

Cell Complexes and Membrane Computing for Thinning 2D and 3D Images

Raúl Reina-Molina, Daniel Díaz-Pernil and Miguel A. Gutiérrez-Naranjo

Abstract—In this paper, three different research areas are brought together in order to propose a new point of view on a classical problem: Digital Image Analysis, Algebraic Topology and Membrane Computing. The chosen problem is the thinning of multi-dimensional black and white digital images. Thinning is usually considered as a preprocess in Digital Image Analysis, but it has important applications in the study of fingerprints, vessels and arteries or cartography. It does not consist on a mere removal of outer pixels, since the thinning process must preserve, in some sense, the *meaning* of the image. This means that the most important features concerning a shape are its *topology* (represented by connected components, holes, etc.) and its *geometry* (elongated parts, ramifications, etc.), thus they must be preserved.

Recently, a new algorithm for thinning multi-dimensional black and white digital images by using cell complexes was presented. Such cell complexes allow a discrete partition of the space and the algorithm preserves topological and geometrical properties of the complex. Algebraic Topology allows us to split the space in discrete units and to define mechanical operators on such space units of different dimensions.

As a third research area, a recent bio-inspired computational model, Membrane Computing, is used to handle with such operators and provide an alternative point of view for handling with them. Algebraic Topology and Membrane Computing deal with compartments of the Euclidean space on their foundations, but their inspiration and motivation are quite different. Membrane Computing techniques are inspired in the flow of metabolites between cells of a living tissue or between the organelles in an eucaryotic cell. This flow of metabolites takes place in parallel in Nature and can be interpreted as a flow of information for computational purposes. Instead of a set of few instructions with complex data structures, the computation steps in a Membrane Computing device are regulated by a set of rules with a notation close to biochemical reactions. From a computational point of view, such reactions can be read as a set of *if A then B* rules where *A* and *B* are very simple data.

In this paper, we present a parallel bio-inspired Membrane Computing implementation of an algorithm working with cubical complexes which solves the problem of thinning digital images.

Index Terms—Digital Image Analysis, Thinning Algorithm, Membrane Computing, Algebraic Topology

I. INTRODUCTION

Computer vision [3] is a vivid research area that tries to acquire, process, analyze and understand images from the real world in order to produce useful information. From the computational side, a 2D digital image can be roughly defined

as a function from a two dimensional surface which maps each point from the surface onto a set of attributes as bright or color. Analogously, a 3D image maps a region of a three-dimensional space onto a set of attributes. Different algorithms on such mappings provide a great variety of useful *information* in computer vision areas as biometrics, surveillance or medical imaging, among others.

In this paper, we focus on the problem of obtaining the so-called *skeleton* of a black and white image. Skeletonization is one of the approaches for representing a shape with a small amount of information, by converting the initial image into a more compact representation and keeping the meaning features. The conversion should remove redundant information, but it should also keep the basic structure. Skeletonization is usually considered as a pre-process in pattern recognition algorithms, but its study is also interesting by itself for the analysis of line-based images, as coronary arteries [4], fingerprints classification [5] or cartography [6]. The concept of skeleton was introduced by Blum in [7], under the name of medial axis transform. There are many different definitions of the skeleton of a black and white image and many skeletonizing algorithms¹, but in general, the image *B* is a skeleton of the image *A*, if the former has fewer black pixels than the latter, preserves its topological properties and, in some sense, keeps its *meaning*. The most important features concerning a shape are its *topology* (represented by connected components, holes, etc.) and its *geometry* (elongated parts, ramifications, etc.), thus they must be preserved. When the skeletonizing process is made by the iterative removal of non-significant elements of the image, the process is known as *thinning*. In this paper, we present a bio-inspired implementation of the algorithm presented by Liu in [1], [2] for thinning images based on Membrane Computing techniques.

The key notion of this algorithm is the *cell complex*, a basic concept in Algebraic Topology which can be seen as a mathematical abstraction of a space unit. In the original work, the thinning algorithm is designed for cell complexes of any kind, however we restrict to cubical complexes.

The main contribution of this paper is to bridge an emergent bio-inspired research area as Membrane Computing with real-world Image Analysis via well-known concepts of Algebraic Topology. This is not the first time in which life-based methods are applied to Algebraic Topology. J. Chao and J. Nakayama connected Natural Computing and Algebraic Topology using Neural Networks [9]. Some years after, K.G. Subramanian

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¹A detailed description of skeletonizing algorithms is out of the scope of this paper. For a survey in this topic, see e.g., [8].

et al. linked in [10] Digital Image and Natural Computing. Christinal *et al.* started a new way where algebraic-topological processes have been parallelized via Membrane Computing (see [11]–[14]).

As it will be shown below, Membrane Computing² techniques are inspired in the movement of metabolites between cells of a living tissue or between the organelles in an eucaryotic cell. This flow of metabolites takes place in parallel in Nature and can be interpreted as a flow of information for computational purposes. Instead of a set of few instructions with complex data structures, the computation steps in a Membrane Computing device are regulated by a set of many rules with a notation close to biochemical reactions. From a computational point of view, such reactions can be read as a set of *if A then B* rules where *A* and *B* are very simple data.

Many problems in the processing of digital images have features which make it suitable for techniques inspired by nature. The subset of the integer plane or space taken to be the *support* of the image and the set of possible features associated to each 2D or 3D point can be considered finite and hence, the transformation of an image into another can be performed in a *discrete* way. Other of such features is that the treatment of the image can be parallelized and locally modified. Regardless how large is the picture, the process can be performed in parallel in different local areas of it. Another interesting feature is that the information of the image can also be easily encoded in the data structures used in Natural Computing. In the literature, we can find many examples of the use of Natural Computing techniques dealing with problems associated to the treatment of digital images. One of the classic examples is the use of cellular automata [17]. Other efforts are related to artificial neural networks as in [18]. In this paper, we use *tissue P systems*, a computational model in the framework of Membrane Computing.

The paper is organized as follows: Firstly, we recall the computational bio-inspired model used in this paper (Section II) and some basics on Algebraic Topology (Section III). In Section IV, a brief description of Liu's algorithm is given and, in Section V, our bio-inspired implementation is provided. Next, an outline of the computation is presented, finishing with conclusions and future work.

II. MEMBRANE COMPUTING

Nature is a big source of inspiration for new computational paradigms. It *acts* by performing changes (from microscopic biochemical reactions to ecological global variations) which can be interpreted as *computations*. Natural Computing³ abstracts the way Nature operates, providing ideas for new computing models. It involves research where the physical support is non standard, as *DNA-based Molecular Computing* or *Quantum Computing*; but almost all the research lines in Natural Computing are currently supported in silicon-based computers. Among them, we can cite *Artificial Neural*

Networks, *Genetic Algorithms*, *Swarm Intelligence*, *Artificial Immune Systems*, *Amorphous Computing*, *Membrane Computing* or *Cellular Automata*. All these computational paradigms have in common the use of an alternative way of encoding the information and the use of intrinsic parallelism of natural processes. In this paper, we will use the theoretical framework of Membrane Computing for handling digital images.

Membrane Computing [16] is a model of computation inspired by the structure and functioning of cells as living organisms able to process and generate information. In particular, it focuses on membranes, which are involved in many reactions taking place inside various compartments of a cell. They act as selective channels of communication between different compartments as well as between the cell and its environment. The computational devices in Membrane Computing are called *P systems*. Roughly speaking, a P system consists of a membrane structure, in whose compartments one places multisets of objects which evolve according to given rules. These multisets encode the information and the rules dealing with them perform the computation. Following a biological inspiration, the multisets of objects represent the chemicals placed in a vesicle of a living being. Such chemicals are sent to other vesicles or transformed according to biochemical reactions, represented here by computational rules. These rules are usually applied in a synchronous non-deterministic maximally parallel manner. In this paper, the so-called (because of their membrane structure) *tissue P Systems* [20] are considered.

A. Tissue P systems

The chosen P system model for our Membrane Computing implementation of Liu's algorithm is the *tissue P systems model*. This Membrane Computing model has been widely used to solve computational problems in other areas (see e.g. [21], [22]), but recently, they have been also used in the study of digital images (see, e.g., [14], [23], [24] and references therein).

In the tissue P systems used in this paper, the flow of information (the application of the rules) are regulated by *promoters* and *inhibitors* (see [25]) and a (weak) priority relation between rules is considered. Below, a formal description of the computational model is provided but, roughly speaking, a *tissue P system* of the type used in this paper can be seen as a set of enclosed regions of the space, called *cells*, where some objects are placed, together with a set of communication rules which settles which objects can be interchanged between cells. The cells are the nodes of a virtual graph, where the edges connecting the cells are determined by the communication rules of the P system, i.e., as usual in tissue P systems, the edges linking cells are not provided explicitly. Rules are syntactically expressed as a tuple $(pro \neg inh \mid i, u/v, j)$. If a rule $(pro \neg inh \mid i, u/v, j)$ is given, then cells *i* and *j* are considered linked. The application of a rule $(pro \neg inh \mid i, u/v, j)$ consists of trading the multiset *u* (initially in the cell *i*) against the multiset *v* (initially in *j*). After the application of the rule, the multiset *u* disappears from the cell *i* and it appears in the cell *j*. Analogously, the multiset *v* disappears from the cell *j* and

²We refer to [15] for basic information in this area, to [16] for a comprehensive presentation and the web site <http://ppage.psystems.eu> for the up-to-date information.

³An introduction on Natural Computing can be found in [19].

it appears in the cell i . The trade can also be between one cell and the environment, labeled by 0. The rule is applied if in the cell with label i the objects of the multiset pro are present in the cell i (*promoters*), while any of the objects in the multiset inh do not appear in the cell (*inhibitors*). The promoters or the inhibitors are not modified by the application of the rule. If the promoters and inhibitors are empty, we will write $(i, u/v, j)$ instead of $(\emptyset - \emptyset | i, u/v, j)$. Finally, we write $(pro | i, u/v, j)$ or $(-inh | i, u/v, j)$ when only promoters or inhibitors appear, respectively.

The biological inspirations of this model are intercellular communication and cooperation between neurons. A rule $(pro - inh | i, u/v, j)$ is applied if the reactants u and v are present, but it is also necessary the presence of all the promoters pro and none of the inhibitors inh in the cell i . The promoters are not *consumed* nor *produced* by the application of the rule, but if they are not in the cell, the rule cannot be applied. In one step, each reactant in a membrane can only be used for one rule, but if several rules need the presence of the same promoter, then the presence of one unique copy of the promoter suffices for the application of the rules.

Before giving the formal definition of such P systems, let us recall some technical preliminaries. An *alphabet*, Σ , is a non empty set, whose elements are called *symbols*. An ordered sequence of symbols is a *string*. The set of all the strings on Σ is denoted by Σ^* . The number of symbols in a string u is the *length* of the string, and it is denoted by $|u|$. As usual, the empty string (with length 0) will be denoted by λ . A *multiset* m over a set A is a pair (A, f) where $f : A \rightarrow \mathbb{N}$ is a mapping. If $m = (A, f)$ is a multiset then its *support* is defined as $supp(m) = \{x \in A \mid f(x) > 0\}$ and its *size* is defined as $\sum_{x \in A} f(x)$. A multiset is empty (resp. finite) if its support is the empty set (resp. finite). If $m = (A, f)$ is a finite multiset over A , and $supp(m) = \{a_1, \dots, a_k\}$, then it will be denoted as $m = a_1^{f(a_1)}, \dots, a_k^{f(a_k)}$. That is, superscripts indicate the multiplicity of each element, and if $f(x) = 0$ for any $x \in A$, then this element is omitted.

Formally, a *tissue P system with promoters, inhibitors and priorities* of degree $q \geq 1$ is a tuple of the form

$$\Pi = (\Gamma, \Sigma, \mathcal{E}, w_1, \dots, w_q, \mathcal{R}, Pri, i_{in}, i_{out})$$

where q is the number of cells (or membranes) of the P system and

- 1) Γ is a finite alphabet, whose symbols will be called objects. These objects can be placed in the cells or in the surrounding space (called the *environment*);
- 2) $\Sigma \subseteq \Gamma$ is the input alphabet. The *input* of the computation performed by the P system is encoded by using this alphabet;
- 3) $\mathcal{E} \subseteq \Gamma$ is a finite alphabet representing the set of the objects in the environment. Following a biological inspiration, the objects in the environment are available in an arbitrary large amount of copies;
- 4) w_1, \dots, w_q are strings over Γ representing the multisets of objects placed inside the cells at the starting of the computation;
- 5) \mathcal{R} is a finite set of rules of the following

form: $(pro - inh | i, u/v, j)$ for $0 \leq i \neq j \leq q$, $pro, inh, u, v \in \Gamma^*$

- 6) Pri is a finite set of relations $R_i > R_j$, where R_i and R_j are rules from \mathcal{R} . It means that if R_i and R_j can be applied, then the application of R_i has *priority* on the application of R_j .
- 7) $i_{in} \in \{1, 2, \dots, q\}$ denotes the input cell, i.e., the cell where the *input* of the computation will be placed.
- 8) $i_{out} \in \{1, 2, \dots, q\}$ denotes the output cell, i.e., the cell where the *output* of the computation will be placed.

Rules are used as usual in the framework of membrane computing, that is, in a maximally parallel way (a universal clock is considered). A *configuration* is an instantaneous description of the tissue P system and it is represented as a tuple (w_1, \dots, w_q) . Given a configuration, we can perform a computation step and obtain a new configuration by applying the rules in a parallel manner as it is shown above. A configuration is *halting* when no rules can be applied to it. A *computation* is a sequence of computation steps such that either it is infinite or it is finite and the last step yields a halting configuration (i.e., no rules can be applied to it). Then, a computation halts when the P system reaches a halting configuration. The output of a computation is the multiset placed in the output cell in a halting configuration collected from its halting configuration by reading the objects contained in the output cell.

B. Example

Let us consider the following tissue P system with promoters, inhibitors and priorities of degree 3

$$\Pi = (\Gamma, \Sigma, \mathcal{E}, w_1, w_2, w_3, \{R_1, \dots, R_5\}, \{R_1 > R_2\}, 2, 1)$$

where $\Gamma = \{a, b, c, d, e, f\}$, $\Sigma = \{b, c\}$ and $\mathcal{E} = \{f\}$. The multisets in the initial configuration are $w_1 = df$, $w_2 = a^2e$ (two copies of a and one copy of e) and $w_3 = a^2e^2cd$. The rules are $R_1 \equiv (b, \neg d | 2, a^2/d, 1)$, $R_2 \equiv (b | 2, a^2/f, 1)$, $R_3 \equiv (d, \neg b | 3, a^2/c, 2)$, $R_4 \equiv (a, \neg c | 2, be/c, 3)$ and $R_5 \equiv (c, \neg f | 3, e/f^2, 0)$. Finally, the input cell is $i_{in} = 2$ and the output cell is $i_{out} = 1$. Notice that the rules determine a virtual graph with the cells as nodes. From R_1 and R_2 we can consider an edge between cell 1 and cell 2. Analogously, R_3 and R_4 determine an edge between cell 2 and cell 3. We also know by rule R_5 that cell 3 can trade some objects with the environment.

Let us consider as input of our computation the multiset bc will be placed in the input cell 2. By considering the input, the initial configuration has three multisets $C_0 = \langle w_1, w_2, w_3 \rangle$ with $w_1 = df$, $w_2 = a^2bce$ and $w_3 = a^2e^2cd$. By inspection of the occurrence of the objects in the cells, the *applicable* rules from this first configuration are R_1 , R_2 , R_3 and R_5 , but R_1 and R_2 cannot be applied together, because only two copies of a occur in the cell 2. By considering the priority relation, R_1 has priority on R_2 and R_1 , R_3 and R_4 are applied. After trading the corresponding objects, we get the configuration $C_1 = \langle w'_1, w'_2, w'_3 \rangle$ with $w'_1 = a^2f$, $w'_2 = a^2bde$ and $w'_3 = c^2df^4$. From this configuration, R_2 and R_4 satisfy the conditions and both are applied. The new configuration

is $C_2 = \langle w_1'', w_2'', w_3'' \rangle$ with $w_1'' = a^4$, $w_2'' = cdf$ and $w_3'' = bcde f^4$. No more rules can be applied and C_2 is the halting configuration. The multiset a^4 in the output cell 1 in the halting configuration is the output of this tissue P system for the input bc .

III. CUBICAL COMPLEXES

As pointed above, the key concept in our adaptation of Liu's algorithm is the *cell complex* which can be seen as a mathematical abstraction of a space unit. This space unit is built in some n dimensional space and embedded in a space of higher dimension, as a 2-dimensional square can be embedded in a 3D space. On such abstractions, several operators are defined, as the *border* one, which associates, for example, a 3D cell (cube) with six 2D cells (squares), or properties to define *free cells* or *isolated cells*.

Before presenting Liu's algorithm, let us recall some basics on the combinatorial structure on a topological space (see [26]). An *elementary closed interval* is a closed interval $\bar{I} \subset \mathbb{R}$ of the form $\bar{I} = [l, l+1]$ or $\bar{I} = [l, l]$ for some $l \in \mathbb{Z}$. The former are called *non-degenerated*, while the latter are called *degenerated*. The interval $[l, l]$ that contains only one point will be denoted by $[l]$. Degenerated elementary closed intervals are simply points with 0 dimensions. Non-degenerated elementary closed intervals are segments (objects with one dimension). An elementary cube $\bar{\sigma}$ is a finite product of elementary closed intervals:

$$\bar{\sigma} = \bar{I}_1 \times \bar{I}_2 \times \cdots \times \bar{I}_d \subset \mathbb{R}^d$$

where \bar{I}_j is an elementary closed interval, $j \in \{1, \dots, d\}$. The set of all elementary cubes in \mathbb{R}^d is denoted by \mathcal{K}^d . The set of all elementary cubes is

$$\mathcal{K} = \bigcup_{d=1}^{\infty} \mathcal{K}^d$$

Given an elementary cube $\bar{\sigma} = \bar{I}_1 \times \bar{I}_2 \times \cdots \times \bar{I}_d$ in \mathbb{R}^d , its *embedding number* d is denoted by $\text{emb } \bar{\sigma}$. The dimension of $\bar{\sigma}$ is defined to be the number of non-degenerated intervals in its definition and it is denoted by $\dim \bar{\sigma}$. The set of all elementary cubes with dimension p is denoted by \mathcal{K}_p . The set of all elementary cubes in \mathbb{R}^d with dimension p is denoted by \mathcal{K}_p^d . Elementary cubes can be decomposed into lower-dimensional objects. If $\bar{\delta}$ and $\bar{\sigma}$ are two elementary cubes in \mathbb{R}^d of any dimension and $\bar{\delta} \subset \bar{\sigma}$, then $\bar{\delta}$ is a *face* of $\bar{\sigma}$. If $\bar{\delta}$ is a face of $\bar{\sigma}$ and $\bar{\delta} \neq \bar{\sigma}$, then $\bar{\delta}$ is a *proper face* of $\bar{\sigma}$. If it is a face of $\bar{\sigma}$ and $\dim \bar{\delta} = \dim \bar{\sigma} - 1$ then $\bar{\delta}$ is a *primary face* of $\bar{\sigma}$. Given an elementary cube $\bar{\sigma} \in \mathcal{K}_p^d$, the set of all primary faces of $\bar{\sigma}$ is called the *border* of $\bar{\sigma}$ and it is denoted by $\partial \bar{\sigma}$.

Let \bar{I} be an elementary closed interval. The associated *elementary interval* is

$$I = \begin{cases} (l, l+1) & \text{if } I = [l, l+1], \\ [l] & \text{if } I = [l]. \end{cases}$$

If $\bar{\sigma} = \bar{I}_1 \times \bar{I}_2 \times \cdots \times \bar{I}_d \subset \mathbb{R}^d$ is an elementary cube, then the associated *elementary cell* is $\sigma = I_1 \times I_2 \times \cdots \times I_d$. The *dimension* of an *elementary cell* σ is defined as $\dim \bar{\sigma}$, i.e., the dimension of the associated elementary cube. The border

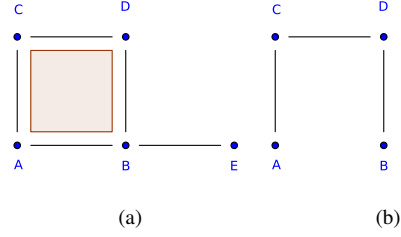


Fig. 1. (a) Example of cubical complex. (b) Elementary collapse example: E collapses onto BE and AC collapses onto $ABDC$ in the image at the left, producing the image at the right.

for an elementary cell σ can also be defined as the set $\partial \sigma = \{\delta : \bar{\delta} \in \partial \bar{\sigma}\}$. A *cubical complex* is a set of elementary cells such that, given an elementary cell σ in the complex, all of its principal faces (the cells in $\partial \sigma$) are in the complex.

For the sake of simplicity, hereafter we will say *cells* instead of elementary cells, bearing in mind that we refer to such kind of objects. When a cell is not a proper face of any cell in a given cell complex, it will be called *isolated cell*. A cell that is a proper face of exactly one cell in the complex is called *free face*. It is known that if δ is a free face in a cell complex and δ is a proper face of σ , then σ is an isolated cell and $\dim \delta = \dim \sigma - 1$.

If K is a cubical complex and δ a free cell in K , σ is the only cell in K such that δ is a proper face of σ and $K' = K \setminus \{\delta, \sigma\}$, then K' is obtained from K via a process called *elementary collapse* of σ by δ . A pair of cells $\langle \delta, \sigma \rangle$ in a cubical complex K is said to be a *simple pair* if δ is a free cell in K and σ is the only cell such that $\delta \in \partial \sigma$. The cell σ is called the *facet* of the simple pair. Simple pairs removal does not change the topology of the given cell complex.

A. Example

Figure 1 (left) shows the cubical complex

$$K = \{ABCD, AC, CD, BD, AB, BE, A, B, C, D, E\}$$

This cubical complex has 1 cell of dimension 2 ($ABCD$), 5 cells of dimension 1 (AC, CD, BD, AB, BE) and 5 cells of dimension 0 (A, B, C, D, E). The cells $ABCD$ and BE are isolated. The cells AC, CD, BD, AB and E are free faces, but A, B, C and D are not free faces, since they are proper faces of more than 1 cell complex. The cell E is a free face of BE and, hence, we can consider the elementary collapse of BE by E . The effect of such elementary collapse is the removal of E and BE from the cell complex K . Analogously, AC is a free face of $ABCD$. The elementary collapse of $ABCD$ by AC is the removal of both cells ($ABCD$ and AC) from K . Figure 1 (right) shows the final cubical complex obtained after both collapses.

IV. LIU'S ALGORITHM

As shown in the introduction, finding a skeleton for a given d -D image is an useful tool in various areas. In this paper we will introduce a bioinspired parallel solution to the problem

Algorithm 1 Cell complex thinning algorithm

Require: K cell complex, $\varepsilon_a, \varepsilon_r > 0$

for all $\sigma \in K$ **isolated do**

$I(\sigma) \leftarrow 0$

end for

iter $\leftarrow 1$

repeat

Let $S = \{\langle \delta, \sigma \rangle : \langle \delta, \sigma \rangle \text{ is a simple pair}\}$

Let $S' = \{\langle \delta, \sigma \rangle \in S : I(\sigma) = 0 \vee \text{MP}_{abs}(\sigma) < \varepsilon_a \vee \text{MP}_{rel}(\sigma) < \varepsilon_r\}$

Let S'' be the greatest subset of S' such that

$\forall \langle \delta_1, \sigma_1 \rangle, \langle \delta_2, \sigma_2 \rangle \in S'', \delta_1 = \delta_2 \rightarrow \sigma_1 = \sigma_2$

for all $\sigma \in \pi_2(S)$ **do**

$R(\sigma) \leftarrow \text{iter}$

end for

$K \leftarrow K \setminus \{\sigma, \delta : \langle \delta, \sigma \rangle \in S''\}$

for all $\sigma \in K$ **new isolated cell do**

$I(\sigma) \leftarrow \text{iter}$

end for

iter $\leftarrow \text{iter} + 1$

until $S' = \emptyset$

of finding a skeleton for a given d -D image, denoted as d – **ThCub**.

In [1], a cell complex is processed in order to obtain another complex with the same topology, and the same geometry. The input cell complex is processed by consecutive parallel removal of certain cells. The removal process is a simple homotopy equivalence and the set of non-removed cells will make the skeleton.

We use π_2 for the second coordinate projection, defined as $\pi_2(\langle \delta, \sigma \rangle) = \sigma$ for the pair $\langle \delta, \sigma \rangle$.

Let K be a cubical cell complex and let ∂ be its border operator. As previously seen, if only simple pairs of cells are removed, the topology is kept. Nonetheless, in order to keep the *information* of the image, some geometrical features must be preserved. For such a geometry preservation it is necessary to require some additional properties to those cells to be removed.

The basic idea of the algorithm is to define an iterative process where *outer* cells are removed. Here, the idea of *outer* cells makes reference to simple pairs, since in a simple pair $\langle \delta, \sigma \rangle$ the cell δ is a *terminal* cell as it does not lie in the border of any other one rather than σ .

In the process of iterative thinning, given a cell σ , we will denote the later iteration when σ is the facet of a simple pair by $R(\sigma)$. The earlier iteration when σ becomes isolated will be denoted by $I(\sigma)$. Liu *et al.* describe in [2] the relation between $I(\sigma)$ and $R(\sigma)$, and the maximum *isotropic* elongation in $p+1$ and p directions, respectively, since $\dim \sigma = p$. Thus, if σ is a p -cell in a cell complex, $I(\sigma)$ measures the shortest discrete distance from σ to the object boundary. This distance gives an idea of the size of the maximum disk centered at σ and inscribed in the object. On the other hand, $R(\sigma)$ measures the longest distance from σ to the object boundary going along the skeleton $(p-1)$ -cells.

From the observation of the behaviour of previous measures, Liu *et al.* defined two *difference measures*. The absolute one, $R(\sigma) - I(\sigma)$, is called *absolute medial persistence* and is denoted by MP_{abs} . On the other hand, *relative medial persistence* is defined as $1 - \frac{I(\sigma)}{R(\sigma)}$ and denoted by MP_{rel} . Both of them measure the duration in which a cell remains isolated during thinning process.

In Algorithm 1, we present a sketch of the process of thinning implemented by Liu *et al.* in [1]. It starts by initializing the isolated cells. Next, the thinning iterations start. In each iteration, all simple pairs are selected, all the pairs where the facet cell has one of the medial persistence measures less than given thresholds or are initially isolated are chosen, taking into account that, if $\langle \delta, \sigma_1 \rangle$ and $\langle \delta, \sigma_2 \rangle$ are simple pairs that can be removed, only one of them is selected to be deleted from the complex, non-deterministically chosen. Finally, the cells in selected simple pairs are removed from the cell complex.

If no cell has been removed, the thinning iterations stop, otherwise, the iteration counter increases and the thinning iterations continue. When the algorithm halts, a cell complex representing the skeleton for the initial one is obtained.

V. MEMBRANE COMPUTING ALGORITHM

In previous sections, two key concepts have been reviewed. On the one hand, cell complexes provide an useful link between continuous spaces and discrete structures where combinatorial algorithms may be developed using well-established properties and results by continuous topology. On the other hand, it has been settled a theoretical framework for working with images, considering them as a function from a topological discrete space to a set of *features*.

The main goal is, starting from a d -dimensional binary image, to build other image which represents a skeleton for the original one. In this process, a cell complex from the original image is obtained, it is skeletonized and a new image from the last skeleton is obtained. Of course, no topological or shape information must be lost. The set of points⁴ for our source images will be the set $T_n^d = \{0, 1, \dots, n-1\}^d \subset \mathbb{Z}^d$ equipped with a *cubic* neighbourhood function, described as follows: Two points $\mathbf{i} = (i_1, \dots, i_p, \dots, i_d)$ and $\mathbf{j} = (j_1, \dots, j_p, \dots, j_d)$ are to be said $(2d)$ -adjacent if $i_l = j_l$ for $l \neq p$ and $|i_p - j_p| = 1$, for some $p \in \{1, 2, \dots, d\}$. More formally, the neighbourhood function is given by

$$N(i_1, \dots, i_d) = \left\{ (j_1, \dots, j_d) \in T_n^d : \sum_{p=1}^d |j_p - i_p| = 1 \right\}$$

This neighborhood function, when restricted to $d = 2$, gives the 4-adjacency, and 6-adjacency when $d = 3$.

Let $I : T_n^d \rightarrow \{0, 1\}$ be a d -D binary image of size n^d , where the set of points in the object (or black points) is $I^{-1}(1)$. Let $K = K(I)$ be the cubic cell complex built from I . In K , the 0-cells represent points in the object, the 1-cells represent pairs of $(2d)$ -adjacent points, the 2-cells represent unit squares where its edges are pairs of $(2d)$ -adjacent points, and so on. In

⁴The reader is supposed to be familiar with concepts of Image Algebra. For a detailed text, see [27].

general, each p -cell is a p -dimensional unit hypercube whose edges are pairs of $(2d)$ -adjacent points.

As we have shown in section III, a (cubical) cell σ is a product of intervals $I_1 \times I_2 \times \dots \times I_d$. We will encode σ as the tuple

$$\mathbf{i} = (\inf I_1 + \sup I_1, \dots, \inf I_d + \sup I_d)$$

Let us denote $\mathcal{K}^d(n)$ the set of cubic cells such that the extremes of its elementary intervals are integers in $\{0, 1, \dots, n-1\}$.

Lemma 1. *The function $\mathbb{T} : \mathcal{K}^d(n) \rightarrow T_{2n-1}^d$ defined by*

$$\mathbb{T}(I_1 \times I_2 \times \dots \times I_d) = (\inf I_1 + \sup I_1, \dots, \inf I_d + \sup I_d)$$

is a bijection.

Proof. First of all, \mathbb{T} is well defined as, for all $\sigma = I_1 \times \dots \times I_d$ is $0 \leq \inf I_p, \sup I_p \leq n-1$ for each $p \in \{1, \dots, d\}$, therefore $0 \leq \inf I_p + \sup I_p \leq 2(n-1)$ and $\mathbb{T}(\sigma) \in T_{2n-1}^d$.

Let us prove that \mathbb{T} is injective. Let $\sigma_1, \sigma_2 \in \mathcal{K}^d(n)$ be two different cells. Let us denote $I_p(\sigma)$ as the p -th elementary interval in σ . As $\sigma_1 \neq \sigma_2$, let us suppose that $I_p(\sigma_1) \neq I_p(\sigma_2)$ for some $p \in \{1, \dots, d\}$. If both elementary intervals are degenerated, as they are different, both $\sup I_p(\sigma_1)$ and $\sup I_p(\sigma_2)$ are different too, so $\mathbb{T}(\sigma_1) \neq \mathbb{T}(\sigma_2)$. If both intervals are non-degenerated, as they are different and its extremes are pairs of consecutive integers, $\sup I_p(\sigma_1) \neq \sup I_p(\sigma_2)$ and $\mathbb{T}(\sigma_1) \neq \mathbb{T}(\sigma_2)$. If one elementary interval is degenerated and the other is non-degenerated, $\mathbb{T}(\sigma_1) \neq \mathbb{T}(\sigma_2)$ as the p -th coordinate of one of them is even as the corresponding coordinate of the other is odd. Note that $\sup(l, l+1) + \inf(l, l+1) = 2l+1$ while $\sup[l] + \inf[l] = 2l$. Therefore, \mathbb{T} is a bijection.

To prove that \mathbb{T} is surjective, it is enough to find a cell $\sigma \in \mathcal{K}^d(n)$ for a given tuple $(l_1, \dots, l_d) \in T_{2n-1}^d$. Let us denote $\mathbf{I}(s)$ as follows

$$\mathbf{I}(s) = \begin{cases} \left(\frac{s-1}{2}, \frac{s+1}{2} \right) & \text{if } s \equiv 1 \pmod{2} \\ \left[\frac{s}{2} \right] & \text{if } s \equiv 0 \pmod{2} \end{cases}$$

Let us define $\sigma = \mathbf{I}(l_1) \times \dots \times \mathbf{I}(l_d)$. As $0 \leq l_p \leq 2n-2$ for each $p \in \{1, \dots, d\}$, $\sigma \in \mathcal{K}^d(n)$. If l_p is odd, $\sup \mathbf{I}(l_p) + \inf \mathbf{I}(l_p) = l_p$ and, if l_p is even then $\sup \mathbf{I}(l_p) + \inf \mathbf{I}(l_p) = l_p$ too. Hence, $\mathbb{T}(\sigma) = (l_1, \dots, l_d)$. \square

Note that, as the dimension of a cell is the number of non-degenerated intervals in its definition, the dimension of a cell given as a tuple is the amount of odd coordinates.

As an example, the cell $\sigma = (0, 1) \times [2] \times (5, 6)$ in \mathbf{R}^3 is encoded as $\mathbb{T}(\sigma) = (0+1, 2+2, 5+6) = (1, 4, 11)$.

As we have defined above, the border operator $\partial : \mathcal{K}^d \rightarrow 2^{\mathcal{K}^d}$ returns the set $\partial\sigma$ of principal faces for any cell σ . More over, it is easy to prove that $\partial(\mathcal{K}^d(n)) \subset 2^{\mathcal{K}^d(n)}$. Hence the following commutative diagram gives us an appropriate definition of border operator for cubical cells represented as tuples:

$$\begin{array}{ccc} \mathcal{K}^d(n) & \xrightarrow{\mathbb{T}} & T_{2n-1}^d \\ \downarrow \partial & & \downarrow \partial \\ 2^{\mathcal{K}^d(n)} & \xrightarrow{2^{\mathbb{T}}} & 2^{T_{2n-1}^d} \end{array}$$

where $2^{\mathbb{T}}(K) = \{\mathbb{T}(\sigma) : \sigma \in K\}$

The operator $\partial : T_{2n-1}^d \rightarrow 2^{T_{2n-1}^d}$ can be defined, hence, as follows:

$$\partial \mathbf{i} = 2^{\mathbb{T}} \circ \partial \circ \mathbb{T}^{-1}(\mathbf{i})$$

where its explicit expression is given by

$$\begin{aligned} \partial \mathbf{i} = \{ & \mathbf{d}_p^-(\mathbf{i}) : 1 \leq p \leq d \wedge i_p \equiv 1 \pmod{2} \} \cup \\ & \{ \mathbf{d}_p^+(\mathbf{i}) : 1 \leq p \leq d \wedge i_p \equiv 1 \pmod{2} \} \end{aligned}$$

where $\mathbf{i} = (i_1, \dots, i_d)$; \mathbf{d}_p^- and \mathbf{d}_p^+ are defined as follows

$$\mathbf{d}_p^-(i_1, \dots, i_p, \dots, i_d) = (i_1, \dots, i_p - 1, \dots, i_d)$$

$$\mathbf{d}_p^+(i_1, \dots, i_p, \dots, i_d) = (i_1, \dots, i_p + 1, \dots, i_d)$$

Therefore, as an example,

$$\partial(1, 4, 11) = \{(0, 4, 11), (1, 4, 10)\} \cup \{(2, 4, 11), (1, 4, 12)\}$$

Next, we provide the description of the P systems used as a Membrane Computing implementation of the Algorithm 1, all of them consisting on five membranes. The first membrane is used as input and for marking the isolated cells before starting the thinning iterations. The second membrane is used to mark simple pairs. The third membrane selects the cells to be removed, removes the marked cells and updates the facet counter. The fourth membrane marks new isolated cells, updates counters I and R . The fifth one is used as output membrane. Next, our tissue P system is formally described.

In order to properly denote the number of facets of a cell we will use the *star* operator, defined as follows:

$$\text{st } \sigma = \{\mu \in K : \sigma \in \partial\mu\}$$

for any cell σ in a cell complex K . Making a little abuse of notation, we will also represent as st to the operator

$$2^{\mathbb{T}} \circ \text{st} \circ \mathbb{T}^{-1} : T_{2n-1}^d \rightarrow 2^{T_{2n-1}^d}$$

determined in the following commutative diagram:

$$\begin{array}{ccc} \mathcal{K}^d(n) & \xrightarrow{\mathbb{T}} & T_{2n-1}^d \\ \downarrow \text{st} & & \downarrow \text{st} \\ 2^{\mathcal{K}^d(n)} & \xrightarrow{2^{\mathbb{T}}} & 2^{T_{2n-1}^d} \end{array}$$

Recall that, given a finite set A , $\#A$ denotes the number of elements in A .

Next, a family of tissue P systems to solve the d -ThCub problem is defined. This P systems find an skeleton for a cubical complex K while the topology (in terms of homotopy equivalence) and shape is kept. The P systems are designed to work following the next sequence of steps:

- Step 1: Mark initially isolated cells in the input membrane.
- Step 2: Move objects from the first membrane to second one.
- Step 3: Mark simple pairs.
- Step 4: Move objects from the second membrane to the third.
- Step 5: Mark cells to be removed.
- Step 6: Remove marked cells and update some counters.

Step 7: Move objects from the third membrane to the fourth.

Step 8: If no cell has been removed in previous steps, send skeleton cells to the output membrane and halts. In other case, move objects from fourth membrane to fifth.

Step 9: Update counters.

Step 10: Move objects from the fifth membrane to the second and continue from Step 3.

Let I be a d -D binary image of size n^d , let K be the cubical cell complex built from I , let $\varepsilon_{abs} \in \{1, 2, \dots, n\}$ and $\varepsilon_{rel} \in \{\tau_1, \dots, \tau_m\} \subset (0, 1) \cap \mathbb{Q}$, where $\tau_j < \tau_{j+1}$ for $1 \leq j < m$. For every tuple $\langle n, \varepsilon_{abs}, \varepsilon_{rel} \rangle$ we will define a tissue P system with promoters, inhibitors, priorities and input, denoted by $\Pi(n, \varepsilon_{abs}, \varepsilon_{rel})$ and defined as follows:

$$\Pi(n, \varepsilon_{abs}, \varepsilon_{rel}) = (\Gamma, \Sigma, \mathcal{E}, w_1, \dots, w_6, \mathcal{R}, Pri, i_{in}, i_o)$$

where:

- $\Gamma = \{\mathbf{i} : \mathbf{i} \in T_{2n-1}^d\} \cup \{(R, \mathbf{i}, z), (I, \mathbf{i}, z) : \mathbf{i} \in T_{2n-1}^d, 1 \leq z \leq 2n\} \cup \{F_i : \mathbf{i} \in T_{2n-1}^d\} \cup \{I_i, \bar{I}_i, S_i, \bar{S}_i, R_i, \bar{R}_i, U_i : \mathbf{i} \in T_{2n-1}^d\} \cup \{R, H\}$
- $\Sigma = \{\mathbf{i}, (R, \mathbf{i}, 1), (I, \mathbf{i}, 0), F_i^{\#st \mathbf{i}} : \mathbf{i} \in K\}$
- $w_1 = \dots = w_6 = \emptyset$
- $\mathcal{E} = \Gamma \setminus \Sigma$

The rules in the P system will be determined by the use of rule schemes. A rule scheme is a set of rules where some elements inside them are given as parameters in some set. From this point of view, a rule scheme is a function from some set of parameters to the set of rules of the system.

The set of rules \mathcal{R} is provided in the Appendix.

- $Pri = \{R_6 > R_7, R_{13} > R_{15}, R_{14} > R_{15}, R_{36} > R_{34}\}$
- $i_{in} = 1$ is the input cell.
- $i_{out} = 5$ is the output cell.

Theorem 1. *The family of P Systems $\Pi(n, \varepsilon_{abs}, \varepsilon_{rel})$ defined above builds an skeleton for the cubical complex built from a d -D image, $d = 2, 3$, in logarithmic time (measured as number of computation steps with respect to the size of the input image n^d).*

Proof. (Sketch). Let $K \subset T_{2n-1}^k$ be the cubical cell complex encoded as described above for a given d -D image.

The evolution of the P systems follows the steps sequenced below.

Step 1 In this step only rules establishing communication with the first membrane can be selected, hence only rules from R_1 to R_5 can be taken up. However, rules R_4 and R_5 require the presence of isolation marks, either positive (I_i) or negative (\bar{I}_i), and those marks are not present yet into the first membrane. Consequently, only rules R_1 , R_2 and R_3 can be selected. The application of the selected rules in the scheme R_1 , import from the environment an isolation mark I_i for each isolated cell \mathbf{i} . Also, the application of rules in R_2 take one non isolation mark \bar{I}_i for every non isolated cell \mathbf{i} . The application of the rules in the scheme R_3 moves every facet counter F_i to the second membrane.

Step 2 In this situation, only rules in the five first schemes can be selected. However, the presence of isolation marks blocks the selection of rules R_1 and R_2 . On the other hand, facet counters have been moved to the second membrane in the prior step. Henceforth, only rules R_4 and R_5 can be selected.

After the application of the selected rules, all the available objects are in the second membrane.

Step 3 In this computation stage, with all the objects inside the second membrane, only rules in the schemes R_6 to R_{12} can be selected. Rules in R_9 through R_{12} cannot be selected as there is no simple pair marks, either positive or negative. Rules R_6 , R_7 and R_8 determine a maximal set of rules bearing in mind that rules in R_6 has greater priority than rules in R_7 and both of them take all the available cells objects \mathbf{i} .

After applying the selected rules, every cell in the second membrane has a simple pair mark, either positive or negative. In addition, the application of rules in R_8 moves the facet counters to the third membrane.

Step 4 Only rules in the schemes R_6 to R_{12} can be selected, as any other one has not enough elements in the corresponding membrane to be chosen. On the other hand, in the present stage the presence of simple pair marks prevent the selection of rules R_6 or R_7 and the absence of facet counters F_i avoid the selection of rules in R_8 . Hence, only rules from R_9 to R_{12} can be selected. The application of selected rules leads the system to the following computation, with all the objects in the third membrane.

Step 5 In this step, only rules in R_{13} through R_{15} can be selected. All of them determine a maximal set of rules that marks every cell \mathbf{i} as removable or not removable, bearing in mind the priority relation $R_{13} > R_{14} > R_{15}$. After the application of selected rules, all the cells are marked with remove mark, either positive or negative. Although, for each cell marked as removable, one object R and one object H have been imported from the environment.

Step 6 In this computation stage, all the cells have been marked as removable or not removable. Here we can find two different possibilities, depending on the existence of objects R_i , R and H , marking the remove of some cells.

1) If no cell has been marked as removable, there are no objects R_i , R or H in the third membrane. In this situation, rules R_{16} through R_{27} cannot be selected as all of them require the presence either of objects R_i , R or H . On the other hand, rules in R_{29} and R_{30} cannot be selected as they require the absence of objects \mathbf{i} . The rules from R_{31} above cannot be neither selected, as the take objects from the fourth membrane and, in the present situation, there are no objects inside it.

The unique rules that can be selected are those in the scheme R_{28} . After its application, all the cells are inside the output membrane, the fifth one. This situation prevent the selection of any rule, resulting in a halting computation.

2) Let us suppose now that there are some cells marked to be removed inside the third membrane. Henceforth, there are objects R_i , R and H for each cell marked as removable. The unique rules that can be selected are those in R_{16} through R_{21} . The rules in R_{22} up to R_{30} cannot be selected in this stage as they require the absence of objects R . Rules from R_{31} above cannot be selected as they require the presence of objects inside the fourth membrane and, in the current stage, this membrane is empty.

The application of the selected rules updates the facet counter (R_{16}), removes cells, simple pair marks and facet counters for each removable cell (R_{17} to R_{21}).

Step 7 To reach this configuration it is essential to have removed some cells in prior steps. After removing all the cells marked to be removed, all the objects R have been removed too. This allows the rules R_{22} up to R_{27} to be selected. The other rules cannot be selected as they require the presence of objects that there are not present inside the corresponding membrane.

The application of the selected rules leads the system to the next configuration, where all the objects are inside the fourth membrane.

Step 8 In this computation, only the rules in R_{29} up to R_{35} can be selected. Rules R_{34} and R_{35} have been designed to make a maximal set of rules, as every object i can be used by only one of them.

The application of the selected rules erases auxiliary objects (R_{29} to R_{33}), marks new isolated cells and update counter I (R_{34}).

Step 9 In this stage, the unique rules that can be selected are those in R_{36} due to priority reasons ($R_{36} > R_{35}$). Rules in R_{37} up to R_{39} require the presence of objects U_i and those objects are not present yet inside the fourth membrane.

The application of the selected rules updates the counter R .

Step 10 Now, only the rules R_{37} up to R_{39} can be selected. After the application of the selected rules, all the available objects are inside the second membrane, allowing the system to evolve repeating the steps shown below.

□

Table I shows the required computational resources for the family of tissue P systems defined in this paper.

A. Example

In order to proper understanding of the parallel bioinspired algorithm presented in this paper, we will show here a simple example. We will start with a binary 2-D image shown in Figure V-A(a), where each pixel is represented as a square. In Figure V-A(b) the cubical complex associated to the input image is shown over the source image and, finally, the associated cubical complex is presented alone in Figure V-A(c). In all of the previous figures a coordinate system is presented. The last one presents the encoding used in this paper to represent a

d -D binary image thinning problem (d – ThCub)	
Complexity	
Number of steps of computation	$\leq 8d(n+1) + 6$
Resources needed	
Size of the alphabet	$O(n^{d+1})$
Initial number of cells	5
Initial number of objects	$O(K)$
Number of rules	$O(n^{d+2})$
Upper bound for the length of the rules	8

TABLE I

COMPLEXITY ASPECTS, WHERE THE SIZE OF THE INPUT DATA IS $O(n^d)$, $|K|$ IS THE NUMBER OF CELLS IN THE INPUT CELL COMPLEX K .

cubic cell by a tuple of integers. For example, the cell $(3, 5)$ in Figure V-A(c) is the unique square in the complex. In the following paragraphs we show the thinning process using thresholds $\varepsilon_{abs} = 2$ and $\varepsilon_{rel} = 0.5$.

Figure V-A(d) presents one of the simple pairs that can be selected for removal, e.g. $\langle (2, 8), (2, 7) \rangle$ and $\langle (4, 5), (3, 5) \rangle$. Recall that, when a cell belongs to many simple pairs, one of them is non-deterministically chosen. The counters I and R for the facets of those simple pairs are $I(2, 7) = I(3, 5) = 0$ as both cells are initially isolated and $R(2, 7) = R(3, 5) = 1$ as they are selected to be removed in the first iteration. Therefore, the medial persistence measures are given by

$$\begin{aligned} MP_{abs}(2, 7) &= 1, MP_{rel}(2, 7) = 1, \\ MP_{abs}(3, 5) &= 1, MP_{rel}(3, 5) = 1 \end{aligned}$$

Hence, both simple pairs can be removed resulting in the complex given in Figure V-A(e).

In the next iteration, the simple pairs are shown (in red) in Figure V-A(f). The values for isolation and remove counter for the facet of the unique simple pair are, respectively, $I(3, 6) = 1$ as it became isolated in the first iteration and $R(3, 6) = 2$ as it was selected for removal in the second iteration. Medial persistence values are given by

$$MP_{abs}(3, 6) = 1, MP_{rel}(3, 6) = 0.5$$

Hence, the unique simple pair can be removed, obtaining the complex in Figure V-A(g).

Once again, there is a unique simple pair (shown in Figure V-A(h)) where isolation and removal counters values are, respectively, $I(2, 5) = 1, R(2, 5) = 3$. Therefore, the medial persistence measures are given by

$$MP_{abs}(2, 5) = 2, MP_{rel}(2, 5) \approx 0.67$$

so it cannot be removed, as this simple pair retains shape information.

Finally, the algorithm halts returning the complex in Figure V-A(i).

VI. CONCLUSIONS AND FUTURE WORK

In this paper, three different research areas are brought together in order to propose a new point of view on a classical problem: Algebraic Topology, Membrane Computing and Digital Image Analysis. Firstly, Algebraic Topology allows us to split the space in discrete units and to define mechanical

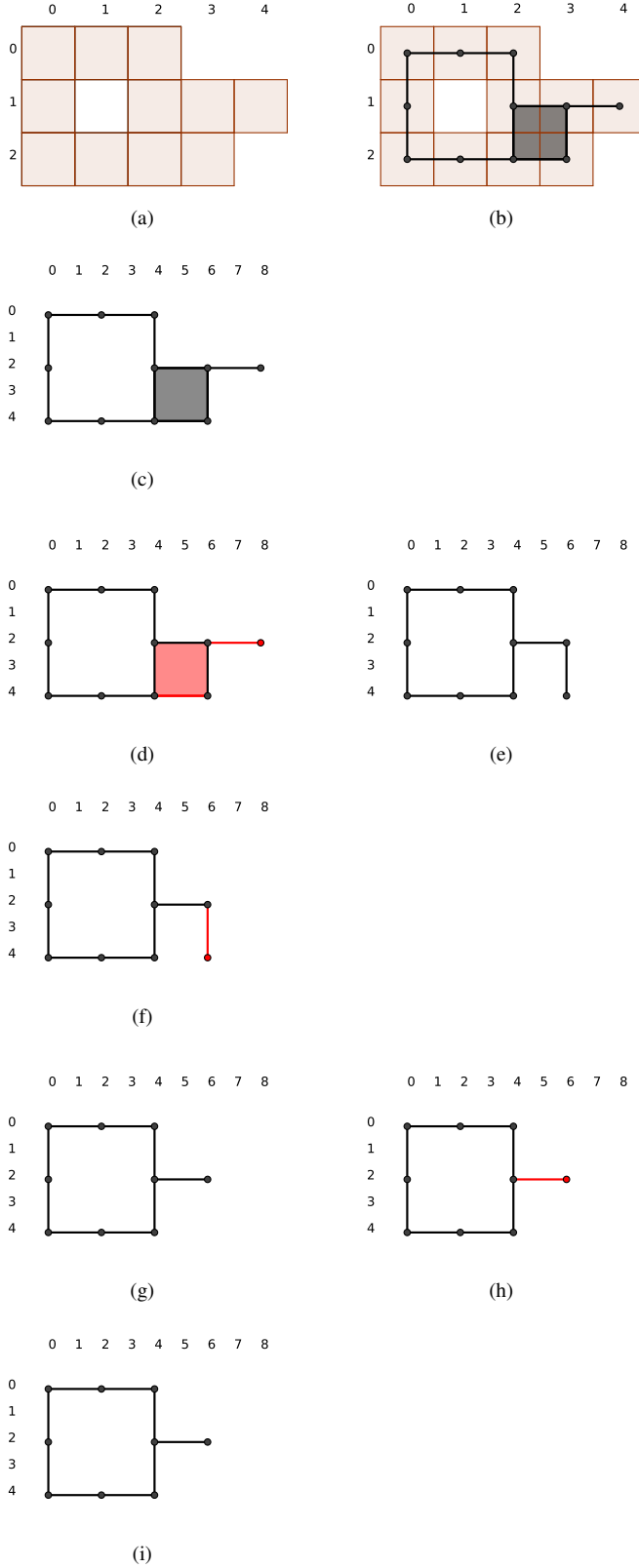


Fig. 2. Construction of a cubical complex associated to a binary image (a). In (b) the complex is shown over the image. In (c) the complex is shown alone. From (e) to (i) the thinning process is sketched.

operators on such space units of different dimensions. Secondly, a recent bio-inspired computational model, Membrane Computing, is used to handle with such operators and provide an alternative point of view for handling with them. Both disciplines deal with compartments of the Euclidean space on their foundations, but their inspiration and motivation are quite different. The former is born as a tool for handle concepts of Algebraic Topology and the latter is a computation model inspired in the functioning of living cells and tissues.

The case of study for applying the Membrane Computing implementation of the Algebraic Topology concepts has been the skeletonization problem, a classical case study in Digital Image Analysis. This implementation provides a new proof that the Membrane Computing framework is flexible enough to adapt to unexpected situations. This paper follows the research line open with [12], but, to the best of our knowledge, this is the first work which put together Membrane Computing, cells complexes and thinning processes. In this way, this is a pioneer work that open a new research line that can be followed at different levels.

Further research can be made by exploring other P system models with new computational features, the application of Membrane Computing techniques to further Digital Image Analysis or a deeper theoretical study of the compartmental view of the Euclidean space by Membrane Computing and Algebraic Topology as the starting point for a deeper study of their common properties.

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APPENDIX

Next, the set of rules \mathcal{R} is provided:

- $R_1 \equiv (\neg\{F_i, I_i\} | 1, i/i I_i, 0)$
for $i \in T_{2n-1}^d$.
- $R_2 \equiv (\{F_i^l\} \neg\{F_i^{l+1}, \bar{I}_i\} | 1, i/i \bar{I}_i, 0)$
for $i \in T_{2n-1}^d, 1 \leq l \leq 2d$
These rules mark every isolated cell i with *isolation mark* I_i and every non isolated cell i with *non isolation mark* \bar{I}_i .
- $R_3 \equiv (1, F_i/\lambda, 2)$
for $i \in T_{2n-1}^d$.
- $R_4 \equiv (1, i(R, i, 1) (I, i, 0) I_i/\lambda, 2)$
for $i \in T_{2n-1}^d$.

- $R_5 \equiv (1, i(R, i, 1) (I, i, 0) \bar{I}_i/\lambda, 2)$
for $i \in T_{2n-1}^d$
These rules move cells, *remove counters*, *isolation counters*, *facet counters* and isolation marks to the second membrane, in order to detect simple pairs.
- $R_6 \equiv (\neg\{F_j^2, S_i, S_j\} | 2, i j F_j / i j F_j S_i S_j, 0)$
for $i \in T_{2n-1}^d, j \in \partial i$
These rules detect simple pairs $\langle j, i \rangle$ and mark its elements using *simple pair mark* S_i and S_j .
- $R_7 \equiv (\neg\{\bar{S}_i\} | 2, i/i \bar{S}_i, 0)$
for $i \in T_{2n-1}^d$.
These rules mark every cell i that not belongs to any simple pair with the *non simple pair mark* \bar{S}_i .
- $R_8 \equiv (\{i\} | 2, F_i/\lambda, 3)$
for $i \in T_{2n-1}^k$.
- $R_9 \equiv (2, i(R, i, z) (I, i, Z) \bar{I}_i S_i/\lambda, 3)$
for $i \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$.
- $R_{10} \equiv (2, i(R, i, z) (I, i, Z) I_i \bar{S}_i/\lambda, 3)$
for $i \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$.
- $R_{11} \equiv (2, i(R, i, z) (I, i, Z) \bar{I}_i \bar{S}_i/\lambda, 3)$
for $i \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$.
- $R_{12} \equiv (2, i(R, i, z) (I, i, Z) I_i S_i/\lambda, 3)$
for $i \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$
These rules send cells, counters and markers to the third membrane, where cells with not enough shape information will be removed.
- $R_{13} \equiv (\{I_i, (I, i, 0), S_i, S_j\} \neg\{R_i, R_j\} | 3, i j / i j R_i R_j R H, 0)$
for $i \in T_{2n-1}^d, j \in \partial i$.
These rules mark every simple pair $\langle j, i \rangle$ as removable where the facet cell i is initially isolated.
- $R_{14} \equiv (\{S_i, S_j, (R, i, z), (I, i, Z)\} \neg\{R_i, R_j\} | 3, i j / i j R_i R_j R H, 0)$
for $i \in T_{2n-1}^d, j \in \partial i, MP_{abs} < \varepsilon_{abs}$ or $MP_{rel} < \varepsilon_{rel}$ and $0 \leq z, Z \leq 2n$, where $MP_{abs} = z - Z$ and $MP_{rel} = 1 - \frac{Z}{z}$.
These rules mark those cells i and j such that $\langle j, i \rangle$ is a simple pair, none of them has been marked to be removed and the cell i has not enough shape meaning, measured in terms of medial persistence, absolute or relative.
- $R_{15} \equiv (\neg\{\bar{R}_i\} | 3, i/i \bar{R}_i, 0)$
for $i \in T_{2n-1}^d$.
These rules mark any cell i as non removable, with *non removable mark* \bar{R}_i , if it has not been previously marked as removable.
- $R_{16} \equiv (\{i, j, R_i, \bar{R}_j\} | 3, F_j/\lambda, 0)$
for $i \in T_{2n-1}^d, j \in \partial i$
These rules update facet counters.
- $R_{17} \equiv (3, i(R, i, z) (I, i, Z) R_i R I_i/\lambda, 0)$
for $i \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$.
- $R_{18} \equiv (3, i(R, i, z) (I, i, Z) R_i R \bar{I}_i/\lambda, 0)$
for $i \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$.
- $R_{19} \equiv (\{R_i\} | 3, S_i/\lambda, 0)$
for $i \in T_{2n-1}^d$.
- $R_{20} \equiv (\{R_i\} | 3, \bar{S}_i/\lambda, 0)$
for $i \in T_{2n-1}^d$.
- $R_{21} \equiv (\{R_i\} | 3, F_i/\lambda, 0)$
for $i \in T_{2n-1}^d$.
These rules remove those cells marked for removal.

- $R_{22} \equiv (\{H, \bar{R}_i\} \neg \{R\} | 3, \mathbf{i}(R, \mathbf{i}, z) (I, \mathbf{i}, Z) I_i / \lambda, 4)$
for $\mathbf{i} \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$
- $R_{23} \equiv (\{H, \bar{R}_i\} \neg \{R\} | 3, \mathbf{i}(R, \mathbf{i}, z) (I, \mathbf{i}, Z) \bar{I}_i / \lambda, 4)$
for $\mathbf{i} \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$
- $R_{24} \equiv (\{H, \bar{R}_i\} \neg \{R\} | 3, F_i / \lambda, 4)$
for $\mathbf{i} \in T_{2n-1}^d$
- $R_{25} \equiv (\{H, \bar{R}_i\} \neg \{R\} | 3, S_i / \lambda, 4)$
for $\mathbf{ij} \in T_{2n-1}^d$
- $R_{26} \equiv (\{H, \bar{R}_i\} \neg \{R\} | 3, \bar{S}_i / \lambda, 4)$
for $\mathbf{i} \in T_{2n-1}^d$
- $R_{27} \equiv (\neg \{R\} | 3, H / \lambda, 4)$
These rules move remaining objects to the fourth membrane.
- $R_{28} \equiv (\{\bar{R}_i\} \neg \{H, R\} | 3, \mathbf{i} / \lambda, 5)$
for $\mathbf{i} \in T_{2n-1}^d$
These rules send output to the fifth membrane.
- $R_{29} \equiv (\neg \{\mathbf{i}, R\} | 3, R_i / \lambda, 0)$
for $\mathbf{i} \in T_{2n-1}^d$
- $R_{30} \equiv (\neg \{\mathbf{i}, R\} | 3, \bar{R}_i / \lambda, 0)$
for $\mathbf{i} \in T_{2n-1}^d$
- $R_{31} \equiv (4, S_i / \lambda, 0)$
for $\mathbf{i} \in T_{2n-1}^d$
- $R_{32} \equiv (4, \bar{S}_i / \lambda, 0)$
for $\mathbf{i} \in T_{2n-1}^d$
- $R_{33} \equiv (4, H / \lambda, 0)$
These rules remove auxiliary objects for each removed cell.
- $R_{34} \equiv (\{(R, \mathbf{i}, z)\} \neg \{F_i\} | 4, \mathbf{i} \bar{I}_i (I, \mathbf{i}, Z) / \mathbf{i} I_i (I, \mathbf{i}, z), 0)$
for $\mathbf{i} \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$
- $R_{35} \equiv (\{F_i\} \neg \{U_i\} | 4, \mathbf{i} / \mathbf{i}, 0)$
for $\mathbf{i} \in T_{2n-1}^d$
These rules update isolation counter and isolation marks.
- $R_{36} \equiv (\neg \{U_i\} | 4, \mathbf{i}(R, \mathbf{i}, z) / \mathbf{i}(R, \mathbf{i}, z + 1) U_i, 0)$
for $\mathbf{i} \in T_{2n-1}^d, 1 \leq z \leq 2n - 1$
These rules update removal counter.
- $R_{37} \equiv (\{U_i\} | 4, \mathbf{i}(R, \mathbf{i}, z) (I, \mathbf{i}, Z) I_i / \lambda, 2)$
for $\mathbf{i} \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$
- $R_{38} \equiv (\{U_i\} | 4, \mathbf{i}(R, \mathbf{i}, z) (I, \mathbf{i}, Z) \bar{I}_i / \lambda, 2)$
for $\mathbf{i} \in T_{2n-1}^d, 0 \leq z, Z \leq 2n$
- $R_{39} \equiv (\{U_i\} | 4, F_i / \lambda, 2)$
for $\mathbf{i} \in T_{2n-1}^d$
These rules move objects to the second membrane.