# On Catalytic P Systems with One Catalyst

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**Summary.** In this paper we address the possibility of studying the computational capabilities of catalytic P systems with one catalyst by the means of iterated finite state transducers. We also give a normal form for catalytic P systems.

### 1 Introduction

P systems are a computational model introduced by G. Păun in [4]. One of the basic variant considered there was P systems with catalysts and priorities; these systems where shown to be computationally universal. In [2], Sosík and Freund proved that priorities among the rules can be discarded from the model without any loss of computational power. Moreover, it was shown that for extended P systems only one membrane and two catalysts are enough for reaching computational universality. However, the computational power for P systems with only one catalyst was not established. The present paper characterize these systems in terms of iterated finite state transducers hence it converts an open problem from P system framework to an open problem from string rewriting theory. Additionally, a normal form for catalytic P systems is presented.

## 2 Preliminaries

We assume the reader is acquainted with the basic notions and notations from the formal language theory (see [3] for more details). Here we only recall the definitions and the results which are useful for the present work.

If FL is a family of languages, then NFL denotes the family of length sets of languages in FL. We denote by REG, CF, REC, and RE the family of regular, context-free, recursive, and recursively enumerable languages, respectively. It is known that  $NREG = NCF \subsetneq NREC \subsetneq NRE$ .

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#### 2.1 Iterated Finite State Transducers

An iterated (finite state) sequential transducer (IFT) is a construct  $\gamma = (K, V, q_0, a_0, F, P)$ , where K is a finite set of states, V is a finite set of symbols (the alphabet of  $\gamma$ ),  $K \cap V = \emptyset$ ,  $q_0 \in K$  is the initial state,  $a_0 \in V$  is the starting symbol,  $F \subseteq K$  is the set of final states, and P is a finite set of transition rules of the form  $q_a \to xp$ , for  $q, p \in K$ ,  $a \in V$ , and  $x \in V^*$ .

For  $q, p \in K$  and  $u, v, x \in V^*$ ,  $a \in V$ , a *direct transition* step of  $\gamma$  is defined as  $uqav \vdash uxpv$  if and only if  $qa \rightarrow xp \in P$ . The reflexive and transitive closure of the relation  $\vdash$  is denoted by  $\vdash^*$ . In general, for  $\alpha, \beta \in V^*$  we say that  $\alpha$  *derives* into  $\beta$  and we write  $\alpha \Longrightarrow \beta$ , if and only if  $q_0 \alpha \vdash^* \beta p$  for some  $p \in K$ . By  $\Longrightarrow^*$  we denote the reflexive transitive closure of  $\Longrightarrow$ . If  $q_0 \alpha \vdash^* \beta p$  such that  $p \in F$ , then we write  $\alpha \Longrightarrow_f \beta$ .

The language generated by  $\gamma$  is  $L(\gamma) = \{\beta \in V \mid a_0 \Longrightarrow^* \alpha \Longrightarrow_f \beta$ , for some  $\alpha \in V^*\}$ .

If for each pair  $(q, a) \in K \times V$ , there is at most one transition rule  $qa \vdash xp \in P$ , then  $\gamma$  is called *deterministic* (otherwise, it is called *nondeterministic*). The family of languages generated by nondeterministic IFTs with at most  $n \geq 1$  states is denoted by  $IFT_n$ . It is known from [1] that  $CF \subset IFT_2 \subseteq IFT_3 \subseteq IFT_4 = RE$ . Moreover, there are non-semilinear languages belonging to  $IFT_2$ , and there are non-recursive languages belonging to  $IFT_3$ . Consequently, if we denote by  $NIFT_n$ ,  $n \geq 1$ , the family of length sets of languages from  $IFT_n$ , then we have that  $NREG = NCF \subsetneq NIFT_2 \subseteq NIFT_3 \subseteq NIFT_4 = NRE$ .

#### 2.2 Membrane Systems

A catalytic P system of degree  $m \ge 1$  is a construct

$$\Pi = (O, C, \mu, w_1, \dots, w_m, R_1, \dots, R_m, i_0)$$

where

• O is an alphabet of *objects*;

•  $C \subseteq O$  is the set of *catalysts*;

•  $\mu$  is a hierarchical tree structure of  $m \ge 1$  uniquely labelled membranes (which delimit the regions of  $\Pi$ ); typically, the set of labels is  $\{1, \ldots, m\}$ ;

•  $w_i \in O^*$ , for  $1 \le i \le m$ , are the multisets of objects initially present in the m regions of  $\mu$ ;

•  $R_i$ ,  $1 \leq i \leq m$ , are finite sets of evolution rules; these rules can be noncooperative  $a \to v$  or catalytic  $ca \to cv$ , where  $a \in O \setminus C$ ,  $v \in ((O \setminus C) \times \{here, out, in\})^*$ , and  $c \in C$ ;

•  $i_0 \in \{1, \ldots, m\}$  is the label of the *output region* of  $\Pi$ .

A configuration of  $\Pi$  is a vector  $C = (\alpha_1, \ldots, \alpha_m)$ , where  $\alpha_i \in O^*$ ,  $1 \le i \le m$ , is a multiset of objects present in the region *i* of  $\Pi$ . The vector  $C_0 = (w_1, \ldots, w_m)$ is the *initial configuration* of  $\Pi$ . Starting from the initial configuration and always applying in all membranes a maximal multiset of evolution rules in parallel, one gets a sequence of consecutive configurations. By  $\Rightarrow$  is denoted the *transition* between two consecutive configurations. A sequence (finite or infinite) of transitions starting from  $C_0$  represents a *computation* of  $\Pi$ . A computation of  $\Pi$  is a halting one if no rules can be applied to the last configuration (the *halting configuration*). The result of a halting computation is the number of objects from O contained in the output region  $i_0$ , in the halting configuration. A non-halting computation yields no result. By collecting the results of all possible halting computations of a given P system  $\Pi$ , one gets  $N(\Pi)$  – the set of all natural numbers generated by  $\Pi$ . The family of all sets of numbers computed by catalytic P systems with at most mmembranes and k catalysts is denoted by  $NOP_m(cat_k)$ . The above definition can be relaxed such that in a halting configuration one counts only the symbols from a given alphabet  $\Sigma \subseteq O$ . In particular, one can consider  $\Sigma = O \setminus C$ ; correspondingly, the family of all sets of numbers computed by such particular P systems will be denoted by  $NO_{-C}P_m(cat_k)$ .

It is known (see [7], for instance) that  $NO_{-C}P_m(cat_k) = NO_{-C}P_1(cat_k)$ . Moreover, in [2] it is shown that  $NO_{-C}P_1(cat_2) = NRE$ .

### 3 A Normal Form for P Systems with Catalysts

The following result states that any catalytic P system is equivalent with a catalytic P system having a restriction on the form of the rules.

**Theorem 1.** For any P system  $\Pi$  with catalysts there exists an equivalent P system  $\overline{\Pi}$  with one region and whose rules are of the form  $a \to \alpha$ , with  $|\alpha| \leq 2$ , or  $ca \to c\beta$ , with  $|\beta| \leq 1$ .

*Proof.* As we already stated in Section 2.2, for any P system with catalysts and n > 1 membranes one can construct an equivalent P system with the same number of catalysts and one membrane. Consequently, without loss of generality, we might assume that  $\Pi$  has only one membrane, that is  $\Pi = (O, C, []_1, w_1, R_1, i_0)$ .

Let  $O \setminus C = \{a_1, a_2, \ldots, a_p\}$  and let  $m = max\{|\alpha| \mid a \to \alpha \in R_1 \text{ or } ca \to c\alpha \in R_1\}$ . In addition, assume for our convenience that the rules of  $\Pi$  are labeled in an unique manner with numbers from the set  $\{1, \ldots, card(R_1)\}$ .

Then one can construct an equivalent P system  $\overline{\Pi} = (\overline{O}, C, []_1, w_1, \overline{R_1}, i_0)$ where

$$\overline{O} = O \cup \{a_{(i,j)} \mid 1 \le i \le p, 1 \le j \le m\}$$
$$\cup \{X_{(i,j)} \mid i : a \to \alpha_i \in R_1, 1 \le j \le m-2\}.$$

The set  $\overline{R_1}$  is defined as follows (for the simplicity of the explanations, we will only consider the rules in  $\overline{R_1}$  that are useful for simulating a non-cooperative rule from  $R_1$ ; the rules corresponding to a catalytic rule are defined similarly, therefore we will not present them here). Let  $i : a \to a_{j_1} a_{j_2} \dots a_{j_k} \in R_1$  and let m - k = t. Then we add to  $\overline{R_1}$  the rules:

$$a \to X_{(i,1)}$$
(1)  

$$X_{(i,1)} \to X_{(i,2)}$$
...  

$$X_{(i,t-1)} \to X_{(i,t)}$$
(2)  

$$X_{(i,t+1)} \to a_{(j_1,k-1)}X_{(i,t+1)}$$
(2)  

$$X_{(i,t+1)} \to a_{(j_2,k-2)}X_{(i,t+2)}$$
...  

$$X_{(i,t+k-3)} \to a_{(j_{k-2},2)}X_{(i,t+k-2)}$$
...  

$$X_{(i,t+k-2)} \to a_{(j_{k-1},1)}a_{(j_k,1)}$$
(3)  

$$a_{(i,m-1)} \to a_{(i,m-2)}$$
...  

$$a_{(i,1)} \to a_{i}$$
(3)

The proof is based on the existence of the universal global clock that governs the functioning of the P system (the clock marks equal time units for the whole system, hence synchronization is possible). While trying to simulate the application of an arbitrary non-cooperative rule with several rules of type  $a \to \alpha$ , with  $|\alpha| \leq 2$ , one has to accomplish two conditions. Firstly, one has to guarantee that all the objects from  $\alpha$  will eventually be produced. Secondly, these objects must be produced at the "proper" time: all of them in the same moment (a local synchronization) and according with the simulation of other rules that were started at the same time with  $a \to \alpha$  (a global synchronization).

Consequently, the rules presented above are grouped according with their function in the simulation. The first group represents a set of "delaying" rules (they are used while simulating the rules with a shorter right hand side in order to synchronize their executions with those that have the longest right hand side). These rules are "chained", hence, staring from an object a, an object  $X_{(i,t)}$  is produced in exactly t computational steps. The second group is responsible for producing in consecutive computational steps the objects  $a_{(j_1,k-1)}, a_{(j_2,k-2)}, \ldots, a_{(j_{k-1},1)}, a_{(j_k,1)}$ (in order of their production, the last two being produced in the same time). For an object  $a_{(i,l)}$  in this sequence, the index l represents the number of computational steps that  $\overline{\Pi}$  will perform, starting from its production and until the object  $a_i$  is produced (see the third group of rules). Finally, one can remark that the objects  $a_{j_1}, a_{j_2}, \ldots, a_{j_k}$  are produced in the same computational step by  $\overline{\Pi}$  (while simulating the rule  $i: a \to a_{j_1} a_{j_2} \dots a_{j_k} \in R_1$ ). Moreover, all the other rules from  $\Pi$  that stated at the same moment as  $i: a \to a_{j_1}a_{j_2}\ldots a_{j_k}$ , are simulated in the same manner by  $\overline{\Pi}$  and their output is produced in the same computational step as mentioned above. Consequently  $\overline{\Pi}$  correctly simulates any computation of  $\Pi$ , hence the theorem holds true.

### 4 Catalytic P Systems with One Catalyst and IFTs

In what follows we prove that the family of sets of numbers computed by catalytic P systems with only one catalyst is included in the family of the length sets of the languages generated by iterated finite state transducers with at most 3 states.

**Theorem 2.**  $NIFT_3 \supseteq NOP_1(cat_1)$ .

*Proof.* Given an arbitrary catalytic P system  $\Pi = (O, C, w_1, R_1, i_1)$  such that  $C = \{c\}$ , then one can construct an iterated finite transducer  $\gamma = (K, V, q_0, a_0, F, P)$  which simulates  $\Pi$  as follows.

Without loss of generality we assume that the initial configuration of  $\Pi$  is  $w_1 = ca_0$ .

Let  $w = a_1 a_2 \dots a_m$  be a string. We denote by

$$Perm(w) = \{a_{i_1}a_{i_2}\dots a_{i_m} \mid 1 \le i_j \le m, 1 \le j \le m, \text{ with } i_j \ne i_l, 1 \le j, l \le m\}$$

the set of all permutations of string w, i.e., the set of all strings that can be obtained from w by changing the order of symbols.

In addition, let us consider the following sets of objects from O:  $X = \{A \in O \mid (\exists) A \to \alpha \in R_1 \text{ and } (\nexists) cA \to c\beta \in R_1\};$   $Y = \{A \in O \mid (\exists) A \to \alpha \in R_1 \text{ and } cA \to c\beta \in R_1\};$   $Z = \{A \in O \mid (\exists) cA \to c\alpha \in R_1 \text{ and } (\nexists) A \to \beta \in R_1\};$   $T = \{A \in O \mid (\nexists) A \to \alpha \in R_1 \text{ and } (\nexists) cA \to c\beta \in R_1\}.$ One can remark that  $O = X \cup Y \cup Z \cup T \cup \{c\}.$ Based on the above settings the IFT  $\gamma$  is defined as follows:

$$K = \{q_0, q_1, q_2\} V = O \setminus \{c\}, F = \{q_0\},$$

and the set of rules P is constructed in the following manner:

• for any  $a \in T$  we add to P the rule  $q_0 a \rightarrow a q_0$ ;

• for any  $a \in X \cup Y$  and  $a \to \alpha \in R_1$  we add to P the rules  $q_0 a \to \overline{\alpha} q_1$ , where  $\overline{\alpha} \in \operatorname{Perm}(\alpha)$ ;

• for any  $a \in T$  we add to P the rule  $q_1 a \to a q_1$ ;

• for any  $a \in X \cup Y$  and  $a \to \alpha \in R_1$  we add to P the rules  $q_1 a \to \overline{\alpha} q_1$ , where  $\overline{\alpha} \in \operatorname{Perm}(\alpha)$ ;

• for any  $a \in Y \cup Z$  and  $ca \to c\alpha \in R_1$  we add to P the rules  $q_1a \to \overline{\alpha}q_2$ , where  $\overline{\alpha} \in \operatorname{Perm}(\alpha)$ ;

• for any  $a \in T \cup Z$  we add to P the rule  $q_2a \to aq_2$ ;

• for any  $a \in X \cup Y$  and  $a \to \alpha \in R_1$  we add to P the rules  $q_2 a \to \overline{\alpha} q_2$ , where  $\overline{\alpha} \in \operatorname{Perm}(\alpha)$ ;

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• for any  $a \in Y \cup Z$  and  $ca \to c\alpha \in R_1$  we add to P the rules  $q_0 a \to \overline{\alpha} q_2$ , where  $\overline{\alpha} \in \operatorname{Perm}(\alpha)$ .

The construction was designed such that each string processed by  $\gamma$  during its computation will correspond to a configuration of  $\Pi$ . Moreover, one iteration of  $\gamma$  simulates the maximal parallel applications of the rules of  $\Pi$ .

If the current string (say w) processed by the IFT is composed only by the symbols from T, then  $\gamma$  remains in  $q_0 \in F$  and stops, accepting the string. This situation corresponds to the halting configuration of  $\Pi$  (that is,  $\Pi$  contains in its region the multiset cw and no rules can be further applied).

In case w contains symbols form  $X \cup Y \cup Z$ , then  $\gamma$  starts the simulation of the maximal parallel applications of the rules of  $\Pi$ . Since  $\gamma$  processes strings at each iteration, then the simulation of  $\Pi$  has to accomplish the following task: all the symbols which are the subject of a rule of  $\Pi$  have to be processed also by  $\gamma$ . Recall that  $\gamma$  processes strings and in these strings there might be symbols from T (which are not the subject of any rule) in any position. Consequently, one has be sure that any symbol in a configuration of  $\Pi$  that is a subject of a rule (non-cooperative or catalytic) has to have the opportunity to be rewritten in the corresponding string processed by  $\gamma$  (by the corresponding rule). This is why,  $\gamma$  uses the rules  $q_i a \to aq_i$  for  $q_i \in Q$ ,  $1 \leq i \leq 3$ , and  $a \in T$  (that is, while processing the string,  $\gamma$  "skips" all the symbols that are not the subject of any rule).

In one iteration of  $\gamma$  one can apply at most once a rule corresponding to a catalytic rule of  $\Pi$  (recall that the P system functioning semantics define such behaviour). More precisely, assuming that w is the current processed string, we have

- either  $\gamma$  is in state  $q_0$  and executes a rule of type  $q_0 a \to \overline{\alpha} q_2$  for  $q_1, q_2 \in Q$ ,  $a \in Y \cup Z$ ,  $ca \to c\alpha \in R_1$ , and  $\overline{\alpha} \in \operatorname{Perm}(\alpha)$ . This situation occurs when  $\gamma$  processes  $w = w_1 a w_2$ ,  $w_1 \in T^*$ , and  $w_2 \in (X \cup Y \cup Z \cup T)^*$  (w has the prefix  $w_1$  composed only by symbols from T, followed by the symbol  $a \in Y \cup Z$  that triggers the simulation of the catalytic rule; the symbols from  $w_2$  that belong to  $X \cup Y$  will trigger only the simulation of the non-cooperative rules).
- either  $\gamma$  is in state  $q_1$  and executes a rule of type  $q_1a \to \overline{\alpha}q_2$  for  $q_1, q_2 \in Q$ ,  $a \in Y \cup Z$ ,  $ca \to c\alpha \in R_1$ , and  $\overline{\alpha} \in \operatorname{Perm}(\alpha)$ ). This situation occurs when  $\gamma$  processes  $w = w_1 a w_2$ , where  $w_1$  is described by the regular expression  $T^*(X|Y)(X|Y|T)^*$ ,  $w_2 \in (X \cup Y \cup Z \cup T)^*$  (the symbols from  $w_1$  and  $w_2$  that belong to  $X \cup Y$  will trigger only the simulation of the non-cooperative rules).

One can also remark that if a configuration w of  $\Pi$  contains at least one object  $a \in Z$ , then in the current computational step a catalytic rule will be executed (because of the maximal parallel applications of the rules); in contrary, if w does not contain any symbol  $a \in Z$  then it is not guaranteed that a catalytic rule will be executed (even if w contains symbols from Y, then, because of the nondeterminism, it might happen that all the rules selected for application are non-cooperative). On the other hand,  $\gamma$  simulates  $\Pi$  by processing strings (hence the order of symbols is precisely defined). The design of  $\gamma$  guarantees that, if applicable, a rule corresponding to a catalytic rule of  $\Pi$  is executed at most once. The only issue that could appear regards the presence of multiple symbols from  $Y \cup Z$  in the current string processed by  $\gamma$  (in order to perform a correct simulation, one has to be sure that any of these symbols has a "chance" to be rewritten). This is why for any rule  $a \to \alpha \in R_1$  or  $ca \to c\alpha \in R_1$ , the IFT  $\gamma$  will use for the simulation a set of rules of the type  $qa \to p \operatorname{Perm}(\alpha)$ .

Based on the above theorem, the following result holds true.

**Corollary 1.** If  $NIFT_3 \subset NRE$  then  $NOP_1(cat_1) \subset NRE$ 

### 5 Conclusions

In this paper we gave a normal-form theorem for catalytic P systems. We also investigated the relation between P systems with one catalyst and iterated finite transducers. This last topic is of a particular interest because it converts an open problem from the P system framework to an open problem from the string rewriting theory. In addition, the simplicity of the construction gives hopes for solving an open problem stated from the introduction of P systems.

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