MR-NOM: MULTI-SCALE RESOLUTION OF NEURONAL CELLS IN NISSL-STAINED HISTOLOGICAL SLICES VIA DELIBERATE OVER-SEGMENTATION AND MERGING

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ABSTRACT

In comparative neuroanatomy, the characterization of brain cytoarchitecture is critical to a better understanding of brain structure and function, as it helps to distill information on the development, evolution, and distinctive features of different populations. The automatic segmentation of individual brain cells is a primary prerequisite and yet remains challenging.

A new method (MR-NOM) was developed for the instance segmentation of cells in Nissl-stained histological images of the brain. MR-NOM exploits a multi-scale approach to deliberately over-segment the cells into superpixels and subsequently merge them via a classifier based on shape, structure, and intensity features. The method was tested on images of the cerebral cortex, proving successful in dealing with cells of varying characteristics that partially touch or overlap, showing better performance than two state-of-the-art methods.

Index Terms—cell segmentation, histological images, brain, nissl, multi-scale, laplacian of gaussian, superpixels

1. INTRODUCTION

Comparative neuroanatomy studies investigate anatomical changes between the brains of populations defined by factors such as sex, age, pathology, or species. The characterization of brain cytoarchitecture holds special significance in such studies, as it can provide insights into the links between the specific structure of the brain and the animal morphology, behavior, or environment [1–3].

In a typical analysis pipeline, tissue sections (i.e., histological slices) of brain specimens are processed with Nissl stain to label neuronal cells [4] and are fixed for digitization as Whole Slide Images (WSI) for subsequent examination. Due to their size and complexity, WSIs are preferably processed by computerized methods, which can ensure reproducibility and speed in high-throughput pipelines, while a manual examination would be prohibitively time-consuming as well as impacted by inter- and intra-observer bias.

A critical prerequisite in such pipelines is the challenging instance segmentation of cells. Fig. 1a shows 3 tiles extracted from Nissl-stained histological slices of the auditory cortex from different brain specimens of *Tursiops truncatus* (Montagù, 1821), an example of which is shown in Fig. 1b, with the highlighted area representing the primary (A1) and secondary (A2) auditory cortex. Segmentation of individual cells is complicated by their dishomogeneity in shape, texture, and size, due to the co-existence of large and small neurons, glia and endothelial cells, the presence of touching or overlapping cells with ambiguous boundaries, and background clutter.

In the wider field of digital pathology and microscopy, many segmentation methods have been proposed to segment cells/nuclei. The vast majority rely on a set of underlying algorithms: intensity thresholding, morphology operations, watershed transform, deformable models, clustering, graph-based approaches, and supervised classification [5]. Few,
however, are designed for the segmentation of cells in Nissl-stained histological slices of the brain [6–8], and most are conceived for cells of uniform characteristics.

In light of the above, a new method called MR-NOM was developed to be used in an active learning fashion to facilitate the construction of ground truth annotations and to subsequently segment cells in WSIs. This method will be exploited for the characterization of brain cytoarchitecture in comparative neuroanatomy studies, and in particular as an enabler of solid morphometric analyses aimed at objective tissue screening in the field of diseases affecting brain structure and functionality (e.g., neurodegeneration and neuroinflammation).

This paper is organized as follows: Section 2 describes the dataset; Section 3 details the steps of the method; Section 4 illustrates the results; Section 5 draws the conclusions.

2. DATASET

Brain tissues were sampled from 20 specimens of Tursiops truncatus archived in the Mediterranean Marine Mammals Tissue Bank (http://www.marinemammals.eu) of the University of Padova. The brains originated from stranded cetaceans with a decomposition and conservation code (DCC) of 1 and 2, according to the guidelines for cetacean post-mortem investigation [9].

The images used in this study are 27 2048x2048 tiles extracted from Nissl-stained 40x magnification WSIs of the auditory cortex of Tursiops truncatus, also known as the bottlenose dolphin, originating from different subjects (newborn, adult, old). The tiles were annotated via QuPath [10] software, leading to 13,986 annotated cells. 4 and 3 tiles were used as validation and test set, respectively.

3. METHODS

3.1. Pre-processing

Each image was converted to grayscale and filtered with a 2-D Gaussian smoothing filter with standard deviation (SD) of 1. Contrast-limited adaptive histogram equalization was applied to enhance the contrast while avoiding noise amplification. The mean neuropil (area between cell bodies) intensity was standardized by applying a correcting factor as

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Contrast-limited adaptive histogram equalization was applied to enhance the contrast while avoiding noise amplification. A Gaussian smoothing filter with standard deviation (SD) of 2 was applied to the standardized image $I_1(x, y)$ to get $I_2(x, y)$. A combined foreground multi-scale map $R_{FG}(x, y)$ (Fig. 2b) was then computed as

$$R_{FG}(x, y) = \sum_{i=1}^{n_{FG}} R_{FG,i}(x, y),$$

where $n_{FG} = 10$, $R_{FG,i}(x, y) = \sigma_{FG,i}^2 I_i * I_2(x, y)$, with $\gamma_{FG} = 1$ and $\{\sigma_{FG,i}\}_{i=1}^{n_{FG}} = \{5, 6, ..., 14\}$. $R_{FG}$ was rescaled to the intensity range 0–255, normalized to match the mean of $I_2(x, y)$, and summed to $I_2(x, y)$ to enhance the neuronal cells. The resulting map was thresholded via the triangle method [11] to extract the foreground, which was refined through hole-filling, morphological opening and closing, and removal of connected components smaller than 70 pixels, H-connected, and spur pixels. The final foreground map $FG(x, y)$ (Fig. 2c) was obtained by removing most of the poorly focused or too bright objects, using two morphological reconstructions starting from the maps $EM(x, y)$ and $NS(x, y)$ defined in Section 3.1.
The first term corresponds to the complement of $R$ weighing gradient information by adding the gradient map cells) appear dark on a bright background. The second term as defined in Section 3.3, so that objects of interest (neuronal information. It was defined as follows:

$$W(x, y) = R_{MK}(x, y)^c + \alpha_1 \sigma_{MK} \gamma_{MK}$$

The first term corresponds to the complement of $R_{MK}(x, y)$ as defined in Section 3.3, so that objects of interest (neuronal cells) appear dark on a bright background. The second term weighs gradient information by adding the gradient map $GM$ defined in Section 3.1, with $\alpha_1$ controlling the importance given to gradient cues (set to 0.15 via hyperparameter validation). $\alpha_2 = R_{MK}^{\gamma_{MK}}/GM$ is a standardization factor to make the maps comparable by matching the mean of $GM$ to the mean of $R_{MK}$. Prior to any other operation, $R_{MK}$ and $GM$ were rescaled to the range $0 \rightarrow 1$.

$W$ was modified using morphological reconstruction to impose the markers from Section 3.3 and the Skeleton by Influence Zones (SKIZ) of the FG map as regional minima [15] (Fig. 2e). The watershed transform was then applied to get the label image $LB(x, y)$ (Fig. 2f), where a different label, i.e., an integer value, is assigned to each identified region. All the pixels of $LB$ in the background were set to 0.

3.5. Supervised superpixels merging

Ideally, each cell would correspond to a single region in the label image $LB$. However, due to cell variability, more than one marker is often associated with larger non-circular cells with diverse characteristics. These cells are represented by a set of multiple regions (or superpixels) in $LB$. Drawing inspiration from [16–18], a classifier was trained to decide whether a pair of adjacent superpixels has to be merged.

For every candidate merge, let $S_1$ and $S_2$ be the two superpixels to be merged, $S_{1+2}$ the resulting superpixel, $e$ the edge segment between $S_1$ and $S_2$ (denoted as $S$ in the descriptions): (1a) size, (2a) solidity, (3a) extent, (4a) eccentricity, (5a) circularity, (6a) axes ratio, (7a) portion of the perimeter of $S$ touching the background, (8a) ratio between the length of $e$ and the perimeter of $S$, (9a) ratio between the length of $e$ and the minor axis of the ellipse with the same second-moments as $S$, (10a) maximum, (11a) minimum, (12a) mean intensity in $I_e(x, y)$ for pixels in $S$, (13a) SD of the intensity, (14a) intensity SD to mean ratio, (15-20a) $1^{st}$, $3^{rd}$, $5^{th}$, $10^{th}$, $50^{th}$ and $75^{th}$ intensity percentiles, (21a) maximum, (22a) minimum and (23a) mean intensity in the gradient map $GM$ (range $0 \rightarrow 1$) for pixels in
For the resulting superpixel \( S_{1+2} \), some features were computed as above (all except 8-9a), and others were added: (1b) feret ratio, (2b) maximum, (3b) minimum and (4b) mean distance from the centroid of \( S_{1+2} \) to boundary points, (5b) distance SD, (6b) distance SD to mean ratio, (7b) length of e, (8b) ratio between the number of pixels in the intersection between the edge map EM and e, and the length of e, (9b) ratio between the orientation of \( S_1 \) and \( S_2 \), (10b) ratio between the mean intensity value in \( GM \) (range \( 0 - 1 \)) for pixels in e and the mean intensity value in \( GM \) for pixels in \( S_1 \) and \( S_2 \).

The training dataset was built by processing the training images up to the marker-controlled watershed step. Pairs of adjacent superpixels in \( LB \) were then considered iteratively for merging. Specifically, two iterations were performed for each connected component of the \( FG \) map, typically corresponding to a single cell or a cluster of 2 to 10 cells. During each iteration, for each superpixel \( S_1 \) in the connected component, the adjacent superpixels \( S_2 \) were considered sequentially. For each candidate merge given by a pair \((S_1, S_2)\), \( 25 \times 2 + (25 - 2) + 10 = 83 \) features were extracted from \( S_1, S_2 \), and \( S_{1+2} \), as detailed in the previous two paragraphs, and inserted into the dataset, along with the respective class (1 if “to be merged”, 0 otherwise), set according to ground truth. If \( S_1 \) and \( S_2 \) were to be merged, \( S_1 \) was replaced by the merge \( S_{1+2} \) before continuing. The obtained dataset was used to train a random forest classifier.

Test images were treated with the same procedure as above, with the only difference that the class of a pair of adjacent superpixels was defined by the output of the classifier.

3.6. Post-processing

Hole-filling, morphological opening and reconstructions were applied to the revised \( LB \), followed by the removal of objects smaller than 70 pixels and 20 iterations of the Chan-Vese model for active contours [19] to refine the cell shapes according to \( I_1(x, y) \). Boundaries between touching cells were forced as defined in \( LB \). Finally, a second random forest classifier was trained on 28 features (1-6a, 10-25a, 1-6b) of candidate cells to filter out false positive findings.

4. RESULTS

On test images, predictions were matched to the ground truth masks at different thresholds of matching precision based on the standard intersection over union metric (IoU). We evaluated performance with the average precision metric (AP), derived from the number of true positives (TP), false positives (FP), and false negatives (FN) as \( AP = TP/(TP+FP+FN) \).

For comparison, we considered two state-of-the-art open-source solutions: Ilastik [20] and the generalist CellPose model [21]. Ilastik instance segmentation is attained by interactive training of a classifier to separate foreground from background, followed by hysteresis thresholding. CellPose is a deep learning-based method where a neural network is trained to predict the gradients of a topological map. These are followed via the gradient tracking process to route pixels toward the centres of the cells and group them accordingly. The generalist model is trained on over 70,000 objects.

As shown in Fig. 3, MR-NOM outperformed Ilastik and CellPose at all thresholds. In particular, the AP@0.5 was 0.7 for MR-NOM trained on 24 images (qualitative results in Fig. 2g), 0.51 for Ilastik, and 0.45 for CellPose. It can also be observed that MR-NOM provided satisfactory results when trained on a smaller dataset. The AP@0.5 was 0.69 and 0.68 for MR-NOM trained on 12 and 6 images, respectively (6159 and 3236 cells, respectively).

5. CONCLUSIONS

Few techniques have been designed for the instance segmentation of neuronal cells in Nissl-stained histological slices of the brain. We proposed a new segmentation method called MR-NOM, which exploits a multi-scale approach to deliberately over-segment the cells into superpixels to be merged via a classifier. MR-NOM dealt effectively with cells of varying characteristics that partially touch or overlap, even with a small training dataset. It was used in an active learning mode to aid the annotation process and will be exploited to segment WSIs of the auditory cortex of Tursiops truncatus. It is also expected to be adopted with suitable refinements (e.g., more annotations and deep learning-based marker definition) to process WSIs of different species for the characterization of brain cytoarchitecture in comparative neuroanatomy studies aimed in particular at a better understanding of neurodegenerative and neuroinflammatory disorders.
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