DEFORMING TESSELLATIONS FOR THE SEGMENTATION OF CELL AGGREGATES

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ABSTRACT

We present a new active contour to segment cell aggregates. We describe it by a smooth tessellation that is attracted toward the cell membranes. Our approach relies on subdivision schemes that are tightly linked to the theory of wavelets. The shape is encoded by control points grouped in tiles. The smooth and continuously defined boundary of each tile is generated by recursively applying a refinement process to its control points. We deform the smooth tessellation in a global manner using a ridge-based energy that we have designed for that purpose. By construction, cells are segmented without overlap and the tessellation structure is maintained even on dim membranes. Leakage, which afflicts usual image-processing methods (*e.g.*, watershed), is thus prevented. We validate our framework on both synthetic and real microscopy images, showing that the proposed method is robust to membrane gaps and to high levels of noise.

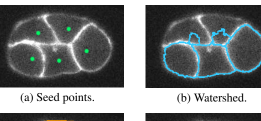
Index Terms— Active contour, *C. elegans*, cell aggregate, Deslauriers-Dubuc, segmentation, subdivision, tessellation.

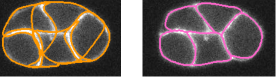
1. INTRODUCTION

We address the segmentation of the cell aggregates that appear in images of several biological specimen such as *C. elegans* embryo or cornea endothelium (Figures 1 and 8). This task is challenging because of the proximity of the cells and the presence of gaps in the membranes.

Over the past decade, automated methods were proposed for the segmentation of cell membranes, including intensity thresholding, morphological operations, Voronoi-based methods [1], labeling procedures [2], or watershed transform [3-7]. Those methods have three main limitations. First, they suffer from leakage in case of dimmed membranes. Second, they are sensitive to noise and might result in over-segmentation. Third, it is not easy to introduce prior knowledge to improve the accuracy of the segmentation [8]. Recent approaches are based on the detection of membrane patterns coupled with graph-cut [9], or on deep learning [10]. If these methods are better suited for incorporating prior knowledge, they do not allow for easy and user-friendly interaction. Moreover, they may provide non-continuous cell boundaries, which complicates the extraction of quantitative measurements. Topology adaptive methods, such as level sets or T-snakes [11, 12], are not required in this context as the topological structure of the cell aggregate is generally known.

Parametric active contours—a.k.a. snakes—are popular models for the interactive segmentation of bioimages [13–17]. They consist in a curve that evolves from an initial position toward the boundary of the object of interest through the minimization of an energy term [16, 18]. The underlying shape has a continuous representation in terms of basis functions [16], which facilitates the incorporation





(c) Five classical snakes. (d) Active tessellation, J = 0.98.

Fig. 1. Segmentation of a *C. elegans* embryo; Source: R. Jankele and P. Gönczy, EPFL. (a) Seed points used for the initialization of each method; (b) watershed method; (c) five classical snakes [16]; (d) active tessellation.

of prior knowledge. They are built to ensure continuity and smoothness, which prevents leakage. They are parametrized by only a few control points, which results in fast optimization and provides robustness to noise. However, parametric snakes are not well suited to segment objects that are close to each other: They might yield overlapping segmentations (Figure 1 (c)) as the snakes are optimized independently. In the literature, only few works regarding active contours address the segmentation of touching objects. Extension of traditional parametric snakes to track non-occluding objects that transiently touch each other are presented in [19, 20]. These methods could only segment cell aggregates with thick membranes or non-touching cells. Networks of active contours were introduced in [21, 22]. However, they involve many parameters (nodal points).

In this paper, we propose a new active contour with a geometric representation that keeps the advantages of parametric snakes while addressing globally the segmentation problem of cell aggregates. The model consists in a smooth tessellation, so-called *active tessellation*, that is globally deformed towards the cell membranes through the minimization of a suitable ridge-based energy. The smooth tessellation is encoded by a set of control points and generated through a geometric subdivision scheme, a computer-graphics tool for representation and modeling. By construction, the segmented tiles are non-overlapping and the tessellation structure bridges membrane gaps. After optimization, each cell of the aggregate can be individually extracted to compute statistical descriptors.

The main contributions of this work are: 1) the construction of a smooth tessellation to describe an active contour; 2) the derivation of an oriented ridge-based energy functional; 3) the implementation of the whole framework. We illustrate the benefits of the proposed active tessellation on real biological applications in the context of semi-interactive segmentation.

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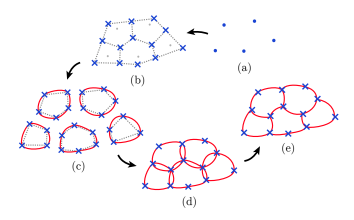


Fig. 2. Construction of the smooth tessellation. Given seed points (a), we first generate control points (blue crosses) grouped in tiles (b). Then, we specify the smooth boundary of each tile with a continuous closed curve by applying a subdivision scheme to its control points (c). Finally, we keep one of the common edges (d) to obtain the final smooth tessellation (e).

2. CONSTRUCTION OF A SMOOTH TESSELLATION

Our active contour model is a smooth tessellation, as described in Figure 2 (e). Its shape is parametrized by control points that are grouped in tiles (Figure 2 (b)). Each tile is associated with a closed curve that specifies its smooth boundary (Figure 2 (c)). This smooth curve is generated from the control points via a subdivision scheme (Section 2.1). The smooth tessellation is the union of the closed curve of each tile. As two control points that belong to two adjacent tiles are connected by two continuous edges (Figure 2 (d)), we keep only the edge of the largest tile (Figure 2 (e)). The remaining edges are then optimized by fitting them to the image data via energy optimization (Section 3).

2.1. A Tiling Made of Closed Subdivision Curves

A subdivision scheme allows one to describe a contour by an initial set of control points which, by the iterative application of a refinement rule, becomes continuous in the limit [23–26]. The smoothness and geometric-reproduction properties of the limit curve depend on the particular choice of the subdivision mask. We use this powerful mathematical method to generate the closed curve of each tile.

2.1.1. Notations

We represent by $\mathbf{p}_{(k)}[\cdot]$ a $(2^k M)$ -periodic sequence of points $\mathbf{p}_{(k)}[m] = (p_{1(k)}[m], p_{2(k)}[m])$, indexed by $m \in \mathbb{Z}$, where $p_{1(k)}$ and $p_{2(k)}$ are the coordinates. The discrete convolution of $\mathbf{p}_{(k)}[\cdot]$ with a scalar mask $h[\cdot]$ is defined as $(h * \mathbf{p}_{(k)})[m] = \sum_{n \in \mathbb{Z}} h[m-n]\mathbf{p}_{(k)}[n]$.

2.1.2. Closed Subdivision Curves

Let $\mathbf{p}_{(0)}$ be the *M*-periodic sequence that contains the control points of a tile. The closed curve associated to this tile is generated as follows (Figure 3): We apply a refinement rule to $\mathbf{p}_{(0)}$ to double the number of points. We repeat this operation k times to obtain the $(2^k M)$ -periodic sequence of subdivision points $\mathbf{p}_{(k)}$. When $k \to \infty$, the set of points converges to the continuous curve $\mathbf{r} = (r_1, r_2)$ with $r_1, r_2 \in \mathcal{C}^1$. In practice, k = 6 iterations are enough for the contour points to be sufficiently dense. The subdivision points at the *k*th iteration ($k \ge 1$) are directly obtained from $\mathbf{p}_{(0)}$ by

$$\mathbf{p}_{(k)} = h_{0 \to k} * \mathbf{p}_{(0)}_{\uparrow_{-k}},\tag{1}$$

where \uparrow_{2^k} denotes an upsampling by a factor of 2^k , $h_{0\to k} = h_{\uparrow_{2^{k-1}}} * h_{\uparrow_{2^{k-2}}} * \cdots * h_{\uparrow_2} * h$ [17], and h is the subdivision mask whose z-transform is

$$H(z) = -\frac{1}{16}z^{-3} + \frac{9}{16}z^{-1} + 1 + \frac{9}{16}z - \frac{1}{16}z^3.$$
 (2)

This subdivision mask corresponds to the convergent subdivision scheme introduced by Deslauriers and Dubuc in [27], which generates C^1 -continuous functions and reproduces polynomials up to degree 3 [27]. This scheme is interpolating [27, 28], which means that the subdivision points, at each level of the process, interpolate the limit curve **r**. We thus have that

$$\mathbf{r}(t)|_{t=\frac{m}{2^k}} = \mathbf{p}_{(k)}[m]. \tag{3}$$

This is an advantage when user interaction is involved, as it facilitates the editing of the curve. Moreover, it is affine invariant, so that the geometry of the limit curve changes in synchrony with any affine transformation that would be applied to the initial set of control points. This is an important property that facilitates the design of active contours.

3. AN ORIENTED RIDGE-BASED ENERGY

The energy functional E drives the deformation of the smooth tessellation. Usually, this energy is divided into an internal energy, which ensures the smoothness of the curve, and an external energy, which is purely data driven. In our formulation, the smoothness of the curve is ensured by the Deslauriers-Dubuc subdivision scheme that produces C^1 -continuous curve. Since we want the external term to attract the smooth tessellation toward the cell membranes, we reduce E to a ridge-based energy term.

To detect ridges, a common approach is to compute the Hessian matrix U at location $\mathbf{x} = (x_1, x_2)$ as

$$\mathbf{U} = \begin{pmatrix} \frac{\partial^2 u(x_1, x_2)}{\partial x_1^2} & \frac{\partial^2 u(x_1, x_2)}{\partial x_1 \partial x_2} \\ \frac{\partial^2 u(x_1, x_2)}{\partial x_2 \partial x_1} & \frac{\partial^2 u(x_1, x_2)}{\partial x_2^2} \end{pmatrix}, \tag{4}$$

where $u(x_1, x_2) = f(x_1, x_2) * (g(x_1)g(x_2))$ with f the input image and $g(x_i) = \frac{1}{\sigma\sqrt{2\pi}} e^{\frac{-(x_i)^2}{2\sigma^2}}$ for i = 1, 2. High values of the standard deviation $\sigma \in \mathbb{R}$ increase the basin of attraction of the active contour. We define the ridge strength at location x by

$$\xi(\mathbf{x}) = \sqrt{|\lambda_{\min}|} \cdot \sqrt{|\lambda_{\min} - \lambda_{\max}|}, \qquad (5)$$

where λ_{\min} and λ_{\max} are the minimum and maximum eigenvalues of U, respectively. On the ridge, the ridge strength is maximum and the eigenvector \mathbf{v}_{\min} is normal to the ridge.

Let c be the control points of the active tessellation. They generates a set of N points $\mathbf{p}_{tess} = \mathbf{p}_{tess}(\mathbf{c})$ that delineates the smooth tessellation (Figure 3). The oriented ridge-based energy that we propose is

$$E(f, \mathbf{p}_{\text{tess}}) = -\frac{1}{N} \sum_{m=0}^{N-1} \xi(\mathbf{p}_{\text{tess}}[m]) \\ \times \frac{|\mathbf{v}_{\min}^{\mathsf{T}}(\mathbf{p}_{\text{tess}}[m]) \cdot \mathbf{n}(\mathbf{p}_{\text{tess}}[m])|}{\|\mathbf{v}_{\min}(\mathbf{p}_{\text{tess}}[m])\| \|\mathbf{n}(\mathbf{p}_{\text{tess}}[m])\|},$$
(6)

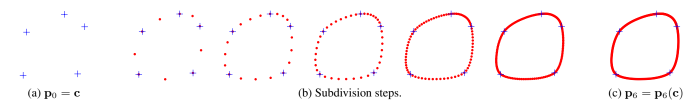


Fig. 3. Interpolating Deslauriers-Dubuc subdivision scheme. (a) Control points \mathbf{p}_0 ; (b) subdivision steps that converge to the continuous curve (c) which is encoded by the five control points \mathbf{p}_0 (blue crosses).

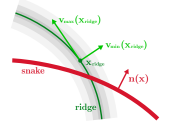


Fig. 4. Ridge feature. When minimizing the ridge-based energy (6), the active tessellation (red line) tends to align with the center of the ridge (green line).

where $\mathbf{n}(\mathbf{p}_{tess}[m])$ is the normal vector at the *m*th point. The energy (6) is minimal when the vectors \mathbf{v}_{min} and \mathbf{n} are aligned and when the tessellation lies on the ridge (Figure 4). The optimization is computed as a function of the control points \mathbf{c} . The optimum is obtained as

$$\mathbf{c}_{\text{opt}} = \arg\min E(f, \mathbf{p}_{\text{tess}}(\mathbf{c})). \tag{7}$$

4. IMPLEMENTATION

We implemented the proposed framework as a user-friendly plugin for the bioimaging platform Icy [29].¹

We initialize the active tessellation as follows: We construct a Voronoi diagram around seed points and then apply a mask to shrink the Voronoi tiles in the neighborhood of the cell aggregate. This mask is obtained by constructing the convex hull of the seed points. The vertices of the resulting tiles are the control points of the active tessellation. Note that the seed points are manually specified through the interface or automatically detected from a provided image of the cells' nuclei.

We follow a coarse-to-fine optimization strategy. We first optimize the active tessellation made of the few control points of the Voronoi tiling using a Powell-like line-search method. We obtain a rough segmentation that is less likely to be stuck in local minima. We then double the number of control points to increase the flexibility of the active tessellation and we optimize it once again. This strategy makes our active contour less sensitive to initialization.

5. EXPERIMENTS AND VALIDATION

We perform three experiments to evaluate the performance of our active tessellation. We first investigate its robustness with respect to noise and dim membranes on synthetic data. Then, we illustrate applications on real data.

5.1. Synthetic Data

We compare our approach in term of accuracy against the watershed method [30] implemented by I. Arganda-Carreras and D. Legland for the bioimage platform Fiji². We use the Jaccard index (J) to measure the similarity between the segmentation result Ω and the corresponding ground truth $\Omega_{\rm GT}$. It is defined as $0 \le J = \frac{|\Omega \cap \Omega_{\rm GT}|}{|\Omega \cup \Omega_{\rm GT}|} \le 1$. In the experiments, we compute the Jaccard index for each cell and take the average. It is this average value that we refer to as Jaccard index in the next Sections.

We created a test image that simulates the fluorescence microscopy of a *C. elegans* embryo with 5 cells (Figure 5 (a)). We use the same seed points to initialize the two methods. The initial configuration of the active tessellation is illustrated in Figure 5 (a). Its initial similarity with the ground truth corresponds to J = 0.64.

5.1.1. Robustness with Respect to Noise

We corrupted the test image by different levels of additive Gaussian noise (20 realizations per level of noise, Figure 5 (d)). The signal-tonoise ratio (SNR) corresponding to a given noise level and Jaccard index were computed. The SNR that we use is the ratio of the mean value of the signal and the standard deviation of the noise. The results are given in Figure 6 and illustrated in Figures 5 (e) and (f). We observe that the active tessellation is robust with respect to noise since it is able to give a proper segmentation outcome even for low SNRs. On the contrary, the accuracy of the watershed method degrades significantly for a SNR below 2.4.

5.1.2. Robustness with Respect to Dim Membranes

We progressively dimmed the fluorescence signal on the membranes of the test image (Figure 5 (g)). We computed the Jaccard index as a function of the information-loss percentage. This dimming percentage corresponds to the ratio of the mean intensity on the membrane of the test image over the one of the corrupted image. The resulting plot is given in Figure 7 and we illustrate results in Figures 5 (h) and (i). The active tessellation accurately segments the cells until 49% of information loss while the watershed method can tolerate no more than 15% of information loss, then it quickly decreases. As this model is only based on intensity, it leaks through dim membranes. Due to the structure and smoothness of the active tessellation, the proposed framework does not suffer from leakage.

5.2. Segmentation on Real Data

We applied our active tessellation on real biomedical images. These images are challenging because of the presence of noise and gaps in the membranes. For each segmentation, the initial configuration of

¹The source code and a demo of the plugin are available at http://bigwww.epfl.ch/demo/active-tessellation/, as of October 2018.

²The source code is available at https://imagej.net/Classic_Watershed.

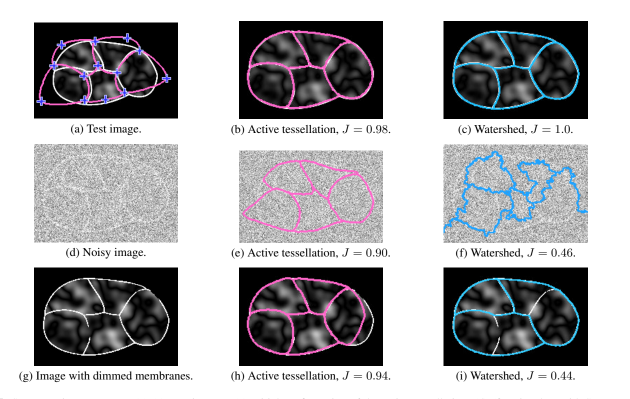


Fig. 5. Segmentation outcomes. (a)-(c) Test images. (a) Initial configuration of the active tessellation; (d)-(f) noisy data with SNR= 0.81; (g)-(i) image with 23.95% of membrane information loss.

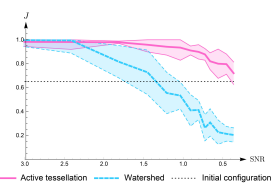


Fig. 6. Segmentation of noisy data. Evolution of the Jaccard index as a function of the SNR. Filled area: standard deviation across the 20 realizations.

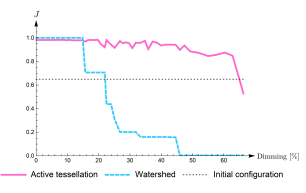


Fig. 7. Evolution of the Jaccard index as a function of the dimming percentage.

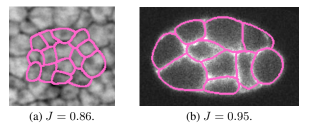


Fig. 8. Cell segmentation of (a) cornea endothelium; (b) *C. elegans* embryo in a light-sheet fluorescence-microscopy image. Source: R. Jankele and P. Gönczy, EPFL.

the active tessellation has 2.6 control points per cell in average. We compute the Jaccard index of each outcome considering a manual segmentation as ground truth. The results obtained are satisfactory in most cases (Figures 1 and 8).

6. CONCLUSION

We have presented a new subdivision-based active contour for the segmentation of cell aggregates. We have modeled the active contour by a smooth tessellation and designed an oriented-ridge-based energy term to efficiently attract the curve toward the membranes. The tessellation structure prevents from overlapping segmentation of the cells and from leakage issues. Moreover, each cell of the segmentation outcome can be easily extracted as a continuous closed curve making possible the computation of cell metrics. We have demonstrated the robustness of our method under noisy conditions and to dim membranes. We have also illustrated its behavior on real bioimages.

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