Multi-crop Convolutional Neural Networks for lung nodule malignancy suspiciousness classification

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\section{Abstract}

We investigate the problem of lung nodule malignancy suspiciousness (the likelihood of nodule malignancy) classification using thoracic Computed Tomography (CT) images. Unlike traditional studies primarily relying on cautious nodule segmentation and time-consuming feature extraction, we tackle a more challenging task on directly modeling raw nodule patches and building an end-to-end machine-learning architecture for classifying lung nodule malignancy suspiciousness. We present a Multi-crop Convolutional Neural Network (MC-CNN) to automatically extract nodule salient information by employing a novel multi-crop pooling strategy which crops different regions from convolutional feature maps and then applies max-pooling different times. Extensive experimental results show that the proposed method not only achieves state-of-the-art nodule suspiciousness classification performance, but also effectively characterizes nodule semantic attributes (subtlety and margin) and nodule diameter which are potentially helpful in modeling nodule malignancy.

\section{Introduction}

Lung cancer is an aggressive disease carrying a dismal prognosis with a 5-year survival rate at 18\% [1]. Despite the development of multi-modality treatments over the past decade, lung cancer remains the leading death of cancer and accounts for approximately 27\% of all cancer deaths [2]. Technological advances in Computed Tomography (CT) have been routinely used in lung cancer detection, risk assessment, and clinical management. In particular, the increasing quantity of CT image assays has created a unique avenue for data-driven analysis to capture underlying cancer characteristics at a macroscopic level, allowing identification of prognostic imaging biomarkers [3].

In this study, we investigate the problem of automatic lung nodule malignancy suspiciousness classification using CT imaging data. The annotation of nodule malignancy suspiciousness has permitted a chance to evaluate consensus assessments from different experienced radiologists. Specifically, the automatic classification of malignancy suspiciousness on CT studies is a worthy task, because it would facilitate radiologists to assess early risk factors which is essential in lung cancer research [4,5]. A typical implication of such analysis is to provide useful cues for subsequent therapeutic plannings and holds promise for improving individualized patient management. For example, distinct malignancy likelihood derived from imaging can be used to recommend follow-up treatments including CT surveillance (e.g. low likelihood score) or biopsy test and surgical resection (e.g. high likelihood score) [6]. Despite different approaches were proposed for lung nodule diagnosis, novel data-driven techniques are required to advance the predictive power with CT imaging, especially for the prediction on malignancy suspiciousness.

Image-based techniques for analyzing lesions are normally performed with detection [7,8], segmentation [9–12], hand-crafted feature engineering [13,14], and category labelling [15–18]. Zinovev et al. [19] adopted a belief decision tree approach to predict nodule semantic attributes. Chen et al. [20] proposed to use a neural network ensemble scheme to distinguish probably benign, uncertain and probably malignant lung nodules. Han et al. [16] used a 3-D image-based texture feature analysis for nodule diagnosis. More recently, Balagurunathan et al. [14] and Aerts et al. [13] extracted a number of nodule image features to investigate their prognostic
power. Related studies on the Lung Image Database Consortium and Image Database Resource Initiative (LIDC-IDRI) dataset [21] are shown in Table 1. However, all these methods rely on nodule segmentation as a prerequisite. Notably, automatic nodule segmentation may affect classification results since methods such as region growing and level set typically depend on initialization. Working on these segmented regions may yield inaccurate features that lead to erroneous outputs. To derive a suspiciousness-sensitive descriptor in CT imaging, we need to overcome at least two major obstacles: the difficulty of nodule delineation caused by a large range of nodule morphology variation, and the challenge posed by the nodule radiological heterogeneity for computational models to capture qualitative characteristics.

Image patch-based approaches provide an alternative way for the region of interest (ROI) definition [22,23]. Researchers are seeking visual feature descriptors, such as Local Binary Patterns (LBP) [24] and Histogram of Oriented Gradients (HOG) [25], to refine measurement on lung cancer imaging. Nevertheless, the yielded textural features are largely determined by the parameter setting. Thus, using them to accurately describe the variability of lung nodules is difficult. In response to these challenges, we utilize the Convolutional Neural Network (CNN) [26–28] to build an end-to-end computational architecture which is robust in lung nodule image feature extraction and malignancy suspiciousness classification. We propose a computational architecture—the Multi-crop Convolutional Neural Network (MC-CNN)—to learn high-level suspiciousness-specific features for lung nodule classification. As outlined in Fig. 1, our approach automatically classifies nodule malignancy suspiciousness by extracting a set of highly compact features. It is an end-to-end architecture which embeds nodule feature extraction into a hierarchical network. The proposed method simplifies conventional lung nodule malignancy suspiciousness classification by removing nodule segmentation and hand-crafted feature (e.g., texture and shape compactness) engineering work. Our main contributions can be summarized as follows:

1. We demonstrate that even without nodule segmentation and hand-crafted feature engineering which are time-consuming and subjective, the high-level features extracted by our MC-CNN from the detected nodule patches are able to present high inter-class variations related to nodule malignancy suspiciousness (Fig. 2), bridging the gap between the raw nodule image and the malignancy suspiciousness.

2. We propose a multi-crop pooling operation which is a specialized pooling strategy for producing multi-scale features to surrogate the conventional max-pooling operation. Without using multiple networks to produce multi-scale features, the proposed approach applying on a single network is effective in computational complexity (Section 4.2).

3. Beyond nodule malignancy suspiciousness classification, we extend the proposed approach to quantify nodule semantic labels as well as to estimate nodule diameter that may potentially assist researchers in evaluating malignancy uncertainty (Section 4.5). Our results showed the possible applications of the proposed method in other lung nodule-relevant analysis that may potentially assist researchers in evaluating malignancy uncertainty.

Applying a supervised learning scheme in deep feature extraction, our approach is in contrast with an auto-encoder approach [30] that applied an unsupervised learning method without prior labeling information. The proposed method also differs from our previous work based on the multi-scale CNN model [31] which utilized multiple CNNs in parallel with different scales of nodule images. In [31], a resampling strategy was used to uniformly represent nodule patches. However, multiple networks become the main burden for training CNNs efficiently since they involve more computational costs, especially when dealing with high-resolution images. As opposed to the design of multiple CNNs [31], the proposed model simplified the training process by replacing multiple CNNs with the multi-crop pooling architecture that is specially tailored to lung nodule malignancy suspiciousness classification. Furthermore, our model underscored the knowledge extraction from feature space rather than image space. In other words, the computation is specified on the intermediate convolutional features (i.e., feature space), rather than different scales of raw input signals (i.e., image space).

The rest of the paper is organized as follows. Section 2 introduces the proposed multi-crop CNN architecture. Section 3 presents the detail of the dataset and data augmentation. Section 4 describes the experimental setup and results. Section 5 is the discussion and Section 6 concludes the paper.

Table 1 Some classification results on LIDC-IDRI dataset from literatures. “NA” denotes “nodule attributes” and “MS” denotes “malignancy suspiciousness”.

<table>
<thead>
<tr>
<th>Related work</th>
<th>Label</th>
<th>Accuracy</th>
<th>AUC</th>
<th>Sample size</th>
</tr>
</thead>
<tbody>
<tr>
<td>Zinoviev et al. [19]</td>
<td>NA</td>
<td>54.12%</td>
<td>–</td>
<td>914</td>
</tr>
<tr>
<td>Chen et al. [20]</td>
<td>MS</td>
<td>78.70%</td>
<td>–</td>
<td>47</td>
</tr>
<tr>
<td>Han et al. [16]</td>
<td>MS</td>
<td>–</td>
<td>0.927</td>
<td>1356</td>
</tr>
</tbody>
</table>

Fig. 1. The proposed MC-CNN architecture for lung nodule malignancy suspiciousness classification. The numbers along each side of the cuboid indicate the dimensions of the feature maps. The inside cuboid represents the 3-D convolution kernel and the inside square stands for the associated 2-D pooling region. The dimension of the final hidden feature layer is marked at the bottom. The output layer is a softmax layer that predicts the probability of the class of nodule malignancy suspiciousness, i.e., low malignancy suspiciousness and high malignancy suspiciousness. The pink arrow indicates a multi-crop pooling layer that serves as a surrogate of a max-pooling layer for improving classification performance. (For interpretation of the references to color in this figure caption, the reader is referred to the web version of this paper.)

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2. Methods

In recent studies [26,32,33], the CNN architecture has been brought to the forefront in the image processing field. The core computation seeks a feature representation, also known as the activation of the final hidden layer in the network, that is transformed from high-dimensional features in $\mathbb{R}^{d \times h \times w}$ and remains well separated in a low-dimensional $\mathbb{R}^{p \times 1}$ space ($p < M \times N$). Specifically, two computational units including convolutional layers and pooling layers are used to quantify the mechanism. The network defines a feature-extraction cascade consisting of concatenated convolutional layers and pooling layers (i.e., “Conv+Pool”). Thus, the formed hierarchical network can learn high-level compact features from signal activations of high layers. As shown in Fig. 1, our MC-CNN also consists of “Conv+Pool” layers. However, a proposed multi-crop pooling layer (Section 2.2) is used to surrogate the max-pooling layer to extract multi-scale features.

Given the lung nodule CT images, our goal is to discover a set of discriminative features from the proposed hierarchical neural networks and thus to capture the essence of suspiciousness-specific nodule information. The challenge is that the image space is heterogeneous including both healthy tissues and nodules at different visual scales. Compared to the conventional feature extraction [13,14,34], we propose an integrated computational deep learning architecture. The major components which form the basis of our multi-crop CNN are presented from Sections 2.1 to 2.3.

2.1. Conv+pool layer design

The CNN starts from a convolutional layer where we adopt the Randomized Leaky Rectified Linear Units (RReLU) [35,36] as a non-linear transformation. Formally, the convolution operation is defined by

$$ y^i = \text{RReLU}\left( \sum_j c^i_j \cdot h^j + b^i \right), $$

where $h^j$ and $y^j$ are the ith input map and the jth output map, respectively. We define $c^i_j$ as the convolution kernel between the ith input map and the jth output map (denotes the 2-D convolution), $b^i$ is the bias of the jth output map. $h^j$, $y^j$ and $c^i_j$ are all 2-D. The entire input, output and convolution kernel of a convolutional layer are a stack of $h^i$, $y^j$, and $c^i_j$. As seen in Fig. 1, there are 64 CT slices $(h^i)$ in the input layer and the convolutional layer outputs 64 convolutional feature maps ($y^j$). Accordingly, the number of convolution kernels is 64 with dimension of $3 \times 3 \times 64$ voxels. Both $c^i_j$ and $b^i$ are continuously learned in the network training process. The non-linear transformation function RReLU($x$) [35] is expressed as

$$ \text{RReLU}(x) = \begin{cases} 
  x & \text{if } x \geq 0 \\
  x - a & \text{if } x < 0, \ a \sim U(b_l, b_u) 
\end{cases} $$

where $a$ is a random factor sampled from a uniform distribution $U(b_l, b_u)$, and $b_l$ and $b_u$ are the lower and higher bounds of the distribution respectively. RReLU allows for a small, non-zero gradient initialization for unit activation that has been proven to be less prone to overfit the dataset [35,36] than conventional ReLU [26] in a classification task, especially when the training samples are limited.

Following the convolutional layer, a max-pooling layer is commonly introduced to select feature subsets. It is defined as

$$ y^j_{(j,k)} = \max_{0 \leq m \leq n - s} (h^i_{j + s, m + k + s, n + s}), $$

where $s$ is the size of pooling region, $y^j_{(j,k)}$ represents the neuron at position $(j,k)$ in the ith output map $h^i_{j + s, m + k + s, n + s}$, denotes the neuron at position $(j\cdot s + m, k\cdot s + n)$ in the ith input map where $m$ and $n$ are the offsets of the position. The advantage of using the max-pooling layer is its translation invaribility even when different nodule images are not well-aligned. In the following section, we introduce our multi-crop pooling strategy which can surrogate the traditional max-pooling operation.

2.2. Multi-crop pooling strategy

We extend the traditional max-pooling layer into our multi-crop pooling layer which allows the capture of nodule-centric visual features. Traditional max-pooling layers in the network play a role of selecting feature subsets and reducing the size of the feature maps. However, the max-pooling operation performs
uniformly on each feature map, and thus the max-pooling layer is a single-level feature reduction operation. Such a setting hinders it from capturing accurate object-specific information when the size of objects varied largely in the images. As seen in Fig. 3, a large variability of nodule sizes steers us to pursue an alternative strategy for capturing nodule sensitive information.

We proposed a multi-crop layer to surrogate the conventional max-pooling layer. It is a strategy with repetitive pooling features, enabling a multi-scale feature extraction from the input feature maps on nodule samples. Given a stack of feature maps from a previous convolutional layer, a multi-crop strategy is designed to fully capture nodule-centric features. As shown in Fig. 4, the concatenated nodule-centric feature $f = \{f_0, f_1, f_2\}$ is formed from three nodule-centric feature patches $R_0, R_1, R_2$ respectively. Specifically, let the size of $R_0$ be $l \times l \times n$, where $l \times l$ is the dimension of the feature map and $n$ is the number of feature maps:

$$f_i = \max - pooling_2 \times 2(R_i), \quad i = \{0, 1, 2\},$$

where $R_0, R_1$ are two center regions with size of $(l/2) \times (l/2) \times n$ and $(l/4) \times (l/4) \times n$. The superscript of “max-pool” indicates the frequency of the utilized max-pooling operation on $R_i$. In Fig. 4, the input of multi-crop pooling operation is the convolutional features $R_0$ obtained from either the original image or the pooled features. $R_1$ is the center region cropped from $R_0$ and $R_2$ is the center region cropped from $R_1$. Then, $R_0$ is max-pooled twice and $R_1$ is max-pooled once to generate pooled feature maps $f_0$ and $f_1$. $R_2$ serves as $f_2$ without any pooling operation. The final multi-crop feature is made up with the concatenation of $f_0, f_1$, and $f_2$. Specifically, the strategy on targeting nodule-specific patches allows us to feed multi-scale nodule sensitive information into the following convolutional layers. The functionality of the multi-crop pooling layer is similar to that of the max-pooling layer since they both pool the features. Thus, it can surrogate any max-pooling layers for the purpose of extracting multi-scale features. The effectiveness of multi-crop features is discussed in Section 4.2.

The objective of multi-crop pooling strategy is to extract multi-scale features from a single network. The strategy draws inspiration from spatial pyramid pooling network (SPPNet) [37] which concatenated the feature pyramid as the final feature vector. Although both SPPNet and the proposed method share a similarity in extracting features at different scales, several remarkable differences are recognized: (1) the pooling frequency of multi-crop pooling relies on feature location in the feature map, while spatial pyramid pooling strategy pools features at different location equally; (2) the output features of our multi-crop pooling layer at different scales have the same dimension while the feature dimensions from spatial pyramid pooling are determined by their pooling levels; (3) a third and more important distinction is that the output of our multi-crop pooling at each scale can be concatenated to feed into the following convolutional layer, while the output of spatial pyramid pooling can only be placed at the top of a CNN.

Speaking of computational complexity, unlike conventional multi-scale features [31] with multiple parallel networks in image space, the multi-crop pooling layer could generate multi-scale features from a singular MC-CNN pipeline, which greatly simplifies the training process and shortens the training time without sacrificing the classification accuracy (Section 4.2).

2.3. Loss function

In general, multiple pairs of concatenated Conv+Pool layers consist of a major network architecture and the last pooling layer is usually connected to a fully-connected layer. The output layer of the entire network is a 2-way softmax layer (see Fig. 1) predicting the probability distribution over low malignancy suspiciousness and high malignancy suspiciousness:

$$p_j = \frac{\exp(y_j)}{\exp(y_0) + \exp(y_j)}, \quad j = \{0, 1\},$$

where $y_j = \sum_{i=1}^{52} h_i w_{ij} + b_j$ is the linear combination of the input $h_i$ (the activations of the final hidden features in Fig. 1). $w_{ij}$ is the weight and $b_j$ is the scalar.

The network is learned by minimizing the cross entropy loss, which can be expressed as

$$\text{LOSS} = -(q \log p_i + (1 - q) \log p_0),$$

where $q$ indicates suspiciousness label with the value 1 or 0 corresponding to being high suspiciousness or low suspiciousness respectively. The network is trained using Stochastic Gradient Descent (SGD) with a standard backprop [38,39].

2.4. Prediction modeling and model evaluation

In addition to classify nodule malignancy suspiciousness category, we also predict nodule attributes including nodule subtlety, margin, and diameter (Section 4.5), which could potentially be used to model nodule malignancy uncertainty. For malignancy suspiciousness, subtlety and margin, we model them as a binary classification problem and predict whether the nodule belongs to the high score category or the low score category. For nodule diameter estimation, we modify our MC-CNN to be a regression model by replacing the last softmax layer with a single neuron which predicts the estimated diameter in a real value. Balanced datasets, obtained by sample selection, are prepared for the classification tasks. For diameter estimation model, the entire dataset without any balancing process is used. More detailed discussions on the validation setting are given in Section 3.

In order to do model selection for predicting outcomes, we split the dataset into the training set, validation set and test set. Each network model was trained for 5000 iterations, and we saved the trained model at every 100 iterations. After the entire training process, the associated validation scores obtained from the validation set were sorted in a descending order. We then selected the top 3 models as the final trained models and the prediction outcome of a test patch was the average of the ensemble probability scores.

The performance of classifying malignancy suspiciousness (Section 4.2), subtlety and margin (Section 4.5.1), and of estimating nodule diameter (Section 4.5.2) were evaluated via five-fold cross validation. In each experiment, three folds were used as the training set. One fold was used as the validation set and the rest one as the test set. We reported the classification performance by averaging the classification accuracies and the area under the curve (AUC) scores across 30 times tests.

3. Dataset description

3.1. Dataset

The dataset used in this work is the LIDC-IDRI dataset [21], consisting of 1010 patients with lung cancer thoracic CT scans as well as mark-up annotated lesions. We included nodules along with their annotated centers from the nodule collection report. The diameters of the nodules range from 3 mm to 30 mm. Since the resolution of the images varied, we resampled images using

spline interpolation to have a fixed resolution with 0.5 mm/voxel along all three axes. Each nodule patch was cropped from the resampled CT image based on the annotated nodule center.

The malignancy suspiciousness of each nodule is rated from 1 to 5 by four experienced thoracic radiologists, indicating an increasing degree of malignancy suspiciousness. We chose the averaged malignancy rating for each nodule as [40,31,16]: for those with an average score lower than 3, we labelled them as low malignancy-suspicious nodules (LMNs); for those with an average score higher than 3, we labelled them as high malignancy-suspicuous nodules (HMNs). We removed nodule samples with ambiguous IDs. Overall, there were 880 LMN and 495 HMN cases included for performance evaluation. For nodules with an average rating of 3, we followed the study in [16] by conducting two additional experiments of excluding them from the evaluation and including them in another experiment, respectively. The number of nodules with an average rating of 3 was 1243 in total. These nodules will be referred to as uncertain nodules (UNs) in the following sections since they do not fall to any distinct categories.

Similarly, for nodule subtlety and margin attributes analysis in Section 4.5, we selected equal numbers of positive nodule samples (average attribute rating >3) and negative nodule samples (average attribute rating <3) from the LIDC-IDRI dataset, resulting in 756 nodules for subtlety classification and 658 nodules for margin classification. For nodule diameter estimation in Section 4.5, we used the entire dataset of 2618 nodules for the regression modeling.

3.2. Data augmentation

We sought to train the MC-CNN model with augmented training samples that complemented the learning process given limited training samples. We augmented nodules by random image translation, rotation and flip operations as in [26,41,42]. The translation was in a range of [−6, 6] voxels; the rotation was done by first swapping the three axes in 3-D followed by a 2-D rotation of [90°, 180°, 270°]. Nodule patches were augmented when fed into the input layer. Such augmentation helped the MC-CNN to capture nodule features invariant to image-level translation, rotation and flip operations.

4. Experiments and results

In this section, we evaluate our MC-CNN performance by measuring the classification accuracy and the AUC score of computed deep features. We perform a systematic evaluation against different parameters including the number of convolution kernels,
the position of multi-crop pooling layer in the whole architecture, and dataset sample sizes. We first describe the experimental setup. Then, we report results of our MC-CNN and compare it with the state-of-the-art approaches. We also conduct an exploratory analysis on modeling nodule malignancy uncertainty. Finally, we demonstrate the effectiveness of our MC-CNN in nodule subtlety prediction, nodule margin prediction and nodule diameter estimation.

4.1. Experimental setup

To observe performance with regard to the network configuration, we investigated different configurations including the number of convolution kernels of each convolution layer and the position of a multi-crop pooling layer. The MC-CNN which had \( n_{ker} \) convolution kernels for each convolutional layer and a multi-crop pooling layer surrogating the \( i \)th max-pooling layer (the position of the multi-crop pooling layer) was named as MC – CNN\(^{n_{ker}}\)\(_{i}\), where \( n_{ker} = (16, 32, 64) \) and \( i = (1, 2, 3) \). Thus, there were 9 different configurations in total. For simplicity, MC – CNN, was used to indicate a general MC-CNN with an \( i \)th max-pooling layer surrogated by a multi-crop pooling layer but with an arbitrary \( n_{ker} \).

The number of neurons \( n_{h} \) in the final hidden feature layer was fixed to 32. All results of these parameter settings are discussed in Section 4.2.

To capture a majority of nodule morphology, the input nodule patch size was set to 64 \( \times \) 64 \( \times \) 64 voxels. We set the learning rate to be \( 1.0 \times 10^{-3} \). In order to relieve the risk of overfitting, we added an L-2 norm weight decay during the training process and the weight decay coefficient was \( 5 \times 10^{-4} \). We let \( b_{1} = 3 \) and \( b_{2} = 8 \) in Eq. (2) as [35]. The pooling region size \( s \) was 3 with pooling stride of 2 in the first two pooling layers, while \( s = 4 \) with a stride of 4 in the third pooling layer to decrease the feature dimension. The size of convolution kernel was \( 3 \times 3 \). These chosen parameters were commonly used as discussed in [26,43].

Our MC-CNN implementation was based on CAFFE [44], HOG and LBP descriptors, which were implemented in the scikit-image package [45], were compared for the performance evaluation of segmentation-free classification methods with our method (Section 4.3). The classifier used was the Support Vector Machine (SVM) classifier from the scikit-learn package [46].

4.2. MC-CNN classification performance

In this section, we perform a systematic evaluation against different parameters including the number of convolution kernels, the position of multi-crop pooling layer, and the dataset sample size. During each round of the five-fold cross validation, there were originally 825 nodules (528 LMNs and 297 HMNs) in the training set and 275 nodules (176 LMNs and 99 HMNs) in either of the validation set and test set. We oversampled HMN samples to approximately balance the training set.

4.2.1. Results with different network configurations

Following the description in Section 4.1, there were 9 different network configurations in total with respect to the number of convolution kernels \( (n_{ker} = (16, 32, 64)) \) and the \( i \)th position \( (i = (1, 2, 3)) \) of the multi-crop layer. As shown in Fig. 5, our MC-CNN was stable to different configurations with all being above 86% in accuracy with a maximum standard variation of 0.27% and above 0.90 for the AUC score with a maximum standard variation of 0.0016. The highest classification accuracy obtained was 87.14% from MC – CNN\(^{64}\)\(_{2}\), and the highest AUC score was 0.93 from MC – CNN\(^{64}\)\(_{2}\). Besides \( n_{ker} \) and \( i \), we also evaluated the effect of the number of neurons (\( n_{h} \)) in the hidden feature layer. We trained MC – CNN\(_{2}\) with different \( n_{h} \) from \( (16, 32, 64) \). Classification accuracy comparison is shown in Fig. 6. It was obvious that the performance was quite stable and the variation was less than 0.3% for a certain \( n_{ker} \), indicating \( n_{h} \) is not a crucial parameter to the performance of our MC-CNN.

4.2.2. Effectiveness of multi-crop pooling features

We justify the effectiveness of the multi-crop pooling features by comparing our MC-CNNs with three other networks without applying multi-crop pooling operation in the feature space. First, since multi-crop pooling on feature maps helped achieve high classification accuracy, we also applied multi-crop pooling operation directly to the image space (i.e., the input nodule patches). Instead of using max-pooling inside the multi-crop pooling layer, we used average-pooling which simulated the image

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The SVM classifier was used for classification. We extracted HOG descriptors and LBP descriptors with three scales on nodule patches, i.e., 32 × 32 × 32 voxels, 64 × 64 × 64 voxels, and 96 × 96 × 96 voxels. Accuracies of HOG and LBP descriptors were shown in Table 4. We found that HOG descriptor was quite sensitive to the size of the cell window \( s_w \). For LBP descriptor, we observed that the number of neighborhood points \( n_{pt} \) was positively related to the performance probably because sophisticated neighborhood structures led to improved results. However, when competing with the best results among these two descriptors, our method outperformed them by 12.96% and 19.79% respectively. Overall, our observation confirmed that parametric textural descriptors were sensitive to various parameters.

4.3.2. Comparison with segmentation-dependent classification

We have reported related results on the LIDC-IDRI dataset in the literature in Table 1. Although we noticed that different number of samples were used which made the fair comparison difficult, the results from our MC-CNN were still quite competitive in terms of both classification accuracy and the AUC score. In this section, we included two more metrics: sensitivity and specificity. The sensitivity and the specificity of our method were 0.77 and 0.93.

4.3.1. Comparison with HOG and LBP based classification

We first compared our results with commonly used descriptors including HOG and LBP descriptors. For HOG descriptor, we used different cell window sizes, \( s_w = \{8, 16, 32\} \) with the number of orientations \( n_g = 8 \). For LBP descriptor, the uniform LBP descriptor was computed with different neighbourhood points \( n_{pt} = \{8, 16, 24\} \). The SVM classifier was used for classification. We extracted HOG descriptors and LBP descriptors with three scales on nodule patches, i.e., 32 × 32 × 32 voxels, 64 × 64 × 64 voxels, and 96 × 96 × 96 voxels. Accuracies of HOG and LBP descriptors were shown in Table 4. We found that HOG descriptor was quite sensitive to the size of the cell window \( s_w \). For LBP descriptor, we observed that the number of neighborhood points \( n_{pt} \) was positively related to the performance probably because sophisticated neighborhood structures led to improved results. However, when competing with the best results among these two descriptors, our method outperformed them by 12.96% and 19.79% respectively. Overall, our observation confirmed that parametric textural descriptors were sensitive to various parameters.

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4.2.3. Performance with varying data samples

We evaluate the performance of MC – CNN\(^4\) on datasets with different sizes by randomly sampling different numbers of nodules from the original dataset including three sub-datasets: a quarter, a half and the entire dataset. The classification accuracies are shown in Table 3. Adding more training data improved the model performance from 83.09% to 87.14%, leading to a performance increase by around 4%. Although Table 3 demonstrated empirical success of the MC-CNN with regard to different sizes of samples, we would expect to collect more nodule samples to further improve and validate the stability of the proposed approach.

4.3. Competing with state-of-the-art approaches

We compare our method with both segmentation-free and segmentation-dependent classification methods in this section. Segmentation-free methods included LBP and HOG descriptors working on nodule patches. Segmentation-dependent methods relied on nodule image segmentation for feature engineering.

<table>
<thead>
<tr>
<th>Dataset size</th>
<th>Accuracy (%)</th>
<th>Sensitivity</th>
<th>Specificity</th>
</tr>
</thead>
<tbody>
<tr>
<td>340</td>
<td>83.09</td>
<td>0.73</td>
<td>0.85</td>
</tr>
<tr>
<td>1030</td>
<td>86.36</td>
<td>0.87</td>
<td>0.78</td>
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<tr>
<td>1375</td>
<td>87.14</td>
<td>0.93</td>
<td>0.93</td>
</tr>
</tbody>
</table>

Table 4

Classification accuracies using HOG descriptor with different \( s_w \) and LBP descriptor with different \( n_{pt} \).

<table>
<thead>
<tr>
<th>Descriptor</th>
<th>Parameter</th>
<th>32 (%)</th>
<th>64 (%)</th>
<th>96 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HOG</td>
<td>( s_w = 8 )</td>
<td>74.18</td>
<td>66.69</td>
<td>64.07</td>
</tr>
<tr>
<td></td>
<td>( s_w = 16 )</td>
<td>63.27</td>
<td>66.40</td>
<td>65.16</td>
</tr>
<tr>
<td></td>
<td>( s_w = 32 )</td>
<td>49.82</td>
<td>56.15</td>
<td>56.58</td>
</tr>
<tr>
<td>LBP</td>
<td>( n_{pt} = 8 )</td>
<td>64.58</td>
<td>49.24</td>
<td>36.00</td>
</tr>
<tr>
<td></td>
<td>( n_{pt} = 16 )</td>
<td>66.40</td>
<td>59.93</td>
<td>52.22</td>
</tr>
<tr>
<td></td>
<td>( n_{pt} = 24 )</td>
<td>67.35</td>
<td>59.20</td>
<td>54.84</td>
</tr>
</tbody>
</table>

Fig. 7. The receiver operating characteristic curve (ROC curve) of our MC – CNN\(^4\), the Massive-feat method and the Autoencoder method. It can be seen that the ROC of our MC – CNN\(^4\) (AUC = 0.93) is very competitive compared to the Autoencoder method (AUC = 0.86) and the Massive-feat method (AUC = 0.89).
0.93 respectively. We implemented two approaches in the literature for additional comparison with the same number of nodules (see Table 5 and Fig. 7). The first one is the autoencoder-based method (Autoencoder) [30] in an unsupervised learning scheme. We tested it on the same dataset and achieved the classification accuracy of 80.29% with an AUC score of 0.86. The sensitivity and the specificity were 0.73 and 0.85 respectively. The lack of prior label information in the unsupervised learning may lead to a suboptimal feature learning, causing its outcomes lower than that of the MC-CNN. Second, we implemented a massive feature mining pipeline (Massive-feat) following the strategy in [13]. Four types of nodule image features were included: first order statistics, shape and size features, textural features, and wavelet features. After feature selection using the mRMR score [47], the top 54 features were chosen from the extracted 319 dimensional features. Applying the SVM classifier led to the best accuracy of 83.21% with an AUC score of 0.89. The sensitivity and specificity were 0.87 and 0.78 respectively. Both the classification accuracies and the AUC scores from two implemented approaches were shown inferior to the proposed MC-CNN. Though the sensitivity of our method was lower than that of the Massive-feat method, the specificity of our MC-CNN was higher compared to those of the other two methods. More importantly, the reporting figures of our approach here were not meant to lead a significant improvement over the current literature. Instead, we sought to demonstrate an alternative feature extraction pipeline that can complement state-of-the-art architectures for lung nodule analysis.

4.4. Exploratory analysis on modeling nodule malignancy uncertainty

Since our prior results were based on a binary setting of malignancy suspiciousness classification, in this section, we extended to estimate nodule malignancy uncertainty by taking into account nodules with a moderate score of 3. The uncertainty estimation on nodule malignancy suspiciousness is challenging because of ambiguous assessment from human experts. We provided exploratory evidence to model uncertain nodules by analyzing inclination of uncertain samples to the distinct group of LMNs or HMNs, so that we may be able to gain insight into better patient sub-group stratification.

Two tasks were designed to quantify nodule malignancy uncertainty by applying the model of MC – CNN [64]. First, the uncertain nodules with a score 3 were either categorized into LMNs or HMNs respectively. Second, we additionally treated them as an independent category and performed classification on three groups. Table 6 showed the classification results with uncertain nodules included. Comparing the first row in Table 6 with our best results, we found that uncertain nodules made the model slightly inferior to that trained without uncertain nodules, probably since inclusion of uncertain nodules introduced variation within each category. Comparing first and second columns, we found that incorporating uncertain cases into LMNs led to better results than incorporating them into HMNs.

The observation indicated that uncertain cases shared more similarities with LMNs. The finding presented that radiologists seemed to have a biased scoring towards classifying some LMS cases into uncertain nodules. Our observation was consistent with the study [16]. Also, the dropped accuracy was observed when the uncertain nodules were regarded as an independent category. The result is not surprising since nodules with a moderate score present heterogeneous characteristics in nature, leading to deteriorate performance in classification during both training and test phases. The evidence data here suggests that a more sophisticated comparison will be needed to investigate between subtle sub-groups in the future.

4.5. Nodule semantic prediction and diameter estimation

Beyond nodule malignancy suspiciousness classification, we quantify nodule semantic prediction including subtlety and margin and nodule diameter estimation using the MC – CNN [64].

4.5.1. Nodule subtlety and margin prediction

We performed semantic label prediction including two attributes: subtlety and margin. Subtlety indicates the difficulty in nodule detection which refers to the contrast between the lung nodule and its surroundings, while margin describes how well-defined the margins of the defining nodule [48]. The model we used here is the classification model of MC – CNN [64] and the evaluation method is also the five-fold cross validation. The classification accuracy of subtlety is 74.32% and that of margin is 76.99%.

4.5.2. Nodule diameter estimation

The diameter of a solitary pulmonary nodule (SPN) can be a useful predictor of malignancy—larger diameter indicates the increasing suspiciousness of nodule malignancy [49]. In this experiment, given the five-fold cross validation, we used the regression version of MC – CNN [64] (Section 2.4). The metric used to evaluate the estimation performance was the relative estimation error

\[ E_r = \frac{|d_{est} - d_{truth}|}{d_{truth}}, \]

where \(d_{est}\) is the estimated diameter and \(d_{truth}\) indicates the ground truth diameter. The distribution of \(E_r\) is shown in Fig. 8. The population of \(E_r\) less than 0.2, 0.3 and 0.4 respectively occupy 73.78%, 84.54% and 90.15% of the entire dataset. The results suggest an alternative way of estimating the nodule diameter, indicating a strong correlation of the learned deep features with lung nodule diameter distribution.

---

Table 6  
Classification accuracies including uncertain nodules. “UNs” indicate uncertain nodules. “IC” indicates an independent category.

<table>
<thead>
<tr>
<th>Settings</th>
<th>UNs as LMNs (%)</th>
<th>UNs as HMNs (%)</th>
<th>UNs as IC (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>in training set</td>
<td>86.12</td>
<td>85.60</td>
<td>–</td>
</tr>
<tr>
<td>in both training and test sets</td>
<td>87.29</td>
<td>72.57</td>
<td>62.46</td>
</tr>
</tbody>
</table>

Fig. 8. The distribution of the relative estimation error \(E_r\). The population of \(E_r\) within 0.2, 0.3 and 0.4 occupy 73.78%, 84.54% and 90.15% of the entire dataset.
Table 7

<table>
<thead>
<tr>
<th>Methods</th>
<th>Segmentation</th>
<th>Feature extraction</th>
<th>Classifier</th>
<th>Training time</th>
<th>Test time</th>
</tr>
</thead>
<tbody>
<tr>
<td>Massive-feat</td>
<td>Manual</td>
<td>10h15min25s</td>
<td></td>
<td>0.09 s</td>
<td>32.76 s</td>
</tr>
<tr>
<td>Autoencoder</td>
<td>Manual</td>
<td>7.29 s</td>
<td></td>
<td>0.14 s</td>
<td>0.01 s</td>
</tr>
<tr>
<td>MC – CNN64</td>
<td>–</td>
<td>47min01s</td>
<td></td>
<td>0.23 s</td>
<td></td>
</tr>
</tbody>
</table>

5. Discussion

In this paper, we proposed a deep learning computational architecture, called MC-CNN, to classify nodule malignancy suspiciousness using CT images. It is an end-to-end architecture which embeds nodule feature extraction into a hierarchical network and simplifies conventional lung nodule malignancy suspiciousness classification by removing nodule segmentation and hand-crafted feature engineering. Providing early suspiciousness estimation from imaging allowed a strategy of non-invasively identifying patient sub-groups before treatments of needle biopsy or surgical resection. Thus, it has the potential to facilitate radiologists to discern the underlying risk factors for better individualized patient management. Experimental results demonstrated that our proposed method achieved promising results in both classification accuracy (87.14%) and the AUC score (0.93). Additional semantic prediction and diameter estimation reaffirmed the strength of the proposed approach in characterizing nodule-related information. To further assimilate diagnostic values of the proposed approach, we would expect to validate the deep learning architecture by incorporating additional lung cancer imaging studies with both radiologists’ opinions and follow-up pathologic scores. As vast quantity of clinical imaging sequences are becoming increasingly available, our data-driven model holds promise for early diagnosis with more rapid clinical translation.

Although accurate processing time comparison of different methods is difficult, we list the time consumed by the Massive-feat method [13], the Autoencoder method [30] and our MC – CNN64 in Table 7. All the methods run on the same machine with 12GB memory and a 6-core Intel Xeon CPU. Nvidia Tesla K40 GPU was enabled for our MC – CNN64 model and the Autoencoder method. We did not migrate the feature extraction code in the Massive-feat method to GPU. The manual segmentation time for the Massive-feat method and the Autoencoder method were not included because the segmentation was provided in the dataset. We found that hand-crafted feature extraction in the Massive-feat method was very time-consuming which took more than 10 h and the test time for one single nodule was also much longer than that of the other two methods. The input data of the Autoencoder method was the 2-D nodule slice while that of our method was 3-D which brought much more computational cost. This could explain why our method took more time than the Autoencoder method. It was also obvious that our method could simplify the traditional nodule analysis pipeline by removing nodule segmentation and feature extraction.

The rationale for seeking ‘deep features’ is that deep learning networks would make mostly correct assumptions about the nature of images by varying the depth and breadth of the network capacity [26]. As the results in [50], through hierarchical networks, the CNN can produce a dimensional reduction that is particularly useful for image-related classification. Our MC-CNN prioritized a repetitive pooling strategy for nodule-centric feature extraction. By considering different regions of the feature maps independently, the strategy preserved more details of the salient region of nodules. Thus, features from small nodules were also well kept and forwarded to the following layers, indicating that our MC-CNN was able to capture a variety of nodule dynamic structures.

Dropout [51] is a known strategy to prevent the CNN from over-fitting. However, we did not observe much test difference between networks with or without dropout on the LIDC-IDRI dataset. The reason may be ascribed to that the LIDC-IDRI dataset is quite different from generic image datasets (e.g., the imagenet dataset) having thousands of categories, where the learning models are prone to overfit decision boundaries easily. The use of 3-D augmentation created augmented training samples that preserve intra-class variation to minimize the potential over-fitting issue. Additionally, results on semantic prediction and diameter estimation revealed the generalized performance of the proposed method.

The proposed study complemented the traditional approaches. The only prerequisite of our method is the identification of the nodule central location, which is a substitute for conventional nodule image segmentation. Both multi-crop pooling and max-pooling can tolerate a small amount of shift of a nodule center point, thus the process actually does not require an accurate localization of nodule centers. This suggests an appealing strategy of approach initialization. However, when dealing with a growing number of clinical imaging centers, an extra automatic nodule detection process might be needed for both our method and the traditional nodule classification methods in order to speed up the diagnosis process.

6. Conclusion

Deep learning architecture is a rising computational paradigm in developing predictive models of diseases. In this paper, we introduced a deep learning model of MC-CNN to tackle the challenging problem of lung nodule malignancy suspiciousness classification. We demonstrated that the learned deep features were able to capture nodule salient information by the multi-crop pooling strategy. The encouraging results on nodule malignancy suspiciousness classification showed the effectiveness of our MC-CNN. Additional experiments on nodule semantic prediction and nodule diameter estimation revealed that the proposed method could be potentially helpful to other aspects of nodule-relevant characteristics analysis. In general, the extracted deep features can be considered to be integrated with conventional image features to further improve the precision performance for lung cancer patients.

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