

Reversible Molecular Computation in Ciliates

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Abstract

We prove that a reversible model for the guided homologous recombinations that take place during gene rearrangement in ciliates has the computational power of a Turing machine, the accepted formal model of computation. This indicates that, in principle, these unicellular organisms may have the capacity to perform any computation carried out by an electronic computer.

1 Gene unscrambling as computation

Ciliates are a diverse group of 8000 or more unicellular eukaryotes (nucleated cells) named for their wisp-like covering of cilia. They possess two types of nuclei: an active *macronucleus* (soma) and a functionally inert *micronucleus* (germline) which contributes only to sexual reproduction. The somatically active macronucleus forms from the germline micronucleus after sexual reproduction, during the course of development. The genomic copies of some protein-coding genes in the micronucleus of hypotrichous ciliates are obscured by the presence of intervening non-protein-coding DNA sequence elements (internally eliminated sequences, or *IESs*). These must be removed before the assembly of a functional copy of the gene in the somatic macronucleus. Furthermore, the protein-coding DNA segments (macronuclear destined sequences, or *MDSs*) in species of *Oxytricha* and *Stylonychia* are sometimes present in a permuted order relative to their final position in the macronuclear copy. For example, in *O. nova*, the micronuclear copy of three genes (Actin I, α -telomere binding protein, and DNA polymerase α) must be reordered and intervening DNA sequences removed in order to construct functional macronuclear genes. Most impressively, the gene encoding DNA polymerase α (DNA pol α) in *O. trifallax*

Research partially supported by Grant R2824AO1 of the Natural Sciences and Engineering Research Council of Canada to L.K. and a Burroughs-Wellcome Fund New Investigator Award in Molecular Parasitology to L.F.L.

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is apparently scrambled in 50 or more pieces in its germline nucleus [10]. Destined to unscramble its micronuclear genes by putting the pieces together again, *O. trifallax* routinely solves a potentially complicated computational problem when rewriting its genomic sequences to form the macronuclear copies.

This process of unscrambling bears a remarkable resemblance to the DNA algorithm Adleman [1] used to solve a seven-city instance of the Directed Hamiltonian Path Problem. The developing ciliate macronuclear “computer” (Figures 2-3) apparently relies on the information contained in short direct repeat sequences to act as minimal guides in a series of homologous recombination events. These guide-sequences act as splints, and the process of recombination results in linking the protein-encoding segments (MDSs, or “cities”) that belong next to each other in the final protein coding sequence. As such, the unscrambling of sequences that encode DNA polymerase α accomplishes an astounding feat of cellular computation. Other structural components of the ciliate chromatin presumably play a significant role, but the exact details of the mechanism are still unknown.

2 The path towards unscrambling

Typical IES excision in ciliates involves the removal of short (14 - 600bp) A-T rich sequences flanked by direct repeats of 2 to 14 bp. IESs are often released as circular DNA molecules [20]. The choice of which sequences to remove appears to be minimally “guided” by recombination between direct repeats of only 2 to 14 base pairs.

Unscrambling is a particular type of IES removal in which the order of the MDSs in the micronucleus is often radically different from that in the macronucleus. For example, in the micronuclear genome of *Oxytricha nova*, the MDSs of α -telomere binding protein (α -TP) are arranged in the cryptic order 1-3-5-7-9-11-2-4-6-8-10-12-13-14 relative to their position in the “clear” macronuclear sequence 1-2-3-4-5-6-7-8-9-10-11-12-13-14. This particular arrangement predicts a spiral mechanism in the path of unscrambling which links odd and even segments in order (Figure 4; [14]).

Homologous recombination between identical short sequences at appropriate MDS-IES junctions is implicated in the mechanism of gene unscrambling, as it could simultaneously remove the IESs and reorder the MDSs. For example, the DNA sequence present at the junction between MDS n and the downstream IES is generally the same as the sequence between MDS $n+1$ and its upstream IES, leading to correct ligation of MDS n to MDS $n+1$, over a distance. However the presence of such short repeats (average length 4 bp between non-scrambled MDSs, 9 bp between scrambled MDSs [17]) implies that although these guides are necessary, they are certainly not sufficient to guide accurate splicing. Hence it is likely that the repeats satisfy more of a structural requirement for MDS splicing, and less of a role in substrate recognition. Oth-

erwise, incorrectly spliced sequences (the results of promiscuous recombination) would dominate, especially in the case of very small (2-4 bp) repeats present thousands of times throughout the genome. This incorrect hybridization could be a driving force in the production of newly scrambled patterns in evolution. However during macronuclear development only unscrambled molecules which contain 5' and 3' telomere addition sequences would be selectively retained in the macronucleus, ensuring that most promiscuously ordered genes would be lost.

Alternate splicing at the RNA level, as well as other forms of programmed DNA rearrangements, could also be viewed as solutions to path finding problems in nature. Dynamic processes, such as maturation of the immune response, provide examples of genuine evolutionary computation in cells, whereas the path finding problems here may follow a more deterministic algorithm. Current effort is directed toward understanding how cells unscramble DNA, how this process has arisen, and how the “programs” are written and executed. Do they decode the message by following the shortest unscrambling path or by following a more circuitous but equally effective route, as in the case of RNA editing ([12])? Also, how error prone is the unscrambling process? Does it actually search through several plausible unscrambled intermediates or follow a strictly deterministic pathway?

3 The formal model

Before introducing the formal model, we summarize our notation. An alphabet Σ is a finite, nonempty set. In our case $\Sigma = \{A, C, G, T\}$. A sequence of letters from Σ is called a string (word) over Σ and in our interpretation corresponds to a linear strand. The words are denoted by lowercase letters such as u, v, α_i, x_{ij} . The length of a word w is denoted by $|w|$ and represents the total number of occurrences of letters in the word. A word with 0 letters in it is called an empty word and is denoted by λ . The set of all possible words consisting of letters from Σ is denoted by Σ^* , and the set of all nonempty words by Σ^+ . We also define circular words over Σ by declaring two words to be equivalent if and only if (iff) one is a cyclic permutation of the other. In other words, w is equivalent to w' iff they can be decomposed as $w = uv$ and $w' = vu$, respectively. Such a circular word $\bullet w$ refers to any of the circular permutations of the letters in w . Denote by $\Sigma^\bullet$ the set of all circular words over Σ .

With this notation, we define intramolecular recombination using set theoretical notation as:

$$\{uxwxv\} \implies \{uxv, \bullet wx\}$$

where u, w, x , and v are words in Σ^* , and x , the junction sequence that guides unscrambling, is nonempty.

Thus the defined operation models the process of intramolecular recombination. After x finds its second occurrence in $uxwxv$, the molecule undergoes

a strand exchange in x that leads to the formation of two new molecules: uxv and a circular DNA molecule $\bullet wx$.

Intramolecular recombination also accomplishes the deletion of either sequence wx or xw from the original molecule $uxwxv$. The fact that $\bullet wx$ is circular implies that we can use any circular permutation of its sequence as an input for a subsequent operation.

In this model, the effects of intramolecular recombination can be reversed. Note that the operation in the forward direction is formally intramolecular recombination, whereas the operation in the reverse direction is intermolecular recombination. The intermolecular recombination

$$\{uxv, \bullet wx\} \Longrightarrow \{uxwxv\}$$

also accomplishes the insertion of the sequence wx or xw in the linear string uxv .

The above operations resemble the “splicing operation” introduced by Head in [7] and “circular splicing” ([8], [19], [16]). [15], [3] and subsequently [21] showed that these models have the computational power of a universal Turing machine. (See [9] for a review.)

The process of gene unscrambling entails a series of successive or possibly simultaneous intra- and inter-molecular homologous recombinations. This is followed by excision of all sequences $\tau_s y \tau_e$, where the sequence y is marked by the presence of telomere addition sequences τ_s for telomere “start” (at its 5’ end), and τ_e for telomere “end” (at its 3’ end). Thus from a long sequence $u \tau_s y \tau_e v$, this step retains only $\tau_s y \tau_e$ in the macronucleus. Lastly, the enzyme telomerase extends the length of the telomeric sequences (usually double-stranded TTTTGGGG_{*n*} repeats in these organisms) from τ_s and τ_e to protect the ends of the DNA molecule.

We now make the assumption that, by a clever structural alignment, such as the one depicted in Figure 4, or other biological factors, the cell decides which sequences are non-protein-coding (IESs) and which are ultimately protein-coding (MDSs), as well as which sequences x guide homologous recombination. Moreover, such biological shortcuts are presumably essential to bring into proximity the guiding sequences x . Each of the n MDSs, denoted primarily by α_i , $1 \leq i \leq n$ is flanked by the guiding sequences $x_{i-1,i}$ and $x_{i,i+1}$. Each guiding sequence points to the MDS that should precede or follow α_i in the final sequence. The only exceptions are α_1 , which is preceded by τ_s , and α_n which is followed by τ_e in the input string or micronuclear molecule. Note that although present generally once in the final macronuclear copy, each $x_{i,i+1}$ occurs at least twice in the micronuclear copy – once after α_i and once before α_{i+1} .

We denote by ϵ_k an internal sequence that is deleted; ϵ_k does not occur in the final sequence. Thus, since unscrambling leaves one copy of each $x_{i,i+1}$ between α_i and α_{i+1} , an IES is nondeterministically either $\epsilon_k x_{i,i+1}$ or $x_{i-1,i} \epsilon_k$, depending on which guiding sequence $x_{i,i+1}$ is eliminated. Similarly an MDS is technically either $\alpha_i x_{i+1}$ or $x_{i-1,i} \alpha_i$. For this model, either choice is equivalent.

Removal of nonscrambled IESs in *Euplotes crassus* actually leaves extra sequences (including a duplication of x_{ij}) at the junctions between ϵ_k 's in the resulting non-protein-coding products. This may result when the x_{ij} 's are as short as two nucleotides [11]. It is unknown whether unscrambling also introduces extra sequences, since it uses considerably longer x_{ij} 's on average. However, since the extra sequences have always been found at junctions between ϵ_k 's, this would not affect our unscrambling model.

The following example models unscrambling of a micronuclear gene that contains MDSs in the scrambled order 2-4-1-3:

$$\begin{aligned}
 & \{u \ x_{12} \ \alpha_2 \ x_{23} \ \epsilon_1 \ x_{34} \ \alpha_4 \ \tau_e \ \epsilon_2 \ \tau_s \ \alpha_1 \ x_{12} \ \epsilon_3 \ x_{23} \ \alpha_3 \ x_{34} \ v\} \implies \\
 & \{u \ x_{12} \ \epsilon_3 \ x_{23} \ \alpha_3 \ x_{34} \ v, \quad \bullet \alpha_2 \ x_{23} \ \epsilon_1 \ x_{34} \ \alpha_4 \ \tau_e \ \epsilon_2 \ \tau_s \ \alpha_1 \ x_{12} \} = \\
 & \{u \ x_{12} \ \epsilon_3 \ x_{23} \ \alpha_3 \ x_{34} \ v, \quad \bullet \epsilon_1 \ x_{34} \ \alpha_4 \ \tau_e \ \epsilon_2 \ \tau_s \ \alpha_1 \ x_{12} \ \alpha_2 \ x_{23} \} \implies \\
 & \{u \ x_{12} \ \epsilon_3 \ x_{23} \ \epsilon_1 \ x_{34} \ \alpha_4 \ \tau_e \ \epsilon_2 \ \tau_s \ \alpha_1 \ x_{12} \ \alpha_2 \ x_{23} \ \alpha_3 \ x_{34} \ v\} \implies \\
 & \{u \ x_{12} \ \epsilon_3 \ x_{23} \ \epsilon_1 \ x_{34} \ v, \quad \bullet \alpha_4 \ \tau_e \ \epsilon_2 \ \tau_s \ \alpha_1 \ x_{12} \ \alpha_2 \ x_{23} \ \alpha_3 \ x_{34} \} = \\
 & \{u \ x_{12} \ \epsilon_3 \ x_{23} \ \epsilon_1 \ x_{34} \ v, \quad \bullet \tau_s \ \alpha_1 \ x_{12} \ \alpha_2 \ x_{23} \ \alpha_3 \ x_{34} \ \alpha_4 \ \tau_e \ \epsilon_2 \} \implies \\
 & \{\tau_s \ \alpha_1 \ x_{12} \ \alpha_2 \ x_{23} \ \alpha_3 \ x_{34} \ \alpha_4 \ \tau_e, \quad \epsilon_2, \quad u \ x_{12} \ \epsilon_3 \ x_{23} \ \epsilon_1 \ x_{34} \ v\}
 \end{aligned}$$

Note that the process is nondeterministic in that, for example, one could start by replacing the first step, between homologous sequences x_{12} , by recombination between the homologous sequences x_{34} instead, obtaining the same result in the same number of steps.

Once the cell has “decided” which are the α_i 's, $x_{i,i+1}$'s and ϵ_i 's, the process that follows is simply sorting, requiring a linear number of steps (possibly fewer than n if some of the recombination events take place simultaneously). Part of this “decision” process entails finding the correct “path” linking the pieces of protein-coding regions in the correct order, with the occurrence of $\alpha_i x_{i,i+1}$ and $x_{i,i+1} \alpha_{i+1}$ in the micronuclear sequence providing the link between α_i and α_{i+1} in the macronuclear sequence.

A computational difficulty is the presence of multiple copies of the sequences $x_{i,i+1}$ which may direct the formation of incorrect “paths”. Indeed, throughout the genome, such simple sequences may be present in extreme redundancy. Some of the $x_{i,i+1}$ even overlap with each other. For example, in the *O. trifallax* gene encoding DNA polymerase α , $x_{24,25} = \text{GAGAGATAGA}$ contains $x_{1,2} = \text{AGATA}$ as a subsequence. The search for the proper junction sequences thus amounts to finding the correct “path” and is potentially the most costly part of the computation. Production of incorrect paths will not necessarily lead to the production of incorrect proteins unless the path sequences start and end with the correct telomere addition sites (τ_s and τ_e), since these ensure survival of the genes in the macronucleus. The role of telomeres here is thus to preserve those strands that start and end with the correct origin and final destinations.

4 Computational power of gene rearrangement

In this section we define the notion of a *reversible guided recombination system* that models the process taking place during gene rearrangement, and prove that such systems have the computational power of a Turing machine, the most widely used theoretical model of electronic computers.

The following strand operations generalize the intra- and intermolecular recombinations defined in the preceding section by assuming that homologous recombination is influenced by the presence of certain contexts, i.e., either the presence of an IES or an MDS flanking a junction sequence x_{ij} . The observed dependence on the old macronuclear sequence for correct IES removal in *Paramecium* suggests that this is the case ([13]). This restriction captures the fact that the guide sequences do not contain all the information for accurate splicing during gene unscrambling.

We define the contexts that restrict the use of recombinations by a *splicing scheme*, [7], [8], a pair $(\Sigma, \sim)$ where Σ is the alphabet and $\sim$, the pairing relation of the scheme, is a *commutative* binary relation between triplets of nonempty words satisfying the following condition: If $(p, x, q) \sim (p', y, q')$ then $x = y$.

In the splicing scheme $(\Sigma, \sim)$ pairs $(p, x, q) \sim (p', x, q')$ now define the contexts necessary for a recombination between the repeats x . Then we define *contextual recombination* as

$$\{uxwxv\} \iff \{uxv, \bullet wx\}, \text{ where } u = u'p, w = qw' = w''p', v = q'v'.$$

This constrains recombination to occur only if the restrictions of the splicing scheme concerning x are fulfilled, i.e., the first occurrence of x is preceded by p and followed by q and its second occurrence is preceded by p' and followed by q' .

Intramolecular recombinations can also take place within a circular strand, and intermolecular recombinations can occur between two circular strands. This leads to the definition of *circular contextual recombination* as

$$\{\bullet uxwxv\} \iff \{\bullet uxv, \bullet wx\}, \text{ where } u = u'p, w = qw' = w''p', v = q'v'.$$

Note that sequences p, x, q, p', q' are nonempty, and that the binary relation of the splicing scheme is commutative.

The operations defined in the preceding section are particular cases of contextual recombinations, where all the contexts are empty, i.e, $(\lambda, x, \lambda) \sim (\lambda, x, \lambda)$ for all $x \in \Sigma^+$. This would correspond to the case where recombination may occur between every repeat sequence, regardless of the contexts.

If we use the classical notion of a set, we can assume that the strings entering a recombination are available for multiple operations. Similarly, there would be no restriction on the number of copies of each strand produced by recombination. However, we can also assume some strings are only available in a limited number of copies. Mathematically this translates into using *multisets*, where one keeps

track of the number of copies of a string at each moment. In the style of [6], if $\mathbf{N}$ is the set of natural numbers, a multiset of Σ^* is a mapping $M : \Sigma^* \longrightarrow \mathbf{N} \cup \{\infty\}$, where, for a word $w \in \Sigma^*$, $M(w)$ represents the number of occurrences of w . Here, $M(w) = \infty$ means that there are unboundedly many copies of the string w . The set $\text{supp}(M) = \{w \in \Sigma^* \mid M(w) \neq 0\}$, the *support of M* , consists of the strings that are present at least once in the multiset M . The definition can be naturally extended to a multiset consisting of linear as well as circular strands.

We now define a *reversible guided recombination system* that captures the series of dispersed homologous recombination events that take place during these gene rearrangements in ciliates.

Definition *A reversible guided recombination system is a triple $R = (\Sigma, \sim, A)$ where $(\Sigma, \sim)$ is a splicing scheme, and $A \in \Sigma^+$ is a linear string called the axiom.*

A reversible guided recombination system R defines a *derivation relation* that produces a new multiset from a given multiset of linear and circular strands, as follows. Starting from a “collection” (multiset) of strings with a certain number of available copies of each string, the next multiset is *derived from* the first one by an intra- or inter-molecular recombination between existing strings. The strands participating in the recombination are “consumed” (their multiplicity decreases by 1) whereas the products of the recombination are added to the multiset (their multiplicity increases by 1).

For two multisets S and S' in $\Sigma^* \cup \Sigma^\bullet$, we say that S derives S' and we write $S \Longrightarrow_R S'$, iff one of the following two cases hold:

- (1) there exist $\alpha \in \text{supp}(S)$, $\beta, \bullet\gamma \in \text{supp}(S')$ such that
 - $\{\alpha\} \Longrightarrow \{\beta, \bullet\gamma\}$ according to an intramolecular recombination step in R ,
 - $S'(\alpha) = S(\alpha) - 1$, $S'(\beta) = S(\beta) + 1$, $S'(\bullet\gamma) = S(\bullet\gamma) + 1$;
- (2) there exist $\alpha', \bullet\beta' \in \text{supp}(S)$, $\gamma' \in \text{supp}(S')$ such that
 - $\{\alpha', \bullet\beta'\} \Longrightarrow \{\gamma'\}$ according to an intermolecular recombination step in R ,
 - $S'(\alpha') = S(\alpha') - 1$, $S'(\bullet\beta') = S(\bullet\beta') - 1$, $S'(\gamma') = S(\gamma') + 1$.

In the above definition of the derivation relation α and β are either both linear or both circular strands in $\Sigma^* \cup \Sigma^\bullet$.

Those linear strands which, by repeated recombinations with initial and intermediate strands eventually produce the axiom, form the language of the reversible guided recombination system. Formally,

$$L_k(R) = \{w \in \Sigma^* \mid \{w\} \Longrightarrow_R^* S \text{ and } A \in \text{supp}(S)\},$$

where the the multiplicity of w equals k . Note that $L_k(R) \subseteq L_{k+1}(R)$ for any $k \geq 1$.

In a Turing machine (TM), a read/write head scans an infinite tape composed of discrete “squares”, one square at a time. The read/write head communicates with a control mechanism under which it can read the symbol in the current square or replace it by another. The read/write head is also able to

move on the tape, one square at a time, to the right and to the left (note the analogy to the action of RNA or DNA polymerase). The set of words which make a Turing machine finally halt is considered its language.

Formally, [18], a rewriting system $TM = (S, \Sigma \cup \{\#\}, P)$ is called a *Turing machine* iff:

- (i) S and $\Sigma \cup \{\#\}$ (with $\# \notin \Sigma$ and $\Sigma \neq \emptyset$) are two disjoint alphabets referred to as the *state* and the *tape* alphabets.
- (ii) Elements s_0 and s_f of S , and B of Σ are the *initial* and *final* state, and the *blank symbol*, respectively. Also a subset T of Σ is specified and referred to as the *terminal* alphabet. It is assumed that T is not empty.
- (iii) The productions (rewriting rules) of P are of the forms

- (1) $s_i a \longrightarrow s_j b$ (overprint)
- (2) $s_i a c \longrightarrow a s_j c$ (move right)
- (3) $s_i a \# \longrightarrow a s_j B \#$ (move right and extend workspace)
- (4) $c s_i a \longrightarrow s_j c a$ (move left)
- (5) $\# s_i a \longrightarrow \# s_j B a$ (move left and extend workspace)
- (6) $s_f a \longrightarrow s_f$
- (7) $a s_f \longrightarrow s_f$

where s_i and s_j are states in S , $s_i \neq q_f$, $s_j \neq q_f$, and a, b, c are in Σ . For each pair (s_i, a) , where s_i and a are in the appropriate ranges, P either contains no productions (2) and (3) (resp. (4) and (5)) or else contains both (3) and (2) for every c (resp. contains both (5) and (4) for every c). There is no pair (s_i, a) such that the word $s_i a$ is a subword of the left side in two productions of the forms (1), (3), (5).

A configuration of the TM is of the form $\#w_1 s_i w_2 \#$, where $w_1 w_2$ represents the contents of the tape, $\#$ s are the boundary markers, and the position of the state symbol s_i indicates the position of the read/write head on the tape: if s_i is positioned at the left of a letter a , this indicates that the read/write head is placed over the cell containing a . The TM changes from one configuration to another according to its rules. For example, if the current configuration is $\#w s_i a w' \#$ and the TM has the rule $s_i a \longrightarrow s_j b$, this means that the read/write head positioned over the letter a will write b over it, and change its state from s_i to s_j . The next configuration in the derivation will be thus $\#w s_j b w' \#$.

The Turing machine TM *halts* with a word w iff there exists a derivation that, when started with the read/write head positioned at the beginning of w eventually reaches the final state, i.e. if $\#q_0 w \#$ derives $\#q_f \#$ by successive applications of the rewriting rules (1) - (7). The language $L(TM)$ *accepted* by TM consists of all words over the terminal alphabet T for which the TM halts. Note that TM is *deterministic*: at each step of the rewriting process, the application of at most one production is possible.

Theorem. *Let L be a language over T^* accepted by a Turing machine $TM = (S, \Sigma \cup \{\#\}, P)$ as above. Then there exist an alphabet Σ' , a sequence $\pi \in \Sigma'^*$,*

depending on L , and a reversible guided recombination system R such that a word w over T^* is in L if and only if $\#^6 q_0 w \#^6 \pi$ belongs to $L_k(R)$ for some $k \geq 1$.

Proof. Consider that the rules of P are ordered in an arbitrary fashion and numbered. Thus, if TM has m rules, a rule is of the form $i : u_i \longrightarrow v_i$ where $1 \leq i \leq m$.

We construct a reversible guided recombination system $R = (\Sigma', \sim, A)$ and a sequence $\pi \in \Sigma'^*$ with the required properties. The alphabet is $\Sigma' = S \cup \Sigma \cup \{\#\} \cup \{\$i \mid 0 \leq i \leq m+1\}$. The axiom, i.e., the target string to be achieved at the end of the computation, consists of the final state of the TM bounded by markers:

$$A = \#^{n+2} q_f \#^{n+2} \$0 \$1 \dots \$m \$_{m+1},$$

where n is the maximum length of the left-side or right-side words of any of the rules of the Turing machine.

The sequence π consists of the catenation of the right-hand sides of the TM rules bounded by markers, as follows:

$$\pi = \$0 \$1 e_1 v_1 f_1 \$1 \$2 e_2 v_2 f_2 \$2 \dots \$m e_m v_m f_m \$m \$_{m+1},$$

where $i : u_i \longrightarrow v_i$, $1 \leq i \leq m+1$ are the rules of TM and $e_i, v_i \in \Sigma \cup \{\#\}$.

If a word $w \in T^*$ is accepted by the TM, a computation starts then from a strand of the form $\#^{n+2} q_0 w \#^{n+2} \pi$, where we will refer to the subsequence starting with $\$0$ as the “program”, and to the subsequence at the left of $\$0$ as the “data”.

We construct the relation $\sim$ such that:

(i) The right-hand sides of rules of TM can be excised from the program as circular strands which then interact with the data.

(ii) When the left-hand side of a TM rule appears in the data, the application of the rule can be simulated by the insertion of the circular strand encoding the right-hand side, followed by the deletion of the left hand side.

To accomplish (i), for each rule $i : u \longrightarrow v$ of the TM, we introduce in $\sim$ the pairs

$$(C) \quad (\$_{i-1}, \$i, evf) \sim (evf, \$i, \$_{i+1}),$$

for all $e, f \in \Sigma \cup \{\#\}$.

To accomplish (ii) for each rule $i : u \longrightarrow v$ of the TM, add to the relation $\sim$ the pairs

$$(A) \quad (ceu, f, \$i ev) \sim (\$i ev, f, d),$$

$$(B) \quad (c, e, uf \$i) \sim (uf \$i, e, vfd),$$

for all $c \in \{\#\}^* \Sigma^*$, $d \in \Sigma^* \{\#\}^*$, $|c| = |d| = n$, $e, f \in \Sigma \cup \{\#\}$.

Pair (C) allows recombinations of the type

$$\{x\$_{i-1}\$_i\text{evf}\$_i\$_{i+1}y\} \iff \{x\$_{i-1}\$_i\$_{i+1}y, \bullet\$_i\text{evf}\}$$

i.e., allows excision of right-hand-sides or TM rules from the program.

Pair (A) allows recombinations of the type

$$\{x\text{ceufdy}, \bullet\$_i\text{evf}\} \iff \{x\text{ceuf}\$_i\text{evfdy}\}$$

that insert the right-hand-side of a TM rule next to its left-hand-side occurring in the data part.

Finally, pair (B) allows recombinations of the type

$$\{x\text{ceuf}\$_i\text{evfdy}\} \iff \{x\text{cevdy}, \bullet\$_i\text{euf}\}$$

that excise the left-hand-side of the TM rule in the data part.

Note that (C) serves the role of excising the necessary right-hand-side of a TM rule from the program while the tandem (A)-(B) accomplishes the replacement of a left-hand-side of a TM rule in the data by its left-hand-side, simulating thus a TM rewriting step. Indeed, for any $x, y \in \Sigma'$ we can simulate a derivation step of the TM as follows:

$$\begin{aligned} &\{x\text{ceufdy}\$ _0 \dots \$ _{i-1}\$ _i\text{evf}\$ _i\$ _{i+1} \dots \$ _{m+1}\} \implies_R \\ &\{x\text{ceufdy}\$ _0 \dots \$ _{i-1}\$ _i\$ _{i+1} \dots \$ _{m+1}, \bullet\$ _i\text{evf}\} \implies_R \\ &\{x\text{ceuf}\$ _i\text{evfdy}\$ _0 \dots \$ _{i-1}\$ _i\$ _{i+1} \dots \$ _{m+1}\} \implies_R \\ &\{x\text{cevdy}\$ _0 \dots \$ _{i-1}\$ _i\$ _{i+1} \dots \$ _{m+1}, \bullet\$ _i\text{euf}\}. \end{aligned}$$

The first step is an intramolecular recombination using contexts (C) around the repeat $\$ _i$ to excise $\bullet\$ _i\text{evf}$. Note that if the current strand does not contain a subword $\$ _i\text{evf}\$ _i$, this can be obtained from another copy of the original linear strand, which is initially present in k copies. The second step is an intermolecular recombination using contexts (A) around the repeat f , to insert $\$ _i\text{evf}$ after ceuf . The third step is an intramolecular recombination using contexts (B) around the direct repeat e to delete $\$ _i\text{euf}$ from the linear strand. Thus, the “legal” insertion/deletion succession that simulates one TM derivation step claims that any u in the data, that is surrounded by at least $n + 1$ letters on both sides may be replaced by v . This explains why in our choice of axiom we needed $n + 1$ extra symbols $\#$ to provide the contexts allowing recombinations to simulate all TM rules, including (3) and (5).

From the fact that a TM derivation step can be simulated by recombination steps we deduce that, if the TM accepts a word w , then we can start a derivation in R from

$$\#^{n+2}q_0w\#^{n+2}\pi = \#^{n+2}q_0w\#^{n+2}\$ _0\$ _1 \dots \$ _i\text{eivif}_i\$ _i \dots \$ _m\$ _{m+1}$$

and reach the axiom by only using recombinations according to R . This means that our word is accepted by R , that is, it belongs to $L_k(R)$ for some k . Note that if some rules of the TM have not been used in the derivation then they can be excised in the end, and that k should be large enough so that we do not exhaust the set of rewriting rules.

For the converse implication, it suffices to prove that starting from the strand $\#^{n+2}q_0w\#^{n+2}\pi$, no other recombinations except those that excise rules of TM from the program and those that simulate steps of the TM (forwards or backwards) in the data are possible in R .

In the beginning of the derivation we start with no circular strands and k copies of the linear strand

$$\#^{n+2}q_0w\#^{n+2}\$ _0\$ _1e_1v_1f_1\$ _1\dots \$ _{i-1}\$ _ie_iv_if_i\$ _i\$ _{i+1}\dots \$ _{m+1}, w \in T^*,$$

where $i : u_i \longrightarrow v_i$ are TM rules, $e_i, f_i \in \Sigma \cup \{\#\}$, $1 \leq i \leq m$.

Assume now that the current multiset contains linear strands of the form $\delta_0\pi$, where $\delta_0 \in \Sigma'^*$ contains only one state symbol and no $\$ _i$ symbols and

$$\pi = \$ _0r_1r_2\dots r_m\$ _{m+1},$$

with r_i either encoding the right-hand side of a rule or being the remnant of a rule, i.e., $r_i \in \{\$ _ie_iv_if_i\$ _i\} \cup \{\$ _i\}$, $1 \leq i \leq m$. Moreover, assume that the circular strands present in the multiset are of the form $\bullet\$ _ie_iv_if_i$, or $\bullet\$ _ie_iu_if_i$ with $i : u_i \longrightarrow v_i$ a rule of the TM and $e_i, f_i \in \Sigma \cup \{\#\}$.

Then,

(i) Due to the way we constructed the recombination system, circular strands cannot interact with each other.

Indeed, an insertion or deletion using (C) would require the presence of more than one $\$$ -marker in a circular strand, which contradicts our assumptions. An insertion/deletion using (A) would require the presence in some circular strand of a subword of the form $ceuf$ the length of which is at least $n+3$, greater than the length of any $\$$ -free subword of existent circular strands. An insertion/deletion using (B) would require the presence in a circular strand of a subword $evfd$ whose length of $n+3$ contradicts again the assumptions about the length of circular strands.

(ii) We cannot use (A) or (B) to insert or delete in the program.

Indeed, an insertion in π using (A) could happen in one of the following cases. In the first case, π contains a word $ceufd$ of minimum length of $2n+3$. This, however, is more than the length of the longest subword over $\Sigma \cup \{\#\}$ in π . The other possibility would be that π contains the subsequence $\$ _ievf\$ _iev$, but this contradicts the fact that no marker $\$ _i$ appears alone in π , as π always contains at least two consecutive markers. Consequently, insertions in π using (A) cannot happen.

A deletion from π using (A) would imply the presence in π of $ceuf\$_i evfd$, which contradicts the fact that no marker $\$ _i$ appears in π flanked by letters in $\Sigma \cup \{\#\}$.

An insertion in π according to (B) would require either the presence in π of the word $cevf d$ which is longer than $2n + 3$, or the presence of a $\$$ -marker flanked by letters in $\Sigma \cup \{\#\}$, both of which contradict the assumptions about the form of π . Deletion according to (B) requires the presence of $ceuf$ in π which is again too long of a $\$$ -free subword to appear in π .

(iii) We cannot use (C) to insert or delete in the data. Indeed such recombinations would require the presence in either δ_0 or in a circular strand of more than one $\$$ -marker, which is impossible.

Arguments (i) - (iii) show that the only possible recombinations are either insertions/deletions in the program using (C) or insertions/deletions in the data using (A) and (B).

Deletions in π according to (C) result in the release of circular strands $\bullet\$_i evf$ that encode right-sides of TM rules. Insertions according to (C) only mean that circular strands encoding right sides of a TM rule can be inserted back after their excision, pointing out the reversibility of the constructed system. Indeed the only other possibility of inserting into π using (C) would require the presence in π of $evf\$_i evf$ - contradiction with the form of π .

The next step is to show that the only possible insertions/deletions in the data are those simulating a rewriting step of TM using rule i .

Indeed,

(1) It is not possible to delete in δ_0 using (A), or (B), as all these operations would require the presence of a $\$ _i$ in δ_0 . Therefore only insertions in δ_0 using (A) or (B) are possible.

An insertion is possible only if δ_0 contains the left- or right-hand-side of some rewrite rule $u \longrightarrow v$. In the first case, an insertion using (A) is possible, in the second case, an insertion using (B) is possible.

In any case, after the insertion in the data part, $xceufdy\pi$ or $xcevf dy\pi$ has become $xceuf\$_i evfdy\pi$ There is no read-write head outside this segment.

Let δ_1 be the new data part, i.e., the current linear strand is of the form

$$\delta_1 \pi = xceuf\$_i evfdy \pi.$$

Note that, as δ_0 contains only one state symbol and no marker $\$ _i$, the newly formed word δ_1 contains only two state symbols (read/write heads), one in u and one in v , and only one marker $\$ _i$. (Here we use the fact that every rule $u \longrightarrow v$ of the TM has exactly one state symbol on each side.)

(2) Starting now from $\delta_1 \pi$,

(2a) No insertion in δ_1 using (A) or (B) may take place.

Indeed, another insertion in δ_1 using (A) can happen in one of two cases. The case requiring that $\$_i evf \$_i ev$ be present in δ_1 immediately leads to a contradiction as δ_1 does not contain two $\$$ -markers.

The other case is also impossible as the marker $\$_i$ “breaks” the contexts necessary for further insertions. Indeed, this second case requires the presence in δ_1 of a word of the form $ceufd$. This implies that the read/write head symbol should be both followed and preceded by at least $(n + 1)$ letters different from $\$_i$. In δ_1 , the first read/write head is in u and the number of letters in $\Sigma \cup \{\#\}$ following it is at most $|u| - 1 + |f| \leq n - 1 + 1 = n$, which is not enough. The second read/write head is in v and the number of letters preceding it is at most $|e| + |v| - 1 \leq 1 + n - 1 = n$, which is not enough.

Also, no insertion in δ_1 using (B) is possible. Indeed, such an insertion can happen in two cases. The case where δ_1 contains $uf \$_i euf \$_i$ immediately leads to a contradiction with the form of δ_1 . The other case requires the presence in δ_1 of a word of the form $cevfd$. This also leads to a contradiction, as one can show using an argument like the one above).

(2b) We can delete in δ_1 using (B) or (A).

There is only one possible location for either deletion, and they result in replacing $xceuf \$_i evfdy\pi$ by $xceufdy\pi$ or $xcevfdy\pi$.

All in all, in two steps (1)-(2), the only possible recombinations on the data part either replace u with v , v with u or keep the strand unchanged, where $u \rightarrow v$ is a rewrite rule of the TM.

The arguments above imply that the only possible operations on the data simulate legal rewritings of the TM by tandem recombination steps that necessarily follow each other.

Together with the arguments that the only operations affecting the program are excisions of circular strands encoding TM rules, and that the circular TM rules do not interact with each other, this proves the converse implication.

From the definition of the Turing machine we see that n , the maximum length of a word occurring in a TM rule, equals 4, which completes the proof of the theorem.

□

The preceding theorem implies that if a word $w \in T^*$ is in $L(TM)$, then $\#^6 q_0 w \#^6 \pi$ belongs to $L_k(R)$ for some k and therefore it belongs to $L^i(R)$ for any $i \geq k$. This means that, in order to simulate a computation of the Turing machine on w , any sufficiently large number of copies of the initial strand will do. The assumption that sufficiently many copies of the input strand are present at the beginning of the computation is in accordance with the fact that there are multiple copies of each strand available during the (polytene chromosome) stage where unscrambling occurs.

The proof that a reversible guided recombination system can simulate the computation of a Turing machine suggests that the micronuclear gene, present in multiple copies, consists of a sequence encoding the input data, combined with a sequence encoding a program, i.e., a list of encoded computation instructions. The “computation instructions” can be excised from the micronuclear gene and become circular “rules” that can recombine with the data. The process continues then by multiple intermolecular recombination steps involving the linear strand and circular “rules”, as well as intramolecular recombinations within the linear strand itself. The resulting linear strand, which is the functional macronuclear copy of the gene, can then be viewed as the output of the computation performed on the input data following the computation instructions excised as circular strands.

The last step, telomere addition and the excision of the strands between the telomere addition sites, can easily be added to our model as a final step consisting of the deletion of all the markers, rule delimiters and remaining rules from the output of the computation. This would result in a strand that contains only the output of the computation (macronuclear copy of the gene) flanked by end markers (telomere repeats).

The conclusion that the process of gene unscrambling in ciliates can simulate a Turing machine suggests that such processes may have many potential uses in solving computational problems that occur in complex evolving systems.

Acknowledgements. Rani Siromoney, Gilles Brassard for suggestions, Gheorghe Păun for comments. Grzegorz Rozenberg, Richard Lipton, David Prescott, Hans Lipps, and Ed Curtis for discussion.

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4.1 Figure legends

Figure 1: DNA hybridization in a molecular computer. PCR primers are indicated by arrows.

Figure 2: Overview of gene unscrambling. Dispersed coding MDSs 1-7 reassemble during macronuclear development to form the functional gene copy (top), complete with telomere addition to mark and protect both ends of the gene.

Figure 3: A ciliate computer? Correct gene assembly in *Stylonychia* (inset) requires the joining of many segments of DNA guided by short sequence repeats, only at the ends. Telomeres, indicated by thicker lines, mark the termini of correctly assembled gene-sized chromosomes. Note the similarities to DNA computations that specifically rely on pairing of short repeats at the ends of DNA fragments, as in Adleman's experiment.

Figure 4: Model for unscrambling in α -TP (adapted from [14]).