Imperial College London Department of Computing

# Automatic Cell Tracking in Noisy Images for Microscopic Image Analysis

by

Pedro Damian Kostelec

September 2014

Supervised by Ben Glocker

Submitted in part fulfilment of the requirements for the MSc degree in Computer Science (Artificial Intelligence) of Imperial College London

# Contents

1	Intr	roduction DRAFT I	7
	1.1	Motivation DRAFT I	7
	1.2	Objectives DRAFT I	8
	1.3	Contributions DRAFT I	8
	1.4	Report structure DRAFT I	9
2	$\mathbf{Rel}$	ated work DRAFT I	10
	2.1	Cell detection DRAFT I	10
		2.1.1 Cell segmentation using the Watershed technique DRAFT I	10
		2.1.2 Cell segmentation using level sets DRAFT I	11
		2.1.3 Cell detection by model learning DRAFT I	11
		2.1.4 Cell detection by image restoration DRAFT I	12
	2.2	Cell tracking DRAFT I	12
		2.2.1 Tracking by model evolution DRAFT I	13
		2.2.2 Tracking by frame-by-frame data association DRAFT I	13
		2.2.3 Tracking with a dynamics filter DRAFT I	14
		2.2.4 Cell tracking by global data association DRAFT I	14
	2.3	Conclusion DRAFT I	15
3			
3	Det	section of cells DRAFT I	17
3	<b>Det</b> 3.1	$egin{array}{cccc} egin{array}{cccc} egin{array}{cccc} egin{array}{ccccc} egin{array}{cccccc} egin{array}{cccccc} egin{array}{cccccc} egin{array}{cccccc} egin{array}{cccccc} egin{array}{ccccccccc} egin{array}{ccccccccc} egin{array}{ccccccccc} egin{array}{ccccccccc} egin{array}{ccccccccc} egin{array}{cccccccccc} egin{array}{cccccccccc} egin{array}{cccccccccc} egin{array}{cccccccccccccccccccccccccccccccccccc$	17 17
3			
3	3.1	Method overview DRAFT I	17
3	3.1 3.2	Method overview DRAFT I	17 18
3	3.1 3.2 3.3	Method overview DRAFT I	17 18 19
3	3.1 3.2 3.3 3.4	Method overview DRAFT I	17 18 19 20
<b>3 4</b>	3.1 3.2 3.3 3.4 3.5 3.6	Method overview DRAFT I	17 18 19 20 20
	3.1 3.2 3.3 3.4 3.5 3.6	Method overview DRAFT I  Detection of candidate regions DRAFT I  Inference under the non-overlap constraint DRAFT I  Learning the classifier DRAFT I  Feature selection DRAFT I  Performance improvements DRAFT I	17 18 19 20 20 21
	3.1 3.2 3.3 3.4 3.5 3.6	Method overview DRAFT I  Detection of candidate regions DRAFT I  Inference under the non-overlap constraint DRAFT I  Learning the classifier DRAFT I  Feature selection DRAFT I  Performance improvements DRAFT I  cking of cells DRAFT I	177 188 199 200 200 211
	3.1 3.2 3.3 3.4 3.5 3.6 <b>Tra</b> 4.1	Method overview DRAFT I  Detection of candidate regions DRAFT I  Inference under the non-overlap constraint DRAFT I  Learning the classifier DRAFT I  Feature selection DRAFT I  Performance improvements DRAFT I  cking of cells DRAFT I  Method overview DRAFT I	177 188 199 200 210 244 244
	3.1 3.2 3.3 3.4 3.5 3.6 <b>Tra</b> 4.1 4.2	Method overview DRAFT I  Detection of candidate regions DRAFT I  Inference under the non-overlap constraint DRAFT I  Learning the classifier DRAFT I  Feature selection DRAFT I  Performance improvements DRAFT I  cking of cells DRAFT I  Method overview DRAFT I  Joining cell detections into robust tracklets DRAFT I	177 188 199 200 210 244 244
	3.1 3.2 3.3 3.4 3.5 3.6 <b>Tra</b> 4.1 4.2 4.3	Method overview DRAFT I  Detection of candidate regions DRAFT I  Inference under the non-overlap constraint DRAFT I  Learning the classifier DRAFT I  Feature selection DRAFT I  Performance improvements DRAFT I  cking of cells DRAFT I  Method overview DRAFT I  Joining cell detections into robust tracklets DRAFT I  Global data association DRAFT I	177 188 199 200 211 244 246 277
	3.1 3.2 3.3 3.4 3.5 3.6 <b>Tra</b> 4.1 4.2 4.3 4.4	Method overview DRAFT I  Detection of candidate regions DRAFT I  Inference under the non-overlap constraint DRAFT I  Learning the classifier DRAFT I  Feature selection DRAFT I  Performance improvements DRAFT I  cking of cells DRAFT I  Method overview DRAFT I  Joining cell detections into robust tracklets DRAFT I  Global data association DRAFT I  Implementation using linear programming DRAFT I	177 188 199 200 210 244 244 266 277 299
	3.1 3.2 3.3 3.4 3.5 3.6 <b>Tra</b> 4.1 4.2 4.3 4.4	Method overview DRAFT I  Detection of candidate regions DRAFT I.  Inference under the non-overlap constraint DRAFT I  Learning the classifier DRAFT I  Feature selection DRAFT I  Performance improvements DRAFT I  cking of cells DRAFT I  Method overview DRAFT I  Joining cell detections into robust tracklets DRAFT I  Global data association DRAFT I  Implementation using linear programming DRAFT I  Hypotheses likelihood definitions DRAFT I	177 188 199 200 210 244 244 266 277 299 300
	3.1 3.2 3.3 3.4 3.5 3.6 <b>Tra</b> 4.1 4.2 4.3 4.4 4.5	Method overview DRAFT I  Detection of candidate regions DRAFT I  Inference under the non-overlap constraint DRAFT I  Learning the classifier DRAFT I  Feature selection DRAFT I  Performance improvements DRAFT I  cking of cells DRAFT I  Method overview DRAFT I  Joining cell detections into robust tracklets DRAFT I  Global data association DRAFT I  Implementation using linear programming DRAFT I  Hypotheses likelihood definitions DRAFT I  Computing the likelihoods DRAFT I	177 188 199 200 211 244 246 277 299 300 311

		4.7.3 Best feature selection NEW	35		
5	Data acquisition and annotation DRAFT I				
	5.1	Data acquisition and example datasets	36		
		5.1.1 Datasets DRAFT I	37		
		5.1.2 Imaging analysis challenges DRAFT I	41		
	5.2	The annotation tool DRAFT I	42		
6	Exp	perimental results NEW	45		
	6.1	Cell detector NEW	45		
		6.1.1 Speed NEW	45		
		6.1.2 Detection accuracy NEW	47		
	6.2	Cell tracker NEW	47		
		6.2.1 Performance metrics NEW	47		
		6.2.2 Speed NEW	47		
		6.2.3 Tracking accuracy	47		
7	Conclusions and future work DRAFT I				
	7.1	Conclusion DRAFT I	48		
	7.2	Future work DRAFT I	49		
$\mathbf{A}_{\mathbf{l}}$	Appendices				
$\mathbf{A}$	Use	r Guide for the Annotation Tool	<b>52</b>		
В	Use	r Guide for the Interactive Annotation Viewer	53		
$\mathbf{C}$	Use	r Guide for the Cell Detector and Tracker	<b>54</b>		
Bi	Bibliography				

## 5 Data acquisition and annotation DRAFT I

This chapter describes the data that influenced the decisions of selecting the cell detection and tracking methods. In section 5.1 we briefly describe the imaging method used to acquire the image sequences and present some example datasets. Section 5.1.2 describes some of the challenges of tracking cells in these datasets. Section 5.2 presents the data annotation tool that was developed to ease the annotation process.

## 5.1 Data acquisition and example datasets

As discussed in the concluding section of chapter 2, the datasets heavily influenced the choice of algorithms for cell detection and tracking. Many computer vision algorithms rely on heuristics to improve their accuracy. In cell detection methods, this is obvious from the fact that a algorithm developed for a certain type of imaging method will likely perform poorly on an different types of images (e.g. different shape of cells). In cell tracking heuristics help adjust the algorithms to the specific behaviour of cells that are being analysed. For example, a different tracking method could be used for images with cells that move slowly (and there is a large overlap between cells in consecutive frames) than for cells that move quickly (and there is little overlap between cells in consecutive frames).

The data acquisition process is not part of this research. However, for the reasons stated above, it is important to know the characteristics of the datasets. Because these are directly influenced by the imaging method, we follow by outlining the image acquisition process, and then present the datasets used in the evaluation of our methods.

The image sequences that inspired the development of the methods describe in this thesis were acquired *in vivo*. This means that the images are obtained on living mice, in contrast to *in vitro* where cells are analysed on a tissue sample in a standard laboratory environment using petri dishes and other instruments. *In vivo* analysis is preferred over *in vitro* because it is better suited for observing the behaviour of cells in their natural environment.

Recent advances in intravital microscopy of the lung [28] and other organs allow us to image neutrophils and other cells of interest simultaneously by combining fluorescent reporter mice and intravenously-injected fluorescently-labelled antibodies. Briefly, the mouse is anaesthetized, mechanically ventilated and the surface of the lung is exposed surgically. Then a specialized "vacuum chamber" (as in [28]) is used to stably hold a glass coverslip against the surface of the lung. The alveoli, alveolar capillaries and associated cells can then be visualized by laser scanning confocal

microscopy. The same apparatus can be used to interrogate other organs without mechanical ventilation, e.g. the liver.

All the data was provided by Dr. Leo Carlin from the Leukocyte Biology Section at the National Heart and Lung Institute (NHLI)<sup>1</sup>.

## 5.1.1 Datasets DRAFT I

From the datasets provided by Dr. Leo Carlin, five have been selected to use in the evaluation of this work because of their distinct characteristics. The original dimensions of the datasets were 512-by-512 pixels, but some have been cropped to reduce the number of cells that had to be annotated to train the cell detector and tracker.

Below we describe the main characteristics of the datasets, together with the number of frames that were annotated to train the cell detector. Because of the large number of trajectories in certain datasets, not all the trajectories have been annotated. Instead, in each dataset a few trajectories were chosen and annotated. We aimed to select the longest trajectories, as these could give the most insight in the analysis of cell behaviour.

#### Dataset A

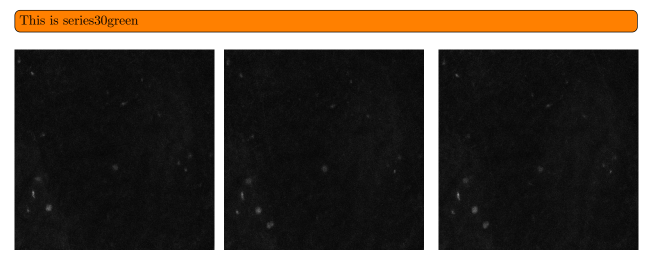


Figure 5.1: Three consecutive frames from dataset A.

Dataset A, seen in fig. 5.1, contains 66 frames of dimensions 512-by-512 pixels. All the frames have been manually annotated for the purpose of training the cell detector. The dataset only has an average population of 1.7 cells per frame (with a standard deviation  $\sigma = 0.7$ ), and all four main trajectories have been annotated, although three of them can only be tracked in a dozen frames. The dataset is obtained from the liver. There is a good contrast between the cells and the background. The background is almost uniform but still contains some information about the architecture of the liver sinusoids, which is difficult to discern without adjusting the image contrast. The cells appear

<sup>1</sup>http://www1.imperial.ac.uk/nhli/

as medium dark shades of grey of circular shape and their boundaries smoothly blend with the background. The cells move slowly. The images are of constant quality, and there are few significant camera artefacts.

Question for Leo: What is the difference between these cells and the ones in dataset B? They are taken simultaneously... one is series 30 red the other series 30 green (they were just different channels in the provided dataset)

#### Dataset B

# This is series30red

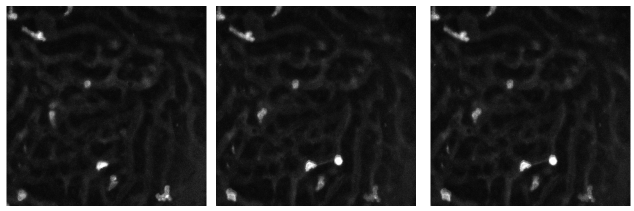


Figure 5.2: Three consecutive frames from dataset B.

Figure 5.2 shows three consecutive frames from dataset B. The dataset contains 66 frames of dimensions 512-by-512 pixels and an average population of 7.5 cells per frame ( $\sigma = 1.6$ ). The entire dataset has been annotated for the purpose of training the cell detector and (TODO 5) of the longest trajectories for the purpose of training the cell tracker. This dataset is also obtained from the lung

#### Lung for sure?

. The cells appear brighter than in dataset A, but their shapes vary. Some are round and others elongated and deformed to fit into the tight blood vessels

#### Are these blood vessels?

in which they move. Their brightness varies from medium to very bright. The brighter ones have clearer boundaries than the medium bright ones, whose boundaries are softer. In the background we can clearly discern the blood vessels in a darker grey colour. Cells in this dataset are active. Some move relatively fast and change direction frequently. The images are of constant quality, and there are few significant camera artefacts.

Although datasets A and B were obtained simultaneously, cells in dataset A were marked with a fluorescent reporter (green fluorescent protein (GFP)) that the mouse has been genetically modified to produce just in a type of immune cells termed 'NK' cells, while those in dataset B were marked by diffuse staining (unbound Ab in circulation), which is weaker and colors the cells more diffusely.

#### Dataset C

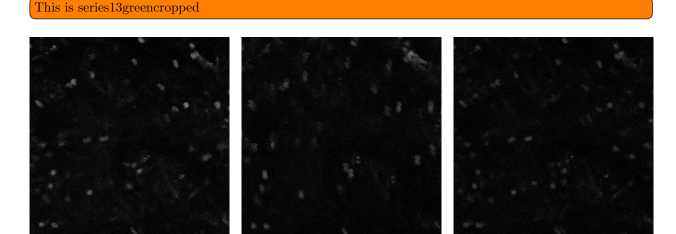


Figure 5.3: Three consecutive frames from dataset C.

Dataset C, seen in fig. 5.3, is composed of 126 frames which have been cropped to 251-by-251 pixels. The first 55 frames have been annotated with dot-annotation for the purpose of training the cell detector, and 5 of the longest trajectories were annotated for the purpose of training the cell tracker. This dataset is also obtained from the lung

#### Lung for sure?

. The dataset contains an average population of 19.7 cells per frame ( $\sigma=4.1$ ) as estimated from the first 55 annotated frames that were manually annotated. The background is dark, but contains several very faint cells which we are not interested in tracking. We are interested in tracking the brighter cells. The colours of the cells are dark to medium grey. The cell boundaries blend smoothly with the background. Most cells appear as round patches. Although the cells are mostly stagnant, but the lung tissue is moving due to the breathing of the mice. For this reason, the dataset contains many motion artefacts. Some frames are blurred, and many appear to contain the duplicates of the cells shifted in position by a small amount, as seen in the middle image in fig. 5.3.

#### Dataset D

#### This is series14croppedclean

Dataset D, seem in fig. 5.4, is likely to be the most challenging for the tracking module. It was originally composed of 682 frames. Unfortunately, the dataset contained large sequences of frames that were of exceptionally low quality, such that even a human could not track the cells accurately. Thus, the dataset has been manually pruned to 377 frames and cropped to dimensions 199-by-199

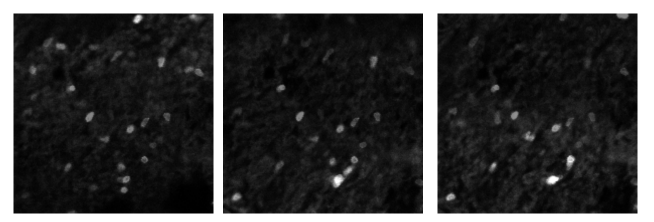


Figure 5.4: Three consecutive frames from dataset D.

pixels. 50 frames have been annotated with dot-annotations and 7 of the longest trajectories. Dataset D is obtained from the liver

#### Liver for sure?

. The cell density is 10.6 cells per frame ( $\sigma=2.3$ ). The background contains some noise, but it is still relatively different to the texture of the cells. The cells are mostly elliptical and have a strong boundaries. This dataset contains several fast moving cells. As mentioned before, the dataset contains many motion artefacts, the contrast of the images is constantly changing, and often about half of the cells in each frame become invisible for a few frames, even after manually eliminating the most affected.

#### Dataset E

# This is seriesm170\_13cropped

Figure 5.5: Three consecutive frames from dataset E.

The last dataset, seen in figure fig. 5.5, contains 194 frames which have been cropped to dimensions 277-by-277 pixels. 65 frames were annotated for the purpose of training and evaluating the quality of the cell detection module. (TODO) of the longest trajectories were annotated for training the tracker.

#### Where is the dataset from?

The density of the cells is of 6.6 cells per frame ( $\sigma = 2.4$ ). The background contains some texture, that could sometime be confused with the cell texture. The cells in this dataset appear smaller than in the other datasets and are less elliptical. They contain more sharp edges. The intensity of the cells varies from dark to bright. The cells appear to move very slowly. The images contain many artefacts that make tracking more difficult. First, the images become darker in the later parts of the image sequence. Second, there is an alternating dark/bright area in all the frames probably caused by the camera shutter.

## 5.1.2 Imaging analysis challenges DRAFT I

In this section we are going to present two phenomena caused by moving tissue and the camera shutter that make this datasets especially hard to track for frame-by-frame tracking methods.

The first challenge is the artefacts caused by the camera shutter. These appear as alternating dark/bright horizontal lines which seem to move from the top to the bottom of the images. The effect can be seen in fig. 5.6. This makes the tracking problem difficult, because the cells behind the dark areas appear fainter or disappear completely for the few frames that they are covered.

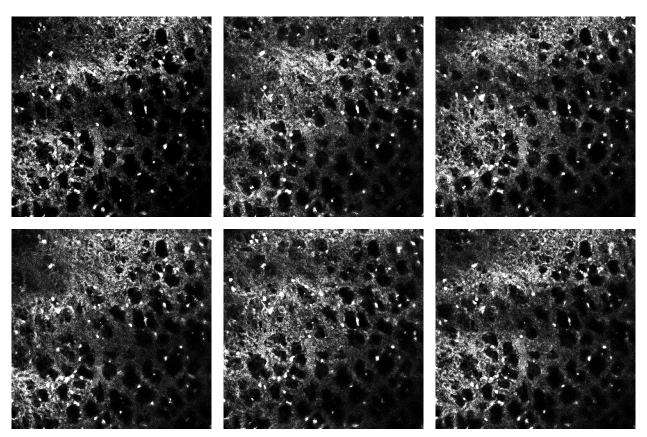


Figure 5.6: Six consecutive frames from a sequence displaying the alternating dark/bright regions caused by the camera shutter. The contrast and sharpness of the images has been adjusted to dramatize the effect.

The second challenge is the movement of the tissue, especially in the case of lung imaging. The

shaking can be in the x-y plane, but sometimes, as in the case shown in figure fig. 5.7 it can be primarily in the z-direction. Displacement in the x-y direction causes the cells to jiggle. Displacement of the tissue in the z-direction causes part of all of the image to become out-of-focus for a few frames.

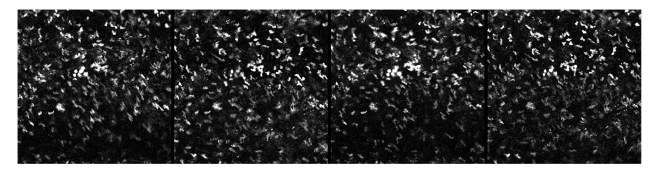


Figure 5.7: Four consecutive frames taken from the lung displaying varying cell clarity due to the displacement caused by the lung movement. The contrast of the images has been adjusted to dramatize the effect.

Several of the datasets present these phenomena to a lesser or larger extend. These artefacts where the primary reason for choosing a global optimization method for associating detection responses into trajectories instead of a frame-by-frame approach.

## 5.2 The annotation tool DRAFT I

In order to train the cell detector and tracker it is necessary to annotate a few frames from the datasets. The cell detector requires dot annotations - a single dot within each cell - that indicate whether an extremal region corresponds to a cell. The detector is then trained to recognize extremal regions similar to the ones that were annotated. Similarly, the cell tracker needs annotations that indicate whether two cells from different frames belong to the same tracklet. The cell tracker is then trained based on the similarity of the appearance and spatio-temporal features extracted from these cells.

To train a good detector and tracker it is important that the annotations are consistent. For example, several datasets contain cells that appear bright and some that are darker. If we are only interested in tracking the cells that consistently appear bright, the annotations should only include the bright cells. The models may not be trained correctly if only part of the bright cells were annotated. Note that this paragraph makes observations about brightness because the concept is easy to understand for a reader. However, the detector and tracker rely on many more features for detection and tracking.

To facilitate the annotations of datasets a new annotation tool has been developed. The graphic user interface of the tool is visible in fig. 5.8. The tool is able to load a sequence of images, display them and permits the user to annotate them. The user is able to preform the following actions: adding a dot detection, deleting a dot detection, adding a link between cells, deleting a link between cells. The tool also features zooming and panning.

Furthermore, the tool includes a broad set of filters that can be applied on the images to increase

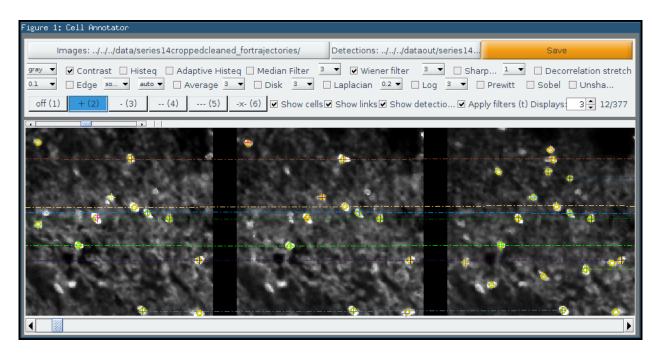


Figure 5.8: Screenshot of the annotation tool. User annotations are shown as crosses and dashed links between them, while detection responses are are shown as yellow circles.

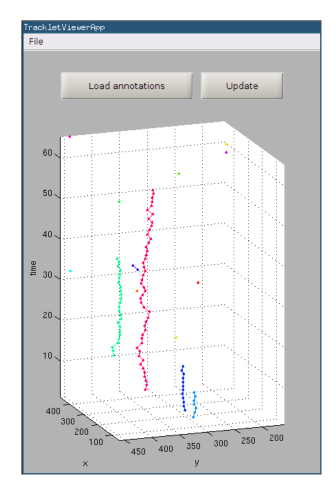


Figure 5.9: Screenshot of the interactive 3D tracklet viewer showing annotations for Dataset A.

the clarity of the cells and ease the annotation process. These filter include contrast adjustments, smoothing, sharpening, edge detection, etc. To reduce the clutter on densely populated datasets, the user is able to only display annotations from a specific part of the image, hide trajectories that include more than three dot annotations, hide the part of the GUI containing the buttons and filters in order to focus on just the images (full-screen mode), and more.

The annotation tool includes detection of potentially bad annotations. These include a warning system that displays a triangle next to a cell if two annotations are very close together and a different display style for links that deviate too much from the average (in terms of between-frame displacement). These help the eliminate accidental human errors, that would be very hard to discover otherwise.

Finally, the tool can also be used to compare the detection responses obtained from the cell detector by overlaying them over the images. In fig. 5.8 these detections are shown as yellow circles. This can be very useful to visually evaluate the quality of the detections, and possibly reveal cells that the user might have missed when annotating.

In addition to the annotation tool, another small GUI has been developed that shows the annotations in an interactive 3D view, where each tracklet colour is different. This tool is shown in image fig. 5.9. It is helpful to recognize possibly bad annotations (for example an outlier cell within a trajectory) or discover missed links.

All the annotations have been reviewed by Dr. Leo Carlin, who provided the datasets. However, due to the noisy nature of the images, some of the dataset are difficult to correctly annotate even for a human. It is expected that a small number of annotations are erroneous.

Both tools were developed and packaged as standalone executables with MATLAB 8.3.0.532 (R2014a). The tools do not depend on MATLAB to run, but require the free MATLAB Compiler Runtime 8.3<sup>2</sup> which is automatically downloaded and installed when the applications are installed on the system. User guides for both applications are provided in appendices A and B.

Will I include user guides in the appendix?

<sup>&</sup>lt;sup>2</sup>http://www.mathworks.co.uk/products/compiler/mcr/

Appendices

## Bibliography

- [1] P. K. Elzbieta Kolaczkowska, "Neutrophil recruitment and function in health and inflammation," 2013. 7
- [2] J. Pillay, I. den Braber, N. Vrisekoop, L. M. Kwast, R. J. de Boer, J. A. M. Borghans, K. Tesselaar, and L. Koenderman, "In vivo labeling with 2h2o reveals a human neutrophil lifespan of 5.4 days," *Blood*, vol. 116, no. 4, pp. 625–627, 2010.
- [3] P. S. Tofts, T. Chevassut, M. Cutajar, N. G. Dowell, and A. M. Peters, "Doubts concerning the recently reported human neutrophil lifespan of 5.4 days," *Blood*, vol. 117, no. 22, pp. 6050–6052, 2011. 7
- [4] C. Arteta, V. Lempitsky, J. A. Noble, and A. Zisserman, "Learning to detect cells using non-extremal regions," in *Proceedings of the 15th International Conference on Medical Im*age Computing and Computer-Assisted Intervention - Volume Part I, MICCAI'12, (Berlin, Heidelberg), pp. 348–356, Springer-Verlag, 2012. 8, 9, 11, 17, 18, 19, 20, 21, 22, 48
- [5] Y. Chen, K. Biddell, A. Sun, P. Relue, and J. Johnson, "An automatic cell counting method for optical images," in [Engineering in Medicine and Biology, 1999. 21st Annual Conference and the 1999 Annual Fall Meetring of the Biomedical Engineering Society] BMES/EMBS Conference, 1999. Proceedings of the First Joint, vol. 2, pp. 819 vol.2—, Oct 1999. 10
- [6] X. Chen, X. Zhou, and S.-C. Wong, "Automated segmentation, classification, and tracking of cancer cell nuclei in time-lapse microscopy," *Biomedical Engineering*, *IEEE Transactions on*, vol. 53, pp. 762–766, April 2006. 10, 13
- [7] L. Vincent, "Morphological grayscale reconstruction in image analysis: applications and efficient algorithms," *Image Processing, IEEE Transactions on*, vol. 2, pp. 176–201, Apr 1993. 10
- [8] J. Serra, Image Analysis and Mathematical Morphology. Orlando, FL, USA: Academic Press, Inc., 1983. 10
- [9] D. Mukherjee, N. Ray, and S. Acton, "Level set analysis for leukocyte detection and tracking," Image Processing, IEEE Transactions on, vol. 13, pp. 562–572, April 2004. 11, 13
- [10] C. Tang, Y. Wang, and Y. Cui, "Tracking of active cells based on kalman filter in time lapse of image sequences of neuron stem cells." 11, 14
- [11] D. Xu and L. Ma., "Segmentation of image sequences of neuron stem cells based on level-set

56 Bibliography

algorithm combined with local gray threshold.," Master's thesis, Harbin Engineering University, 2010. 11

- [12] C. Arteta, V. S. Lempitsky, J. A. Noble, and A. Zisserman, "Learning to detect partially overlapping instances.," in *CVPR*, pp. 3230–3237, IEEE, 2013. 11, 12, 19
- [13] J. Matas, O. Chum, M. Urban, and T. Pajdla, "Robust wide baseline stereo from maximally stable extremal regions," in *Proceedings of the British Machine Vision Conference*, pp. 36.1–36.10, BMVA Press, 2002. doi:10.5244/C.16.36. 11
- [14] T. Joachims, T. Finley, and C.-N. J. Yu, "Cutting-plane training of structural syms," Mach. Learn., vol. 77, pp. 27–59, Oct. 2009. 11
- [15] R. Bise, T. Kanade, Z. Yin, and S. il Huh, "Automatic cell tracking applied to analysis of cell migration in wound healing assay," in *Engineering in Medicine and Biology Society, EMBC*, 2011 Annual International Conference of the IEEE, pp. 6174–6179, Aug 2011. 12, 25
- [16] S. Huh, Toward an Automated System for the Analysis of Cell Behavior: Cellular Event Detection and Cell Tracking in Time-lapse Live Cell Microscopy. PhD thesis, Robotics Institute, Carnegie Mellon University, Pittsburgh, PA, March 2013. 12, 13
- [17] D. House, M. Walker, Z. Wu, J. Wong, and M. Betke, "Tracking of cell populations to understand their spatio-temporal behavior in response to physical stimuli," in *Computer Vision and Pattern Recognition Workshops*, 2009. CVPR Workshops 2009. IEEE Computer Society Conference on, pp. 186–193, June 2009. 13
- [18] B. Xu, M. Lu, P. Zhu, Q. Chen, and X. Wang, "Multiple cell tracking using ant estimator," in Control, Automation and Information Sciences (ICCAIS), 2012 International Conference on, pp. 13–17, Nov 2012. 14
- [19] K. Li and T. Kanade, "Cell population tracking and lineage construction using multiple-model dynamics filters and spatiotemporal optimization," in *Proceedings of the 2nd International* Workshop on Microscopic Image Analysis with Applications in Biology (MIAAB), September 2007. 14
- [20] A. Massoudi, D. Semenovich, and A. Sowmya, "Cell tracking and mitosis detection using splitting flow networks in phase-contrast imaging," in *Engineering in Medicine and Biology* Society (EMBC), 2012 Annual International Conference of the IEEE, pp. 5310–5313, Aug 2012.
- [21] L. Zhang, Y. Li, and R. Nevatia, "Global data association for multi-object tracking using network flows," in Computer Vision and Pattern Recognition, 2008. CVPR 2008. IEEE Conference on, pp. 1–8, June 2008. 15, 27
- [22] C. Huang, B. Wu, and R. Nevatia, "Robust object tracking by hierarchical association of detection responses," in *Computer Vision ECCV 2008* (D. Forsyth, P. Torr, and A. Zisserman,

Bibliography 57

- eds.), vol. 5303 of *Lecture Notes in Computer Science*, pp. 788–801, Springer Berlin Heidelberg, 2008. 15, 27
- [23] R. Bise, Z. Yin, and T. Kanade, "Reliable cell tracking by global data association.," in *ISBI*, pp. 1004–1010, IEEE, 2011. 15, 25, 27, 30, 48
- [24] H. Kuhn, "The hungarian method for the assignment problem," Naval Research Logistics Quarterly, vol. 2, pp. 83–97, 1955. 15
- [25] J. Matas, O. Chum, M. Urban, and T. Pajdla, "Robust wide-baseline stereo from maximally stable extremal regions," *Image and Vision Computing*, vol. 22, no. 10, pp. 761 767, 2004. British Machine Vision Computing 2002. 18
- [26] I. Tsochantaridis, T. Hofmann, T. Joachims, and Y. Altun, "Support vector machine learning for interdependent and structured output spaces," in *Proceedings of the Twenty-first International* Conference on Machine Learning, ICML '04, (New York, NY, USA), pp. 104–, ACM, 2004. 20
- [27] K. Li, E. D. Miller, M. Chen, T. Kanade, L. E. Weiss, and P. G. Campbell, "Cell population tracking and lineage construction with spatiotemporal context," *Medical Image Analysis*, vol. 12, no. 5, pp. 546 566, 2008. Special issue on the 10th international conference on medical imaging and computer assisted intervention {MICCAI} 2007. 25, 50
- [28] M. Looney, E. Thornton, D. Sen, W. Lamm, R. Glenny, and M. Krummel, "Stabilized imaging of immune surveillance in the mouse lung.," *Nature Methods*, vol. 8, no. 5, pp. 91–6, 2011-01-01 00:00:00.0. 36