

Comprehensive Modality Integration for Diabetic Retinopathy Image Analysis

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Abstract:

Diabetic retinopathy (DR) is frequently discovered in the eyes of diabetics and this has been found to be a contributing factor to vision loss. Early intervention, along with regular fundus photography monitoring, is the most effective way to treat the condition. There are many patients with diabetes who need intensive screening; hence, there is an increase in computer-assisted and fully automated methods for diagnosing DR. Neural networks have come a long way in recent years in various domains. As a result of automating diagnosis of DR and giving customized suggestions to DR patients, it may be seen that precise as well as intricate DR classification is important. Here we have showed cross-modality feature fusion framework for diabetic retinopathy (DR) images categorization. Cross mode here means an RGB image having green channel. A multi-scale and multi-receptive feature extraction block has thus been presented initially so as to learn local and global features from both modalities. Moreover, the learnt features at different scales are successfully fused with current multi-level feature fusion block for image classification task. Our existing system has been compared against state-of-the-art (SOTA) deep learning frameworks for categorizing DR images using the MESSIDOR and IDRID databases. Results analysis shows clearly that the ongoing cross-modality feature fusion based classification framework outperforms all the present SOTA frameworks according to some evaluation metrics.

Keywords: Modality Integration, Non-proliferative and proliferative diabetic retinopathy, Green channel, Classification.

Introduction

Diabetic retinopathy often develops without any noticeable symptoms in its early stages. However, as the condition progresses, the symptoms like blurred or distorted vision, floaters (small dark spots), impaired color vision, fluctuating vision, loss of central vision may become apparent. In advanced stages, you may lose central vision, which is critical for tasks like reading, recognizing faces, and driving. Historically, the assessment of diabetic retinopathy (DR) grade has been conducted by considering a combination of various structural characteristics observed in color fundus images. These features encompass the presence of microaneurysms, exudates, hemorrhages, and neovascularization, among others [1]. Over the past two decades, image classification has evolved into a proficient and high-demand research area, particularly within the realms of automated medical, scientific, and educational sectors, as well as various computer vision applications. Inter-active media, such as images and videos, are consistently being generated and shared on social media platforms, thanks to the

proliferation of advanced data-capturing devices like smart- phones and high-speed internet connections. Consequently, extensive digital databases or repositories have been established. Likewise, within the field of healthcare, substantial medical databases have been created, fueled by advanced techniques for patient diagnosis, treatment planning, and the evaluation of treatment responses.

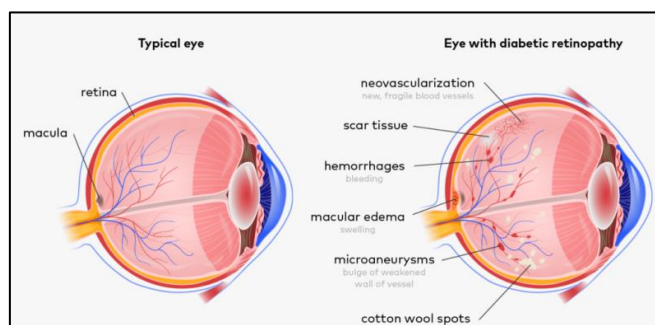


Fig. 1. Sample eyes with and without diabetic retinopathy (*left part: Healthy eye and right part: Diabetic eye*). Diabetic retinopathy is a condition that arises as a consequence of the harm inflicted by diabetes on the tiny blood vessels situated within the retina of the eye.

According to the World Health Organization (WHO), in 2014, there were 422 million individuals diagnosed with diabetes, and of that number, 35% developed some form of retinopathy due to the accumulation of damage to small blood vessels in the retina [2]. Assessing diabetic neovascularization and macular edema in laboratory animals has posed challenges. Because most commonly used species lack a macula and fail to exhibit the characteristics seen in advanced diabetic retinopathy in human patients, especially in terms of retinal neovascularization and thickening. Figure 1 illustrates the differences between diabetic retinopathy in a healthy eye and a compromised one, featuring various types of DR lesions. Diabetic retinopathy is a medical condition that affects the retina of the eye and can lead to vision problems, including blindness. The DR patient with high blood pressure, high hemoglobin A1C are considered at highest risk. Immediate screening with medical expertise is required to recover from the various issues [3]. Therefore, regular screening is essential for detection or DR at early stage. Generally, DR has four different types of lesions *i.e.*, Microaneurysms (MA), Haemorrhages (HM), soft and hard exudates [4]. The detailed explanation for each type of DR images is given below:

- **Microaneurysms (MA)** represent an initial stage in the progression of diabetic retinopathy. These are characterized by the presence of small, red, round dots on the retina, which occur due to the weakening of the blood vessel walls. Early detection and management of microaneurysms are crucial in preventing the advancement of diabetic retinopathy and preserving vision [4].
- **Hemorrhages (HM)** in the context of diabetic retinopathy are identified by the presence of larger spots on the retina. These spots typically have irregular margins and can be larger than 125 micrometers in size, which distinguishes them from microaneurysms. Recognizing and monitoring hemorrhages is important in assessing the severity of diabetic retinopathy as it progresses to more advanced stages [5].
- **Hard exudates** are a result of plasma leakage in the context of diabetic retinopathy. These

exudates are visible as yellow spots on the retina and are caused by the leakage of plasma from damaged blood vessels. The presence of hard exudates is an important clinical sign, and their detection and management play a role in assessing and treating diabetic retinopathy [6].

- **Soft exudates**, also known as cotton wool spots, are a consequence of nerve fiber swelling in the context of diabetic retinopathy. These exudates appear as white, oval-shaped areas on the retina. They are associated with localized damage to the nerve fibers and are a significant sign of retinal involvement in diabetic retinopathy. Detecting and monitoring soft exudates is crucial in managing the condition and preventing further vision impairment [7].

Microaneurysms and hemorrhages typically manifest as red lesions on the retina, while both types of exudates (hard and soft exudates) appear as bright lesions. These differences in color and appearance are significant characteristics used in diagnosing and categorizing diabetic retinopathy, aiding healthcare professionals in determining the stage and severity of the condition. Diabetic retinopathy detection typically involves identifying five distinct stages of the condition, which are: No Diabetic Retinopathy, Mild Diabetic Retinopathy, Moderate Diabetic Retinopathy, Severe Diabetic Retinopathy and Proliferative Diabetic Retinopathy [8]. To address this issue, image classification has emerged as an effective solution, enabling efficient access to medical image data. Traditional approaches have relied on manually crafted techniques for classifying images, but these methods prove ineffective when dealing with extensive databases. Thus in this paper, we have proposed a deep learning based DR classification approach. The major contributions are:

- 1) A novel approach with cross modality (*RGB and Green channel*) feature fusion is presented for diabetic retinopathy image classification.
- 2) The detailed experimental analysis with present and existing approaches is performed on MESSIDOR [9] and IDIRD [10] databases.

I. LITERATURE SURVEY

The significant number of individuals diagnosed with diabetes and the high prevalence of diabetic retinopathy (DR) among them have spurred a growing demand for automated DR diagnosis systems. Over time, considerable progress has been achieved, and satisfactory results have been obtained in several sub-problems, such as vessel segmentation and lesion detection. However, it's essential to note that these outcomes have primarily been derived from relatively small datasets and are still a considerable distance from practical real-world applications. The choice of the most suitable treatment for patients with diabetic retinopathy can vary depending on the disease stage. For patients with no DR or mild non-proliferative diabetic retinopathy (NPDR), regular screening is typically sufficient. The sample images for various classes are provided in the Figure 2. However, for patients with moderate NPDR or more severe conditions, treatment options can range from scatter laser therapy to vitrectomy. Therefore, accurately grading the severity of a patient's DR is a crucial initial step in providing them with the appropriate and timely treatment [11].

Seepthi *et al.* [12] proposed a methodology that employs morphological operations and segmentation procedures to detect blood vessels, exudates, and microaneurysms in retinal fundus images. The retinal fundus image is divided into four sub-frames, and various features are extracted. Wavelet trans-

formations are applied to these extracted features, followed by principal component analysis to enhance feature quality. Neural network backpropagation and the one-rule classifier methods are employed to classify the images as diabetic or non-diabetic. In the work of Amin *et al.* [13], a DR model was developed to automatically differentiate retinal images into regions with exudates and non-exudates. This technique involves pre-processing, starting with lesion extraction, feature extraction, and image classification. Jiang *et al.* [14] proposed gradient-weighted class activation mapping for multi-label classification of diabetic retinopathy images. The correlation between diabetic retinopathy and its complication *i.e.*, diabetic macular edema with cross stage attention is studied in [15]. In [16], as part of simulating the diagnostic process, a novel approach is introduced. It involves the use of a two-stream binocular network designed to capture subtle correlations between the left and right eyes. This innovative design allows the model to effectively leverage information from both eyes to enhance the diagnostic capabilities, potentially leading to more accurate and robust results. The authors have introduced a two-stream model in [17]. This model employs the image itself as one input stream and also incorporates one of its individual channels as a complementary input stream. This approach aims to harness both the full image information and specific channel data to enhance the model's performance and capture a broader range of features and characteristics in the input data.

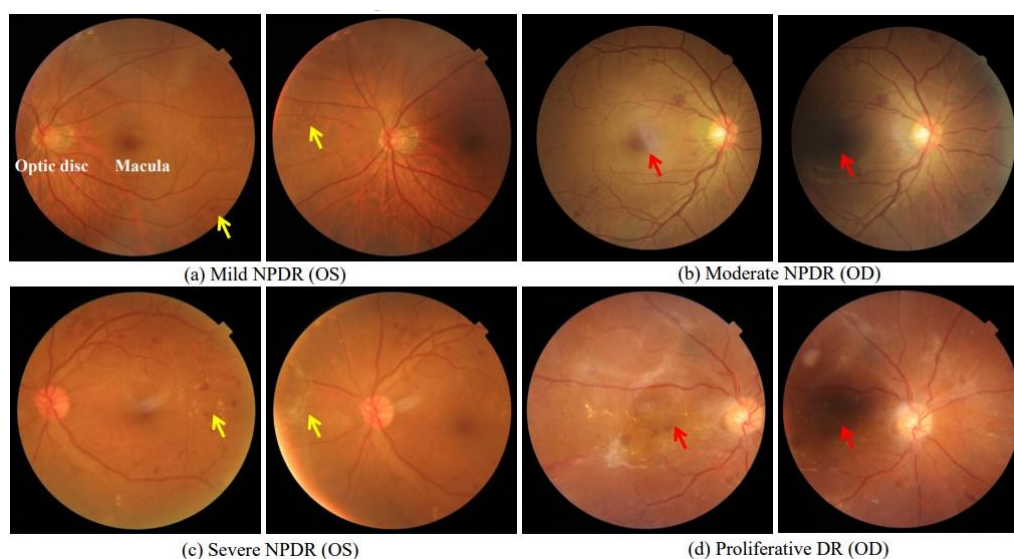


Fig. 2. Sample images for mild, moderate, severe non-proliferative and proliferative diabetic retinopathy classes. *Here, the degraded part on each retina image is marked with arrow. We can observe that the each class has different level of degradation.*

He *et al.* [18] introduce a novel component known as the Category Attention Block (CAB). This block is designed to delve into region-specific features that are more discriminative for each diabetic retinopathy (DR) grade. Importantly, it ensures that each category, or DR grade, is treated with equal importance, allowing the model to focus on the unique features associated with each grade for more accurate and balanced classification. Authors proposed a joint learning of multi-level tasks for diabetic retinopathy grading approach on low-resolution fundus images [19]. This approach likely involves the simultaneous learning of multiple aspects or tasks related to DR grading at different levels, with a focus on effectively handling low-resolution fundus images. The goal is to improve the accuracy and robustness of DR grading under conditions where image quality may be limited. The

framework is designed to facilitate the joint training of sub- networks involved in tasks related to image quality assessment, image enhancement, and diabetic retinopathy (DR) disease grading [20]. This unified approach allows these sub-networks to work together and learn from one another, potentially improving the overall performance of these tasks by leveraging their interdependencies. Along with two-field database, authors have proposed the cross-field transformerto capture the correspondence between two fields as well as the long-range spatial correlations within each field in [21]. Structural and angiographic optical coherence tomography based diabetic retinopathy classification approach at multiple levels is proposed in [22]. Here, to enhance the accuracy and reliability of classification, a new convolutional neural network architecture was developed. This architecture is founded on the principles of dense and continuous connections, complemented by adaptive rate dropout. These features are integrated to improve the network's performance in tasks such as image classification, making it more robust and effective in handling complex data. Adarsh *et al.* [23] employed Support Vector Machines (SVM) as a classifier for DR classification. This classification was based on utilizing features extracted from fundus images, specifically focusing on characteristics related to blood vessels and exudates. SVM is a popular machine learning algorithm known for its effectiveness in binary and multiclass classification tasks, making it a suitable choice for tasks like DR classification based on image features. These research efforts contribute to the development of effective methods for identifying and classifying different characteristics of diabetic retinopathy, which is crucial for early diagnosisand treatment.

II. PROPOSED FRAMEWORK

The complete overview of the proposed diabetic retinopathy image classification network is provided in Fig. 3. As one of the studies [17] proves that the green channel often provides good contrast for blood vessels in retinal images. This can be helpful in the detection of microaneurysms, hemorrhages, and other vascular abnormalities associated with diabetic retinopathy. The detailed visualization of each channel is providedin Fig. 4. From Fig. 4, it can be observed that the greenchannel often provides good contrast for blood vessels in retinal images, less sensitive to hemoglobin compared to the red channel and changes in color or variations in the green channel can indicate abnormalities in the retinal tissue. Totake advantages of these features, we have processed the Greenchannel independently with RGB image.

Along with the advantage of Green channel, to consider local and global level features, we have proposed multi- receptive multi-scale feature extraction block. The proposed block is defined as:

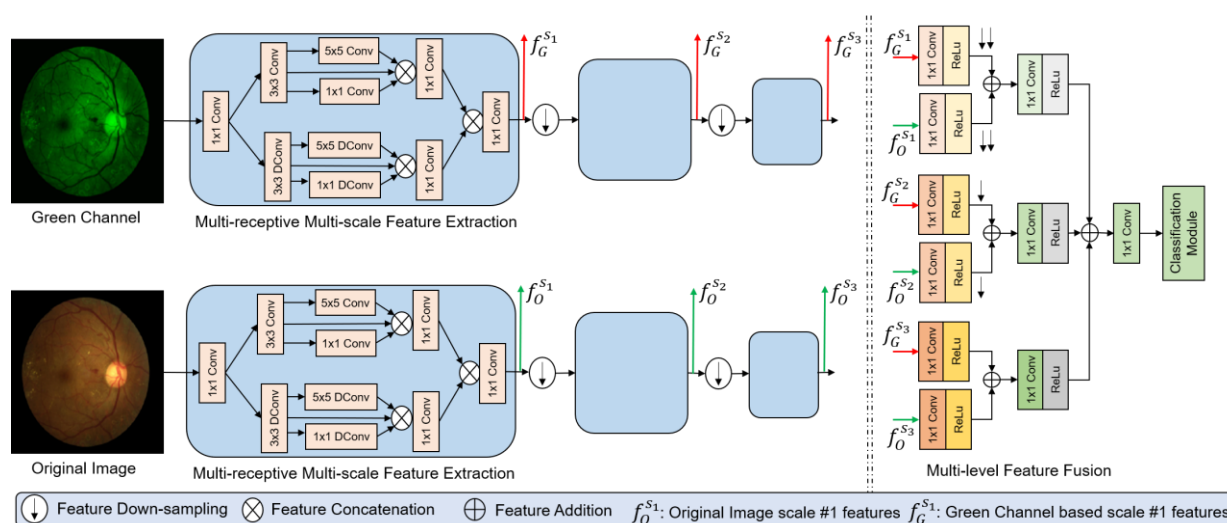


Fig. 3. Overview of the proposed network for diabetic retinopathy image classification. *Initially, the RGB image is pre-processed and green channel is extracted. Further, the independent features are extracted from RGB image and green channels at various scales. Finally, the various level features are fused with proposed multi-level feature fusion block for classification.*

incorporate both global and local channel information are fused with the proposed multi-level feature fusion block. The iterative processing and incorporation of both global and local information at different levels aim to enhance the model's ability to make accurate and robust classifications. Therefore, all the features from green channel ($f_G^{s1}, f_G^{s2}, f_G^{s3}$) and RGB image ($f_o^{s1}, f_o^{s2}, f_o^{s3}$) fused with the proposed multi-level feature fusion block. Initially, the RGB and green channel features at scale level are merged as:

$$f^{s1} = C_1(f_G^{s1}) + C_1(f_o^{s1}) \quad (4)$$

$$f^{s2} = C_1(f_G^{s2}) + C_1(f_o^{s2}) \quad (5)$$

$$f^{s3} = C_1(f_G^{s3}) + C_1(f_o^{s3}) \quad (6)$$

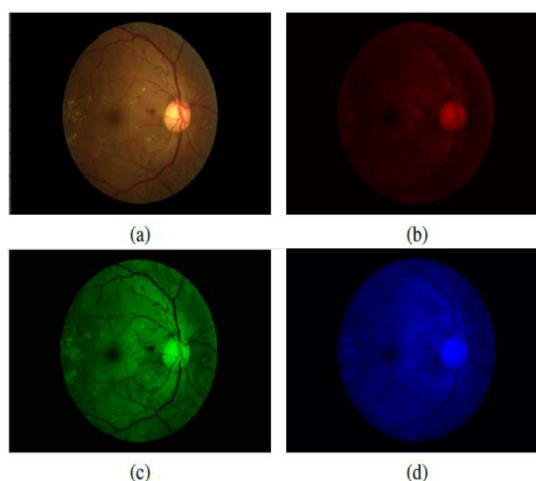


Fig. 4. Original RGB and channel-level image representation of diabetic retinopathy image.

$$f^s = C_1 \{f_{mr}, f_{ms}\}; \quad i \in 1, 2, 3 \quad (1)$$

where, C_1 is convolution operation with 1X1 kernel size, .

indicates concatenation operation, f_{mr} and f_{ms} are the multi-receptive and multi-scale features respectively. These features are extracted as:

$$f_{ms} = C_1 \{C_3(f), C_5(C_3(f)), C_1(C_3(f))\} \quad (2)$$

$$f_{mr} = C_1 \{DC_3(f), DC_5(DC_3(f)), DC_1(DC_3(f))\} \quad (3)$$

where, C_s and DC_s are the convolution and dilated convolution operations with $s \times s$ kernel size, respectively and f is the input feature after passing through C_1 .

After repeating the above operation three times ($s; i1, 2, 3$) at different scales, which likely involves a process of feature extraction or refinement, the resulting features that where, C_1 represents convolution operation with 1 \times 1 kernel size followed by ReLu activation function. Further, these features are merged as:

$$f = C_1[C_1(f^1) + C_1(\downarrow(f^2)) + C_1(\downarrow\downarrow(f^3))] \quad (7)$$

where, \downarrow and $\downarrow\downarrow$ indicate down-sampling operation with factor of 2 and 4, respectively. The resulting features that incorporate both global and local information at different levels. These feature are flatten to 1-dimensional layer. Further, fully connected layers are used for classification.

III. TRAINING DETAILS

A. Databases

MESSIDOR [9]: The MESSIDOR database is a publicly available dataset that contains 1,200 retinal images captured from patients with diabetic retinopathy. This dataset is a valuable resource for researchers and medical professionals working on the diagnosis and management of diabetic retinopathy. It provides a substantial collection of retinal images, which can be used for various purposes, including the development and evaluation of algorithms and models for

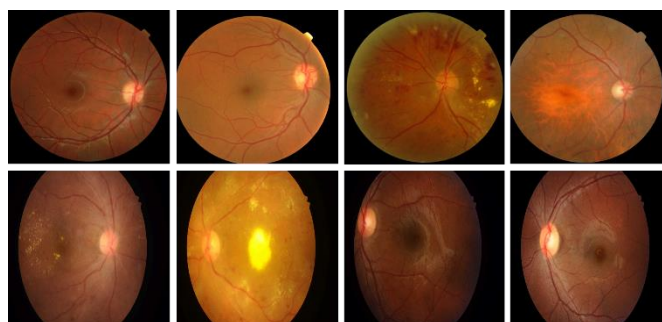


Fig. 5. Sample images from MESSIDOR [9] (first row) and IDRID [10] (second row) database used for classification.

the detection and classification of diabetic retinopathy. The division of the retinal images in the MESSIDOR database is done into four groups based on the severity of diabetic retinopathy is as: Group 1: 546 images, Group 2: 153 images, Group 3: 247 images and Group 4: 254 images. Each of these groups likely represents different stages or severity levels of diabetic retinopathy, ranging from mild to severe. The specifications of retinopathy grading are based on number of micro-aneurysms,

hemorrhage, and the sign of neovascularization. The images in which the above abnormalities are absent are considered as normal images. For experimental analysis of the proposed and existing network, we have divided the total 1200 images as 900 training and 300 testing splits.

IDRID [10]: The Indian Diabetic Retinopathy Image Dataset (IDRID) is a significant database representing an Indian population, specifically tailored for diabetic retinopathy research. This dataset is valuable because it includes retinal images from individuals in India, making it relevant for the study of diabetic retinopathy. Such region-specific datasets are essential for ensuring that the research and diagnostic tools are applicable and effective for the unique characteristics of the Indian population. This database has 413 and 103 training and testing images respectively. The ground truth labels for diabetic retinopathy and diabetic macular edema severity grade is also provided for training and testing analysis.

B. Implementation Details

All the images are resized to 256×256 for training the proposed network. While training the proposed network on IDRID [10] database, we have performed the data augmentation like horizontal and vertical flipping as number of training images are less. We train the network on Google Colab with batch size of 16 and learning rate 0.00002. We have kept 1×4 (four classes) and 1×5 (five classes) dimensional fully connected layer at last while training on IDRID and MESSIDOR database respectively.

V. RESULT ANALYSIS

The effectiveness of the proposed and existing architectures is analysed on testing splits of MESSIDOR [9] and IDRID [10] database. The parameters like Accuracy, Sensitivity and Specificity are calculated

Table I Classification Accuracy Analysis Of Diabetic Retinopathy On Messidor [9] Database.

Method	Accuracy	Sensitivity	Specificity
VGG-19 [24]	89.51%	79.21%	91.74%
ResNet [25]	90.01%	79.65%	92.35%
GoogleNet [26]	90.25%	80.87%	93.18%
InceptionNet [27]	91.12%	81.36%	93.87%
Proposed	91.67%	82.99%	94.63%

Table II Classification Accuracy Analysis Of Diabetic Retinopathy On Idrid [10] Database.

Method	Accuracy (%)	Sensitivity (%)	Specificity (%)
VGG-19 [24]	84.79	78.69	89.09
ResNet [25]	85.22	79.26	90.36
GoogleNet [26]	85.99	79.89	91.79
InceptionNet [27]	86.35	80.17	91.54
Proposed	83.56	80.81	92.61

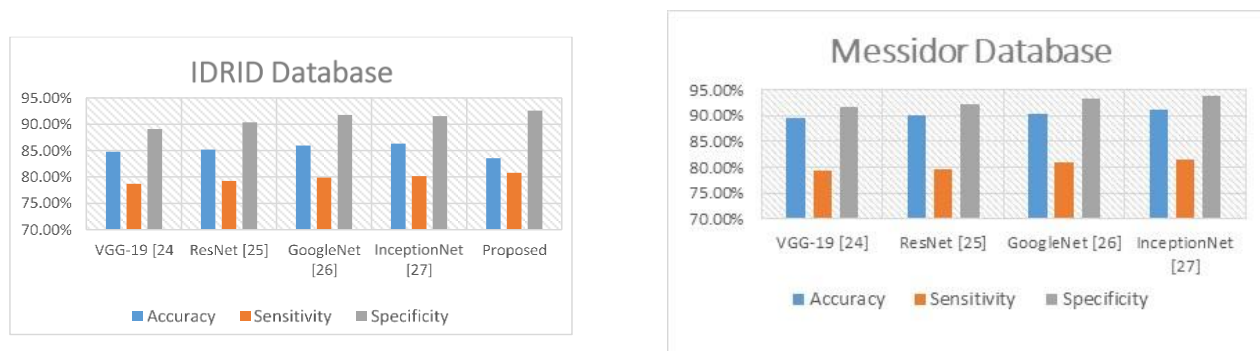


Fig. 6. Analysis with Accuracy, Sensitivity and Specificity on MESSIDOR (*top graph*) and IDRID [10] (*bottom graph*).

Specificity are calculated in terms of True Positive (TP), True Negative (TN), False Positive (FP), and False Negative(FN).

Mathematical expression are given below:

$$\text{Accuracy} = \frac{TP+TN}{TP+TN+FP+FN} \quad (8)$$

$$\text{Sensitivity} = \frac{TP}{TP+FN} \quad (9)$$

$$\text{Specificity} = \frac{TN}{TN+FP} \quad (10)$$

The quantitative analysis on MESSIDOR and IDRID

databases is provided in TABLE I and II respectively. Also, the graphical analysis on both the databases is provided in Figure

6. From these results, it is clear that the proposed network works effectively as compared for existing state-of-the-art methods for diabetic retinopathy classification.

IV. CONCLUSION

A modality integration feature fusion network is used in the current study project to classify images of diabetic retinopathy. The current work uses a cross-channel feature fusion approach to effectively extract information from several modalities. A cross-channel attention module that is coupled with a two-stream model is used to accomplish this method. This method greatly enhances the model's performance and accuracy in an image classification task by enabling it to concentrate on and collect data from several channels. The empirical findings demonstrate a performance parameter analysis, Accuracy (83.33%), Sensitivity (86.67%), and Specificity (94.44%) on MESSIDOR database and Accuracy (87.56%), Sensitivity (92.61%), and Specificity (80.81%) on IDRID database. The present work attains 85.45% average accuracy which is significantly better as compared to an existing state-of-the-art deep learning architectures for DR image classification.

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