

Multiscale Convolutional Neural Networks for Vision-Based Classification of Cells

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Abstract. We present a Multiscale Convolutional Neural Network (MCNN) approach for vision-based classification of cells. Based on several deep Convolutional Neural Networks (CNN) acting at different resolutions, the proposed architecture avoid the classical handcrafted features extraction step, by processing features extraction and classification as a whole. The proposed approach gives better classification rates than classical state-of-the-art methods allowing a safer Computer-Aided Diagnosis of pleural cancer.

1 Introduction

Computer-Aided Diagnosis (CAD) through image processing [1] [2] is a domain of ever increasing interest. Indeed, automatic salient feature extraction by a computer can save time to a pathologist, and with the ever-increasing amount of virtual biomedical data, CAD has now become a crucial point for enabling safer and faster diagnosis.

Our work deals with cytological pleural cancer cells analysis. Body fluids obtained by needle aspiration can be examined as liquid-based preparations or cytospin preparations under a microscope. The aim of our study is to provide pathologists with an automatic classification of cells. This will help them to distinguish abnormal and dystrophic mesothelial cells that are very important in the early diagnosis of cancers due to asbestos [3].

We deal with preparations under Feulgen stain where chromatin takes a stoichiometric pink color while cytoplasm is not colored or few. In conjunction with other preparations, Feulgen stain can provide useful information to the cytopathologist for establishing a diagnosis. Classification of cells under Feugen stain has already been studied in [4]. Our work proposes to deal with whole virtual slides scanned with a Leica SCN400 scanner 40× resolution, rendering at full resolution images of size 80000 × 90000.

Fluids contain several types of cells that may take different chromatin spatial arrangements [3]. The aim of an automatic processing tool is to classify cells contained in a virtual slide into predefined classes: malignant cells, dystrophic cells, normal mesothelials, macrophages and all the inflammatory cells such as polynuclears or lymphocytes. Indeed, the analysis of chromatin distribution over a cell is critical to distinguish between malignant, dystrophic and normal mesothelials.

A fully automatic virtual slides processing scheme is typically decomposed into 3 steps:

- Segmentation and extraction of the cells contained in a virtual slide,
- Feature extraction from the detected cells,
- and Classification of the cells according to the extracted features.

In this paper, we propose a vision-based method for the classification of cells. The last two steps are merged by the use of deep neural networks that performs features extraction and classification as a whole. Moreover, a multiscale approach is proposed that uses different networks at each scale, and fuse their outputs.

The rest of the paper is organized as follows: Section 2 describes the proposed Multiscale Convolutional Neural Network (MCNN) approach, while experiments and results are detailed in Section 3. We discuss the proposed approach and the results in Section 4 and draw some conclusions in Section 5.

2 Multiscale CNN

Convolutional Neural Networks (CNNs) are multi-layered neural networks that are specialized in pattern recognition tasks [5], [6]. They are well-known for robustness to small inputs variations, minimal pre-processing and do not require any specific feature extractor choice. The proposed architecture relies on several deep neural networks that alternate convolutional and pooling layers. These deep neural networks belong to a wide class of models generally termed *Multi-stage architectures* reminiscent Hubel and Wiesel's 1962 work on the cat's primary visual cortex [7]. This architecture of convolution interlaced with pooling layers is dedicated to the automatic feature extraction. The final classification is performed by some classical fully connected layers stacked on top.

2.1 Convolution Layer

A convolution layer C^i (layer i of the network) is parametrized by its number N of convolution maps M_j^i ($j \in \{1, \dots, N\}$), the kernels size $K_x \times K_y$ (often squares) and the connection scheme to previous layer L^{i-1} . Each map M_j^i is the result of a sum of convolution of previous layer's maps M_j^{i-1} by their respective kernel. A bias b_j^i is added and the result is passed through a non-linear squashing function $\phi(x) = 1.7159 \tanh(\frac{2}{3}x)$ [6]. In the case of a full connected convolution map, the result is computed as

$$M_j^i = \phi \left(b_j^i + \sum_{n=1}^N M_n^{i-1} * K_n^i \right)$$

2.2 Max-Pooling Layer

In classical CNNs, convolution layers are followed by a subsampling layer. This layer reduces the effective maps size, and introduces some invariance to distorted

or shifted inputs. A max–pooling layer is a variant which has shown some merit in the literature [8]. The output of a max–pooling layer is given by the maximum activation over non–overlapping regions of size $K_x \times K_y$, instead of averaging the inputs as in a classical subsampling layer. A bias is added to the resulting pooling and the output map is passed through the squashing function ϕ defined above.

2.3 Classification Layer

Parameters of convolution and pooling layers are chosen such that output maps of the last convolutional layer are downsampled to 1 pixel per map, resulting in a 1D vector of attributes. Classical feed–forward fully connected layers are then added to perform the classification. The last layer, in the case of supervised learning, contains as many neurons as the number of classes. In our work, this last layer contains 6 neurons, and a softmax activation function is used to turn outputs into probabilities.

2.4 Multiscale CNN (MCNN)

It is well admitted that human vision is a multiscale process. In this work, we create N CNNs with different retina sizes. A given input pattern is rescaled N times to fit the retina of the CNNs. The question of optimizing the outputs of classifiers is a recurrent question in the pattern recognition field. For handwritten digits, it was shown [9] that a simple average gives better classification results than a linear combination whose weights learned over a cross–validation set [10]. However, in [9] there was no reason to weight the classifiers since they act at the same resolution and only differ from the distortions applied to the training set. In our case, a reasonable hypothesis is that the classifier at the lowest resolution may be less salient than the one at full resolution. Final outputs are then computed as a linear combination of the outputs of the N CNNs. The overall architecture of the MCNN is shown in Figure 3.

3 Experiments

In this section, we detail the conducted experiments. In order to compare the performances of our approach to other classification schemes (where features are mainly handcrafted), we first proceed to a manual segmentation of the cells.

3.1 Data Acquisition and Segmentation

We manually acquire a database of annotated cells (samples shown at the first row of Figure 1) composed of :

- 215 *abnormal mesothelials* (noted C_1 in the rest of the paper),
- 209 *dystrophic mesothelials* (C_2),

- 201 *normal mesothelials* (C_3),
- 195 *macrophages* (C_4),
- 198 *polynuclears* (C_5),
- and 196 *lymphocytes* (C_6).

This database (and the segmentation of each cell) is made available for research purposes¹. It contains some difficulties and maybe some annotation errors. Figure 6 shows some difficult cases where cells belonging to different classes look similar.

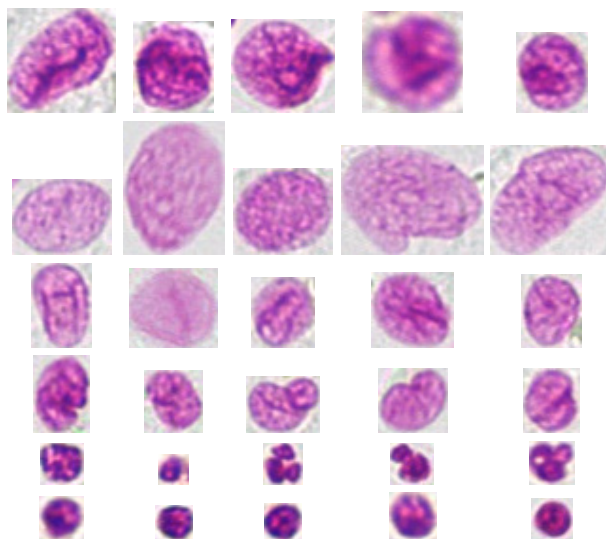


Fig. 1. Samples of the database: First row: abnormal mesothelials (C_1), second row: dystrophic mesothelials (C_2), third row: normal mesothelials (C_3), fourth row: macrophages (C_4), fifth row: polynuclear (C_5), and sixth row: lymphocyte (C_6)

A manual segmentation of the cells of the database is performed that separates foreground pixels belonging to the cell and background pixels. The original color images vary in size and are not necessarily square. After a visual inspection of the database, we retain the size of 80×80 as a base size for the inputs. The images are padded with background pixels equally distributed on each side, such that the cell appears in the center of the image. Background pixels are then set to 127.5 to ensure they will be close to 0 in the further preprocessing step.

3.2 Networks Architectures

We construct 4 different CNNs that act at different resolutions. The size of the images at full resolution is 80×80 . This size is successively divided by a factor

¹ <http://www.greyc.ensicaen.fr/~pbyssen/feulgenodb.html>

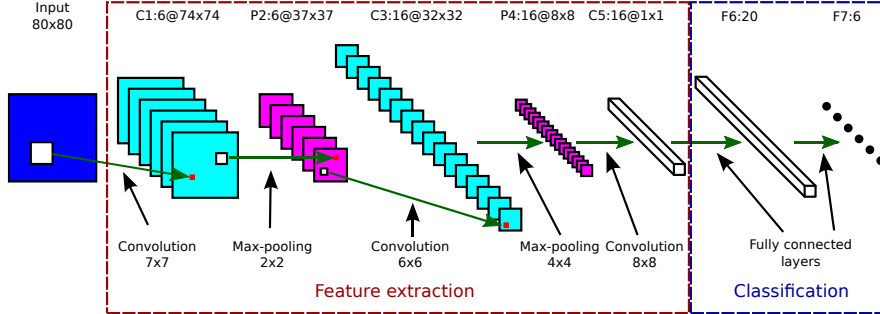


Fig. 2. Architecture of the CNN₈₀

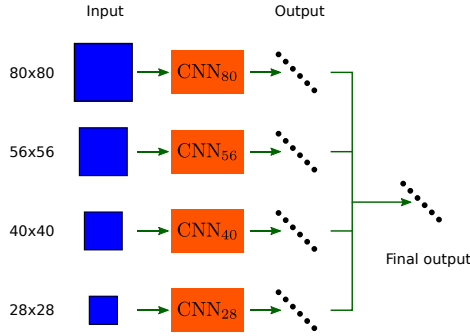


Fig. 3. Overall architecture of the proposed MCNN approach

$\sqrt{2}$, $\sqrt{2}^2$ and $\sqrt{2}^3$ along each dimension giving images of size 56×56 , 40×40 and 28×28 respectively. The constructed CNNs, named CNN₈₀, CNN₅₆, CNN₄₀ and CNN₂₈, are built in a similar way. They differ essentially from the size of their respective retinas and convolution masks. The detailed architecture of the full resolution CNN is shown in Figure 2. The architectures of the other CNNs are summarized in Table 1.

3.3 Existing State-of-the-Art Approaches

Classical approaches in the literature for the classification of cells rely on the extraction of shape, photometric and texture features from cells. Shape features include *surface*, *perimeter*, *compactness* or *stretching* [11]. Photometric features include *mean* and *standard deviation* of the color according to a chosen spectrum. *Integral Optical Density* (IOD) is also an important feature designed to reflect specific visual criteria usually used by cytopathologists [11]. It is computed as:

$$IOD = \sum_{x,y} OD(x,y) \quad \forall x,y \in \text{Cell}$$

Table 1. CNNs architectures. In each cell: Top: kernel size, Bottom: map size.

	CNN ₈₀	CNN ₅₆	CNN ₄₀	CNN ₂₈
Retina size	80 × 80	56 × 56	40 × 40	28 × 28
C1	7 × 7 74 × 74	7 × 7 50 × 50	5 × 5 36 × 36	5 × 5 24 × 24
P2	2 × 2 37 × 37	2 × 2 25 × 25	2 × 2 18 × 18	2 × 2 12 × 12
C3	6 × 6 32 × 32	6 × 6 20 × 20	5 × 5 14 × 14	5 × 5 8 × 8
P4	4 × 4 8 × 8	4 × 4 5 × 5	2 × 2 7 × 7	2 × 2 4 × 4
C5	8 × 8 1 × 1	5 × 5 1 × 1	7 × 7 1 × 1	4 × 4 1 × 1

where OD denotes the Optical Density defined as

$$OD(x, y) = -\log\left(\frac{I(x, y)}{I_0}\right)$$

with I_0 the mean background color value, and (x, y) belonging to the cell. Finally, texture analysis aims to extract important texture features, especially features concerning the chromatin distribution contained into the cell, which indicates a possible abnormal cell. Such analysis includes morphologic features [4], wavelet features [1], shape and photometric features computed on sub-regions [11], and hierarchical texture models [12] based on textons [13].

For comparison purposes with the state-of-the-art, we implemented most of these features, and extracted them from all the cells of the database at full resolution. Some of these features are computed on the whole cell, while others are computed on parts of it. Some auxiliary images used to compute these features are shown at Figure 4. We especially compute some photometric features on images of partitions (Figures 4(c) and 4(f)) to reflect the chromatin distribution within the cell: for an abnormal mesothelial, chromatin indeed tends to concentrate near the border of the cell. Cells have then been divided into concentric rings (Figure 4(c)) or into graph-based geodesic regions (Figure 4(f)) via a Region Adjacency Graph (RAG, Figure 4(e)) computed according to a watersheded version of the cell (Figure 4(d)).

A ten-fold cross validation is then processed on these features via a SVM carefully tuned with a Gaussian kernel. The final classification error rate obtained with this method is shown in Table 2.

A similar ten-fold cross validation is also processed via a simple feed-forward neural network (NN), containing 103 inputs (the size of a feature vector), 80 neurons on the hidden layer, and 6 neurons as outputs (one per class). The classification error rate are reported in Table 2.

Finally, these features are also classified via a Multi-Kernel SVM (MKL) approach [14]. This classification scheme aims at simultaneously learning a kernel

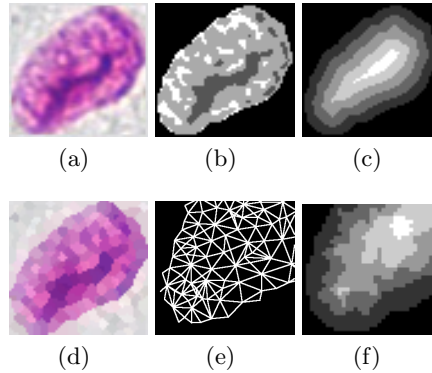


Fig. 4. (a) Extracted cell, (b) textons image, (c) concentric rings image, (d) partition image, (e) part of the computed RAG, and (f) part of the graph-based concentric partition image

per feature (or group of features) and weighting them. The MKL classification error rates are also reported in Table 2.

3.4 Classification Results

Prior to the training of the CNNs, the images are preprocessed (First row at Figure 5). Since the complementary color of the pink colored cells is green, only this channel is kept. Pixel values are then normalized such that they lie in the range $[-1, 1]$ (background pixel values are then close to 0). Images are then resized to fit the CNNs retina sizes.

A ten-fold cross validation is processed to test the MCNN. Since the classification has to be rotation invariant, the training set is augmented with rotated images with a angle step of 10 degrees (second row at Figure 5). The training sets are then composed of about 39000 images. Initial weights of the networks are initialized at random with small values, and we apply a stochastic learning with an annealing learning step and second order methods to speed up the training. Training ends when the error on a small subset of the training set does not decrease anymore (usually after 50 epochs). Classification results for each CNN are shown in Table 2. The error rates for the CNNs decrease while the resolution of the inputs increase since they are able to capture more information. We tested also two merging scheme of the outputs of the CNNs:

- The simplest one (MCNN in Table 2) averages the outputs of each CNN;
- A weighted fusion scheme (w -MCNN in Table 2).

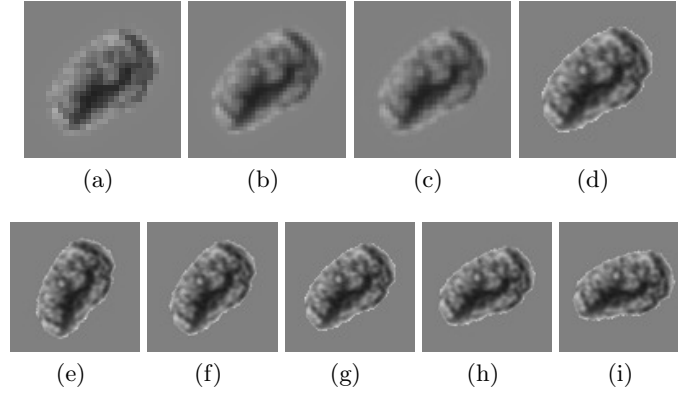


Fig. 5. First row: An abnormal mesothelial at increasing resolution. Second row: artificial rotations of a cell.

Table 2. Evaluation of the different methods (Standard deviation in parenthesis)

<i>Method</i>	<i>Error rate</i>
CNN ₂₈	9.04% (1.07)
CNN ₄₀	8.02% (1.02)
CNN ₅₆	8.01% (0.98)
NN	7.85% (0.98)
SVM	7.75% (0.95)
CNN ₈₀	7.55% (0.89)
MV /w CNN ₂₈	6.13% (0.84)
MKL	6.12% (0.83)
MCNN	6.02% (0.82)
_w MCNN	5.90% (0.80)
<u>_wMCNN /w CNN₂₈</u>	<u>5.74% (0.79)</u>

For this last scheme, the weights are computed for each CNN as:

$$w(CNN_x) = \frac{mean(cr_x)}{std(cr_x)}$$

where cr_x is the classification rate of the CNN _{x} . This weighting gives a low weight to a CNN which has a low classification rate and/or a big standard deviation of its results. The weights indeed increases with the resolution giving more saliency to the highest resolution. A majority voting scheme (MV) has also been tested and error rate is reported in Table 2. The best combination is found without considering results of CNN₂₈. The lowest error rate has finally been found with the second weighting scheme (see above) without considering the results of CNN₂₈. The fusion of CNNs outputs (with or without CNN₂₈) gives an error rate that is lower or equal to the state-of-the-art considered approaches.

Table 3 shows the confusion matrix for the best approach (w MCNN /w CNN₂₈). One can see that misclassified cells are mainly confused with their surrounding classes (non zeros values of the confusion matrix close to its diagonal). The good classification rate of the *abnormal mesothelials* (class C_1), that are the most important for establishing a diagnosis of pleural cancer, are encouraging. The worst classification rate is for the *normal mesothelials* (class C_3). Misclassified examples are mainly detected as *dystrophic mesothelials* (C_2) or *macrophages* (C_4). Differences between these three classes may indeed be very mild, see Figure 6 for some examples of such difficult cases.

4 Discussion

This approach avoid the design and/or the extraction of handcrafted features. Classical approaches are typically confronted to difficult tasks such as:

- The design of robust and easy-to-compute features,
- The evaluation of these features for their class separability capacities,
- and the more general problem of features selection.

Our Convolutional Neural Network approach bypasses these issues by avoiding the step related to the features.

Table 3. Confusion matrix of the classification (in percentage) for the best method. In row: real class, in column: found class.

	C_1	C_2	C_3	C_4	C_5	C_6
C_1	97.41	0.90	0.77	0.90	0	0
C_2	2.39	93.75	3.85	0	0	0
C_3	1.24	12.58	80.22	5.94	0	0
C_4	0.14	0	2.56	97.15	0.14	0
C_5	0	0	0	0.42	99.14	0.42
C_6	0	0	0	0.28	1.82	97.89

The approach of a committee of Convolutional Neural Networks has already been studied in [9]. The basic idea is to have multiple CNN that learn different features from the same dataset. To ensure such a learning, authors of [9] applied different distortions to the training set for each CNN. We cannot use this approach in this work, since applying distortions to cells may change their size or their chromatin distribution, which is not acceptable for the classification. A simple way to have more information from a simple cell is then to apply some multiresolution transformations and to build CNNs accordingly.

A not-so-easy problem is also the need of a robust segmentation tool, since handcrafted features are mostly computed on a masked version of the cell. For example, if a cell is poorly segmented and its mask contains some pixels belonging to the background, the computed surface will be higher. In this case, a simple

normal mesothelial could be confused with a dystrophic mesothelial or worse with an abnormal mesothelial. The segmentation has also to be fast since a virtual slide at full resolution can contain hundreds of thousands cells. For the purposes of this work, a relatively small number of cells have been manually segmented by a cytopathologist. A further work will be to extend this MCNN approach to cells that have not been segmented so as to avoid the segmentation step and replace it by a simpler cell detection module.

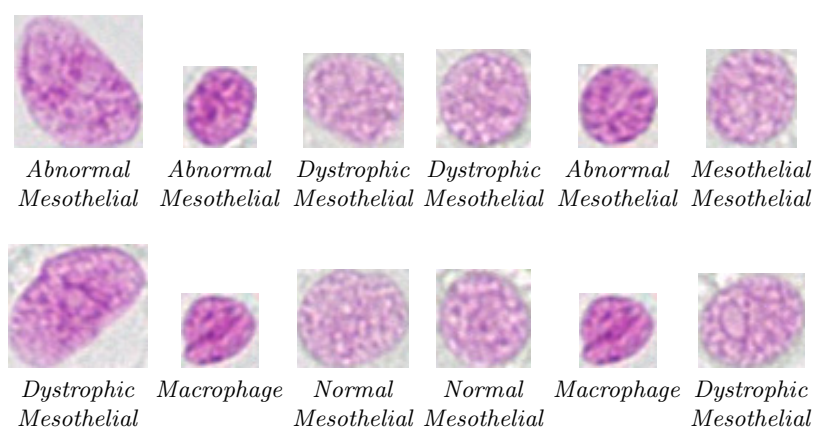


Fig. 6. In columns, some visually similar cells belonging to different classes. Class name below each cell.

5 Conclusion

This paper presents a Multiscale Convolutional Neural Networks (MCNN) approach for a vision-based classification of cells. Relying on several CNNs acting at different resolution that do not need any handcrafted features, the proposed architecture achieves better classification rates than classical state-of-the-art approaches.

Further work will involve a vision-based cell detection module in order to avoid the segmentation step. Moreover, an implementation of MCNN on a Graphics Processing Unit (GPU) will fasten the whole process in order to approach a real-time diagnosis.

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