**DFU\_MultiNet: A deep neural network approach for detecting diabetic foot ulcers through multi-scale feature fusion using the DFU dataset**

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**Abstract**

Diabetic foot ulcer (DFU) is a common problem among people with diabetes that can result in amputation of the affected limb. Modern DFU treatment and diagnosis methods are expensive and time-consuming. Today, the development of the computer-aided diagnosis (CAD) method makes it possible for pathologists to diagnose DFU more swiftly and accurately. This has led to a rise in interest in deep learning (DL) approaches based on CAD. In this study, we introduce a novel framework called "DFU\_MultiNet," which focuses on the transfer learning approach to classify healthy and ulcer skin images using publicly available repositories. The proposed framework is developed to offer an efficient and robust method for DFU classification that determines the distinction between healthy and ulcerated skin. The proposed approach extracts features from foot samples using three well-known pre-trained CNN models: VGG19, DenseNet201, and NasNetMobile. Finally, these extracted results are merged through a summing layer to create a powerful hybrid network. Through obtaining impressive accuracy (99.06%), precision (100.00%), recall (98.18%), specificity (100.00%), F1-score (99.08%), and AUC (99.09%) the proposed "DFU\_MultiNet" framework holds great potential as a diagnostic tool in healthcare and clinical settings.

**Keywords**: diabetic foot ulcer; CAD; VGG19; NasNetMobile; DenseNet201; feature fusion.

**1 Introduction**

Diabetic foot ulcers (DFUs) are a fateful issue of diabetes, identified by foot injuries. According to reports, the global population of diabetic individuals was 151 million in the year 2000, surged to over 422 million in 2014, and is now estimated to be around 537 million in 2021. In the past two decades, there has been a significant 10.5% increase in the prevalence of diabetes among adults aged 18 years and older. [1]

It is worth noting that around 80% of these diabetic patients reside in developing countries, which often lack adequate healthcare facilities and resources, leading to a lower level of awareness about patient health conditions [**2**]. A significant proportion of diabetic patients (ranging from 15% to 25%) suffer from DFUs, which can progress to a severe stage, necessitating lower limb amputation, hospitalization, and even death if left untreated [**3, 4**]. Infection with DFUs is a common cause of limb or foot amputation [**5**].

It reduces patients’ survival rates and diminishes the power of human life, impacting their ability to earn a livelihood and participate in social activities [**6**]. Conditions such as gangrene and tissue death from disease can lead to the need for amputation, and the problem of DFUs is expected to rise in the future [**7**]. The foot is a vital part of human body, but sadly, approximately a million patients who have high blood sugar will lose this vital part every year. After analyzing several studies, we observed that every 20 seconds, a diabetic foot undergoes an operation.

Doctors need thorough information for a correct diagnosis and better treatment of DFU. Traditional diagnostic techniques require a lot of manual labor and are prone to mistakes. Computer-assisted diagnostic (CAD) techniques improve performance as well as lower expenses. Moreover, recent advancements in wearable health and mobile technologies help treat diabetes and related complications. They can help enhance the patient’s quality of life and extend remission by detecting and managing detrimental inflammation and foot pressure [**8**]. Sensors are devices that detect various types of signals, such as physical, chemical, and biological signals, and record and measure these signals in a special process. These sensor technologies are widely used in current medical systems. When new sensors and sensor-related technologies are schematically developed, non-medical areas will adjust them for utilization in their business sectors. The evolution of new medical sensor generations points to a wider application of these devices in the healthcare sector [**9**]. Medical imaging [**10–12**] plays a critical role in diagnosing and treating various medical conditions. The performance of machine learning (ML) and deep learning (DL) approaches in this field heavily relies on the use of advanced feature extraction and selection methods that can accurately capture important visual characteristics such as color, shape, and size. Previous studies utilizing ML and CNN approaches have achieved remarkable outcomes in accurately diagnosing DFUs. However, there is still a need for further research to ensure the efficacy of these techniques in real-world settings with various functions. Accurate diagnosis and proper management of DFUs play a crucial role in improving the patient's prognosis. DFU management [**13**] involves various procedures such as vascular resection, wound removal, and infection treatment. The DFU treatment regimen depends on the type and condition of the foot wound. The challenges associated with DFU care involve a series of educational challenges that require the evaluation and comparison of various classification, detection, and segmentation [**14**] techniques to determine the pre-trained techniques and their potential applications [**15**].

Transfer learning refers to the reuse of a pre-trained CNN model. This technique is highly popular in DL due to its ability to train deep CNN networks with a small dataset. This technique works better for problems with large datasets. However, most real-world problems consist of large datasets (i.e., the medical sector) that cannot be tackled by traditional transfer learning. To address this major issue, the multi-scale transfer learning (MTL) technique is applied to solve these problems.

MTL enhances transfer learning by incorporating multiple pre-trained CNN models within the neural network, whereas traditional transfer learning uses only one pre-trained CNN model. These models process samples at various scales and capture features at different levels of detail. Once features are extracted from each model, they are harmoniously combined to create a comprehensive representation.

MTL is particularly beneficial for data scientists to extract features from the multiple pre-trained CNN models at multiple scales. Both MTL and transfer learning can be employed in medical image classification, treatment planning, disease diagnosis, and anomaly identification. In the healthcare sector, these techniques can be applied to improve diagnostic tools and increase their precision and dependability. Processing medical images at multiple scales enhances the ability to capture the fine features of lesions effectively. MTL contributes to early detection in medical images, facilitating prompt interventions, and ultimately enhancing patient outcomes. Additionally, these models excel at rapidly analyzing vast quantities of medical images, enabling clinicians to make quicker and more accurate decisions.

In this study, we proposed a hybrid "DFU\_MultiNet" framework-based automatic DFU categorization system that is capable of distinguishing between normal and ulcer foot skin from the DFU dataset. The DFU data were first preprocessed and partitioned into train-test sets. Then the training set is fed into the multi-scale transfer learning (MTL) model. Three popular pre-trained CNN models, namely DenseNet201, NasNetMobile, and VGG19, make up this MTL model, which is applied to extract features from samples of foot skin. After that, we integrated all the extracted results through a summing layer and fine-tuned those using two dropout layers, two dense layers, and two batch normalization (BN) layers. Finally, the final dense layer is employed for the DFU classification task.

The crucial points of this research work are described as follows:

1. The “DFU\_MultiNet” framework propose a multi-scale feature fusion technique which outperform the existing models.
2. Proposed framework demonstrates a remarkable performance on classifying all the ulcer and healthy images with omittable miss classification rate.
3. “DFU\_MultiNet” provides a segmentation-free feature extraction technique on a large DFU dataset and show the proficiency in acquiring the high results in DFU dataset. In this technique, no manual feature extraction methods are necessary, distinguishing it from conventional ML approaches. On the other hand, it directly captures pertinent features from the entire image, eliminating the need for a distinct segmentation process.

The other parts of this paper have been structured into several sections. In **Section 2**, we described the literature review. After that, the proposed ‘‘DFU\_MultiNet” framework has been explained in detail, including data pre-processing and MTL model-building strategies in **Section 3**. The experimental setup and consequences are presented comprehensively in **Section 4**. **Section 5** contains a comparative analysis of the outcomes and suggestions for future direction. The final findings of this study are summarized in **Section 6**.

**2 Literature review**

Numerous studies have been published on the classification and detection of DFU images to distinguish between healthy skin and ulcer skin. Most of the authors propose ML and image processing approaches that analyze different features like patterns, hue, and morphological textures. The proposed work's performance depends on the adopted models and training approaches. Various types of DFU-related works are briefly described in this section.

Kaselimi et al. [**16**] (**2022**) introduced a comprehensive review of the existing research on the use of artificial intelligence (AI) in monitoring DFUs, highlighting the advantages of these methods while also acknowledging the challenges in implementing them effectively for remote patient care. Their analysis focused on the imaging methods and optical sensors employed to detect DFUs. The study contemplates both the characteristics of sensors and the physiological aspects of the patients. The image data source recommended a number of monitoring tactics, which limits the use of AI algorithms [**17**].

Thotad et al. [**18**] (**2022**) proposed a DL approach named EfficientNet, for the early forecast and detection of diabetic foot ulcers (DFU). EfficientNet was employed for a sample set whose image size was 844 feet, consisting of diabetic ulcers and healthy skin. In this approach, they built up a robust network by adjusting three crucial properties (resolution, width, and depth) of the CNN model to classify diabetic and normal feet. Their method attained excellent results compared to modern algorithms like GoogleNet, VGGNet (VGG16 and VGG19), and AlexNet. It gave the utmost accuracy, recall, precision, and f1-score of 98.97%, 98%, 99%, and 98%, respectively.

In another method, Juan et al. [**19**] (**2022**) proposed a novel deep (CNN) classifier named DFU\_VIRNet, for an automatic DFU skin classification task. Furthermore, their method focused on estimation maps to identify the probability of risk areas responsible for developing DFU. Two types of samples, namely, visible and invisible, were fed to the proposed scheme for training and testing purposes. The DFU\_VIRNet provided the highest AUC score (0.99301) and ACC (0.97750), beating the recent outcomes.

Doulamis et al. [**20**] (**2021**) proposed a valuable non-invasive device that utilizes photonic-based technology for the treatment of DFUs in diabetic patients. This device employs hyperspectral and thermal imaging to evaluate the ulcer's condition and estimates the biomarkers deoxyhemoglobin and oxyhemoglobin using the imaging technique. Additionally, this novel device was improved by incorporating signal processing methods utilizing DL for improving pixel accuracy and reducing noise using super-resolution approaches.

Das et al. [**21**] (**2021**) proposed a unique framework (DFU\_SPNet), which was constructed from stacked parallel (SP) convolution layers. DFU\_SPNet employed three distinct kernel size modules of SP convolution layers to extract the feature map. Obtaining ACC (97.4%), the DFU\_SPNet outperformed the existing ultra-modern works after evaluating the DFU test dataset utilizing the optimizer (SGD) with a learning rate (1-e2).

Alzubaidi et al. [**22**] (**2021**) proposed four hybrid CNN models for classifying abnormal skin vs. normal skin. The models they developed incorporated traditional CNN layers along with parallel convolutional layers (PCL). Each model has six modules of PCL, but the range of the PCL branches is from 2 to 5. All models extracted the features from the same input, utilizing PCL with different kernel sizes, and then merged the extracted results. Among the models tested, the model with four branches achieved the highest F1 score (95.8%).

Alzubaidi et al. [**23**] (**2020**) proposed a study that utilized a 754-foot DFU dataset from both healthy and diabetic ulcer patients. To automate the categorization of these images, a deep CNN named DFU\_QUTNet was proposed, which differed from conventional CNNs. DFU\_QUTNet was wider but not necessarily deeper. This approach improved the gradient propagation problem since errors were remitted over several different channels.

Tan and Le et al. [**24**] (**2019**) demonstrated that optimizing network resolution, width, and depth can enhance performance and revealed an extensive description of model scaling. They introduced a new scaling mechanism that utilized a compound coefficient to uniformly scale these parameters. By applying this mechanism to a baseline model, they constructed the EfficientNets family of CNN models. This study surpassed other existing ConvNets in accuracy, efficacy, and speed of inference while obtaining an impressive ACC (84.3%) on ImageNet [**25**].

Manu Goyal et al. [**26**] (**2017**) endorsed the potential of conventional computer vision (CCV) features as a practical and cost-effective method for detecting foot ulcers in diabetic patients. They developed a new CNN-based framework called DFUNet to differentiate between DFUs and healthy skin by extracting an image feature map. DFUNet achieved the AUC (0.962) by applying a 10-fold cross-validation approach. It demonstrated better performance than utilizing traditional ML and DL approaches.

Wang et al. [**27**] (**2017**) employed a special capture box to capture DFU images and, on the other hand, identified the exact location of DFU by employing a two-stage SVM classification approach. These two stages were: (i) the segmentation stage, which extracts super-pixels; and (ii) the feature extraction stage, which extracts important features from the image.

Manu et al. [**28**] (**2017**) implemented a method for segmenting DFUs from whole-foot images. Though the method exhibited promising results, the approach had limitations. The first limitation was the impracticality of using a box surface for capturing DFU images due to concerns about potential infection risks in a healthcare setting, which emphasized the importance of exploring alternative approaches for image collection and analysis. The second was its inability to effectively process a large DFU dataset. **Table 1** describes an overview of the published paper.

**Table 1:** A summery table of all the approaches.

|  |  |  |  |
| --- | --- | --- | --- |
| **Paper** | **Dataset size** | **Approach** | **Performance** |
| Thotad et al. [**18**] | 1,688 | EfficientNet | Accuracy = 98.97%, Precision = 99%, Recall = 98%, F1-score = 98% |
| Juan et al. [**19**] | 13,200 | DFU\_VIRNet | Accuracy = 0.9775, Sensitivity = 0.98167, Specificity = 0.97333, Precision = 0.97355, Recall = 0.98167, F1-score = 0.97759, AUC = 0.99301 |
| Doulamis et al.[**20**] | \_ | Hyperspectral and thermal imaging techniques | Early prediction and prognosis of a DFU, understanding  the effect of the biomarkers on DFU |
| Das et al. [**21**] | 3,827 | DFU\_SPNet | Accuracy = 0.964, Sensitivity = 0.984, Specificity = 0.951, Precision = 0.926, Recall = 0.984, F1-score = 0.954, AUC = 0.974 |
| Alzubaidi et al. [**22**] | 17,053 | Hybrid CNN | Precision = 97.3%, Recall = 94.5%, F1-score = 95.8% |
| Alzubaidi et al. [**23**] | 17,053 | DFU\_QUTNet+  SVM | Precision = 0.954, Recall = 0.936, F1-score = 0.945 |
| Tan and Le et al. [**24**] | ImageNet | ConvNet | Accuracy = 84.3% |
| Manu Goyal et al. [**26**] | 22,777 | DFUNet | Accuracy = 0.925, Sensitivity = 0.934, Specificity = 0.911, Precision = 0.945, F1-score = 0.939, AUC = 0.961 |
| Wang et al. [**27**] | 100 | SVM | Sensitivity = 73.3%, Specificity = 94.6% |
| Manu et al. [**28**] | 705 | FCN-16s | Dice for ulcer region (UR) = 0.794 and surrounding skin (SS) = 0.851, Specificity for SS = 0.994, Sensitivity for UR = 0.789 and SS = 0.874 and MCC for UR = 0.785 and SS = 0.852 |

**3 Methodology**

Developing a precise diagnosis system for classifying DFU is a challenging task for clinical purposes. This section explains how we proposed a hybrid ‘‘DFU\_MultiNet” framework for the DFU classification challenge by merging various pre-trained CNN models. **Figure 1 shows the framework of the proposed** ‘‘DFU\_MultiNet”. The framework performs the following operations: extracting DFU samples, loading labels, sample preprocessing, splitting the DFU dataset, data augmentation, training the MTL model, and finally, analyzing statistical parameters utilizing the test set. The overall classification task of our framework is shown step by step in **Algorithm 1**.

|  |
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| C:\Users\shuvo\Downloads\out.png |
| **Figure 1:** Proposed DFU\_MultiNet Framework for DFU classification tasks M1, M2, and M3 are three CNN models: DenseNet201, VGG19, and NasNetMobile, respectively. These three models built a hybrid MTL (multi-scale transfer learning) model. |

**Algorithm 1** Automated diabetic foot ulcer (DFU) classification and detection.

***Input:*** *DFU Training dataset γ1 (70%), Validation dataset γ2 (10%), and Testing dataset γ3 (20%)*

*β = batch size*

*σ = epochs*

*λ = optimizer*

*ƞ = learning rate*

*ε = the number of samples converted into mini-batch size*

***Output:*** *Ω = weight of pre-trained CNN algorithms*

***Begin:***

*1: Convert each sample in the training dataset into a size of 224x224.*

*2: Apply the data augmentation method to enhance the sample size.*

*3: Extract the feature maps from the sample utilizing three pre-trained CNN algorithms, namely, VGG19, NasNetMobile, and DenseNet201.*

*4: Merge the extracted maps by applying the concatenation layer.*

*5: Initialize four fine-tuned CNN layers: batch normalization, dense, dropout, and softmax.*

*6: Set the training parameters: ƞ, σ, λ, β, and ε.*

*7: Train the ‘‘DFU\_MultiNet” framework and calculate the primary weights.*

*8:****for****σ = 1 to σ****do***

*9: Select a mini-batch size ε.*

*10: Forward propagation and evaluation of the binary loss function.*

*11: Backpropagation and improving the weight Ω.*

*12:****end for***

***End***

**3.1 DFU dataset**

In this study, the working dataset named DFU dataset was collected from publicly accessible online repository [**29**], which contains four folders that are original images, patches, test set and transfer-learning images. To train and test our hybrid model, we selected the patch folder from these four folders, which included a total of 1055 skin patches. Out of these patches, 512 were identified as abnormal (ulcers), while the remaining 543 were classified as normal (healthy skin). **Figure 2** displays sample of skin patches. The DFU dataset is split into two sets, train (80%) and test (20%), with the help of the train\_test\_split function. The train\_test\_split function is imported from the "sklearn.model\_selection" package in Python. Then, for the validation set, 10% of the data is split from the train set with the help of the train\_test\_split function. Finally, the DFU dataset (i.e., 1055 samples) has been divided into three distinct phases: 70% (i.e., 760 samples) for training, 20% (i.e., 211 samples) for testing, and 10% (i.e., 84 samples) for validation. **Table 2** provides further details regarding the DFU dataset.

**3.2 Data Preprocessing**

Before inputting the DFU images into the multi-scale transfer learning (MTL) model, various pre-processing stages were implemented. According to transfer learning principles, each image in the DFU dataset was in .jpg format with a size of 224 x 224 pixels and channel RGB. The images were converted into Numpy arrays to enable faster training and reduce memory usage. Additionally, we have also shuffled the dataset for training unordered samples. The DFU dataset has been divided into distinct phases (see **Figure 03(a)**), with 70% (i.e., 760 samples) allocated to training, 20% (i.e., 211 samples) to testing, and 10% (i.e., 84 samples) to validation, respectively. The bar chart (**Figure 3(b))** indicates the number of samples after splitting the data into three phases. Deep networks require a large number of training samples due to their numerous parameters. This amount is increased by applying a powerful technique called data augmentation. This technique can serve various purposes, such as improving the performance of the DFU\_MultiNet framework, handling overfitting matters, and enhancing the robustness of the model. Finally, applying this technique, we enhanced the DFU dataset from 1055 to 6963. In data augmentation techniques involving non-binary (true or false) parameters like rotation angle, shift, zooming, or shearing, these parameter values are usually selected randomly from predefined ranges or distributions. For instance, in image rotation, the rotation\_range argument allows random selection of any degree between 0 and 360. When zooming an image, it is typically scaled within the range of [1 - zoom\_range, 1 + zoom\_range]. Shearing involves selecting a floating-point value from a uniform distribution in the range between 0 and 1. In contrast, flipping an image is binary, represented as True or False. All the augmentation parameters that were used in this study are given in **Table 3**. **Table 4** reveals detailed information about the DFU dataset after the augmentation approach. **Figure 4** shows some examples of augmented images.

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| --- |
| samples334 |
| **Figure 2:** Sample skin images (Healthy and Ulcer) of the DFU dataset. |

**Table 2** Detail information about working data before applying augmentation techniques.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Dataset | Label | Training | Validation | Testing |
|  | Healthy skin | 390 | 43 | 110 |
| DFU | Ulcer skin | 370 | 41 | 101 |
|  | Total samples | 760 | 84 | 211 |

|  |  |
| --- | --- |
| C:\Users\shuvo\Downloads\new1 fig.png C:\Users\shuvo\Downloads\WhatsApp Image 2023-05-13 at 1.55.05 AM (3).jpeg | |
| **Figure 3(a):** Data splitting ratio. | **Figure 3(b):** Number of samples after splitting. |

**Table 3** Data augmentation techniques and parameters.

|  |  |  |
| --- | --- | --- |
| **Number** | **Data techniques** **strategies** | **Parameter values** |
| 1 | Zooming range | 2 |
| 2 | Rotation range | 90 |
| 3 | Shearing range | 0.4 |
| 4 | Width shift range | 0.2 |
| 5 | Height shift range | 0.2 |
| 6 | Horizontal flip | True |
| 7 | Vertical flip | True |

**Table 4** Detail information about working data after applying augmentation techniques.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| Dataset | Label | Training | Validation | Testing |
|  | Healthy skin | 3120 | 344 | 110 |
| DFU | Ulcer skin | 2960 | 328 | 101 |
|  | Total samples | 6080 | 672 | 211 |

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| E:\thesis shuvo\DFU_Thesis\DFU_MultiNet\samples11.png |
| **Figure 4:** Sample augmented images. |

**3.3 Building the MTL Model**

There is a growing interest in applying pre-trained neural networks to a wide range of tasks beyond their initial domains [**30**]. This is particularly valuable in the medical field, where obtaining sufficient labeled data for training DL networks can be challenging [**31**]. To address this, researchers have harnessed the power of networks trained on ImageNet [**32**], a vast database containing over 14 million (M) images in 20,000 categories but used 1.2M images spanning more than 1,000 classes for benchmarking. These classes may be abstract concepts, scenes, objects, and animals. The utilized pre-trained models in this experiment were pre-trained on this vast database and the weights of these networks are determined by this database.

In DL, these pre-trained networks have already acquired valuable representations and features from this vast database. These learned features enhance a model's proficiency in handling unseen data but this advantage is not obtained from the re-trained model. Conversely, training DL models from scratch necessitates computational resources and powerful hardware but these resources can be saved by using pre-trained models. These models are typically constructed upon well-structured architectures that have been greatly fine-tuned and tested. Given these advantages, in this study we opting pre-trained models rather than re-training them from scratch.

To build the MTL model, firstly three pre-trained models DenseNet201, NasNetMobile, and VGG19 are each fed the DFU color images with a size of 224x224 as input so that each model can extract the features of the images separately. Then each of the three GlobalAveragePooling2D layers is separately applied to each model, flattening the respective layers into a vector by averaging the features of each input. Subsequently, these individual vectors are combined into a unified vector by employing the concatenate layer. After that, the integrated features are fine-tuned using six CNN layers. Within these six layers, the first is the dropout layer (dropout rate 0.4), the second is the batch normalization layer, the third is the dense layer (128 units and ReLU activation function), the fourth is the dropout layer with a dropout rate of 0.6, the fifth is the batch normalization layer, and the final layer is the dense layer (2 units and softmax activation function). **Figure 5** depicts the MTL (multi-scale transfer learning) model for classifying DFU images. The MTL model contains 43,077,398 parameters after merging all of the collected features, which is roughly 2, 2, and 10 times more compared to the VGG19, DenseNet201, and NasNetMobile models. The basic explanation of these three adopted models and the fine-tuning approach are given in the following sub-sections.

|  |
| --- |
| C:\Users\shuvo\Desktop\multinet\WhatsApp Image 2023-04-13 at 2.07.01 PM.jpeg |
| **Figure 5:** MTL (multi-scale transfer learning) model to extract features from samples and merge them for diagnosis of ulcer skin. M1, M2, and M3 are three pre-trained DenseNet201, VGG19, and NasNetMobile models, respectively. GAP indicates the global average pooling layer, Concat indicates the concatenation layer, BN indicates the batch-normalization layer, and Drop indicates the dropout layer. |

**3.3.1 DenseNet**

Huang et al. (**2017**) [**33**], initially developed the best pre-trained CNN classification model called DenseNet for obtaining the best accuracy on the ImageNet, CIFAR10, and CIFAR-100 datasets. This model was created using a feed-forwarding model similar to the ResNet model. Such a connection enables the architecture to exchange crucial data within the network, improving model performance and increasing the effectiveness of model training [**34**]. In this study, we employ DenseNet201 as our first feature detector network. This network has 201 deep neural layers in total, each of which is constructed to address overfitting difficulties while working with unordered samples. Additionally, after training this model, it contains a total of 18,321,984 parameters.

**3.3.2 NasNetMobile**

Zoph et al. [**35**] (**2018**) first proposed NasNetMobile, our second feature extractor pre-trained model. On the CIFAR-10 dataset, the NasNetMobile model achieved a 2.4% error rate using a novel regression method called ScheduledDropPath. In this study, this model is trained and tested on 224 x 224 DFU images using approximately 5.3M training parameters. According to Saxena et al. [**36**] (**2019**), there exists an optimized model comprised of core building blocks that have been optimized through reinforcement learning. These blocks incorporate various pooling, convolution, and separable-convolution functions that enhance the overall reliability of the model.

**3.3.3 VGGNet**

Simonyan et al. [**37**] (**2014**) proposed VGGNet, which achieved high performance in image localization and classification, ranking first and second, respectively, at the ILSVRC competition. Compared to the AlexNet architecture, VGGNet exhibited an impressive error rate (8.1%) that is much better than the AlexNet architecture. For this study, we utilized VGG19 as the last feature extractor to build our DFU\_MultiNet framework. The popular VGG19 model is organized by 16 convolution layers and three FC (fully connected) layers. The filter sizes for each convolution layer range from 64 to 512, with a 3x3 window size for all of them. Five blocks make up this model, with the first four convolution layers located in the first two blocks and the remaining twelve located in the next three. An MP (max pooling) layer after each block with a 2x2 window size detects the most important features from the modified activation maps [**38**]. The activation function (ReLU) is applied to each convolution layer. Finally, this model obtained a total of 20,024,384 parameters after training on the DFU dataset, which is more than the other two models.

**3.4 Fine-tuning process**

**Figure 5** illustrates how to integrate three pre-trained CNN models to build the MTL model for categorizing the DFU dataset, utilizing various FC (fully connected) layers. These three models employ GlobalAveragePooling2D simultaneously to flatten into a vector, which is done by computing the average value of the input images. A concatenate layer is then used to merge each vector into a single vector and fed through six additional layers with an activation function (softmax) for fine-tuning purposes for categorizing the DFU dataset. The explanations of each DL layers are given below.

The DL model faces a significant problem known as overfitting, which happens when it over trains on training data and performs poorly on test data [**39**]. We employ two dropout regularization layers to overcome the overfitting situation. During the DFU\_MultiNet framework training, these layers excluded 40 percent and 20 percent of the samples, while also significantly improving training time. Additionally, such a procedure significantly speeds up the DFU dataset training task [**40**].

On the other hand, the inclusion of two BN (batch normalization) [**41**] layers is crucial for the success of our DFU\_MultiNet framework. The main operations of this layer are to rescale and normalize the DFU samples, which makes the model more robust and reliable.

The dense layer also called the FC (fully connected) [**42**] layer connects all neurons between the two layers (previous and current). The main task of this special layer is to process input samples and generate the classification result. In our approach, we employ two FC layers, where the first use ReLU [**43**] and the second uses softmax as an activation function. This final layer predicts the length of the class and generates the DFU prediction. The softmax determines the most relevant features to predict the normal/ulcer class, whose outcome value ranges from 0 to 1, and triggers the neuron accordingly. It can be expressed as the following equation:

Softmax(w)p = …………………………………… (1)

The results of combining various pre-trained models with FC (fully connected) layers are reported in table (see **Appendix A**). This table was obtained during the construction of the "DFU\_MultiNet" framework for DFU classification. That’s why, in this framework, the final FC layer contains two neurons.

**4 Dataset description, performance metrics, and results analysis**

In the experimental setup, hyperparameters utilized in the study, and outcomes achieved by the "DFU\_MultiNet" framework from the DFU dataset are presented in this section. Moreover, a comprehensive comparative discussion between the "DFU\_MultiNet" framework and individual state-of-the-art CNN frameworks is conducted to evaluate the effectiveness of the proposed framework.

**4.1 Dataset description**

The dfu-dataset, which is available online [**29**], was used for training and testing. This DFU dataset has four folders. The "original images" folder contains 493 images of different patients' feet with healthy feet and diabetic ulcers. The photos are from the diabetes center of the Nasiriyah Hospital in southern Iraq [**23**], and it's noteworthy that written consent and ethical approval were diligently obtained from all relevant patients and persons involved in the data collection process. These photos were taken by the experts with an iPad and a Samsung Galaxy Note 8 in a variety of lighting and viewing conditions. The "patches" folder contains 543 normal (healthy skin) and 512 abnormal (ulcer) skin patches, which were cropped from the samples of the "original images" folder with resolution 224x224 pixels. In this experiment, we used "patches" folder samples to train and test our model.

**4.2 Experimental setup**

The proposed ‘‘DFU\_MultiNet” framework was developed with the help of Keras [**44**], for connecting Python [**45**] to the NN (neural network). The experimental setup with parameters described in **Table 5**.

**Table 5** Experimental setup with parameters.

|  |  |
| --- | --- |
| **Item** | **Performance** |
| Platform | Google Colab |
| GPU | Tesla K80 |
| RAM | 64 GB |
| CPU | Intel Core i5-12600K @ 3700 MHz |

**4.3 Performance metrics**

The performance evaluation of the ‘‘DFU\_MultiNet” framework involved the use of various statistical parameters, such as accuracy (ACC), Kappa statistic, F1-score (FS), precision (PRE), Matthews correlation coefficient (MCC), specificity (SPE), sensitivity (SEN), and recall (REC). These parameters were calculated based on the values of false negative (FN), true positive (TP), true negative (TN), and false positive (FP) in the confusion matrix.

In the evaluation of the "DFU\_MultiNet" framework, **TP** refers to the correct identification of positive foot skin. **TN** refers to the accurate identification of negative foot skin. On the other hand, **FP** denotes inaccurate identification of positive foot skin and **FN** refers to its incorrect identification of negative foot skin.

The followings are the performance metrics of the "DFU\_MultiNet" framework:

**Accuracy (ACC):** ACC refers to the relationship between appropriately identified samples and the total samples of the DFU dataset.

……………………… (2)

**Recall (REC):** REC refers to the ability of the “DFU\_MultiNet” framework to successfully identify TP (true positive) samples by calculating the ratio of total positive samples in the DFU dataset.

..………… ……………………….. (3)

**Specificity (SPE):** SPE refers to the ability of the “DFU\_MultiNet” framework to successfully identify TN (true negative) samples by calculating the division of total negative samples of the DFU dataset, also called the TNR (true negative rate).

………………….…..……… (4)

**Precision (PRE):** PRE refers to the ability of the “DFU\_MultiNet” framework to successfully identify TP (true positive) samples by calculating the proportion of successfully predicted positive samples to all the predicted positive examples of the DFU dataset.

…………………..…………… (5)

**F1-Score** **(FS):** The harmonic mean of REC and PRE is known as the FS.

………………………….… (6)

**Matthews correlation coefficient (MCC)**: MCC is a statistical parameter that is applied for binary labeling. The value is bounded from -1 (worst outcome) to 1 (best outcome). It can be expressed as the following equation:

………………….. (7)

**Cohen’s Kappa Coefficient (Kappa):**Kappa is utilized to compare the predicted classes from the DFU\_MultiNet framework with the actual classes in the DFU data. The value is bounded from -1 (worst outcome) to 1 (best outcome). It can be expressed as the following equation:

………………….……… (8)

**4.4 Training and parameter optimization**

**Figure 6**presents the simulation outcomes of the proposed “DFU\_MultiNet” framework, which are extracted during the training phase of the framework.To train the ‘‘DFU\_MultiNet”, specific hyper-parameter values were employed, as outlined in **Table 6**. Optimizer and gradient descent loss functions are two crucial parts for selecting hyperparameters during the framework training. In selecting the optimizer function for our framework, we opted for Adam [**42**] due to its ability to effectively manage sparse gradients in large datasets by combining the desirable properties of RMSProp and AdaGrad optimizers. Considering our work's focus on binary classification, we employed a loss function named binary cross-entropy. Determining an appropriate learning rate is crucial to minimizing the loss function, but this task can be very challenging. In the DL approach, a small learning rate makes the CNN classifier training slower; for this reason, the weight of the model updates is very minimal. To mitigate these issues, we set learning rate = 0.0001, epochs = 50, and batch size = 32 for demonstrating an effective model. **Figure 6** clearly shows that after training the 12th epoch, the "DFU\_MeltiNet" framework offered to achieve 98.68% training accuracy, 97.65% validation accuracy, 6.17% training loss, and 10.03% validation loss, respectively. **Figure 6(a)** further confirms that overfitting was not observed during the DFU\_MeltiNet training process. **Figure 6(b)** confirmed that the curve showed a rapid decrease in the loss value. However, some fluctuations occurred when selecting the narrow batch size.

Using a separate validation set for hyperparameter tuning is crucial as it allows for the selection of the best-performing model by systematically evaluating different hyperparameters. Additionally, it enables efficient searching for optimal hyperparameters and works as an indicator for detecting overfitting problems during the tuning process. If the test set is used for hyperparameter tuning, it may introduce data leakage, potentially affecting the model's performance. Furthermore, such tuning can result in overfitting to the test set. That's why hyperparameters shouldn't be tuned during the evaluation of the test set.

For hyperparameter tuning, the term "Factor" parameter signifies a scaling factor that is applied to adjust a hyperparameter's value. There isn't a fixed range for the "Factor" parameter. For instance, in DL, during tuning learning rates, the range for this parameter is [0.1, 10]. As we have integrated three models, we select smaller learning rates (0.0001) to induce finer updates to the weights during each iteration, which helps to integrate models. When we train our model for too few epochs (like 10) then underfitting occurs, again while training for too many epochs (like 80) then overfitting occurs. For this reason, we select the epoch value 50 based on analyzing the validation accuracy and training loss (see **Figure 6**). In DL, the batch size (BS) of 32 is a good initial point and a thumb rule. It strikes a balance between computational efficiency and model accuracy. Large BSs can expedite training but may risk overfitting and reduced accuracy, whereas smaller BSs can be time-consuming and computationally expensive.

**Table 6:** Training parameter with value for the ‘‘DFU\_MultiNet” framework.

|  |  |  |
| --- | --- | --- |
| **Number** | **Parameter** | **Value** |
| 1 | Optimizer | adam |
| 2 | Learning Rate | 0.0001 |
| 3 | Minimum Learning Rate | 1e-7 |
| 4 | Decay | 0.00001 |
| 5 | Patience | 5 |
| 6 | Factor | 0.2 |
| 7 | Loss Function | binary\_crossentropy |
| 8 | Metrics | accuracy |
| 9 | Batch Size | 32 |
| 10 | Epochs | 50 |

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| **Figure 6.** Training improvement of the proposed DFU\_MultiNet framework: (a) training versus validation accuracy plot (higher values indicate better performance), and (b) training versus validation loss plot (lower values indicate better performance). |

**4.5 Results analysis**

**Figure 7** presents the CM (confusion matrix) and ROC (receiver operating characteristic) curves for the DFU dataset, utilizing the "DFU\_MultiNet" framework. The framework merged three renowned transfer learning algorithms, namely DenseNet201, VGG19, and NasNetMobile. By leveraging fusion features, the proposed approach effectively categorizes whether a diabetic foot is ulcerated or healthy. **Figure 7(a)** reveals that the proposed "DFU\_MultiNet" framework accurately classifies 101 ulcer skin images and 108 healthy skin images. Remarkably, the framework only misclassifies two instances of healthy skin. Notably, a significant advantage of the framework is its flawless performance in misclassifying no ulcer skin within the DFU dataset. The "DFU\_MultiNet" framework demonstrates remarkable consistency and stability, as evidenced by the achieved AUC (0.99091) in **Figure 7(b)**. This high AUC score indicates the model's strong performance. Additionally, individual evaluations of all models on the DFU dataset further enhance the robustness of the "DFU\_MultiNet" framework. **Table 7** proves the superiority of the 'DFU\_MultiNet' framework between the 'DFU\_MultiNet' framework and five additional CNN models. The results indicate that the "DFU\_MultiNet" framework attains impressive metrics such as precision of 1.00, recall of 0.982, f1-score of 0.991, kappa of 0.981, and MCC of 0.981, surpassing all other ultramodern models. Notably, DenseNet201 and VGG16 also demonstrate strong performance with accuracy values of 0.976 and 0.981, respectively. Meanwhile, VGG19 and MobileNet exhibit comparable performance in their ability to detect foot skin.

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| **Figure 7.** (a) Confusion matrix (b) ROC curve for the ‘‘DFU\_MultiNet” framework. | |

**Table 7:** Results attained from the ‘‘DFU\_MultiNet” framework and five individual fine-tuned algorithms on DFU Dataset

|  |  |  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- | --- | --- |
| **Model** | **Accuracy** | **Precision** | **Recall** | **F1score** | **Specificity** | **AUC** | **Error Rate** | **MCC** | **Kappa** |
| VGG19 | 0.867 | **1.00** | 0.745 | 0.854 | **1.00** | 0.873 | 0.133 | 0.764 | 0.737 |
| VGG16 | 0.981 | **1.00** | 0.964 | 0.981 | **1.00** | 0.982 | 0.019 | 0.963 | 0.962 |
| NasNetMobile | 0.773 | **1.00** | 0.564 | 0.721 | **1.00** | 0.782 | 0.227 | 0.618 | 0.553 |
| DenseNet201 | 0.976 | **1.00** | 0.955 | 0.977 | **1.00** | 0.977 | 0.024 | 0.954 | 0.953 |
| MobileNet | 0.867 | 0.894 | 0.845 | 0.869 | 0.891 | 0.868 | 0.133 | 0.736 | 0.735 |
| DFU\_MultiNet | **0.991** | **1.00** | **0.982** | **0.991** | **1.00** | **0.991** | **0.009** | **0.981** | **0.981** |

**5 Discussion**

The incidence of diabetic foot infections (DFI) and Diabetes-related problems can be caused by not obeying a healthy diet and no adequate safety precautions among individuals with diabetes. Ensuring proper guidance is provided to diabetic patients and their caregivers becomes imperative in addressing these challenges. Additionally, new techniques for diagnosis, therapy, and forecasting have been created as a result of applying technology to control diabetes. This framework has been built based on the heterogeneous parallel ensemble DL architecture that has learned features in parallel from input samples through three pre-trained models (i.e. DenseNet121, VGG19, and NasNetMobile). This powerful technique can be leveraged across other clinical machine-learning applications. By utilizing this framework, healthcare professionals can make better decisions regarding patient enrollment in clinical trials, optimize drug development processes, and seamlessly integrate data from various sources. As a result, this network will play a vital role in enhancing the efficiency and effectiveness of clinical research and healthcare systems. As shown in **Table 7**, the results from the suggested framework are more dependable and robust than those from the current models. This study suggests a novel method for classifying the skin of diabetic-affected feet by employing a hybrid "DFU\_MultiNet" framework on DFU images. **Table 8** provides a summary of the performance of the "DFU\_MultiNet" framework in comparison to previous research that employed the same dataset but with various structures, depths, and parameters. Our approach, which combines predictions from multiple pre-trained models, offers several advantages for handling imbalanced datasets. It effectively mitigates overfitting and minimizes the possibility of noise impacting the minority label. This technique also assigns greater weight to minority class, resulting in improved classification for the imbalanced dataset. Since we used three different models, if one model fails to extract features from some data points in the dataset, others extract those features, which can enhance the ability to adapt to changes in the imbalanced dataset. Additionally, training these models in parallel on the same dataset significantly reduces the overall training time when dealing with an imbalanced dataset. **Table 8** makes it obvious that, when compared to the earlier studies, the suggested framework offers the highest accuracy for the diagnosis of diabetic foot ulcers (DFU). Remarkably, the combination of all the pre-trained algorithms enables the framework to achieve a classification accuracy of 99% for the DFU dataset. The ROC curve presented in **Figure 8** compares the performance of the DFU\_MultiNet, a proposed framework, with several transfer learning networks such as DenseNet201, VGG16, VGG19, NasNetMobile, MobileNet, and DFU\_MultiNet, using the same data partition. The results demonstrate that the DFU\_MultiNet achieves an outstanding result over the standard transfer learning networks in accurately classifying ulcers versus healthy samples. **Figure 9** exhibits some of the diabetic-affected foot skin samples that are accurately predicted with the help of the novel DFU\_MultiNet framework.

Adapting the "DFU\_MultiNet" approach for clinical use involves several crucial steps and considerations to ensure its effectiveness and safety within a healthcare setting. Initially, the dataset must undergo annotation with ground truth labels to distinguish healthy feet from those with diabetic foot ulcers (DFUs). To enhance transparency and understanding, various explainable AI techniques (e.g. LIME and SHAP) are employed to interpret the framework, providing clinicians with insights into the model's feature utilization for classification. Subsequently, integration of the "DFU\_MultiNet" model into existing clinical systems or diagnostic tools utilized by healthcare professionals is imperative. The model's predictions must be presented in a clear and actionable format, ensuring that clinicians can readily comprehend and act upon the diagnostic results. Continuous monitoring of the model's performance is essential, encompassing diagnostic accuracy, treatment planning, and patient outcomes assessment within real clinical scenarios.

In this clinical context, clinicians can employ the framework as a prospective Computer-Aided Diagnostic (CAD) tool. They can effortlessly input DFU images into the CAD tool's interface, where the "DFU\_MultiNet" framework undertakes feature extraction by analyzing the images. These extracted features are then amalgamated through a summing layer, generating a comprehensive representation of the image for diagnostic evaluation and decision-making.

|  |
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| C:\Users\Hp\Downloads\ROC curve1.png |
| **Figure 8**. ROC curve for several transfer learning algorithms. |

**Table 8:** Comparison outcome of the ‘‘DFU\_MultiNet” framework with SOTA techniques on the DFU dataset.

|  |  |  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- | --- | --- |
| **Paper** | **Accuracy (%)** | **Precision (%)** | **Recall (%)** | **F1-score (%)** | **Classification** **Method** | **Training and validation**  **Sample** | **Testing**  **Sample** |
| Thotad et al. (2022) [**18**] | 98.97 | 99 | 98 | 98 | EfficientNet | 1,350 | 338 |
| Juan, et al. (2021) [**19**] | 97.8 | 97.4 | 98.2 | 97.8 | DFU\_VIRnet | 12,600 | 600 |
| K. Das et al. (2021)[**21**] | 96.4 | 92.6 | **98.4** | 95.4 | DFU\_SPNet | 3,491 | 336 |
| Alzubaidi et al. (2021) [**22**] | - | 97.3 | 94.5 | 95.8 | Hybrid CNN | 16,731 | 322 |
| Alzubaidi et al. (2020) [**23**] | - | 95.4 | 93.6 | 94.5 | DFU\_QUTNet+  SVM | 16,731 | 322 |
| Goyal et al. (2017)[**26**] | 92.5 | 94.5 | - | 93.9 | DFUNet | 22,605 | 172 |
| **Proposed Framework** | **99.1** | **100** | 98.2 | **99.1** | **DFU\_MultiNet** | 6,752 | 211 |

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| **Figure 9.** ‘‘DFU\_MultiNet” successfully examined some DFU samples. |

**5.1 Limitations of the Study**

In this framework, it is feasible to diagnose whether the conferred sample is an ulcer or healthy only. It cannot provide a real-time assessment of pain severity or complexity levels.

To maximize the potential of this framework in clinical areas with limited computing power, future works should prioritize the development of distributed training algorithms and federated learning, alongside the creation of interpretable AI systems. Additionally, the development of DL models that operate efficiently on edge devices and mobile platforms will be crucial for clinical areas where there is not significant computing power available.

To test the performance of the DFU\_MultiNet approach for DFU classification, we have trained this approach as well as five individual pre-trained models (i.e. DenseNet121, VGG19, VGG16, MobileNet, and NasNetMobile) on the same dataset with the same splitting ratio. These five pre-trained models are trained as follows: firstly, input images with a 224x224 size are fed into these models to produce a set of output feature maps by extracting features from the input image. A GlobalAveragePooling2D layer is then applied after each model to reduce the produced output feature map to a one-dimensional vector. This is done by taking the mean of all the values in the feature map. After that, six fine-tuning layers are added after the GlobalAveragePooling2D layer, one after another which follows in dropout->batch\_normalization->dense and again dropout->batch\_normalization->dense manner. These six layers are described in **Section 3.3**, which is also used to build the MTL model.

Though the proposed model exhibits good performance on this DFU dataset, it will provide a better and more accurate result on a larger dataset. During training on a large dataset, this framework will learn unique feature maps from different samples, which makes it a powerful ulcer detector tool. In healthcare and medical imaging, this powerful tool promises to detect disease more precisely and earlier, enabling better treatment planning and patient outcomes. In the future, this framework will be applicable for accurate quantification of different parameters within DFU samples, such as ulcer size, tissue density, volume, or growth rate. This quantitative data will facilitate ongoing patient monitoring and enhance treatment planning.

**6 Conclusion**

Regular foot examinations are essential for individuals with diabetes to identify any potential lesions, in addition to undergoing comprehensive assessments for peripheral arterial and neuropathy problems, as these conditions have the potential to result in the formation of ulcers or wounds. DFU can be prevented with regular foot exams, glucose control, patient concealment, suitable footwear, and timely treatment for pre-ulcerative infections. In this task, we have offered an innovative hybrid framework named ‘‘DFU\_MultiNet” to diagnose diabetic-affected skin from foot samples more accurately and consistently. The ‘‘DFU\_MultiNet” framework is based on a feature extraction and fine-tuning approach that permits various pre-trained CNN algorithms to extract and merge feature maps in parallel for DFU classification. It is developed in a balanced way that can control diverse DFU datasets. The outcomes of the experiments exhibit that the "DFU\_MultiNet" framework, which surpasses both separate CNN pre-trained algorithms and all other modern algorithms reported in the published work, attains 99% accuracy. This hybrid framework outperformed the most recent DFU classification methods. Considering the promising outcomes, we have strong confidence in the potential of our "DFU\_MultiNet" framework as an excellent tool to aid doctors in efficiently detecting and diagnosing DFU. Additionally, it performs admirably in locating ulcer skins, improving the likelihood of survival.

In future studies, this hybrid framework should be expanded to detect and classify the DFU into ischemia, neuropathy, osteomyelitis, or Charcot arthropathy.

**Declaration**

**Data Availability**

The data supporting this study's findings are available from the corresponding author upon reasonable request.

**Funding**

None

**Declaration of Interests**

The authors declare that there are no conflicts of interest.

**Ethical Approval**

Not required

**Consent to participate**

Not required

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**Appendix A**

Details of the ‘‘DFU\_MultiNet” framework.

|  |  |  |  |
| --- | --- | --- | --- |
| **Layer (type)** | **Output Shape** | **Param #** | **Connected to** |
| input\_1 | (224, 224, 3) | 0 |  |
| densenet201 | (7, 7, 1920) | 18321984 | input\_1[0][0] |
| NASNet | (7, 7, 1056) | 4269716 | input\_1[0][0] |
| vgg19 | (7, 7, 512) | 20024384 | input\_1[0][0] |
| global\_average\_pooling2d | (1920) | 0 | densenet201[0][0] |
| global\_average\_pooling2d | (1056) | 0 | NASNet[0][0] |
| global\_average\_pooling2d | (512) | 0 | vgg19[0][0] |
| concatenate\_4 | (3488) | 0 | global\_average\_pooling2d[0][0] global\_average\_pooling2d\_1[0][0]  global\_average\_pooling2d\_2[0][0] |
| dropout | (3488) | 0 | concatenate\_4[0][0] |
| batch\_normalization | (3488) | 13952 | dropout[0][0] |
| dense | (128) | 446592 | batch\_normalization[0][0] |
| dropout\_1 | (None, 128) | 0 | dense[0][0] |
| batch\_normalization\_1 | (None, 128) | 512 | dropout\_1[0][0] |
| dense\_1 | (None, 2) | 258 | batch\_normalization\_1[0][0] |
| Total params: 43,077,398  Trainable params: 42,804,372  Non-trainable params: 273,026 | | | |