An Iterated Local Search Algorithm for Cell Nuclei Detection from Pap Smear Images

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Abstract: In this work, we propose an Iterated Local Search (ILS) approach to detect cervical cell nuclei from digitized Pap smear slides. The problem consists in finding the best values for the parameters to identify where the cell nuclei are located in the image. This is an important step in building a computational tool to help pathologists to identify cell alterations from Pap tests. Our approach is evaluated by using the ISBI Overlapping Cervical Cytology Image Segmentation Challenge (2014) database, which has 945 synthetic images and their respective ground truth. The precision achieved by the proposed heuristic approach is among the best ones in the literature; however, the recall still needs improvement.

1 INTRODUCTION

According to the World Health Organization (WHO), 14.1 million new cases of cancer are diagnosed every year worldwide. In addition, an estimate is that this is the cause of 13% of all deaths in the world per year, which corresponds to 8.2 million people. WHO also reports that, according to the scientists, a number of cancer cases increase 70% in the coming decades, reaching 21.4 million in 2032.

The Pap smear is a screening test performed to detect alterations in cervix cells. The British Medical Journal (BMJ) published a study showing that patients who discovered the disease through the Pap smear test had a survival rate of 92%, while those who discovered it through symptoms had a 66% rate. It happens due to late symptoms when the disease is already at a more advanced stage. However, the test allows the lesions to be detected early, even at the beginning of the disease, which eases treatment.

There are two ways to perform the Pap smear. In the first one, the cells are initially collected. For this, an instrument called speculum is introduced into the vagina. With a wooden spatula, the collector performs a cervix visual inspection intending to collect its internal and external surface. The harvested cells are placed on a slide, called a smear, and are sent for analysis to laboratories specialized in cytopathology. The second method is initiated analogously to the previous one, but before placing the cell samples on a slide, they are placed in a liquid medium which is treated to remove other elements, such as mucus and red blood cells, in order to increase the sensitivity and specificity of the examination. Although the liquid medium concentrates the cells in smaller areas, speeding the manual reading, this method is costly, which determines its less frequent employment.

A sample collection of the Pap test generates around 15,000 fields for analysis on a slide. Since this analysis is performed manually, the volume of data is extensive. Thus, difficulties such as professionals’ physical and mental fatigue arise. In addition, once the procedure requires great technical knowledge, the labor becomes more expensive, reducing the number of trained people capable to perform it. Thus, we pursue a way to help these professionals to reduce...
the number of false positives (cases in which the examination detects a false lesion) and false negatives (cases in which a cell lesion is not detected), once that this interferes with the physical and/or psychological health of a patient.

The first step to identify if a cell has malignant changes is the detection and segmentation of its nuclei since the morphological and texture characteristics of the nucleus present significant variations when they are altered. Cytopathologists identify the lesion by altering the nucleus / cytoplasm ratio, accompanied by changes in chromatin distribution, hyperchromasia, and nuclear membrane shape. On the other hand, computer scientists hypothesize that only the identification of nuclear irregularity, texture difference and hyperchromasia, or irregular chromatin condensation would be sufficient to identify a suspected cell. However, many researchers (Moshavegh et al., 2012; Samsudin et al., 2016) have been arguing that the detection and segmentation of their nuclei (Plissiti and Nikou, 2012b; Plissiti and Nikou, 2012a; Lorenzo-Ginori et al., 2013) is enough for lesion detection, since the morphological and texture characteristics of the nucleus present significant variations when altered. What may happen, for example, is an increase in the size of the nucleus, the irregularity of its nucleic acid form, texture difference, and hyperchromasia, or irregular chromatin condensation (Plissiti et al., 2011).

Cells and nuclei segmentation is a working progress enabling different approaches, from region-based segmentation to Convolutional Neural Network (CNN) (Song et al., 2014; Lee and Kim, 2016; Tareef et al., 2017; Araujo et al., 2018). One of those studies use the concept of superpixel (Song et al., 2014) as a clustering stage to generate the superpixels that were used to train a CNN. They have been used to classify what was background in the image, cytoplasm or nucleus. (Lee and Kim, 2016) also used superpixel for partitioning and cell-bound refinement to perform the detection. (Tareef et al., 2017) proposed a method based on local distinctive characteristics and guided shape deformation that are embedded and classified by a Support Vector Machine (SVM). They have used it to segment the image into nuclei, cell clusters, and background. In addition, they use a structure based on the theory of sparse coding (SC) and guided by representative characteristics of the form to construct the cytoplasm of each cell.

Another frequently used technique is the nucleus-detection one, called Maximally Stable Extremal Region (MSER) (Lu et al., 2015). (Nosrati and Hamarneh, 2014) used it combined with Random Forest (RF) classifier.

This work focuses on cervical cell nuclei segmentation obtained in Pap smears images. The objective is to maximize the number of true positive results found and to minimize the number of false positive results. In other words, the aim is to maximize the number of nuclei found and to minimize the number of nuclei found that do not exist in the image. In the proposed method, we used a superpixel clustering algorithm, simple linear iterative clustering (SLIC) to oversegment the image, the density-based spatial clustering of applications with noise (DBSCAN) to cluster generated superpixels, and an Iterated Local Search (ILS) heuristic to calibrate the parameters that are used to identify whether or not a superpixel is a nucleus.

The remainder of this paper is organized as follows. Section 2 introduces the database that is used in tests. Section 3 presents the preprocessing of database images. Section 4 details a heuristic approach, based on the Iterated Local Search metaheuristic, for solving the problem. Then, the results are discussed in Section 5 and the conclusions are exposed in Section 6.

2 DATABASE

The database used to implement the methods was the “Overlapping Cervical Cytology Image Segmentation Challenge”, proposed during a challenge at International Symposium on Biomedical Imaging (ISBI) in 2014. It contains 945 synthetic images that were generated from real images obtained from a Pap smears test. All images are $512 \times 512$, grayscale, with a different number of cells, ranging from one to ten, and varying overlapping levels of cells. These images are divided into two groups: 45 for training and 900 for testing. An example of these images is shown in Figure 1.

3 IMAGE PREPROCESSING

As shown in Figure 2, image preprocessing consists of two steps. In the first step, the initial image that was obtained in the database shown in Section 2 (see Figure 2(a)) is used as input of the SLIC algorithm - a segmentation superpixel algorithm proposed by (Kovesi, 2000), and in the second one, DBSCAN is used to cluster similar superpixels.

The idea of the SLIC algorithm is to generate superpixels by clustering pixels based on their color
similarity and proximity. The algorithm is based on
the k-means clustering approach (MacQueen, 1967; Duda et al., 2000), and considers a space of five
dimensions \([labxy]\), where \(l\), \(a\) and \(b\) are CIELAB
space values, and \(x\) and \(y\) are the coordinates of
the pixels. Moreover, a new distance is used to
considering the size of the superpixels, which will be
explained below.

Consider an image with \(N\) pixels and an input
parameter \(K\) estimating the desired amount of
superpixels. So, it is known that the size of each
superpixel generated by SLIC has \(N/K\) pixels. Besides
that, in order for all superpixels to have approximately
the same size, a center \(C_k = [l_k, a_k, b_k, x_k, y_k]^T\) is
placed at each distance of \(S = \sqrt{N/K}\) pixels.

Since the spatial extent of any superpixel is
approximately \(S^2\), it is known that the pixels
associated with the center are within an area \(2S \times 2S
around it in the xy plane, which is the grouping area.

The distance \(D_s\) can be described according to
Equation (1):

\[
\begin{align*}
d_{lab} & = \sqrt{(l_k-l)^2 - (a_k-a)^2 - (b_k-b)^2} \\
d_{xy} & = \sqrt{(x_k-x)^2 - (y_k-y)^2} \\
D_s & = \frac{m}{S^2}d_{xy}
\end{align*}
\]

where \(D_s\) is the sum of the distance \(lab\) and the
normalized distance \(xy\) by the interval \(S\). The variable
\(m\) is used to control the superpixel compaction.

With the defined distance, it is possible to realize
the generation of the superpixels, as shown in
Algorithm 1. The algorithm response is described in
Figure 2(b).

Once we obtained the superpixels segmented
image, we can move on to the second preprocessing
stage. It consists in grouping similar superpixels.
Figure 2: Preprocessing flow chart of images.

\[ s = \langle 0.50, 1.10, 70, 120, 600 \rangle \]

Figure 3: Example for a solution \( s \) of the problem.

4.2 Solution Evaluation

For each image \( I \), a resulting mask \( X \) is generated containing only the clusters within the intervals delimited by all the CIA parameters of solution \( s \). In this way, image \( I \) may have fewer nuclei than the actual ones.

The process of constructing a mask \( X \) is shown in Figure 4. As we can see, in the image \( I \) the are five clusters candidates to nuclei, but only two (in green color) have been identified as a nucleus. Thus, mask \( X \) generated has only these two clusters.

Figure 4: An example of the construction of a mask \( X \).

If mask \( X \) contains more than one nucleus, it is decomposed into new masks so that each one contains only one of its nucleus. For example, as shown in Figure 5, the two-nuclei mask is decomposed into two new masks \( X_1 \) and \( X_2 \), each one containing only one of its nucleus. The same procedure is carried out for each template \( Y \) in the ground truth image. It is noted that the template of an image \( I \) may contain a different number of nuclei of the mask of the same image \( I \).

Each mask \( X_i \) relative to a resulting image \( I \) is compared to all ground truth \( Y_j \) of that image under evaluation to determine the method’s assertiveness level.

To that end, the Similarity Coefficient Dice (Dice, 1945), also known as Sørensen-Dice Coefficient, is used. The coefficient is calculated by Equation (2), which is a statistical metric used to compare the similarity between two samples \( X_i \) and \( Y_j \):

\[
\text{Dice}(X_i, Y_j) = \frac{2|X_i \cap Y_j|}{|X_i| + |Y_j|} \tag{2}
\]

The coefficient value in Equation (2) gives in a real interval \((0, 1)\). We consider that a coefficient greater than 0.6 indicates that the similarity between them is greater than 60%. Consequently, the analyzed cluster is considered a nucleus. If there is a ground truth \( Y_j \) such as \( \text{Dice}(X_i, Y_j) \geq 0.6 \), then the procedure informs that it correctly detected that \( X_i \) is a nucleus (true positive). In different circumstances, it is said that the algorithm has mistakenly detected that \( X_i \) is a nucleus, that is, a false positive result is provided.

In Figure 5 we have a mask \( X \) with two nuclei decomposed into two masks \( X_1 \) and \( X_2 \), each with a single nucleus. Also, we have a ground truth \( Y \)
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with three nuclei relative to an image $I$. This ground truth was decomposed into three other templates - $Y_1$, $Y_2$ and $Y_3$, each with a single nucleus. By applying the Similarity Coefficient, it is observed that $X_1$ and $Y_1$ were considered equivalent (see the green arrow). Since the $\text{Dice}(X_1, Y_1)$ value was greater than 0.6, we can say that the nucleus in $X_1$ has been detected correctly. Also in the same example, since the mask $X_2$ was not considered equivalent with no ground truth $Y_j$, with $j = 1, 2, 3$ (see red arrows), we can affirm that the nucleus in $X_2$ was detected incorrectly.

Thus, in order to evaluate the solution $s$, we have maximized the function $F_1$ (Manning and Schütze, 1999), given by Equation (3), which corresponds to the harmonic mean between precision and recall metrics:

$$F_1(s) = 2 \times \frac{\text{prec}(s) \times \text{rec}(s)}{\text{prec}(s) + \text{rec}(s)} \quad (3)$$

where precision and recall of the solution $s$ are calculated according to Equations (4) and (5), respectively:

$$\text{prec}(s) = \frac{\sum_{I \in \text{DataBase}} TP(I, s)}{\sum_{I \in \text{DataBase}} [TP(I, s) + FP(I, s)]} \quad (4)$$

$$\text{rec}(s) = \frac{\sum_{I \in \text{DataBase}} TP(I, s)}{\sum_{I \in \text{DataBase}} [TP(I, s) + FN(I, s)]} \quad (5)$$

In these equations, $TP(I, s)$, $FP(I, s)$ and $FN(I, s)$ represent the number of true positive, false positive and false negative results in each database image $I$ detected by applying Equation (2) to all masks $X_j$ related to the decomposed image $I$ that were generated from solution $s$.

Therefore, we want to maximize both precision and recall. Consequently, we are looking for maximizing the number of true positive results (TP) and minimizing the number of false positive (FP) and false negative (FN) results.

### 4.3 Initial Solution and Neighborhood Structure

A solution to the problem is obtained by randomly choosing the CIA parameter values, following the limits defined in Table 1. First, the algorithm constructs 10 solutions but only the best is chosen as the initial one.

In order to explore the solution space, we define a simple neighborhood structure that consists in increasing or decreasing a chosen position of the solution according to a step $r$. For the first two parameters, the step value is set to $r = 0.01$ units, whereas for the last three parameters, the step value varies by up to $r = 5$ units. In both, the step value can be either positive or negative, respecting the limits established in Table 1. Therefore, a solution $s$ has $N_{\text{Neigh}} = 6 \times r + 4$ neighbor solutions.

Figure 6 shows an example where the fourth position of the solution $s$ was chosen to be decremented in 3 units, generating a neighbor $s'$ of the solution $s$.

$$s = (0.50, 1.10, 70, 120, 600)$$

$$s' = (0.50, 1.10, 70, 117, 600)$$

This neighborhood was the only one used because by applying this move it is possible to explore all the problem-solution space using local search methods.

### 4.4 Iterated Local Search

The Iterated Local Search method – ILS (Stützle, 1998; Lourenço et al., 2010) explores the solution space through perturbations in local optima. These perturbations must be strong enough to prevent the algorithm from being trapped in a local optimum and thus explore different solutions, but weak enough to prevent random restarts.

As shown in Algorithm 2, the perturbation consists in applying $p + 1$ consecutive moves according to Section 4.3, where $p$ corresponds to the perturbation level to be performed. Algorithm 3 shows the pseudo-code of the ILS method. It starts with an initial solution (line 1) and applies a local search in it (line 2). In order to avoid getting stuck in this solution (which is possibly a local optimum), the current solution $s$ is disturbed, generating an intermediate solution $s'$ and a new local search is made (lines 8 and 9, respectively). If the solution $s''$ resulting from this local search is better than
According to function $F_1$, given by Equation (3), then $s^{**}$ becomes the new current solution and the perturbation level is restarted (lines 10-13); otherwise, the perturbation level is incremented (line 15). As previously explained, the perturbation level represents the intensity of the perturbation that will be performed. This whole process is repeated until the maximum number of iterations ($ILS_{Max}$) is reached without an improvement in the current solution.

Algorithm 3: Iterated Local Search.

```
Input: $F_1(.)$, $ILS_{Max}$

1. $s_0 \leftarrow$ Initial Solution
2. $s \leftarrow LocalSearch(s_0)$
3. $iter \leftarrow 0$
4. $bestIter \leftarrow iter$
5. $p \leftarrow 1$
6. while $iter - bestIter < ILS_{Max}$ do
7.     $iter \leftarrow iter + 1$
8.     $s' \leftarrow Perturbation(s, p)$
9.     $s'' \leftarrow LocalSearch(s')$
10.    if $F_1(s'') > F_1(s)$ then
11.        $s \leftarrow s''$
12.        $bestIter \leftarrow iter$
13.        $p \leftarrow 1$
14.    else
15.        $p \leftarrow p + 1$
16. end
17. return $s$
```

The local search used in Algorithm 3, lines 2 and 9, is a Random Descent. It works as follows:

**Step 1:** A random neighbor of the current solution is generated;

**Step 2:** If this neighbor is better than the current solution, then it becomes the new current solution.
If not, this neighbor is discarded and another random neighbor is generated, returning to Step 1;

**Step 3:** The process is repeated until a maximum number of consecutive neighbors without improvement ($RD_{Max}$) in the current solution is reached, where $RD_{Max} = pRD \times NNeigh$, $pRD \in (0, 1)$, and $NNeigh$ is the maximum number of neighbors in the current solution.

5 COMPUTATIONAL EXPERIMENTS

The proposed preprocessing method (Section 3) was performed in MATLAB language, whereas the proposed algorithm (Section 4) was developed in Python. All experiments were performed on an Intel Core i7-8700 processor with a 3.2GHz processor, with 16GB RAM, and running on Windows 10 at 64-bits.

During the preprocessing stage, it was necessary to define SLIC and DBSCAN parameters. These values were obtained empirically using the training database. To explain the parameter calibration that was made it was necessary to zoom in the image (see Figure 7) because it has a lot of details to be observed. In Figure 7, the red square shows the zoomed image area used to explain SLIC parameters (Figures 8, 9 and 10), whereas the green one shows zoomed image area for DBSCAN (Figure 11).

![Figure 7: Zoomed image areas for description of SLIC (in red) and DBSCAN (in green) parameters.](image)

Figure 8: Number of desired superpixels in SLIC.

(a) 500  (b) 2000  (c) 3500

![Figure 8: Number of desired superpixels in SLIC.](image)

The second SLIC parameter calibrated was the weighting factor between color and spatial differences. (Kovesi, 2000) suggests values from 5 to
As shown in Figure 9, if we increase too much the parameter it does not bring good results, once the clusters that have a nucleus part, also have others noise parts of the image too. So, after some tests, the best value found for this parameter was 11.

Finally, the last SLIC parameter calibrated was the cluster size threshold. In other words, regions morphologically smaller than the threshold are merged into an adjacent cluster. (Kovesi, 2000) suggests a value of 1 or 1.5 and 0 to disabled this merge. The results of applying this parameter are shown in Figure 10. As we can see below, the difference between them is not relevant. But, as a nucleus can not be so small, we use 1.5 as the cluster size threshold just to exclude the small ones.

Following, the only DBSCAN calibrated parameter was the threshold that controls which superpixels should be clustered. (Kovesi, 2000) suggests values from 5 to 10. As shown in Figure 11, a bigger threshold returns less clusters, without decreasing the quality of clusters found. In this way, a higher threshold can reduce the processing time of the heuristic approach to analyze the clusters that are nuclei candidates. So, the value chosen for the threshold was 10, completing the calibration of the preprocessing parameters.

As stated in Section 2, the database used provides 45 images for training and 900 for testing. The training images were used to estimate the intervals for each of the five parameters described in Table 1 so that the allowed limits correspond to actual values found in cells.

After executing the proposed algorithm for the solution $s$, we used the precision and recall measures, given by Equations (4) and (5), to determine the quality of the nuclei detection. Table 2 presents the precision and recall values obtained from the literature methods and the proposed heuristic approach.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Minimum</th>
<th>Maximum</th>
</tr>
</thead>
<tbody>
<tr>
<td>Minimum Circularity</td>
<td>0.48</td>
<td>0.63</td>
</tr>
<tr>
<td>Maximum Circularity</td>
<td>1.08</td>
<td>1.23</td>
</tr>
<tr>
<td>Minimum Intensity</td>
<td>57</td>
<td>158</td>
</tr>
<tr>
<td>Minimum Area</td>
<td>114</td>
<td>213</td>
</tr>
<tr>
<td>Maximum Area</td>
<td>511</td>
<td>611</td>
</tr>
</tbody>
</table>

Table 2: Results for nuclei detection.

<table>
<thead>
<tr>
<th>Method</th>
<th>Precision</th>
<th>Recall</th>
</tr>
</thead>
<tbody>
<tr>
<td>(Nosrati and Hamarneh, 2014)</td>
<td>0.903</td>
<td>0.893</td>
</tr>
<tr>
<td>(Lu et al., 2015)</td>
<td>0.977</td>
<td>0.883</td>
</tr>
<tr>
<td>(Ushizima et al., 2014)</td>
<td>0.959</td>
<td>0.895</td>
</tr>
<tr>
<td>(Saha et al., 2016)</td>
<td>0.918</td>
<td>0.915</td>
</tr>
<tr>
<td>(Tareef et al., 2017)</td>
<td>0.990</td>
<td>0.940</td>
</tr>
<tr>
<td>(Braz and Lotufo, 2017)</td>
<td>0.929</td>
<td>0.917</td>
</tr>
<tr>
<td>Our method with ILS</td>
<td>0.985</td>
<td>0.879</td>
</tr>
</tbody>
</table>

As can be seen in Table 2, the proposed algorithm was able to generate a solution with the second best measure of precision. However, the recall was lower than all methods to which it was compared.
6 CONCLUSIONS

This paper introduces an ILS-based algorithm to detect cell nuclei from cervical cell images. Each image was analyzed with respect to the values of circularity (minimum and maximum), intensity (minimum) and area (minimum and maximum) parameters. The main purpose is to simulate cytopathologists analysis, since the pap smear test uses morphological and chromatin distribution in nucleus to detect anomalies.

The proposed algorithm produced adequate results, according to the precision standards proposed in the literature, and when compared to other algorithms; in fact, the ILS-based algorithm showed the second best measure of precision. However, its performance regarding recall was not satisfactory. It is known that the recall is related to the number of nuclei that the algorithm failed to find. Therefore, it is important that the recall of Pap smears tests are as high as possible since failing to detect a lesion might influence prognosis. On the other hand, as a computer cannot diagnose, then the images should be analyzed later by a pathologist. Thus, the method is not required to offer perfect precision, that is, all clusters detected as nuclei are nuclei, indeed.

In this way, the study on the influence of other parameters and the reasons why these nuclei were not found are considered future work aiming to improve the recall while maintaining high precision.

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