

Cell Nuclei Segmentation in Cytological Images Using Convolutional Neural Network and Seeded Watershed Algorithm

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Abstract

Morphometric analysis of nuclei is crucial in cytological examinations. Unfortunately, nuclei segmentation presents many challenges because they usually create complex clusters in cytological samples. To deal with this problem, we are proposing an approach, which combines convolutional neural network and watershed transform to segment nuclei in cytological images of breast cancer. The method initially is preprocessing images using color deconvolution to highlight hematoxylin-stained objects (nuclei). Next, convolutional neural network is applied to perform semantic segmentation of preprocessed image. It finds nuclei areas, cytoplasm areas, edges of nuclei, and background. All connected components in the binary mask of nuclei are treated as potential nuclei. However, some objects actually are clusters of overlapping nuclei. They are detected by their outlying values of morphometric features. Then an attempt is made to separate them using the seeded watershed segmentation. If the attempt is successful, they are included in the nuclei set. The accuracy of this approach is evaluated with the help of referenced, manually segmented images. The degree of matching between reference nuclei and discovered objects is measured with the help of Jaccard distance and Hausdorff distance. As part of the study, we verified how the use of a convolutional neural network instead of the intensity thresholding to generate a topographical map for the watershed improves segmentation outcomes. Our results show that convolutional neural network outperforms Otsu thresholding and adaptive thresholding in most cases, especially in scenarios with many overlapping nuclei.

Keywords Convolutional neural networks · Watershed · Nuclei segmentation · Breast cancer · Oversegmentation · Mathematical morphology

Introduction

According to global cancer project (GLOBOCAN), breast cancer is the most common cancer among women worldwide. It was estimated that in 2012, nearly 1.7 million new cases were diagnosed (second most common cancer overall) and 521,907 cases of deaths due to breast cancer occurred (fifth cause of death from cancer overall). This represents about 12% of all new cancer cases and 25% of all cancers in women [3].

Breast cancer is mostly diagnosed by three medical examinations usually occurring in the following order: palpation, ultrasonography or mammography, and fine

 Michał Żejmo mzejmo@uz.zgora.pl needle biopsy (FNB). In this work, we concentrate on the analysis of results of FNB examination in an automatic way. In FNB method, a cytological material is gathered directly from tumor using a fine needle. Next, it is fixed and stained with hematoxylin and eosin. Finally, glass slide with the cellular material is examined by the pathologist under the microscope. FNB is less traumatic and much safer for a patient than an open surgical biopsy. However, detection of cancer cells on the slide glass is not an easy task. Novice pathologists to gain experience and knowledge to become professional specialist must spend a lot of time browsing various cytological samples. Moreover, analysis of entire slide is a time-consuming process, even for experienced pathologists.

During the examination of cytological images, pathologists evaluate morphometric features of cells and their nuclei in order to distinguish tumor type. In recent years, we have seen a very intensive development of techniques dedicated to microscopic digital imaging. More and more specialists browse virtual slides on a computer screen instead of examining glass slides under a microscope.



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This situation opens up the possibility of supporting the pathologist's work using modern image processing and machine learning techniques. They can speed up diagnosis and increase their accuracy. However, we must remember that the accuracy of nuclei segmentation is critical to the performance of computer-assisted cytology. Unfortunately, cytological images are rather challenging for existing segmentation methods because nuclei have the tendency to create complex structures like clumps or nests.

The most common approaches for nuclei segmentation are based on active contours, intensity thresholding, mathematical morphology, region growing, watershed, and deep learning [4, 6, 10, 11, 18]. Last years brought an enormous progress in classification and object recognition using Convolutional Neural Networks (CNN) [2, 29]. It seems to be a promising technique for semantic segmentation of cytological images [24, 25]. The most considerable advantage of CNN is its ability to classify a single pixel based on its neighborhood described by highlevel features. That is why excellent results are obtained while detecting the nuclei and cytoplasm even when their staining is strongly heterogeneous. Problems arise when nuclei overlap and there are no clear boundaries between them. In this situation, ordinary CNN networks are not able to separate such nuclei. It seems that watershed method deals better with such cases. However, the results obtained by the watershed are strongly dependent on the quality of the topographical map of distances. To obtain precise topographical map, we need a precise mask of nuclei region. Usually, intensity thresholding or extended h-minima are used for this purpose [7, 23]. Unfortunately, intensity thresholding usually generates nuclei mask that contain objects with jagged contours and clumped objects. As an effect, watershed usually is affected by the oversegmentation. On the other hand, the results of extended h-minima can vary substantially with regard to the value of chosen h. According to the results presented in [16], the method can miss finding some nuclei for some values of h.

To tackle these problems, we propose an approach that combines the advantages of CNN and the watershed method. CNN is employed to detect precise nuclei mask, which then is used to generate topographical map and nuclei seeds for watershed. Watershed is applied to separate overlapping nuclei. Experimental studies have shown that the use of CNN instead of the usual thresholding to determine the nuclei mask significantly increases the accuracy of nuclei segmentation.

The remainder of this paper is organized as follows. Section "Data" presents material used for experiments. The details of the proposed approach are described in the "Methods" and "Data Preprocessing" sections. Section "Experiment" presents the details of the experiments carried out. Their results are shown in the "Results" section. The

paper ends with conclusions and future research in the "Conclusions and Further Research" section.

Data

Cytological images used in this study came from 40 patients of the University Hospital in Zielona Góra, Poland. Half of the cases are malignant, half are benign. All tumors were histologically confirmed, and all patients who had a benign disease were biopsied or followed up for a year. Cellular material was acquired from affected tissue using 0.5-mmdiameter needle under the control of an ultrasonograph. The material was fixed and then dyed with hematoxylin and eosin (H+E). Glass slides were scanned using VS120 Olympus Virtual Microscopy System. The system consists of a 40x lens and 2/3" CCD camera giving 0.17 μm resolution. As a result, 40 slides were generated. The average size of slide is approximately $200k \times 100k$ pixels. For each virtual slide 2 regions of interests (ROI) of size, 1583 × 828 pixels representing malignant or benign cells were selected and saved as 8 bit/channel RGB TIFF image. Example of ROI selected by the pathologist from the virtual slide is shown in Fig. 4.

In total, we have collected 80 images. We divided them into subset 1 (40 images coming from 10 benign cases and 10 malignant cases) and subset 2 (40 images coming from the rest 10 benign cases and the rest 10 malignant cases). Subset 1 was used to train and validate CNN, subset 2 was used as a test subset to verify the accuracy of the proposed approach. To train CNN, we have divided subset 1 (40 images) into training subset (20 images coming from 5 malignant cases and 5 benign cases) and validation subset (20 images coming from 5 malignant cases and 5 benign cases). Thus, images used for training CNN were never used to the validation procedure. Moreover, the experiment conducted to verify the effectiveness of the segmentation procedure was carried out using test images. They do not include images coming from patients that were used for training and validation of CNN.

Methods

In this section, we describe the main steps of the proposed segmentation method: (1) semantic segmentation of nuclei and background using CNN; (2) determining connected object on the semantic map generated by CNN; (3) detection of connected clusters of objects (clustered nuclei) based on their area and roundness; (4) applying conditional erosion to determine nuclei seeds among clumped objects; (5) separation overlapping nuclei using seeded watershed; (6) aggregating segmentation results for overlapping and



non-overlapping nuclei. The first step is implemented using Python and Keras library, the rest of the steps are implemented using Matlab.

The watershed transform is one of the most often used segmentation method to separate touching or overlapping objects. However, the method is effective if proper seeds of objects are given. Here, we are proposing conditional erosion to detect centers (seeds) of prospective nuclei. Erosion is carried out on nuclei mask. Therefore, it is crucial for the method to determine precise mask of the nuclei. If nuclei overlap, we need at least to see some parts of their silhouettes to determine their centers using conditional erosion. Overlapping nuclei on inaccurate nuclei mask usually create huge clumps and thus even conditional erosion will fail to find their centers. To deal with this challenge, we propose CNN as a tool for recognizing nuclei regions. We also used to segment nuclei regions two other methods based on intensity thresholding: Otsu thresholding (GO) and adaptive thresholding (AT). Finally, we compared the accuracy of CNN with both intensity thresholding approaches.

Convolutional Neural Network

In recent years, CNN has gained a lot of popularity as a tool for image segmentation and object recognition [12]. Typical CNN is usually comprised of at least with two convolutional layers combined with pooling layers and ended by at least one fully connected layer (Fig. 6).

- **a)** Convolutional layer is a core part of CNN, composed of a set of learnable filters. Each filter extracts different features from the input image. Filter parameters (weights) are tuned during the learning procedure.
- **b) Pooling layer** is used to progressively reduce the spatial size of the input in order to extract higher level features. Spatial size reduction is usually done by max pooling using window of size 2×2 pixels.
- c) Fully connected layer is at the end of CNN and is connected to all activations in the previous layer. The input of this layer is a one-dimensional feature

vector. The task of this layer is to capture the complex relationships between high-level features and output labels.

Trained CNN model was used for semantic segmentation. It classifies each pixel from the input image into one of four categories (nuclei, cytoplasm, nuclei edge, background). In fact, the output of CNN contains probability distribution over four classes. Therefore, each pixel is always labeled by the class which gained the highest probability. As a result, we get a semantic mask for input image.

Segmentation of Nuclei Region

In this step, semantic mask generated by CNN model is transformed into nuclei mask. Pixels belonging to nuclei are labeled by 1, while others by 0. Therefore, such mask is very similar to binary masks generated by GO or AT method. Figure 1 presents sample results for CNN, GO, and AT. We can visually asses that CNN is much more precise in nuclei segmentation than GO and AT. It can be observed that CNN separates the nuclei that are touching and overlapping much better than two other techniques.

Detection of Overlapping Nuclei

All cytological images used in this study were manually annotated by marking nuclei contours. Therefore, it is possible to compute morphometric features of nuclei and determine their distributions (see Fig. 2). We decided to describe nuclei by their area and roundness [20]:

$$Roundness = \frac{4 \times Area}{(Perimeter)^2}. (1)$$

Based on 4447 manually annotated nuclei, it can be concluded that nuclei have area in the range from 309 to 7801 pixels and roundness in the range from 0.31 to 0.99. Based on these findings, we are able to distinguish the nucleus from the object which consists of many clumped nuclei.



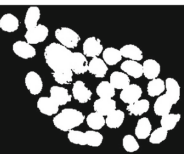




Fig. 1 Segmentation of nuclei region: CNN (left), AT (middle), and GO (right)

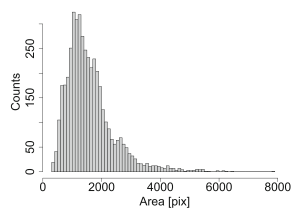


Fig. 2 Histograms of the nuclei areas and roundness

Nuclei mask determined by CNN usually contain two types of regions that we call non-overlapping or overlapping. In non-overlapping regions, objects are easily recognized as nuclei because they are well separated, and thus their areas and roundness are within allowed limits. These objects are immediately classified as nuclei and they do not require further processing. The rest of the objects that do not meet the requirements of being single nucleus must be subjected to a separation procedure by seeded watershed.

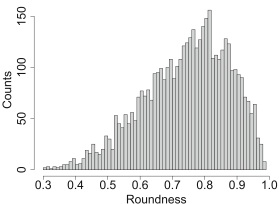
Separating Overlapping Nuclei Using Seeded Watershed

The classical watershed transform treats the image to be segmented as a topographic surface. It segments image by flooding basins from the seeds until basins attributed to different seeds meet on watershed lines. The input of the algorithm is usually a binary mask of the image. It is transformed by the Euclidean distance transform and local maxima from this transform are used as seeds. Unfortunately, algorithm in this form tends to create many micro-segments. Such over-segmentation makes the results of the watershed method completely useless. To deal with this problem, we used extended version of watershed that uses nuclei seeds generated by the conditional erosion [9, 30]. The method process binary mask to detect centers of objects. In our case, it was applied to nuclei mask I_{NM} to find nuclei centers I_S (seeds). We applied this processing only to the objects classified as nuclei clumps.

In our approach, conditional erosion is based on classical erosion defined as the operation of structuring element B on image I_{NM} :

$$I_{NM} \ominus \check{B} = \{ x \in \mathbb{R} | (B + x) \subset I_{NM} \}, \tag{2}$$

where \check{B} is a reflection of set B. Conditional erosion is conducted in two steps. First, coarse erosion using structuring element M_c reduces the size of objects. To prevent objects from disappearing, coarse phase switches



to fine phase for objects with area below T_1 . Fine erosion uses structuring element M_f which is less likely to make the nucleus disappear. It tries to separate clustered nuclei. Element M_f is used iteratively until all objects have area below T_2 . The size of both structuring elements should be significantly smaller than the size of the processed objects to not to reduce them too rapidly. Based on the reference nuclei (manually segmented), we know that the size of nuclei can vary from 300 to 8000 pixels, thus we can use the coarse structuring element M_c and fine structuring element M_f proposed in [30]:

$$M_c = \begin{pmatrix} 0 & 0 & 0 & 1 & 0 & 0 & 0 \\ 0 & 0 & 1 & 1 & 1 & 0 & 0 \\ 0 & 1 & 1 & 1 & 1 & 1 & 0 \\ 0 & 1 & 1 & 1 & 1 & 1 & 0 \\ 0 & 1 & 1 & 1 & 1 & 1 & 0 \\ 0 & 0 & 1 & 1 & 1 & 0 & 0 \\ 0 & 0 & 0 & 1 & 0 & 0 & 0 \end{pmatrix}, \qquad M_f = \begin{pmatrix} 0 & 1 & 0 \\ 1 & 1 & 1 \\ 0 & 1 & 0 \end{pmatrix}.$$

To determine thresholds T_1 and T_2 , we conducted a segmentation test on a few chosen images. Segmentation results were compared with those obtained by using classical watershed segmentation. We determined experimentally that the best segmentation is obtained for thresholds $T_1 = 350$ and $T_2 = 50$.

The prototype of topographic map I_{TM} is determined by Euclidean distance transform. Next, seeds I_S are used to refine the topographic map. Seeds I_S are combined with the original topographic map I_{TM} by morphological reconstruction $\rho_{I_{TM}}(I_S)$ [28]. The algorithm is based on repeated dilations of a seed mask I_S until the contour of the seed mask fits under a topographic map I_{TM} :

$$I'_{TM} = \rho_{I_{TM}}(I_S) = \bigcup_{n \ge 1} \delta_{I_{TM}}^{(n)}(I_S).$$
 (3)

The grayscale geodesic dilation of size n is then given by:

$$\delta_{I_{TM}}^{(n)}(I_S) = \underbrace{\delta_{I_{TM}}(\dots \delta_{I_{TM}}(\delta_{I_{TM}}(I_S)))}_{n}, \tag{4}$$



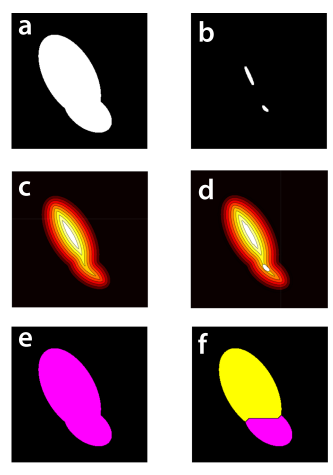


Fig. 3 Segmentation of overlapping ellipses using seeded watershed and classical watershed: **a** input mask; **b** result of conditional erosion; **c** distance transform; **d** modified distance transform with imposed seeds; **e** watershed using distance transform; **f** watershed using seeded distance transform

and the elementary geodesic dilation is described by the following relationship:

$$\delta_{I_{TM}}(I_S) = (I_S \oplus B) \cap I_{TM},\tag{5}$$

where $(I_S \oplus B)$ is a standard dilation of size one followed by an intersection (pointwise minimum \cap) and B is 4connected neighborhood structural element with pair of horizontal and vertical connected pixels [28]. In Fig. 3, we can observe how the seeds generated by conditional erosion modify original topographic map and the positive effect of such approach on the segmentation of overlapping objects with the elliptic shape.

The approach described above is used to separate nuclei clusters found in nuclei mask by procedure described in the "Detection of Overlapping Nuclei" section. The whole separating procedure employing conditional erosion and watershed is applied to each object classified as overlapping objects (clumped nuclei). Therefore, we obtain separate segmentation results for each nuclei cluster. After the separation procedure by seeded watershed, objects

detected are again classified as overlapping or nonoverlapping nuclei using the method described in the "Detection of Overlapping Nuclei" section. We do this to check if the separation was successful. All nuclei that successfully pass the test of being single nucleus are included in the final segmentation results. Clusters of nuclei that have failed to be separated are rejected and do not participate in further processing.

Data Preprocessing

Input images are preprocessed by color separation procedure and cutting into blocks of fixed size (patches). Patches are further processed and augmented to prepare training data for CNN.

Color Separation

Cytological samples are subjected to a staining process to precisely visualize the cellular material that is being analyzed. We usually use hematoxylin and eosin for this purpose. Hematoxylin is mainly absorbed by the cell nuclei and dyes them blue. Eosin dyes the cellular material in red and deposits mainly in the cytoplasm. Unfortunately, the process of staining and digitizing glass slides is not standardized. In effect, cytological samples coming from different laboratories may differ in color. Color variation can arise due to different staining protocols, different stain brands, a shelf life of stains, or due to using different microscopy scanners. It has been shown that the performance of segmentation algorithms deteriorates substantially when the color of processed images differs from the color of training images [8, 15, 17]. To tackle this problem, various color normalization methods have been proposed. They can be generally categorized into histogram matching methods, color transfer methods, and spectral matching methods (For a complete overview of the state of the art color normalization methods please see [22]).

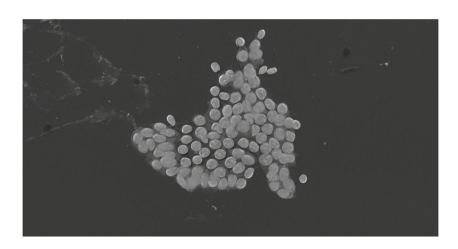
Color normalization is usually preceded by the stain separation because different cellular structures absorb all stains to some extent. The stain concentration is closely related to the attenuation of the light transmitted through the stained material. In turn, light transmission through the cytological sample can be described by the Beer Lambert law:

$$I = I_0 \exp\left(-WH\right),\tag{6}$$

where I is the intensity of the light that passed the sample $(3 \times n \text{ matrix}, n \text{ - number of pixels})$, I_0 is the intensity of light entering the sample (matrix of the same size as I), W



Fig. 4 Image representing hematoxylin concentration



is a stain color matrix of size $3 \times k$ (k = number of stains), and matrix H ($k \times n$) represents the stain concentrations. Stain separation approaches can be divided into supervised and unsupervised methods. The well-known supervised method is based on color deconvolution [21]. In this method matrix, W is determined empirically and it uses pseudo inverse transform to obtain H. By contrast, unsupervised methods estimate both W and H. Usually these methods are based on independent component analysis (ICA) or non-negative matrix factorization (NMF) [1, 19]. Another approach is based on a model where image colors are linear combinations of stain vectors which describe the proportion of absorbed light in RGB channels. Due to the fact that absorption weights are non-negative, thus every value may exist between stain vectors. The method exploits this fact to find them using two largest singular values determined by singular value decomposition (SVD) [14].

In this study, we applied a supervised method of stain separation proposed in [21]. The effectiveness of this algorithm is confirmed by numerous research publications and moreover it is easily available for many scientific computing environments. Our implementation is based on built-in stain vectors for Hematoxylin/Eosin taken from ImageJ color deconvolution plugin. Absorption spectra of hematoxylin and eosin overlap in RGB space, but mentioned color separation allows us to some extent evaluate the contribution of hematoxylin and eosin at

Table 1 Collections of patches

Number of images	Training 20	Validation 20
Total patches	18,315,912	18,961,569
 nuclei border 	1,739,200	1,968,232
- nuclei center	5,105,059	5,788,305
- cytoplasm	5,588,398	5,770,039
background	5,883,255	5,434,993

each pixel. Three separate intensity images are created as a result of deconvolution, the first represents the hematoxylin concentration, second eosin concentration, and third residuals. For further processing, we are using images of hematoxylin concentration. They emphasize nuclei and suppress cytoplasm which absorbs mainly eosin (Fig. 4).

After color separation, hematoxylin image is subjected to feature-wise standardization. Each pixel in the image array is treated as a separate feature. Based on the randomly chosen sample of 10,000 training image patches, we can determine the mean and standard deviation for every feature (pixel). They are stored and then used to standardize images. Every pixel in the image is standardized using a mean and standard deviation determined exactly for his position in the image

Manual Segmentation

To train CNN, we needed ground truth images with labeled objects. Therefore, all images used in this study were subjected to manual segmentation. The procedure was carried out using ImageJ software¹ and involved selection of four types of objects: nuclei interior, nuclei contour, cytoplasm, and background. In the case of ambiguity, e.g., if overlapping nuclei cannot be separated, no objects were marked. But, if the separation was possible, then nuclei were marked as separate objects.

Based on selected regions, semantic maps were created for each image. On a semantic map, each pixel is given the label and can belong to the following classes: indeterminate, background, cytoplasm, nuclei, or nuclei border. Set of pixels belonging to nuclei border was determined automatically by extracting pixels from nuclei regions lying on the contours of nuclei.



¹https://imagej.nih.gov/ij/

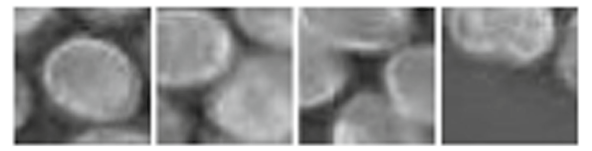


Fig. 5 Example of training patches (hematoxylin concentration). From left: nuclei, nuclei border, cytoplasm, background

Extraction of Patches

The images were cut into patches of size 43×43 pixels. The size of the patch was chosen arbitrary so that the patches contain large fragments of cell nuclei. For each pixel, a single patch is extracted. The class of each patch is assigned based on the label of central pixel. Label is read from semantic map. Patches corresponding to unlabelled pixels are excluded from further processing and thus are not used to train or validate CNN. As in many other studies concerning semantic segmentation, CNN is deciding about the class of the pixel based on the patch centered in that pixel. Set of all patches was divided into training and validation subsets, their sizes, and distributions are presented in Table 1. Examples of patches used for training CNN are shown in Fig. 5.

Patches Augmentation, Transformation, and Preprocessing

The raw patches generated in the previous step were augmented and preprocessed to improve the learning process of the CNN model. It is important for learning procedure to have balanced number of training samples within each class. In our case, the class of patches describing

nuclei borders is underrepresented (see Table 1). To overcome this problem, a set of nuclei border patches was augmented by artificial patches. They were generated using some well-known transformations applied to actual patches. To generate new patches, each original nuclei border patch was subjected to three randomized transformations: scaling by a factor from range 0.8 to 1.2, rotating using random angle, flipping vertically and/or horizontally. The set of patches was enlarged four times using augmentation technique. To increase the diversity and variability of patches coming form other classes, they were also randomly subjected to these transformations. In order not to increase the size of these classes, the original patch after being transformed was replaced by the new one.

Experiment

Network Architectures and Training Parameters

A lot of different CNN architectures have been already described and tested in the scientific literature [12, 13, 27]. They vary in the layer configuration and depth of the structure. The structure of CNN used in this study is shown in Fig. 6. Our network consists of four convolutional

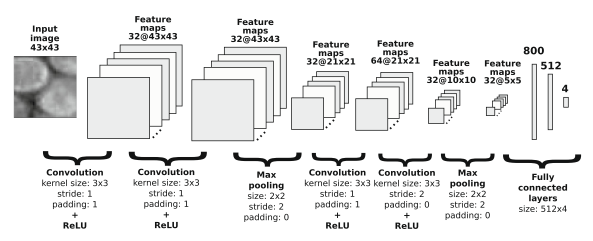


Fig. 6 The structure of CNN



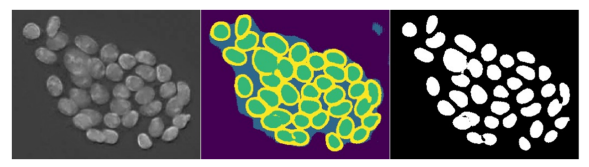


Fig. 7 Results of semantic segmentation using CNN: input image: hematoxylin concentration (left), semantic mask generated by CNN classifier (middle), and nuclei mask (right)

layers, which are separated by two max-pooling layers. At the top of the network, we placed two fully connected layers (512 neurons and 4 neurons respectively). All convolutional layers are followed by rectified linear units (ReLU). Weights in convolutional layers were initialized using Xavier method, learning rate was set to 10^{-4} , and weight decay was set to 10^{-6} [5]. Training was conducted using stochastic gradient descent, mini-batch was set to 256, and training process was finished after 20 epochs. We applied dropout technique to prevent the network from over-fitting [26]. This allowed us to achieve the 91.33% classification accuracy for the validation set. All experiments were realized with the help of GeForce GTX TITAN X with 12GB of RAM.

Semantic Segmentation

Trained CNN model was used to predict classes for all pixels in the images from the test set. The input of the network is a single patch. The output is a class probability distribution for a central point of the patch. To segment the whole image, classification procedure must be repeated for each pixel in the image. As a result, we obtain a class probability distribution for every pixel. Finally, pixel is

labeled by the class that achieved the highest probability. In this way, a semantic mask was obtained for a given input image. From the semantic mask, it is possible to extract nuclei mask to generate a topographic surface for the watershed transform. Sample semantic mask and nuclei mask are shown in Fig. 7.

Results

Evaluation Procedure

In order to verify the effectiveness of the proposed approach, it was applied to detect nuclei in 20 test images. The accuracy of CNN was compared with the accuracy of GO and AT method. These methods were used to determine the nuclei mask but final segmentation was always carried out using seeded watershed (see "Segmentation of Nuclei Region"). Thus, three methods used to extract nuclei mask were compared with respect to watershed segmentation accuracy.

To measure the accuracy of automatic segmentation, we compare nuclei segmented automatically with reference nuclei segmented manually. We are given a list of manually

Table 2 Results for Hausdorff distance

	TP			FP		
	CNN	AT	GO	CNN	AT	GO
Benign						
Mean	83.4%	52.4%	51.3%	5.1%	14.6%	14.9%
Sd	12.8%	21.6%	19.0%	4.1%	5.0%	5.8%
Max	98.2%	86.1%	77.2%	15.5%	23.1%	30.8%
Min	41.0%	7.7%	10.3%	0.0%	3.8%	6.3%
Malignant						
Mean	78.1%	54.4%	56.6%	21.4%	27.9%	19.2%
Sd	11.7%	10.8%	12.1%	13.4%	12.4%	7.2%
Max	93.7%	77.8%	85.2%	51.9%	63.6%	32.7%
Min	56.3%	38.7%	37.0%	3.2%	14.5%	8.1%

The best results are italicized



Table 3 Results for Jaccard distance

	TP			FP		
	CNN	AT	GO	CNN	AT	GO
Benign						
Mean	77.6%	50.1%	49.6%	10.9%	16.9%	16.5%
Sd	14.2%	22.6%	20.0%	6.5%	6.5%	7.6%
Max	93.3%	81.9%	76.8%	24.6%	27.4%	38.5%
Min	38.5%	7.7%	10.3%	1.3%	7.1%	7.5%
Malignant						
Mean	73.2%	55.1%	57.8%	26.3%	27.2%	18.0%
Sd	11.4%	10.9%	13.7%	11.2%	11.9%	6.7%
Max	90.5%	81.5%	92.6%	52.7%	54.5%	29.3%
Min	53.1%	38.7%	37.5%	12.0%	10.9%	3.7%

The best results are italicized

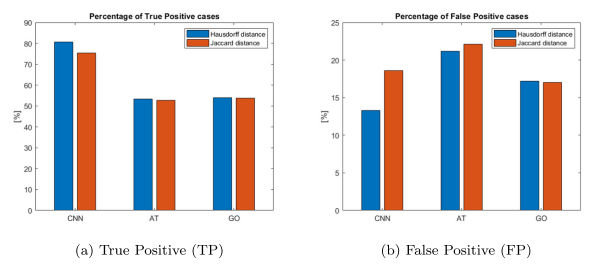


Fig. 8 Aggregated TP rates and FP rates with regard to methods of segmentation

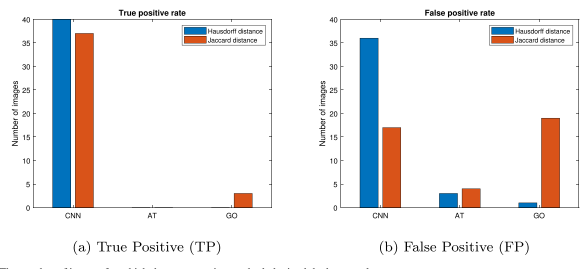


Fig. 9 The number of images for which the segmentation method obtained the best result



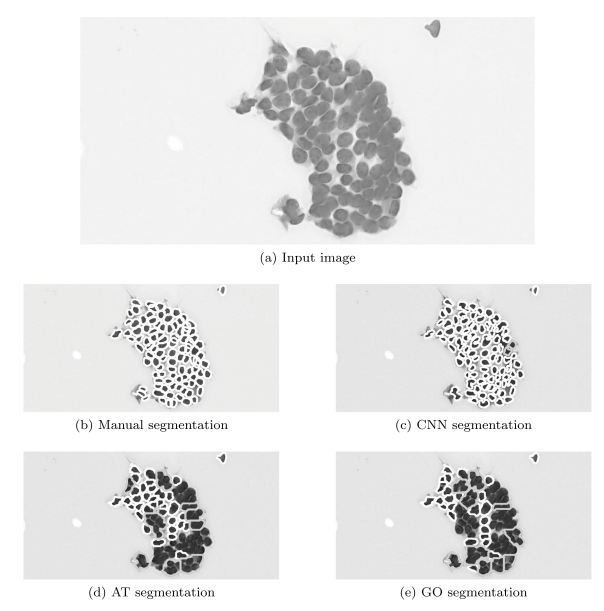


Fig. 10 Sample segmentation results: TP (white), FP (gray)

segmented nuclei in the form of binary masks. Nuclei segmented automatically can be also presented in the form of binary masks. Therefore, it is possible to measure distances between reference nuclei and automatically segmented nuclei. To do that, we are using Hausdorff distance (HD) and Jaccard distance (JD). For each segmentation method, (CNN + seeded watershed, GO + seeded watershed, AT + seeded watershed), two distance matrices were determined using Hausdorff distance and Jaccard distance respectively. The distance matrix stores the distances between all pairs of manually segmented nuclei and nuclei segmented using a chosen automatic segmentation method. Using distance matrices, we are trying to pair all manually segmented nuclei with the closest nuclei segmented automatically. We assumed that

the nuclei between which Hausdorff distance is greater than 30 or Jaccard distance is greater than 0.5 are so different that must be considered as two separate objects. The single manually segmented nucleus can be paired with the only one, nearest nucleus segmented automatically and the distance between them must be below the predefined threshold. As a result, 3 scenarios are possible: manually segmented nucleus can be matched with the nearest automatically segmented nucleus and such case is classified as true positive (TP), no automatically segmented nucleus can be found to match with the manually segmented nucleus and such case is classified as false negative (FN), and automatically segmented nucleus can stay without corresponding manually segmented nucleus and thus it is classified as false positive (FP).



Experimental Results

Tables 2 and 3 summarize basic statistics for TP and FP coefficients determined for all test images. The results are presented separately for benign and malignant cases. Aggregated (for benign and malignant cases) mean values of TP and FP rates are shown in Fig. 8.

We can observe that CNN method outperforms GO and AT for benign cases. On average, CNN-based approach can extract 83.4% of benign nuclei according to HD and 77.6% according to JD. Whereas the methods based on GO and AT are able to extract from 51.3 to 52.4% of benign nuclei according to HD and from 49.6 to 50.1% according to JD. We also see that CNN-based segmentation outperforms other methods for malignant cases. For malignant cases, CNN-based method can correctly detect 78.1% of nuclei according to HD and 73.2% according to JD, while for the other methods average values are in range 54.4%–56.6% for HD and 51.1%–57.8% for JD.

The CNN method received on average lower FP error for benign cases than the other methods, while for malignant cases FP error is slightly larger than for GO method. The obtained values of standard deviations indicate that for all methods results are distributed in similar range around the mean.

In Fig. 9, we have presented results that indicate how many times each segmentation method reached the highest TP rate and the lowest FP rate. CNN clearly outperforms two other approaches, because it gained the best results for most test images.

Classical intensity thresholding methods (AT, GO) are very effective for detecting areas occupied by the nuclei if nuclei are well separated. Unfortunately, these methods usually fail if overlapping nuclei are present in an image. This problem is much less visible if we use CNN instead of GO or AT. Sample segmentation results confirming this fact are shown in Fig. 10. The CNN method is more effective than AT and GO in detecting overlapping objects, but the side effect is the detection of redundant objects. Thus, the value of FP for some images can slightly increase if we use CNN segmentation. However, on average CNN method performs much better than GO and AT.

Conclusions and Further Research

Methods of nuclei segmentation based on intensity thresholding, edge detection, watershed transform, active contours and artificial neural networks usually have problems with cytological images. Mainly because of complex and heterogeneous nature of these images which often contain overlapping and touching nuclei. In this paper, we are presenting an alternative approach which uses CNN

classifier to pre-segment nuclei and then seeded watershed to deal with overlapping nuclei. We are showing that our approach outperforms classical watershed method. Moreover, we determined that the proposed method can detect about 81% of real nuclei (on average) and, at the same time, have a low rate of false positive detection (13%).

Despite the fact that obtained result looks quite promising, there are some points of our project which can be improved. Semantic segmentation using a classical convolutional neural network turned out to be very computationally expensive. Especially training process was time consuming. In future research, we are going to use for semantic segmentation fully convolutional network (e.g., U-Net network) which will allow us to significantly reduce the computational burden of the method.

Another drawback of the method is that it needs manually segmented nuclei to train CNN. Unfortunately, the process of manual segmentation is very time consuming and tedious. Therefore, we plan to develop a software for semi-automatic segmentation of nuclei, an operator will have to only correct the errors introduced by the automatic method. Such approach allows us to prepare richer set of training images and save time.

We also plan to modify the step responsible for detection of overlapping nuclei coming from semantic segmentation. We are going formally determine the thresholds for roundness and area based on hypothesis testing framework.

Our system can be used to segment and detect nuclei. Therefore, we plan to develop a computer-aided cytology system to help diagnose breast cancer and lung cancer. Based on the segmentation results of our system, we plan to compute morphometric, textural, and colorimetric features of nuclei and apply machine learning techniques to make predictions about the type of the cancer.

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