A Robust Algorithm for Segmenting and Tracking Clustered Cells in Time-Lapse Fluorescent Microscopy

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Abstract—We present herein a robust algorithm for cell tracking in a sequence of time-lapse 2-D fluorescent microscopy images. Tracking is performed automatically via a multiphase active contours algorithm adapted to the segmentation of clustered nuclei with obscure boundaries. An ellipse fitting method is applied to avoid problems typically associated with clustered, overlapping, or dying cells, and to obtain more accurate segmentation and tracking results. We provide quantitative validation of results obtained with this new algorithm by comparing them to the results obtained from the established CellProfiler, MTrack2 (plugin for Fiji), and LSetCellTracker software.

Index Terms—Cell segmentation and tracking, ellipse fitting, fluorescent microscopy, multiphase active contours.

I. INTRODUCTION

HE ability of cells to change shape, divide, migrate, and undergo apoptosis in response to a specific stimuli is of paramount importance in cell biology. The study of cell and

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nuclear dynamics is crucial both to the understanding of tissue morphogenesis in development and to deciphering the complex events leading to the onset of neoplastic transformations.

The monitoring of living cells with high spatial and temporal resolution is made possible by modern techniques in optical fluorescent microscopy which allow the parallel tracking of multiple cellular parameters over long periods of time (hours to days). Data obtained through high-throughput microscopy approaches contain a significant amount of information.

The position of an individual cell over time can be individuated by addressing a fluorescent marker to its nucleus. Nuclear markers include fluorescent dyes [1] and proteins [2], yielding a bright signal limited to the nuclear compartment.

In this paper, we address two main problems that appear in the automatic analysis of living cells in time-lapse 2-D fluorescent microscopy: 1) segmentation of highly clustered nuclei with surrounding cell compartments (e.g., cytoplasm); and 2) tracking over time to provide information on the cell cycle stage and its migration strategy [3] and the effects of the surrounding extracellular environment on cell activities. Main challenges are related to problems arising from overlapped and clustered nuclei which are difficult to split in 2-D imaging and by the nonuniform and noisy signal emitted by fluorescent markers. Moreover, the boundary between migrating or dividing cells in fluorescence microscopy is very often not visible and the only a priori knowledge useable to separate cells is the assumption that the shapes of the nuclei are similar to ellipses. In our approach, we strongly exploit this assumption to analyze image to obtain optimal segmentation of the nucleus or cytoplasm. It is opposite to other approaches in this field recently presented in [4] and [5]. In these papers, authors explore the color, gradient, or texture information for nucleus and cytoplasm/membrane segmentation in immunohistochemical tissue image. The approach presented in [4] aims at using color values for remerging of objects that were oversegmented by watershed. The color values are obtained via a color deconvolution technique. The method introduced in [5] models the cell borders based on image gradient information contained in regions associated with cell membranes. To overcome the common problem of cell overlapping and touching with nonuniform signals in fluorescent microscopy, we incorporate into our segmentation approach an ellipse fitting method based on [6], [7], and [8]. It enables us to preprocess every object tracked with active contour to obtain optimal segmentation at a single time point with much smaller sensitivity to changes of gradient or texture.

Generally, the existing approaches to solve the cell-tracking problem can be divided into two classes: segmentation-based tracking methods and model evolution approaches. The first and most common class is based on minimizing the distances between previously segmented nuclei in two successive frames according to earlier predefined criteria (see, e.g., [9] or [10]). In the second class, active contour models are widely used (see, e.g., [11]–[15] and latest [16], [17]). In these approaches, the contours of nuclei in the previous image frame are the initial positions for evolving the contours of nuclei in the current one. For general information on active contours models, we refer the reader to [18] and [19].

Our approach to cell tracking is based on the multiphase active contour model, which in the context of image processing applications was first presented in [20]. The main advantage of this approach is the possibility of combining cell segmentation with tracking. Our extension includes the new method of segmentation assuming ellipse-like shapes of nuclei, that is combined with the evolution model represented by active contours. The proposed method is able to segment highly clustered and dividing objects (see, e.g., Fig. 2) frame-by-frame.

An interesting approach for the same goal was proposed in [16], where a Radon-based method was used to separate touching objects. We have made a performance comparison of that method implemented in *LSetCellTracker* software [17] with our method. In addition, we provide comparisons with two open source programs widely used in biology: *CellProfiler* [21] and *MTrack2* software [22], implemented as plugin for *Fiji*.

We designate our main algorithm as *robust* for the following reasons. It works well with blurred images having poor signal-to-noise ratio (SNR). Moreover, it allows us to segment nuclei with nonhomogeneous structure and changing light intensity. The proposed algorithm also handles possible difficulties in cell tracking. It detects the appearance of a new nucleus that may occur as a result of: 1) cell division; 2) separation of two initially overlapping nuclei; 3) appearance of a new nucleus in a frame. Furthermore, it detects the disappearance of nucleus that may be the result of: 1) leaving a frame and 2) the fading of the fluorescent marker (death). We note that our algorithm does not distinguish automatically between the cell division and separation of two initially overlapping nuclei. Both cases in this paper are within our *parent-child* definition.

We tracked the migration and proliferation of HeLa-Kyoto cells expressing a histone H2BeGFP construct (H2BeGFP [2]), HeLa cells which expressed a histone H2B-mCherry and underwent cell division, and HeLa cells stained with Hoechst 33258 dye. In addition, we used highly clustered human erythroleukemia line (HEL) cells to illustrate the segmentation abilities of the algorithm. To illustrate the variability of noise in the input data, we measured the SNR of all images. All input and resulting images, as well as the measured SNR can be found online [23].

The paper is organized as follows. In Section II, we introduce the active contour algorithm for cell tracking in two successive microscopy images from a given image sequence, and present the modification of this algorithm to overcome problems in the segmentation of nuclei with obscure boundaries. In Section III, we present the ellipses fitting method that we use to improve

the accuracy of nuclei detection and address problems associated with cell overlap and division. In Section IV, we give short summary and propose the full algorithm for cell tracking. In Section V, we compare results obtained with our algorithm to those obtained with *CellProfiler*, *MTrack2*, and *LSetCellTracker*.

II. MULTIPHASE ACTIVE CONTOURS MODEL

Let Ω be a subset of \mathbb{R}^2 . Assume that two successive nuclei images are given. Let $f^{s-1}:\Omega\to\mathbb{R}$ be an image corresponding to time T_{s-1} and $f^s:\Omega\to\mathbb{R}$ be an image corresponding to time T_s , where $T_{s-1}< T_s$ and 0< s< S. We denote by N the number of nuclei in the image f^{s-1} and by C_i^{s-1} the approximate contour of the ith nucleus in the image f^{s-1} , for $i=1,\ldots,N$. Our aim is to find contours C_i^s of nuclei in the image f^s given C_i^{s-1} , $i=1,\ldots,N$. We note that a solution to this problem will, by itself, not allow us to determine cell division. Further steps are needed to yield the full cell-tracking algorithm.

The active contour model that we consider in this section was originally introduced by Chan and Vese [24]. Its multiphase extension suitable for cell-tracking purposes was proposed by the same authors in [20] and taken up by others, e.g., in [11]–[17]. Its extension to analysis of 3-D microscopy image sequences was shown in [16], and an efficient numerical method to solve the cell-tracking problem on 3-D active meshes was introduced in [25]. A similar model was proposed in [26] to detect cytoplasm contours.

In the next part of this section, we briefly introduce the multiphase active contour model and propose some modifications that are used in our cell-tracking algorithm. In the general multiphase active contour model, it is assumed that the contour C_i^s of the ith nucleus in image f^s is the zero level set of the function $\phi_i^s:\Omega\to\mathbb{R}$ which minimizes the functional

$$F_{\varepsilon}(\phi_{i}) = \sum_{i=1}^{N} \left(\mu \int_{\Omega} g(|\nabla f^{s}|) \, \delta_{\varepsilon}(\phi_{i}(x)) \, |\nabla \phi_{i}(x)| \, dx \right.$$

$$\left. + \lambda_{o} \int_{\Omega} H_{\varepsilon}(\phi_{i}(x)) (f^{s}(x) - c_{i}(\phi_{i}))^{2} \, dx \right.$$

$$\left. + \lambda_{b} \int_{\Omega} \prod_{j=1, j \neq i}^{N} (1 - H_{\varepsilon}(\phi_{j}(x))) (f^{s}(x) - c_{b}(\phi_{i}))^{2} \, dx \right.$$

$$\left. + \omega \int_{\Omega} \sum_{j=1, j \neq i}^{N} H_{\varepsilon}(\phi_{i}(x)) H_{\varepsilon}(\phi_{j}(x)) \, dx \right). \tag{1}$$

In the aforementioned definition, $\mu, \lambda_o, \lambda_b$, and ω are positive parameters, and H_ε is a slightly regularized Heaviside function. Variables c_b and c_i are the averages of the image f^s in $\Omega \setminus \bigcup_{i=1}^N in(C_i^s)$ and $in(C_i^s)$, respectively, where $in(C_i^s)$ denotes a region inside of the contour C_i^s . Finally, the function g is a typical edge-detector function (see, e.g., [26]).

The role of the first three terms in the functional F_{ε} is the same as the corresponding ones in the original model of Chan and Vese [24]. That is, the regularization term weighted by μ penalizes the length of a contour C_i^s , and terms weighted by λ_b and λ_o penalize the L^2 norm of the difference between the average intensity of f^s in $\Omega \setminus \bigcup_{i=1}^{N} in(C_i^s)$ and $in(C_i^s)$,

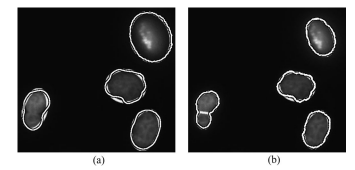


Fig. 1. Nucleus segmentation results for nuclei with obscure boundaries using a) the standard multiphase active contour model and (b) the here proposed modified approach. The emphasized contours are the results of the respective active contour algorithm initialized with ellipses shown as smoothed white lines.

respectively. Finally, the term weighted by the parameter ω penalizes the intersection of regions inside two contours.

Minimizing the functional F_{ε} with respect to ϕ_i results in the Euler–Lagrange equation for ϕ_i^s with the associated Neumann boundary condition and initial condition. Next, parametrization by an artificial time t>0 yields the evolution equation that is considered with the initial condition $\phi_i^s(x,t=0)=\psi_i^s(x;C_i^{s-1})$ for all $x\in\Omega$, where

$$\psi_{i}^{s}(x; C_{i}^{s-1}) \text{ for all } x \in \Omega, \text{ where}$$

$$\psi_{i}^{s}(x; C_{i}^{s-1}) := \begin{cases} d(x, C_{i}^{s-1}), & \text{if } x \in in(C_{i}^{s-1}) \\ -d(x, C_{i}^{s-1}), & \text{if } x \in \Omega \setminus in(C_{i}^{s-1}). \end{cases}$$
(2)

In the aforementioned definition, $d(x, C_i^{s-1})$ denotes the Euclidean distance of the point x to the contour C_i^{s-1} . In order to obtain more stable results, we rescale all values of ψ_i^s by dividing them by the maximum of $|\psi_i^s|$ over all $x \in \Omega$.

We propose to introduce some modification in the definition of the functional F_{ε} to avoid problems with an inaccurate detection of nuclei boundaries in cases where they are not sufficiently sharp. This problem is illustrated in Fig. 1.

The idea that we propose is based on a simple observation. The difference between the arithmetic average and standard deviation of image f^s gray intensity inside the contour C^s_i is greater when it contains a nucleus with a blurred boundary than the case when it contains one with a sharp boundary. Therefore, we modify the second term in the definition of the functional F_ε , scaling the average $c_i(\phi_i)$ by the factor

$$p_i(\phi_i) = \frac{2}{1 + c_i(\phi_i)/s_i(\phi_i)}$$

where $s_i(\phi_i)$ denotes the standard deviation of image f^s gray intensity inside of the contour C_i^{s-1} . The quantity p_i is defined to be greater than 1 in the case of an object with blurred edges. This increases the force that moves the contour toward the perceived boundary. In Fig. 1, we compare the segmentation results obtained with the proposed modified version of the multiphase active contour algorithm to those obtained with its standard version as used in, e.g., [15] and [26]. The emphasized contours of nuclei shown in Fig. 1(a) were determined with the standard multiphase active contour model (1). In Fig. 1(b), the results of the modified method are also shown as emphasized contours, i.e., where the average $c_i(\phi_i)$ in the standard algorithm (1) was scaled with the factor $p_i(\phi_i)$. It is evident that the modified al-

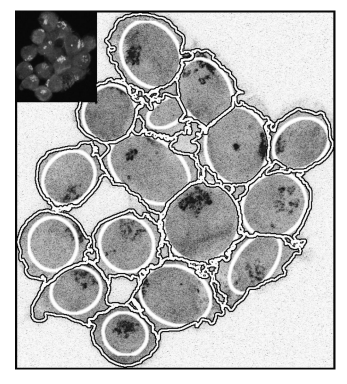


Fig. 2. Result of the WAFR algorithm applied to an image showing clustered HEL cells. The original image is shown in the top left corner, the larger image is inverted to improve visibility. The emphasized contours are obtained with the MMAC algorithm. The ellipses are the output of WAFR.

gorithm yields superior results. We will refer to it henceforth as modified multiphase active contour (MMAC) algorithm.

III. CELL SEGMENTATION BASED ON H-MINIMA TRANSFORM AND ELLIPSE FITTING

In our full cell-tracking algorithm (described in Section IV), there is a need for a tool that can segment objects and, moreover, if it detects a cell division, will allow tracking of the *parent-child* relation.

In this section, we describe such an algorithm. Although it was mainly designed for tracking purposes, it is also of interest as a stand-alone segmentation tool. The algorithm assumes that the objects to be tracked or segmented are of elliptical shape. The underlying idea is based on Jung and Kim [7]. With that algorithm, the tracking of *parent*—*child* relations is not possible, since it does not retain information on the relation of objects. In contrast, our algorithm segments each object separately, allowing us to track relations between objects. This further enables efficient parallelization and implementation on GPUs. Finally, we introduce weights that significantly improve segmentation results.

Watershed is one of the most widely used algorithms for cell segmentation. However, it often yields oversegmentation. This is because regional minima are employed for segmenting objects directly. One of the remedies that suppresses undesired minima is H-minima transform (for a detailed description, see [27] or [7]), which is performed by

$$H_h(\gamma) = R^{\varepsilon}(\gamma + h),$$

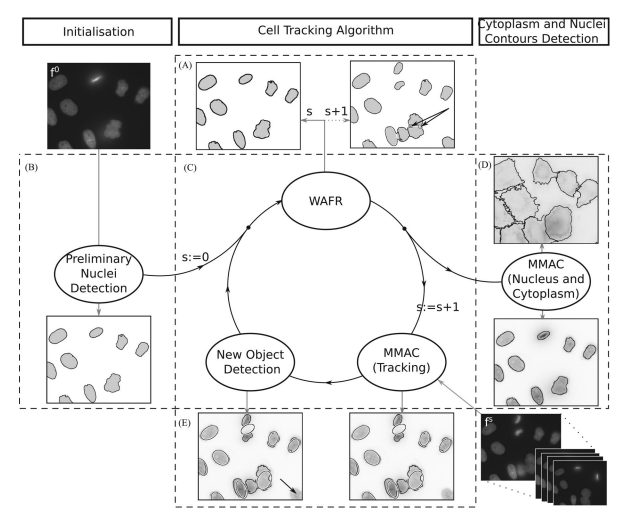


Fig. 3. Schematic diagram of the full tracking algorithm. A detailed description of the parts A–E is given in Section IV. Images are presented as negatives for better visualization. The images in the top-left and bottom-left corners represent the first input image from the sequence and the following images in the time sequence, respectively.

where γ is the inverse distance map of an object, R and ε are the reconstruction and erosion operators, respectively, and h is a given depth. The key is to find the optimal h (typically, it is between 2 and 10). We propose instead of finding a global h for the entire image to determine it for each object separately. We assume from the very beginning that nuclei have elliptical shapes, and we use ellipse fitting for finding the best choice of h for each object. Concretely, we apply H-minima transform with a fixed hand the watershed algorithm. Suppose that the watershed algorithm splits a nucleus into m objects: $w_{h,i}$, $i = 1, \ldots, m$. We fit an ellipse to each partial original boundary using the effective direct least-squares fitting algorithm by Fitzgibbon et al. [6]. Let $n_{h,i}$ denote the number of points of the boundary of $w_{h,i}$. Let $Q_{h,i}$ denote the ellipse fitted to the object, and $b_{h,i,j}$ be the jth point of the boundary. Finally, let $T_{h,i}$ be the transformation that maps the points of the ellipse to the unit circle, and let $d(x, Q_{h,i})$ denote the distance from x to $Q_{h,i}$. Employing weighted average fitting residuals (WAFR)

WAFR
$$(w_{h,i}) = W_i^{(1)} \frac{1}{n_{h,i}} \sum_{i=1}^{n_{h,i}} d(T_{h,i}(b_{h,i,j}), T_{h,i}(Q_{h,i})) + W_i^{(2)}$$

where $W_{i}^{\left(1\right)}$ and $W_{i}^{\left(2\right)}$ are weights to be defined later, yields the distortion

$$S_h = \frac{1}{m} \sum_{i=1}^m \text{WAFR}(w_{h,i}). \tag{3}$$

The optimal h is then the one with the smallest associated distortion.

Weights can improve the result significantly. Both $W_i^{(1)}$ and $W_i^{(2)}$ play roles of penalties: $W^{(1)}$ increases the value of WAFR if the ratio of the area of the fitted ellipse to the area of the object within the ellipse is small, and $W_i^{(2)}$ increases WAFR if there are concativities on the boundary of an object. (For a detailed description of concativity, see Kothari $et\ al.\ [8]$ and compare the regions of Fig. 3, part A, indicated by two arrows.) Specifically,

$$W_i^{(1)} = \left(1 - \frac{\text{Area of } Q_i}{\text{Area of } Q_i \cap w_{h,i}}\right)^2.$$

For $W_i^{(2)}$, we look for concativities on the partial original boundary of $w_{h,i}$, disregarding points near division returned by watershed. If the concativity is larger than a threshold, then we set $W_i^{(2)}$ to a positive constant, otherwise it is set to 0. This way

there is extra penalty to WAFR, if large concativities are found. Although we set both the threshold and the constant manually, we note that for reasonable values the result can only improve: If the threshold is too small, then simply a fixed $W^{(2)}$ is added to each WAFR and thus has no influence on the choice of h. We henceforth refer to the above as the WAFR algorithm. For each object on the binary image, it returns a division with smallest S_h , objects' division, and fitted and filled ellipses.

Note that since we find an optimal h for each object separately, we immediately gain information on object division. This information is exploited in the final cell-tracking algorithm. Moreover, rather than returning the division obtained by watershed, one can simply receive fitted and filled ellipses as new objects. In this case, we also need to find intersections of ellipses and plot them in an image (since they must be separate objects—see white contours of ellipses in Fig. 1). The final cell-tracking algorithm using fitted and filled ellipses outperforms the one using objects returned by the watershed algorithm.

Also, to assess the reliability of the WAFR algorithm, we applied it to images of HEL clustered cells shown in Fig. 2, top left. In the same figure, the results of WAFR are given as smoothed ellipses, and the emphasized contours are the result of the MMAC algorithm (which started with the ellipses).

IV. FULL CELL-TRACKING ALGORITHM

In this section, we combine the active contours algorithm described in Section II with the segmentation method based on H-minima transform and ellipse fitting described in Section III into a full algorithm for cell tracking in the time-lapse microscopy images sequence.

We assume that the sequence $\{f^s\}_0^{S-1}$ of gray-scale images with nuclei is given as input (see Fig. 3, bottom right). Each image f^s of this sequence presents the location of nuclei at time T_s for $s=0,\ldots,S-1$. Furthermore, we assume that areas occupied by the same nucleus in two successive time steps overlap. In other words, each nucleus cannot cover a distance larger than its diameter in one time step.

The main steps of our algorithm are presented in Fig. 3.

In the initialization step, we binarize the gray-scale image f^0 (see Fig. 3, left top; the result is shown in part B). We note that the segmentation of nuclei in the first frame can be either automatic, e.g., using simple thresholding based on the c-means method, or manual in extended feature space based on textures.

Next, for each binary object detected in the initialization step, we apply the WAFR algorithm to divide overlapping objects and fit ellipses (see the resulting gray object in part A of Fig. 3). However, what is returned by this algorithm is not the optimal division yielded by the H-minima transform and watershed algorithm, but filled ellipses, fitted to such division (separated by their intersections), i.e., the contour C_i^0 of ith nucleus in f^0 is approximated by ellipse Q_i^0 for $i=1,\ldots,N^0$. In the next step, we define ψ_i^0 as the scaled signed distance function to the ellipse Q_i^0 for all $i=1,\ldots,N^0$. Next, for each $i=1,\ldots,N^0$, we apply the MMAC algorithm with ψ_i^1 and image f^1 as the input [this process is depicted in Fig. 3 as MMAC (Tracking)]. This algorithm returns level set functions ϕ_i^1 , where the zero level set corresponds to contours C_i^1 of the detected objects (see black

contours in Fig. 3, part E, right image). In the next step, the algorithm detects new objects that appear in the current frame f^s which are brighter than the minimal intensity from all objects tracked thus far (see the object indicated by the arrow in Fig. 3, part E, left image). The result of this step is indicated by arrows in the left image in part E of Fig. 3.

We repeat this three-step scheme for successive images from the given sequence $\{f^s\}_0^{S-1}$ with the following difference: Positive parts of $\phi_i^s, i=1,\dots,N^s$ (the number of objects at time point s) are now the inputs for the WAFR algorithm. Note that at this point, we still formally have N^s objects, although some ϕ_i^s can be positive in two disjoint areas. Then, if for some object WAFR returns more than one ellipse, these are treated as children.

If in addition images with labeled cytoplasms are given, then we can apply at each time point the MMAC algorithm to find corresponding contours of cytoplasm with ellipses returned by WAFR as the input. This is illustrated as *MMAC* (*Nucleus and Cytoplasm*) in part D of Fig. 3. Additionally, to find the exact shapes of nuclei detected by the WAFR algorithm, the MMAC algorithm is applied again (see Fig. 3, part D, lower image).

Depending on the application of the proposed algorithm, one can also store several quantities that describe cells in each image f^s for $s=0,\ldots,S-1$. These quantities can be, for instance, the mass center of a nucleus, average intensity of a cell, or some of its texture characteristics. We denote by \mathcal{F}^s_i the list of interesting features of the ith cell in the image f^s with $i=1,\ldots,N_s$ and $s=0,\ldots,S-1$. We present the full algorithm for cell tracking as follows.

Cell-tracking algorithm (part C in Fig. 3)

```
Require: Images \{f^s\}_{s=0}^{S-1}
 1: Initialisation with image f^0 (part B in Fig. 3)
 2: Apply WAFR algorithm to segment the binary image from
    the previous step resulting in N^0 ellipses Q^0_1, \ldots, Q^0_{N^0}
    (results in Fig. 3, part A)
 3: for s=1 \rightarrow S do
       Compute \psi_i^s(\cdot,Q_i^{s-1}) for i=1,\dots,N^{s-1}. for i=1\to N^{s-1} do
          Apply MMAC algorithm with \psi_i^s(\cdot, Q_i^{s-1}) and image
          f^s resulting in the level set function \phi_i^s.
          Apply the procedure of New Object Detection
          Apply WAFR algorithm with binary image obtained
          from \phi_i^s resulting in m(i) ellipses
          if m(i) > 1 then store new objects (as children)
10:
          Update N^s. Compute the set of features \mathcal{F}_i^s
11:
       end for
13: end for
14: return tracked objects with parent-child relation, list of
```

V. RESULTS OF NUMERICAL EXPERIMENTS

features \mathcal{F}_i^s , $i = 1 \dots N^s$, $s = 0 \dots S - 1$.

In this section, we present our experimental results of tracking cells in seven series of benchmark images, including five series of *HeLa-Kyoto* cell images expressing a histone H2BeGFP construct (H2BGFP [2]), one series of *HeLa mCherry* images, and

Feature	Dataset 1	Dataset 2	Datasets 3-7
Labeling	HeLa mCherry	HeLa Hoechst	HeLa-Kyoto series 1-5
channels	nucleus	nucleus	nucleus/cytoplasm
Image size	592 × 592	512 × 512	1344 × 1024
Resolution [pix/cm ²]	55452 ²	35840 ²	43404 ²
Cells per image (average)	10	12	16
Time-points	13	16	65
Time interval [min]	8	10	15
SNR mean and st. dev.	$\mu = 12.06, \sigma = 0.197$	$\mu = 5.72, \sigma = 0.041$	$\mu = 10.89, \sigma = 0.135$

TABLE I
CHARACTERISTICS OF DATASETS USED FOR ANALYSIS OF TRACKING PERFORMANCE

TABLE II Numerical Results (Average Over All Frames)

Method	sensitivity*	specificity*	precision*	F-score*		
Data set: HeLa mCherry						
CellProfiler	0.82	0.64	0.80	0.81		
Fiji	0.80	0.63	0.77	0.78		
LSetCellTracker	0.96	0.83	0.96	0.97		
Proposed	0.97	0.60	0.91	0.94		
Data set: HeLa Hoechst						
CellProfiler	0.37	0.19	0.32	0.34		
Fiji	0.58	0.38	0.51	0.53		
LSetCellTracker	0.91	0.66	0.88	0.90		
Proposed	0.93	0.67	0.89	0.91		
Data set: HeLa-Kyoto series 1						
CellProfiler	0.92	0.42	0.87	0.89		
Fiji	0.90	0.31	0.81	0.84		
LSetCellTracker	0.88	0.74	0.87	0.88		
Proposed	0.96	0.53	0.93	0.94		
Data set: HeLa-Kyoto series 2						
CellProfiler	0.95	0.58	0.92	0.94		
Fiji	0.95	0.53	0.90	0.92		
LSetCellTracker	0.91	0.85	0.91	0.91		
Proposed	0.97	0.62	0.94	0.96		
Data set: HeLa-Kyoto series 3						
CellProfiler	0.95	0.85	0.94	0.94		
Fiji	0.87	0.72	0.84	0.85		
LSetCellTracker	0.91	0.81	0.91	0.91		
Proposed	0.97	0.81	0.95	0.96		
Data set: HeLa-Kyoto series 4						
CellProfiler	0.91	0.69	0.89	0.90		
Fiji	0.92	0.64	0.88	0.90		
LSetCellTracker	0.85	0.74	0.85	0.85		
Proposed	0.96	0.67	0.93	0.95		
Data set: HeLa-Kyoto series 5						
CellProfiler	0.96	0.78	0.95	0.95		
Fiji	0.96	0.80	0.94	0.95		
LSetCellTracker	0.96	0.93	0.97	0.97		
Proposed	0.97	0.71	0.95	0.96		

one series of *HeLa Hoechst* (for details see Table I and [23]). For each input image, the SNR was calculated as SNR = $\mu_{\rm sig}/\sigma_{\rm bg}$, where $\mu_{\rm sig}$ is the mean value of pixels within objects, and $\sigma_{\rm bg}$ is the standard deviation of the background.

The images in *HeLa mCherry* and *HeLa Hoechst* series have lower SNR, nonhomogeneous cells, and varying light intensity. The tracking results are summarized in Table II as arithmetic means of sensitivity* (see definition further below), specificity*, and F-score* over all time points in the respective series. For each dataset the highest value of the respective measure is shown in bold.

In addition, to illustrate the quality of segmentation returned by the WAFR algorithm, we used four z-stacks of HEL cells. For all of these images, the WAFR algorithm detected the correct number of cells, whereas *CellProfiler*, *MTrack2*, and *LSetCell-Tracker* gave inferior results (see [23]).

We tested different sets of parameters and then used the one with the best tracking performance for all sets of images. It is worth noting that the algorithm is not very sensitive to parameters: Changing parameters by approximately 20% results in no more than 5% change in performance.

Quantitative performance evaluation was performed relative to ground truth, which was established independently by manual cell tracking by two experienced biologists.

We used experts' knowledge to define the best possible procedure for cell tracking in these programs. The most important steps include the following.

- 1) *In CellProfiler*: identify primary objects block with options: typical diameter (min, max) = (50, 250), try to merge small objects with larger, use Kapur Global for thresholding method; and TrackObjects block with the tracking method set as distance" and maximum pixel distance to consider matches set to 50.
- 2) In MTrack2: it needs 8-bit thresholded image; first, Gaussian blur with radius 2 is applied, then automatic threshold, and the last step is to launch MTrack2 module with options: minimum object size set to 200 pixels, maximum velocity set to 40, and minimum track length set to 1.
- 3) *In LSetCellTracker*: we tried different sets of parameters and finally used default ones which gave the best results.

Consider two consecutive frames f^{s-1} and f^s with all tracking information. We have N^{s-1} objects in frame s-1 and N^s in sth frame and also identification information. A tracked nucleus can be in solid state from frame s-1 to the next one, or can change its state (be nonsolid). That is, it can disappear, hide or divide. For each frame, we have four counters: TP, TN, FP, and FN (true positive, true negative, false positive, and false negative, respectively), all of which are initially equal to 0. For each object that occurs in one or in both adjacent frames, we increment the correct counter:

- 1) TP—if correct tracking of solid nucleus is detected;
- 2) TN—if correct tracking of nonsolid nucleus is detected;
- 3) FP—if incorrect tracking of solid nucleus is detected;
- 4) FN—if incorrect tracking of nonsolid nucleus is detected.

The solid or nonsolid state of a nucleus is determined by the ground truth as a positive or negative condition, respectively. Then, good or bad tracking can be treated as positive or negative test outcome. For measurement of the quality of tracking, we

used commonly known measures:

$$\begin{aligned} \text{sensitivity} &= \frac{\text{TP}}{\text{TP+FN}}, \text{specificity} &= \frac{\text{TN}}{\text{TN+FP}} \\ \text{precision} &= \frac{\text{TP}}{\text{TP+FP}}. \end{aligned}$$

In addition, we used the F-score measure which is defined as

$$F\text{-score} = \frac{2 \cdot \text{precision} \cdot \text{sensitivity}}{\text{precision} + \text{sensitivity}}.$$

To incorporate the quality of segmentation of clustered objects into the measurement of the tracking quality, we multiply the aforementioned measures by the term $\rho=1-\frac{|k-o|}{k+o}$, where k is a real number of cells (ground truth) and o is the number of cells considered by the algorithm. This term penalizes wrong segmentation of highly clustered cells. We identify measures multiplied by ρ with asterisk superscripts.

The results given in Table II show that the proposed algorithm significantly outperforms *CellProfiler* and *MTrack2* software. It slightly outperforms *LSetCellTracker* (the latter one is only better under specificity* measure), which moreover is much more sensitive to parameters' changes.

VI. CONCLUSION

We have presented here a method for extracting the position of individual cells from a sequence of time-lapse 2-D gray-scale microscopy images of cell nuclei and cytoplasms.

The contributions of this paper include: 1) the modification of a multiphase active contours method, where the force term is multiplied by a nonlinear scaling term (improved blurred boundaries detection); 2) improved watershed approach based on H-minima transform, which is integrated in the cell-tracking algorithm (including two new weights); 3) algorithm construction for easy parallelization, e.g., for GPU implementation. To demonstrate the latter, we produced a CUDA-optimized implementation on an NVIDIA GTX 295 GPU card. This led to an acceleration by a factor of approximately 13 compared to the CPU implementation (one core of an Intel Core i7 860 processor with 2.79 GHz clock speed). The robustness of the presented method, which can be classified as a model evolution approach for cell segmentation and tracking, is demonstrated by: 1) low sensitivity to SNR, e.g., see the performance of other methods in Table II for images with low SNR; 2) high efficiency for highly clustered overlapping cells by introducing WAFR algorithm with time complexity independent from the degree of cell clustering; 3) high efficiency of segmentation of nuclei with obscure boundaries caused by excessive blurring due to out-offocus light signal by introducing a nonlinear scaling in the force term.

An interesting alternative is to extend this approach for 3-D image data (e.g., z-stack obtained in confocal microscopy) which is on the agenda of our current research pursuits. In parallel, we also work on delivering for research community the specialized software library including the proposed algorithm in computational cloud using "Software as a Service" referred to as "on-demand software".

REFERENCES

- P. Vitorino and T. Meyer, "Modular control of endothelial sheet migration," *Genes Dev.*, vol. 22, no. 23, pp. 3268–3281, 2008
- [2] M. P. Stewart, J. Helenius, Y. Toyoda, S. P. Ramanathan, D. J. Muller, and A. A. Hyman, "Hydrostatic pressure and the actomyosin cortex drive mitotic cell rounding," *Nature*, vol. 469, no. 7329, pp. 226–230, 2011.
- [3] M. H. Schmitz and D. W. Gerlich, "Automated live microscopy to study mitotic gene function in fluorescent reporter cell lines," *Methods Mol. Biol.*, vol. 545, pp. 113–134, 2009.
- [4] S. Di Cataldo, E. Ficarra, A. Acquaviva, and E. Macii, "Achieving the way for automated segmentation of nuclei in cancer tissue images through morphology-based approach: A quantitative evaluation," *Comput. Med. Imag. Graph.*, vol. 34, no. 6, pp. 453–461, 2010.
- [5] E. Ficarra, S. D. Cataldo, A. Acquaviva, and E. Macii, "Automated segmentation of cells with IHC membrane staining," *IEEE Trans. Biomed. Eng.*, vol. 58, no. 5, pp. 1421–1429, May 2011.
- [6] A. Fitzgibbon, M. Pilu, and R. B Fisher, "Direct least square fitting of ellipses," *IEEE Trans. Pattern Anal. Mach. Intell.*, vol. 21, no. 5, pp. 476– 480, May 1999.
- [7] C. Jung and C. Kim, "Segmenting clustered nuclei using H-minima transform-based marker extraction and contour parameterization," *IEEE Trans. Biomed. Eng.*, vol. 57, no. 10, pp. 2600–2604, Oct. 2010.
- [8] S. Kothari, Q. Chaudry, and M. Wang, "Automated cell counting and cluster segmentation using concavity detection and ellipse fitting techniques," in *Proc. IEEE Int. Symp. Biomed. Imag.*, 2009, pp. 795–798.
- [9] X. Yang, H. Li, and X. Zhou, "Nuclei segmentation using marker-controlled watershed, tracking using mean-shift, and Kalman filter in time-lapse microscopy," *IEEE Trans. Circuits Syst. I, Reg. Papers*, vol. 53, no. 11, pp. 2405–2414, Nov. 2006.
- [10] J. Klein, S. Leupold, I. Biegler, R. Biedendieck, R. Münch, and D. Jahn, "TLM-Tracker: Software for cell segmentation, tracking and lineage analysis in time-lapse microscopy movies," *Bioinformatics*, vol. 28, no. 17, pp. 2276–2277, 2012.
- [11] F. Leymarie and M. D. Levine, "Tracking deformable objects in the plane using an active contour model," *IEEE Trans. Pattern Anal. Mach. Intell.*, vol. 15, no. 3, pp. 617–634, Jun. 1993.
- [12] N. Ray, S. Acton, and K. Ley, "Tracking leukocytes in vivo with shape and size constrained active contours," *IEEE Trans. Med. Imag.*, vol. 21, no. 10, pp. 1222–1235, Oct. 2002.
- [13] O. Debeir, I. Camby, R. Kiss, P. V. Ham, and C. Decaestecker, "A model-based approach for automated in vitro cell tracking and chemotaxis analyzes," *Cytometry*, vol. 60A, no. 1, pp. 29–40, 2004.
- [14] C. Zimmer, E. Labruyere, V. Meas-Yedid, N. Guillen, and J. Olivo-Marin, "Segmentation and tracking of migrating cells in videomicroscopy with parametric active contours: A tool for cell-based drug testing," *IEEE Trans. Med. Imag.*, vol. 21, no. 10, pp. 1212–1221, Oct. 2001.
- [15] C. Zimmer and J. Olivo-Marin, "Coupled parametric active contours," IEEE Trans. Pattern Anal. Mach. Intell., vol. 27, no. 11, pp. 1838–1842, Nov. 2005.
- [16] O. Dzyubachyk, W. A. van Cappellen, J. Essers, and W. J. Niessen, "Advanced level-set-based cell tracking in time-lapse fluorescent microscopy," *IEEE Trans. Med. Imag.*, vol. 29, no. 3, pp. 852–867, Mar. 2010.
- [17] O. Dzyubachyk, J. Essers, W. A. van Cappellen, C. Baldeyron, A. Inagaki, W. J. Niessen, and E. Meijering, "Automated analysis of time-lapse fluorescence microscopy images: From live cell images to intracellular foci," *Bioinformatics*, vol. 26, no. 19, pp. 2424–2430, 2010.
- [18] N. Paragios and R. Deriche, "Geodesic active contours and level sets for the detection and tracking of moving objects," *IEEE Trans. Pattern Anal. Mach. Intell.*, vol. 22, no. 3, pp. 266–280, Mar. 2000.
- [19] V. Caselles, R. Kimmel, and G. Sapiro, "Geodesic active contours," *Int. J. Comput. Vis.*, vol. 22, no. 1, pp. 61–79, 1997.
- [20] T. F. Chan and L. A. Vese, "A multiphase level set framework for image segmentation using the Mumford and Shah model," *Int. J. Comput. Vis.*, vol. 50, no. 3, pp. 271–293, 2002.
- [21] M. R. Lamprecht, D. M. Sabatini, and A. E. Carpenter, "CellProfilerTM free, versatile software for automated biological image analysis," *BioTechniques*, vol. 42, no. 1, pp. 71–75, 2007.
- [22] MTrack2 Software. (2003). [Online]. Available:http://valelab.ucsf. edu/~nico/IJplugins/MTrack2.html/
- [23] (2012). [Online]. Available:http://robusttracker.vratis.com/
- [24] T. F. Chan and L. A. Vese, "Active contours without edges," *IEEE Trans. Image Process.*, vol. 10, no. 2, pp. 266–277, Feb. 2001.
- [25] A. Dufour, R. Thibeaux, E. Labryere, N. Guillen, and J-Ch. Olivo-Martin, "3D Active meshes: Fast discrete deformable models for cell tracking in

- 3D time-lapse microscopy," IEEE Trans. Image Process., vol. 20, no. 7, pp. 1925-1937, Jul. 2011.
- [26] P. Yan, X. Zhou, and M. Shah, "Automatic segmentation of highthroughput RNAi fluorescent cellular images," IEEE Trans. Inf. Technol. Biomed., vol. 12, no. 1, pp. 109-117, Jan. 2008.
- [27] P. Soille, Morphological Image Analysis. New York, NY, USA: Springer-Verlag, 1998.



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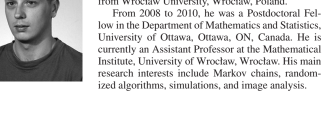
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