

# Self-Evolution in a Constructive Binary String System

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**Abstract** We examine the qualitative dynamics of a catalytic self-organizing system of binary strings that is inspired by the chemical information processing metaphor. A string is interpreted in two different ways: either (a) as raw data or (b) as a machine that is able to process another string as data in order to produce a third one. This article focuses on the phenomena of evolution whose appearance is notable because no explicit mutation, recombination, or artificial selection operators are introduced. We call the system self-evolving because every variation is performed by the objects themselves in their machine form.

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## Keywords

artificial chemistry, autocatalytic reaction system, molecular computing, prebiotic evolution, self-organization, self-programming

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## 1 Introduction

In recent years computer simulations have been used to study the phenomena of life, not by simulating life as it is (weak AL) but by instantiating life as it could be (strong AL) [22]. AL systems have been used, for instance, for studying questions concerning the emergence and quality of organizations [13, 19, 26], the process of diversification [27], the origin of replicators [25], or morphological evolution [21].

An important property of most strong AL systems is that they contain the ability for self-reference. For instance, Ray's Tierra organisms are able to read, copy, and modify their own code [27]. In Fontana's algorithmic chemistry every object is a character string able to process other objects by using the lambda-calculus that maps the character string into an (active) function [13]. The dualism inherent in those systems can be traced back to Gödel [15] who defined a mapping of mathematical statements into natural numbers that allowed self-reference, to Turing's universal machine [31], and to von Neumann's stored program computer [7].

The system discussed here is inspired by a chemical reaction dynamics, where molecules collide and interact to create new molecules forming metabolic networks. This contribution tries to give insight into the origin and evolution of these networks [2, 3]. Related work [4, 13, 20, 23, 30] focuses on the static structure of the emerged networks. Here, we will concentrate on dynamic phenomena, especially on the emergence of prebiotic evolution [2, 17]. Evolutionary phenomena can be observed, although no explicit fitness, mutation, recombination, or selection operators are used. Variations are only performed by the objects (molecules, binary strings) themselves when they act in machine form.

It seems worthwhile to start our considerations with a short note on self-organization and what we mean by "self-evolution" in the context of this article. *Self organization* is a process where a system is organized while the components directing this process are part of the system. This kind of view of "self-organization" implies a) every organization phenomenon becomes a result of self-organization provided we enlarge the system

boundaries; and b) self-organization need not to be directed, for example, toward higher complexity.

When transforming this linguistic abstract description of self-organization into the language of computer science it seems to be straightforward to represent an organizer as a program and the data a program is working on as the target for organization. To close the loop inherent in self-organization we need a mapping from data to machines or vice versa, yielding the dualism of the system components. The information residing in the system should either act as *active* machines or be processed as *passive* data.

The organization process can also be expressed in the terms of mathematics as the application of (active) operators to (passive) data. An example of this approach is [4], where the operands are binary strings. Operators are formed out of binary strings by “folding” them into a matrix.

The term *self-evolution* should refer to an evolutionary process within a population system where the components responsible for the evolutionary behavior are (only) the individuals of the population system itself. Every variation is carried out by the individuals. Selection pressure is generated implicitly through interaction among the individuals and not by external agents. The system must not contain a component that can be identified as a fitness function or global operators performing selection or variation (e.g., crossover).

The outline of this article is as follows: Section 2 gives a general introduction to algorithmic reaction systems. In Section 3 we briefly describe the special reaction mechanisms used for this contribution. Section 4 summarizes methods for investigation and visualization. In Section 5 we discuss four qualitative types of dynamic self-organization phenomena by showing typical simulation results, namely (a) extinction, (b) emergence of an organization structure, (c) exploration/innovation, and (d) evolution.

## 2 Algorithmic Reaction Systems

The system used for the experiments in this article is in general called an *artificial chemistry* or *algorithmic reaction system*, because the interaction between its elements is specified by an algorithm. It consists of the following three components:

1. a set of objects  $S$ : These objects may be abstract symbols [32], character sequences [20], lambda-expressions [13], binary strings [4, 30], numbers [5], or proofs [14].

Here, we use binary strings with a constant length of 32 bits, thus  $S = \{0,1\}^{32}$ . In a basic setting, the soup has no topological structure so that its state can be noted as a concentration vector.

2. a collision or reaction rule: The collision rule defines the interaction between two objects  $s_1, s_2 \in S$ , which may lead to the production of a new object  $s_3 \in S$ . This is denoted as

$$s_1 + s_2 \implies s_3$$

Note that the operator “+” is *not* commutative, in general.

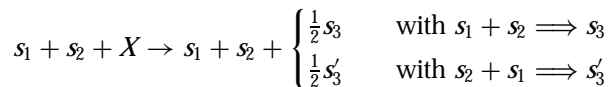
3. an algorithm simulating the reaction vessel: In this contribution we use the following algorithm that simulates a well-stirred tank reactor with mass-action kinetics, which assures that the probability of a collision is proportional to the product of the concentration of the colliding objects (Equation 1, below). The algorithm can also be found with minor modifications in [4, 13, 20, 30].

### 2.1 Reactor Algorithm

1. Select two objects  $s_1, s_2$  from the soup randomly, without removing them.
2. If there exists a reaction  $s_1 + s_2 \implies s_3$  and the filter condition  $f(s_1, s_2, s_3)$  holds, replace a randomly selected object of the soup by  $s_3$ .

The replaced objects form the *dilution flux* of the system. The *filter*  $f$  can be used to block lethal objects and to introduce elastic collisions easily. The term *collision* refers to one execution of the two steps of the reactor algorithm. A collision is called *elastic* if no product  $s_3$  is inserted into the soup. So, a collision of two objects  $s_1, s_2$  is elastic if no product is defined by the reaction rule or if the filter condition prohibits the insertion of the product. An object is said to be *lethal* if it is able to replicate in an unproportionally large number in almost any ensemble configuration [4].

The interaction scheme of the algorithm may be written as a chemical reaction equation [13]:



In other words,  $s_1$  and  $s_2$  are not consumed but act as catalysts of the reaction. The *raw material*  $X$  is used to balance the equation. It does not appear explicitly in our system and could be interpreted as computational resources such as processing time or memory [27]. Reaction systems exhibiting this kind of reaction scheme are able to form *hypercyclic* organizations [10].

To measure the running time of the algorithm we call  $M$  iterations (collisions) a *generation*, where  $M$  is the soup or reactor size. Using “generations” rather than “number of iterations” allows an easier comparison of runs with different soup sizes.

### 2.2 A Model for the Algorithmic Reactor

In a setting corresponding to a well-stirred tank reactor the state of the system can be described by a concentration vector  $\mathbf{x} = (x_1, \dots, x_n)$  with  $x_1 + \dots + x_n = 1$  and  $x_i > 0$ , where  $x_i$  is the concentration of string type  $s_i$ . For a large and constant soup size the behavior of the reactor algorithm is modeled by the *catalytic network equation* [29]:

$$\frac{dx_k}{dt} = \sum_{i=1}^n \sum_{j=1}^n \alpha_{ij}^k x_i x_j - x_k \sum_{i,j,k=1}^n \alpha_{ij}^k x_i x_j \quad k = 1, \dots, n \tag{1}$$

with second-order rate constant  $\alpha_{ij}^k$  for the reaction  $i + j \xrightarrow{\alpha_{ij}^k} k$ . Here the rate constants become

$$\alpha_{ij}^k = \begin{cases} 1 & \text{if } s_1 + s_2 \implies s_3 \text{ and the filter condition } f(s_1, s_2, s_3) \text{ holds,} \\ 0 & \text{otherwise.} \end{cases} \tag{2}$$

Equation 1 becomes the famous *replicator equation* [17, 29] if  $\forall i, j, k \in \{1, \dots, n\} : \alpha_{ij}^k > 0 \implies i = k \vee j = k$  holds.

Simulating systems with large number  $n$  of different objects becomes difficult. For the special case of only a small fraction of all possible objects being present in the reactor, a technique called *meta dynamics* is used elsewhere, where the ordinary differential equation (ODE) system (Equation 1) is treated dynamically and updated under certain conditions [2, 12, 20]. For example, equations are removed when the corresponding

concentrations fall below a given threshold. The advantage of this method is that a potentially infinite number of objects can be handled by still using an ODE framework. Problems arise, however, if the diversity (e.g., the number of different objects in the reactor) becomes high, as will be the case here.

### 3 Special 32-bit Reaction Mechanisms

We will now briefly describe the special reaction mechanisms used for this contribution, namely the AND reaction and the automata reaction.

#### 3.1 AND Reaction

The AND reaction is simply the bitwise logic AND of strings  $s_1$  and  $s_2$ .

$$\forall i \in \{0, 31\} : s_3^{(i)} = s_1^{(i)} \wedge s_2^{(i)}$$

where  $s^{(i)} \in \{0, 1\}$  denotes the  $i$ th bit of string  $s$ . This reaction is used for test and reference, because its behavior can be easily described and understood.

#### 3.2 Automata Reaction

The *automata reaction* is based on a finite state automaton that is a mixture of a Turing machine and a register machine. It has also been inspired by the Typogenetics of Hofstadter [18].

The automata reaction instantiates a deterministic reaction  $s_1 + s_2 \implies s_3$ , where  $s_1, s_2, s_3 \in \{0, 1\}^{32}$ . To calculate the product  $s_3$ , string  $s_1$  is “folded” into an automaton  $A_{s_1}$ , which receives  $s_2$  as input. The construction of  $A_{s_1}$  ensures that the automaton will halt after a bounded finite number of steps. Because  $A_{s_1}$  is a deterministic finite automaton, the automata reaction defines a functions  $\{0, 1\}^{32} \times \{0, 1\}^{32} \implies \{0, 1\}^{32}$ .

Figure 1 shows the structure of the automaton. It contains 32-bit registers, the *IO register* and the *operator register*. At the beginning, operator string  $s_1$  is written into the operator register and operand  $s_2$  into the IO register. The program is generated from  $s_1$  by simply mapping successive 4-bit segments into instructions. Note that two slightly different mappings are used (Figure 1, right). The resulting program is executed sequentially, starting with the first instruction. The automaton halts after the last instruction has been executed or in case of STOP instruction. After halting, the IO register contains the output  $s_3$ . There are no control statements for loops or jumps in the instruction set.

Each 32-bit register has a pointer, referring to a bit location. These are the *IO pointer*, referring to a bit  $b'$  in the IO register and the *operator pointer*, referring to a bit  $b$  in the operator register, respectively. For each collision both pointers are initialized with position 0, as shown in Figure 1. Bit  $b$  and  $b'$  are inputs to the *arithmetic logic unit* (ALU), which is able to perform logic operations on two bits. The ALU result is stored at the IO pointer location, therefore replacing  $b'$ .

The execution of a logic instruction (ID, NOT, OR, AND, EXOR, or EQ) is divided into two steps: a) The ALU processes bit  $b$  and  $b'$  and replaces  $b'$  by its output. b) The pointers are moved one bit left or right, according to the *direction register*. The *move-mode register* determines whether only the IO pointer, only the operator pointer, or both pointers are moved. The move-direction and the move-mode can be toggled by TDIR and TMM, respectively.

The MOV instruction is the most complicated one. Its execution depends on all control registers. If the pattern register is set to *none* and the copy-mode is switched off, the pointers are moved one bit according to the direction and move-mode register.

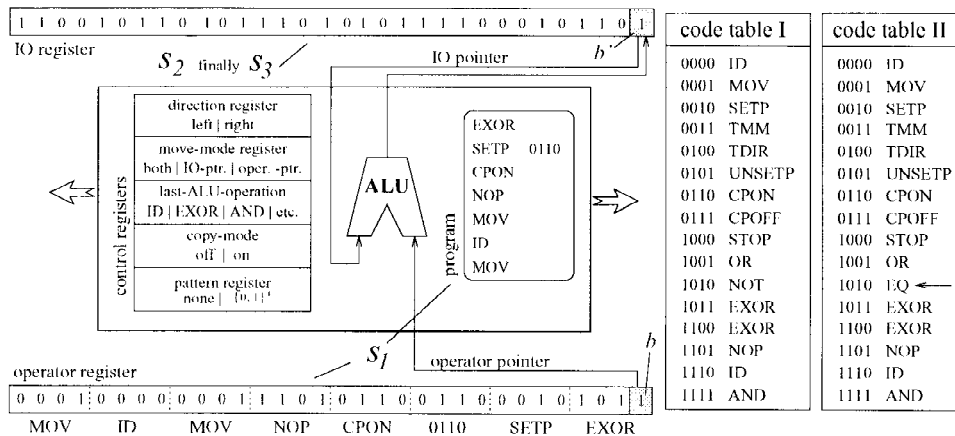


Figure 1. Automaton, resulting from folding  $s_1$ . It carries out the reaction  $s_1 + s_2 \implies s_3$ . String  $s_1$  is written into the operator register and specifies the program. The IO register is initialized with  $s_2$  and contains the result  $s_3$  after running the program.

This is equal to the second step of the execution of a logic instruction. If the pattern register is set to a pattern (a 4-bit bit string) the pointers are moved until the pattern is found in the operator register (max. 32 steps). If the copy-mode is switched on, the last ALU operation is executed before each step. Thus, one MOV instruction may trigger many ALU operations. The pattern register is set by the 8-bit instruction SETP (pattern) and cleared by UNSETP. The copy-mode is switched on and off by CPON and CPOFF, respectively.

Table 1 shows the operation of the automaton for the following example:

$$s_1 = 0001\ 0000\ 0001\ 1101\ 0110\ 0110\ 0010\ 1011$$

$$s_2 = 1100\ 1011\ 0101\ 1101\ 0101\ 1110\ 0010\ 1101$$

$$s_3 = 1100\ 1011\ 0101\ 1101\ 0101\ 0110\ 0000\ 0110$$

In hex notation, this reads:  $101d662b + cb5d5c2d \implies cb5d5606$ . Table 2 shows more examples in hex notation that should give an impression of the static properties of the automata reaction. The first noticeable property is that the structure of the product  $s_3$  is similar to its "parents"  $s_1, s_2$ . This indicates that there is a correlation between  $s_1, s_2$ , and  $s_3$  that is a prerequisite for evolution. Secondly, variations of string sequences appear usually at the edges of a sequence, mostly on the right side and sometimes on the left. Modification of the core bits only is very rare. This is due to the fact that the pointers are always initialized to bit position 0 and the direction register is always set to *left*. Passive replication<sup>1</sup> is common. About 30% of all randomly generated strings are passive replicators. Active self-replication<sup>2</sup> is rare. About 0.004% randomly generated sequences are active replicators. As we will demonstrate later, the system evolves toward active replication, provided the soup is large enough for an explorative or evolutionary behavior.

1 A string  $s_1$  is called a *passive replicator* if  $\forall s_2 \in S: s_1 + s_2 \implies s_2$ .  
 2 A string is called an *active self-replicator* if  $\forall s_2 \in S: s_1 + s_2 \implies s_1$ .

Table 1. Example for the automata reaction.

$s_1$	command	comment
1011	EXOR	The exclusive-or product of $b'$ (referred to by the IO pointer) and $b$ (referred to by the operator pointer) is written at the position of $b'$ . Therefore, the 0th bit of the IO register becomes 0. Then both pointers are moved one bit position to the left.
0010 0110	SETP 0110	The pattern register is set to 0110. If the pattern is set, a subsequent MOV operation will move the pointers until the pattern is found in the operator register (max. 32 steps).
0110	CPON	The copy-mode is switched on. Every time the pointers are moved the ALU operation is performed (here, EXOR).
1101	NOP	No operation. The registers are not modified.
0001	MOV	The pointers are moved to the left until the pattern 0110 is found in the operator register, which happens at the 8th bit position. Because the copy-mode is activated (by CPON), the last ALU operation (here: EXOR) is executed for every bit passed.
0000	ID	The 8th bit of the operator register is copied into the IO register, and both pointers are moved one step to the left.
0001	MOV	Both pointer are moved to the next appearance of 0110, namely to the 12th bit position. This time the ID operation is executed at each step.

#### 4 Investigation and Visualization of System Behavior

Designing a system that shows behavior of enormous complexity is surprisingly easy. But to visualize and to explain the behavior in detail is much more difficult. Methods for investigation can be divided into macroscopic, mesoscopic, and microscopic approaches.

A *macroscopic method* observes the whole system, by mapping its high-dimensional state into a low-dimensional space, where every component of the system should be considered. An example from thermodynamics is “temperature.” A *mesoscopic method* considers only a small subset of the system’s components. It still accumulates many microscopic events into low dimensional values (e.g., the concentration of a specific substance in a region). *Microscopic methods* visualize elementary details of the system, for instance, the history of a specific object or memory cell.

Here, we shall apply only macroscopic methods, which are explained in the following. The soup (population)  $P = \{s_1, \dots, s_M\}$  is defined as a multi set on  $S$ .

**Diversity** Various measures for the diversity of a population exist. In ecology, diversity is often defined as the number of different groups or taxa. The dynamic behavior of this diversity depends on the definition of the grouping. A fine-grained grouping results in a diversity fluctuating over time. This is due to the fact that the number of individuals belonging to one group is not considered. Here, we use a simple, fine-grained grouping, where every possible genotype forms a group. Then, the resulting (absolute) diversity  $\text{Div}_{\text{abs}}$  becomes the number of different string types in  $P$ :

$$\text{Div}_{\text{abs}}(P) = |\{s \mid s \in P\}|.$$

Table 2. Examples for the automata reaction.

No.	$s_1$	$s_2$	$s_3$	comment
	operator	operand	product	
1	101d662b	cb5d5c2d	cb5d5606	typical random reaction
2	f82710fb	73fbc890	73fbc891	typical random reaction
3	98ac35f1	4c22bc78	4c22bc7c	typical random reaction
4	c1126a24	f9c8a25e	c1c8a25e	typical random reaction with TDIR
5	9a716b98	9878fa02	9878fa02	passive replication
6	9a716b98	00000000	00000000	passive replication
7	1e1ca260	078f21ff	1e1ca260	active replication
8	1e1ca260	00000000	1e1ca260	active replication

To get a relative value  $\text{Div}(P)$  in the range  $[0, 1]$  the absolute diversity is divided by the soup size  $M$ . In the future, the term “diversity” refers to the *relative diversity*

$$\text{Div}(P) = \frac{\text{Div}_{\text{abs}}(P)}{M}.$$

**Distance distribution complexity (DDC)** One problem of the diversity measure mentioned before is that the distance between different groups is not taken into account. For example, an ecology with 10 horses and 10 zebras would have the same diversity as an ecology with 10 horses and 10 amoeba. The DDC, recently introduced by Kim [21], is able to distinguish these cases: Given a discrete distance measure  $D : P \times P \Rightarrow \mathbb{N}$  the *distance distribution* is defined here as the relative frequency of the distance value  $d \in \mathbb{N}$ :

$$f(d) = \frac{2}{M(M-1)} |\{(s_i, s_j) | s_i, s_j \in P, i < j, D(s_i, s_j) = d\}|.$$

The *distance distribution complexity* is then defined as the Shannon entropy of the distribution of distance values:

$$\text{DDC}(P) := - \sum_d f(d) \log(f(d)).$$

For the following analysis  $D$  is defined as the Hamming-distance. This simple and fast measure is sufficient, because no shifting and no insertion or deletion of bits is taking place. Note, a high DDC indicates a high diversity and vice versa, but the DDC can increase even if the diversity decreases (e.g., Figure 2).

**Productivity** The *productivity*  $\text{Prod}(P)$  is the probability that a collision of two strings is reactive. If the productivity is zero, no new string is produced and inserted into the soup. To measure the productivity during a simulation, the number of iterations in which a string  $s_3$  is inserted into the soup is counted for  $M$  steps. The result is divided by  $M$ . If the soup size  $M$  is constant the productivity is equal to the dilution flux.

**Innovativity** The *innovativity* is the probability that a collision produces an object that is new, with respect to a given time window  $[t_1, t_2]$ . Let  $P_t$  be the soup after the  $t$ th iteration of the reactor algorithm. Then the (*total*) *innovativity* is defined for the time

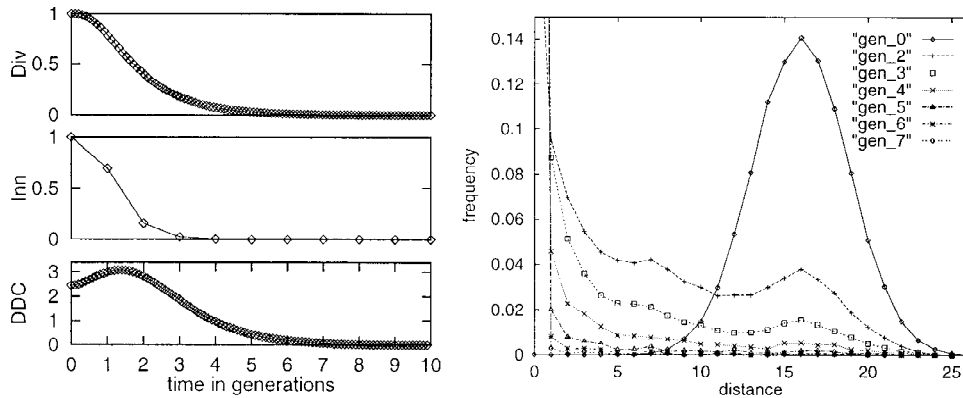


Figure 2. Example for extinction. Parameters: AND reaction, no filter condition,  $M = 10^4$ . Left: Diversity and DDC drop to zero because the system is exploited by the destructor 00000000, which remains as the only surviving string. Right: Distance distribution for the generation 0–7.

window  $[0, t]$  as

$$\text{Inn}(P_t) = p \left( s_3 \text{ is inserted into the soup} \quad \text{and} \quad s_3 \notin \bigcup_{t'=0}^t P_{t'} \right).$$

The total innovativity has been measured by counting the number of completely new strings that are inserted into the soup during  $M$  iterations of the reactor algorithm. The result is divided by  $M$ .

**Reaction table** Given a set of objects  $A = \{s_1, \dots, s_{n_a}\}$  then the corresponding reaction table  $\text{RT} : \{1, \dots, n_a\}^2 \implies \{1, \dots, n_a, -, *\}$  is defined as

$$\text{RT}(i, j) = \begin{cases} - & \text{if the collision of } s_i, s_j \text{ is elastic,} \\ k & \text{if } s_i + s_j \implies s_k \text{ and } s_k \in A, \\ * & \text{if } s_i + s_j \implies s_k \text{ and } s_k \notin A. \end{cases}$$

The term “the collision of  $s_i, s_j$  is elastic” refers to Step 2 of the reactor algorithm, where  $s_i, s_j$  are selected but no product is inserted into the soup. For the reaction tables shown in this article, the set of objects is sorted according to their concentration:  $\forall s_i, s_j \in A : [s_i] \geq [s_j] \iff i \geq j$ .

### 5 Qualitative Types of Dynamic Self-Organization Phenomena

In this section we show experimental results to discuss four qualitative types of self-organization processes. In all experiments the objects are binary strings of fixed size (32-bits). The soup is always initialized with random strings, resulting in an initial diversity of approximately 1. The phenomena will be illustrated by visualizing “typical” runs, because averaging over a series of runs with the same parameter set but different random numbers is not feasible.

#### 5.1 Extinction

Figure 2 shows the behavior of the AND reaction. The diversity drops quickly to  $1/M$  (only one string type left) and the DDC to zero, indicating that the soup is exploited



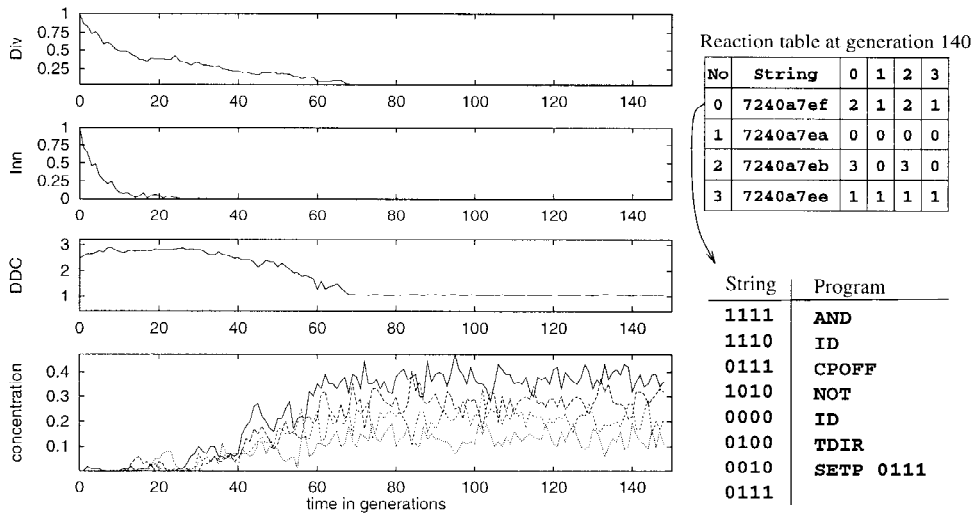


Figure 3. Emergence of an organization. Parameters: Automata reaction with code table 1,  $M = 100$ , no filter condition, soup seeded with  $M$  random strings. For the four surviving strings the reaction table and their concentration over time is given.

by a single type, the lethal string 00000000. It is able to replicate with every other string and is furthermore produced through many reaction pathways, for example,  $f120000 + 0004711a \implies 00000000$ .

At the outset the DDC increases, because intermediate products appear that are characterized by a small number of ones (e.g., 00000200), resulting in a higher complexity of the distribution of distances during generation 0–2 (Figure 2, right).

Extinction is a typical phenomenon where lethal strings appear likely [4]. The resulting organization is so simple that further examination is normally not warranted. Extinction can be circumvented by filtering lethal objects [4] or by prohibiting replication reactions [13] (Equation 3, below).

### 5.2 Emergence of an Organization

Figure 3 shows the emergence of a simple organization comprising four different string types, now in a simulation of the automata reaction in a soup of size  $M = 100$ . Diversity and innovativity drop continuously until a closed self-maintaining<sup>3</sup> organization dominates the soup with only a few distinct objects. Through drift the diversity of this organization is slowly reduced until an organization remains that does not contain a closed, self-maintaining subset.

### 5.3 Exploration and Innovation

If the soup size is increased to  $M = 10^4$  a new qualitative phenomenon appears. Figure 4 shows a typical simulation of the automata reaction with instruction Table 1 and without filter condition. After a quick drop of the diversity at the beginning (generations 0–2) the system enters an *explorative phase* (generations 2–40), which is characterized by high innovativity, productivity, and diversity. Many new objects are generated constantly and there are no string types with exceptionally high concentrations. The high innovativity indicates that the sequence space is intensively explored.

<sup>3</sup> A set  $\mathcal{A}$  of object species is *closed* if every interaction within  $\mathcal{A}$  produces only objects already in  $\mathcal{A}$ . A set  $\mathcal{A}$  of object species is *self-maintaining* if every object is produced by at least one interaction within  $\mathcal{A}$  [13].

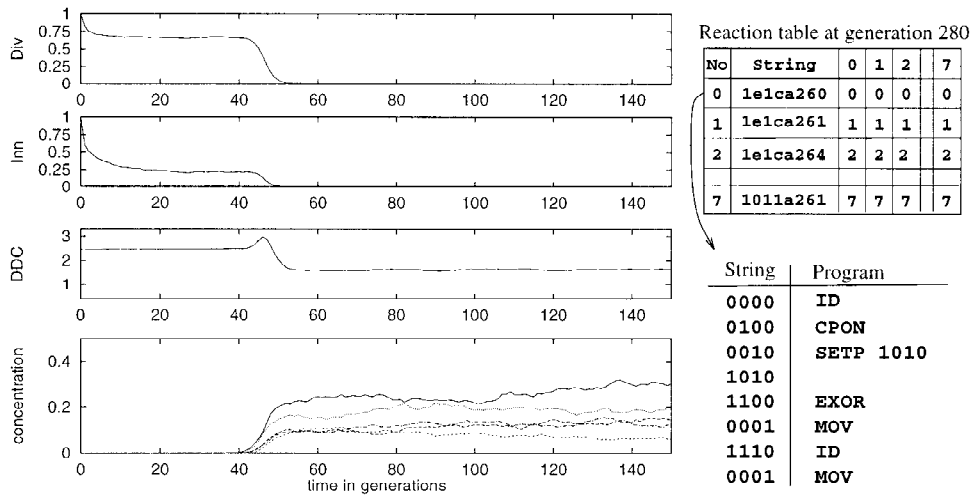


Figure 4. Exploration and innovation. During the explorative phase (generations 2–40) the diversity is constantly high and many totally new string types are produced. Parameters: soup size  $M = 10^4$ , automata reaction with Table 1, no filter condition  $f$ , soup seeded with  $M$  random strings.

The explorative phase comes to a sudden end when string types appear (around generation 40) that are replicating with every other string (active replicators), thus quickly dominating the soup. The appearance of active replicators may be called an innovation, which leads—in this case—to an end of the explorative phase. The process that follows after generation 40 is similar to the extinction process shown in Figure 3. The system becomes simple and enters a stable phase where the organization structure is reduced only through undirected drift, with every string being an active replicator. The qualitative behavior is different from the AND reaction and the process leading to the final string is more gradual and continuous.

### 5.4 Evolution

This section describes experiments that showed evolutionary behavior in the absence of external variation operators (i.e., mutation) and the absence of explicit selection<sup>4</sup> and fitness function. The following setup has been used for the experiments in this section:

- Soup size  $M = 10^5$  or  $M = 10^6$ . These large soup sizes are used to get an explorative behavior and to reduce the effects of the nondeterministic components of the reactor algorithm.
- Elastic collisions introduced through the filter condition

$$f_1(s_1, s_2, s_3) = (s_1 \neq s_3 \wedge s_2 \neq s_3). \tag{3}$$

By using this filter condition the exact replication is disabled. Every collision that normally produces a string identical to one of the reactants is considered to be elastic [13].

<sup>4</sup> The selection of two objects in the reactor algorithm is not selection in the sense of evolution theory. The random selection simulates only random collisions among objects.

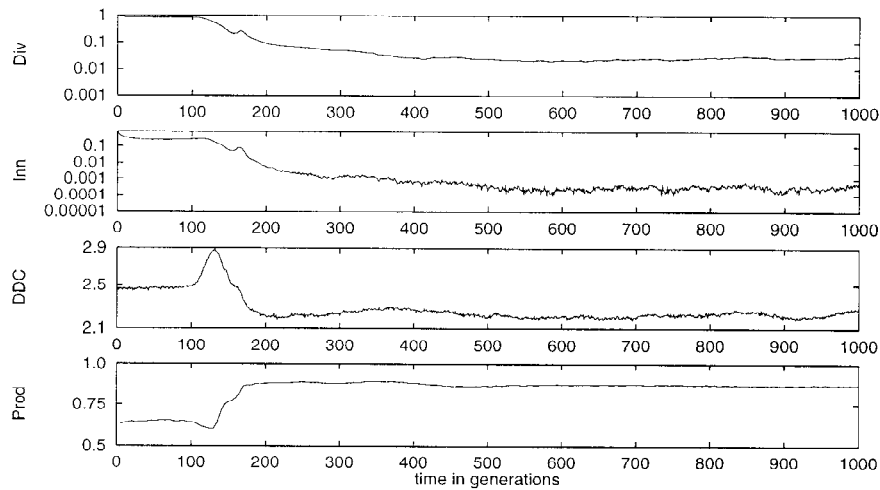


Figure 5. Short-time evolutionary behavior. Parameters: soup size  $M = 10^5$ , automata reaction with Table 2, filter condition  $f_1$ , soup seeded with  $M$  random strings.

- Reaction mechanism: automata reaction with Code Table 2. Code Table 2 does not contain a NOT operation. Without a NOT operation it is harder to construct a self-maintaining organization of strings under filter condition  $f_1$ .
- The soup is seeded with random strings from  $\{0, 1\}^{32}$ , resulting in an initial diversity approximately equal to one.

Figure 5 shows the short-time behavior.<sup>5</sup> As in Figure 4 an explorative phase with very high diversity can be observed (generations 0–110). A lot of completely new strings are produced. The productivity (fraction of collisions that are not elastic) is similar to the productivity of a soup of randomly generated strings.

This changes when a self-maintaining though not closed organization emerges (generations 110–200). This does not happen as fast as in Figure 4. Here, the process takes a much longer time and seems to be more complex. Many different “species” are competing for space. Over a short period of time new strings are generated that increase their concentrations and are replaced by “better” ones. Figure 6 shows this early phase for some representative strings of a run with soup size  $M = 10^6$ .

Figure 7 shows the long-time behavior. After the first early evolutionary phase (generations 110–200) an organization has emerged whose gene diversity is massively reduced. It consists of approximately 50,000 different string types where only about 3,000 are present at a specific time. This organization still generates completely new strings but at a much lower frequency than the soup in the explorative phase (generations 0–110).

The productivity in Figure 7 (generations 500–7,000) shows that even this (over a long period stable) organization is able to develop further. The character of this evolutionary process is different from that shown in Figure 6. There, evolution is taking place on the binary string (better: quasi-species) level. Here, the core set of strings is not changed. The organization is enhanced by adding/absorbing “useful” new strings that are “invented.”

<sup>5</sup> Although this is called “short-time behavior,” a rather long time is shown compared to Figure 2.

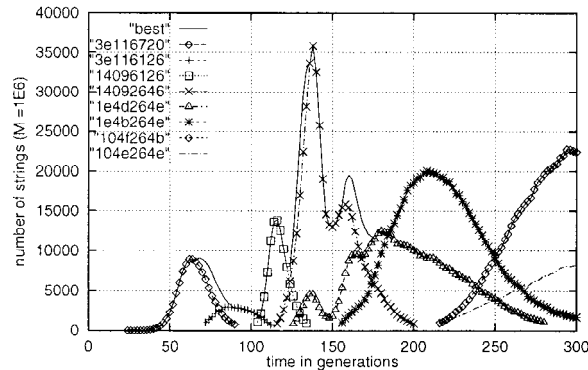


Figure 6. Early evolutionary behavior. The concentration of some representative strings are shown. Parameters: soup size  $M = 10^6$ , automata reaction with Table 2, filter condition  $f_1$ , soup seeded with  $M$  random strings.

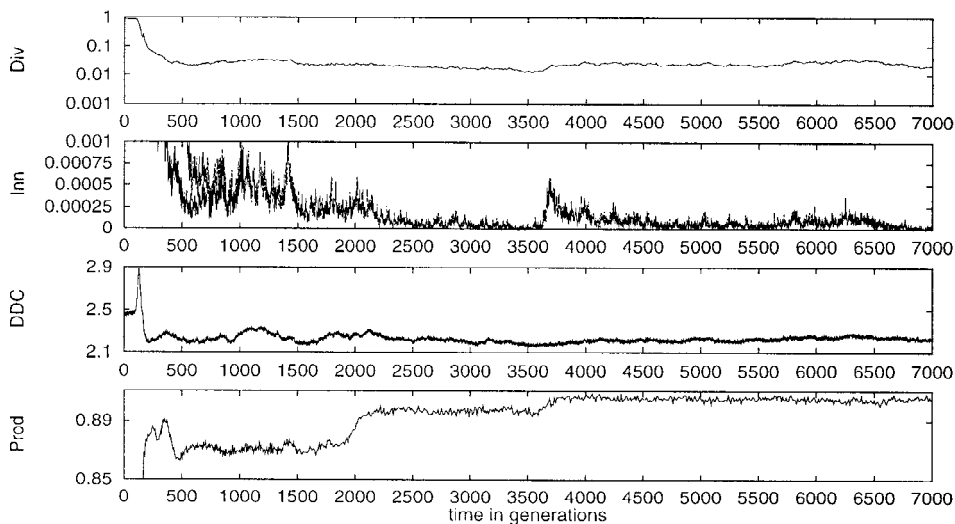


Figure 7. Long-time evolutionary behavior. Parameters: soup size  $M = 10^5$ , automata reaction with Table 2, filter condition  $f_1$ , soup seeded with  $M$  random strings. Same run as in Figure 5.

The phenomenon that long quasi-stable periods are interrupted by rapid changes is often referred to as punctuated equilibrium [11]. It has been observed in natural as well as in artificial systems (e.g., DNA world [28] and multi-agent systems [33]).

For a more detailed analysis of the organization structure the reaction table can be used. Figures 8 and 9 show a part of the reaction table of the 90 most frequent strings at different stages of the evolutionary process. They demonstrate how the coupling between strings becomes closer and the amount of elastic collisions is reduced.

An interesting detail is that a crossover mechanism has emerged. Figure 10 shows an example where 8 “crossovering” strings form a closed organization. The organization is so small and closed because the crossover is performed at a specific point.

generation 75											generation 110												
No.	String	0	1	2	3	4	5	6	7	88	89	No.	String	0	1	2	3	4	5	6	7	88	89
0	8140b6fa	29	*	*	*	*	*	*	*	*	*	0	c11a260b	-	-	-	-	-	-	-	-	-	-
1	140da6e8	-	-	-	-	-	-	-	-	-	-	1	011a2614	02	-	-	-	-	02	02	02	02	02
2	a625ffb3	29	22	-	*	*	-	-	*	-	-	2	011a2615	-	-	-	01	01	-	-	-	-	-
3	140ca6f2	-	14	-	-	14	14	14	14	14	14	3	c11a260e	-	-	-	-	-	-	-	-	-	-
4	77ae526e	-	*	*	-	-	*	*	-	*	*	4	c11a260a	-	00	-	00	00	-	-	-	-	-
5	c46b12a1	*	*	*	*	*	*	*	*	*	*	5	8211e267	27	28	28	27	27	-	28	27	28	27
6	4a8c05c9	*	*	*	*	*	*	*	*	*	*	6	1211e267	23	-	-	23	23	23	-	23	-	23
7	223be8ae	-	-	*	-	-	*	*	-	82	*	7	d211e267	-	06	06	-	-	23	-	-	*	-
88	7ecbf629	*	*	*	*	*	*	*	*	*	*	88	211a260d	49	-	-	49	49	49	-	49	-	49
89	70507121	*	-	*	*	*	-	-	*	-	-	89	c611e267	-	*	*	-	-	*	*	-	*	-

Figure 8. Left: Reaction table during the explorative phase (generation 75, Figure 5). The objects are loosely coupled. Many reaction pathways leave the reaction table (denoted by “\*”). Right: Reaction table shortly after the explorative phase (generation 110, Figure 5). The objects are coupled more closely. Still many reaction pathways leave the reaction table. “-” means an elastic collision (no reaction product).

## 6 Discussion

Our experiments demonstrate that complex forms of evolution can take place in systems without any explicit variation operator, like mutation or recombination, and without any explicit (artificial) selection. The setup used here is one of the simplest ways of letting machines act on each other. We think that an important concept concerning natural evolution is that the way evolution is carried out is itself under the influence of evolution, a concept called *meta-evolution*. Meta-evolution has long been applied in evolutionary algorithms. For example, in evolution strategies (ES) parameters are used that determine the variance and covariance of a generalized  $n$ -dimensional normal distribution for mutations. The strategy parameters themselves are adapted during the optimization process. Only the operators used for adaptation are fixed. In our system, the variation of an object alters the way other objects are modified, because the object will subsequently interact with others. We can interpret such a system as having many levels of meta-evolution, maybe a potentially infinite number of levels.

Bagley, Farmer, and Fontana have studied the emergence and evolution of autocatalytic metabolic networks by using a constructive artificial chemistry, where the molecules are character sequences and the reactions are concatenation and cleavage [2]. They have shown how small, spontaneous fluctuations can be amplified by an autocatalytic network, possibly leading to a modification of the entire network. A sequence of such modifications has been identified as an evolutionary process. The spontaneous fluctuations are explicitly induced by the meta-dynamical algorithm, which can be interpreted as explicit mutation.

In contrast to the metadynamical ODE approach in [2], we have used a discrete soup of objects. The diversity in the kinetic parameters is low. Only two different values for rate constants are needed (e.g., 0 and 1, see Equation 2). There are only catalyzed reactions. No spontaneous reactions are needed to generate an evolutionary behavior. The evolutionary steps are not explicitly initiated by the simulation algorithm. The algorithm for simulating the kinetics is much simpler than the stochastic metadynamics proposed in [2].

Disadvantages of our approach are that (a) the number of molecules inside the soup is finite and bounded by the memory size, (b) reactions with rate constants that differ in several orders of magnitude can not be simulated easily, and (c) the computational effort is much higher in the case of low diversity. Advantages are that (a) the number

generation 3400

No.	String	0	1	2	3	4	5	6	7	8	9	10	11	12	13	14	15	16	17	67	68	69
0	104e264b	01	-	03	02	05	04	07	06	01	-	03	02	01	-	02	-	03	-	64	65	-
1	104e264a	-	00	03	02	05	04	07	06	-	00	03	02	-	00	02	00	03	00	64	65	00
2	104f264b	01	00	03	-	05	04	07	06	01	00	03	-	01	00	-	00	03	00	64	65	00
3	104f264a	01	00	-	02	05	04	07	06	01	00	-	02	01	00	02	00	-	00	64	65	00
4	140e264b	-	00	-	02	-	-	-	06	00	00	02	02	00	00	02	00	02	00	-	65	00
5	140e264a	01	-	03	-	-	-	07	-	01	01	03	03	01	01	03	01	03	01	64	-	01
6	140f264b	-	00	-	02	-	04	-	-	00	00	02	02	00	00	02	00	02	00	-	65	00
7	140f264a	01	-	03	-	05	-	-	-	01	01	03	03	01	01	03	01	03	01	64	-	01
8	104e246b	09	-	11	10	30	24	31	28	09	-	11	10	09	-	10	-	11	-	*	*	-
9	104e246a	-	08	11	10	30	24	31	28	-	08	11	10	-	08	10	08	11	08	*	*	08
10	104f246b	09	08	11	-	30	24	31	28	09	08	11	-	09	08	-	08	11	08	*	*	08
11	104f246a	09	08	-	10	30	24	31	28	09	08	-	10	09	08	10	08	-	08	*	*	08
12	104e206b	-	00	-	02	00	00	02	02	-	08	-	10	-	-	16	21	-	18	00	02	25
13	104e206a	01	-	03	-	01	01	03	03	09	-	11	-	-	-	-	14	-	-	01	03	23
14	104f206a	01	-	03	-	01	01	03	03	09	-	11	-	13	-	-	-	-	-	01	03	23
15	104e261a	01	-	03	-	01	01	03	03	09	-	11	-	13	-	-	-	14	-	01	03	23
16	104f206b	-	00	-	02	00	00	02	02	-	08	-	10	-	12	-	21	-	18	00	02	25
17	104e260a	01	-	03	-	01	01	03	03	09	-	11	-	13	-	-	-	14	-	01	03	23
67	1046264b	01	00	03	02	05	04	07	06	*	*	*	*	*	*	*	00	*	00	64	65	00
68	10af264a	59	59	-	-	59	59	-	-	*	*	*	*	*	*	*	*	*	*	59	-	*
69	104e265e	01	-	03	-	01	01	03	03	09	-	11	-	13	-	-	-	14	-	01	03	-

Figure 9. Reaction table of a converged soup (generation 3,400, Figure 5). The objects are coupled very closely. Only a few reaction pathways are leaving the reaction table. The objects are very similar.

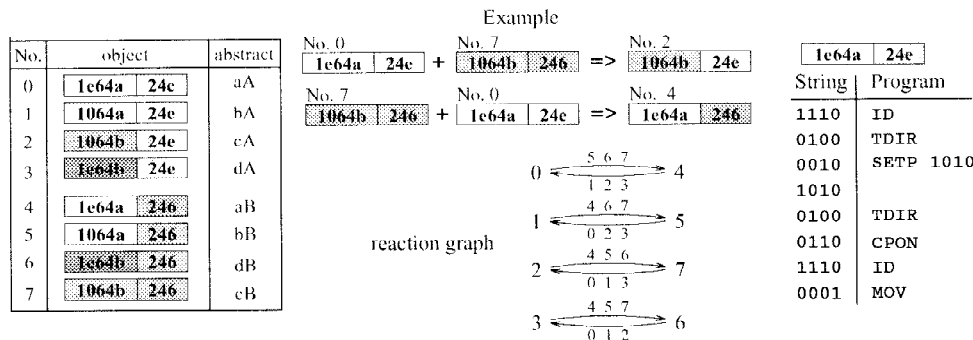


Figure 10. Emergence of crossover. The strings are able to insert a part of their sequence into the sequence of an arbitrary string. The result is a crossover of both, for example, 1e64a24e + 00000000 ⇒ 0000024e.

of different objects inside the soup can be very high (e.g., 10<sup>6</sup> here versus 75 types in [2]), (b) the time evolution is not influenced or disturbed by the algorithm simulating the chemical dynamics, and (c) implementation is much simpler.

### 7 Summary and Future Work

We have extended our earlier work by introducing a reaction mechanism inspired by techniques of computer science. It has been demonstrated that the building blocks

used in computer science to describe hardware elements and functions can be easily used to construct self-organizing and even evolving systems.

For exploring generalizations about evolving systems the high speed at which the reactions are carried out is helpful, because it allows interactive and large experiments. The disadvantages are the (so far) fixed length of strings and that the formalization of the reaction mechanism is more complicated than using lambda-calculus [13] or matrix multiplication [4].

The complex behavior shown here results from a simple, ad hoc, hand-written interaction mechanism working on small objects in an environment with no spatial structure. Introducing larger, variable-length objects and topological structures, allowing the formation of niches, will result in an increase in complexity. Then we have to face the danger that in order to grasp a world we do not understand, we create a system we do not understand [6, 24]. To circumvent this, powerful analyzation and visualization methods are required; these should be general (like the DDC), so that they can be applied to different systems [9].

Reaction systems—like the system discussed here—are able to perform computations [5]. For example, a chemical reaction system has been used to instantiate an artificial neural network [16] or to solve a variant of the Hamilton-path problem [1]. So far, the reaction pathways have been chosen carefully by hand. The integration of self-evolution into molecular computing systems is a promising goal for future research.

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### Appendix

The Appendix contains a description for all registers and instructions of the automata reaction.

#### The Control Registers

In addition to the two 32-bit registers the automaton has five smaller *control registers*. The control registers are initialized with the first value given in the following description:

direction $r_D$	$r_D \in \{\text{left, right}\}$ The direction register determines the direction the pointers are moving. The operation TDIR toggles the direction.
move-mode $r_M$	$r_M \in \{\text{both, IOpointer, operator-pointer}\}$ This register controls which pointer is moved in case of a pointer movement. The operation TMM toggles the movement mode.
last-ALU-operation $r_{ALU}$	$r_{ALU} \in \{ID, NOT, AND, OR, EXOR, EQ\}$ This register stores the last ALU operation. If the copy mode is switched on, the operation stored in $r_{ALU}$ will be executed before each movement step.

copy-mode $r_C$	$r_C \in \{\text{off}, \text{on}\}$ If the copy mode is switched on, the last ALU operation stored in $r_{ALU}$ is executed before each movement step of the IO pointer and the operator-pointer.
pattern $r_P$	$r_P \in \{\text{none}\} \cup \{0, 1\}^4$ With “SETP $\mathbf{p}$ ” this register will be set to the pattern $\mathbf{p}$ ( $\mathbf{p} \in \{0, 1\}^4$ ). The operation UNSETP clears the pattern by setting $r_P$ to none. If the pattern is set, each MOV operation will move the pointers unless the pattern is found in the operator register (max. 32 steps). If only the IO pointer is moved the pattern is searched in the IO register. If $r_P = \text{none}$ the pointers are moved one step.

### Arithmetic and Logic Instructions

During an arithmetic or logic operation bit  $b$  and  $b'$  are processed by the ALU (Figure 1). The result overwrites  $b'$ . Then the pointers are moved accordingly to the direction register  $r_D$  (left or right) and the move-mode register  $r_M$  (both, IO pointer only, or operator pointer only).

ID	identity: $b' := b, r_{ALU} := ID$ .
NOT	negation: $b' := \neg b, r_{ALU} := NOT$ .
AND	logic and: $b' := b \wedge b', r_{ALU} := AND$ .
OR	logic or: $b' := b \vee b', r_{ALU} := OR$ .
EXOR	exclusive or: $b' := b \neq b', r_{ALU} := EXOR$ .
EQ	equality: $b' := b = b', r_{ALU} := EQ$ .

After the execution of a logic instruction, register  $r_{ALU}$  is set to the operation executed. It is used only during the execution of the MOV instruction, if the copy mode is switched on ( $r_C = \text{on}$ ).

### Movement and Control Instructions

MOV	Moves the pointers accordingly to the move-mode, direction and pattern register. If no pattern is set, ( $r_P = \text{none}$ ) the pointers are moved one step (one bit). If a pattern is set, the pointers are moved unless the pattern is found (max. 32 steps). If $r_M = \text{IOpointer}$ the pattern is searched in the IO register, otherwise in the operator register. Again, the direction of the movement is given by the direction register $r_D$ . If the copy mode is switched on ( $r_C := \text{on}$ ) by CPON, the last ALU operation is executed before each step.
CPON	Switches the copy-mode on ( $r_C := \text{on}$ ).
CPOFF	Switches the copy-mode off ( $r_C := \text{off}$ ).
SETP	Sets the pattern $\mathbf{p} \in \{0, 1\}^4$ for the move operation ( $r_P := \mathbf{p}$ ). This is the only operation with an argument.
UNSETP	Clears the pattern ( $r_P := \text{none}$ ).



TDIR Toggles the move direction for the pointers.

$$r_D := \begin{cases} \text{right} & \text{if } r_D = \text{left}, \\ \text{left} & \text{if } r_D = \text{right} \end{cases}$$

TMM Toggles the move-mode:

$$r_M := \begin{cases} \text{both} & \text{if } r_M = \text{operator-pointer}, \\ \text{IOpointer} & \text{if } r_M = \text{both}, \\ \text{operator-pointer} & \text{if } r_M = \text{IOpointer} \end{cases}$$

NOP No operation.

STOP Stops the automaton before the whole program is executed.

For a precise formal specification of the automata reaction the source code is available [8].

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