. More details are given in [8] and [12].

³ We express by these configurations restrictions that need to be satisfied by each of the current multisets in their respective regions so that the overall configuration can be satisfied.

As an example for the C_{start} and C_{stop} configurations, let us give the following restriction⁴ $C = b^3 d^7 (O - \{a, b, d\})^*$ for a single membrane (the proofs obtained below need only one membrane, thus we can simplify the notation by not using the symbol \$). This means that in the region delimited by the only membrane in the system, the configuration C is satisfied if and only if we do not have any symbol of type a, we must have exactly 3 symbols of type b and exactly 7 symbols of type d. Any other symbol not mentioned is not restricted, e.g. we can have any number of symbols of type c.

We emphasize the fact that in the definition of Π we assume that C_{start} and C_{stop} are regular. Other, more restrictive, cases can be of interest but we do not discuss them here.

We can now denote the systems as above based on time with $NTOP_m(pro_r; list-of-types-of-rules)$ the family of sets of numbers $N(\Pi)$ generated by systems Π with at most m membranes, using rules as specified in the list-of-types-of-rules, and with at most r proteins present on a membrane. When parameters are not bounded we replace them by *.

3 Register Machines

In the proofs from the next sections we will use register machines as devices characterizing NRE, hence the Turing computability.

Informally speaking, a register machine consists of a specified number of registers (counters) which can hold any natural number, and which are handled according to a program consisting of labeled instructions; the registers can be increased or decreased by 1 - the decreasing being possible only if a register holds a number greater than or equal to 1 (we say that it is non-empty) –, and checked whether they are non-empty.

Formally, a (non-deterministic) register machine is a device $M = (m, B, l_0, l_h, R)$, where $m \ge 1$ is the number of counters, B is the (finite) set of instruction labels, l_0 is the initial label, l_h is the halting label, and R is the finite set of instructions labeled (hence uniquely identified) by elements from B (R is also called the *program* of the machine). The labeled instructions are of the following forms:

- $l_1: (ADD(r), l_2, l_3), 1 \le r \le m$ (add 1 to register r and go non-deterministically to one of the instructions with labels l_2, l_3),
- l_1 : (SUB $(r), l_2, l_3$), $1 \le r \le m$ (if register r is not empty, then subtract 1 from it and go to the instruction with label l_2 , otherwise go to the instruction with label l_3),
- l_h : HALT (the halt instruction, which can only have the label l_h).

We say that a register machine has no ADD instructions looping to the same label (or *without direct loops*) if there are no instructions of the form

⁴ C can be written also in the following form $C = (a^0 b^3 d^7)$

 l_1 : (ADD $(r), l_1, l_2$) or l_1 : (ADD $(r), l_2, l_1$) in R. For instance, an instruction of the form l_1 : (ADD $(r), l_1, l_2$) can be replaced by the following instructions, where l'_1 is a new label: l_1 : (ADD $(r), l'_1, l_2$), l'_1 : (ADD $(r), l_1, l_2$). The generated set of numbers is not changed.

A register machine generates a natural number in the following manner: we start computing with all m registers being empty, with the instruction labeled by l_0 ; if the computation reaches the instruction l_h : HALT (we say that it halts), then the values of register 1 is the number generated by the computation. The set of numbers computed by M in this way is denoted by N(M). It is known (see [11]) that non-deterministic register machines with three registers generate exactly the family NRE, of Turing computable sets of numbers. Moreover, without loss of generality, we may assume that in the halting configuration all registers except the first one, where the result of the computation is stored are empty.

4 Previous Results

In [14] the following results were proved:

Theorem 1.

$$\begin{aligned} NOP_1(pro_2; 2cpp) &= NRE. \quad (Theorem \ 5.1 \ in \ [14]) \\ NOP_1(pro_*; 3ffp) &= NRE. \quad (Theorem \ 5.2 \ in \ [14]) \\ NOP_1(pro_2; 2res, 4cpp) &= NRE. \quad (Theorem \ 6.1 \ in \ [14]) \\ NOP_1(pro_2; 2res, 1cpp) &= NRE. \quad (Theorem \ 6.2 \ in \ [14]) \\ NOP_1(pro_*; 1res, 2ffp) &= NRE. \quad (Theorem \ 6.3 \ in \ [14]) \end{aligned}$$

As an extension of the work reported in [14], a significant amount of energy was devoted to the flip-flopping variant of these membrane systems which resulted in the paper [9]. S.N Krishna was able to prove several results in [9] improving Theorem 5.2, and Theorem 6.3 from [14]:

Theorem 2.

 $NOP_1(pro_7; 3ffp) = NRE.$ (Theorem 1 in [9]) $NOP_1(pro_7; 2ffp, 4ffp) = NRE.$ (Theorem 2 in [9]) $NOP_1(pro_7; 2ffp, 5ffp) = NRE.$ (Corollary 3 in [9]) $NOP_1(pro_1; 1res, 2ffp) = NRE.$ (Theorem 4 in [9]) $NOP_1(pro_7; 1ffp, 2ffp) = NRE.$ (Theorem 6 in [9]) $NOP_1(pro_9; 1ffp, 2res) = NRE.$ (Theorem 7 in [9]) $NOP_1(pro_9; 2ffp, 3res) = NRE.$ (Theorem 9 in [9]) $NOP_1(pro_9; 1ffp, 3res) = NRE.$ (Theorem 10 in [9]) $NOP_1(pro_9; 3res, 4ffp) = NRE.$ (Theorem 11 in [9])

$$NOP_1(pro_8; 2ffp, 5res) = NRE.$$
 (Theorem 13 in [9])

A close reading of the theorems mentioned above will yield some improvements that are given in the following section.

5 New Results

We start this section by first discussing the results from [9] which we mentioned in Theorem 2. The main idea in all the proofs reported in [9] was to simulate register machines (it is known that such devices with 3 registers are universal). The novelty of the proof technique in [9] was to consider for all ADD instructions associated with a particular register a single protein, similarly we use one protein for all the SUB instructions associated with a specific register. Thus in the proofs of the results mentioned in Theorem 2 we will have 6 proteins used for the simulation of the instructions in the register machine, (both ADD and SUB instructions for the 3 registers in the machine) the other(s) protein(s) being needed mainly for the test with zero processing in the simulation of SUB instructions.

The main observation that we want to make at this point is the fact that register machines with three registers out of which one (the output register) is non-decreasing are still universal, thus all the results from [9] are better by one protein without any major changes in their proofs. This is due to the fact that we only need two proteins to simulate the SUB instructions, and also the proof technique allows for such a modification. Subsequently, the following results were shown in [9]:

Theorem 3.

$$NOP_1(pro_6; 3ffp) = NRE.$$
 (Theorem 1 in [9])
 $NOP_1(pro_6; 2ffp, 4ffp) = NRE.$ (Theorem 2 in [9])
 $NOP_1(pro_6; 2ffp, 5ffp) = NRE.$ (Corollary 3 in [9])
 $NOP_1(pro_9; 1res, 2ffp) = NRE.$ (Theorem 4 in [9])
 $NOP_1(pro_6; 1ffp, 2ffp) = NRE.$ (Theorem 6 in [9])
 $NOP_1(pro_8; 1ffp, 2res) = NRE.$ (Theorem 7 in [9])
 $NOP_1(pro_8; 2ffp, 3res) = NRE.$ (Theorem 9 in [9])
 $NOP_1(pro_7; 1ffp, 3res) = NRE.$ (Theorem 10 in [9])
 $NOP_1(pro_7; 2ffp, 5res) = NRE.$ (Theorem 11 in [9])
 $NOP_1(pro_7; 2ffp, 5res) = NRE.$ (Theorem 13 in [9])

We will proceed now to consider the same framework, but with the extra feature of the output based on time. We show that we can improve the result from Theorem 11 from [9]:

Theorem 4. $NRE = NTOP_1(pro_7, 3res, 4ffp)$.

Proof. We consider a register machine $M = (m, B, l_0, l_h, R)$ and we construct the system

$$\Pi = (O, P, [1]_1, \{l_0, b\}/P, E, R_1, C_{start}, C_{stop})$$

with the following components:

 C_s

$$O = \{a_r, a'_r \mid 1 \le r \le 3\} \cup \{i, i', l_i, l'_i, l''_i, l'''_i, l_i^{iv}, L_i, L'_i \mid 0 \le i \le h\}$$

$$\cup \{o, o_1, o_2, b, h, \dagger\}.$$

$$E = \{a_r, a'_r \mid 1 \le r \le 3\} \cup \{i \mid 0 \le i \le h\} \cup \{o\}.$$

$$P = \{p_1, p_2, p_3, s_2, s_3, p, t\}.$$

$$t_{art} = l''_h (O - \{l''_h, \dagger\})^*, \text{ in other words, } l''_h$$

$$= p_{art} = a_{ar} (A_r - \{l''_h, \dagger\})^*, \text{ in other words, } l''_h$$

appears exactly once and there are no copies of \dagger in the membrane, and the rest of the symbols can

appear in any multiplicity as they are ignored.

 $C_{stop} = (O - \{a_1\})^*$, in this case a_1 does not appear in the membrane.

The proteins p and t are of the type 3res while all the others are of the type 4ffp. Proteins p and p_i are used in the simulation of ADD instructions of register i, proteins p, t and s_i are used in the simulation of SUB instructions of register i, and protein p, t, s_2 and s_3 are used in the simulation of the instructions for counting or termination.

The system has the following rules in R_1 :

For an **ADD** instruction l_1 : $(ADD(r), l_2, l_3) \in R$, we consider the rules as shown in Table 1.

Step	Rules	Type	Environment	Membrane
1	$a_r'[{}_1p_r \mid l_1 \to l_1[{}_1p_r' \mid a_r'$	4ffp	El_1	ba'_r
2 or	$a_r[_1p'_r \mid a'_r \rightarrow a'_r[_1p_r \mid a_r \text{ and } l_1[_1p \mid \rightarrow [_1p \mid l_2])$	4ffp, 3res	E	bl_2a_r
2	$ a_r[_1p'_r \mid a'_r \to a'_r[_1p_r \mid a_r \text{ and } l_1[_1p \mid \to [_1p \mid l_3] $	4ffp, 3res	E	bl_3a_r

 Table 1. Steps for ADD instruction for Theorem 4.

We simulate the work of the ADD instruction in two steps. First we send out the current instruction label l_1 and bring in a copy of the (padding) symbol a'_r using the protein p_r . Next we simultaneously apply the rules to replace a'_r with a_r using the protein p'_r and we bring in the next instruction label l_2 or l_3 according to the currently simulated rule l_1 . Of course, l_1 uniquely identifies which rule was simulated, thus there is no ambiguity about which symbols l_i are able to enter the membrane at this time. Let us now consider the case of the SUB instructions:

For a **SUB instruction** l_1 : (SUB $(r), l_2, l_3$) $\in R$ we consider the rules as shown in Table 2.

Step	Rules	Type	Environment	Membrane
1	$\begin{bmatrix} p & l_1 \rightarrow l_1' \end{bmatrix} p$	3res	El'_1	ba_r
2	$ l_1'[_1p \to [_1p \mid l_1'']$	3res	E	$bl_1''a_r$
3	$o[_1s_r \mid l_1'' \xrightarrow{\rightarrow} l_1''[_1s_r' \mid o$	4ffp	El''_1	boa_r
4	$l_1''[_1p \mid \rightarrow [_1p \mid l_1''' \text{ and } [_1t \mid o \rightarrow o_1[_1t]$	3res, 3res	Eo_1	$bl_1^{\prime\prime\prime}a_r$
5	$\begin{bmatrix} 1 \\ 1 \end{bmatrix} p \mid l_1^{\prime\prime\prime} \rightarrow l_1^{iv} \begin{bmatrix} 1 \\ 1 \end{bmatrix} p \mid \text{and } o_1 \begin{bmatrix} 1 \\ 1 \end{bmatrix} t \mid \rightarrow \begin{bmatrix} 1 \\ 1 \end{bmatrix} t \mid o_2$	3res, 3res	El_1^{iv}	bo_2a_r
	Register r is non-empty			
6	$l_1^{iv}[_1s'_r \mid a_r \to a_r[_1s_r \mid l_1^{iv} \text{ and } [_1t \mid o_2 \to o[_1t \mid $	4ffp, 3res	Eoa_r	bl_1^{iv}
7	$\left \begin{bmatrix} p & i \\ 1 & p \end{bmatrix} l_1^{iv} \to 2' \begin{bmatrix} p \\ 1 & p \end{bmatrix} \right $	3res	E2'	b
8	$2^{\overline{\prime}}[_{1}p \mid \rightarrow [_{1}p \mid \overline{l}_{2}$	3res	E	bl_2
	Wrong Computation			
6	$l_1^{iv}[_1t \mid \rightarrow [_1t \mid L'_3 \text{ and } [_1p \mid o_2 \rightarrow \dagger [_1p \mid $	3res, 3res	E^{\dagger}	$bL'3a_r$
7	$\dagger [_{1}t \mid \rightarrow [_{1}t \mid \dagger$	3res	E	$b \dagger a_r$
	Register r is empty			
6	$\begin{bmatrix} 1 \\ t \end{bmatrix} o_2 \to o \begin{bmatrix} 1 \\ t \end{bmatrix}$	3res	Eo	b
7	$l_1^{\bar{i}v}[_1t \mapsto [_1t] L_3'$	3res	E	bL'_3
8	$\left 3\left[{}_{1}s_{r}^{\prime} \mid L_{3}^{\prime} \rightarrow L_{3}^{\prime}\right] {}_{1}s_{r} \mid 3 \right.$	4ffp	EL'_3	b3
9	$\left \left[\begin{smallmatrix} 1\\ p \end{smallmatrix}\right 3 \to 3'\left[\begin{smallmatrix} 1\\ p \end{smallmatrix}\right]\right $	3res	E3'	b
10	$3'[_1p \mid]ra[_1p \mid] l_3$	3res	E	bl_3

 Table 2. Steps for SUB instruction for Theorem 4.

We simulate the work of the SUB instruction in several steps (eight if the register is not empty and ten if it is empty). We first send out the current label as l'_1 using the protein p. At the next step the symbol l'_1 is brought in as l''_1 . Next we exchange l''_1 and o using the protein s_r (the protein s_r is moved in its primed version of the flip-flop). We can now apply two rules in parallel and bring in l''_1 as l'''_1 while sending out o as o_1 . Next, l'''_1 is sent out as l^{iv}_1 while we bring in o_1 as o_2 in parallel.

In this moment our system will perform the checking of the contents of the register r. If the register is not empty, then l_1^{iv} will enter the membrane, decreasing the register and at the same time another marker o_2 is sent outside as o to help identify the correct case later. At the next stage l_1^{iv} will be sent out as 2' using protein p. Finally 2' will return as the next instruction label to be brought in (in this case l_2 as the register is not empty). If l_1^{iv} comes back in the membrane through the protein t instead of s'_r , we will have a wrong computation. In this case we can send out o_2 as symbol \dagger in parallel using the protein p (as this is the only channel available at this time to o_2 , t being used by l_1^{iv}). Next we can bring in a copy of the symbol \dagger into the membrane. The application of this rule will never satisfy the starting configuration; hence, we will not be able to use the time counter.

If the register is empty, after step 5 we have l_1^{iv} in the environment and o_2 in the membrane, and the protein associated with the subtract rule for the register $r(s_r)$ is primed. At this moment l_1^{iv} cannot enter the membrane through the protein s'_r as there are no a_r objects in the membranes with which it must be exchanged.

There are two choices: either l_1^{iv} enters the membrane through t (and we get the wrong computation case as above) or t is used by o_2 , and then l_1^{iv} sits one step in the environment. At the next step we have the "branching point": rather than exchanging with a_r (which will be present in the membrane in the case when the register is not empty), l_1^{iv} comes into the membrane as L'_3 through t. Next we use the protein s'_r to exchange L'_3 and 3, and then send out 3 as 3' using protein p. Now we bring in 3' as the next instruction to be simulated l_3 .

Terminating/counting work: It is clear that at the end of the simulation, if the register machine has reached the final state, we will have the halting instruction symbol in the membrane along with one copy of the symbol b and multiple copies of the three different objects associated with their respective registers. At that time we will have the computed value encoded as the multiplicity of the object a_1 that is associated with the output register. We will also have in the system the label of the halting instruction, l_h ; thus, the rule $([_1p | l_h \rightarrow l'_h[_1p |)$ can be applied only when the simulation is performed correctly. At the next step, using the protein s_2 we exchange l'_h and b.

The terminating/counting work is done by the rules as shown in Table 3.

Step	Rules	Type	Env	Membrane
1	$\left[{}_{1}p \mid l_{h} \rightarrow l_{h}^{\prime} \right]_{1}p \mid$	3res	El'_h	$ba_1^{n_1}a_2^{n_2}a_3^{n_3}$
2	$l_h^{}[_1s_2 \mid b ightarrow b[_1s_2' \mid l_h']$	4ffp	Eb	$l'_{h}a_1^{n_1}a_2^{n_2}a_3^{n_3}$
3	$b[_1p] \mapsto [_1p \mid l_h^{\bar{n}} \text{ and } [_1t \mid l_h' \to l_h^{\prime\prime}[_1t \mid]$	3res, 3res	El''_h	$l_h'' a_1^{n_1} a_2^{n_2} a_3^{n_3}$
4	$h[_{1}s_{2} \mid l_{h}'' \rightarrow l_{h}''[_{1}s_{2}' \mid h \text{ or } h[_{1}s_{2}' \mid l_{h}'' \rightarrow l_{h}''[_{1}s_{2} \mid h]$	4ffp	El''_ha_1	$ l_h'' a_1^{n_1} a_2^{n_2} a_3^{n_3} h $
	and $l_h''[_1s_3 \mid a_1 \rightarrow a_1[_1s_3' \mid l_h'']$	4ffp		
	$\mathbf{or} \ l_h''[_1 s_3^{\vec{i}} \mid a_1 \to a_1[_1 s_3^{\vec{i}} \mid l_h'']$			

Table 3. Steps for terminating/counting instructions for Theorem 4.

Next we apply two rules in parallel and bring in b as l''_h while sending out l'_h as l''_h , satisfying the C_{start} configuration. One can note that if there are no copies of a_1 in the membrane, then also the configuration C_{stop} is satisfied at the same time, thus our system would compute the value zero in that case. Next we exchange h from the environment with l''_h and l''_h from the environment with a_1 until we reach the stopping configuration. For any other value encoded in the multiplicity of a_1 it will take exactly the same number of steps to push the number of copies of object a_1 from the membrane. \Box

An interesting observation is the fact that the object b is used for the counting at the end of the computation. If one considers the same construct for membrane systems with proteins as defined in [14] (the "classical" systems with the output the multiplicity of objects in the membrane), then our construction is still valid even in the case of systems without time, thus we have the following theorem also proven:

Theorem 5. $NRE = NOP_1(pro_7, 3res, 4ffp)$.

The theorem above is valid as one can restrict the register machine to be simulated (without loss of generality) to the case when the machine halts with the non-output registers empty.

Thus it can be seen that we are able to improve the result shown in Theorem 10 in [9] both for systems based on multiplicity output and also for systems based on time. The next result improves significantly Theorem 11 from [9], in this case for systems based on time, and later one we will discuss also about the non-timed systems.

Theorem 6. $NRE = NTOP_1(pro_3, 2ffp, 5res).$

Proof. We consider a register machine $M = (m, B, l_0, l_h, R)$ and we construct the system

$$\Pi = (O, P, [1]_{1}, \{l_{0}, b, e\}/P, E, R_{1}, C_{start}, C_{stop})$$

with the following components:

$$O = \{l_i, l'_i \mid 0 \le i \le h\} \cup \{a_1, a_2, a_3, b, o, y\}.$$

$$E = \{a_1, a_2, a_3, o\}.$$

$$P = \{p, q, s\}.$$

 $C_{start} = (O - \{b\})^*$, in other words, there are no copies of b in the membrane, and the rest of the symbols

can appear in any multiplicity as they are ignored.

 $C_{stop} = (O - \{a_1\})^*$, in this case a_1 does not appear in the membrane.

Protein q is of type 5res while all the others are of the type 2ffp. Proteins p and q are used in the simulation of the ADD instruction, proteins q and s are used in the simulation of the SUB instruction, and protein q is used in the simulation of the instructions for counting or termination.

The system has the following rules in R_1 :

For an **ADD** instruction l_1 : $(ADD(r), l_2, l_3) \in R$, we consider the rules as shown in Table 4.

Step	Rules	Type	Environment	Membrane
1	$a_r[_1q \mid l_1 \to l_1'[_1q \mid a_r]$	5res	El'_1	bea_r
2	$l_1'[\ _1^{ \prime} q \mid e \to e[\ _1^{ \prime} q \mid l_2$	5res	Ee	bl_2a_r
2	$l_1' []_1 q \mid e \to e []_1 q \mid l_3$	5res	Ee	bl_3a_r
3	$e[_1p] \mapsto [_1p'] e \text{ or } e[_1p'] \mapsto [_1p] e$	2ffp, 2ffp	E	bea_r

 Table 4. Steps for ADD instruction for Theorem 6.

We simulate the work of the ADD instruction in two steps. First we send out the current instruction label l_1 as l'_1 and bring in a copy of the symbol a_r using the protein q. Next we apply the rule to send out e using the protein q and we bring l'_1 in as the new instruction label. To simulate the non-deterministic behavior of these machines we have two rules that do the same job, the only difference being the next instruction label being brought back in the system. It is clear that the simulation of the ADD instruction is performed correctly. The work is finished in this case by the rule $(e_1p \mid \rightarrow [_1p' \mid e)$ or $(e_1p' \mid \rightarrow [_1p \mid e)$.

For a **SUB instruction** $l_1 : (SUB(r), l_2, l_3) \in R$ we consider the rules as shown in Table 5.

Step	Rules	Type	Environment	Membrane
1	$\left[{_1}s \mid l_1 \to l_1 [_1s' \mid \right]$	2ffp	El_1	bea_r
	Register r is non-empty			
2	$o[_1s' \mapsto [_1s \mid o \text{ and } l_1[_1q \mid a_r \to a_r[_1q \mid l'_1]$	2ffp, 5res	Ea_r	$beol'_1$
3	$o[_1q \mid l_1' \to l_1'[_1q \mid l_2$	5res	El'_1	$beol_2$
4	$l_1' [_q \mid o \to o [_q \mid y$	5res	Eo	bey
	Register r is empty			
2	$o[_1s' \mapsto [_1s \mid o$	2ffp	El_1	beo
3	$l_1[_1q \mid o \to o[_1q \mid l_3$	5res	Eo	bel_3

Table 5. Steps for SUB instruction for Theorem 6.

We simulate the work of the SUB instruction in several steps (four if the register is not empty and three if it is empty). At step 1 we first send out the current label l_1 using the protein s. If the register is not empty, at step 2, l_1 will enter the membrane, decreasing the register and at the same time the symbol o is brought in. At the next stage (step 3) l'_1 will be sent out using protein q, and o will return as the next instruction label to be brought in (in this case l_2 as the register is not empty). Finally l'_1 will return as the symbol y while sending out o, so that no extra copies of o are left in the membrane so that future SUB simulations will be performed correctly. The symbols y will accumulate in the membrane.

In the case when the register to be decremented is empty, we perform the same initial step, sending out the current label using the protein s. This time l_1 cannot enter the membrane at the step 2 as there is no a_r in the membrane to help bring it in. So l_1 will wait for one step in the environment. o is entering the membrane at step 2, so at the step 3 l_1 can now come into the membrane through q and is changed into the label of the next instruction to be simulated l_3 .

The **terminating/counting work** stage is done by the rules as shown in Table 6.

Step	Rules	Type	Environment	Membrane
1	$o[_1q \mid l_h \to l_h[_1q \mid y]$	5res	El_h	$bea_1^{n_1}a_2^{n_2}a_3^{n_3}$
2	$l_h[_1q \mid b \to l'_h[_1q \mid y]$	5res	Ebl'_h	$ea_1^{n_1}a_2^{n_2}a_3^{n_3}$
3	$\left l_h' \right _1^{\uparrow} q \mid a_1 \to l_h' \left _1^{\uparrow} q \mid y \right $	5res	El'_ha_1	$ea_1^{n_1}a_2^{n_2}a_3^{n_3}$

Table 6. Steps for terminating/counting instructions for Theorem 6.

It is clear that at the end of the simulation, if the register machine has reached the final state, we will have the halting instruction symbol in the system membrane, along with one copy of the symbol b and multiple copies of the three different objects associated with the respective registers and the symbol y. At that time we will have the computed value encoded as the multiplicity of the object a_1 that is associated with the output register. We will also have in the system the label of the halting instruction, l_h , thus the rule $(o[_1q \mid l_h \rightarrow l_h[_1q \mid y))$ can be applied only when the simulation is performed correctly. At the next step, using the protein q we bring in l_h as y while sending out b as l'_h , satisfying the C_{start} configuration. One can note that if there are no copies of a_1 in the membrane, then also the configuration C_{stop} is satisfied at the same time, thus our system would compute the value zero in that case. Next we bring in l'_h as y while sending out a_1 as l'_h until we reach the stopping configuration. For any other value encoded in the multiplicity of a_1 it will take exactly the same number of steps to push the a_1 -s out of the membrane. \Box

Thus it can be seen that by using time as the output, we are able to improve the result shown in Theorem 13 from [9], where seven proteins were required for universality, as opposed to the three used in the above proof.

If one wants to still restrict the discussion to only the case of the non-timed systems, with the price of one protein we can remove the objects y and e from the membrane (by first modifying them into some other symbols such as y' and o' and then expelling them to the environment). In this way it is easy to see that our proof for Theorem 6 leads to the following theorem:

Theorem 7. $NRE = NOP_1(pro_4, 2ffp, 5res).$

Membrane systems with proteins on membranes are universal for one membrane and rules of the type 2ffp and 5res using only four proteins.

6 Final Remarks

We have shown that previous results about membrane systems with proteins on membranes can be improved in what concerns the number of proteins, we have also extended the model to have the output encoded as the time between two configurations and this has lead to a significant improvement as opposed to the previous results reported in [9]. Additional similar improvements are under investigation.

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Rewriting P Systems with Conditional Communication: Improved Hierarchies

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Summary. We consider here a variant of rewriting P systems [1], where communication is controlled by the contents of the strings, not by the evolution rules used for obtaining these strings. Some new characterizations of recursively enumerable languages are obtained by means of P systems with a small number of membranes, which improves some of the known results from [1] and [4].

1 Introduction

P systems are a class of distributed parallel computing models inspired from the way the living cells process chemical compounds, energy, and information. Many variants of P systems use string objects and context-free rules for processing them. *Rewriting P systems* with string objects were introduced in [5]. Several variants of P systems with string objects have also been investigated extensively. In this work, we concentrate on rewriting P systems with conditional communication introduced in [1].

In this variant of rewriting P systems, the communication is controlled by the contents of the strings, not by the evolution rules themselves. This is achieved by considering certain types of *permitting* and *forbidding* conditions, based on the symbols or the substrings (arbitrary, or prefixes/suffixes) which appear in a given string. Several characterizations of recursively enumerable languages were obtained in [1]. In [4], some of these results were improved. Here we give some new characterizations of recursively enumerable languages by means of P systems with a small number of membranes. These results improve some of the results from both [1] and [4]. In [1], there is a characterization of recursively enumerable language by systems, where both prefixes and suffixes are checked, without a bound on the number of membranes. It was also conjectured that the characterization holds also for a reduced number of membranes. We settle this here in an affirmative way by giving the characterization with 8 membranes.

2 Some Prerequisites

In this section we introduce some formal language theory notions which will be used in this paper; for further details, we refer to [7].

For an alphabet V, we denote by V^* the set of all strings over V, including the empty one, denoted by λ . By RE we denote the family of recursively enumerable languages. The set of symbols appearing in a string x is denoted by alph(x) and the substrings of x is denoted by Sub(x).

In our proofs in the following sections we need the notion of a matrix grammar with appearance checking. Such a grammar is a construct G = (N, T, S, M, F), where N, T are disjoint alphabets, $S \in N$, M is a finite set of sequences of the form $(A_1 \rightarrow x_1, \ldots, A_n \rightarrow x_n), n \ge 1$, of context-free rules over $N \cup T$, and F is a set of occurrences of rules in M (N is the nonterminal alphabet, T is the terminal alphabet, S is the axiom, while the elements of M are called matrices).

For $w, z \in (N \cup T)^*$ we write $w \Rightarrow z$ if there is a matrix $(A_1 \to x_1, \ldots, A_n \to x_n)$ in M and the strings $w_i \in (N \cup T)^*, 1 \le i \le n+1$, are such that $w = w_1, z = w_{n+1}$, and, for all $1 \le i \le n$, either (1) $w_i = w'_i A w''_i, w_{i+1} = w'_i x_i w''_i$, for some $w'_i, w''_i \in (N \cup T)^*$, or (2) $w_i = w_{i+1}, A_i$ does not appear in w_i , and the rule $A_i \to x_i$ appears in F. (The rules of a matrix are applied in order, possibly skipping the rules in Fif they cannot be applied - one says that these rules are applied in the *appearance checking mode*).

The languages generated by G is defined by $L(G) = \{w \in T^* \mid S \Rightarrow^* w\}$. The family of languages of this form is denoted by MAT_{ac} . It is known that $MAT_{ac} = RE$

A matrix grammar G = (N, T, S, M, F) is said to be in the *binary normal form* if $N = N_1 \cup N_2 \cup \{S, \#\}$, with these three sets mutually disjoint, and the matrices in M are in the following forms:

- 1. $(S \to XA)$, with $X \in N_1, A \in N_2$,
- 2. $(X \to Y, A \to x)$, with $X, Y \in N_1, A \in N_2, x \in (N_2 \cup T)^*, |x| \le 2$,
- 3. $(X \to Y, A \to \#)$, with $X, Y \in N_1, A \in N_2$,
- 4. $(X \to \lambda, A \to x)$, with $X \in N_1, A \in N_2$, and $x \in T^*, |x| \leq 2$.

Moreover, there is only one matrix of type 1 and F consists exactly of all rules $A \rightarrow \#$ appearing in matrices of type 3; # is a trap symbol - once introduced, it is never removed. A matrix of type 4 is used only once, in the last step of derivation.

According to [2], for each matrix grammar there is an equivalent matrix grammar in the binary normal form.

For an arbitrary matrix grammar G = (N, T, S, M, F), let us denote by ac(G) the cardinality of the set $\{A \in N \mid A \to \alpha \in F\}$. It was proved that each recursively enumerable language can be generated by a matrix grammar G such that $ac(G) \leq 2$. Consequently, to the properties of a grammar G in the binary normal form we can add the fact that $ac(G) \leq 2$. We will say that this is the *strong binary normal form* for matrix grammars.

There are several normal forms for type 0 grammars. We use the Penttonen normal form in our proofs. A type 0 grammar G = (N, T, S, P) is said to be

in *Penttonen normal form* if the rules from P are of one of the following forms: $A \to \lambda$, $A \to a$, $A \to BC$, $AB \to AC$, for $A, B, C \in N$ and $a \in T$.

3 Rewriting P Systems with Conditional Communication

An extended rewriting P systems (of degree $m \ge 1$) with conditional communication is a construct

$$\Pi = (V, T, \mu, M_1, \dots, M_m, R_1, P_1, F_1, \dots, R_m, P_m, F_m),$$

where:

- 1. V is the alphabet;
- 2. $T \subseteq V$ is the terminal alphabet;
- 3. μ is the membrane structure;
- 4. M_1, \ldots, M_m are finite languages over V, representing the strings initially present in the m regions;
- 5. R_1, \ldots, R_m are finite sets of context-free rules over V present in the regions of μ ;
- 6. P_i and F_i are permitting and forbidding conditions associated with the regions.

The conditions can be of the following forms:

- 1. *empty*: no restriction is imposed on strings, they either exit the current membrane or enter any of the directly inner membrane freely (but they cannot remain in the current membrane); we denote an empty permitting condition by $(true, X), X \in \{in, out\}$, and an empty forbidding condition by $(false, notX), X \in \{in, out\}$.
- 2. symbols checking: each P_i is a set of pairs $(a, X), X \in \{in, out\}$, for $a \in V$, and each F_i is a set of pairs $(b, notX), X \in \{in, out\}$, for $b \in V$; a string w can go to a lower membrane only if there is a pair $(a, in) \in P_i$ with $a \in alph(w)$ and for each $(b, notin) \in F_i$ we have $b \notin alph(w)$; similarly for sending the string w out of membrane i it is necessary to have $a \in alph(w)$ for at lease one pair $(a, out) \in P_i$ and $b \notin alph(w)$ for all $(b, notout) \in F_i$.
- 3. substring checking: each P_i is a set of pairs (u, X), $X \in \{in, out\}$, for $u \in V^+$, and each F_i is a set of pairs (v, notX), $X \in \{in, out\}$, for $v \in V^+$; a string w can go to a lower membrane only if there is a pair $(u, in) \in P_i$ with $u \in Sub(w)$, and for each $(v, notin) \in F_i$ we have $v \notin Sub(w)$; similarly for sending the string w out of membrane i it is necessary to have $u \in Sub(w)$ for at lease one pair $(u, out) \in P_i$ and $v \notin Sub(w)$ for all $(v, notout) \in F_i$.
- 4. *prefix/suffix checking:* exactly as in the case of substrings checking, with the checked string being a prefix or a suffix of the string to be communicated.

We say that we have conditions of the types *empty*, *symb*, *sub_k*, *pref_k*, *suff_k*, respectively, where k is the length of the longest string in all P_i , F_i .

A systems is said to be *non-extended* if V = T.
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The transitions of the system are defined in the following way. In each region, each string which can be rewritten is rewritten by a rule from that region. The rule to be applied and the nonterminal it rewrites are non-deterministically chosen. The string obtained in this way is checked against the conditions P_i , F_i from that region. If it fulfills the required conditions, then it will be immediately sent out of the membrane or to an inner membrane, if any exists; if it fulfills both *in* and *out* conditions, then it is sent to a membrane non-deterministically choosing the direction - and non-deterministically choosing the inner membrane in the case when several directly inner membranes exist. If a string does not fulfill any condition, or it fulfills only *in* conditions and there is no inner membrane, then the string remains in the same region. If a string cannot be rewritten, then it is directly checked against the communication conditions. That is, the rewriting has priority over communication.

A sequence of transitions form a computation and the result of a halting computation is the set of strings over T sent out of the system. In the case of nonextended systems, all strings sent out are accepted. A computation does not yield a result if it does not halt. A string which remains inside the system or, in the case of extended systems, which exits but contains nonterminal symbols does not contribute to the generated language. The language generated by a system Π is denoted by $L(\Pi)$.

We denote by $RP_n(rw, \alpha, \beta)$, $n \geq 1$, $\alpha, \beta \in \{empty, symb\} \cup \{sub_k \mid k \geq 2\} \cup \{prefsuff_k \mid k \geq 2\}$, the family of languages generated by P system of degree at most n and with permitting and forbidding conditions of type α and β respectively.

4 Improved Universality Results

In [1], it was proved that P systems of degree 4 with permitting conditions of type sub_2 and forbidding conditions of type symb are computationally universal. This result has been improved from 4 to 3 membranes in [4]. We improve this result and show that universality can be achieved with 2 membranes in this case.

Theorem 1. $RE = RP_2(rw, sub_2, symb)$.

Proof. Let us consider a type 0 grammar G = (N, T, S, P), in Penttonen normal form, with the non context-free rules from P labeled in a one-to-one manner and construct the system

$$\Pi = (V, T, [1[2]_2]_1, \{S\}, \emptyset, (R_1, P_1, F_1), (R_2, P_2, F_2)),$$

with the following components :

$$V = N \cup T \cup \{(B, r) \mid r : AB \to AC \in P\};$$

$$R_1 = \{A \to x \mid A \to x \in P\}$$

$$\cup \{B \to (B,r) \mid r : AB \to AC \in P\};$$

$$P_1 = \{(A(B,r),in) \mid r : AB \to AC \in P\}$$

$$\cup \{(true, out)\};$$

$$F_1 = \{(false, notout)\};$$

$$R_2 = \{(B,r) \to C \mid r : AB \to AC \in P\};$$

$$P_2 = \{(true, out)\};$$

$$F_2 = \{((B,r), notout) \mid r : AB \to AC \in P\}.$$

The system works as follows:

The initial configuration of the system is $[{}_{1}S[_{2}]_{2}]_{1}$. The context-free rules from P are present in R_{1} as rewriting rules, hence we can simulate them without any difficulty. Let us assume that we have a string $w_{1}ABw_{2}$ in membrane 1. In order to simulate a rule $r: AB \to AC \in P$, we apply the rule $B \to (B, r)$ on the string. The string is sent to membrane 2 only if it has a substring of the form A(B,r) for some $r: AB \to AC \in P$. Otherwise, the string is sent out, but it is not a terminal one. In membrane 2, we replace the symbol (B, r) with C and send the resulting string to the skin membrane. In this way, we complete the simulation of the non context-free rule.

The process can be iterated until no nonterminal is present in the sentential form. Hence, each derivation in G can be simulated in Π and, conversely, all halting computations in Π correspond to correct derivations in G. Therefore, the computation in Π can stop only after reaching a terminal string with respect to G. Thus, we have $L(G) = L(\Pi)$. \Box

In [1], it was proved that P systems of degree 4 with permitting conditions of type sub_2 and forbidding conditions of type empty can characterize recursively enumerable languages. We improve this result by proving the universality with 3 membranes.

Theorem 2. $RE = RP_3(rw, sub_2, empty)$.

Proof. We start again from a type 0 grammar G = (N, T, S, P) in Penttonen normal form, with the non context-free rules in P labeled in a one-to-one manner, and we construct the P system

$$\Pi = (V, T, [1[2]_2[3]_3]_1, \emptyset, \{S\}, \emptyset, (R_1, P_1, F_1), \dots, (R_3, P_3, F_3)),$$

with the following components:

$$V = N \cup T \cup \{(B, r) \mid r : AB \to AC \in P\}$$
$$\cup \{A', A'' \mid A \in N\} \cup \{f, Z\};$$
$$R_1 = \{f \to \lambda, C'' \to Z\}$$
$$\cup \{C' \to C'' \mid C \in N\};$$
$$P_1 = \{(\lambda, out)\} \cup \{(C'', in) \mid C \in N\}$$

$$\cup \{ (A(B,r),in) \mid r : AB \to AC \in P \};$$

$$R_2 = \{ B \to (B,r) \mid r : AB \to AC \in P \}$$

$$\cup \{ A \to x, \ A \to xf \mid A \to x \in P \}$$

$$\cup \{ C'' \to C \mid C \in N \};$$

$$P_2 = \{ (f,out) \}$$

$$\cup \{ ((B,r),out) \mid r : AB \to AC \in P \};$$

$$R_3 = \{ (B,r) \to C' \mid r : AB \to AC \in P \}$$

$$\cup \{ C'' \to Z \mid C \in N \};$$

$$P_3 = \{ (C',out) \mid C \in N \}.$$

All sets of forbidding conditions consist of the pairs (false, notin), (false, notout).

This system works as follows. We start in membrane 2 with the axiom of G. The context-free rules of G can be simulated here. If a terminal rule $A \to xf$ is used in membrane 2, then the string goes to membrane 1 and from here out of the system. If it is not terminal, it is not accepted in the generated language. If the string is terminal, then it is introduced in $L(\Pi)$.

Suppose that a string w is rewritten in membrane 2 by a rule $B \to (B, r)$ associated with a rule $r : AB \to AC \in P$. It exits; if the symbols A and (B, r)are not associated with the same rule from P, then the string is sent out, but it is not a terminal one. Assume that the string is of the form $w_1A(B,r)w_2$, for some $r : AB \to AC \in P$. No rule can be applied in membrane 1, but the string can be sent to a lower membrane. If it arrives back in membrane 2, then it will exit either unchanged or after introducing one more symbol of the form (B, r). The process is repeated; eventually; the string will arrive in membrane 3 (otherwise we either continue between membrane 1 and 2 or we send the string out of the system and it is not a terminal one). Here in membrane 3 we replace the symbol (B, r) with C' and the string is sent back to the skin membrane. In the skin membrane, the symbol C' is replaced with C''.

Now there are two cases. If we had at least two symbols of the form (B, r) and (B_1, r_1) in the string, then before finishing the simulation of the rule r, we can start the simulation of the rule r_1 . But then the trap symbol Z will be introduced. So we have to finish the simulation of the rule r first. In the other case, the string can be sent to one of membrane 2 and 3. If the string arrives back to 3, then the trap symbol will be introduced. Thus, we have to send the string to membrane 2. We have two cases here. If in membrane 2 we use the rule $C'' \to C$, then we have again a string $(N \cup T)^*$, and the process can be iterated. If before using the rule $C'' \to C$, we use a rule $B \to (B, r)$, then the string should go to membrane 1 where we introduce the trap symbol Z by the rule $C'' \to Z$. Thus $L(G) = L(\Pi)$. \Box

The universality result for P systems with both permitting and forbidding conditions of type symb has been improved from 6 [1] to 5 membranes in [4]. Here

we give a universality result with only 3 membranes. We use the same idea as in [3].

Theorem 3. $RE = RP_3(rw, symb, symb)$.

Proof. Consider a matrix grammar with appearance checking G = (N, T, S, M, F)in the strong binary normal form with $N = N_1 \cup N_2 \cup \{S, \#\}$. Assume that ac(G) = 2, and let $B^{(1)}$ and $B^{(2)}$ be the two objects in N_2 for which we have rules $B^{(j)} \to \#$ in matrices of M. Let us assume that we have k matrices of the form $m_i : (X \to \alpha, A \to x), X \in N_1, \alpha \in N_1 \cup \{\lambda\}, A \in N_2, \text{and } x \in (N_2 \cup T)^*$. We replace each matrix of the form $(X \to \lambda, A \to x)$ by $(X \to f, A \to x)$ where f is a new symbol. We continue to lable the obtained matrices in the same way as the original one. The matrices of the form $(X \to Y, B^{(j)} \to \#)$, are labeled by m_i with $i \in lab_j$, for j = 1, 2 such that lab_1, lab_2 and $lab_0 = \{1, 2, \ldots, k\}$ are mutually disjoint sets.

We construct the P system (of degree 3)

$$\Pi = (V, T, \mu, M_1, \dots, M_3, R_1, P_1, F_1, \dots, R_3, P_3, F_3),$$

with the following components:

$$V = N_1 \cup N_2 \cup T \cup \{X_{i,j} \mid X \in N_1, 1 \le i \le k, 0 \le j \le k\}$$

$$\cup \{A_i, A_{i,j} \mid A \in N_2, 1 \le i \le k, 0 \le j \le k\}$$

$$\cup \{X', X'', X^{(1)}, X^{(2)} \mid X \in N_1 \cup \{f\}\},$$

$$\mu = [1[2[3]_3]_2]_1,$$

$$M_1 = \{XA\}, \text{ for } (S \to XA) \text{ being the initial matrix of } G,$$

$$M_2 = M_3 = \emptyset,$$

and with the following triples $(R_i, P_i, F_i), 1 \le i \le 3$:

$$\begin{split} R_{1} &= \{X \to Y^{(1)} \mid m_{i} : (X \to Y, B^{(1)} \to \#\} \\ &\cup \{X \to Y^{(2)} \mid m_{i} : (X \to Y, B^{(2)} \to \#\} \\ &\cup \{A \to A_{i,0} \mid m_{i} : (X \to \alpha, A \to x), 1 \leq i \leq k, \alpha \in N_{1} \cup \{f\}\} \\ &\cup \{A_{i,j} \to \# \mid 1 \leq j < i \leq k\} \\ &\cup \{A_{i} \to x \mid m_{i} : (X \to \alpha, A \to x), 1 \leq i \leq k, \alpha \in N_{1} \cup \{f\}\} \\ &\cup \{\alpha' \to \alpha \mid \alpha \in N_{1}\} \cup \{f' \to \lambda\}; \\ P_{1} &= \{(A_{i,0}, in) \mid i \leq i \leq k\} \\ &= \{(X^{(1)}, in), (X^{(2)}, in) \mid X \in N_{1}\} \\ &= \{(a, out) \mid a \in T\}; \\ F_{1} &= \{(X, notout) \mid X \in N_{1} \cup N_{2}\} \\ &\cup \{(A_{i, n}, notin) \mid A \in N_{2}, 1 \leq i \leq k\} \\ &\cup \{(A_{i, j}, notout) \mid A \in N_{2}, 1 \leq i, j \leq k\} \end{split}$$

$$\cup \{(\alpha', notin) \mid \alpha \in N_1 \cup \{f\}\}; \\ R_2 = \{X \to X_{i,0} \mid m_i : (X \to \alpha, A \to x), 1 \le i \le k, \alpha \in N_1 \cup \{f\}\} \\ \cup \{B^{(1)} \to \#, \ \# \to \#\} \\ \cup \{Y^{(1)} \to Y, \ Y'' \to Y \mid Y \in N_1\} \\ \cup \{X_{i,j} \to X_{i,j+1} \mid X \in N_1, 1 \le j < i \le k\} \\ \cup \{X_{i,i} \to \alpha' \mid m_i : (X \to \alpha, A \to x), 1 \le i \le k, \alpha \in N_1 \cup \{f\}\}; \\ P_2 = \{(Y, out), (Y^{(2)}, in) \mid Y \in N_1\} \\ \cup \{(X_{i,j}, in) \mid 1 \le j < i \le k\} \\ \cup \{(\alpha', out) \mid \alpha \in N_1 \cup \{f\}\}; \\ F_2 = \{(B^{(1)}, notout) \mid B \in N_2\} \\ \cup \{(\alpha'', notin) \mid \alpha \in N_1 \cup \{f\}\}; \\ R_3 = \{Y^{(2)} \to Y''\} \cup \{B^{(2)} \to \#, \# \to \#\} \\ \cup \{A_{i,j} \to A_{i,j+1} \mid A \in N_2, 1 \le j < i \le k\} \\ \cup \{A_{i,i} \to A_i, \ A_i \to \# \mid 1 \le i \le k\}; \\ P_3 = \{(Y', out) \mid Y \in N_1\} \\ \cup \{(A_{i,j}, out) \mid 1 \le j < i \le k\}; \\ F_3 = \{(B^{(2)}, notout) \mid B \in N_2\}.$$

Only strings over T are accepted in the generated language; # is a trap symbol. From the skin membrane in any moment we can send out a string if it contains at least one terminal symbol, but the string is not accepted in the generated language if it contains any symbol not in T.

Simulation of the matrix $m_i : (X \to \alpha, A \to x), 1 \le i \le k$:

We start the simulation by the rule $A \to A_{i,0}$. The string can be sent to membrane 2 where we apply the rule $X \to X_{j,0}$. The obtained string is sent to membrane 3. From now on, the string will go back and forth between membranes 2 and 3, and the second subscript of the symbols $X_{i,s}$ and $Y_{j,t}$ is alternatively increased. Now we have three cases here:

Case 1: i < j. This means that at some step in membrane 3 we have a string of the form $X_{j,i}w_1A_{i,i-1}w_2$. We replace $A_{i,i-1}$ with $A_{i,i}$ and no communication is possible. So we use the rule $A_{i,i} \rightarrow A_i$ and the string is sent out. In membrane 2, we replace $X_{j,i}$ with $X_{j,i+1}$ and sent the string back to membrane 3, where the trap symbol # is introduced (the rewriting has priority over communication).

Case 2: i > j. At some moment we have a string of the form $X_{j,j}w_1A_{i,j-1}w_2$ in membrane 2 which is sent to membrane 3. Here we replace $A_{i,j-1}$ with $A_{i,j}$ and send the string out. In membrane 2 we replace $X_{j,j}$ with α' , and the string is sent out. In the skin membrane, we can apply $A_{i,j} \to \#$, hence the string will never lead to a terminal one.

Case 3: i = j. At some moment we pass from membrane 2 to membrane 3 a string

 $X_{i,i}w_1A_{i,i-1}w_2$. In membrane 3, we replace $A_{i,i-1}$ with $A_{i,i}$ and, because we cannot exit, we replace $A_{i,i}$ with A_i and send out the string. Here in membrane 2 we replace $X_{i,i}$ with α' and send the string to the skin membrane. In the skin membrane we have to replace α' with α and A_i with x before starting the simulation of the next matrix.

Simulation of the matrices $(X \to Y, B^{(j)} \to \#), j = 1, 2.$

The simulation of a matrix of this form starts by a rule $X \to Y^{(1)}$ or $X \to Y^{(2)}$ in the skin membrane. If we are simulating a rule $(X \to Y, B^{(1)} \to \#)$, then in membrane 2 we use the rule $Y^{(1)} \to Y$. Now the string can be sent to the skin membrane only if $B^{(1)}$ is not present. Similarly, if we are simulating $(X \to Y, B^{(2)} \to \#)$, then in membrane 2 there is no rule we can apply. So we send the string to membrane 3, where we replace $Y^{(2)}$ with Y'' and the resulting string can be sent out if $B^{(2)}$ is not present. Back in membrane 2, we replace Y'' with Y and send the string to the skin membrane.

If at any moment we get a string of the form f'w, for $w \in T^*$, in the skin membrane, then we remove f' and send the string out. Consequently, $L(G) = L(\Pi)$. \Box

There is a characterization of recursively enumerable languages by P systems with permitting conditions of type $prefsuff_2$ and forbidding conditions of type empty in [1] without a bound on the number of membranes. It was conjectured that such a characterization holds also for a reduced number of membranes. We settle this conjecture in the positive here and show that *eight* membranes are enough for achieving the universality.

Theorem 4. $RE = RP_8(rw, prefsuff_2, empty)$.

Proof. Let us consider a type 0 grammar G = (N, T, S, P) in Penttonen normal form, with the non context-free rules in P labeled in an injective manner, and assume that $N \cup T \cup \{\$\} = \{E_1, E_2, \ldots, E_n\}$. We construct the P system Π , of degree 8, with the following components:

$$V = N \cup T \cup \{A' \mid A \in N\}$$

$$\cup \{X, Y, Y', Z, \$\}$$

$$\cup \{X_i, Y_i, X_{i,j}, Y_{i,j} \mid 1 \le i \le n, 0 \le j \le n\}$$

$$\cup \{(B, r) \mid r : AB \to AC \in P\},$$

$$\mu = [1[2[3[4[5[6]6]5]4]3[7[8]8]7]2]1,$$

$$M_i = \emptyset, 1 \le i \le 8, i \ne 2,$$

$$M_2 = X\$SY,$$

and with the following sets of rules and associated permitting conditions. All forbidding condition sets are of the form $\{(false, notin), (false, notout)\}$:

$$R_1 = \{ X \to \lambda, \ Y \to \lambda, \ \$ \to \lambda \};$$

$$P_1 = \{ (a, out) \mid a \in T \};$$

$$\begin{split} R_2 &= \{E_i \to E'_i, \ Y \to Y_{i,0} \mid 1 \leq i \leq n\} \\ &\cup \{B \to (B, r) \mid r : AB \to AC \in P\} \\ &\cup \{A \to x \mid A \to x \in P\} \\ &\cup \{(B, r) \to Z \mid r : AB \to AC \in P\} \\ &\cup \{E'_i \to Z, \ Y_{i,0} \to Z \mid 1 \leq i \leq n\}; \\ P_2 &= \{(\$Y, out)\} \\ &\cup \{(E'_i Y_{i,0}, in) \mid r : AB \to AC \in P, \ 1 \leq i \leq n\} \\ &\cup \{((B, r)Y, in) \mid r : AB \to AC \in P\}; \\ R_3 &= \{E'_i \to \lambda, Y_i \to Y \mid 1 \leq i \leq n\} \\ &\cup \{(B, r) \to Z \mid r : AB \to AC \in P\}; \\ P_3 &= \{(Y_{i,0}, in) \mid 1 \leq i \leq n\} \\ &\cup \{(Y, out)\}; \\ R_4 &= \{X \to X_{i,0}E_i, \ X_i \to X \mid 1 \leq i \leq n\} \\ &\cup \{Y_{i,j} \to Z \mid 1 \leq j < i \leq n\}; \\ P_4 &= \{(X_{i,0}, in) \mid 1 \leq i \leq n\} \\ &\cup \{X_{i,i} \to X_i \mid 1 \leq i \leq n\}; \\ P_5 &= \{X_{i,j} \to X_{i,j+1} \mid 0 \leq j < i \leq n\} \\ &\cup \{X_{i,i} \to X_i \mid 1 \leq i \leq n\}; \\ R_6 &= \{Y_{i,j} \to Y_{i,j+1} \mid 0 \leq j < i \leq n\} \\ &\cup \{Y_{i,i} \to Y_i, \ Y_i \to Z \mid 1 \leq i \leq n\}; \\ R_7 &= \{Y \to \lambda, \ Y' \to Y\} \\ &\cup \{Y_{i,0} \to Z \mid 1 \leq i \leq n\}; \\ R_7 &= \{(A(B, r), in), \ (CY, out) \mid r : AB \to AC \in P\}; \\ R_8 &= \{(CY', out)\}. \end{split}$$

We start from the string X\$SY, initially present in membrane 2. We plan to simulate the non context-free rules from P in the right end of the strings of Π and to this aim we use the so-called *rotate-and-simulate* technique much used in the DNA computing area. If Xw_1 \$ w_2Y is a sentential form of Π , then w_2w_1 is a sentential form of G. The symbol \$ indicates the actual beginning of strings from G. Z is a trap symbol, once introduced, it cannot be removed, hence the string will never become a terminal one. In membrane 2, we can simulate any context-free rule from P and the string will remain in the same region. We start the procedure of circularly permuting the string with one symbol by using the rules $E_i \to E'_i$ and $Y \to Y_{i,0}$. If the primed symbol is the right most one, then the condition to send the string to a lower membrane is fulfilled. If we did not use both the rules or the primed symbol is not the rightmost one, then the trap symbol is introduced. Now we can either send the string to membrane 7 or 3. If it enters membrane 7, then we introduce the trap symbol.

In membrane 3, we remove E'_i and send the string to membrane 4. In membrane 4, we replace X with $X_{i,0}E_i$ and send the string to membrane 5. From now on, the string will go back and forth between membranes 5 and 6, and the second subscript of the symbols $X_{i,s}$ and $Y_{j,t}$ is alternatively increased. Now there are three cases:

Case 1: i < j. This means that at some step in membrane 6 we have a string $X_{j,i}wY_{i,i-1}$. We replace $Y_{i,i-1}$ with $Y_{i,i}$ and no communication is possible, hence one more rewriting is necessary. We replace $Y_{i,i}$ with Y_i and the string is sent out. In membrane 5 we replace $X_{j,i}$ with $X_{j,i+1}$ and the string is sent back to membrane 6, where we introduce the trap symbol.

Case 2: i > j. At some moment we have a string of the form $X_{j,j}wY_{i,j-1}$ in membrane 5, which is sent to membrane 6. We replace $Y_{i,j-1}$ with $Y_{i,j}$ in membrane 6 and the string exits. In membrane 5 we replace $X_{j,j}$ with X_j and the string is sent out. Back in membrane 4 we can apply $Y_{i,j} \to Z$, hence the string will never lead to a terminal one.

Case 3: i = j. At some moment we pass from membrane 5 to membrane 6 a string $X_{i,i}wY_{i,i-1}$. In membrane 6 we replace $Y_{i,i-1}$ with $Y_{i,i}$ and, because we cannot exit, we replace $Y_{i,i}$ with Y_i and sent the string out. In membrane 5 we replace $X_{i,i}$ with X_i and the string is sent out. We replace the symbols X_i and Y_i with X and Y respectively in membrane 4 and 5 and the string is sent to membrane 2. The process of circularly permuting the symbol will end successful if we add the symbol E_i in the left end of the string corresponding to the symbol E'_i which was removed from the right end of the string

We simulate the non context-free rules $r : AB \to AC$, in the following way. A symbol B is replaced by (B, r) in membrane 2, if this is not done in the right most position, then the symbol Z is introduced. If the string is of the form Xw(B, r)Y, then it has to go to membrane 7. In membrane 7 we replace the symbol Y and send the string to membrane 8, if the string is of the form $Xw_1A(B, r)$ corresponding to some rule $r : AB \to AC \in P$. In membrane 8 we replace (B, r) with CY' and send out the resulting string . In membrane 7 we replace Y' with Y and send the string to membrane 2.

The process can be iterated. Consequently, $L(G) = L(\Pi)$. \Box

5 Conclusion

In this paper we gave some improved results about rewriting P systems with conditional communication. We believe that the result of Theorem 3 cannot be improved further. It is an open problem whether or not the result of Theorem 4 can be improved.

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Characterizing Membrane Structures Through Multiset Tree Automata^{*}

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Summary. The relation between the membrane structures of P systems and an extension of tree automata which introduces multisets in the transition function has been proposed in previous works. Here we propose two features of tree automata which have been previously studied (namely, reversibility and local testability) in order to extend them to multiset tree automata. The characterization of these families will introduce a new characterization of membrane structures defined by the set of rules used for membrane creation and deletion.

1 Introduction

The relation between membrane structures and tree languages has been explored in previous works. So, Freund et al. [4] proved that P systems are able to generate recursively enumerable sets of trees through their membrane structures. Other works have focused on extending the definition of finite tree automata in order to take into account the membrane structures generated by P systems. So, in [13], the authors propose an extension of tree automata, namely multiset tree automata, in order to recognize membrane structures. In [7], this model is used to calculate editing distances between membrane structures. Later, a method to infer multiset tree automata from membrane observations was presented in [14].

In this work we introduce two new families of multiset tree automata, by using previous results taken from tree language theory. We propose a formal definition of reversible multiset tree automata and local testable multiset tree automata. These features have been widely studied in previous works [6, 8].

The structure of this work is simple: first we give basic definitions and notation for tree languages, P systems and multiset tree automata and we define the new families of multiset tree automata. Finally, we give some guidelines for future research.

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2 Notation and Definitions

In the sequel we will provide some concepts from formal language theory, membrane systems and multiset processing. We suggest the books [12], [10] and [2] to the reader.

Multisets

First, we will provide some definitions from multiset theory as exposed in [15].

Definition 1. Let D be a set. A multiset over D is a pair $\langle D, f \rangle$ where $f : D \longrightarrow \mathbb{N}$ is a function.

Definition 2. Suppose that $A = \langle D, f \rangle$ and $B = \langle D, g \rangle$ are two multisets. The removal of multiset B from A, denoted by $A \ominus B$, is the multiset $C = \langle D, h \rangle$ where for all $a \in D$ h(a) = max(f(a) - g(a), 0).

Definition 3. Let $A = \langle D, f \rangle$ be a multiset; we will say that A is empty if for all $a \in D$, f(a) = 0.

Definition 4. Let $A = \langle D, f \rangle$ and $B = \langle D, g \rangle$ be two multisets. Their sum, denoted by $A \oplus B$, is the multiset $C = \langle D, h \rangle$, where for all $a \in D$ h(a) = f(a) + g(a).

Definition 5. Let $A = \langle D, f \rangle$ and $B = \langle D, g \rangle$ be two multisets. We will say that A = B if the multiset $(A \ominus B) \oplus (B \ominus A)$ is empty.

The size of any multiset M, denoted by |M| will be the number of elements that it contains. We are specially interested in the class of multisets that we call *bounded multisets*. They are multisets that hold the property that the sum of all the elements is bounded by a constant n. Formally, we will denote by $\mathcal{M}_n(D)$ the set of all multisets $\langle D, f \rangle$ such that $\sum_{a \in D} f(a) = n$.

A concept that is quite useful to work with sets and multisets is the *Parikh* mapping. Formally, a Parikh mapping can be viewed as the application $\Psi : D^* \to \mathbb{N}^n$ where $D = \{d_1, d_2, \dots, d_n\}$. Given an element $x \in D^*$ we define $\Psi(x) = (\#_{d_1}(x), \dots, \#_{d_n}(x))$ where $\#_{d_i}(x)$ denotes the number of occurrences of d_i in x.

P systems

We will introduce basic concepts from membrane systems taken from [10]. A general P system of degree m is a construct

$$\Pi = (V, T, C, \mu, w_1, \cdots, w_m, (R_1, \rho_1), \cdots, (R_m, \rho_m), i_0),$$

where:

- V is an alphabet (the *objects*)
- $T \subseteq V$ (the *output alphabet*)

- $C \subseteq V, C \cap T = \emptyset$ (the *catalysts*)
- μ is a membrane structure consisting of m membranes
- $w_i, 1 \leq i \leq m$, is a string representing a multiset over V associated with the region i
- $R_i, 1 \le i \le m$, is a finite set of *evolution rules* over V associated with the *i*th region and ρ_i is a partial order relation over R_i specifying a *priority*.

An evolution rule is a pair (u, v) (or $u \to v$) where u is a string over V and v = v' or $v = v'\delta$ where v' is a string over

$$\{a_{here}, a_{out}, a_{in_j} \mid a \in V, 1 \le j \le m\}$$

and δ is a special symbol not in V (it defines the membrane dissolving action). From now on, we will denote the set tar by {here, out, $in_k : 1 \le k \le m$ }.

• i_0 is a number between 1 and m and it specifies the *output* membrane of Π (in the case that it equals to ∞ the output is read outside the system).

The language generated by Π in external mode $(i_0 = \infty)$ is denoted by $L(\Pi)$ and it is defined as the set of strings that can be defined by collecting the objects that leave the system by arranging them in the leaving order (if several objects leave the system at the same time then permutations are allowed). The set of vector numbers that represent the objects in the output membrane i_0 will be denoted by $N(\Pi)$. Obviously, both sets $L(\Pi)$ and $N(\Pi)$ are defined only for halting computations.

One of the multiple variations of P systems is related to the creation, division and modification of membrane structures. There have been several works in which these variants have been proposed (see, for example, [1, 9, 10, 11]).

In the following, we enumerate some kind of rules which are able to modify the membrane structure:

- 1. 2-division: $[ha]_h \rightarrow [h'b]_{h'}[h''c]_{h''}$
- 2. Creation: $a \to [hb]_h$
- 3. Dissolving: $[{}_{h}a]_{h} \rightarrow b$

The power of P systems with the previous operations and other ones (e.g., *exocytosis, endocytosis,* etc.) has been widely studied in the membrane computing area.

Tree automata and tree languages

Now, we will introduce some concepts from tree languages and automata as exposed in [3, 5]. First, let a *ranked alphabet* be the association of an alphabet V together with a finite relation r in $V \times \mathbb{N}$. We denote by V_n the subset $\{\sigma \in V \mid (\sigma, n) \in r\}$.

The set V^T of trees over V, is defined inductively as follows:

$$a \in V^T$$
 for every $a \in V_0$
 $\sigma(t_1, ..., t_n) \in V^T$ whenever $\sigma \in V_n$ and $t_1, ..., t_n \in V^T$, $(n > 0)$

and let a *tree language* over V be defined as a subset of V^T .

Given the tuple $l = \langle 1, 2, ..., k \rangle$ we will denote the set of permutations of l by perm(l). Let $t = \sigma(t_1, ..., t_n)$ be a tree over V^T . We denote the set of permutations of t at first level by $perm_1(t)$. Formally, $perm_1(t) = \{\sigma(t_{i_1}, ..., t_{i_n}) \mid \langle i_1, i_2, ..., i_n \rangle \in perm(\langle 1, 2, ..., n \rangle) \}$.

Let \mathbb{N}^* be the set of finite strings of natural numbers, separated by dots, formed using the catenation as the composition rule and the empty word λ as the identity. Let the prefix relation \leq in \mathbb{N}^* be defined by the condition that $u \leq v$ if and only if $u \cdot w = v$ for some $w \in \mathbb{N}^*$ $(u, v \in \mathbb{N}^*)$. A finite subset D of \mathbb{N}^* is called a *tree domain* if:

> $u \leq v$ where $v \in D$ implies $u \in D$, and $u \cdot i \in D$ whenever $u \cdot j \in D$ $(1 \leq i \leq j)$

Each tree domain D could be seen as an unlabeled tree whose nodes correspond to the elements of D where the hierarchy relation is the prefix order. Thus, each tree t over V can be seen as an application $t : D \to V$. The set D is called the *domain of the tree* t, and denoted by dom(t). The elements of the tree domain dom(t) are called *positions* or *nodes* of the tree t. We denote by t(x) the label of a given node x in dom(t).

Let the level of $x \in dom(t)$ be |x|. Intuitively, the level of a node measures its distance from the root of the tree. Then, we can define the depth of a tree t as $depth(t) = max\{|x| : x \in dom(t)\}$. In the same way, for any tree t, we denote the size of the tree by |t| and the set of subtrees of t (denoted with Sub(t)) as follows:

$$Sub(a) = \{a\} \text{ for all } a \in V_0$$

$$Sub(t) = \{t\} \cup \bigcup_{i=1,\dots,n} Sub(t_i) \text{ for } t = \sigma(t_1,\dots,t_n) \ (n > 0)$$

Given a tree $t = \sigma(t_1, \ldots, t_n)$, the root of t will be denoted as root(t) and defined as $root(t) = \sigma$. If t = a then root(t) = a. The successors of a tree $t = \sigma(t_1, \ldots, t_n)$ will be defined as $H^t = \langle root(t_1), \ldots, root(t_n) \rangle$. Finally, leaves(t) will denote the set of leaves of the tree t.

Definition 6. A finite deterministic tree automaton is defined by the tuple $A = (Q, V, \delta, F)$ where Q is a finite set of states; V is a ranked alphabet with m as the maximum integer in the relation $r, Q \cap V = \emptyset; F \subseteq Q$ is the set of final states and $\delta = \bigcup_{i:V_i \neq \emptyset} \delta_i$ is a set of transitions defined as follows:

$$\delta_n : (V_n \times (Q \cup V_0)^n) \to Q \qquad n = 1, \dots, m$$
$$\delta_0(a) = a \qquad \forall a \in V_0$$

Given the state $q \in Q$, we define the *ancestors* of the state q, denoted by Ant(q), as the set of strings

$$Ant(q) = \{ p_1 \cdots p_n \mid p_i \in Q \cup V_0 \land \delta_n(\sigma, p_1, \dots, p_n) = q \}$$

From now on, we will refer to finite deterministic tree automata simply as *tree automata*. We suggest [3, 5] for other definitions on tree automata.

The transition function δ is extended to a function $\delta: V^T \to Q \cup V_0$ on trees as follows:

$$\delta(a) = a \text{ for any } a \in V_0$$

$$\delta(t) = \delta_n(\sigma, \delta(t_1), \dots, \delta(t_n)) \text{ for } t = \sigma(t_1, \dots, t_n) \ (n > 0)$$

Note that the symbol δ denotes both the set of transition functions of the automaton and the extension of these functions to operate on trees. In addition, you can observe that the tree automaton A cannot accept any tree of depth zero.

Given a finite set of trees T, let the subtree automaton for T be defined as $AB_T = (Q, V, \delta, F)$, where:

$$Q = Sub(T)$$

$$F = T$$

$$\delta_n(\sigma, u_1, \dots, u_n) = \sigma(u_1, \dots, u_n) \qquad \sigma(u_1, \dots, u_n) \in Q$$

$$\delta_0(a) = a \qquad a \in V_0$$

Let \$ be a new symbol in V_0 , and $V_{\T the set of trees $(V \cup \{\$\})^T$ where each tree contains \$ only once. We will name the node with label \$ as *link point* when necessary. Given $s \in V_{\T and $t \in V^T$, the operation s # t is defined as:

$$s\#t(x) = \begin{cases} s(x) \text{ if } x \in dom(s), \ s(x) \neq \$ \\ t(z) \text{ if } x = yz, \ s(y) = \$, \ y \in dom(s) \end{cases}$$

therefore, given $t, s \in V^T$, let the tree quotient $(t^{-1}s)$ be defined as

$$t^{-1}s = \begin{cases} r \in V_{\$}^T : s = r \# t \text{ if } t \in V^T - V_0 \\ t & \text{ if } t \in V_0 \end{cases}$$

This quotient can be extended to consider set of trees $T \subseteq V^T$ as:

$$t^{-1}T = \{t^{-1}s \mid s \in T\}$$

For any $k \ge 0$, let the k-root of a tree t be defined as follows:

$$root_k(t) = \begin{cases} t, & \text{if } depth(t) < k \\ t': t'(x) = t(x), \ x \in dom(t) \land |x| \le k, \text{ otherwise} \end{cases}$$

Multiset tree automata and mirrored trees

We will extend over multisets some definitions of tree automata and tree languages. We will introduce the concept of multiset tree automata and then we will characterize the set of trees that it accepts.

Given any tree automaton $A = (Q, V, \delta, F)$ and $\delta_n(\sigma, p_1, p_2, \ldots, p_n) \in \delta$, we can associate to δ_n the multiset $\langle Q \cup V_0, f \rangle \in \mathcal{M}_n(Q \cup V_0)$ where f is defined by $\Psi(p_1p_2 \ldots p_n)$. The multiset defined in such way will be denoted by $M_{\Psi}(\delta_n)$. Alternatively, we can define $M_{\Psi}(\delta_n)$ as $M_{\Psi}(p_1) \oplus M_{\Psi}(p_2) \oplus \cdots \oplus M_{\Psi}(p_n)$ where $\forall 1 \leq i \leq n \ M_{\Psi}(p_i) \in \mathcal{M}_1(Q \cup V_0)$. Observe that if $\delta_n(\sigma, p_1, p_2, \ldots, p_n) \in \delta$, $\delta'_n(\sigma, p'_1, p'_2, \ldots, p'_n) \in \delta$ and $M_{\Psi}(\delta_n) = M_{\Psi}(\delta'_n)$ then δ_n and δ'_n are defined over the same set of states and symbols but in different order (that is the multiset induced by $\langle p_1, p_2, \cdots, p_n \rangle$ equals the one induced by $\langle p'_1 p'_2 \ldots p'_n \rangle$).

Now, we can define a *multiset tree automaton* that performs a bottom-up parsing as in the tree automaton case.

Definition 7. A multiset tree automaton is defined by the tuple $MA = (Q, V, \delta, F)$, where Q is a finite set of states, V is a ranked alphabet with maxarity(V) = n, $Q \cap V = \emptyset$, $F \subseteq Q$ is a set of final states and δ is a set of transitions defined as follows:

$$\delta = igcup_{1 \le i \le n} \delta_i \ i : V_i
eq \emptyset$$

$$\delta_i : (V_i \times \mathcal{M}_i(Q \cup V_0)) \to \mathcal{P}(\mathcal{M}_1(Q)) \qquad i = 1, \dots, n$$

$$\delta_0(a) = M_{\Psi}(a) \in \mathcal{M}_1(Q \cup V_0) \qquad \forall a \in V_0$$

We can observe that every tree automaton A defines a multiset tree automaton MA as follows

Definition 8. Let $A = (Q, V, \delta, F)$ be a tree automaton. The multiset tree automaton induced by A is defined by the tuple $MA = (Q, V, \delta', F)$ where each δ' is defined as follows: $M_{\Psi}(r) \in \delta'_n(\sigma, M)$ if $\delta_n(\sigma, p_1, p_2, ..., p_n) = r$ and $M_{\Psi}(\delta_n) = M$.

Observe that, in the general case, the multiset tree automaton induced by A is non deterministic.

As in the case of tree automata, δ' could also be extended to operate on trees. Here, the automaton carries out a bottom-up parsing where the tuples of states and/or symbols are transformed by using the Parikh mapping Ψ to obtain the multisets in $\mathcal{M}_n(Q \cup V_0)$. If the analysis is completed and δ' returns a multiset with at least one final state, the input tree is accepted. So, δ' can be extended as follows

$$\delta'(a) = M_{\Psi}(a) \text{ for any } a \in V_0$$

$$\delta'(t) = \{ M \in \delta'_n(\sigma, M_1 \oplus \dots \oplus M_n) \mid M_i \in \delta'(t_i) 1 \le i \le n \}$$

for $t = \sigma(t_1, \dots, t_n) \ (n > 0)$

Formally, every multiset tree automaton MA accepts the following language

$$L(MA) = \{ t \in V^T \mid M_{\Psi}(q) \in \delta'(t), q \in F \}$$

Another extension which will be useful is the one related to the ancestors of every state. So, we define $Ant_{\Psi}(q) = \{M \mid M_{\Psi}(q) \in \delta_n(\sigma, M)\}.$

Theorem 1. (Sempere and López, [13]) Let $A = (Q, V, \delta, F)$ be a tree automaton, $MA = (Q, V, \delta', F)$ be the multiset tree automaton induced by A and $t = \sigma(t_1, \ldots, t_n) \in V^T$. If $\delta(t) = q$ then $M_{\Psi}(q) \in \delta'(t)$.

Corollary 1. (Sempere and López, [13]) Let $A = (Q, V, \delta, F)$ be a tree automaton and $MA = (Q, V, \delta', F)$ be the multiset tree automaton induced by A. If $t \in L(A)$ then $t \in L(MA)$.

We will introduce the concept of *mirroring* in tree structures as exposed in [13]. Informally speaking, two trees will be related by mirroring if some permutations at the structural level hold. We propose a definition that relates all the trees with this mirroring property.

Definition 9. Let t and s be two trees from V^T . We will say that t and s are mirror equivalent, denoted by $t \bowtie s$, if one of the following conditions holds:

1.
$$t = s = a \in V_0$$

2. $t \in perm_1(s)$
3. $t = \sigma(t_1, \dots, t_n), s = \sigma(s_1, \dots, s_n)$ and there exists $\langle s^1, s^2, \dots, s^k \rangle$
 $\in perm(\langle s_1, s_2, \dots, s_n \rangle)$ such that $\forall 1 \le i \le n \ t_i \bowtie s^i$

Theorem 2. (Sempere and López, [13]) Let $A = (Q, V, \delta, F)$ be a tree automaton, $t = \sigma(t_1, \ldots, t_n) \in V^T$ and $s = \sigma(s_1, \ldots, s_n) \in V^T$. Let $MA = (Q, V, \delta', F)$ be the multiset tree automaton induced by A. If $t \bowtie s$ then $\delta'(t) = \delta'(s)$.

Corollary 2. (Sempere and López, [13]) Let $A = (Q, V, \delta, F)$ be a tree automaton, $MA = (Q, V, \delta', F)$ the multiset tree automaton induced by A and $t \in V^T$. If $t \in L(MA)$ then, for any $s \in V^T$ such that $t \bowtie s, s \in L(MA)$.

The last results were useful to propose an algorithm to determine whether two trees are mirror equivalent or not [13]. So, given two trees s and t, we can establish in time $\mathcal{O}((\min\{|t|, |s|\})^2)$ if $t \bowtie s$.

3 k-testable in the Strict Sense (k-TSS) Multiset Tree Languages

In this section, we will define a new class of multiset tree languages. The definitions related to multiset tree automata come from the relation between mirrored trees and multiset tree automata which we have established in the previous section. So, whenever we refer to multiset tree languages we are taking under our consideration the set of (mirrored) trees accepted by multiset tree automata.

We refer to [6] for more details about reversibility and local testability in tree languages.

First, we define k-TSS multiset tree languages for any $k \geq 2$.

Definition 10. Let $T \subseteq V^T$ and the integer value $k \geq 2$. T is a k-TSS multiset tree language if and only if, given whatever two trees $u_1, u_2 \in V^T$ such that $root_{k-1}(u_1) = root_{k-1}(u_2), u_1^{-1}T \neq \emptyset$ and $u_2^{-1}T \neq \emptyset$ implies that $u_1^{-1}T = u_2^{-1}T$

Any multiset tree automaton as in the definition given before will be named a k-TSS multiset tree automaton. Given A a tree automaton, t_1 , t_2 valid contexts over $V_{\T , as an extension of a result concerning k-TSS tree languages, we give the following definition:

Definition 11. Let A be a multiset tree automaton over $V_{\T . Let $u_1, u_2 \in V^T$ be two trees such that $root_{k-1}(u_1) = root_{k-1}(u_2)$ and $t_1 \# u_1, t_2 \# u_2 \in L(A)$ for some valid contexts t_1 and t_2 . If A is a k-TSS mirror tree automaton then $\delta(u_1) = \delta(u_2)$.

We can give the following characterization of such automata.

Corollary 3. Let A be a k-TSS multiset tree automaton. There does not exist two distinct states q_1, q_2 such that $root_k(q_1) \cap root_k(q_2) \neq \emptyset$

The previous result can be easily deduced from the definition of k-TSS multiset tree automata and the definitions given in section 2 about tree automata and tree languages.

Example 1. Consider the multiset tree automaton with transitions:

$$\begin{split} \delta(\sigma, aa) &= q_1 \\ \delta(\sigma, a) &= q_2 \\ \delta(\sigma, aq_2) &= q_2 \\ \delta(\sigma, q_1q_1) &= q_1 \\ \delta(\sigma, aq_2q_1) &= q_3 \in F \end{split}$$

Note that the multiset tree language accepted by the automaton is k-TSS for any $k \geq 2$.

Note also that the following one does not have the k-TSS condition for any $k \ge 2$:

F

$$\begin{split} \delta(\sigma, aa) &= q_1 \\ \delta(\sigma, bb) &= q_2 \\ \delta(\sigma, q_2 q_2) &= q_2 \\ \delta(\sigma, q_1 q_1) &= q_1 \\ \delta(\sigma, q_2 q_1) &= q_3 \in \end{split}$$

because both the states q_1 and q_2 (and q_3) share a common k-root.

4 Reversible Multiset Tree Automata

We also extend a previous result concerning k-reversible tree languages (for any $k \ge 0$) to give the following definition.

Definition 12. Let $T \subseteq V^T$ and the integer value $k \geq 0$. T is a k-reversible multiset tree language if and only if, given whatever two trees $u_1, u_2 \in V^T$ such that $root_{k-1}(u_1) = root_{k-1}(u_2)$, whenever there exists a context $t \in V_{\T such that both $u_1 \# t, u_2 \# t \in T$, then $u_1^{-1}T = u_2^{-1}T$

Definition 13. Let A be a multiset tree automaton over $V_{\T . Let $p_1, p_2 \in Q$ be two states such that $root_k(L(p_1)) \cap root_k(L(p_2)) \neq \emptyset$. A is order k reset free if the automaton does not contain two transitions such that

$$\delta(\sigma, q_1 q_2 \dots q_n p_1) = \delta(\sigma, q_1 q_2 \dots q_n p_2)$$

where $q_i \in Q$, $1 \leq i \leq n$.

Definition 14. Let A be a multiset tree automaton. A is k-reversible if A is order k reset free and for any two distinct final states f_1 and f_2 the condition $root_k(L(f_1)) \cap root_k(L(f_2)) \neq \emptyset$ is fulfilled.

Example 2. Consider the multiset tree automaton with transitions:

$$\begin{split} &\delta(\sigma, aa) = q_1 \\ &\delta(\sigma, a) = q_2 \\ &\delta(\sigma, q_2 q_2) = q_2 \\ &\delta(\sigma, aa q_1) = q_1 \\ &\delta(\sigma, q_1 q_1) = q_3 \in F \\ &\delta(\sigma, q_2 q_1) = q_3 \in F \end{split}$$

The multiset tree language accepted by this automaton is k-reversible and it is also an example of non k-TSS multiset tree language.

A relation between the previous classes

Finally, we can relate the two families of multiset tree languages that we have previously defined with the following result.

Theorem 3. Let $T \subseteq V^T$ and an integer value $k \geq 2$. If T is k-TSS then T is (k-1)-reversible.

Proof.

Let $t\#t_1$ and $t\#t_2$ belong to T, with $t \in V_{\T and $root_k(t_1) = root_k(t_2)$. Trivially, $t_1^{-1}T \neq \emptyset$ and $t_2^{-1}T \neq \emptyset$. If T is a k-TSS tree language, then by previous definitions, $t_1^{-1}T = t_2^{-1}T$, and also T is (k-1)-reversible.

5 From Transitions to Membrane Structures

Once we have formally defined the two classes of multiset tree automata, we will translate their characteristics in terms of membrane structures. First we will give a meaning to the concept of $root_k(t)$. Observe that in a membrane structure t, which is represented by a set of mirrored trees $\{t' \mid t \bowtie t'\}$, the meaning of $root_k(t)$ is established by taking into account the (sub)structure of the membranes from the top region up to a depth of length k. Another concept that we have managed before is the operator #. Observe that this is related to membrane creation of P systems. So, we can go from membrane configuration t to t#s by creating a new membrane structure s in a predefined region of t (established by #).

So, k testability implies that, whenever we take two membrane structures u_1 and u_2 at any level of the P regions, if they are part of a common structure then they are part of the same set of structures (u_1 cannot appear as a substructure of a membrane structure if u_2 does not appear as a substructure of the same membrane structure at the same level).

On the other hand, k-reversibility implies that whenever two membrane structures u_1 and u_2 share the same substructure up to length k - 1, if u_1 and u_2 have a common structure t such that $u_1 \# t$ and $u_2 \# t$ are valid configurations of the P system, then $u_1 \# s$ is a valid configuration of the P system iff so is $u_2 \# s$.

6 Conclusions and Future Work

We have introduced two new families of multiset tree languages. These classes have characterized the membrane structures defined by P systems. We think that other classes of tree languages will imply new classes of membrane structures. So, all the theory that has been previously established on tree languages can enrich the way in which we look up to the membrane structures. In addition, there is another way to explore the relation between the membrane structures of P systems and the languages that they can accept or generate. So, a natural question arises: How is affected the structure of the language by the structure of the membranes?

This issue will be explored in future works.

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$OPERAS_{CC}$: An Instance of a Formal Framework for MAS Modelling Based on Population P Systems

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Summary. Swarm-based systems are biology-inspired systems which can be directly mapped to multi-agent systems (MAS), possessing characteristics such as local control over the decisions taken by the agents and a highly dynamic structure which continuously changes. This class of MAS is of a particular interest because it exhibits emergent behaviour through self-organisation and finds itself applicable to a wide range of domains. In this paper, we present OPERAS, an open formal framework that facilitates modelling of MAS, we describe how a particular instance of this framework, namely $OPERAS_{CC}$, could employ existing biological computation systems, such as Population P Systems, and demonstrate how the resulting method can be used to formally model a swarm-based system of autonomous spacecrafts.

1 Introduction

Lately, there has been an increasing interest toward biological and biology-inspired systems. From the smallest living elements, the cells, and how they form tissues in organisms to entire ecosystems and how they evolve, there is growing investigation on ways of specifying such systems. The intention is to create software that mimics the behaviour of their biological counterparts. Examples of biological systems of interest also include insect colonies (of ants, termites, bees etc.), flocks of birds, tumours growth—the list is endless. The understanding of how nature deals with various situations has inspired a number of problem solving techniques [13] that are applicable to a wide range of situations that had been puzzling computer scientists for decades. Swarm Intelligence [15, 16], Ant Colony Optimisation techniques [10] for example, has been successfully applied to robotics [11], network routing [8, 28] and data mining [1] and has inspired agent-based modelling platforms [19].

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The promising feature is that these systems can be directly mapped to multiagent systems (MAS) by considering each entity as an agent, with its own behavioural rules, knowledge, decision making mechanisms and means of communication with the other entities and with the environment. The overall system's behaviour is merely the result of the agents' individual actions, the interactions among them and between them and the environment. This also points to the issue of selforganisation and how collective behavioural patterns emerge as a consequence of individuals' local interactions in the lack of knowledge of the entire environment or global control.

An additional modelling key aspect of MAS has not received much attention so far; it is the dynamic nature of MAS and how their structure is constantly reconfigured. By structure we imply (i) the changing number of agents in a MAS, and (ii) either their physical placement in the environment or, more generally, the structure that is dictated by the communication channels among them. Most modelling methodologies assume a fixed, static structure that is not realistic since in a dynamic MAS, communication between two agents may need to be established or ceased at any point and also new agents may appear in the system while existing ones may be removed. One additional issue that the inherent dynamic nature of these systems raises has to do with distinguishing between the modelling of the individual agents (behaviour) and the rules that govern the communication and evolution of the collective MAS (control). By 'control' we do not imply central control, as this would cancel any notion of self-organisation. Rather, we refer to the part of the agent that takes care of non-behavioural issues. A modelling method that allows such a distinction, would greatly assist the modeller by breaking down the work into two separate and independent activities, modelling the behaviour and modelling the control.

Population P Systems with active membranes [4], a class of variants of P Systems [20] are membrane structures composed of membranes configured in an arbitrary graph and naturally possess the trait of reconfiguring their own structure through rules that restructure the graph and allow membranes to divide and die. Inspired by this appealing characteristic, in this paper we present a formal framework, called *OPERAS*, that facilitates the development of dynamic MAS of the nature of many biology and biology-inspired systems. The next section introduces *OPERAS* formal definition, while section 3 presents an instance of this framework, namely *OPERAS_{CC}* which utilises Population P Systems in order to model MAS. A brief description of a representative case study dealing with a swarm-based system follows in Section 4 which also deals with the formal model for the case problem in question. Finally, Section 5 discusses issues arising from our attempt and concludes the paper.

2 OPERAS: Formal Modelling of MAS

In an attempt to formally model each individual agent as well as the dynamic behaviour of the overall system, we need a formal method that is capable of rigorously describing all the essential aspects, i.e. knowledge, behaviour, communication and dynamics. It is also important that the level of abstraction imposed by a formal method is appropriate enough to lead toward the implementation of a system. New computation approaches as well as programming paradigms inspired by biological processes in living cells, introduce concurrency as well as neatly tackle the dynamic structure of multi-component systems (P Systems, Brane Calculus, Gamma, Cham, MGS) [2, 5, 20]. In agent-oriented software engineering, there have been several attempts to use formal methods, each one focusing on different aspects of agent systems development [3, 6, 9, 12, 21]. Other formal methods, such as π -calculus, mobile ambients and P Systems with mobile membranes [7, 17, 18], successfully deal with the dynamic nature of systems and concurrency of processes but lack intuitiveness when it comes to the modelling of an individual agent (lack of primitives and more complex data structures). An interesting comparison of various formal methods for the verification of emergent behaviours in swarm-based systems is reported in [22].

2.1 OPERAS Definition

We start this section by providing the definition of a model for a dynamic MAS in its general form.

A Multi-Agent System can be defined by the tuple (O, P, E, R, A, S) containing:

- a set of reconfiguration rules, *O*, that define how the system structure evolves by applying appropriate reconfiguration operators;
- a set of percepts, P, for the agents;
- the environment's model / initial configuration, E;
- a relation, R, that defines the existing communication channels;
- a set of participating agents, A, and
- a set of definitions of types of agents, S, that may be present in the system.

More particularly:

- the rules in O are of the form *condition* ⇒ *action* where *condition* refers to the computational state of agents and *action* involves the application of one or more of the operators that create/remove a communication channel between agents or introduce/remove an agent into/from the system;
- *P* is the distributed union of the sets of percepts of all participating agents;
- $R: A \times A$ with $(A_i, A_j) \in R$, $A_i, A_j \in A$ meaning that agent A_i may send messages to agent A_j ;
- $A = \{A_1, \ldots, A_n\}$ where A_i is a particular agent defined in terms of its individual behaviour and its local mechanism for controlling reconfiguration;
- $S_k = (Behaviour_k, Control_k) \in S, k \in Types$ where Types is the set of identifiers of the types of agents, $Behaviour_k$ is the part of the agent that deals with its individual behaviour and $Control_k$ is the local mechanism for control-ling reconfiguration; each participating agent A_i of type k in A is a particular instance of a type of agent: $A_i = (Beh_k, Ctrl_k)_i$.

2.2 OPERAS as an open framework

The general underlying idea is that an agent model consists of two parts, its behaviour and its control. The behaviour of an agent can be modelled by a formal method with its computation being driven by percepts from the environment. The control can be modelled by a set of reconfiguration rules which given the computation states of agents can change the structure of the system. The MAS structure is determined through the relation that defines the communication between the agents. The set of participating agents are instances of agent types that may participate in the system. This deals with the fact that an agent may be present at one instance of the system but disappear at another or that a new agent comes into play during the evolution of the MAS. This assumes that all agent types that may participate in the system should be known in advance.

There are still some open issues which, however, make the *OPERAS* approach a framework rather than a formal method. These are: (i) Which are the formal methods that can be used in order to model the behaviour? (ii) Which are the formal methods that can to use in order to model the control? (iii) Could the methods in (i) and (ii) be different? (iv) Should the agents' behaviour models communicate directly with other agents' behaviour models? (v) Should the agents' control models communicate with other agents' control models? (vi) Could communication be established implicitly through percepts of the environment? (vii) Which method chosen from (i) or from (ii) drives the computation of the resulting system? There is no unique answer to these questions but the modelling solution will depend on the choice of formal methods which are considered suitable to model either behaviour or control.

It is therefore implied that there are several options which could instantiate OPERAS into concrete modelling methods. Regarding the modelling of each type of agent S_k , there are more than one options to choose from in order to specify its behavioural part and the same applies for its control mechanism. We have long experimented with various formal methods, such as X-machines with its communicating counterpart and Population P Systems with active cells. In this paper we present an instance of the framework that employs ideas from the latter, using a PPS to model both the behaviour as well as the control part of the agent.

$3 OPERAS_{CC}$

3.1 Population P Systems with active cells

A Population P System (PPS) [4] is a collection of different types of cells evolving according to specific rules and capable of exchanging biological / chemical substances with their neighbouring cells (Fig. 1). More formally, a PPS with active cells [4] is defined as a construct $\mathcal{P} = (V, K, \gamma, \alpha, w_E, C_1, C_2, \ldots, C_n, R)$ where:

• V is a finite alphabet of symbols called objects;

- *K* is a finite alphabet of symbols, which define different types of cells;
- $\gamma = (\{1, 2, \dots, n\}, A)$, with $A \subseteq \{\{i, j\} | 1 \le i \ne j \le n\}$, is a finite undirected graph;
- α is a finite set of bond-making rules of the form $(t, x_1; x_2, p)$, with $x_1, x_2 \in V^*$, and $t, p \in K$ meaning that in the presence of objects x_1 and x_2 inside two cells of type t and p respectively, a bond is created between the two cells;
- $w_E \in V^*$ is a finite multi-set of objects initially assigned to the environment;
- $C_i = (w_i, t_i)$, for each $1 \le i \le n$, with $w_i \in V^*$ a finite multi-set of objects, and $t_i \in K$ the type of cell i;
- *R* is a finite set of rules dealing with communication, object transformation, cell differentiation, cell division and cell death.



Fig. 1. An abstract example of a Population P System; C_i : cells, R_i : sets of rules related to cells; w_i : multi-sets of objects associated to the cells.

All rules present in the PPS are identified by a unique identifier, r. More particularly:

Communication rules are of the form $r : (a; b, in)_t, r : (a; b, enter)_t, r : (b, exit)_t$, for $a \in V \cup \{\lambda\}, b \in V, t \in K$, where λ is the empty string, and allow the moving of objects between neighbouring cells or a cell and the environment according to the cell type and the existing bonds among the cells. The first rule means that in the presence of an object a inside a cell of type t an object b can be obtained by a neighbouring cell non-deterministically chosen. The second rule is similar to the first with the exception that object b is not obtained by a neighbouring cell but by the environment. Lastly, the third rule denotes that if object b is present it can be expelled out to the environment.

Transformation rules are of the form $r : (a \to b)_t$, for $a \in V$, $b \in V^+$, $t \in K$, where V^+ is the set of non-empty strings over V, meaning that an object a is replaced by an object b within a cell of type t.

Cell differentiation rules are of the form $r : (a)_t \to (b)_p$, with $a, b \in V$, $t, p \in K$ meaning that consumption of an object a inside a cell of type t changes the cell, making it become of type p. All existing objects remain the same besides a which is replaced by b. 556 I. Stamatopoulou, P. Kefalas, M. Gheorghe

Cell division rules are of the form $r: (a)_t \to (b)_t (c)_t$, with $a, b, c \in V, t \in K$. A cell of type t containing an object a is divided into two cells of the same type. One of the new cell has a replaced by b while the other by c. All other objects of the originating cell appear in both new cells.

Cell death rules are of the form $r : (a)_t \to \dagger$, with $a \in V, t \in K$ meaning that an object a inside a cell of type t causes the removal of the cell from the system.

PPS provide a straightforward way for dealing with the change of a system's structure and this is the reason why we have chosen them to define an instance of the OPERAS framework, namely $OPERAS_{CC}$.

3.2 Definition of $OPERAS_{CC}$

In $OPERAS_{CC}$, each agent (individual behaviour) is modelled as a PPS cell, and has a membrane wrapped around it, that is responsible for taking care of structure reconfiguration issues (control). In essence, this may be considered as a usual Population P System in which each cell is virtually divided in two regions, inner (for behaviour) and outer (for control), that deal with different sets of objects and have different kinds of rules that may be applied to them. An abstract example of an $OPERAS_{CC}$ model consisting of two agents is depicted in Fig. 2.

Additionally, when using a PPS for modelling purposes, we consider all objects to be attribute-value pairs of the form att : v so that it is clear to which characteristic of the agent an object corresponds to.



Fig. 2. An abstract example of a $OPERAS_{CC}$ consisting of two agents.

A MAS in $OPERAS_{CC}$ is defined as the tuple $(O, P, E, R, (A_1, \ldots, A_n), S)$ (in correspondence to $P = (R, V, w_E, \gamma, (C_1, \ldots, C_n), k)$ of a PPS) where:

- $A_i = (w_{beh}, w_{ctrl}, t), w_{beh}$ being the objects of the agent behaviour cell, w_{ant} the objects of the control cell (these objects possibly hold information about the w_{beh} objects (computation states) of neighbouring agent cells) and $t \in k$ the type of the cell;
- $O = O_A \cup O_C$.
 - The rules in O_A (to be applied only by the behaviour cells on the w_{beh} objects) are the transformation rules of a PPS that rewrite the objects, as well as the communications rules that move objects between cells that are linked with a bond (both kinds of rules do not affect the structure of the system).
 - The rules in O_C (to be applied only by the control cells on the w_{ctrl} objects) are the birth, death, differentiation and bond-making rules of a PPS (the kinds of rules that affect the structure of the system) as well as environment communication rules (receiving/sending objects from/to the environment) so that there is indirect communication between the control cells.
- $P = P_A \cup P_C$, the set of percepts of all participating agents where P_A is the set of inputs perceived by the behaviour cells and P_C is the set of inputs perceived by the control cells.
- E is the set of objects assigned to the environment holding information about the computation states of all the participating agents;
- R is the finite undirected graph that defines the communication links between the behaviour cells;
- S is the set of possible types of cells.

It should be noted that although the agent descriptions' set A appears fifth in *OPERAS* definition tuple, from a practical perspective it is the first element being defined; the other tuple elements and their form are naturally dependent on the particular method(s) chosen to define the behavioural and control part of the agents.

Computation

In every computation cycle:

- In all the cells modelling the behaviour of the agent, all applicable object rules in O_A (transformation and communication) are applied;
- All control cells expel in the environment the w_{beh} objects (computation states of behaviour cells) along with the cell identity;
- All control cells import the computation states, w_{beh} , of neighbouring agents;
- All rules in O_C (bond-making, birth, death, differentiation) are triggered in the control cells, (if applicable) reconfiguring the structure of the system.

Since the model follows the computation rules of a PPS system, the overall system's computation is synchronous. Asynchronous computation may be achieved with the use of other methods for modelling the agents' behaviour and/or control. In [27] we present another instance of the framework, namely $OPERAS_{XC}$, which uses X-machines for the behavioural part of the agent and membranes wrapped around the machines for the control part, and apply it on the same swarm-based

system that we present hereafter. Because in that version of the framework computation is driven by the computation of the participating X-machines, overall computation is asynchronous.

4 $OPERAS_{CC}$ for a Swarm-based system

4.1 Autonomous Spacecrafts for Asteroid Exploration

A representative example of a system which clearly possesses all the aforementioned characteristics of a dynamic MAS is the NASA Autonomous Nano-Technology Swarm (ANTS) system [22]. The NASA ANTS project aims at the development of a mission for the exploration of space asteroids with the use of different kinds of unmanned spacecrafts. Though each spacecraft can be considered as an autonomous agent, the successful exploration of an asteroid depends on the overall behaviour of the entire mission, as the latter emerges as a result of self-organisation. We chose this case study because relevant work on the particular project included research on and comparison of a number of formal methods [22, 23].



Fig. 3. An instance of the ANTS mission, L: Leader, W: Worker, M: Messenger.

The ANTS mission uses of three kinds of unmanned spacecrafts: L_i , leaders (or rulers or coordinators), W_i , workers and M_i , messengers (Fig. 3). The leaders are the spacecrafts that are aware of the goals of the mission and have a non-complete model of the environment. Their role is to coordinate the actions of the spacecrafts that are under their command but by no means should they be considered to be a central controlling mechanism as all spacecrafts' behaviour is autonomous. Depending on its goals, a leader creates a team consisting of a number of workers and at least one messengers. Workers and messengers are assigned to a leader upon request by (i) another leader, if they are not necessary for the fulfillment of its goals, or (ii) earth (if existing spacecrafts are not sufficient in number to cover current needs, new spacecrafts are allocated to the mission). A worker is a spacecraft with a specialised instrument able, upon request from its leader, to take measurements from an asteroid while flying by it. It also possesses a mechanism for analysing the gathered data and sending the analysis results back to its leader in order for them to be evaluated. This in turn might update the view of the leader, i.e. its model of the environment, as well as its future goals.

The messengers, finally, are the spacecrafts that coordinate communication among workers, leaders and the control centre on earth. While each messenger is under the command of one leader, it may also assist in the communication of other leaders if its positioning allows it and conditions demand it.

What applies to all types of spacecrafts is that in the case that there is a malfunctioning problem, their superiors are being notified. If the damage is irreparable they need to abort the mission while on the opposite case they may "heal" and return back to normal operation.

4.2 The $OPERAS_{CC}$ approach to the ANTS mission

The swarm-based system in the ANTS mission can be directly mapped into the OPERAS framework (Fig. 4). A number of agents of three different types (workers, W, leaders, L, and messengers, M) compose the MAS system. System configuration is highly dynamic due to its nature and unforeseen situations that may come up during the mission.



Fig. 4. (a) An instance of MAS structure corresponding to ANTS in Fig. 3 with an *OPERAS* agent (W_4) consisting of separate Behaviour and Control components. (b) A change in the structure of MAS after possible events (e.g. destruction of worker W_2 , leader L_1 employs worker W_6 etc.).

Leader: Formal Modelling of Behaviour in $OPERAS_{CC}$

For the modelling of the leader agent, one has to identify the internal states of the agent, its knowledge as well as the inputs it is capable of perceiving, so that they are represented as objects of the corresponding PPS.

The state of a leader can be either one of the three: *Processing* for an leader that is fully operational, *Malfunctioning* for one that its facing problems and *Aborting* for one that is either facing irreparable problems or has been commanded by the control centre on earth to abort the mission.

Object	Description
status	The current operational state of the leader
existingWorkers	The set of IDs and statuses of the workers under its com-
	mand
existingMsgs	The set of IDs and statuses of the messengers under its
	command
results	The set containing analysis results it has gathered
model	The current model of the agent's surroundings
goals	The agent's goals

 Table 1. Objects representing the knowledge of the Leader agent.

The knowledge of the agent consists of the objects presented in Table 1 along with their description.

Similarly, the leader type of cell will be able to also perceive other objects representing input from the environment or from other agents. The most prominent ones are summarised in Table 2.

Object	Description
abrt	A request from the control centre that the agent
	should abort the mission
worker	A new worker that joins the team under the leaders
	command
messenger	A new messenger that joins the team under the lead-
	ers command
requestForWorker	A request for a worker, made by another leader (so
	that the worker is reallocated)
requestForMsg	A request for a messenger, made by another leader
	(so that the messenger is reallocated)
message	An object representing a message sent by another
	agent

Table 2. Objects representing the percepts of the Leader agent.

Indicatively, two of the operations that a leader may perform in the form of transformation rules follow.

The rule representing the joining of a worker w_i to the leader's team of *Workers* is specified as:

workerJoining:

 $(status: processing worker: w_i existing Workers: Workers$

 \rightarrow status : processing existing Workers : $\{w_i\} \cup Workers)_L$

The newly allocated worker w_i may be received by another leader with the use of a communication rule of the form:

 $receiveWorker: (message: canSendYouAWorker; worker: w_i, in)_L$

which assumes that a *canSendYouAWorker* message has been previously sent by the other leader informing that it is willing to reallocate one of its workers.

Similarly, the rule representing the reallocation of the messenger m_i to another leader is:

 $\begin{aligned} reAllocatingMessenger: \\ (status: processing \ percept: requestForMsg \ existingMsgs: Messengers \\ & \rightarrow status: processing \ existingMsgs: Messengers \ \{m_i\})_L, \\ & \text{if } isMessengerNeeded(m_i) == false \end{aligned}$

Worker: Formal Modelling of Behaviour in $OPERAS_{CC}$

Similarly for a worker agent, the internal states in which a it may be in are *Measuring*, when taking measurements from an asteroid, *Analysing*, when analysing the measurements in order to send results to its leader, *Idle*, *Malfunctioning* and *Aborting*.

The knowledge of the agent consists of the objects presented in Table 3 along with their description.

Object	Description
status	The current operational state of the worker
myLeader	the identity of its commanding leader,
teamWorkers	The set of other coworkers belonging to the same team
teamMsgs	The set of messengers belonging to the same team
Target	The target asteroid
Data	The set of data collected from the asteroid
Results	The set of the data analysis results

Table 3. Objects representing the knowledge of the Worker agent.

The worker type of cell will also be able to also perceive other objects that represent either environmental stimuli or messages from other agents. Indicative ones are being summarised in Table 4. 562 I. Stamatopoulou, P. Kefalas, M. Gheorghe

Indicatively, some of the operations that a worker may perform in the form of transformation rules follow.

The rule representing the measurements' analysis mechanism of the worker is: analysingData :

 $(status : Analysing \ data : Data \rightarrow status : Idle \ results : Results)_W$

The rule that informs a worker that it is being reallocated to another leader is defined as:

reAllocating:

```
(status : Idle myLeader : Leader reassignedTo : NewLeader 

ightarrow status : Idle myLeader : NewLeader)_W
```

Object	Description
abrt	A request from the control centre that the agent should abort the
	mission
reassigned To	The identifier of the new leader the worker is being reassigned to
data	The set of measurements taken from the asteroid

Table 4. Objects representing the percepts of the Worker agent.

4.3 Formal Modelling of Control in OPERAS_{CC}

According to $OPERAS_{CC}$, for the definition of the given system as a dynamic MAS, we need to assume an initial configuration. To keep the size restricted for demonstrative purposes, let us consider an initial configuration that includes one leader L_1 , one messenger M_1 and two workers W_1, W_2 . According to $OPERAS_{CC}$ the above system would be defined as follows.

The set O contains all the aforementioned transformation rules that model the agents' behaviour as well as the reconfiguration rules (birth, death and bondmaking) regarding (i) the generation of a new worker when the control centre on earth decides it should join the mission, (ii) the destruction (i.e. removal from the system) of any kind of agent in the case it must abort the mission, (iii) the establishment of a communication channel between a leader and all members of its team. More particularly O additionally contains the following rules.

The following birth rules create a new worker w_i or messenger m_i under the command of a leader L_i when the leader has received the corresponding messages (objects *earthSendsWorker* and *earthSendsMsg*) from the control centre on earth.

 $\begin{array}{l} newWorkerFromEarth: \\ (status: Processing \ earthSendsWorker: w_i \ existingWorkers: Workers)_{L_i} \\ \rightarrow (status: Processing \ existingWorkers: Workers \cup \{w_i\})_{L_i} \\ (status: Idle \ myLeader: L_i)_{W_i} \end{array}$

newMessengerFromEarth:

 $(status : Processing \ earthSendsMsg : m_i \ existingMsgs : Messengers)_{L_i} \rightarrow (status : Processing \ existingMsgs : Messenger \cup \{m_i\})_{L_i} (status : Idle \ myLeader : L_i)_{M_i}$

Inputs, such as $earthSendsMsg : m_i$, from the environment are perceived with the use of communication rules of the form:

 $receiveInput: (\varepsilon; earthSendsMsg: m_i, enter)_L$

The death rule below removes agent instances that have aborted the mission from the model ($t \in S$ stands for any type of agent).

abortion:

 $(status: aborting)_t \rightarrow \dagger$

Finally, the following bond-making rules ensure the creation of a communication bond between a leader agent (ε stands for the empty multi-set, i.e. no object is necessary) and any messenger or worker that belongs to this leader's team. *workerBondMaking*:

 $(L_i \ \varepsilon \ ; \ myLeader : L_i \ W)$

messengerBondMaking:

 $(L_i \varepsilon; myLeader: L_i M)$

The set P contains all objects recognised by the Population P System.

Regarding the environment E, it should initially contain objects representing the initial percepts for all agents.

Since in the assumed initial configuration we consider to have one group of spacecrafts under the command of one leader, all agents should be in communication with all others and so:

 $R = \{ (L_1, W_1), (L_1, W_2), (W_1, W_2), (M_1, L_1), (M_1, W_1), (M_1, W_2) \}$ Finally, the set S that contains the agent types is: $S = \{L, W, M\}.$

5 Conclusions and Further Work

We presented *OPERAS* with which one can model multi-agent systems that exhibit dynamic structure, emergent and self-organisation behaviour. The contributions of *OPERAS* can be summarised in the following:

- A formal framework for MAS modelling.
- The behaviour and the control of an agent are separate components which imply distinct modelling mental activities.
- Flexibility on the choice of formal methods to utilise and option to combine different formal methods.

It is because of this distinct separation between behaviour and control that *OPERAS* provides this flexibility of choosing different methods for modelling

these two aspects; while some methods are better at capturing the internal states, knowledge and actions of an agent, others focusing on the dynamic aspect of a MAS are more suitable for capturing the control mechanisms.

In this paper, we employed Population P Systems with active cells to define $OPERAS_{CC}$, an instance of the general framework. We presented the $OPERAS_{CC}$ model of a swarm-based system of a number of autonomous spacecrafts. It could easily be spotted that an $OPERAS_{CC}$ model does resemble (as a final outcome) a model which could be developed if one used Population P Systems with active cells from scratch [24]. However, in the current context we have the following advantages:

- PPS can be viewed as a special case for *OPERAS*.
- The distinction of modelling behaviour and control as separate, offers the ability to deal with transformation/communication rules separately from cell birth/division/death and bond making, with implications both at theoretical as well as practical level.
- Practically, during the modelling phase, one can find advantages and drawbacks at any of the behaviour or control component and switch to another formal method for this component if this is desirable.

As far as the last point is concerned, we verified our initial findings [24] in which it was stated that modelling the behaviour of an agent with PPS rewrite and communication rules may be rather cumbersome. Especially in this rather complex case study, although the modelling of the control is absolutely straightforward, we had difficulties to establish the necessary peer to peer communication between agents by employing just the communication rules. That gave us the opportunity to consider alternatives. For example, we have experimented with Communicating Xmachines which have a number of advantages in terms of modelling the behaviour of an agent. The resulting model, $OPERAS_{XC}$ [27], seems to ease the modelling process in complex MAS. It is worth noticing that none of the two formal methods (X-machines and Population P Systems) by itself could successfully (or at least intuitively) model a MAS [14, 26]. This is true for other formal methods too, which means the current framework gives the opportunity to combine those methods that are best suited to either of the two modelling tasks.

We would like to continue the investigation of how *OPERAS* could employ other formal methods that might be suitable for this purpose. In the near future, we will focus on theoretical aspects of the framework. Towards this direction, we are also currently working on various types of transformations that could prove its power for formal modelling as well as address legacy issues concerned with correctness.

Finally, efforts will also be directed towards enhancing existing animation tools on Population P Systems in order to come up with a new version of the tool that will be able animate $OPERAS_{CC}$ specified models. More particularly, the PPS-System [25] is a tool that generates Prolog executable code from Population P Systems models written in a particular notation. Future work will involve extending the notation and the system in order to integrate the necessary OPERAS features, allowing us to gain a deeper understanding of the modelling issues involved with $OPERAS_{CC}$ and helping us investigate the practicability of our approach.

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Algorithm of Rules Application Based on Competitiveness of Evolution Rules

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Summary. Transition P systems are a parallel computational model based on the notion of the cellular membrane system. The distributed architectures for P system implementation in digital device must deal jointly with the time for active rules application over multisets and the time for communication among membranes. Analysis of these architectures shows that it is very important to improve the time used in the rules application step since this allows for reducing the evolution step time and the number of processors needed in the system. This work introduces the concepts of competitiveness relationship among active rules and competitiveness graph. For this, it takes into account the fact that some active rules in a membrane can consume disjointed object sets. Based on these concepts this paper presents a new evolution rules application algorithm that improves throughput of active rules elimination algorithms (sequential and parallel).

1 Introduction

Computation with membranes was introduced by Gheorghe Păun in 1998 [10] through a definition of transition P systems. This new computational paradigm is based on the observation of biochemical processes. The region defined by a membrane contains chemical elements (multisets) which are subject to chemical reactions (evolution rules) to produce other elements. Transition P systems are hierarchical, as the region defined by a membrane may contain other membranes. Multisets generated by evolution rules can be moved towards adjacent membranes (parent and children). This multiset transfer feeds back into the system so that new products are consumed by further chemical reactions in the membranes.

These systems perform computations through transition between two consecutive configurations. Each transition or evolution step goes through two sequential steps: rules application and objects communication. First, the evolution rules are applied simultaneously to the multiset in each membrane. This process is performed by all membranes at the same time. Then, also simultaneously, all membranes communicate with their destinations. The objective of this paper is to present an improvement of the algorithm of active rules elimination [12] used in the rules application step. To achieve this, the paper is structured as follows: first, related works are presented; then, the basic ideas of the active rules elimination algorithm are summarized, which is followed by a definition of the concept of competition between rules; next, the optimization of the algorithm is specified and, finally, conclusions are drawn.

2 Related works

No line of research for the implementation of P system in digital devices has managed to achieve the massively parallel nature of these systems. However, significant advances have been made. First, specifications have been made of faster algorithms for application of evolutions rules for both sequential devices [12] [3] and parallel ones [4] [5]. Moreover, the problem of network congestion in the multiset communications step described by Ciobanu [2] has been solved in the works of Tejedor [11] and Bravo [1]. In short, today a number of implementations can be made that achieve a certain degree of parallelism that depends on communications and the application of evolution rules.

Viable communications architectures for the P systems implementation proposed in [11] and [1] use a number of processors on the order of the square root of the number of membranes and place several membranes in each processor. Moreover, these solutions achieve an evolution pass time, at worst, on the order of the square root of the number of membranes and eliminate network congestion by means of access control to the common medium. However, these architectures need to know the time needed to perform rules application. This information is needed to be able to optimally distribute membranes among processors. In these P system implementation architectures, it is important to improve the time used in the rules application step since this allows for reducing the evolution step time and the number of processors needed in the system. Specifically, if rule application time is divided by the factor N, then the evolution time and the number of processors in the system is divided by the square root of N.

Analysis of the rules application algorithms published to date shows that only the algorithm of active rules elimination [12] together with its parallel version [5] meet the viable architectures of Tejedor [11] and Bravo [1]. These 2 algorithms enable prior determination of the maximum execution time, since this value depends on the number of rules rather than on the multiset cardinal to which they are applied, as in other algorithms reported. [2] [3]. In addition, these algorithms are the quickest in their category (sequential and parallel). This paper improves throughput of active rules elimination algorithms and takes into account the fact that some active rules in a membrane can consume disjointed object sets. This improvement is applicable to the algorithm in both its sequential and parallel versions.

3 Algorithm of active rules elimination [12]

The general idea of this algorithm is to eliminate, one by one, the rules from the set of active rules. Each step of rule elimination performs two consecutive actions:

- 1. Iteratively, any rule other than that which is to be eliminated is applied for a randomly selected number of times in an interval from 0 to the maximum applicability benchmark. This action ensures the non-determinism inherent to P systems.
- 2. The rule to be eliminated is applied a number of times which is equal to its maximum applicability benchmark, thus making applicable no longer and resulting in its disappearance from the set of active rules.

We assume that:

- 1. The object multiset to which active rules are applied is ω .
- 2. The active rules set is transformed in an indexed sequence R in which the order of rules is not relevant.
- 3. The object multiset resulting from application of active rules is ω' .
- 4. The multiset of applied rules that constitute the algorithm output is ω_R .
- 5. Operation |R| determines the number of rules in the indexed sequence R.
- 6. Operation $\Delta_{R[Ind]} [\omega']$ calculates the maximum applicability benckmark of the rule R[Ind] over ω' .
- 7. The operation $input (R [Ind]) \cdot K$ performs the scalar product of the antecedent of rules by a natural number.

and thus the algorithm is:

Remember that if rule R[i] is no longer applicable in the elimination step for R[j], it is no longer necessary to perform the elimination step for R[i], the algorithm is greatly improved, as shown in [12].

In each iteration of the algorithm of actives rules elimination, the maximum applicability benchmark of a rule is calculated and then the rule is applied. The number of iterations executed at worst is:

$$\#iterations = \sum_{i=1}^q i = \frac{q \cdot (q+1)}{2}$$

Let q be the cardinal of the indexed sequence of active rules. Thus, this algorithm allows one to know how long it takes to be executed in the worst case, with knowledge of the rules set of a membrane.

It is important to note that, in general, it is essential to perform the first action in each elimination step of a rule. This action is necessary to ensure that any possible result of the rules application to the multiset is produced by the algorithm. In case the action is not performed, the eliminated rule (applied as many times as the value of its maximum applicability benchmark) may consume the objects necessary so that any other rule can be applied. However, the latter does not always occur and the first action in each elimination step can be simplified. For the sake of illustration, let us assume that the antecedents of a set of active rules are shown in figure 1.

 $\begin{array}{ll} input(r_1) = a & input(r_2) = ab \\ input(r_3) = cd & input(r_4) = d \end{array}$

Fig. 1. Antecedents of a set of active rules

In this case, in the elimination step of the rule r_1 only the first action with the rule r_2 has to be taken, as r_1 and r_2 are the only rules with the object a in their antecedents. The same is the case with rules r_3 and r_4 , as these two compete for the object d. Thus, taking into account the competition between rule antecedents, one can adjust the rule elimination algorithm to perform only 6 iterations in the worst case, rather than 10 (2 to eliminate r_1 , 1 to eliminate r_2 , 2 to eliminate r_3 , 1 to eliminate r_4) as shown in figure 2.



Fig. 2. Execution trace of Rules Elimination and Rules Elimination with competitiveness algorithms

4 Definition of competitiveness between rules

Definition 1: Competitiveness graph.

Let R be a set of active rules

$$R = \{r_1, r_2, ..., r_q\}$$
 with $q > 0$

Let C be a binary relationship defined over the set R

$$\forall x, y \in R, \ x \neq y \quad x \ C \ y \ \Leftrightarrow \ input(x) \cap input(y) \neq \emptyset$$

This binary relationship can be represented by a non-directed graph CG = (R, C) called a competitiveness graph, where the rules are related to each other if, and only if its antecedents have an object in common. For example, given the rules inputs shown in figure 3, the competitiveness graph generated by these rules taking into account the relationship C will be as shown in figure 4.

Definition 2: Subgraph resulting from elimination of a rule. Let competitiveness graph be CG = (R, C) and rule $x \in R$. The subgraph resulting from elimination of rule x is defined as:

$$CSG = (R - \{x\}, C \cap R - \{x\} \times R - \{x\})$$

Definition 3: Induced subgraph. Let a competitiveness graph be CG = (R, C) and $R' \subseteq R$. The competitiveness subgraph induced by the subset R' is the graph:







Fig. 4. Competitiveness graph

$$CSG = (R', C \cap R' \times R')$$

Definition 4: Competitiveness chain. For a competitiveness graph CG = (R, C), a competitiveness chain is defined as an ordered sequence of rules pertaining to R

$$s_1, s_2, \ldots, s_n \quad s_i \in \mathbb{R},$$

satisfying:

$$s_i C s_{i+1} \quad \forall i \in \{1, ..., n-1\}$$

By definition, there is always a competitiveness chain composed of a single rule. **Definition 5:** Accessible rule relationship. For a competitiveness graph CG = (R, C), the accessible rule relationship (A) is defined as:

 $x, y \in R$ $x \land y \Leftrightarrow \exists$ a competitiveness chain $s_1, ..., s_n | s_1 = x \land s_n = y$

This is an equivalence relation which divides the rule set R into equivalence classes.

Definition 6: Connected component of competitiveness graph. Let competitiveness graph be CG = (R, C), let the accessible rule relationship be A and let E be an equivalence class produced by A. The connected component of CG is defined as the graph induced by the vertices pertaining to the equivalence class E.

Definition 7: Connected competitiveness graph. Let a competitiveness graph be CG = (R, C) and consider the its rule accessibility relation. We call connected CG if and only if it has a connected component.

Definition 8: Articulation. For a competitiveness graph CG = (R, C) and a rule $x \in R$, it is said that x is an articulation of CG if and only if the subgraph resulting from the elimination of rule x has more connected components than CG.

5 Algorithm based on rules competitiveness

Based on the rules competitiveness relationship of a membrane's rules one can improve the algorithm of elimination of active rules. To do this, an analysis must be made of the evolution rules of each membrane prior to P system evolution. The analysis will determine the order of active rules elimination and what rules set are used in the first action of each elimination step of a given rule. The following optimizations can be made of the algorithm of rule elimination:

5.1 1^{st} optimization

The idea of this optimization is based on the fact that in the elimination step of a rule, the first action of the algorithm must be applied to the rules in the same connected component of the competitiveness graph. This can be done because the antecedents of rules in different connected components do not compete for common objects of the multiset.

The analysis prior to the execution of each P system calculates the competitiveness graph of each membrane. Then the connected components of the graph are calculated. The algorithm of active rule elimination will be applied independently to the rules of each of the connected components, with no need for any change in its codification.

In the worse case of the example of figure 4, the sequential version of this algorithm will need to perform 3 iterations in the connected component consisting of the rules $\{r_1, r_2\}$ and 36 iterations in the connected component consisting of

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the rules $\{r_3, r_4, r_5, r_6, r_7, r_8, r_9, r_{10}\}$. Therefore, this example has gone from 55 iterations in the worst case of the algorithm of active rules elimination to 39 iterations (figure 5), that is, it has been reduced by 71% compared to the active rules elimination algorithm.



Fig. 5. Execution trace of sequential 1^{st} optimization

Making a parallel version of the algorithm is quite simple. One need only apply the algorithm of active rules elimination in parallel to the rules of each connected component on the competitiveness graph. The parallel version would require only 36 iterations (maximum(36,3)) in the worst case, as shown in figure 6), that is, it has been reduced by 65% compared to the active rules elimination algorithm.

5.2 2^{nd} optimization

This optimization is applied in each connected component of the competitiveness graph. If the competitiveness graph of a membrane has articulations the algorithm can be used to eliminate these rules first and cause the appearance of new connected components. Thus, if rule r_6 is eliminated in our example (figure 4) the connected component splits in two: the one composed of $\{r_3, r_4, r_5\}$ and the one composed of $\{r_7, r_8, r_9, r_{10}\}$.



Fig. 6. Execution trace of parallel 1^{st} optimization

When a connected component has no articulations, elimination of more than one rule can break it into more than one connected component. Continuing with the example we have proposed, if we first remove from connected component $\{r_7, r_8, r_9, r_{10}\}$ rules r_7 and r_{10} in two elimination steps, it then splits into two connected components consisting of the rules r_8 and r_9 , respectively.

To perform this optimization, a slight change must be made in the sequential algorithm of active rules elimination. Now, each step of elimination of a rule must eliminate a specific rule and is carried out in a sequence of determinate rules. Moreover, there is a certain partial order in the elimination steps of a rule. Whereas order is irrelevant in previous versions of the active rules elimination algorithm, it is decisive in this version. The set of rules used and the rule being eliminated in each elimination step is calculated for each membrane in the analysis prior to the evolution of the P system; as a result, the calculation does not penalize the execution time of the algorithm.

Figure 7 shows the order in which evolution rules are eliminated and the set of rules used in each elimination step for the example in figure 4. The number of iterations of this algorithm in the worst case is 25, that is, it has been reduced by 45% compared to the active rules elimination algorithm.

The parallel version of the algorithm involves applying the sequential version to each of the connected components that are either in the original competitiveness graph or that are generated as a result of the elimination of a rule.

The execution trace of the parallel algorithm used with the set of rules of the example in figure 4 is shown in figure 8. It may be noted that the number of iterations in the worst case is 16 (maximum(8, 2) + maximum(3, 4, 1) + maximum(1, 1, 3) + maximum(1, 1, 3))



Fig. 7. Execution trace of sequential 2^{nd} optimization

maximum(1,1)) using 5 processes. Hence, the number of iterations is reduced by 29% compared to the active rules elimination algorithm.



Fig. 8. Execution trace of parallel 2^{nd} optimization

Table 1 shows the number of iterations performed by the algorithms in the worst case:

- Actives rules elimination (ARE)[12]
- Sequential competitive rules with 2^{nd} optimization(SCR)
- Delimited massively parallel (DMP)[5]
- Parallel competitive rules with 2^{nd} optimization (PCR)

Applied to P systems defined in:

- Reference#1:Computing with Membranes [8]
- Reference#2:A Guide to Membrane Computing [9]
- Reference#3: P Systems Running on a Cluster of Computers [2]

	Sequential			Parallel		
	ARE	SCR	SCR/ARE	DMP	PCR	PCR/DMP
Ref#1 pag 10	6	4	66%	5	3	60%
Ref#1 pag 13	6	4	66%	5	3	60%
Ref#1 pag 15	3	3	100%	3	3	100%
Ref#2 pag 9	6	4	66%	5	3	60%
Ref#2 pag 14	10	6	60%	8	3	37%
Ref#3 pag 137	10	5	50%	8	3	37%
Ref#3 pag 138	6	4	66%	5	3	60%

Table 1. Number of iterations performed by the algorithms in the worst case

5.3 3^{rd} optimization

This last optimization is based on an analysis of the execution trace of the 2^{nd} optimization. It can occasionally be observed that the elimination step of one rule r_j also eliminates one or more additional rules r_i . This can occur either because r_i is applied a number of times that coincides with the maximum applicability benchmark, or the rules applied prior to r_i consume the objects it needed to continue being active. This can be used mainly in three ways to improve the execution time of the algorithm:

- 1. There is no need to execute the elimination step of the rule r_i eliminated in a previous step. Bearing in mind the execution trace in figure 7, if the elimination step of rule r_6 also eliminates rule r_4 , then it would no longer be necessary to execute the elimination step of r_4 , thus allowing execution of the algorithm to save 3 iterations.
- 2. Rule r_i is not to be applied in the elimination steps of subsequent rules. Bearing in mind the execution trace in figure 7, if the elimination step of rule r_6 also eliminates rule r_8 , it is therefore unnecessary in elimination steps of rules r_7

and r_{10} to try to apply r_8 , thus allowing execution of the algorithm to save 2 iterations.

3. Elimination of rule r_i causes a change in the composition and order of the subsequent elimination steps. Keeping in mind the execution trace in figure 7, if the elimination step of rule r_6 also eliminates rule r_8 , then it is beneficial to execution of the algorithm for r_9 to be the next rule eliminated. This is the case because once r_6 , r_8 and r_9 have been eliminated, r_7 and r_{10} can be eliminated in a single iteration in their elimination step since they do not share objects. Here, 3 iterations would be saved.

To implement this optimization, a determination is necessary of what rules continue to be active whenever an elimination step is performed, and this information is used to calculate the next optimal elimination step to be taken. Logically, calculation of the next optimal elimination step would severely penalize the execution time of the algorithm. Hence, a different solution must be sought. This solution involves making an analysis prior to the execution of each P system, in which we can calculate all the possible active rule sets and assign them the next optimal step of rule elimination. All this information would be reflected in a director graph of the algorithm, the definition of which is as follows:

Definition 9: Director graph of algorithm of rule application. Let R be a set of active rules. The director graph of the algorithm of rule application is composed of a triad DG = (Q, A, T) where:

- 1. Q is the node set of the graph, composed of a subset of parts of R, that is: $\forall q \in Q, , q \in \mathcal{P}(R)$
- 2. A is a correspondence whose initial set is Q and whose final set is a set of sequences of rules composed of rules of the origin element of Q. Thus, each set of active rules has one or more sequences of rules. Each sequence of rules indicates the order in which elimination step rules are applied. So a state can have several elimination steps associated in the analysis prior the evolution of each P system.

 $A: Q \to E$ where E is the set of possible sequences with elements in Q

1. T is a set of transitions. Each transition is composed of a triad $\langle q_i, A(q_i), q_f \rangle$ where $q_i, q_f \in Q$ are the initial and final state, respectively, of the transition and $A(q_i)$ are the elimination step (s) of rules associated to state q_i , which, after being executed, means that active rules are those of state q_f .

Execution of the sequential algorithm of application of competitive rules will involve making a loop that ends when it reaches a state with no active rules. In each iteration, there are 3 steps:

- 1. The elimination steps associated to the state are executed.
- 2. Active rules are calculated.
- 3. The state represented by active rules is transited.

Execution of the parallel algorithm of application of competitive rules will be similar to the sequential one. The difference is that execution of several elimination steps associated to a state is performed in parallel way.

In the worse case, 3^{rd} optimization performs the same iterations as the 2^{nd} optimization. However, the experimental data obtained with the execution of the algorithm with 3^{rd} optimization are better than the 2^{nd} one, as show figure 9. In this figure, the X-axis values represent the relationship between the cardinal of the multiset and the cardinal of the sum of inputs of active rules. The Y-axis values are percentage relationship between the number of iterations with active rules elimination algorithm and sequential competitive rules with 3^{rd} optimization algorithm.



Fig. 9. Comparation between Active Rules Elimination Algorithm and Sequential Competitive Rules with 3^{rd} optimization

6 Conclusions

This paper introduces the concept of a competitiveness relationship among active rules. Based on this concept, a new way of parallelism has been opened towards the massively parallel character needed in rules application in P systems. Moreover, the sequential version of this algorithm has a lower number of operations in its execution than in other sequential algorithms published to date.

Both the sequential and the parallel versions of the algorithm perform a limited number of operations, thus allowing for prior knowledge of the execution time. This characteristic makes both versions of the proposed algorithm appropriate for use in viable distributed architectures of implementation of P systems. This is said because architectures require determining the distribution of the number of membranes to be located in each processor of the architecture in order to obtain minimal evolution step times with the use of minimal resources.

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Direct Simulation of the Oregonator Model by Using a Class of P Systems

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Summary. We propose a simple method of simulating chemical reactions by using a multiset rewriting systems, the Abstract Rewriting System on Multisets (ARMS). We simulate the Oregonator model, a mathematical model of the Belousov-Zhabotinsky and We obtain the same behavior when the differential equations are used. We also investigate the Oregonator with large stochastic fluctuations and confirm that they may affect its oscillational behaviors.

We simulated the Oregonator model [2], which is a mathematical model of the Belousov-Zhabotinsky reaction (BZ reaction) [11]. As for the mathematical model of BZ reaction, the brusselator [7] and Oregonator are well known. In the brusselator, each reaction rule does not correspond to the actual chemical reactions exhibiting the BZ reaction, while in the Oregonator each reaction rule corresponds to actual chemical reactions exhibiting the BZ reaction have been obtained by chemical experiments; these parameters of the BZ reaction have been obtained by chemical experiments; these parameters cannot be used for simulation of the burruselator but can be used for the Oregonator. However the Oregonator is not simple and in order to simulate it, usually simplifications of the model by abstracting dimensions of parameters and neglecting variants are required [10]. So, in the simulation of the simplified model, even if we can find interesting behaviors, we cannot feed back to design chemical experiments to confirm them.

Direct methods such as the *Gillespie* method [3] or the *StochSim* [5] allow us to simulate the Oregonator without simplifying and transforming parameters to dimensionless space. the *Abstract Rewriting system on MultiSets* (ARMS) [8], is a direct method and a variant of P Systems, which is closely related to the *Metabolic Algorithm* (MA) (for example [4]) and the Gillespie method.

The main differences between the ARMS and MA are the ARMS is a stochastic model and the rate constants keep the same during computations, while the MA is a deterministic model and the rate constant can change during a computation. And [1] investigated the relationship between P systems and the Gillespie method. The main differences between the ARMS and Gillespie method is that; the ARMS and Gillespie method based on the rate constants and population size of each chemical species. The probability of one reaction occurring relative to another is obtained by multiplying the rate constant of each reaction with the numbers of its substrate molecules. A random number is then used to choose which reaction will occur, based on relative probabilities. And in the Gillespie method, another random number determines how long the step will last, while in the ARMS such a random number is not used to determine how long the step will last and the steps of intervals between reaction events are given by the rate constants.

1 Abstract Rewriting System on Multisets, ARMS

Formally, in the *ARMS* a chemical solution is a multiset of elements denoted by symbols from a given alphabet, $A = \{a, b, \ldots, \}$; these elements correspond to *chemicals*.Reaction rules that act on the chemicals are specified in the *ARMS* by reaction rules. Let A be an *alphabet* (a finite set of abstract symbols). A *multiset* over a set of objects A is a mapping $M : A \mapsto \mathbf{N}$, where **N** is the set of natural numbers, **N**, 0, 1, 2,...

The number M(a), for $a \in A$, is the *multiplicity* of object a in the multiset M. We denote by $A^{\#}$ the set of all multisets over A, including the B (empty multiset, \emptyset , defined by $\emptyset(a) = 0$ for all $a \in A$). A multiset $M : A \mapsto \mathbf{N}$, for $A = \{a_1, \ldots, a_n\}$ is represented by the vector $w = (M(a_1)M(a_2)\ldots M(a_n))$.

The union of two multisets $M_1, M_2 : A \mapsto \mathbf{N}$ is addition of vectors w_1 and w_2 that represent the each multisets respectively. If $M_1(a) \leq M_2(a)$ for all $a \in A$, then we say that multiset M_1 is included in multiset M_2 and we write $M_1 \subseteq M_2$. A reaction rule $u \to v, u, v \in A^{\#}$ is a vector r, r = -u + v. Note that u and v can also be zero vector (empty). For example, the reaction $a \to c$ is the vector of (-1 -1 1)=-(1 1 0) + (0 0 1).

A reaction is the addition of vectors $M \in A^{\#}$ and $r \in R$, and it can be defined only when $r \subseteq M$. We can define over $A^{\#}$ a relation: (\rightarrow) : for $M, M' \in A^{\#}, r \in R$ we write $M \to M'$ iff M' = (M + r). As for strategy of rule application, one rule will be applied in each step, conventionally. But we can easily realize maximal parallel rule application.

Application of rules in the ARMS

Kinetics of bio-chemical reactions have traditionally been described by the reactiondiffusion (RD) equations based on the mass-action law (MAL). The chemical equation of

$$A + B \to C + D \tag{1}$$

indicates that molecules A and B react together to form molecules C and D. From this chemical equation we can obtain the rate equation. It is important to note that most chemical reactions are assumed to follow the mass action law (MAL) kinetics, meaning that the reaction rate is proportional to the concentration of the molecules. Thus the rate equation of the equation (1) is $-[\dot{A}] = r_a = k[A][B]$, where [A] represents the concentration of molecules A, r_a is the reaction rate and k is the rate constant of the reaction.

The reactions in the ARMS obey the Mass Action Law (MAL), where the frequency of a reaction follows the concentration of chemicals and a rate constant. In the ARMS, reaction rules are selected probabilistically, where each probability of selecting a rule is in proportion to the total number of collisions of chemicals. Concretely, the probability is given by the ratio of the total number of colliding chemicals of a reaction to the sum of the total number of colliding chemicals of every reactions in the rule; for example, there are only two reaction rules $a, b \xrightarrow{k_1} c: (r_1)$ and $c, d \xrightarrow{k_2} a: (r_2)$, the probabilities of selecting r_1 and r_2 are respectively given by,

$$P_{r_1} \equiv \frac{k_1[a][b]}{k_1[a][b] + k_2[c][d]},\tag{2}$$

$$P_{r_2} \equiv \frac{k_2[c][d]}{k_1[a][b] + k_2[c][d]}.$$
(3)

In this contribution, the change of concentration of chemicals are given by the expectation value of selected a rule and its stoichiometric constants, for example, the case when r_1 is selected, the change of concentration a is given as $P_{r_1} \times -1$, b, $P_{r_1} \times -1$, c, $P_{r_1} \times 1$ and d, $P_{r_1} \times 1$, respectively. In this contribution, the changes of concentrations obey those of expectation values.

Oregonator

The Oregonator is proposed by [2] as follows;

$$X, Y, H \xrightarrow{k_1} 2W : (r_1), \tag{4}$$

$$A, Y, 2H \xrightarrow{k_2} X, W : (r_2), \tag{5}$$

$$2X \xrightarrow{k_3} A, W, H: (r_3), \tag{6}$$

$$A, X, H \xrightarrow{k_4} 2X, 2Z : (r_4), \tag{7}$$

$$B, Z \xrightarrow{\kappa_5} 0.5Y : (r_5), \tag{8}$$

where $k_1 \dots k_5$ are obtained through chemical experiments and proposed [2]; $k_1 = 10^6 M^{-2} S^{-1}$, $k_2 = 2M^{-3} S^{-1}$, $k_3 = 2 \times 10^3 M^{-1} S^{-1}$, $k_4 = 10M^{-2} S^{-1}$, $k_5 = B \times 2 \times 10^{-2} S^{-1}$, where M stands for one molar, S stands for a second. And A corresponds to the concentration of $B_r O_3$, B, $CH(COOH)_2$, X, $HB_r O_2$, Y, B_r , Z, C_e^{4+} , W, HOB_r and H, H^+ , respectively.



Fig. 1. Population dynamics of X,Y,Z in the Oregonator, where the vertical axis illustrates the number of chemicals and the horizontal axis illustrates the step

In the Oregonator, chemicals A and B are resources and assumed that they are continuously supplied or largely existed than other chemicals. W is the final product through these reactions and oscillations among X, Y and Z emerge.

2 Results

We confirmed that the ARMS can simulate conventional behavior of the Oregonator by using the differential equations (figure 1), where reactions of generating X (Hb_rO_2) are trigger of oscillations and these reactions increase the concentration of Z (C_e^{4+}) then high concentration of Z leads reactions of generating Y (B_r) , since this reaction required Z, the concentration of Z is decreased.

The reaction mechanism of the Oregonator is illustrated by the usage of reaction rules (figure 2). Basically, the $r_1(X, Y, H \to 2W)$ and $r_5(B, Z \to 0.5Y)$ are used continuously. And $r_2(A, Y, 2H \to X, W)$ is also mainly used, however by the increase of X with $r_4(A, X, H \to 2X, 2Z)$, occasionally applications of r_2 is switched by $r_3(2X \to A, W, H)$. These usages of rules fit to actual chemical mechanism of the BZ reactions. Therefore, this result indicates not only the ARMS exhibits the same behavior that of modeled by the differential equations but also shows the correctness of chemical mechanism of the Oregonator.



Fig. 2. The usage of reaction rules, the When the vertical axis illustrates $r_1, r_2 \dots r_5$ from the top to bottom. The horizontal axis illustrates the steps. Each dot illustrates the usage of a rule

3 Discussion

Next, we consider the case when the system with large stochastic fluctuations. The change of concentration by a reaction is small and it can be approximated as continuous; however, most of the chemical reactions in a living system are performed with few molecules. In the reactions with few molecules, its developments are not approximated as continuous but discrete, where stochastic fluctuations can not be ignore.

In order to introduce discreteness in the developments, we define an exernal parameter $\rho(0.0 < \rho)$, it is multiplied by the denominator of the probability of selecting a rule. Under the same extent of concentration change by a reaction when the value of ρ is small, the probability of selecting a rule will be changed easier compared to the case when ρ is large.

For example, probabilities of selecting $a, b \xrightarrow{k_1} c : (r_1)$ and $c, d \xrightarrow{k_2} a : (r_2)$ are given by

$$P_{r_1} \equiv \frac{k_1[a][b]}{(k_1[a][b] + k_2[c][d])\rho},\tag{9}$$



Fig. 3. un-stable oscillations: population dynamics of X,Y,Z in the Oregonator when ρ is small (ρ is an external parameter), where the vertical axis illustrates the number of chemicals and the horizontal axis illustrates the step

$$P_{r_2} \equiv \frac{k_2[c][d]}{(k_1[a][b] + k_2[c][d])\rho},\tag{10}$$

where when $\rho = 1.0$ the probabilities are the same as the conventional definition, while $\rho < 1.0$, as the denominator is getting small, the probability becomes sensitive to the slight change of the value of the numerator. So, the value of ρ is getting smaller, probabilities are getting easier to suffer from the fluctuations, it corresponds to the case when the system with few molecules. And also, as the value of ρ is getting smaller, the change of concentration by a reaction becomes large and discretely.

System with large fluctuations

We found that there exists three cases of behaviors according to the value of ρ . As decreasing the value of ρ to 0.0, it shows stable oscillations, quasi-stable oscillations and un-stable oscillations, where stable oscillations means that conventional oscillations, quasi-stable oscillations, in some trials oscillations disappear in a lapse of steps while others show conventional oscillations, un-stable oscillations(figure 3 and 4), in every trials oscillations disappear in a lapse of steps. The usage of rules



Fig. 4. The usage of reaction rules the case when un-stable oscillations occur (ρ is an external parameter and this graph indicates when it is small), the vertical axis illustrates $r_1, r_2 \dots r_5$ from the top to bottom. The horizontal axis illustrates the steps. Each dot illustrates usage of a rule. From around 250,000 steps, the usage of reactions indicates a rule between 1 and 2 are used, it means that rule 1 and 2 selected but cannot be applied, because of the concentration of X, Y and Z become under 1.0

(figure 4) indicates that, since the concentration of X, Y and Z become under 1.0, no rule can be applied. This result illustrates that discreteness can be induced by low concentration of chemicals, as for this phenomena, the same mechanism has been reported in another abstract chemical system [9].

4 Final remarks

In this contribution, we report our preliminary results on direct simulation of the Oregonator. We confirm that when the Oregonator is affected by large stochastic fluctuations, the oscillatory behaviors may change. More detailed investigation on it is our future work, where in order to confirm the correctness of the investigation, chemical experiment might be required, it is also our future work.

And also, chemical systems could give a good analogy when we consider the complex interaction systems such as protein-protein interactions etc. We will try to apply this method to investigate such a biological network.

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EIGHTH WORKSHOP ON MEMBRANE COMPUTING

This volume contains the papers presented at the Eighth Workshop on Membrane Computing, WMC8 which has been organized in Thessaloniki, Greece, from June 25 to June 28, 2007. The 2007 edition of WMC was organised at City College, Thessaloniki, by the South-East European Research Centre (SEERC), under the auspices of the European Molecular Computing Consortium (EMCC).

As usual in the last years, a balanced attention was paid to both "theory" and "practice", to mathematical and theoretical computer science topics as well as to applications and implementations. This is especially visible in what concerns the five invited talks, all of them included in the present volume, delivered by Luca Bianco, Pierluigi Frisco, Alberto Leporati, Andrea Maggiolo-Schettini, and Gheorghe Stefan. The volume also contains the 32 accepted papers. Each of them was subject of two or three referee reports.

Details about membrane computing can be found at the area web site (maintained in Milano, Italy) from http://psystems.disco.unimib.it. The workshop web site was http://www.seerc.org/wmc8/, where you will be able to find all papers in electronic form. The workshop was sponsored by City College, Thessaloniki, and the South-East European Research Centre (SEERC). Special thanks to the organising and programme committee, the invited speakers, the authors of the papers, the presenters, the reviewers, and all the participants; it is their effort that made the Eighth Workshop on Membrane Computing (WMC8) a very successful event.







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