

DIABETIC RETINOPATHY CLASSIFICATION USING TRANSFER LEARNING AND EXUDATES DETECTION USING FASTER-RCNN

¹Manojkumar S B, ²Dr. H S Sheshadri

¹Research Scholar, ²Professor

¹Dept. of Electronics

¹PET Research Foundation, PESCE, Mandya, India

Abstract : Here we proposed a deep learning method for detecting diabetic retinopathy (DR). DR is one of the common for prolong standing of diabetics later it leads to vision loss. In our early papers we worked with MATLAB software and applied various algorithms like K-means cluster algorithm, SVM and Random forest algorithm. Now we proposed transfer learning process (inception v3) method of tensor flow software by adding the region proposal networks (RPN) after the global averaging pooling layer of the convolutional networks (CNN) for classification of diabetic retinopathy and Faster R-CNN for edudates identification and extraction. The proposed model can localize the regions of a retina image to show the specific region of interest in terms of its severity level. Our opinion is this deep learning method is highly desired for diabetic retinopathy detection because in practice, users are not only interested with high prediction performance, but also keen to understand the insights of DR detection and transfer learning model works. In the experiments conducted on a large scale of retina image dataset collected from kaggle, a number of pre-trained models are available which perform object classification from an image of the object which is given as an input. However, we can retrain the pre-trained models for the set of classes of interest and in this work; one such model has been retained to measure the severity of Diabetic Retinopathy in eye images (data has been provided by eyepacs as color fundus images) on the scale of 0-5 we show that the proposed faster R-CNN model can achieve high performance on DR detection compared with the state-of-the-art while achieving the merits of providing the RPN to highlight the salient regions of the input image and we achieved 83.3% raw image accuracy A convolutional neural network classifier engineered from Inception V3 network which is trained for ImageNet, for 5-class severity classification performed best with an accuracy of 83.3%.

IndexTerms - Diabetic Retinopathy, Faster R-CNN, Transfer Learning, RPN

I. INTRODUCTION

Diabetes is a common disease in both developed and developing countries, and nearly 98 million people in India may have this disease around 2030[14]. Prolong years of diabetes affecting to retina leads to Diabetic retinopathy (DR). Basically, DR affects blood vessels of retina. DR is one of the leading cause of blindness for middle age adults in the world today [9], and around half of Indian's with diabetes have this disease to some extent. A well-known challenge for DR is that during the early stages it has no warning sign, even for diabetic macular edema. Thus, it is very much important that DR can be detected before it cross early stage. Unfortunately, in present situation DR detection in early stage is nearly infeasible. It requires an ophthalmologist to analyze digital color fundus photographs of retina, and DR is identified by locating the lesions associated with vascular abnormalities affected by diabetes, figure 1 shows the Optic disc and abnormal findings I the eye fundus caused by the diabetic retinopathy. Even though the present solution of ophthalmologist identifying DR disease is effective, but it is time consuming and chances of false positives and highly relies on the expertise of well training practitioners. In order to overcome this problem, in the past few years considerable efforts have been put on developing an automated solution for DR detection.

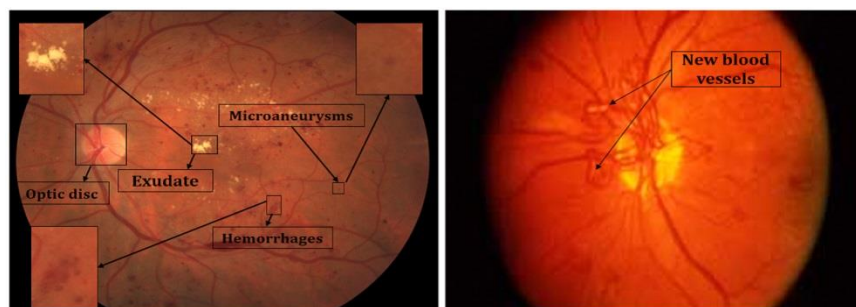


Figure 1: Optic disc and abnormal findings in the eye fundus caused by the diabetic retinopathy.

Among them automated detection of DR consists of two parts: feature extraction and detection/prediction algorithm [10]. Feature extraction is mainly focused as standard machine learning algorithms can be directly used as the detection/prediction algorithm. It is effective to some extent but also suffer from several imperfections. Primarily, as reviewed in this Section, the extracted features are all hand crafted features. Therefore, these features are mainly depend on the parameters of the tools used in feature extraction and they are sensitive to the quality of fundus photography, like object view, artifacts, exposedness, noise, out-of-focus, etc. Secondly, feature extraction may be a separate task instead of embedded into the complete DR detection framework. The above mentioned features extraction methods can be considered as the universal image feature extraction methods that are applicable to

most computer vision tasks, and they are not obligate to the specific task, e.g., In this paper DR detection task is to be considered. It is worth noting that color fundus photography is more challenging than the standard scene or object images that most image feature extraction methods were developed based on, since the key signals are often very tiny in fundus photography and they often look separate from noise and artifacts. Thus, these two challenges make it highly desirable to develop a systematically feature representation approach to effectively characterize the novel approach of features that to be with respect to the DR detection.

II. RELATED WORK

Now, the convolutional neural network (CNN) has achieved tremendous success in computer vision area. It can model high-level abstractions in data relative to specific prediction task [5][13] In CNN, a multiple layers network is built up for automating feature design. Specifically, each and every layer in deep architecture performs a non-linear transformation on the response of the previous layer, so that the data are represented by a hierarchy of features from low-level to high-level. The main attribute of the CNN is conducting different processing units of the signals. Then, the deep learning architecture allows multiple layers of these processing units to be stacked, so that this deep learning model can characterize the salience of signals in different scales. Also, in CNN, feature extraction and prediction algorithm are merged as a single model. Thus, the extracted features own more discriminative power, since the entire CNN model is trained under the supervision of output labels. Briefly telling, the features extracted by the CNN are task dependent and non-handcrafted.

In this paper, we also adopt CNN as the key predictive algorithm, but aim to develop a more efficient CNN architecture that is particularly useful for large-scale dataset. Specifically, the CNN we built has no fully connected layer and only have convolutional and pooling layers. This setting significantly reduces the number of parameters (fully-connected layers often bring more parameters than convolutional layers in the conventional CNN) and provides better conditions for interpretability of neural network as presented below. We show in experiments that with less parameter and no fully-connected layers the proposed CNN architecture can achieve the comparative prediction performance. The key advantage of the proposed network structure is that it can provide a regression activation maps (RAM) of input image to show the contribution score of each pixel of input image for DR detection task. This RAM output, to some extent, somehow mitigates the well-known un-interpretable shortcoming of CNN as a black box method. We believe that this RAM output make the proposed solution more self-explained and can motivate the practitioners to trace the cause of the disease for every patient.

III. ALGORITHM

Description:

In the first phase we classify the fundus retina image into proliferative, non-proliferative mild, non-proliferative moderate and non-proliferative severe. For classification task we use inception V3 model and perform training using transfer training. In the second phase, we detect Exudates and localize exudates. Detection of exudates is achieved by Faster RCNN network.

IV. METHODOLOGY

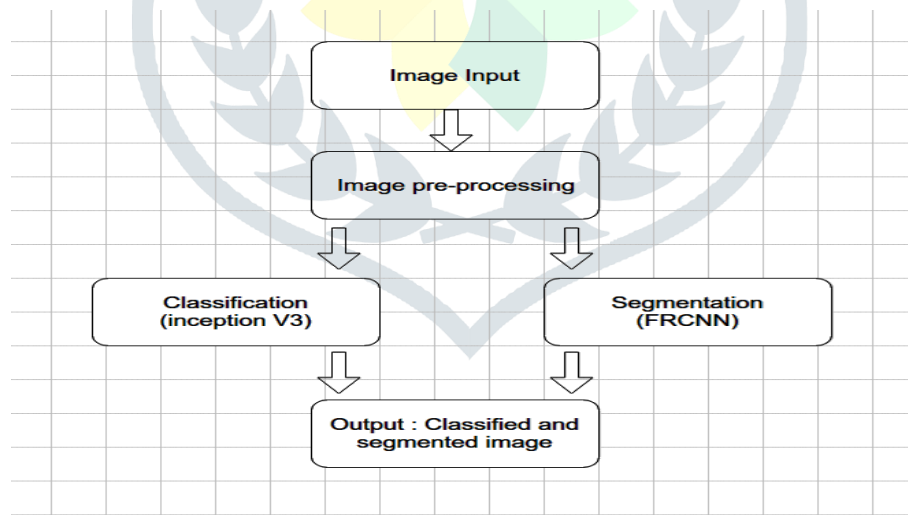


Figure 2: Block Diagram

Figure 2 shows the block diagram of diabetic retinopathy classification using transfer learning process.

A. Image preprocessing

All images were converted to a hierarchical data format for preprocessing, data augmentation, and training. Preprocessing involved several steps [7]: images were cropped using Otsu's method to isolate the circular colored image of the retina. Images were normalized by subtracting the minimum pixel intensity from each channel and dividing by the mean pixel intensity to represent pixels in the range 0 to 1. Figure 3 shows the image preprocessing stage.

B. Data Preprocessing

- Rescale input images to 300x300 or 500x500 pixels.
- Subtract the local average color, and the local average gets mapped to 50% gray scale.
- Clipped the images to 90% size to remove the "boundary effects".

C. Data Augmentation

This step is to increase the variability of training data for enhancing performance.

- Randomly scale the size of images by $\pm 10\%$.
- Randomly rotated by between 0 and 360 degrees.
- Randomly skew the images by ± 0.2 .
- For testing, the images are just rotated randomly.

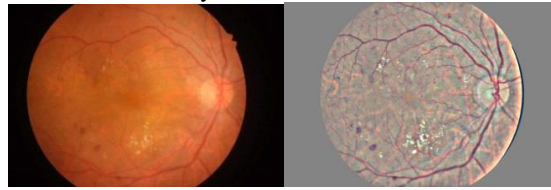


Figure 3 :(a) Input image (b) Input image after preprocessing

D. Classification

1. Transfer learning

The first phase analyzes all the images on disk and calculates and caches the bottleneck values for each of them [1]. This penultimate layer has been trained to output a set of values that's good enough for the classifier to use to distinguish between all the classes it's been asked to recognize. That means it has to be a meaningful and compact summary of the images, since it has to contain enough information for the classifier to make a good choice in a very small set of values. The reason our final layer retraining can work on new classes is that it turns out the kind of information needed to distinguish between all the 1,000 classes in ImageNet[6] is often also useful to distinguish between new kinds of objects.

2. Training

Once the bottlenecks are complete, the actual training of the top layer of the network begins. You'll see a series of step outputs, each one showing training accuracy, validation accuracy, and the cross entropy. The training accuracy shows what percent of the images used in the current training batch were labeled with the correct class. The validation accuracy is the precision on a randomly-selected group of images from a different set. The key difference is that the training accuracy is based on images that the network has been able to learn from so the network can over fit to the noise in the training data. A true measure of the performance of the network is to measure its performance on a data set not contained in the training data -- this is measured by the validation accuracy. If the train accuracy is high but the validation accuracy remains low that means the network is over fitting and memorizing particular features in the training images that aren't helpful more generally. Cross entropy is a loss function which gives a glimpse into how well the learning process is progressing. The training's objective is to make the loss as small as possible, so you can tell if the learning is working by keeping an eye on whether the loss keeps trending downwards, ignoring the short-term noise.

3. Inception v3 model

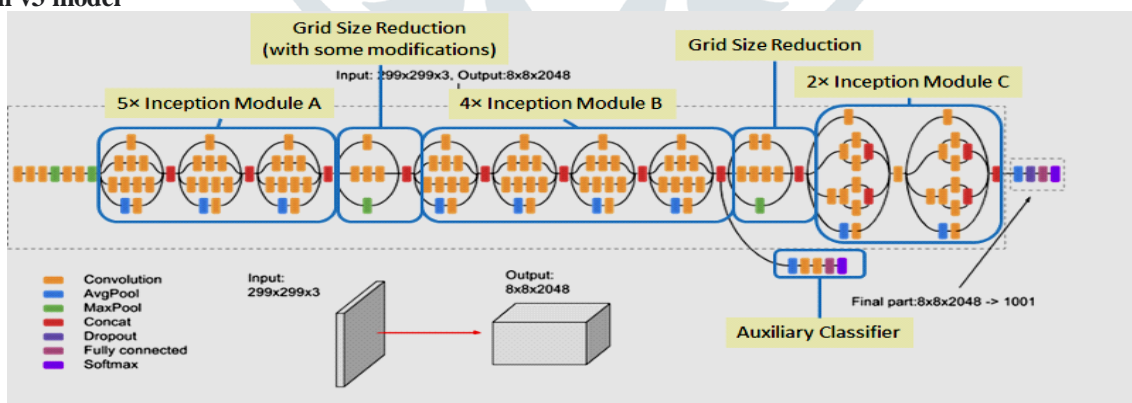


Figure 4: Inception v3 model

Figure 4 shows the Inception v3 model which is one of the widely-used image recognition model and that has been shown to attain greater than 78.1% accuracy on the ImageNet dataset. The model is the culmination of many ideas developed by multiple researchers over the years now a day.

4. Classification architecture

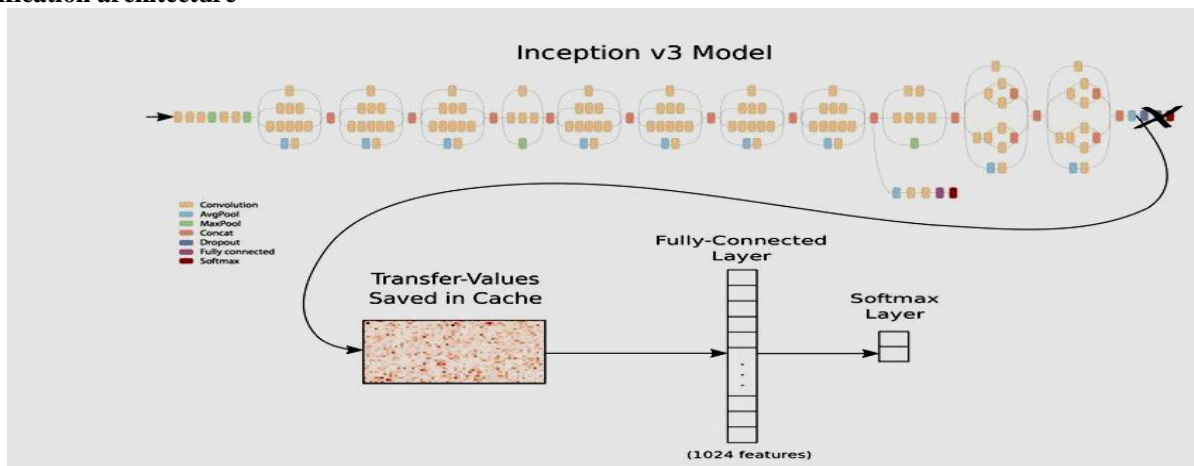


Figure 5: Classification architecture

Figure 5 shows the classification architecture of the inception v3 model. Here transfer learning allows you to retrain the final layer of an existing model, resulting in a significant decrease in not only training time, but also the size of the dataset required. One of the most famous models that can be used for transfer learning is Inception V3. As mentioned above, this model was originally trained on over a million images from 1,000 classes on some very powerful machines. Being able to retrain the final layer means that you can maintain the knowledge that the model had learned during its original training and apply it to your smaller dataset, resulting in highly accurate classifications without the need for extensive training and computational power.

V. EXUDATES DETECTION AND LOCALIZATION.

A .Faster RCNN

Faster R-CNN is composed of two modules. The first module is a deep fully convolution network that proposes regions, and the second module is the Fast R-CNN detector [2] that uses the proposed regions. The entire system is a single, unified network for object detection. Using the recently popular terminology of neural networks with ‘attention’ [31] mechanisms, the RPN module tells the Fast R-CNN module where to look.

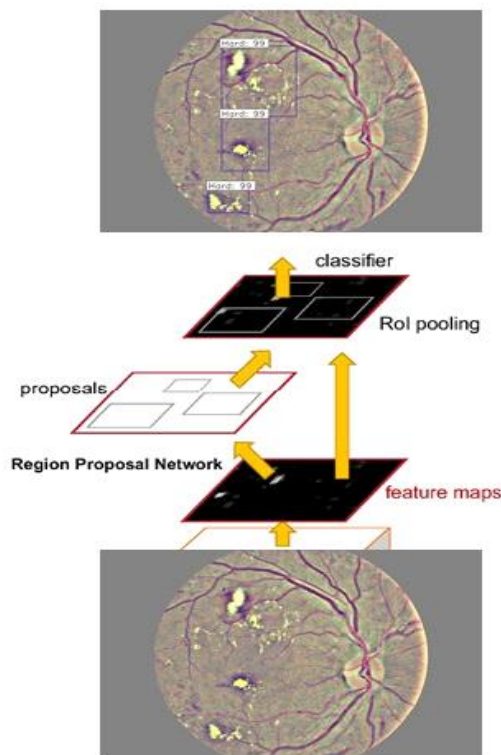


Figure 6: Faster RCNN for Exudates localization.

B. Region Proposal Networks

A Region Proposal Network (RPN) takes an image (of any size) as input and outputs a set of rectangular object proposals, each with an objectness score. To generate region proposals, we slide a small network over the convolutional feature map output by the last shared convolutional layer. This small network takes as input an $n \times n$ spatial window of the input convolutional feature map.

C. Data Preparation

We used 80 images from Indian Diabetic Retinopathy Image Dataset for manual annotation. We created a text file, each line describing path of image, Bounding box [x1, y1, x2, y2] and label name. E.g. image file path, x1, x2, x3, x4, Exudates.

D. Faster RCNN for Exudates localization

Figure 6 shows the faster RCNN for exudates localization. Here faster RCNN network receives preprocessed image as input. The network is configured to 32 ROI's, anchor box scales of {128,256,512} and anchor box ratios of {[1:1],[1:2],[2:1]}

VI RESULTS AND ANALYSIS

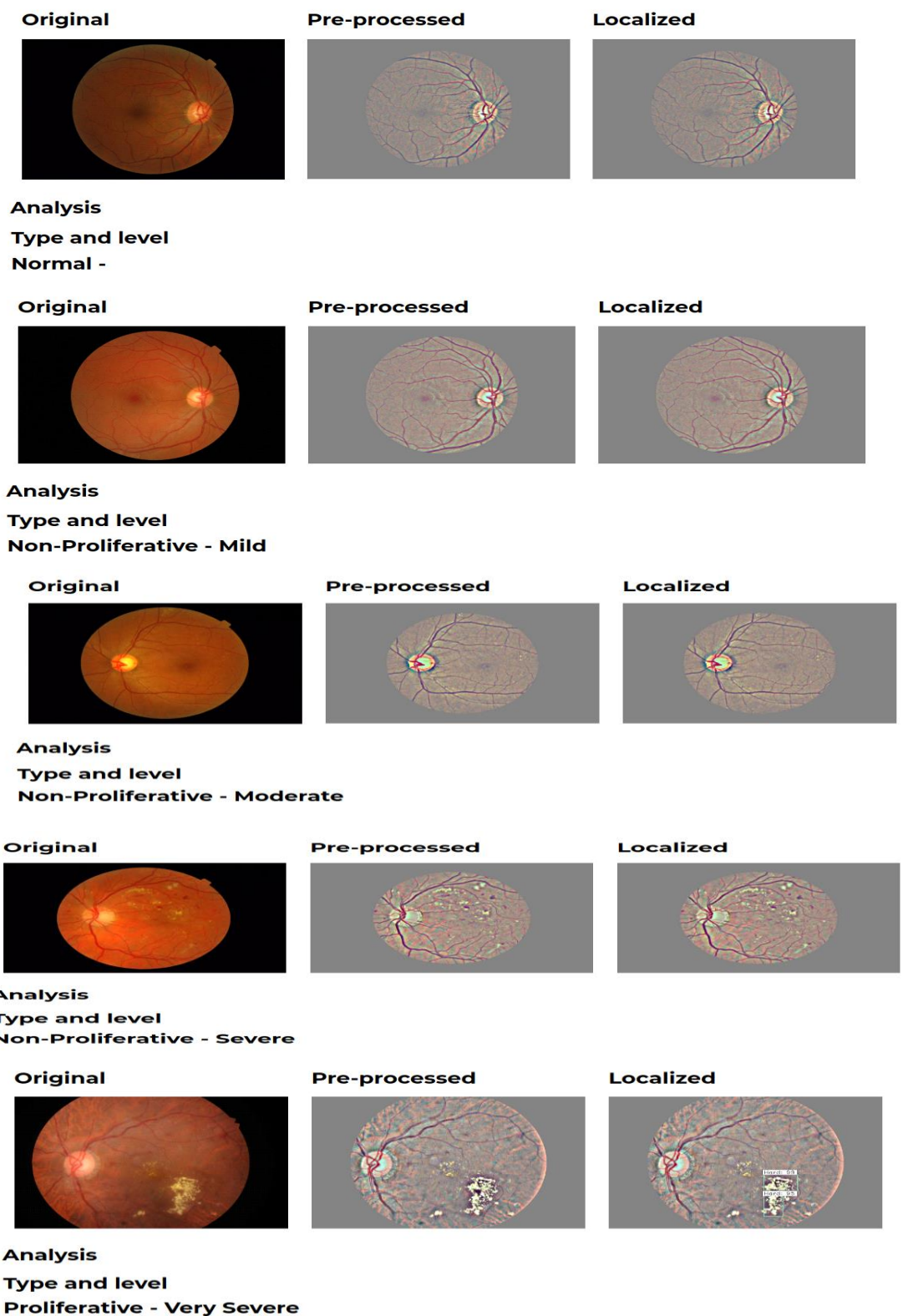


Figure 7: Normal, non-proliferative-mild, moderate, severe and proliferative-very severe results.

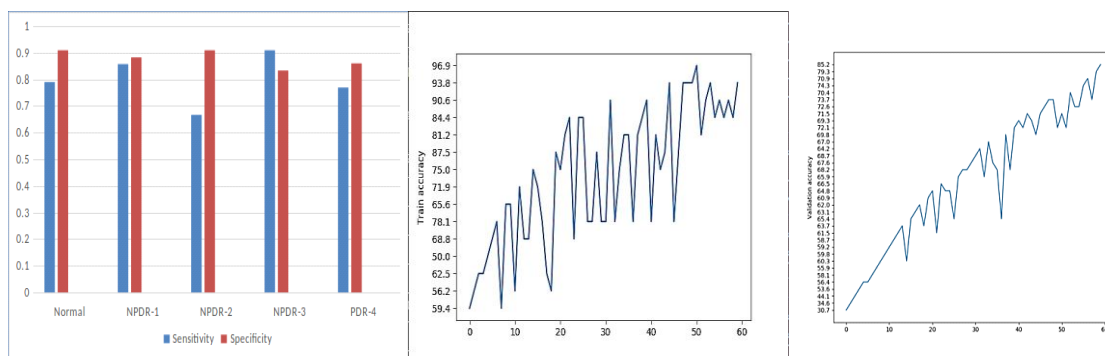


Figure 8: Plot of sensitivity and severity.

Table 1: Diabetic retinopathy raw image classification accuracy.

Diabetic retinopathy symptoms	Level	count	test data	positive results	classification accuracy
Micro aneurysms	1	< 5	1524	1260	82.67%
	2	> 5 & < 15	1482	1104	74.49%
	3	> 15	1520	1200	78.74%
Hemorrhages	1	< 5	1524	1260	82.67%
	2	> 5 & < 15	1482	1104	74.49%
	3	> 15	1520	1200	78.74%
Neovascularization	3	>1	1520	1200	78.74%
Exudates	3	> 1	54	45	83.33%

V. CONCLUSIONS

Practically, ophthalmologist can identify DR by the presence of lesions associated with the vascular abnormalities caused by the disease. While this method is more efficient, its resource demands are high. In this work, we provided a deep learning model that includes RPN and Faster RCNN to identify and analyze the exudates of diabetic retinopathy. The Faster-RCNN provide the robust interpretability of the proposed detection model in a single pass by monitoring the pathogenesis so that the proposed model can be taken as an assistant for ophthalmologist. With this attributes, the proposed methodology can still yield the competitive performance of DR detection, compared with the state-of-the-art methods. In future, we would consider extending the proposed method to other medical application problems.

VI. REFERENCES

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