DeepLung: 3D Deep Convolutional Nets for Automated Pulmonary Nodule Detection and Classification

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Abstract

In this work, we present a fully automated lung CT cancer diagnosis system, DeepLung. DeepLung contains two parts, nodule detection and classification. Considering the 3D nature of lung CT data, two 3D networks are designed for the nodule detection and classification respectively. Specifically, a 3D Faster R-CNN is designed for nodule detection with a U-net-like encoder-decoder structure to effectively learn nodule features. For nodule classification, gradient boosting machine (GBM) with 3D dual path network (DPN) features is proposed. The nodule classification subnetwork is validated on a public dataset from LIDC-IDRI, on which it achieves better performance than state-of-the-art approaches, and surpasses the average performance of four experienced doctors. For the DeepLung system, candidate nodules are detected first by the nodule detection subnetwork, and nodule diagnosis is conducted by the classification subnetwork. Extensive experimental results demonstrate the DeepLung is comparable to the experienced doctors both for the nodule-level and patient-level diagnosis on the LIDC-IDRI dataset.

Introduction

Lung cancer is the most common cause of cancer-related death in men. Low-dose lung CT screening provides an effective way for early diagnosis, which can sharply reduce the lung cancer mortality rate. Advanced computer-aided diagnosis systems (CADs) are expected to have high sensitivities while at the same time maintaining low false positive rates. Recent advances in deep learning enable us to rethink the ways of clinician lung cancer diagnosis.

Current lung CT analysis research mainly includes nodule detection (Dou et al. 2017; Ding et al. 2017), and nodule classification (Shen et al. 2015; 2016; Hussein et al. 2017; Yan et al. 2016). There is few work on building a complete lung CT cancer diagnosis system for fully automated lung CT cancer diagnosis, integrating both nodule detection and nodule classification. It is worth exploring a whole lung CT cancer diagnosis system and understanding how far the performance of current technology differs from that of experienced doctors. To our best knowledge, this is the first work for a fully automated and complete lung CT cancer diagnosis system.

The emergence of large-scale dataset, LUNA16 (Setio et al. 2017), accelerates the nodule detection related research. Typically, nodule detection consists of two stages, region proposal generation and false positive reduction. Traditional approaches generally require manually designed features such as morphological features, voxel clustering and pixel thresholding (Murphy et al. 2009; Jacobs et al. 2014). Recently, deep convolutional networks, such as Faster R-CNN (Ren et al. 2015) and fully convolutional networks (Long, Shelhamer, and Darrell 2015), are employed to generate candidate bounding boxes (Ding et al. 2017; Dou et al. 2017). In the second stage, more advanced methods or complex features, such as more carefully designed texture features, are designed to remove false positive nodules. Due to the 3D nature of CT data and the effectiveness of Faster R-CNN for object detection in 2D natural images (Huang et al. 2016b), we design a 3D Faster R-CNN for nodule detection with U-net-like encoder-decoder structure to effectively learn latent features (Ronneberger, Fischer, and Brox 2015).

Before the era of deep learning, feature engineering followed by classifiers is a general pipeline for nodule classification (Han et al. 2013). After the public large-scale dataset, LIDC-IDRI (Armato et al. 2011), becomes available, deep learning based methods have become dominant for nodule classification research (Shen et al. 2016). Multi-scale deep convolutional network with shared weights on different scales has been proposed for the nodule classification (Shen et al. 2015). The weight sharing scheme reduces the number of parameters and forces the multi-scale deep convolutional network to learn scale-invariant features. Inspired by the recent success of dual path network (DPN) on ImageNet (Chen et al. 2017; Deng et al. 2009), we propose a novel framework for CT nodule classification. First, we design a 3D deep dual path network to extract features. Then we use gradient boosting machines (GBM) with deep 3D dual path features, nodule size and cropped raw nodule CT pixels for the nodule classification (Friedman 2001).

At last, we build a fully automated lung CT cancer diagnosis system, DeepLung, by combining the nodule detection network and nodule classification network together, as illustrated in Fig. 1. For a CT image, we first use the detection subnetwork to detect candidate nodules. Next, we employ the classification subnetwork to classify the detected nodules into either malignant or benign. Finally, the patient-level di-
Figure 1: The framework of DeepLung. DeepLung first employs 3D Faster R-CNN to generate candidate nodules. Then it uses 3D deep dual path network (DPN) to extract deep features from the detected and cropped nodules. Lastly, gradient boosting machine (GBM) with deep features, detected nodule size, and raw nodule pixels are employed for classification. Patient-level diagnosis can be achieved by fusing the classification results of detected nodules in the CT.

Our main contributions are as follows: 1) To our best knowledge, DeepLung is the first work for a complete automated lung CT cancer diagnosis system. 2) Two 3D deep convolutional networks are designed for nodule detection and classification. Specifically, inspired by the effective of Faster R-CNN for object detection and deep dual path network's success on ImageNet (Huang et al. 2016b; Chen et al. 2017), we propose 3D Faster R-CNN for nodule detection, and 3D deep dual path network for nodule classification. 3) Our classification framework achieves superior performance compared with state-of-the-art approaches, and the performance surpasses the average performance of experienced doctors on the largest public dataset, LIDC-IDRI dataset. 4) The DeepLung system is comparable to the average performance of experienced doctors both on nodule-level and patient-level diagnosis.

Related Work

Traditional nodule detection requires manually designed features or descriptors (Lopez Torres et al. 2015). Recently, several works have been proposed to use deep convolutional networks for nodule detection to automatically learn features, which is proven to be much more effective than hand-crafted features. Setio et al. proposes multi-view convolutional network for false positive nodule reduction (Setio et al. 2016). Due to the 3D nature of CT scans, some work propose 3D convolutional networks to handle the challenge. The 3D fully convolutional network (FCN) is proposed to generate region candidates, and deep convolution network with weighted sampling is used in the false positive candidates reduction stage (Dou et al. 2017). Ding et al. uses the Faster R-CNN to generate candidate nodules, followed by 3D convolutional networks to remove false positive nodules (Ding et al. 2017). Due to the effective performance of Faster R-CNN (Huang et al. 2016b; Ren et al. 2015), we design a novel network, 3D Faster R-CNN, for the nodule detection. Further, U-net-like encoder-decoder scheme is employed for 3D Faster R-CNN to effectively learn the features (Ronneberger, Fischer, and Brox 2015).

Nodule classification has traditionally been based on segmentation (El-Baz et al. 2011; Zhu and Xie 2016) and manual feature design (Aerts et al. 2014). Several works designed 3D contour feature, shape feature and texture feature for CT nodule diagnosis (Way et al. 2006; El-Baz et al. 2011; Han et al. 2013). Recently, deep networks have been shown to be effective for medical images. Artificial neural network was implemented for CT nodule diagnosis (Suzuki et al. 2005). More computationally effective network, multi-scale convolutional neural network with shared weights for different scales to learn scale-invariant features, is proposed for nodule classification (Shen et al. 2015). Deep transfer learning and multi-instance learning is used for patient-level lung CT diagnosis (Shen et al. 2016). A comparison on 2D and 3D convolutional networks is conducted and shown that 3D convolutional network is superior than 2D convolutional network for 3D CT data (Yan et al. 2016). Further, a multi-task learning and transfer learning framework is proposed for nodule diagnosis (Hussein et al. 2017; Zhu et al. 2017). Different from their approaches, we propose a novel classification framework for CT nodule diagnosis. Inspired by the recent success of deep dual path network (DPN) on ImageNet (Chen et al. 2017), we design a novel totally 3D deep dual path network to extract features from raw CT nodules. Then, we employ gradient boosting machine (GBM) with the deep DPN features, nodule size, and raw nodule CT pixels for the nodule diagnosis. Patient-level diagnosis can be achieved by fusing the nodule-level diagnosis.

DeepLung Framework

The fully automated lung CT cancer diagnosis system, DeepLung, consists of two parts, nodule detection and nodule classification. We design a 3D Faster R-CNN for nodule detection, and propose gradient boosting machine with deep 3D dual path network features, raw nodule CT pixels and nodule size for nodule classification.
3D Faster R-CNN for Nodule Detection

The 3D Faster R-CNN with U-net-like encoder-decoder structure is illustrated in Fig. 2. Due to the GPU memory limitation, the input of 3D Faster R-CNN is cropped from 3D reconstructed CT images with pixel size $96 \times 96 \times 96$. The encoder network is derived from ResNet-18 (He et al. 2016). Before the max-pooling, two convolutional layers are used to generate features. After that, four residual blocks are employed in the encoder subnetwork. We integrate the U-net-like encoder-decoder design concept in the detection to learn the deep networks efficiently (Ronneberger, Fischer, and Brox 2015). In fact, for the region proposal generation, the 3D Faster R-CNN conducts pixel-wise multi-scale learning and the U-net is validated as an effective way for pixel-wise labeling. This integration makes candidate nodule generation more effective. In the decoder network, the feature maps are processed by deconvolution layers and residual blocks, and are subsequently concatenated with the corresponding layers in the encoder network (Zeiler et al. 2010). Then a convolutional layer with dropout (dropout probability 0.5) is used for the second last layer. In the last layer, we design 3 anchors, 5, 10, 20, for scale references which are designed based on the distribution of nodule sizes. For each anchor, there are 5 parts in the loss function, classification loss $L_{cls}$ for whether the current box is a nodule or not, regression loss $L_{reg}$ for nodule coordinates $x, y, z$ and nodule size $d$.

If an anchor overlaps a ground truth bounding box with the intersection over union (IoU) higher than 0.5, we consider it as a positive anchor ($p^* = 1$). On the other hand, if an anchor has IoU with all ground truth boxes less than 0.02, we consider it as a negative anchor ($p^* = 0$). The multi-task loss function for the anchor $i$ is defined as

$$L(p_i, t_i) = \lambda L_{cls}(p_i, p^*_i) + p^*_i L_{reg}(t_i, t^*_i),$$

where $p_i$ is the predicted probability for current anchor $i$ being a nodule, $t_i$ is the regression predicted relative coordinates for nodule position, which is defined as

$$t_i = (\frac{x - x_a}{d_a}, \frac{y - y_a}{d_a}, \frac{z - z_a}{d_a}, \log(\frac{d^*}{d_a})),$$

where $(x, y, z, d)$ are the predicted nodule coordinates and diameter in the original space, $(x_a, y_a, z_a, d_a)$ are the coordinates and scale for the anchor $i$. For ground truth nodule position, it is defined as

$$t^*_i = (\frac{x^* - x_a}{d_a}, \frac{y^* - y_a}{d_a}, \frac{z^* - z_a}{d_a}, \log(\frac{d^*}{d_a})),$$

where $(x^*, y^*, z^*, d^*)$ are nodule ground truth coordinates and diameter. The $\lambda$ is set as 0.5. For $L_{cls}$, we used binary cross entropy loss function. For $L_{reg}$, we used smooth $l_1$ regression loss function (Girshick 2015).

Gradient Boosting Machine with 3D Dual Path Net Feature for Nodule Classification

Inspired by the success of dual path network on the ImageNet (Chen et al. 2017; Deng et al. 2009), we design a 3D deep dual path network framework for lung CT nodule classification in Fig. 4.

Dual path connection benefits both from the advantage of residual learning and that of dense connection (He et al. 2016; Huang et al. 2016a). The shortcut connection in residual learning is an effective way to eliminate gradient vanishing phenomenon in very deep networks. From a learned feature weights sharing perspective, residual learning enables feature reuse, while dense connection has an advantage of keeping exploiting new features (Chen et al. 2017). And densely connected network has fewer parameters than residual learning because there is no need to relearn redundant feature maps. The assumption of dual path connection is that there might exist some redundancy in the exploited feature maps. The dual path connection splits its feature maps into two parts. One part, $F(x)[d :]$, is used for residual learning, the other part, $F(x)[d :]$, is used for dense connection as shown in Fig. 3. Here $d$ is the hyper-parameter for deciding how many new features to be exploited. The dual path connection can be formulated as

$$y = G(\{F(x)[d :]; F(x)[d :] + x\}),$$

where $y$ is the feature map for dual path connection, $G$ is used as ReLU activation function, $F$ is convolutional layer functions, and $x$ is the input of dual path connection block. Dual path connection integrates the advantages of the two advanced frameworks, residual learning for feature reuse and dense connection for keeping exploiting new features, into a unified structure, which obtains success on the ImageNet dataset.

For CT data, advanced method should be effective to extract 3D volume feature (Yan et al. 2016). We design a 3D deep dual path network for the 3D CT lung nodule classification in Fig. 4. We firstly crop CT data centered at predicted nodule locations with size $32 \times 32 \times 32$. After that, a convolutional layer is used to extract features. Then 4 dual path blocks are employed to learn higher level features. Lastly, the 3D average pooling and binary logistic regression layer are used for benign or malignant diagnosis.

The 3D deep dual path network can be used as a classifier for nodule diagnosis directly. And it can also be employed to learn effective features. We construct feature by...
with the center as the detected center and size as $32 \times 32 \times 32$. The detected nodule size is kept for the classification model as a part of features. The 3D deep dual path network is employed to extract features. We use the GBM and construct features to conduct diagnosis for the detected nodules. For CT pixel feature, we use the cropped size as $16 \times 16 \times 16$ and center as the detected nodule center in the experiments. For patient-level diagnosis, if one of the detected nodules is positive (cancer), the patient is a cancer patient, and if all the detected nodules are negative, the patient is a negative (non-cancer) patient.

**Experiments**

We conduct extensive experiments to validate the DeepLung system. We perform 10-fold cross validation using the detector on LUNA16 dataset. For nodule classification, we use the LIDC-IDRI annotation, and employ the LUNA16’s patient-level dataset split. Finally, we also validate the whole system based on the detected nodules both on patient-level diagnosis and nodule-level diagnosis.

In the training, for each model, we use 150 epochs in total with stochastic gradient descent optimization and momentum as 0.9. The used batch size is relied on the GPU memory. We use weight decay as $1 \times 10^{-4}$. The initial learning rate is 0.01, 0.001 at the half of training, and 0.0001 after the epoch 120.

**Datasets**

LUNA16 dataset is a subset of the largest public dataset for pulmonary nodules, the LIDC-IDRI dataset (Armato et al. 2011). LUNA16 dataset only has the detection annotations, while LIDC-IDRI contains almost all the related information for low-dose lung CTs including several doctors’ annotations on nodule sizes, locations, diagnosis results, nodule texture, nodule margin and other informations. LUNA16 dataset removes CTs with slice thickness greater than 3mm, slice spacing inconsistent or missing slices from LIDC-IDRI dataset, and explicitly gives the patient-level 10-fold cross validation split of the dataset. LUNA16 dataset contains 888 low-dose lung CTs, and LIDC-IDRI contains 1,018 low-dose lung CTs. Note that LUNA16 dataset removes the annotated nodules of size smaller than 3mm.

For nodule classification, we extract nodule annotations from LIDC-IDRI dataset, find the mapping of different doctors’ nodule annotations with the LUNA16’s nodule annotations, and get the ground truth of nodule diagnosis by taking different doctors’ diagnosis averagely (Do not count the 0 score for diagnosis, which means N/A.). If the final average score is equal to 3 (uncertain about malignant or benign), we remove the nodule. For the nodules with score greater than 3, we label them as positive. Otherwise, we label them as negative. For doctors’ annotations, we only keep those of four doctors who label most of the nodules for comparison. We only keep the CTs within LUNA16 dataset, and use the same cross validation split as LUNA16 for classification.

**Preprocessing**

We firstly clip the raw data into $[-1200, 600]$. Secondly, we transform the range linearly into $[0, 1]$. Thirdly, we use the

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**DeepLung System: Fully Automated Lung CT Cancer Diagnosis**

The DeepLung system includes the nodule detection using the 3D Faster R-CNN, and nodule classification using gradient boosting machine (GBM) with constructed feature (deep dual path features, nodule size and raw nodule CT pixels) from network structure design intrinsically.

After we get the detected nodules, we crop the nodule concatenating the learned 3D deep dual path network features (the second last layer, 2,560 dimension), nodule size, and raw 3D cropped nodule pixels. Given complete and effective features, gradient boosting machine (GBM) is a superior method to build an advanced classifier from these features (Friedman 2001). We validate the feature combining nodule size with raw 3D cropped nodule pixels, employ GBM as a classifier, and obtain 86.12% test accuracy averagely. Lastly, we employ GBM with the constructed feature and achieve the best diagnosis performance.

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We firstly clip the raw data into $[-1200, 600]$. Secondly, we transform the range linearly into $[0, 1]$. Thirdly, we use the
LUNA16’s given segmentation ground truth and remove the useless background. It is worth visualizing the processed data during processing because some segmentations are imperfect, and need to be removed. After that, the data can be processed by the deep networks.

**DeepLung for Nodule Detection**

We train and evaluate the detector on LUNA16 dataset following 10-fold cross validation with given patient-level split. In the training, we use the flipping, randomly scale from 0.75 to 1.25 for the cropped patches to augment the data. The evaluation metric, FROC, is the average recall rate at the average number of false positives as 0.125, 0.25, 0.5, 1, 2, 4, 8 per scan, which is the official evaluation metric for LUNA16 dataset.

The FROC performance on LUNA16 is visualized in Fig. 5. The solid line is interpolated FROC based on true prediction, and the dash lines are upper bound and lower bound for the bootstrapped FROC performance. The 3D Faster R-CNN achieves 83.4% FROC without any false positive nodule reduction stage, which is comparable with 83.9% for the best method, which uses two-stage training (Dou et al. 2017).

**DeepLung for Nodule Classification**

We validate the nodule classification performance of the DeepLung system on the LIDC-IDRI dataset with the LUNA16’s split principle, 10-fold patient-level cross validation. There are 1,004 nodules left and 450 nodules are positive. In the training, we firstly pad the nodules of size $32 \times 32 \times 32$ into $36 \times 36 \times 36$, randomly crop $32 \times 32 \times 32$ from the padded data, horizontal flip, vertical flip, z-axis flip the data for augmentation, randomly set $4 \times 4 \times 4$ patch as 0, and normalize the data with the mean and std. obtained from training data. The total number of epochs is 1,050. The learning rate is 0.01 at first, then became 0.001 after epoch 525, and turned into 0.0001 after epoch 840. Due to time and resource limitation for training, we use the fold 1, 2, 3, 4, 5 for test, and the final performance is the average performance on the five test folds. The nodule classification performance is concluded in Table 1.

From the table 1, our 3D deep dual path network (DPN) achieves better performance than that of multi-scale CNN (Shen et al. 2015) because of the power of 3D structure and deep dual path network. Because of the power of gradient boosting machine (GBM), GBM with nodule size and raw nodule pixels with crop size as $16 \times 16 \times 16$ achieves comparable performance as multi-scale CNN (Shen et al. 2015). Finally, we construct feature with 3D deep dual path network features, nodule size and raw nodule pixels, and obtain 90.44% accuracy.

**Compared with Experienced Doctors on Their Individually Confident Nodules** We compare our predictions with those of four experienced doctors on their individually confident nodules (with individual score not as 3). Note that about 1/3 nodules are labeled as 3 for each doctor. Comparison results are concluded in Table 2.

From Table 2, these doctors’ confident nodules are easy to be diagnosed nodules from the performance comparison between our model’s performances in Table 1 and Table 2. To our surprise, the average performance of our model is 1.5% better than that of experienced doctors. In fact, our model’s performance is better than 3 out of 4 doctors (doctor 1, 3, 4) on the nodule diagnosis task. The result validates deep network surpasses human-level performance for image classification (He et al. 2016), and the DeepLung is better suited for nodule diagnosis than experienced doctors.

**DeepLung for Fully Automated Lung CT Cancer Diagnosis**

We also validate the DeepLung for fully automated lung CT cancer diagnosis on the LIDC-IDRI dataset with the same protocol as LUNA16’s patient-level split. Firstly, we employ the 3D Faster R-CNN to detect suspicious nodules. Then we retrain the model from nodule classification model on the detected nodules dataset. If the center of detected nodule is

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### Table 1: Nodule classification comparisons on LIDC-IDRI dataset.

| Models          | 3D DPN | Nodule Size +Pixel+GBM | All feat. +GBM | m-CNN |
|-----------------|--------|------------------------|----------------|-------|
| Acc. (%)        | 88.74  | 86.12                  | 90.44          | 86.84 |

### Table 2: Nodule-level diagnosis accuracy (%) between nodule classification subnetwork in DeepLung and experienced doctors on doctor’s individually confident nodules.

| Doctors | Dr 1 | Dr 2 | Dr 3 | Dr 4 | Average |
|---------|------|------|------|------|---------|
| DeepLung | 93.55 | 93.69 | 91.82 | 86.03 | 91.25   |
| Doctors  | 93.44 | 93.30 | 93.19 | 90.89 | 92.74   |

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Figure 5: Sensitivity (Recall) rate with respect to false positives per scan. The FROC (average recall rate at the false positives as 0.125, 0.25, 0.5, 1, 2, 4, 8) is 83.4%. The 3D Faster R-CNN has a recall rate 94.6% for all the nodules. The dash lines are lower bound and upper bound FROC for 95% confidence interval using bootstrapping with 1,000 bootstraps (Setio et al. 2017). The solid line is the interpolated FROC based on prediction.
within the ground truth positive nodule, it is a positive nodule. Otherwise, it is a negative nodule. Through this mapping from the detected nodule and ground truth nodule, we can evaluate the performance and compare it with the performance of experienced doctors. We adopt the test fold 1, 2, 3, 4, 5 to validate the performance the same as that for nodule classification.

Different from pure nodule classification, the fully automated lung CT nodule diagnosis relies on nodule detection. We evaluate the performance of DeepLung on the detection true positive (TP) set and detection false positive (FP) set individually in Table 3. If the detected nodule of center within one of ground truth nodule regions, it is in the TP set. If the detected nodule of center out of any ground truth nodule regions, it is in FP set. From Table 3, the DeepLung system using detected nodule region obtains 81.42% accuracy for all the detected TP nodules. Note that the experienced doctors obtain 78.36% accuracy for all the nodule diagnosis on average. The DeepLung system with fully automated lung CT nodule diagnosis still achieves above average performance of experienced doctors. On the FP set, our nodule classification subnetwork in the DeepLung can reduce 97.02% FP detected nodules, which guarantees that our fully automated system is effective for the lung CT cancer diagnosis.

**Compared with Experienced Doctors on Their Individually Confident CTs** We employ the DeepLung for patient-level diagnosis further. If the current CT has one nodule that is classified as positive, the diagnosis of the CT is positive. If all the nodules are classified as negative for the CT, the diagnosis of the CT is negative. We evaluate the DeepLung on the doctors’ individually confident CTs for benchmark comparison in Table 4.

From Table 4, DeepLung achieves 81.41% patient-level diagnosis accuracy. The performance is 99% of the average performance of four experienced doctors, and the performance of DeepLung is better than that of doctor 4. Thus DeepLung can be used to help improve some doctors’ performance, like that of doctor 4, which is the goal for computer aided diagnosis system. We also employ Kappa coefficient, which is a common approach to evaluate the agreement between two raters, to test the agreement between DeepLung and the ground truth patient-level labels (Landis and Koch 1977). For comparison, we valid those of 4 individual doctors on their individual confident CTs. The Kappa coefficient of DeepLung is 63.02%, while the average Kappa coefficient of doctors is 64.46%. It shows the predictions of DeepLung are of good agreement with ground truths for patient-level diagnosis, and are comparable with those of experienced doctors.

**Discussion**

In this section, we are trying to explain the DeepLung by visualizing the nodule detection and classification results.

**Nodule Detection**

We randomly pick nodules from test fold 1 and visualize them in the first row and third row in Fig. 6. Detected nodules are visualized in the second, fourth and fifth row. The numbers below detected nodules are with the format (ground truth slice number-detected slice number-detection probability). The fifth row with red detection probabilities are the false positive detections. Here we only visualize the slice of center $z$. The yellow circles are the ground truth nodules or detected nodules respectively. The center of the circle is the nodule center and the diameter of the circle is relative to the nodule size.

From the first four rows in Fig. 6, we observe the 3D Faster R-CNN works well for the nodules from test fold 1. The detection probability is also very high for these nodules. From the detected false positive nodules in the last row, we can find these false positives are suspicious to nodules visually if we observe them carefully. And the sizes of these false positive nodules are very small. Note that the nodules less than 3mm are removed in the ground truth annotations.

**Nodule Classification**

We also visualize the nodule classification results from test fold 1 in Fig. 7. The nodules in the first four rows are those the DeepLung predicted right, but some doctors predicted wrong. The first number is the DeepLung predicted malignant probability. The second number is which doctor predicted wrong. If the probability is large than 0.5, it is a malignant nodule. Otherwise, it is a benign nodule. The rest nodules are those the DeepLung predicted wrong with red numbers indicating the DeepLung predicted malignant probability. For an experienced doctor, if a nodule is big and has irregular shape, it has a high probability to be a malignant nodule.

From the first 4 rows in Fig. 7, we can observe that doctors mis-diagnose some nodules. The reason is that, humans are not good at processing 3D CT data, which is of low signal to noise ratio. In fact, even for high quality 2D natural image, the performance of deep network surpasses that of humans (He et al. 2016). From the doctors’ individual experience point of view, some doctors intend to be optimistic to some nodules, and some doctors intend to be pessimistic to them. And they can just observe one slice each time. Some irregular boundaries are vague. The machine learning based methods can learn these complicated rules and high dimensional features from these doctors' annotations, and avoid such bias.

**Table 3: Comparison between DeepLung’s nodule classification on all detected nodules and doctors on all nodules.**

| Method  | TP Set | FP Set | Doctors |
|---------|--------|--------|---------|
| Acc. (%)| 81.42  | 97.02  | 74.05-82.67 |

**Table 4: Patient-level diagnosis accuracy(%) between DeepLung and experienced doctors on doctor’s individually confident CTs.**

| Doctors | Dr 1   | Dr 2   | Dr 3   | Dr 4   | Average |
|---------|--------|--------|--------|--------|---------|
|         | 83.03  | 85.65  | 82.75  | 77.80  | 82.31   |
| DeepLung| 81.82  | 80.69  | 78.86  | 84.28  | 81.41   |
Figure 6: Visualization of central slices for nodule ground truths and detection results. We randomly choose nodules (the first row and third row) from test fold 1. Detection results are shown in the second row and fourth row. The numbers below detected nodules are with the format (ground truth slice number-detected slice number-detection probability). The last row shows the false positive detected nodules and detection probabilities. Note that the ground truth nodules are of diameter greater than 3mm. The DeepLung performs well for detection.

The rest nodules with red numbers are the nodules the DeepLung predicted wrong for test fold 1 in Fig. 7. If the predicted malignant probability is larger than 0.5, the ground truth label is benign. Otherwise, the ground truth label is malignant. From the central slices, we can hardly decide the DeepLung’s predictions are wrong. Maybe the DeepLung cannot find some weak irregular boundaries as for the bottom right 4 nodules, which is the possible reason why the DeepLung predicts them as benign. From the above analysis, the DeepLung can be considered as a tool to assist the diagnosis for doctors. Combing the DeepLung and doctor’s own diagnosis is an effective way to improve diagnosis accuracy.

**Conclusion**

In this work, we propose a fully automated lung CT cancer diagnosis system, DeepLung. DeepLung consists two parts, nodule detection and classification. For nodule detection, we design a 3D Faster R-CNN with U-net-like encoder-decoder structure to detect suspicious nodules. Then we input the detected nodules into the nodule classification network. The 3D deep dual path network is designed to extract features. Further, gradient boosting machine with different features combined together to achieve the state-of-the-art performance, which surpasses that of experienced doctors. Extensive experimental results on public available large-scale datasets, LUNA16 and LIDC-IDRI datasets, demonstrate the superior performance of the DeepLung.

In the future, we will develop more advanced nodule detection method to further boost the performance of DeepLung. Further, a unified framework for automated lung CT cancer diagnosis is expected for nodule detection and classification. Integrating multi-modality data, such as electronic medical report history, into the DeepLung is another
direction to enhance the diagnosis performance.

References

Aerts, H. J.; Velazquez, E. R.; Leijenaar, R. T.; Parmar, C.; Grossmann, P.; Cavalho, S.; Bussink, J.; Monshouwer, R.; Raike-Kains, B.; Rietveld, D.; et al. 2014. Decoding tumour phenotype by non-invasive imaging using a quantitative radiomics approach. *Nature communications* 5.

Armato, S. G.; McLennan, G.; Bidaut, L.; McNitt-Gray, M. F.; preprint arXiv:1708.03867

online sample filtering and hybrid-loss residual learning. 2017. Automated pulmonary nodule detection via 3d convnets with Dou, Q.; Chen, H.; Jin, Y.; Lin, H.; Qin, J.; and Heng, P.-A. arXiv preprint arXiv:1706.04303

nodule detection in computed tomography images using deep con-

Ding, J.; Li, A.; Hu, Z.; and Wang, L. 2017. Accurate pulmonary nodule detection in computed tomography images using deep convolu-

Ding, J.; Li, A.; Hu, Z.; and Wang, L. 2017. Accurate pulmonary nodule detection in computed tomography images using deep convolu-

Dou, Q.; Chen, H.; Jin, Y.; Lin, H.; Qin, J.; and Heng, P.-A. 2017. Automated pulmonary nodule detection via 3d convnets with online sample filtering and hybrid-loss residual learning. arXiv preprint arXiv:1706.04303.

El-Baz, A.; Nitzken, M.; Khalifa, F.; ElGamal, A.; Gimelfarb, G.; El-Ghar, M. A. 2011. 3d shape analysis for early diagnosis of malignant lung nodules. In Biennial International Conference on Information Processing in Medical Imaging, 772–783. Springer.

Friedman, J. H. 2001. Greedy function approximation: a gradient boosting machine. *Annals of statistics* 1189–1232.

Girshick, R. 2015. Fast r-cnn. In ICCV, 1440–1448.

Han, F.; Zhang, G.; Wang, H.; Song, B.; Lu, H.; Zhao, D.; Zhao, H.; and Liang, Z. 2013. A texture feature analysis for diagnosis of pulmonary nodules using ldc-idi database. In *Medical Imaging, Physics and Engineering (ICMIME), 2013 IEEE International Conference on*, 14–18. IEEE.

He, K.; Zhang, X.; Ren, S.; and Sun, J. 2016. Deep residual learning for image recognition. In *CVPR*, 770–778.

Huang, G.; Liu, Z.; Weinberger, K. Q.; and van der Maaten, L. 2016a. Densely connected convolutional networks. arXiv preprint arXiv:1608.06993.

Huang, J.; Rathod, V.; Sun, C.; Zhu, M.; Korattikara, A.; Fathi, A.; Fischer, I.; Wojna, Z.; Song, Y.; Guadarrama, S.; et al. 2016b. Speed/accuracy trade-offs for modern convolutional object detectors. arXiv preprint arXiv:1611.10012.

Hussein, S.; Cao, K.; Song, Q.; and Bagci, U. 2017. Risk stratification of lung nodules using 3d cnn-based multi-task learning. In *IPMI*, 249–260. Springer.

Jacobs, C.; van Rikxoort, E. M.; Twellmann, T.; Scholten, E. T.; de Jong, P.; A.; Kuhnigk, J.-M.; Oudkerk, M.; de Koning, H. J.; Prokop, M.; Schaefel-Prokop, C.; et al. 2014. Automatic detection of subsolid pulmonary nodules in thoracic computed tomography images. *Medical image analysis* 18(2):374–384.

Landis, J. R., and Koch, G. G. 1977. The measurement of observer agreement for categorical data. *biometrics* 159–174.

Long, J.; Shelhamer, E.; and Darrell, T. 2015. Fully convolutional networks for semantic segmentation. In *CVPR*, 3431–3440.

Lopez Torres, E.; Fiorina, E.; Pennazio, F.; Peroni, C.; Saletta, M.; Camarlinghi, N.; Fantacci, M.; and Cerello, P. 2015. Large scale validation of the m5l lung cad on heterogeneous ct datasets. *Medical physics* 42(4):1477–1489.

Murphy, K.; van Ginneken, B.; Schilham, A. M.; De Hoop, B.; Gietema, H.; and Prokop, M. 2009. A large-scale evaluation of automatic pulmonary nodule detection in chest ct using local image features and k-nearest-neighbour classification. *Medical image analysis* 13(5):757–770.

Ren, S.; He, K.; Girshick, R.; and Sun, J. 2015. Faster r-cnn: Towards real-time object detection with region proposal networks. In NIPS, 91–99.

Ronneberger, O.; Fischer, P.; and Brox, T. 2015. U-net: Convolutional networks for biomedical image segmentation. In *MICCAI*, 234–241. Springer.

Setio, A. A. A.; Ciompi, F.; Litjens, G.; Gerke, P.; Jacobs, C.; van Riel, S. J.; Wille, M. M. W.; Naqibullah, M.; Sánchez, C. I.; and van Ginneken, B. 2016. Pulmonary nodule detection in ct images: false positive reduction using multi-view convolutional networks. *IEEE transactions on medical imaging* 35(5):1160–1169.

Setio, A. A. A.; Traverso, A.; de Bel, T.; Berens, M. S.; van den Bogaard, C.; Cerello, P.; Chen, H.; Dou, Q.; Fantacci, M. E.; Geurts, B.; et al. 2017. Validation, comparison, and combination of algorithms for automatic detection of pulmonary nodules in computed tomography images: the Luna16 challenge. *Medical Image Analysis* 42:1–13.

Shen, W.; Zhou, M.; Yang, F.; Yang, C.; and Tian, J. 2015. Multi-scale convolutional neural networks for lung nodule classification. *IPMI*, 588–599. Springer.

Shen, W.; Zhou, M.; Yang, F.; Dong, D.; Yang, C.; Zang, Y.; and Tian, J. 2016. Learning from experts: Developing transferable deep features for patient-level lung cancer prediction. In *MICCAI*, 124–131. Springer.

Suzuki, K.; Li, F.; Sone, S.; and Doi, K. 2005. Computer-aided diagnostic scheme for distinction between benign and malignant nodules in thoracic low-dose ct by use of massive training artificial neural network. *IEEE Transactions on Medical Imaging* 24(9):1138–1150.

Way, T. W.; Hadjiiski, L. M.; Sahiner, B.; Chan, H.-P.; Cascade, P. N.; Kazerooni, E. A.; Bogot, N.; and Zhou, C. 2006. Computer-aided diagnosis of pulmonary nodules on ct scans: Segmentation and classification using 3d active contours. *Medical Physics* 33(7):2323–2337.

Xie, Y.; Pang, J.; Qi, H.; Zhu, Y.; Bai, C.; Geng, X.; Liu, M.; Terzopoulos, D.; and Ding, X. 2016. Classification of lung nodule malignancy risk on computed tomography images using convolutional neural network: A comparison between 2d and 3d strategies. In *ACCV*, 91–101. Springer.

Zeiler, M. D.; Krishnan, D.; Taylor, G. W.; and Fergus, R. 2010. Deconvolutional networks. In *CVPR*, 2528–2535. IEEE.

Zhu, W., and Xie, X. 2016. Adversarial deep structural networks for mammographic mass segmentation. arXiv preprint arXiv:1612.05970.

Zhu, W.; Lou, Q.; Yang, Y. S.; and Xie, X. 2017. Deep multi-instance networks with sparse label assignment for whole mammogram classification. In *MICCAI*, 603–611. Springer.