Transfer Learning Through Weighted Loss Function and Group Normalization for Vessel Segmentation from Retinal Images

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Abstract—The vascular structure of blood vessels is important in diagnosing retinal conditions such as glaucoma and diabetic retinopathy. Accurate segmentation of these vessels can help in detecting retinal objects such as the optic disc and optic cup and hence determine if there are damages to these areas. Moreover, the structure of the vessels can help in diagnosing glaucoma. The rapid development of digital imaging and computer-vision techniques has increased the potential for developing approaches for segmenting retinal vessels. In this paper, we propose an approach for segmenting retinal vessels that uses deep learning along with transfer learning. We adapted the U-Net structure to use a customized InceptionV3 as the encoder and used multiple skip connections to form the decoder. Moreover, we used a weighted loss function to handle the issue of class imbalance in retinal images. Furthermore, we contributed a new dataset to this field. We tested our approach on six publicly available datasets and a newly created dataset. We achieved an average accuracy of 95.60% and a Dice coefficient of 80.98%. The results obtained from comprehensive experiments demonstrate the robustness of our approach to the segmentation of blood vessels in retinal images obtained from different sources. Our approach results in greater segmentation accuracy than other approaches.

I. INTRODUCTION

Sight is one of the most important senses for humans. It allows us to visualize and explore our surroundings and any conditions that reduce the vision impacts our ability to interact with the surroundings. Several degenerative ocular conditions have been identified, such as glaucoma and diabetic retinopathy. These conditions threaten our sense of sight by causing irreversible visual-field loss [1]. Glaucoma is is a condition that is the world’s second most prominent cause of irreversible vision loss after cataracts, accounting for 12% of cases of blindness worldwide [2].

The structure of retinal blood vessels serves an important role in the diagnosis of retinal conditions such as glaucoma [1]. For instance, one of the indicators of glaucoma is deviation from the inferior, superior, nasal, and temporal (ISNT) rule, which states that the inferior neuro-retinal rim should be thickest, followed by the superior rim, the nasal, and the temporal. Additionally, changes within the optic disc, such as the displacement of blood vessels can be used to help determine if glaucoma is present and if the disease has progressed [3].

With the advancements in the field of image processing and deep learning, it is possible to develop an automated approach for segmenting vessels with reliable precision. Such a method could help in analyzing these vessels when diagnosing retinal conditions such as glaucoma. The process would involve two challenges: first, segmenting the vessels; and second, analyzing the structure by extracting features that are highly correlated with a specific retinal condition.

In this study, we focus on solving the first challenge by proposing a deep-learning approach for segmenting the vessels in retinal images. Our model is based on the U-Net architecture and uses the InceptionV3 [4] model as the encoder. Given the challenges related to having insufficient annotated retina vessels datasets for deep learning, we adopted the idea of using transfer learning and image augmentation. Instead of using random weights to initialize our model, we use weights trained on millions of images for semantic segmentation from the ImageNet dataset, which we then fine-tune to match the object to be segmented (i.e., retinal vessels). To handle the issue of imbalanced classes, we use a customized loss function that tends to penalize incorrect classifications of pixels related to vessels more than incorrect classifications of background pixels.

The proposed model takes the whole retinal image as input and then segments the vessels in a short time (1 second). Instead of using batch normalization on a small number of batches, we used group normalization. To demonstrate the robustness of our proposed approach in segmenting the retinal vessels from images with various resolutions, we tested our approach on six publicly available datasets and one private dataset that we created. Our contributions can be enumerated as follows: (1) We propose a U-Net based deep learning model for vessel segmentation that utilizes a customized InceptionV3 model as an encoder. (2) We demonstrate the effectiveness of using TL and IA for limited data. (3) We handle the issue of imbalanced image classes which may lead to inaccurate results by adopting a weighted loss function. (4) We contribute a new
retinal image dataset for vessels segmentation (ORVS).

II. RELATED WORK

Several approaches have been proposed for retinal vessel segmentation. These approaches can generally be categorized into unsupervised and supervised approaches. The literature reviews conducted by [5] and [11] discuss more than a hundred approaches for vessel segmentation that are either supervised or unsupervised.

Unsupervised approaches segment the vessels using predefined sets of rules. If a pixel passes a set of conditions, it is considered to be related to a vessel; otherwise it is considered to be unrelated. Such approaches use use variety of techniques such as morphological operations [6]; matched filters [7], [8]; multi scale [9]; adaptive thresholding [10], [11]. They may also be model-based [12]. Unsupervised approaches tend to offer faster segmentation than supervised ones; however, they tend to fail when exposed to thin vessels or images with low contrast.

The other category of approaches to retinal vessel segmentation is supervised approaches. In such approaches, a pixel matrix \( I \) is associated with each retinal image, indicating the pixels that belong either to vessel or to background. \( I_{xy} \) represents a pixel at location \((x, y)\) in the retinal image. This pixel has a value of 1 if it belongs to a vessel and 0 if it is a background pixel. The classifier uses these labeled images and the actual retinal image to produce a new image with the same dimensions in which each pixel has a probability between 0 and 1 inclusive of belonging to a vessel. The closer the value is to 1, the higher the model’s confidence is that the pixel belongs to a vessel.

Recently, advancements in the field of deep learning have made it possible to use deep learning-based classifiers in medical-image analysis. Such approaches exhibit superior performance over the hand-crafted ones [1], [13]. Various deep-learning architectures have been proposed for vessel segmentation. For instance, in [13], U-Net architectures were used with residual blocks and binary cross-entropy as the loss functions. In this study they extracted patches from retinal images for the segmentation instead of performing the segmentation from the whole image. In analyzing the images, however, such models face the issue of imbalanced classes. Another model architecture used is fully connected layers, which was adopted by [15] and [16]. Neither handled the issue of imbalanced classes, however. Moreover, they tested their approach with a limited number of images.

Vessel segmentation poses several challenges, such as limited availability of datasets and the imbalance between the number of pixels related to the vessels and those related to other classes. Another challenge is related to the number of images used per batch when training the deep-learning model leading to increasing the batch error and hence inaccurate statistics. In this paper, we show that using image augmentation and transfer learning with a weighted loss function and customized U-Net model can help in achieving high precision when segmenting the retinal vessels. We also created a new dataset that can help researchers working in this field.

III. PROPOSED METHOD

The goal of this study is to segment the vessels in a retinal image. To achieve this, we propose a deep-learning model with the same architecture as the U-Net model [17] and use a customized InceptionV3 model as the encoder. In this section we discuss the architecture of our model, the prepossessing steps, data augmentation, and model initialization.

A. Network Architecture

Instead of creating a new architecture, we adopted the U-Net architecture, which consists of an encoder and a decoder. In the encoder, pooling is applied to summarize neighboring features and create high-level representations. In the decoder, feature maps are reconstructed, combining those from the encoder through skip connections, with those belonging to low-level scales.

We adopted the InceptionV3 model as the encoder, customized the normalization, and reduced the convolutional layers. The architecture of our model is presented in Fig. 1. The original architecture of InceptionV3 requires four instances of block C to be presented in block A (marked as as 3 in Block A in Fig. 1) but we used only three. Another change is that, in the original architecture, block B is required to have four instances of block C, but we used five instead (marked as 3 in Block B in Fig. 1). We also removed the auxiliary classifier. The final concatenation layer and other layers as shown in Fig. 1 were used to construct our encoder. A drop-out was added at the end with a factor of 0.3, and a sigmoid activation was used. The feature map for each convolutional layer uses the ReLU activation method, which applies Eq. 1 to each parameter from the layer, thereby removing all negative pixel values.

\[
f(x) = \max(0, x)
\]

The model is initially fed with a preprocessed image of size \(224 \times 224\). Three channels are used, not only the green channel as used by other approaches [1]. Skip connections were used in several places in the architecture. In Fig. 1, the first skip connection is at the third occurrence of block C. Other skip connections were then added to form the decoder. The selection of blocks to include in the encoder was based on a set of experiments we conducted. After the skip layers had been selected, we performed upsampling and concatenation so that the output image will have a \(224 \times 224\) dimensions. This was used to shorten the distance between the earlier and later layers. Short connections from early to later layers are useful in preserving high-level information about the positioning of the vessels. This is in contrast to the low-level pixel-based information that is transferred across the long pipeline of the architecture in a combination of convolutional and max-pooling or upsampling layers. High-level information tends to be lost as the image is downsampled and the shape and structure of the image is changed.
We maintain this information by using connections to earlier layers in the model.

B. Preprocessing

Retinal images obtained in less than ideal imaging environments tend to be of poor quality. In such images, the appearance of the blood vessels may be affected due to fluctuating pixels intensities or the addition of noise to the image. To handle these issues, we applied some image-processing techniques. These techniques were applied to every image before augmentation.

To deal with the issue of contrast variation, we use the contrast-limited adaptive histogram equalization CLAHE \cite{18} for each channel. This technique can enhance the contrast by stretching the grey level values of low contrast images. Gamma correction is then used to adjust the brightness. To smooth the edges of vessels, we apply a median filter, which computes the median of all the pixels under the kernel window (in our case it is $5 \times 5$), and the central pixel is replaced with this median value. All images are then resized to $224 \times 224$ and normalized to ensure every pixel value is between 0 and 1. Fig. 2 shows an original image before (left) and after (right) these techniques have been applied.

C. Group Normalization layer

In large models such as the one in the present study, normalization, including batch normalization, is important for training and convergence. Batch normalization works well when the number of images per batch is high (greater than 32); however, the number of errors in batch normalization increases as the number of images decreases, leading to inaccurate batch statistics. To overcome this issue, we use group normalization, which has proved to be effective in such situations \cite{19}. The group normalization divides the channels into groups and then calculates the mean and variance among the groups. Hence, group normalization is independent of batch size.

In training we used a batch size of two, because that was the maximum capacity of our machine memory. Group normalization organizes the channels into groups and calculates the mean and variance along the height and width axis and along a group channels using Eq. 2 where $\mu$ and $\sigma$ are computed over a set of pixels defined by $S_i$ where $S_i$ is defined using Eq. 3. In Eq. 3 $G$ represents groups which is in our case is set to 16, $C/G$ represents the number of channels per groups, and the $\lfloor \cdot \rfloor$ represents the floor operation.

\begin{align*}
\mu_i &= \frac{1}{m} \sum_{k \in S_i} x_k, \quad \sigma_i = \sqrt{\frac{1}{m} \sum_{k \in S_i} (x_k - \mu_i)^2 + \epsilon} \quad (2) \\
S_i &= \{k | k_N = i_N, \left\lfloor \frac{k_C}{C/G} \right\rfloor, \left\lfloor \frac{i_C}{C/G} \right\rfloor \} \quad (3)
\end{align*}

D. Loss Function

In order to check if the model had improved or not on the value returned from the loss function during the training
of the network it was run using the validation data. At first the binary cross-entropy function (BCE), shown in Eq. 4 was adopted where the total number of pixels is denoted by \( N \), and the pixels \( y_i \) are labelled by 0 for background and 1 for the vessels. The probability that pixel \( y_i \) belongs to the vessel is \( p_a(y_i) \) and to the background \( p_b(y_i) \). It should be noted that during classification using the BCE both false positives and false negatives can be penalized when working on classifying vessels and background pixels.

\[
BCE = -\frac{1}{N} \sum_{i=1}^{N} y_i \log(p(y_i)) + (1 - y_i) \log(1 - (p(y_i)))
\]  

(4)

Using the BCE loss function will bias the output towards the background resulting in improper vessel segmentation. That is, the output may suggest a misleading result of 90% accuracy. To achieve precise vessel segmentation output therefore require additional processing. This problem was avoided using the Jaccard distance which measures the dissimilarity between two data sets. It is defined as:

\[
L_j = 1 - \frac{|Y_d \cap \hat{Y}_d|}{|Y_d \cup \hat{Y}_d|} = 1 - \frac{\sum_{d \in Y_d} (1 \land \hat{y}_d)}{|Y_d| + \sum_{b \in Y_b} (0 \lor \hat{y}_b)}
\]  

(5)

\( Y_d \) represent the ground truth of the vessels, \( Y_b \) the ground truth of background, \( \hat{Y}_d \) the predicted vessel pixels and \( \hat{Y}_b \) the predicted background pixels. The cardinalities of the vessels \( Y_d \) are \( |Y_d| \) and the cardinality background \( Y_b \) is \( |Y_b| \) and \( \hat{y}_d \in \hat{Y}_d \) and \( \hat{y}_b \in \hat{Y}_b \). The values of \( \hat{y}_d \) and \( \hat{y}_b \) will always be between 0 and 1 since they are both probabilities. We can now approximate this loss function as shown in Eq. 5. The model will then be updated by Eq. 7 where \( j \) represents the the \( j \)th pixel in the input image and \( \hat{y}_j \) represents the predicted value for that pixel.

\[
\hat{L}_j = 1 - \frac{\sum_{d \in Y_d} \min(1, \hat{y}_d)}{|Y_d| + \sum_{b \in Y_b} \max(0, \hat{y}_b)} = 1 - \frac{\sum_{d \in Y_d} \hat{y}_d}{|Y_d| + \sum_{b \in Y_b} \hat{y}_b} \quad \text{for} \quad i \in Y_d
\]

\[
\hat{L}_j = \frac{1 - |Y_d| + \sum_{b \in Y_b} \hat{y}_b}{|Y_d| + \sum_{b \in Y_b} \hat{y}_b} \quad \text{for} \quad i \in Y_b
\]

(6)

(7)

We are now able to balance the emphasis the model gives to the vessel class and the background class using the Jaccard loss function and thus we combine BCE with Jaccard to optimize the results. When both are combined, the model converges faster than when using only the Jaccard still getting better results than either the BCE or the Jaccard loss function by themselves. Our final loss function is therefore with \( \beta_1 = 0.75 \) and \( \beta_2 = 0.25 \):

\[
L_f = \beta_1 \times BCE + \beta_2 \times L_j
\]  

(8)

E. Transfer learning

To handle the challenge faced in the field of medical imaging of not having enough datasets or large enough datasets to train a deep-learning model, we used an approach referred to as transfer learning. Such approaches can alleviate the issues caused by insufficient training data by using weights generated by training on millions of images \[20\]. In our study, we adopted the weights generated when training the InceptionV3 model on the ImageNet dataset which contains around 14 million labeled images. We thus provided a diverse set of images that the model had been exposed to.

By using transfer learning, we could reduce the problem of over-fitting caused when training on limited images and improve the overall performance of the model. Using the ImageNet weights, we initialized the weights of the encoder network component, and other layers were randomly initialized using a Gaussian distribution. We then trained our model using a mini-batch gradient to tune the weights of the whole network. When training, we realized when using transfer learning that the model converged faster than without transfer learning.

F. Image Augmentation

In total there were only 271 retinal images available for training, which is insufficient even when transfer learning is used. We thus augmented the dataset by applying a series of operations. These operations increased the number of images while preserving the characteristics of vessels. Hence, they allowed the model to be exposed to vessels of various appearances.

Unlike the approach in \[14\] we do not split the images into small patches. Doing so would require us to split the images into patches whenever we needed to segment the vessels, perform the segmentation and then re-assemble the images. Instead our approach maintains the appearance of vessels while increasing the number of images. We split each image into two halves vertically and two halves horizontally, producing a total of five images including the original. Each of these images is rotated by 90, 180, and 270 degrees, giving a total of 20 images \((5+5 \times 3)\). We then flipped both vertically and horizontally. Hence, from one image, we can obtain 60 images \((20 + 20 \times 2)\), which increased our dataset from 271 images to 16,260 \((60 \times 271)\). This larger dataset improves the generalization performance of our model.

IV. EXPERIMENTAL ANALYSIS

A. Data sets

1) New dataset: One of the challenges faced when working with retinal images is the limited availability of images. To help increase the number of images available for researchers working in this field, we introduce a new online retinal image for vessel segmentation (ORVS). The ORVS dataset has been newly established as a collaboration between the computer-science and visual-science departments at the University of Calgary\[1\].

This dataset contains 49 images (42 training and seven testing images) collected from a clinic in Calgary-Canada (check Table 1). All images were acquired with a Zeiss Visucam

\[1\]https://tinyurl.com/y3ke704j
200 with 30 degrees field of view (FOV). The image size is 1444×1444 with 24 bits per pixel. Images and are stored in JPEG format with low compression, which is common in ophthalmology practice. All images were manually traced by an expert who has been working in the field of retinal-image analysis and went through training. The expert was asked to label all pixels belonging to retinal vessels. The Windows Paint 3D tool was used to manually label the images.

2) Public datasets: To evaluate our approach on images obtained from different FOVs and in various resolutions, we decided to use six publicly available datasets: namely, the ARIA, CHASE, DR-Hagis, DRIVE, HRF, and STARE. Details about each dataset can be found in Table I. Using these datasets allows us to compare our approach with other state-of-the-art approaches that have evaluated their method with respect to each dataset. Note that, in all these experiments, we used the same approach used by [26] of which we take the first image 5 images from each category (healthy, diabetic retinopathy, and glaucoma) for training and validation and tested on all remaining images. As for DR-Hagis, ARIA, and ORVS we split by selecting randomly 15% of the images to be as testing and the remaining for training and validation.

### Table I

| Dataset     | Images | Dimensions | FOV | Machine           |
|-------------|--------|------------|-----|-------------------|
| ARIA [8]    | 120    | 768×584    | 50  | Canon CR5 non-mydriatic 3CCD |
| CHASE [21]  | 20     | 999×960    | 30  | Handheld Camera NM-200-D |
| DR-Hagis [22] | 34     | 3216×2136, 4752×3168 | 45  | Canon CR DGi, Topcon TRC NW6, Topcon TRC NW8 |
| DRIVE [23]  | 20     | 6301×300   | 35  | Topcon TRV-50     |
| HRF [24]    | 19     | 3504×2336  | 60  | Canon CF 60 UVI with Cannon EOS 20D |
| STARE [25]  | 20     | 700×605    | 45  | Topcon TRC NW6    |
| ORVS        | 42     | 1444×1444  | 30  | Zeiss, Viscuam 200 |

B. Implementation Details

We implemented this model on a Windows machine with a NVIDIA GeForce 2060 RTX with 6 GB dedicated memory. We used the Python language to implement proposed approach using Keras with tensorflow back-end.

Training was performed using the NAdam optimizer with learning rate set to 0.0001, \( \beta_1 = 0.9, \beta_2 = 0.999, \epsilon = 10^{-8} \), and batch size of 2 images. During training, three callbacks were used. First, the model checkpoints would save the model whenever a smaller value was returned on validation data from the custom loss function when comparing to the value at the last checkpoint. Secondly, the learning rate was reduced by a factor of 0.5 whenever 25 epochs passed without any improvement in the validation loss values. Finally the training was stopped if 100 epochs passed without any improvement.

When training our model we split the training and validation set by 85% and 15% respectively. These sets are selected randomly at each epoch. We used four evaluation methods to evaluate and compare our approach: namely, accuracy (Acc), dice coefficient (DC), sensitivity (Sen), specificity (Spec).

C. Effectiveness of TL and IA

To test the impact of using transfer learning (TL) and image augmentation (IA) when training our model, we conducted a series of experiments and then evaluated the model obtained by using the test images for all datasets. In this section, we show the overall performance without showing the performance with respect to each dataset. Note that, in all these experiments, we used the loss function defined in Eq. 8.

We first checked the performance of the model without transfer learning by randomly initializing weights using Gaussian distribution. Then we did an experiment using only rotated images with transfer learning. The third was using transfer learning and rotated and flipped images. The evaluation results...
for each of these experiments are shown in Table II. The results obtained show that using TL and IA together achieves the best results. Flipping helped in improving the results slightly from when using rotation only when augmenting images. Using only rotated images with transfer learning took around 319 epochs to converge while when using rotated and flipped images it needed 50 epochs.

| Experiment                  | Acc  | Sen  | Spec | DC   |
|-----------------------------|------|------|------|------|
| Rotated+Flipped             | 93.98| 56.45| 98.98| 68.14|
| TL+RotatedOnly              | 95.58| 84.57| 96.60| 80.88|
| TL+Rotated+Flipped          | 95.60| 85.18| 96.51| 80.98|

**D. Effectiveness of Loss Functions**

A well-known loss function for binary classification is the binary cross-entropy loss function. This loss function works well when the classes in the image are balanced. However, in our case, the object we are trying to segment represents 20% or less of the image. Hence, we decided to use the Jaccard distance combined with BCE, as noted earlier.

We conducted three experiments to test which configuration would achieve the best results. First, we trained our model using the BCE loss function only, which is a built-in loss function in the Keras library; second, we trained it using the Jaccard distance-loss function only; and third, we trained it by combining both loss functions. The results obtained are shown in Table III. We noted that there is a slight improvement in performance when we combine both loss functions compared to using either one of them alone. We also realized that using BCE and Jaccard distance together would help in converging faster, 50 epochs, than when using BCE with 71 epochs.

| Loss Function     | Acc  | Sen  | Spec | DC   |
|-------------------|------|------|------|------|
| BCE               | 95.25| 86.58| 95.92| 80.23|
| Jaccard Distance  | 95.21| 85.24| 96.12| 79.82|
| Jaccard Distance+BCE | 95.60| 85.18| 96.51| 80.98|

**E. Comparison With Other Approaches**

To evaluate our proposed method, we compared it with approaches which had been tested on some of the same datasets we used, as shown in Table IV. Unfortunately, these other approaches had not been evaluated with respect to all available datasets; hence, when comparing, we split our results per dataset so we could make a fair comparison.

We compared our approach to several studies, all of which used the Stare and DRIVE datasets (check Table IV). Of the 16 approaches, only six used the CHASE dataset (check Table V). Two used the HRF (check table VI), and none used the DR-Hagis and the ARIA datasets (check table VII). For each of these datasets, we show the performance of our model with respect to accuracy, sensitivity, specificity, and the dice coefficient. We also show, in Fig. 3 the vessels segmented by our model for a sample image from each dataset along with ground truth and related retinal images. This figure clearly shows the variation in the orientations and resolutions of the vessels in each dataset.

Our proposed approach achieved the highest sensitivity and dice coefficient among all approaches when tested against DRIVE and Stare. Moreover, our model achieved the highest...
TABLE IV
APPROACHES FOR RETINAL VESSELS SEGMENTATION ON THE DRIVE AND STARE DATASETS.

| Method              | DRIVE | STARE |
|---------------------|-------|-------|
|                     | Acc   | Sen   | Spec | DC   | Acc   | Sen   | Spec | DC   |
| Unsupervised        |       |       |      |      |       |       |      |      |
| Nguyen et al. [9]   | 94.07 | -     | 98.58 | 72.98 | 93.24 | -     | 98.63 | 77.74 |
| Roychowdhury, Sohini, et al. [10] | 94.9 | 73.9 | 97.8 | - | 95.6 | 73.2 | 98.4 | - |
| Memari et al. [12]  | 96.1  | 76.1  | 98.1  | -   | 95.1  | 78.2  | 96.5  | -   |
| Zhao et al. [11]    | 94.7  | 73.54 | 97.89 | -   | 95.09 | 71.87 | 97.67 | -   |
| Khan et al. [13]    | 95.1  | 73.4  | 97.67 | -   | 95.1  | 73.6  | 97.1  | -   |
| Zhang et al. [29]   | 94.7  | 74.3  | 97.6 | -   | 95.4  | 76.7 | 97.6 | -   |
| Bankhead et al. [8] | 93.7  | 70.3  | 97.1  | -   | 93.2  | 75.8  | 95.0  | -   |
| Supervised          |       |       |      |      |       |       |      |      |
| Wang et al. [27]    | 95.41 | 76.48 | 98.17 | 80.93 | 96.40 | 75.23 | 98.85 | 81.25 |
| Hu et al. [16]      | 95.33 | 77.72 | 97.93 | -   | 96.32 | 75.43 | 98.14 | -   |
| Oliveira et al. [15]| 95.76 | 80.39 | 98.04 | -   | 96.94 | 83.15 | 98.58 | -   |
| Xia et al. [14]     | 96.55 | 77.15 | -    | -   | 96.93 | 74.69 | -    | -   |
| Fu, Xu, Wong, et al. [30] | 95.20 | 76.00 | - | - | 95.80 | 74.10 | - | - |
| Yan et al. [26]     | 95.40 | 76.50 | 98.10 | -   | 96.10 | 75.80 | 97.50 | -   |
| Brancati et al. [31] | 94.90 | 78.20 | 97.60 | - | - | - | - | - |
| Orlando et al. [28] | -    | 78.97 | 96.84 | 78.41 | - | 76.80 | 97.38 | 76.44 |
| Jin et al. [32]     | 95.66 | 79.63 | 98.00 | 82.37 | 96.41 | 75.95 | 98.78 | 81.43 |
| Proposed Method     | 95.61 |       |       |      | 95.26 | 85.61 | 96.57 | 84.06 |

TABLE V
APPROACHES FOR RETINAL VESSELS SEGMENTATION ON CHASE DATASET.

| Method              | Acc   | Sen   | Spec | DC   |
|---------------------|-------|-------|------|------|
| Oliveira et al. [15]| 96.53 | 77.79 | 98.64 | -    |
| Wang et al. [27]    | 96.03 | 77.30 | 97.92 | 78.09 |
| Memari et al. [12]  | 93.90 | 73.80 | 96.80 | -    |
| Yan et al. [26]     | 96.10 | 76.33 | 98.09 | -    |
| Orlando et al. [28] | 96.10 | 72.77 | 97.12 | 73.32 |
| Jin et al. [32]     | 96.51 | 74.64 | 98.74 | -    |
| Proposed Method     | 96.83 | 90.21 | 97.34 | 85.46 |

TABLE VI
APPROACHES FOR RETINAL VESSELS SEGMENTATION ON HRF DATASET.

| Method              | Acc   | Sen   | Spec | DC   |
|---------------------|-------|-------|------|------|
| Orlando et al. [28] | -     | 78.74 | 95.84 | 71.58 |
| Jin et al. [32]     | 96.51 | 74.64 | 98.74 | -    |
| Proposed Method     | 95.07 | 91.56 | 95.09 | 81.90 |

TABLE VII
PERFORMANCE OF OUR APPROACH ON THE DR-HAGIS, ARIA, AND ORVS DATASETS.

| Dataset | Acc   | Sen   | Spec | DC   |
|---------|-------|-------|------|------|
| DR-Hagis| 96.87 | 67.13 | 98.57 | 71.49 |
| ARIA    | 95.31 | 81.94 | 96.40 | 77.59 |
| ORVS    | 96.52 | 84.32 | 97.19 | 78.11 |

accuracy, sensitivity, and dice coefficient on the CHASE dataset which had been tested against the same images. Furthermore, we achieved the highest accuracy, sensitivity, precision, and Dice coefficient on the HRF dataset. We realized that, although some approaches performed slightly better than our method in terms of accuracy, we achieved better in terms of dice coefficient and sensitivity, which shows that our approach is more precise in segmenting the vessels than these approaches. For instance, in [14] researchers achieved the highest accuracy with less than 1% difference from what was achieved by our model. However, our model outperformed their approach in terms of sensitivity, with more than a 5% difference between our measurement and theirs. In [32], researchers achieved slightly better scores for accuracy and specificity, but our approach performed better in terms sensitivity and Dice coefficient. Given that the vessels occupy less than 20% of the retinal image, accuracy alone is not good indicator for checking the performance of a model. Sensitivity and Dice coefficient are better indicators against which to evaluate a model when segmenting retinal vessels.

We also noticed that some approaches performed slightly better than ours in terms of accuracy and specificity with respect to some dataset(s) but that we outperformed those approaches when tested against other datasets. For instance, in [15], researchers achieved slightly better accuracy and precision than our approach when tested on the DRIVE datasets, but our model performed better when tested on the CHASE dataset.

Since none of the reported studies reported their performance on the DR-Hagis and ARIA datasets, we just reported the performance of our model with respect to these datasets by including our new public dataset, as shown in Table VII. One reason the ARIA is not commonly used today is that it is no longer available online. However, it can still be retrieved by request, and we accordingly requested it. As for the DR-Hagis dataset, it is newly published. In general, our proposed approach shows superior performance with respect to the approaches reported in this study in terms of precisely segmenting vessels when exposed to images with various resolutions and orientations.

V. CONCLUSION

In this paper, we propose a deep learning based approach for vessels segmentation where we show the effectiveness of our customized model when being combined with transfer learning, image augmentation, and a customized loss function. Our approach achieve state of the art performance on vessels...
segmentation when compared to other modern approaches. We also contribute a new dataset that can be used by researchers for improving vessels segmentation. This will help researchers testing their approaches on images obtained from various sources with diverse data. Our model achieved an average accuracy, sensitivity, specificity, and dice coefficient of 95.60%, 85.18%, 96.51%, and 80.98% respectively.

ETHICS

The necessary ethics approval has been obtained by Health Research Ethics Board of Alberta (HREBA REB17-2302). No identifiable information has been shared and hence is not possible to identify corresponding individuals these images obtained from.

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