A Testing Framework for P Systems

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Testing equivalence was originally defined by De Nicola and Hennessy in a process algebraic setting (CCS) with the aim of defining an equivalence relation between processes being less discriminating than bisimulation and with a natural interpretation in the practice of system development. Finite characterizations of the defined preorders and relations led to the possibility of verification by comparing an implementation with a specification in a setting where systems were seen as black boxes with input and output capabilities, thus neglecting internal undetectable behaviours.

In this paper, we start defining a porting of the well-established testing theory into membrane computing, in order to investigate possible benefits in terms of inherited analysis/verification techniques and interesting biological applications. P Algebra, a process algebra for describing P Systems, is used as a natural candidate for the porting since it enjoys the desirable property of being compositional and comes with other observational equivalences already defined and studied.

We consider P Systems with multiple membranes, dissolution, promoters and inhibitors. Notions as observable and test are conveniently rephrased in the membrane scenario, where they lack as native notions and have a not so obvious mean. At the same time, concepts as promoters, inhibitors, membrane inclusion and dissolution are emphasized and exploited in the attempt of realizing a testing machinery able to formalize several features, which are proper of membranes and, as a consequence, worth being highlighted as basic observables for P Systems. The new testing semantics framework inherits from the original one the ability to define qualitative system properties. Moreover, it results to be suitable also to express quantitative aspects, a feature which turns out to be very useful for the biological domain and, at the same time, puts in evidence an expected high expressive power of the framework itself.

1 Introduction

Membrane computing, the research field initiated by Gheorghe Păun [22, 20], aims to define computational models, called P Systems, which are inspired by the behaviour and structure of the living cell. Since its introduction, the P System model has been intensively studied and developed: many variants of membrane systems have been proposed and regular collective volumes are annually edited.

The most investigated membrane computing topics are related to the computational power of different variants, their capabilities to solve hard problems, decidability, complexity aspects and hierarchies of classes of languages produced by these devices.
In the last years, there have also been significant developments in using the P systems paradigm for modelling, simulation and formal verification [11]. Although such topics have been exercised to different classes of P Systems [3], testing has been quite neglected in this context.

1.1 Testing and P Systems

Testing P Systems has been so far considered by using certain coverage principles. More often the rule coverage is utilised, by taking into account different contexts. Such contexts - typically grammar, automaton and model checking techniques - are described in depth in [14,15]:

- **(Grammar-based methods)** In order to test an implementation developed from a P System specification in a grammar-based method, a test set is built, in a black box manner, as a finite set of sequences containing references to rules. Although there are similarities between context-free grammars utilised in grammar testing and basic P Systems, there are also major differences that pose new problems in defining testing methods and strategies to obtain test sets. Some of the difficulties encountered when some grammar-like testing procedures are introduced, are related to the hierarchical compartmentalisation of the P System model, parallel behaviour, communication mechanisms, the lack of a non-terminal alphabet and the use of multisets of objects instead of sets of strings.

- **(Finite-state machine methods)** Finite state machine-based testing is widely used for software testing. It provides very efficient and exhaustive testing strategies and well investigated methods to generate test sets. In this case it is assumed that a model of the system under test is provided in the form of a finite state machine. In the P System model case, such a machine is typically obtained from a partial computation in a P System.

A different approach uses a special class of state machines, called X-machines. Given that the relationships between various classes of P Systems and these machines are well studied [1] and the X-machine-based testing is well developed, standard techniques for generating test sets based on X-machines can be adapted to the case of P Systems [16].

Specific coverage criteria are defined in the case of finite state machine-based testing. One such criterion, called **transition coverage**, aims to produce a test set in such a way that every single transition of the model is covered.

- **(Model checking-based methods)** The generation of different test sets, according to certain coverage criteria, can be done by utilising some specific algorithms or by applying some tools that indirectly will generate test sets. Such tools, like model checkers, can be used to verify some general properties of a model and when these are not fulfilled then some counter-examples are produced, which act as test sets in certain circumstances.

In the case of P Systems, an encoding based on a Kripke structure associated with the system is provided for model checkers like NuSMV [17] or SPIN [18]. This relies on certain operations defined in [12] and encapsulates the main features of a P System, including maximal parallelism and communication, but within a finite space of values associated with the objects present in the system.

The rule coverage principle is expressed by using temporal logics queries available in such contexts. By negating specific coverage criteria, counter-examples are generated.
1.2 Our contribution: a Process Algebra-based testing machinery for P Systems

The community of Process Algebra taught us that the usefulness of formalisms for the description and the analysis of reactive systems is closely related to the underlying notion of \textit{behavioural equivalence}. Such an equivalence should formally identify behaviours that are informally indistinguishable from each other, and at the same time distinguish between behaviours that are informally different. The authors of \cite{4} go toward such a direction, proposing some observational equivalences on a suitable algebraic notation of P Systems.

One way of determining behavioural equivalences is by observing the systems we are interested in, experimenting on them, and drawing conclusions about the behaviour of such systems based on what we see, e.g., testing the system. Such an approach has been formalized in the Process Algebra setting by a suitable testing machinery \cite{13}, pivoting on a restricted form of context, called \textit{observer}.

The way to exercise (evaluate) a process on a given observer is done by letting the considered process and the given observer to run in parallel and by looking at the computations which the running test can perform.

It is worth noting that internal actions of the process under test do not affect, but the case in which they lead to divergence, the satisfaction of the test: they are not observable as input or outputs, thus they cannot be perceived by an external user that is experimenting on the system. This idea is typical of a testing framework: systems are considered black boxes and only observables matter in their comparison. This characteristic is imported in the testing notion introduced in this paper. Internal production and migration of objects in the P system under test will not be seen by the observer: only the objects injected into the system by the observer and the objects that are returned from the system to the observer will matter.

Another typical characteristic of the testing framework, worth underlining, is that the observer must have the ability to force the system under test to follow certain paths among all the possible ones. This is in order to investigate, for instance, what the system can do after a quite specific sequence of inputs, or after a predefined sequence of inputs/outputs. In order to guarantee this possibility, and thus giving the equivalence a discriminating power similar to the one in the original setting, in the testing framework we introduce in this paper we exploit promoters and inhibitors. Without them the intrinsic nature of P system behaviour, in particular maximal parallelism, would have prevented this central feature, thus invalidating the porting.

The characteristics discussed above are central in the idea of testing equivalence. Bisimulation-based equivalences, even the weak one, are highly discriminating and do not reflect a practical view of “testing” a system: usually, and \textit{this is always the case for biological systems}, the whole internal structure/dynamics of the system is not known, but it is required to check bisimulation. The only way to study such systems is to interact with them and analyse what can be observed from experiments, with the means that are available. Along this idea, we start with this work the definition of a testing framework with the characteristics above, giving initial theoretical results and some simple examples of tests, without any particular biological impact. However, we intend as future work to investigate and exploit the analogy of the defined notion of testing with biological experiments in order to give more evidence of the biological relevance of the work. We can devise, for instance, techniques for experimental planning that could be of great interest for experimental biologists, along the same line of the techniques proposed in \cite{2}.

In \cite{13}, different equivalence relations (e.g., \textit{may} and \textit{must} testing equivalences) between systems are defined. Two systems are considered equivalent if they pass exactly the same set of observers. Such equivalences are further broken down into preorder relations on systems, i.e., relations that are reflexive
and transitive (though not necessarily symmetric). Formally, given a process $P$ and an observer $o$,

- $P \text{ may } o$ means that there exists a successful computation from the test $P\mid o$ (where $\mid$ is the parallel operator, and successful means that there is a state where the special action $\omega$ is enabled);

- $P \text{ must } o$ means that every maximal computation from the test $P\mid o$ is successful;

- The preorder $P \leq_{\text{sat}} Q$ means that for any observer $o$, $P\text{ sat } o$ implies $Q\text{ sat } o$, where sat denotes may or must;

- The equivalence $P \approx_{\text{sat}} Q$ means $P \leq_{\text{sat}} Q$ and $Q \leq_{\text{sat}} P$.

In [23] and in [19] a new testing semantics was proposed to incorporate the fairness notion: the fair-testing (aka should-testing) semantics. In contrast to the classical must-testing (semantics), fair-testing abstracts from certain divergences, e.g., infinite loops of $\tau$ (invisible) actions. This is achieved by stating that the observer $o$ is satisfied if success always remains within reach in the system under observer. In other words, $P \text{ fair } o$ holds if in every maximal computation from $P\mid o$ every state can lead to success after finitely many interactions.

On the basis of $P$ Algebra, the algebraic notation of P Systems introduced in [4] (see Section 2), we adapt for P Systems the testing machinery defined in [13] (see Section 3).

We introduce the concept of context in case of P Systems expressed as P Algebra terms. Using this concept, we then define what we consider an observer, which is again a P Algebra term with certain characteristics. This leads naturally to the definition of computations of a running test, i.e. a tested system running together with an observer. Then, following the classical definition of [13], we define the success of a running test in the two well-known versions of may and must. Finally, the testing preorders are introduced, together with the induced equivalence relations. More in detail:

- An observer consists of a membrane structure in which the skin membrane contains several membranes (with possibly sub-membranes) one of which is a hole, i.e., a place where another fully defined P System can be placed and run. The skin membrane of this tested P System instantiates the hole membrane and becomes a full component of the running test.

- P Systems that are observers are distinguished from P Systems that are normal, testable processes similarly to the classical testing approach, where a particular action, called $\omega$, is used to denote the success of a test and, if the running test is able to perform this action, then the computation under consideration is a successful one. Similarly, in our framework this is easily translated introducing a fresh, particular object $\omega$ that, when sent out of the skin membrane of the running test, denotes the success of the computation.

- As usual in testing frameworks, we consider only the behaviours of the running test in which no output is produced (this corresponds to considering only invisible action (e.g., $\tau$) computations in a CCS-like Process Algebra). This is needed to explore all possible behaviours of the tested system while running together with the observer.

An important result consists of the fact that, differently from the original testing semantics framework, the one proposed here for P Systems results to be suitable to define both qualitative and (above all) quantitative system properties. This is mainly because the formalism of P system is expressive enough to express both qualitative and quantitative aspects. However, these features are crucial for the biological domain. In Section 4 we put quantitative capabilities in evidence defining examples of quantitative tests, both concerning time and number of individuals, of a system modelling a population of individuals that

\footnote{In this paper we do not consider fair testing.}
can reproduce both sexually and asexually. Note that these examples are to be intended only as explanation of the concepts introduced in the paper and not as examples of verifications in the model-checking style. The main direction of continuing this work towards verification is to find finite characterizations of the resulting testing equivalence, possibly with respect to suitable classes of observers, and thus using them to compare the expected behaviour of a finite state system with its actual behaviour.

2 Background

We first briefly recall the definition of P Systems [22] [20]. Then, we give the definition of the syntax and the semantics of the P Algebra as it was presented in [4], where a class of P Systems including rule promoters and inhibitors [10] was considered. The original formulation of the P Algebra, without promoters and inhibitors, can be found in [7] [6] that we refer as a more detailed presentation of the semantics.

2.1 P Systems with Promoters and Inhibitors

A P System consists of a hierarchy of membranes, each of them containing a multiset of objects, representing molecules, a set of evolution rules, representing chemical reactions, and possibly other membranes. Each evolution rule consists of two multisets of objects, describing the reactants and the products of the chemical reaction. A rule in a membrane can be applied only to objects in the same membrane. Some objects produced by the rule remain in the same membrane, others are sent out of the membrane, others are sent into the inner membranes (assumed to exist) which are identified by their labels.

In the original definition of P Systems, rules are applied with maximal parallelism, namely it cannot happen that a rule is not applied when the objects needed for its triggering are available. Here, we assume that at each step at least one evolution rule in the whole system is applied, and also that more than one rule and several occurrences of the same rule can be applied at the same step (to different objects). In other words, we assume that at each step a multiset of evolution rule instances is non-deterministically chosen and applied in each membrane, such that in the whole system at least one rule is applied. This is a general form of parallelism that is better suited than the maximal one to describe events in biological systems.

In P Systems with promoters and inhibitors an evolution rule in a membrane may have some promoters and some inhibitors. Promoters are objects that are required to be present and inhibitors are objects that are required to be absent in the membrane \(m\) in order to enable the application of the rule. Promoters will be denoted simply as objects, namely \(a, b, c, \ldots\), while inhibitors will be denoted as objects preceded by a negation symbol, namely \(\neg a, \neg b, \neg c, \ldots\).

We denote with \(\mathcal{D}_V\) the set of all possible promoters and inhibitors symbols that can be obtained from an alphabet \(V\), namely \(\mathcal{D}_V = V \cup \neg V\). Given a set of promoter and inhibitor symbols \(D\), we denote with \(D^+\) and \(D^-\) the sets of objects containing all the objects occurring in \(D\) as promoters and all the objects occurring in \(D\) as inhibitors, respectively. We remark that \(D^+\) and \(D^-\) are sets of objects, hence elements on \(D^-\) will not be preceded by \(\neg\). Moreover, with \(\neg D\) we denote the set obtained by transforming each promoter in \(D\) into an inhibitor and vice versa. As an example, if \(D = \{a, \neg b, \neg c, d\}\) we have \(D^+ = \{a, d\}\), \(D^- = \{b, c\}\) and \(\neg D = \{\neg a, b, c, \neg d\}\).

We assume that all evolution rules have the following form, where \(u,v_h,v_o,v_1,\ldots,v_n\) are multisets of objects, \(\{l_1,\ldots,l_n\}\) is a set of membrane labels in \(\mathbb{N}\), and \(D\) is a set of promoters and inhibitors:

\[
u \rightarrow (v_h, \text{here})(v_o, \text{out})(v_1, \text{in}_{l_1}) \ldots (v_n, \text{in}_{l_n})||_{D}.
\]
A rule can be applied only if requirements expressed by $D$ are satisfied. When a rule is applied, the multiset of objects $u$ is replaced by $v_h$, multiset $v_j$ is sent to the parent membrane, and each $v_i$ is sent to inner membrane $l_i$. Promoters are not consumed by the application of the corresponding evolution rule and a single occurrence of a promoter may enable the application of more than one rule in each evolution step. Similarly, a single occurrence of an inhibitor forbids the application of all the evolution rules in which it appears. We assume that the set of promoters and inhibitors $D$ of an evolution rule does not contain the same object both as a promoter and as an inhibitor, namely $D^+ \cap D^- = \emptyset$, and that consumed objects $u$ are not mentioned among inhibitors, namely $u \cap D^- = \emptyset$.

**Definition 1.** A P System $\Pi$ is a tuple $(V, \mu, w_1, \ldots, w_n, R_1, \ldots, R_n)$ where:

- $V$ is an alphabet whose elements are called objects;
- $\mu \subseteq \mathbb{N} \times \mathbb{N}$ is a membrane structure, such that $(l_1, l_2) \in \mu$ denotes that the membrane labelled by $l_2$ is contained in the membrane labelled by $l_1$;
- $w_j$ with $1 \leq j \leq n$ are multisets of objects in $V$ associated with the membranes $1, \ldots, n$ of $\mu$;
- $R_j$ with $1 \leq j \leq n$ are finite sets of evolution rules associated with the membranes $1, \ldots, n$ of $\mu$.

**2.2 The P Algebra: Syntax and Semantics**

In this section we recall the $P$ Algebra, the algebraic notation of P Systems we have introduced in [7], with slight modifications introduced in [4]. We assume $V$ to be an alphabet of objects and we adopt the usual string notation to represent multisets of objects in $V$. For instance, to represent \{a,a,b,b,c\} we may write either $aabbc$, or $a^2b^2c$, or $(ab)^2c$. We denote with $\text{Set}(u)$ the support of multiset $u$, namely the set of all the objects occurring in $u$. We denote multiset (and set) union as string concatenation, hence we write $u_1u_2$ for $u_1 \cup u_2$. Moreover, we shall write $u(a)$ for the number of occurrences of a in multiset $u$. For the sake of legibility, we shall write $u \rightarrow v_h v_o \{v_i\}_D$ for the generic evolution rule $u \rightarrow (v_h, \text{here})(v_o, \text{out})(v_1, \text{in}_h) \ldots (v_m, \text{in}_h)_D$.

The abstract syntax of the P Algebra is defined as follows.

**Definition 2 (P Algebra).** The abstract syntax of membrane contents $c$, membranes $m$, and membrane systems $ms$ is given by the following grammar, where $l$ ranges over $\mathbb{N}$ and $a$ over $V$:

$$
c ::= (\emptyset, \emptyset) \mid (u \rightarrow v_h v_o \{v_i\}_D, \emptyset) \mid (\emptyset, a) \mid c \cup c
\$$

$$
m ::= [l c]
\$$

$$
ms ::= ms \mid ms \mid \mu(m, ms) \mid F(m)
\$$

A membrane content $c$ represents a pair $(\mathcal{R}, u)$, where $\mathcal{R}$ is a set of evolution rules and $u$ is a multiset of objects. A membrane content is obtained through the union operation $\_ \cup \_ \_ \_$ from constants representing single evolution rules and single objects, and can be plugged into a membrane with label $l$ by means of the operation $[l \_ \_ \_]_l$ of membranes $m$. Hence, given a membrane content $c$ representing the pair $(\mathcal{R}, u)$ and $l \in \mathbb{N}$, $[l c]$ represents the membrane having $l$ as label, $\mathcal{R}$ as evolution rules and $u$ as objects.

Membrane systems $ms$ have the following meaning: $ms_1 \mid ms_2$ represents the juxtaposition of $ms_1$ and $ms_2$, $\mu(m, ms)$ represents the hierarchical composition of $m$ and $ms$, namely the containment of $ms$ in $m$, and $F(m)$ represents a flat membrane, namely it states that $m$ does not contain any child membrane. Juxtaposition is used to group sibling membranes, namely membranes all having the same parent in a membrane structure. This operation allows hierarchical composition $\mu$ to be defined as a binary operator on a single membrane (the parent) and a juxtaposition of membranes (all the children).
Note that every P System has a corresponding membrane system in the P Algebra, and that there exist membrane systems which do not correspond to any P System.

In what follows we will often write \([\llbracket l \rrbracket c]\) for \(F([\llbracket l \rrbracket c])\). We shall also often write \((\mathcal{R}, u)\) where \(\mathcal{R} = \{r_1, \ldots, r_n\}\) is a set of rules and \(u = o_1 \ldots o_m\) a multiset of objects rather than \((\emptyset, \emptyset) \cup \ldots \cup (r_n, \emptyset) \cup (\emptyset, o_1) \cup \ldots \cup (\emptyset, o_m)\). Moreover, we shall often omit parentheses around membrane contents.

The semantics of the P Algebra is given as a labelled transition system (LTS). The labels of the LTS can be of the following forms:

- \((u, v, v', D, I, O^1, O^1)\), describing a computation step performed by a membrane content \(c\), where:
  - \(u\) is the multiset of objects consumed by the application of evolution rules in \(c\), as it results from the composition, by means of \(\_\cup \_\), of the constants representing these evolution rules.
  - \(v\) is the multiset of objects in \(c\) offered for the application of the evolution rules, as it results from the composition, by means of \(\_\cup \_\), of the constants representing these objects. When operation \([\_\_\_\_]\) is applied to \(c\), it is required that \(v\) and \(u\) coincide.
  - \(v'\) is the multiset of objects in \(c\) that are not used to apply any evolution rule and, therefore, are not consumed, as it results from the composition, by means of \(\_\cup \_\), of the constants representing these objects.
  - \(D\) is a set of promoters and inhibitors required to be present and absent, respectively, by the application of evolution rules in \(c\). More precisely, \(D^-\) contains all the inhibitors of the applied evolution rules in \(c\), whereas \(D^+\) is a subset of the promoters of those rules. Such a subset contains only those objects that are not present in the multiset of objects of \(c\).
  - \(I\) is the multiset of objects received as input from the parent membrane and from the child membranes.
  - \(O^1\) is the multiset of objects sent as an output to the parent membrane.
  - \(O^1\) is a set of pairs \((l_i, v_{il})\) describing the multiset of objects sent as an output to each child membrane \(l_i\).
- \((\mathcal{F}^1, I^1, O^1, O^1, app)\), describing a computation step performed by a membrane \(m\), where: \(\mathcal{F}^1\) is a set containing only the pair \((l, I)\) where \(l\) is the label of \(m\) and \(I\) is the multiset of objects received by \(m\) as input from the parent membrane, \(I^1\) is the multisets of objects received from the child membranes of \(m\), and \(O^1\) and \(O^1\) are as in the previous case. Finally, \(app \in \{0, 1\}\) is equal to 0 if no rule has been applied in \(m\) in the described computation step, and it is equal to 1 otherwise.
- \((\mathcal{F}^1, O^1, app)\), describing a computation step performed by a membrane system \(ms\), where \(\mathcal{F}^1\), \(O^1\) and \(app\) are as in the previous cases.

For the sake of legibility, in transitions with labels of the first form we shall write the first four elements of the label under the arrow denoting the transition and the other elements over the arrow. Now, LTS transitions are defined through SOS rules [21]. We give here a very short explanation of such rules. Please, refer to [7] for more details.

We start by giving in Fig. 1 the transition rules for membrane contents. Rule \((mc1_n)\) describes \(n\) simultaneous applications of an evolution rule for any \(n \in \mathbb{N}\). Rule \((mc2)\) describes the case in which an evolution rule is not applied because a subset \(D'\) of the promoters and inhibitors in \(D\) it requires to be present and absent, respectively, are assumed not to satisfy the requirements. Rules \((mc3), (mc4)\) and \((mc5)\) describe the transitions performed by membrane contents consisting of a single object and the transitions performed by an empty membrane content.

Rule \((u1)\) describes the behaviour of a union of membrane contents. In this transition rule we use some auxiliary notations. Given two sets \(O^1\) and \(O^1\) representing two outputs to inner membranes,
we write $O_1 \cup \in \mathbb{N} O_2$ to denote the set $\{(l, uv) \mid (l, u) \in O_1 \wedge (l, v) \in O_2\} \cup \{(l, u) \mid (l, u) \in O_1 \wedge \exists v. (l, v) \in O_2\} \cup \{(l, v) \mid (l, v) \in O_2 \wedge \exists u. (l, u) \in O_1\}$.

In Fig. 2 we give transition rules for individual membranes, juxtaposition and hierarchical composition. Note that from the transition label of the membrane content we have no information about the objects that have been produced by the applied evolution rules. Rules (m1) and (m2) describe the transitions performed by a membrane with label $l$. In particular, (m1) describes the case in which no objects are received as an input from the external membrane, while (m2) describes the case in which a multisets of objects $I_1 \neq \emptyset$ is received. In these rules app is set to zero if no evolution rule is applied ($u = \emptyset$), and it is set to one if at least one rule is applied ($u \neq \emptyset$). Rule (fm1) allows us to infer the behaviour of a flat membrane $[[c]] = F([[c]])$ from the behaviour of membrane $[c]$. Rule (jux1) allows us to infer the behaviour of a juxtaposition of two membrane structures from the behaviours of the two structures. Finally, rule (h1) describes the behaviour of a hierarchical composition of membranes. In this rule we assume $\approx$ to be an equivalence relation on sets of pairs $(l, u)$ with $l \in \mathbb{N}$ and $u \in V^*$, such that, given two such sets $\mathcal{F}_1$ and $\mathcal{F}_2$, then $\mathcal{F}_1 \approx \mathcal{F}_2$ holds if and only if $\mathcal{F}_1 \setminus \{(l, \emptyset) \mid l \in \mathbb{N}\} = \mathcal{F}_2 \setminus \{(l, \emptyset) \mid l \in \mathbb{N}\}$. In the last two rules app is set to one if at least one between app1 and app2 is equal to one, namely $app = \max(app1, app2)$. This means that at least one rule has been applied in the whole composition.

We conclude by defining a system trace as a sequence of internal information given by an execution of a P Algebra term. We assume that the system can send objects out of the outmost membrane, but cannot receive objects from outside. This requirement corresponds to the fact that in a P System objects cannot be received by the outmost membrane from the external environment. Note that executions containing steps in which no rule is applied, namely those with 0 as last element of the label, are not considered.

**Definition 3** (Trace). A trace of a membrane system $ms_i$ with alphabet $V$ is a (possibly infinite) sequence $w$ of outputs such that, for any $O_1^i$ and $ms_i$ with $i \in \mathbb{N}^+$
Figure 2: Rules for individual membranes and hierarchical composition of membranes

\[ x \xrightarrow{1,0^1,0^2} y \quad \text{app} = \begin{cases} 0 & \text{if } u = \emptyset \\ 1 & \text{otherwise} \end{cases} \quad D^+ = \emptyset \]  
\[ (m1) \]

\[ x \xrightarrow{I_1,I_2,O^1,O^2} y \quad \text{app} = \begin{cases} 0 & \text{if } u = \emptyset \\ 1 & \text{otherwise} \end{cases} \quad D^+ = \emptyset \quad I_1 \neq \emptyset \]  
\[ (m2) \]

\[ x \xrightarrow{\tau^1,\tau^2,\tau,\text{app}} F(x) \xrightarrow{\tau^1,\tau^2,\tau,\text{app}} F(y) \]  
\[ (f1m) \]

\[ x \xrightarrow{\tau_1,\tau_1^1,\text{app},\rho_1,\rho_2,\rho_3} y \xrightarrow{\tau_1,\tau_1^1,\text{app},\rho_1,\rho_2,\rho_3} y_1 \xrightarrow{\tau_2,\tau_2^1,\text{app},\rho_1,\rho_2,\rho_3} y_2 \]  
\[ (jux1) \]

\[ x_1 \xrightarrow{\tau_1^1,\tau_1^1,\text{app},\rho_1,\rho_2,\rho_3} y_1 \xrightarrow{\tau_2,\tau_2^1,\text{app},\rho_1,\rho_2,\rho_3} y_2 \]  
\[ O^1_1 \equiv \tau_1 \quad O^2_2 = I^1_1 \]  
\[ (h1) \]

We denote with \( \mathcal{T} \) the set of all traces.

\section{Testing framework}

We first introduce the concept of context in case of P Systems expressed as P Algebra terms. Using this concept, we then define what we consider an observer, which is again a P Algebra term with certain characteristics. This leads naturally to the definition of computations of a running test, i.e., a tested system running together with an observer. Then, following the classical definition of \cite{13}, we define the success of a test in the two well-known versions of may and must. Finally, the testing preorder is introduced, together with the induced equivalence relation.

\subsection{Contexts and test satisfaction}

At a first look, a natural candidate for context of a P System is another P System, which we call observer, consisting of a membrane structure in which the skin membrane contains several membranes (with possibly sub-membranes) one of which is a “hole”, i.e., a place where another fully defined P System can be placed and run. The skin membrane of this tested P System instantiates the “hole” membrane and becomes a full component of the running test.

\footnote{This instantiation process may require some trivial modifications of the tested P System, such as \( \alpha \)-conversion of the numbers assigned to the skin and inner membranes.}
However, in [6], a result regarding flattening P Systems in P Algebra is presented. A similar result can be found in [8]. The flattening process of [6] reduces any P System, specified in P Algebra with promoters and inhibitors, into a flat one (i.e., with no internal membranes) that is bisimilar to the original one. The notion of bisimulation is the one, based on computation steps, defined in [7]. These results suggested us to simplify, without loss of generality, the notion of context we are defining. For this reason, instead of considering an observer of the form \( \mu(m, ms_1 \mid ms_2 \mid \cdots \mid ms_n \mid \Box) \), we will always consider the equivalent observer \( \mu(m', \Box) \) where \( m' \) results from the flattening process of \( \mu(m, ms_1 \mid ms_2 \mid \cdots \mid ms_n) \).

Another ingredient of the testing framework is needed to distinguish formally P Systems that are observers from P Systems that are normal, testable processes. In classical testing a particular action, usually called \( \omega \), is used to denote the “success” of a test, that is to say, if the running test is able to perform this action then the computation under consideration is a successful one. This easily translates in our framework: we introduction a fresh, particular object, \( \omega \not\in V \) that, when sent out of the skin membrane of the running test, denotes the “success” of the computation.

**Definition 4 (Observer).** Let \( V \) be the alphabet of objects and let \( \omega \not\in V \) be a particular object. An observer system, or simply a test, is a P Algebra term of the form \( \mu(m, \Box) \) where \( m \) is the skin membrane \([1, c]_1\), \( \Box \) is an unspecified membrane \( ms \) numbered 2 and each rule of \( m \) is of the form \( u \rightarrow v_0 \omega v_1 \{v_2\} D \). In other words, \( m \) communicates with its only child membrane 2 and can send out only \( \omega \) objects.

**Definition 5 (Running test).** Let \( V \) be the alphabet of objects and let \( \omega \not\in V \) be a particular object. Let \( \mu(m, \Box) \) be an observer system and let \( ms \) be term in P Algebra denoting a closed membrane (i.e., of the form \( F(-) \) or \( \mu(-, -) \)). The running test is the P Algebra term \( \mu(m, ms) \) where \( ms' \) is the term \( ms \) in which the former skin membrane (numbered 1) is re-labelled in 2 and the other internal membranes numbers are \( \alpha \)-converted in order not to collide with 1 and 2 (the re-labelling of the membrane names of course implies also applying the substitutions to all the references to the membrane names in the term.)

**Definition 6 (Computations).** Let \( \mu(m, ms) \) be a running test. A computation \( c \) of \( \mu(m, ms) \) is any sequence of the form:

- **c is finite:** \( \mu(m, ms) = ms_0 \xrightarrow{\varnothing, \varnothing, 1} ms_1 \xrightarrow{\varnothing, \varnothing, 1} \cdots \xrightarrow{\varnothing, \varnothing, 1} ms_n \xrightarrow{\varnothing, \varnothing, 1} \)

or

- **c is infinite:** \( \mu(m, ms) = ms_0 \xrightarrow{\varnothing, \varnothing, 1} ms_1 \xrightarrow{\varnothing, \varnothing, 1} ms_2 \xrightarrow{\varnothing, \varnothing, 1} \cdots \)

A computation is called successful if there are \( k \in \mathbb{N} \) and \( n \in \mathbb{N}^+ \) such that \( ms_k \xrightarrow{\varnothing, \{\omega\} 1} ms_{k+1} \), i.e., at least a success symbol can be sent out of the skin membrane along the run. A membrane like \( ms_k \), from which a transition can be taken that sends out at least one \( \omega \) object is called success membrane. We may also write \( ms_k \xrightarrow{\omega} \) to indicate that \( ms_k \) is a success membrane.

Note that, as usual in testing frameworks, we consider only the behaviours of the running test in which no output is produced (this corresponds to considering only \( \tau \) computations in a CCS-like Process Algebra). This is needed to explore all possible behaviours of the tested system while running together with the observer. Note that an unsuccessful computation, i.e. one with no success state along it, can be either finite or infinite. In testing theories divergence must have a delicate treatment because it can lead to different testing preorders [13][9]. For simplicity we port here the original formulation of [12]. Thus, we define a unary predicate \( ms \uparrow \) meaning that, with respect to success, the “state” \( ms \) along a computation is “underdefined” because it leads to a divergent computation.
Definition 7 (Divergence). Let $c$ be a computation of a running test $\mu(m, ms)$. $c$ is divergent, which we denote by $c \uparrow$ iff:

- $c$ is unsuccessful, or
- $c$ contains a membrane $ms$ such that $ms \uparrow$ and there is not a success membrane $ms'$ preceding $ms$ in $c$.

We now have all the ingredients to define the success of a test. Following the classical approach, such a definition comes into version: a *may* satisfaction, weaker, and a *must* satisfaction, stronger.

Definition 8 (Satisfaction of a test). Let $\mu(m, \square)$ be a test and $ms$ be a closed membrane. Then we say:

- $ms$ may $\mu(m, \square)$ iff there exists a computation $c$ of $\mu(m, ms)$ and $k \in \mathbb{N}$ such that $c : \mu(m, ms) = ms_0 \xrightarrow{\emptyset, \emptyset, 1} ms_1 \xrightarrow{\emptyset, \emptyset, 1} \cdots \xrightarrow{\emptyset, \emptyset, 1} ms_k$ and $ms_k \rightarrow_\omega$.

- $ms$ must $\mu(m, \square)$ iff for each computation $c$ of $\mu(m, ms)$, $c : \mu(m, ms) = ms_0 \xrightarrow{\emptyset, \emptyset, 1} ms_1 \xrightarrow{\emptyset, \emptyset, 1} ms_2 \xrightarrow{\emptyset, \emptyset, 1} \cdots$ the following conditions hold:
  1. there is $n \in \mathbb{N}$ such that $ms_n \rightarrow_\omega$.
  2. if there is $k \in \mathbb{N}$ such that $ms_k \uparrow$, then there exists $k' \leq k$ such that $ms_{k'} \rightarrow_\omega$.

Note that, in case of *must* satisfaction, computations can be infinite, but it is required that a success state is present just at the beginning of divergence, or before.

3.2 Testing preorders and testing equivalences

Using the definitions of satisfaction of a test, we naturally derive preorders between membrane systems.

Definition 9. Let $ms_1$ and $ms_2$ two closed membrane systems. We define two relations $\sqsubseteq^m_T$ and $\sqsubseteq^T$ between closed membrane systems as follows:

- $ms_1 \sqsubseteq^m_T ms_2$ iff for each observer $\mu(m, \square)$, $ms_1$ may $\mu(m, \square)$ implies $ms_2$ may $\mu(m, \square)$.
- $ms_1 \sqsubseteq^T ms_2$ iff for each observer $\mu(m, \square)$, $ms_1$ must $\mu(m, \square)$ implies $ms_2$ must $\mu(m, \square)$.

Proposition 1. The relations $\sqsubseteq^m_T$ and $\sqsubseteq^T$ are preorders.

Definition 10 (Testing equivalence). We say that $ms_1$ is may testing equivalent to $ms_2$, and write $ms_1 \approx^m_T ms_2$, iff $ms_1 \sqsubseteq^m_T ms_2$ and $ms_2 \sqsubseteq^m_T ms_1$.

Analogously, we say that $ms_1$ is must testing equivalent to $ms_2$, and write $ms_1 \approx^T ms_2$, iff $ms_1 \sqsubseteq^T ms_2$ and $ms_2 \sqsubseteq^T ms_1$.

Since the defined relations are kernels of preorders, it is easy to conclude that they are equivalence relations.

In [7] some equivalence relations are defined between the terms of P Algebra. Among them, we consider bisimulation, denoted by $\approx$, and trace equivalence, denoted by $\approx_{Tr}$. We show that the relationships between these two equivalences with the *must* testing equivalence are the same that hold when classical process calculi, as CCS, are considered [13].

Proposition 2. Let $\approx$, $\approx^m_T$ and $\approx^T$, as above. Then, $\approx \subset \approx^T \subset \approx^m_T$. 

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Figure 3: Two membrane systems, (a) and (c), that are trace equivalent, but not test equivalent, together with the observer (b) that distinguishes them.

Proof. The two set inclusions can be proved rephrasing the argument used for CCS [13]. Regarding the strictness of the inclusions, Figure 3 shows two systems that are trace equivalent, but can be distinguished by the shown observer. Not that both systems perform the set of traces \{∅(α)(β), ∅(α)(γ)\}. However, considering the must testing, the observer shown in Figure 3 (b) is not satisfied by the system (a) because in one computation, after α, only γ can be produced, thus that computation is unsuccessful. Conversely, for the other system (c) all computations are successful. Note that inhibitors and promoters are used in this settings to select only specific paths in the system. In the classical setting of testing for synchronous calculi this role is played by the parallel operator together with the restriction on observing only τ-computations.

Figure 4 shows two systems that are test equivalent, but not bisimilar. Below each system the graph of possible transitions is depicted, in order to show that bisimulation does not hold.

4 Example of testing scenario

In this section we model a population of individuals that can reproduce both sexually and asexually. We define different observers to show the expressiveness of the defined testing framework. It is worth noting that quantitative aspects of systems can be easily expressed. Note that the defined observers are very specific and are intended to only show the capabilities of the framework introduced above. In particular, they should not be intended as examples of verification, because this analysis is to be done by checking the testing equivalence between the system and its expected behaviour, modelled as a simpler “specification” P system.

Most animal species use sexual reproduction to produce offspring, while a minority of species reproduce asexually by producing clones of the mother. Both strategies have advantages and weaknesses.

For the sake of legibility, we omit the third field in the transition labels.
During sexual reproduction genes from two individuals are combined in the offspring that receives genetic material from both parents involving, in diploid populations, recombination among genes. Recombination can break up favourable sets of genes accumulated by selection. Moreover, asexual populations composed by only females can reproduce twice as fast in each generation than sexual populations, because there is no need to produce males for ongoing reproduction.

Despite its considerable cost, sexual reproduction it is still by far the most frequent mode of reproduction in vertebrates. Asexual reproduction has only been described in less than 0.1% of vertebrate species. In general, it is assumed that sexually reproducing populations harbour more genetic variation than asexually reproducing populations, and a high level of genetic variation allows perpetual adaptation to changing environments.

Particularly, populations in heterogeneous habitats, threatened by various parasites or under strong competition, have been shown to have greater genetic variation. For the above reasons a variety of species (essentially among invertebrates) adopted a mixed strategy which tries to combine the advantages of both methods [24].

In this example we model a simple organism able to reproduce either sexually or asexually. We consider the individual of the species as diploid with only a locus (gene), thus each genotype is composed by a pair \((a_1, a_2)\) of alleles which the two chromosomes have for the gene. Moreover, we consider the sex of individuals, thus each of them is represented by a pair of alleles together with the symbol, \(f\) or \(m\), of the sex. The rules controlling the evolution of the population are reproduction rules, either sexual reproduction rules or asexual, and death rules. Each rule has an inhibitor; when the inhibitor is present the rule cannot be applied.

Consider a set of alleles (values for the single gene) of \(k\) elements \(\{v_1, v_2, \ldots, v_k\}\). In the following \(a_1, a_2, a_3, a_4\) belong to \(\{v_1, v_2, \ldots, v_k\}\). The reproduction rules are the following:

1. \(a_1a_2m \ a_3a_4f \rightarrow a_1a_2m \ a_3a_4f \ \neg \text{no_sex_repr} \ (i \in \{1, 2\}, j \in \{3, 4\}, s \in \{m, f\})\)
2. \(a_1a_2f \rightarrow a_1a_2f \ \neg \text{no_asex_repr}\)
Note that each rule has its inhibitor \texttt{no\_sex\_repr} or \texttt{no\_asex\_repr}. The death rules are the following:

3. \(a_1a_2m \rightarrow \lambda \mid \neg\text{no\_male\_death}\)
4. \(a_1a_2f \rightarrow \lambda \mid \neg\text{no\_female\_death}\)

We add also rules for females and males which simply survive, without reproducing or dying:

5. \(a_1a_2m \rightarrow a_1a_2m \mid \neg\text{no\_male\_life}\)
6. \(a_1a_2f \rightarrow a_1a_2f \mid \neg\text{no\_female\_life}\)

Finally we use a rule for sending out individuals from the membrane in which the population evolved. This rule is \textit{promoted} by the promoter \texttt{send\_out}:

7. \(a_1a_2m \rightarrow (a_1a_2m, \text{out}) \mid \text{send\_out}\)
8. \(a_1a_2f \rightarrow (a_1a_2f, \text{out}) \mid \text{send\_out}\)

In the following examples, we consider the membrane system defined above as the system under test. In each example we define a specific observer that is meant to test the evolution forcing certain situations. The observer controls the system by sending into it both the initial individuals and the promoters/inhibitors for constraining the population dynamics.

\textbf{Example 1.} Let us consider a population in which the possible alleles for the single locus are \{0, 1\} and an initial population composed by four males of genotype (00) and four females of genotype (01). Let us control the population dynamics by inhibiting both the sexual reproduction and the death of females. The observer we define, using the \texttt{must} version, is able to analyse the following property of the system: “After two time units, no female in the population can differ from the initial ones, and the number of such females is greater than or equal to the initial female number.”

Note that naturally the tests express quantitative aspects, both on time and on numbers of individuals.

Assume that membrane 1, i.e. the observer, initially contains the element \texttt{a} and that \texttt{Inh} is the set of all inhibitors for constraining the population dynamics.

\begin{align*}
1. \quad & a \rightarrow 1 ((00m)^4, (01f)^4, \texttt{no\_sex\_repr, no\_female\_death},\ \text{in}_2) \\
2. \quad & 1 \rightarrow 2 \\
3. \quad & 2 \rightarrow 3 (\texttt{Inh} \cup \{\texttt{send\_out}\}, \ \text{in}_2) \\
4. \quad & 3 \rightarrow 4 \\
5. \quad & a_1a_2f \rightarrow \texttt{fail} \quad (a_1 \neq 0 \lor a_1 \neq 1) \\
6. \quad & 4 \rightarrow 5 \\
7. \quad & 5 (01f)^4 \rightarrow (\omega, \ \text{out}) \mid \neg\texttt{fail}
\end{align*}

Rule 1 sends into the membrane under test the initial population (four males and four females) and the inhibitors for sexual reproduction and death of females. Rule 2 waits for a time unit, and, after that, Rule 3 sends, during the second step of the populations evolution, all the inhibitors together with the promoter \texttt{send\_out}. Rule 4 waits for a time unit to allow the inner membrane to send out all the individuals. Afterwards, Rule 5 produces a \texttt{fail} if a female different from the initial ones is present. At the same time Rule 6 increases the counter. Finally, Rule 7 sends out the \texttt{\omega} symbol only if \texttt{fail} is absent.

\textbf{Example 2.} Consider the initial population of Example 1 under the same conditions. Again using a \texttt{must} test, we consider the following property: “Given any \(k \in \mathbb{N}\), we can check that it is not possible, in \(n\) time units, for all \(n \in [1, k]\), to produce a female different from the initial ones.” Note that this
property is not a for-all statement: we count until the given k for checking it. This is of course weaker than checking the for-all statement.

The rules in membrane 1 are the following:

1.0. \( a \rightarrow 1 \ block ((00m)^4, (01f)^4, \text{no}_\text{sex}_\text{repr}, \text{no}_\text{female}_\text{death}, \text{in}_2) \)
2.0. \( 1 \rightarrow 2 \)
3.0. \( 2 \rightarrow 3 \)
   ... 
\( k_0. \) \( k - 1 \rightarrow k \)
\( (k + 1)_0. \) \( \text{block} \rightarrow \text{block} |_{-k} \)
\( (k + 2)_0. \) \( \text{block} \rightarrow 1' (\text{Inh} \cup \{\text{send}_\text{out}\}, \text{in}_2) \)
1'0. \( 1' \rightarrow 2' \)
\( 2'_0. \) \( a_1a_2f \rightarrow \text{fail} \quad (a_1 \neq 0 \lor a_1 \neq 1) \)
3'0. \( 2' \rightarrow 3' \)
4'0. \( 3' \rightarrow (\omega, \text{out}) |_{-\text{fail}} \)

Rules from \( 2_0 \) to \( k_0 \) increase the counter until \( k \). At each time unit either Rule \( (k + 2)_0 \) can be executed, stopping the evolution of the population, or Rule \( (k + 1)_0 \) can be fired, allowing the population to evolve for one more step. Rule \( (k + 1)_0 \) is inhibited by \( k \), thus when the counter reaches \( k \) the evolution must terminate. Rules from \( 1'_0 \) to \( 4'_0 \) produce a \( \omega \) if and only if, in the final populations there are only females equal to the initial ones.

Example 3. Consider the initial population of Example 1. This time let us consider a may test expressing the following: “Without initial conditions it is possible to have recombination (offspring with different genotypes with respect to the initial population) after \( k \) steps.”

The rules are the following:

1.0. \( a \rightarrow 1 \ block ((00m)^4, (01f)^4, \text{in}_2) \)
2.0. \( 1 \rightarrow 2 \)
3.0. \( 2 \rightarrow 3 \)
   ... 
\( k_0. \) \( k - 1 \rightarrow k \)
\( (k + 1)_0. \) \( k \rightarrow 1' (\text{Inh} \cup \{\text{send}_\text{out}\}, \text{in}_2) \)
1'0. \( 1' \rightarrow 2' \)
\( 2'_0. \) \( a_1a_2f \rightarrow \omega, \text{out} \quad (a_1 \neq 0 \lor a_1 \neq 1) \)
\( 2'_0. \) \( a_1a_2m \rightarrow \omega, \text{out} \quad (a_1 \neq 0 \lor a_1 \neq 1) \)

Example 4. Consider again the initial population of Example 1. Let us define a may test expressing: “By allowing only the asexual reproduction for \( k \) steps, and then allowing only the sexual reproduction for the following \( k \) steps, it is possible to have recombination in the final population.”

For this example we need the concept of antidote. An antidote is a symbol able to remove the effect of an inhibitor; usually for an inhibitor \( x \), the antidote is denoted by \( \text{anti}_x \). The effect of an antidote is described by particular rules, the antidote rules, which have the form \( \text{anti}_x \ x \rightarrow \lambda \). In this example we assume that, in the membrane under test, there are the antidote rules for the inhibitors no\_sex\_repr and
no_sex_repr. The rules are the following:

\[\begin{align*}
1_0. & \quad a \rightarrow 1 \text{ block } ((00m)^4,(01f)^4,\text{no_sex_repr}, \text{in}_2) \\
2_0. & \quad 1 \rightarrow 2 \\
\cdots & \\
{\sum}. & \quad k - 1 \rightarrow k \text{ (anti_no_sex_repr,no_asex_repr, in}_2) \\
{k+1}_0. & \quad k \rightarrow k + 1 \\
(2k)_{0}. & \quad 2k - 1 \rightarrow 2' \text{ (Inh} \cup \{\text{send_out}\}, \text{in}_2) \\
1'_{0}. & \quad 1' \rightarrow 2' \\
2'_0. & \quad a_1a_2f \rightarrow (\omega, \text{out}) \quad (a_1 \neq 0 \lor a_1 \neq 1) \\
2'_0. & \quad a_1a_2m \rightarrow (\omega, \text{out}) \quad (a_1 \neq 0 \lor a_1 \neq 1)
\end{align*}\]

5 Conclusions

The testing machinery defined in [13] and the P Algebra proposed in [4] inspired us a suitable Process Algebra-based testing environment for P Systems. On the one hand, the new testing environment shares with the original one the concepts of observer, running test, successful and unsuccessful computation, testing preorders/equivalences, allowing us to define qualitative system properties. On the other hand, differently from the original one, it results to be suitable also to express quantitative aspects. Such a feature puts in evidence an expected high expressive power of the framework itself, which needs to be formally studied.

The natural continuation of this work is to find finite decidable characterizations of testing equivalence of finite state P Algebra terms in order to perform verification by comparing a system with its expected behaviour. Moreover, we plan to extend the testing environment also studying a suitable version of fair testing semantics for P Systems, as well as rephrasing the testing environment for Spatial P Systems [5], with the aim of expressing quantitative properties involving spatial information, being crucial in the biological (and not only) domain.

On the biological side, we intend to show as future work the potentials of the testing framework of having a practical impact, for instance on planning both in-silico and wet-lab experiments.

References

[1] J. Aguado, T. Balanescu, T. Cowling, M. Gheorghe, M. Holcombe, and F. Ipati. P Systems with Replicated Rewriting and Stream X-machines (Eilenberg Machines). Fundam. Inf., 49:17–33, January 2002.
[2] J. Ahmad, G. Berrot, J. Comet, D. Lime, and O. Roux. Hybrid Modelling and Dynamical Analysis of Gene Regulatory Networks with Delays. ComPlexUs, 3(4):231–251, 2006.
[3] O. Andrei, G. Ciobanu, and D. Lucanu. A Rewriting Logic Framework for Operational Semantics of Membrane Systems. Theor. Comput. Sci., 373(3):163–181, 2007.
[4] R. Barbuti, A. Maggiolo-Schettini, P. Milazzo, and D. Gruska. A Notion of Biological Diagnosability Inspired by the Notion of Opacity in Systems Security. Fundamenta Informaticae, 102(1):19–34, 2010.
[5] R. Barbuti, A. Maggiolo-Schettini, P. Milazzo, G. Pardini, and L. Tesei. Spatial P Systems. Natural Computing, 2010. Received: 26 October 2009 Accepted: 24 February 2010 Published online: 24 March 2010.
[6] R. Barbuti, A. Maggiolo-Schettini, P. Milazzo, and S. Tini. A P Systems Flat Form Preserving Step-by-step Behaviour. Fundamenta Informaticae, 87(1):1–34, 2008.
[7] R. Barbuti, A. Maggiolo-Schettini, P. Milazzo, and S. Tini. Compositional Semantics and Behavioral Equivalences for P Systems. Theoretical Computer Science, 395(1):77–100, 2008.
[8] L. Bianco and V. Manca. Encoding-Decoding Transitional Systems for Classes of P systems. In Proc. of WMC 2005, 6th International Workshop on Membrane Computing, number 3850 in LNCS, pages 134–143. Springer-Verlag, 2006.

[9] M. Boreale and R. Pugliese. Basic Observables for Processes. Information and Computation, 149(1):77–98, 1999.

[10] P. Bottoni, C. Martín-Vide, G. Páun, and G. Rozenberg. Membrane Systems with Promoters/Inhibitors. Acta Informatica, 38(10):695–720, 2002.

[11] G. Ciobanu, M. J. Pérez-Jiménez, and G. Paun, editors. Applications of Membrane Computing. Natural Computing Series. Springer, 2006.

[12] Z. Dang, C. Li, O. H. Ibarra, and G. Xie. On the Decidability of Model-checking for P Systems. J. Autom. Lang. Comb., 11:279–298, January 2006.

[13] R. De Nicola and M. Hennessy. Testing Equivalences for Processes. Theoretical Computer Science, 34(1-2):83–133, 1984.

[14] M. Gheorghe and F. Ipate. On Testing P Systems. In D. Corne, P. Frisco, G. Paun, G. Rozenberg, and A. Salomaa, editors, Membrane Computing, volume 5391 of Lecture Notes in Computer Science, pages 204–216. 2009.

[15] M. Gheorghe and F. Ipate. Testing Based on P Systems - An Overview. In Proceedings of the 11th International Conference on Membrane Computing, CMC’10, pages 3–6, Berlin, Heidelberg, 2010. Springer-Verlag.

[16] F. Ipate and M. Gheorghe. Testing Non-deterministic Stream X-machine Models and P systems. Electron. Notes Theor. Comput. Sci., 227:113–126, January 2009.

[17] F. Ipate, M. Gheorghe, and R. Lefticaru. Test Generation From P systems Using Model Checking. Journal of Logic and Algebraic Programming, 79(6):350–362, 2010. Membrane computing and programming.

[18] F. Ipate, R. Lefticaru, and C. Tudose. Formal Verification of P Systems using SPIN. International Journal of Foundations of Computer Science, 22(1):133–142, 2011.

[19] V. Natarajan and R. Cleaveland. Divergence and Fair Testing. In Proceedings of the 22nd International Colloquium on Automata, Languages and Programming, ICALP ’95, pages 648–659, 1995.

[20] G. Páun. Membrane Computing. An Introduction. Springer-Verlag, 2002.

[21] G. D. Plotkin. A Structural Approach to Operational Semantics. J. Log. Algebr. Program., 60-61:17–139, 2004.

[22] G. Páun and G. Rozenberg. A Guide to Membrane Computing. Theoretical Computer Science, 287(1):73–100, 2002.

[23] A. Rensink and W. Vogler. Fair testing.

[24] I. Schönh, K. Martens, and P. van Dijk. Lost Sex: The Evolutionary Biology of Parthenogenesis. Springer Verlag, 2009.