Parkinson’s Disease Classification based on Enhanced Ensemble Learning and Brain MRI

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ABSTRACT:
Parkinson’s Disease (PD) is a neurological disorder that causes progressive loss of brain cells. Despite the fact that there is no known cure for this neurodegenerative disease at present, early diagnosis and treatment may improve the quality of life. Magnetic Resonance Imaging (MRI) detects structural changes related to dopamine deficiency in PD. To categorize MRI scans as Healthy Control (HC) or PD, this study proposes an ensemble of Deep Convolution Neural Network (DCNN) models. Initially, we have used DCNN models using augmentation and transfer learning, to classify as PD or HC. In the next stage, we applied a classifier fusion ensemble approach to enhance the overall result of the classification model. The proposed model is trained using the data collected from PPMI database, while assessed on custom dataset which is created using the data collected from Lalitha Super Specialty Hospital (LSSH). Finally, it was observed that the developed ensemble model produced an outstanding performance by plotting an overall accuracy of 99%, while the transfer learned EfficientNet B1 DCNN model stood in the second position, achieving a remarkable accuracy of 98%. This study serves as a significant step forward, providing valuable insights for researchers and clinicians engaged in the domain of clinical image assessment using deep learning techniques.

KEYWORDS: Deep Learning, Ensemble Learning, Neurodegeneration, Parkinson’s Disease, Transfer Learning.

1. INTRODUCTION
Parkinson’s disease (PD) is a progressive neurodegenerative disorder, characterized by the preliminary harm caused in the substantia nigra region of the brain. It occurs when the dopamine-generating neurons become damaged or die, because of various factors such as genetics, environmental factors and lifestyle choices [1]. People with PD begin to experience tremors, and difficulty in speaking, writing, walking, or completing other simple tasks. There are several other symptoms like sweating, soreness, drooling of saliva, pill rolling, anxiety, hysteria and sleep interference. The PD diagnosis can be made using neuroimaging techniques like Magnetic Resonance Imaging (MRI), Single-Photon Emission Computed Tomography (SPECT), and Positron Emission Tomography (PET). These techniques identify the structural and functional alterations that occur in the brain, as the disease progresses. Substantial reduction in grey matter strength is quite common in people with PD, compared to those of Healthy Controls (HC) and it is evident from the bilateral nigral hyperintensity in the substantia nigra (directional arrows) regions of Fig. 1, 2. These methods are able to discover the problem only when 80% of the neurons have degenerated and the person exhibits peculiar indications like tremor, finger tapping, pill rolling, hallucinations, drooling of saliva, posture bent, anxiety, difficulty in speaking, writing, and walking. A dependable, economical, and congenial Computer-Aided Diagnostic (CAD) method is necessary to identify PD and observe its advancement using MRI, which could be valuable for medical professionals. Deep learning algorithms produce contemporary results when several tasks related to computer vision are implemented [5]. Even though DCNNs require a significant portion of data for training the network and powerful GPUs to speed up the learning process, they demonstrated remarkable outcomes in a wide scope of diagnostic imaging applications.
Ensemble learning [58, 59] combines the predictions from multiple models, maximizing the accuracy and robustness by leveraging their unique strengths. It reduces errors, improves generalization, enhances stability, and boosts prediction effectiveness. Therefore, in this work, an image-based classification system using DCNN, transfer learning and ensemble approach for detecting PD and HC is developed on a GPU and tested on a custom data set created from the data obtained from LSSH. The study has several compelling advantages, as follows:

1. The early detection of PD can be accomplished through image-based computer vision techniques, eliminating the need for extensive clinical tests. This approach offers the advantages of reducing the associated costs and saving valuable time in the diagnostic process.
2. Facilitating real-time clinical comparisons in the PD diagnosis process by incorporating a cutting-edge real-time custom dataset created from the data obtained from LSSH, for the testing phase.
3. A meticulous comparison of the proposed approach conducted with state-of-the-art methodologies positions this study at the forefront in the field of research.

1.1. Motivation

The quality of life of a PD person is negatively impacted by both the movement and non-movement indications connected to the process of aging [2, 62]. At present, there are no established blood or laboratory tests that can detect PD and its advancement. In the initial stages of PD, it is essential to take the appropriate preventative measures, which can halt or slow down the progression of the disease. The doctors look at a patient’s medical history and perform a cognitive evaluation using Unified Parkinson’s Disease Rating Scale (UPDRS) or Hoehn and Yahr (H&Y) scale, for initial diagnosis [3]. Skilled therapists play a vital role in the effectiveness of therapeutic tests, but their decisions may be subjective. Obtaining all the required information is a lengthy process that requires the involvement of many individuals.

1.2. Contributions

The major contributions made by this research for improving prediction accuracy are as follows:

1. Data augmentation technique helps to overcome the problem of imbalanced class distribution in the dataset. This approach is superior to the restricted availability of MRIs of PD patients and a noticeable improvement in overall performance can be observed.
2. The proposed DCNN model, based on whole-brain analysis, utilizes spatial structures autonomously, avoiding the need for hand-crafted features, thus making the design as
3. A remarkable increase in performance can be observed when ensemble learning with classifier fusion is applied on the trained DCNN models, compared to individual models.
4. Accuracy enhancement can be observed when the preprocessing operations are applied on images, followed by
DCNN model training, rather than training the raw images directly.
5. Implementing efficient DCNN models, namely EfficientNet B1, ResNet-50, ResNet152V2 and MobileNetV2 using transfer learning and conducting a thorough investigation and comparative analysis to analyze the efficiency of the proposed study, compared to the state-of-the-art works.
6. Facilitating real-time clinical comparisons by incorporating a real-time custom dataset created from the data obtained from LSSH, for the testing phase and also by interpreting the developed model.
7. Providing a comprehensive perspective on classification and analysis applications within the domain of PD diagnosis, intended to benefit PD diagnosis researchers and developers.

2. RELATED WORKS
An early-stage diagnosis approach for PD was suggested by [6], and it involved employing a Joint Function Sample Selection (JFSS) technique to select the most beneficial features to develop a model. For evaluation purposes, fabricated and openly available MRI brain scan files of PD are utilized in order to display a high precision ranking. [7] designed a therapeutic assistance program that makes use of an Artificial Neural Network (ANN). Measurements of DCNN gathered from brain detriments were used to estimate PD, for the purpose of this investigation. DCNN le-Net with transfer learning was used by Sivaranjini et al., [8] to classify MRI Brain scans of persons with PD and HC using PPMI dataset. An accuracy of 88.9% was achieved with this method. Lei et al. [9] came up with the idea of a sparse feature selection model and reported that it had an accuracy of about 80%. To identify distinguishing characteristics, Salvatore et al. [10] looked at MRIs of people with healthy brains, people with PD and people supranuclear palsy. Next, they conducted a Principal Components Analysis (PCA) to identify the pertinent characteristics, and then they fed those features into a SVM classifier for data classification. Experiments to classify PD were carried out by Brahimi et al. [11] using characteristics based on structure and superficial connections, as well as an SVM classifier. An accuracy of 92.6 was achieved from their work. Transfer learning concept and the InceptionV3 model were incorporated by Quan et al. [12], in their experiment of identifying PD. They obtained 98% score of accuracy. From the database of PPMI, Arman et al. [13] selected 64 PD cases, with 38 males and 26 females making up the total of 64. The non-imaging and imaging factors that were utilised to predict motor results were combined using Random Forest (RF) analysis, comprised of 5000 trees. By utilizing this technology, a more accurate early prediction of PD was made possible. The inclusion of radiomic characteristics to traditional measures resulted in a substantial improvement in the prediction of outcome, which in turn resulted in a reduction in the absolute error of predicting MDS-UPDRS-III. The connection of Single Nucleotide Polymorphisms (SNP) with imaging traits increased the statistical power of the study and allowed for the examination of a wider variety for the study that was published by Choi et al. [14]. A DCNN framework along with PD Net model was used of SNP. Data were gathered from PPMI and Seoul National University Hospital (SNUH) cohort, on 431 people diagnosed with PD, 193 HC and 77 patients diagnosed with Scans Without Evidence of Dopaminergic Deficit (SWEDD). Computerized quantification of the DAT Binding Ratio (BR) was carried out as a usual approach for perceptible analysis on the SPECT data. Prashanth et al. [15] used data from the PPMI database. All of the individuals with PD are considered to be in the early stage (Hoehn and Yahr (H&Y) stage 1 or 2), and all of the SWEDD subjects, who are newly identified PD patients based on clinical symptoms but have normal dopaminergic imaging, exhibit early-stage PD symptoms. Binary classification was implemented, employing approaches such as SVM, boosted trees, random forests, and naive Bayes. The classification was early PD versus normal or SWEDD. Using data from the PPMI database, Rumman et al. [16] analyzed 200 SPECT images. Image processing and ANN methodologies were successful in achieving an accuracy of 94%. This method requires a comparatively tiny sum of computing resources, and it can assist a physician in shortening the lengthy procedure of diagnosing PD. Data obtained from the PPMI were utilized in the study of Ortiz et al [17]. A total of 269 DaTSCAN images, were analyzed. DCNN's LeNet-5 and AlexNet designs were used in the process of determining the individuals with PD and HC. Integral Normalization and feature Extraction (INE) Iso-surfaces were used in this investigation. Iso-surfaces resulted in a reduction in the amount of data by 95% while maintaining the highest quality of the information. Chakraborty et al. [18] obtained 3T T1-weighted MRI images from the PPMI database of a total of 406 participants. The DCNN model was utilised in order to train the network. Bayesian Sequential Model-based Optimization (SMBO) method is utilised for the purpose of selecting the ideal collection of hyperparameters, with the goal of preserving the generalizability of the overall model and achieving the highest possible objective score. Magesh and colleagues [19] collected 642 DATSCANS that have been labelled as PD or non-PD and have been trained on a DCNN (VGG-16) via transfer learning. The utilization of intensity normalization resulted in a precision of 95.2%. The model's goals were to make an early diagnosis for PD quicker and more intuitive. Using the Alex-Net Architecture Model with transfer learning and Generative Adversarial Networks (GAN) based data augmentation, Kaur et al. [20] achieved an accuracy of 89.23%. When combined with transfer learned classifier, the data-augmentation strategy is shown to result in an improvement of up to 3% accuracy. Mohammed et al. [21] used AlexNet architecture and 10-fold cross-validation and evaluated model performance. The network was able to achieve
The fact that SPECT imaging technology is not readily available in most hospitals due to the fact that it is nuclear in nature and has a low resolution makes it difficult to visualize the basal ganglia in people with PD. Perhaps, the primary drawback of this study. Pianpanit et al. [22] utilised four distinct DCNN architectures. 159 HC and 448 images of people with dementia were obtained from PPMI. The SShapley Additive exPlanation (SHAP) interpretation approach performed significantly better than the other methods when it came to differentiating between PD and HC individuals and delivered the highest quality heatmap at the location of the uptake depletion. Chien et al. [23] collected 205 DAT-SPECT PD images from one database and 52 images from another. An active contour segmentation to train the network, and a Transfer learned AlexNet for learning and classification, were implemented. PD and non-PD groups were differentiated using the Specific Binding Ratio (SBR) and the ASymmetry Index (ASI) with machine-learning and deep learning approaches. Dhanalakshmi et al. [24] received an aggregate of 1390 DaTscan images from PPMI. The DenseNet-121 model was employed exploring the region of interest (ROI), extracted from the images. The Contour Edge Detection algorithm was utilised to identify the borders of the objects in the images. Table 1 provides a review of the existing research works, for PD detection. The Previous research [6, 7, 9, 12] has examined the progression of PD depending on the selected ROI, which arises the issue of being prone to knowledge loss, limited knowledge and segmentation mistakes, which can be a significant obstacle.

Table 1. Study of Current Methodologies for PD Classification.

<table>
<thead>
<tr>
<th>Ref / author</th>
<th>Data Base</th>
<th>Count</th>
<th>Type of data</th>
<th>Method</th>
<th>Accuracy (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>[7] Amoroso et al.</td>
<td>PPMI</td>
<td>82 PD, 100 HC</td>
<td></td>
<td>ANN</td>
<td>83</td>
</tr>
<tr>
<td>[8] Sivarajnimi et al.</td>
<td>PPMI</td>
<td>123 PD, 56 HC, 29 SWEDD</td>
<td></td>
<td>SVM classifier</td>
<td>86.37</td>
</tr>
<tr>
<td>[12] Quan et al.</td>
<td>PPMI</td>
<td>449 PD, 210 HC</td>
<td></td>
<td>Deep CNN using transfer learning</td>
<td>96.4</td>
</tr>
<tr>
<td>[14] Choi et al.</td>
<td>PPMI, SNUH</td>
<td>PPMI: 379 PD, 170 HC, SNUH: 72 PD, 10 PT</td>
<td></td>
<td>Deep CNN with PD Net</td>
<td>PPMI-96 SNUH-98.8</td>
</tr>
<tr>
<td>[15] Prashanth et al.</td>
<td>PPMI</td>
<td>427 PD, 208 HC, 80 SWEDD</td>
<td></td>
<td>SVM, boosted trees, RF, naive Bayes</td>
<td>97.29</td>
</tr>
<tr>
<td>[16] Rumman et al.</td>
<td>PPMI</td>
<td>100 PD, 100 HC</td>
<td>SPECT / Selected ROI</td>
<td>Image processing and ANN</td>
<td>94</td>
</tr>
<tr>
<td>[22] Pianpanit et al.</td>
<td>PPMI</td>
<td>448 PD, 159 HC</td>
<td></td>
<td>Deep PD Net</td>
<td>96</td>
</tr>
<tr>
<td>[21] Chien et al.</td>
<td>PPMI</td>
<td>105 PD, 100 HC</td>
<td></td>
<td>TL AlexNet</td>
<td>86</td>
</tr>
<tr>
<td>[24] Dhanalakshmi et al.</td>
<td>PPMI</td>
<td>1160PD, 230 HC</td>
<td></td>
<td>DenseNet-121</td>
<td>99.2</td>
</tr>
<tr>
<td>[13] Arman et al.</td>
<td>PPMI</td>
<td>64 PD</td>
<td></td>
<td>RF analysis for prediction.</td>
<td>P&lt;0.001</td>
</tr>
<tr>
<td>[25] Ortiz et al.</td>
<td>PPMI</td>
<td>158 PD, 111 HC</td>
<td></td>
<td>LeNet and AlexNet</td>
<td>95.1</td>
</tr>
<tr>
<td>[26] Majlesi et al.</td>
<td>PPMI</td>
<td>430 PD, 212 HC</td>
<td></td>
<td>VGG16 using TL</td>
<td>95.2</td>
</tr>
<tr>
<td>[27] Castillo et al.</td>
<td>PPMI</td>
<td>168 PD, 194 HC, 26 SWEDD</td>
<td></td>
<td>Ensemble classification</td>
<td>64.5</td>
</tr>
<tr>
<td>[63] Aditi et al.</td>
<td>PPMI</td>
<td>70 Mild PD, 70 Moderate PD, 70 Severe PD, 70 HC</td>
<td></td>
<td>Deep Learning (YOLO v8)</td>
<td>94</td>
</tr>
<tr>
<td>[10] Salvatore et al.</td>
<td>PPMI</td>
<td>56 PD, 28 HC</td>
<td></td>
<td>SVM classifier</td>
<td>85.8</td>
</tr>
<tr>
<td>[18] Chakraborty et al.</td>
<td>PPMI</td>
<td>203 PD, 203 HC</td>
<td></td>
<td>CNN with Bayesian SMBO method</td>
<td>95.29</td>
</tr>
<tr>
<td>[20] Kaur et al.</td>
<td>PPMI</td>
<td>67 PD, 85 HC</td>
<td></td>
<td>TL AlexNet + data augmentation</td>
<td>89.23</td>
</tr>
<tr>
<td>Proposed Approach</td>
<td>PPMI, LSSH</td>
<td>PPMI: 40 PD, 30 HC, LSSH: 20 PD, 10 HC</td>
<td></td>
<td>EL with TL + DCNN + augmentation</td>
<td>99</td>
</tr>
</tbody>
</table>

a EL-Ensemble Learning; TL-Transfer Learning; aug-augmentation; PT-Parkinsonian Tremor; RF-Random Forest;

3. PROPOSED SYSTEM

Transfer learning [26], data augmentation [27], fine-tuning [28] and ensemble learning [29,58,59] are some examples of processes that are often used to decrease the gap as much as possible between training and validation mistakes. This research overcomes the discussed limitations by making use of the subject’s whole brain in the study rather than using segments, thus keeping all the essential details intact. As a novel aspect, an Optimal Weighted Priority Mean (OWPM) ensemble method using classifier fusion mechanism was implemented in this work, for the very first time. The entire process flow of the suggested approach can be observed from Fig. 3. 4. The proposed method uses a neural network, to which brain MR Images are fed as input. In the initial part of the process, the images are pre-processed to remove some of the background noises. In the next stage, data augmentation techniques are carried out to handle the issue of compact size of the dataset as well the data imbalance, besides improving the efficacy of the suggested method. In the next phase,
Fig. 3. The process flow of the suggested approach.

Fig. 4. Transfer-learned DCNN training/testing process.
transfer learned methodology associated with EfficientNetB1, ResNet-50, ResNet152V2, MobileNetV2 are considered and some of the rear layers are transformed to equal the number of classifications, suitable for this application. This is done in preparation for the fourth step, which is the actual classification of the images. In the next step of the process, the suggested model's performance is assessed using test MR scans of HC and people with PD. Finally, the ensemble method is applied to the developed DCNN models and performance analysis is analyzed and compared.

3.1. Materials and Methods

The data for the proposed approach was sourced from the esteemed PPMI database (www.ppmi-info.org/data), which stands as an international landmark and multicenter endeavor aimed at researching biomarkers underlying the progression of PD. 45 images of PD and 25 images of HC were collected from the PPMI, contributing to a total of 70 images. These images belong to the persons, who are aged between 40 and 50 years. These images were used for training the classification models. The data for testing was collected from LSSH. A total of 30 images, of which 15 belongs to PD and 15 belongs to HC categories were obtained, satisfying our criteria that the persons are aged between 40 and 50 years of age. These collected images underwent interpretation by a panel of two experienced nuclear medicine physicians. Table 2 can be observed to find these details. Visual assessment of the images based on the density of Dopamine Transporter (DAT) led to the classification of subjects into a group with reduced DAT density (PD) and another with normal DAT density (HC). The MRI scans selected for this study were meticulously chosen, adhering to specific imaging protocols outlined in Table 3. The training data is augmented using GAN an testing data is augmented using different augmentation mechanisms (rotation, flipping, resizing, etc.).

Table 2. Data Split.

<table>
<thead>
<tr>
<th>Category /Dataset</th>
<th>Real Data</th>
<th>Augmented Data</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>HC</td>
<td>PD</td>
</tr>
<tr>
<td>Training / PPMI</td>
<td>25</td>
<td>45</td>
</tr>
<tr>
<td>Validation / LSSH</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Test / LSSH</td>
<td>5</td>
<td>5</td>
</tr>
<tr>
<td>Total</td>
<td>40</td>
<td>60</td>
</tr>
</tbody>
</table>

Table 3. Specifications of the acquired MRI scans.

<table>
<thead>
<tr>
<th>Image Parameters</th>
<th>PPMI</th>
<th>LSSH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Modality/Weighting</td>
<td>MRI / T1</td>
<td>MRI / T1</td>
</tr>
<tr>
<td>Research Group</td>
<td>HC, PD</td>
<td>HC, PD</td>
</tr>
<tr>
<td>Visit</td>
<td>Initial, Preliminary</td>
<td>Initial, Preliminary</td>
</tr>
<tr>
<td>Dimensions (pixels)</td>
<td>224×224×178</td>
<td>224×224×178</td>
</tr>
<tr>
<td>Interslice Gap</td>
<td>1.0 mm</td>
<td>1.0 mm</td>
</tr>
<tr>
<td>Slice Thickness</td>
<td>1.0 mm</td>
<td>1.0 mm</td>
</tr>
<tr>
<td>Spacing</td>
<td>1.0 × 1.0 × 1.0 mm</td>
<td>1.0 × 1.0 × 1.0 mm</td>
</tr>
<tr>
<td>Acquisition Plane / Type</td>
<td>Axial / 3D</td>
<td>Axial / 3D</td>
</tr>
<tr>
<td>Gender*</td>
<td>PD: M-32, F-13</td>
<td>PD: M-12, F-03</td>
</tr>
<tr>
<td></td>
<td>HC: M-16, F-09</td>
<td>HC: M-09, F-06</td>
</tr>
<tr>
<td>Age</td>
<td>Between 40-50 years</td>
<td>Between 40-50 years</td>
</tr>
<tr>
<td>Disease Stage</td>
<td>H&amp;Y stage-I / Stage-II</td>
<td>H&amp;Y stage-I / Stage-II</td>
</tr>
</tbody>
</table>

* M-Male; F-Female;

3.2. Image Preprocessing

The original datasets were of size 512×512 in Digital Imaging and Communications in Medicine (DICOM) format, which are converted to 3D Neuroimaging InFormatics Technology Initiative (NIIfTI) format using the dcm2nii tool in MRICron software (http://www.mccauslandcenter.sc.edu/mricro/mricron/). Preprocessing involved two main steps: skull stripping using the Brain Extraction Tool (BET) (Smith et al., 2004; Jenkinson et al., 2012) and normalization to Montreal Neurological Institute (MNI) space through co-registration with the MNI template (MNI152 T1MPRAGE-1mm brain) in FSL 4.1 software (Grabner et al., 2006). The preprocessed images were subsequently brought into MATLAB through the utilization of the 'Tools for NIIfTI and ANALYZE Image' toolbox (http://www.mathworks.com/matlabcentral/fileexchange/8797). The final whole-brain volumes were limited to
224×224×178 voxels, which ensures that all the brain images used in the study have the same spatial dimensions as needed for this study, making it easier to analyze and compare them in subsequent data processing and analysis steps. As the obtained MRI datasets contain a large amount of noise, an enhanced Gaussian filter, (modified version of the traditional Gaussian filter) was employed to remove it. It is a high-pass filter which is added to the Gaussian kernel, that amplifies the high-frequency elements of the image while suppressing the low-frequency noise. This kernel is used to preserve the edges, fine details in the image. Reduction in noise and an increase in analytical sensitivity are the outcomes of the filter [45] and hence suits best for the medical image processing [46]. The effect of the preprocessing can be observed from Fig. 5. The pseudocode is displayed in Algorithm-1.

Algorithm-1: Enhanced Gaussian Filter

Input: Image of size 224 × 224 with noise; An empty matrix of size nxn to represent Gaussian Filter kernel GFₙₓₙ; the standard deviation (σ) for the Gaussian distribution; predefined threshold parameter (T);
Output: Enhanced Gaussian Filtered image of size 224 × 224
Begin
3.1 for each position in kernel:
3.1.1 x = row-n / 2
3.1.2 y = col - n / 2
3.1.3 kernel[position] = \( \frac{1}{2\pi\sigma^2} e^{-\frac{x^2 + y^2}{2\sigma^2}} \)
3.1.4 height, width = dimensions of the image (i.e., 224,224)
3.1.5 border = n // 2
3.1.6 for i = 1 to height:
3.1.7 for j = 1 to width:
3.1.8 window = empty matrix (n,n)
3.1.9 for each position in window:
3.1.9.1 window[position] = image[i + row offset - border, j + col offset - border]
3.1.9.2 sum = 0
3.1.9.3 for each value in window:
3.1.9.4 sum += value
3.1.9.5 local_mean = sum / total elements in (window)
3.1.9.6 deviation_sum = 0
3.1.9.7 for each value in window:
3.1.9.8 deviation = value - local_mean
3.1.9.9 deviation_sum += deviation^2
3.1.9.10 local_stddev = \( \sqrt{\text{deviation}_\text{sum} / \text{total elements in (window)}} \)
3.1.9.11 \( W_{ij} = 1 - \frac{\text{local_stddev}}{\text{threshold}} \)
3.1.9.12 enhanced_kernel = empty_matrix (n,n)
3.1.9.13 for each position in enhanced_kernel:
3.1.9.14 enhanced_kernel[position] = kernel[position] * weight
3.1.9.15 \( EGF_{ij} = GF_{ij} * W_{ij} \) ∀ 1 ≤ i ≤ n, ∀ 1 ≤ j ≤ n
3.1.9.16 sum = 0
3.1.9.17 for each value in enhanced_kernel:
3.1.9.18 sum += value
3.1.9.19 enhanced_kernel = \( \frac{\text{enhanced_kernel}}{\text{sum}}, \text{convolved_value} = 0; \)
3.1.9.20 for each value, weight in window, enhanced_kernel
3.1.9.21 convolved_value += value * weight
3.1.9.22 result[i,j] = convolved_value
3.1.9.23 filtered_image = result
End
End
3.3. GANs

GANs used to be considered complex and advanced a few years ago. But they have become a common and widely adopted tool in various fields, now-a-days. They create realistic images and have various applications in image manipulation techniques like image synthesis, augmentation [34], super resolution [30], image-to-image translation [31], image blending and [32], image inpainting. Deep Convolutional Generative Adversarial Networks (DCGAN) are an extension of GANs, that are designed for producing high-quality, realistic images [27]. The distinguishing feature of DCGANs is the incorporation of convolutional layers in both the generator and discriminator networks. This architectural choice enables the networks to effectively capture spatial patterns and dependencies within the data. As a result, DCGANs consistently demonstrate proficiency in generating detailed and visually coherent images across diverse domains. DCGANs make use of two adversarial networks (G(z) and D(x)), of which generator generates a photorealistic image G(z) with the intention of fooling the discriminator, while the discriminator sends back the images that resemble unrealistic (D(x)) back to the generator network i.e., the generator's job is to find the lowest possible value for the cost function V (D, G), while the discriminator's job is to find the highest possible value [33, 34]. As there are very limited labelled images available, the process of analyzing medical images requires the use of an important technique called image augmentation [34]. The DCGAN-based data augmentation [35] was utilized to generate fresh images. A generator network with an architecture that is comprised of seven layers has been chosen, and this design is depicted in Fig. 6. The data that is supplied to the generator network is a vector that has been selected at random from a set of hundreds of invariant values that are arrayed between 0 and 1. The well-connected layer, which consists of 128 × 128 × 64 = 1048576 values, is immediately supplied by the input data and is consequently modified to achieve 64 × 64 × 128 measure, by passing through the convolution layer. A Leaky ReLU activation function is appended to each of the layers, along with batch normalization application. The process is repeated for the consecutive six convolution layers of different sizes of inputs and outputs, to obtain a final block of size 1 × 1 × 512. These 512 units are fed as input to the deconvolution layer, to produce a block of 2 × 2 × 1024. A ReLU activation function is applied to every layer, along with the application of dropout and batch normalization. The process is repeated for the successive two Deconvolution layers of different sizes of inputs and outputs, to obtain an output block of size 8 × 8 × 1024. It is passed through a deconvolution layer, equipped with ReLU activation function and batch normalization. The process is repeated for the consecutive three convolution layers of different sizes of inputs and outputs, to obtain a final block of size 128 × 128 × 128. This block is passed through a deconvolution layer supported with ReLU activation function and Tanh functions to obtain a synthetic image of size 224 × 224 × 3. In addition to the real image samples, fake samples obtained from the generator are fed as input to the first convolution layer of discriminator network, which is of size 128 × 128 × 64. Leaky ReLU function is applied here and this processed block is fed as input to the convolution layer, where Leaky ReLU along with batch normalization is applied. This process is repeated for two times with different sizes of blocks to obtain a block of size 31 × 31 × 512. This block of data is passed through a convolution layer along with sigmoid activation function, thus resulting in generating a block of size 30 × 30 × 1. Sigmoid activation function is applied on this block finally, to produce the final output as ‘0’ or ‘1’ to represent a fake image or real one respectively. If the resultant image resembles fake, it is fed back to the generator, for proper training. Otherwise, added to training set. The iterative process of the DCGAN training is outlined in Algorithm-2 (Appendix). Random numbers are used to determine the values of the weight and bias parameters. All weights were initialized from a zero-centered Normal distribution with standard deviation 0.02. In the Leaky ReLU, the slope of the leak was set to 0.2 in all models. Adam optimizer was used with a suggested learning rate of 0.001. Additionally, when β1 was assigned with a value of 0.9, it resulted in training oscillation and instability while reducing it to 0.6 helped stabilize training.

Assuming total number of batches as $\text{batch count}$, batch size as $\text{batch size}$, total number of epochs as $\text{epoch count}$, total number of images as $\text{img count}$, the total number of images generated during training ($\text{img gen count}$) can be calculated as follows:

First, calculate the number of batches per epoch. Since we have 70 input samples for training, and a batch size of 10, the number of batches per epoch would be:

$$\text{batch count} = \frac{\text{img count}}{\text{batch size}} = \frac{70}{10} = 7$$  \hspace{1cm} (1)

Now, calculate the total number of images generated during training with 160 epochs as follows:

$$\text{image gen count} = \text{batch size} \times (\text{batch count}) \times (\text{epoch count})$$  \hspace{1cm} (2)

$$\text{image gen count} = 10 \times 7 \times 160 = 11,200 \text{ images}$$  \hspace{1cm} (3)
Therefore, a total of 11,200 images (4000 HC images and 7200 PD images) are generated during data augmentation over 160 epochs. These images are saved to a local storage location on the computer and are then organized into folders based on their class labels, to maintain a clear structure. The images are finally fed as input to the DCNN models. The GAN network goes through ‘n’ number of rounds, and the loss matrix is calculated at the end of each iteration. This loss function is used to measure the performance of both the generator and the discriminator. It is a combination of two terms: the generator loss and the discriminator loss.

Generator Loss: It measures how well the generator is able to produce fake samples, similar to the real.

Discriminator Loss: It measures how well the discriminator is able to vary between real, fake samples.

During training, the generator and discriminator are updated alternately to minimize this loss function. Let ‘G’ be the generator function that takes in a random noise vector ‘z’ and outputs a generated sample ‘x’, and let ‘D’ be the discriminator function that takes in a sample ‘x’ and outputs a probability score ‘p’ indicating the probability that ‘x’ is a real sample. The mathematical equation for the overall loss calculation (L) in a GAN, can be represented as

$$L = L_G + L_D$$

Where ‘LG’ is the generator loss, which is calculated using the cross-entropy loss between the discriminator's output on the generated samples and a target vector of all 1’s as

$$L_G = -\sum_{x} y \log(D(x)) + (1 - y) \log(1 - D(G(z)))$$

$$L_D = -\sum_{x} y \log(D(x)) - \sum_{z} (1 - y) \log(D(G(z)))$$

Fig. 6. DCGAN architecture.
and ‘LD’ is the discriminator loss, which is calculated as the sum of the cross-entropy loss between the discriminator's output on the real samples and a target vector of all 1’s, and the cross-entropy loss between the discriminator's output on generated samples and a target vector of all 0’s as:

$$L_D = - \log(D(G(z)))$$  \hspace{3cm} (5)

The weight and bias values are updated, accordingly. This process continues until both networks become stable.

### 3.4. Optimizers

**Stochastic Gradient Descent (SGD):** SGD is an optimization algorithm used to determine the model parameters that best match the predicted and the actual outputs. It calculates the gradient using just a random small part of the observations instead of all of them. The major advantage of SGD is that the update steps are performed very quickly that leads to reach the minimum in a very small amount of time.

**Adam:** Adaptive Moment Estimation (Adam) is an optimization technique for gradient descent. It takes a smaller amount of memory and is more efficient. In this study, the GAN learning rate is configured to be 0.001, the number of iterations to be 160, and batch size as 10. To reach equilibrium, it is necessary for both the Generator and the Discriminator networks to engage in concurrent or the adversarial learning. The variation in the loss function can be observed from Fig. 7, 8. Adam was proved to be better, compared to SGD, for image classification tasks [43, 46, 47].

The loss function transition of the Generator Network gets reduced over time while it gets increased, with adversarial network. This suggests that the modelling level and bias against their most recent data has variable change levels, which is beneficial for the model. During the first round of iteration, the produced images exhibited a high degree of inconsistency. It started to decrease after 150 iterations, and became stable for good, after a total of 160 iterations. When it comes to the replicating periods depicted in Fig. 8, it is easy to demonstrate that the value shift of the adversarial network's loss function starts to climb from roughly 0.65 onwards and continues, until it reaches a maximum. Fig. 9 (x1, y1) demonstrates that the images have become distorted and faulty after 80th iteration, while Fig. 9 (x2, y2) demonstrates that contours and textures have started to take shape after the 120th iteration. When looking at Fig. 9 (x3, y3), it is easy to see that after 150 iterations, the individualized characteristics have been improved and have become, to some extent, more improved and have become, to some extent, more comparable to the actual images. This is evidenced by the fact that they are now more like the genuine images. Succeeding 160 iterations, the suggested strategy is said to be balanced. PSNR is a measurement instrument that is used to determine image quality and its equation can be represented as

$$\text{Peak Signal} - \text{to} - \text{Noise Ratio} = \frac{10 \log ((\text{Count} - 1)^2)}{\text{MSE}}$$  \hspace{3cm} (7)

Where ‘Count’ represents the number of maximum intensity levels and MSE represents Mean Squared Error. If the PSNR number is higher, then the image will have less blurring and other distortions. PSNR>=y, where y=30, makes image, superior to the standard. The objective was not accomplished until more than 150 cycles were carried out and can be seen from Table 4. It can be observed that as the number of iterations increases, the overall image quality that is produced by the study also improves. A high value of PSNR and stability in the images is observed, as the total number of repetitions move towards 160. It has been found that the greatest potential outcomes may be accomplished when the convolution kernel for feature extraction is set to 3 x 3, and the iteration number for DCNN is set to 160.

### 3.5. DCNN

DCNN is a type of deep learning algorithm commonly used for image recognition and processing tasks. The key feature is their ability to automatically learn hierarchical representations of features in images. DCNNs have revolutionized the field of image classification by providing state-of-the-art performance on a wide range of datasets. The role of DCNNs in PD detection is to automatically learn features that may include changes in brain morphology or activity that are associated with the disease. Once trained, the DCNN can be used to classify new medical images as either HC or PD. Different characteristics related to various DCNN models that support image classification tasks are presented in Table 5. It can be observed that Alexnet, GoogLeNet and VGGNet were not considered in this study because of the memory requirements, architecture complexity and efficiency. EfficientNet B1, ResNet-50, ResNet152V2, and MobileNetV2 are all popular and widely used DCNN architectures in image classification tasks. They are chosen for the proposed study because of their image size, number of trainable parameters, and associated merits, demerits.
Fig. 7. Loss function of Generator network.

Fig. 8. Loss function of Discriminator network.

Fig. 9. (x, y): Real images of PD, HC; (x1, y1), (x2, y2), (x3, y3): Synthetic images of PD, HC after 80, 120, 160 iterations respectively.

Table 4. Iterations Vs PSNR Values.

<table>
<thead>
<tr>
<th>Epochs</th>
<th>1</th>
<th>40</th>
<th>80</th>
<th>120</th>
<th>160</th>
</tr>
</thead>
<tbody>
<tr>
<td>PSNR value (dB)</td>
<td>0</td>
<td>19.61</td>
<td>24.75</td>
<td>32.86</td>
<td>35.18</td>
</tr>
</tbody>
</table>
3.6. Transfer Learning

Instead of copying the complete network, including all of the fully connected layers, it is more common practice to merely copy the weights that are included within the convolutional layers. The pre-trained DCNN models (EfficientNetB1, ResNet-50, ResNet152V2 and MobileNetV2) which were trained on ImageNet dataset with 1000 classes were chosen for the classification of brain MRI with two distinct classes, HC and PD, using transfer learning mechanism. Additionally, DCNN would require a significant quantity of MRI to generate upgraded weights and produce new ones. As a result, shifting weights from the pre-trained model into the DCNN will cause it to produce output that is desirable and will speed up the rate at which it converges [20]. The number of output neurons in pre-trained version using ImageNet data set is 1000, but it is only ‘2’ to represent HC and PD classes as per the stated problem. Fig. 10 shows the transfer-learned DCNN architecture.

<table>
<thead>
<tr>
<th>DCNN Model</th>
<th>Image Size</th>
<th>#parameters (millions)</th>
<th>Merits</th>
<th>Demerits</th>
</tr>
</thead>
<tbody>
<tr>
<td>AlexNet</td>
<td>224x224</td>
<td>62.3</td>
<td>Introduced ReLU activation and dropout.</td>
<td>High computational, memory requirements; Prone to overfitting on small datasets.</td>
</tr>
<tr>
<td>VGGNet</td>
<td>224x224</td>
<td>138</td>
<td>Uniform architecture, Strong performance on binary classification problems, Introduced Inception module for feature combination.</td>
<td>Extremely deep with a high number of parameters and not the most parameter efficient architecture.</td>
</tr>
<tr>
<td>GoogLeNet</td>
<td>224x224</td>
<td>5</td>
<td>High accuracy with computational efficiency</td>
<td>Complex architecture; challenging to train.</td>
</tr>
<tr>
<td>ResNet-50</td>
<td>224x224</td>
<td>23.5</td>
<td>Residual connections mitigate vanishing gradient problem, Effective for binary tasks, due to skip connections.</td>
<td>Deeper variants can be computationally expensive; It may require substantial data for training deep variants.</td>
</tr>
<tr>
<td>ResNet152V2</td>
<td>224x224</td>
<td>60.4</td>
<td></td>
<td></td>
</tr>
<tr>
<td>EfficientNet B1</td>
<td>224x224</td>
<td>6</td>
<td>Achieves high accuracy with few parameters</td>
<td>Low performance on very small datasets.</td>
</tr>
<tr>
<td>MobileNetV2</td>
<td>224x224</td>
<td>4.2</td>
<td>Lite weight, highly computational efficient and low memory footprint.</td>
<td>Limited model capacity compared to larger architectures.</td>
</tr>
</tbody>
</table>

3.7. Ensemble Learning

Ensemble techniques are powerful methodologies that leverage a diverse range of models and combine their predictions to create a robust and reliable final model. Instead of relying solely on a single model and hoping for the best accuracy, ensemble learning takes advantage of combining multiple competing models to improve deep learning outcomes. This approach proves to be more effective in predicting future outcomes compared to using a solitary model alone [40, 58, 59]. The proposed ensemble model can be observed from Fig. 3. The dataset under study is experimented with three different ensemble mechanisms of bagging technique [59], that combine the strengths of multiple classifiers, compensating for their individual weaknesses and biases. They are (a) average-voting and (b) majority-voting (c)
OWPM method. It has been shown to improve the prediction accuracy, robustness, and enhance the generalization ability of the ensemble model, by taking into account the outcomes of the various classifiers and make the final choice.

3.7.1 Average Voting method

In the average voting scheme, average function is applied over the predicted outcomes \([\mathbf{O}_1, \mathbf{O}_2, \mathbf{O}_3, \ldots, \mathbf{O}_n]\), generated from different classifiers (\(\text{DCNN}_1, \text{DCNN}_2, \ldots, \text{DCNN}_n\)) of all the DCNN models, where ‘n’ is the total number of models. Let these outcomes belong to multiple classes \([\mathbf{C}_1, \mathbf{C}_2, \mathbf{C}_3, \ldots, \mathbf{C}_j]\), where ‘j’ is the total number of classes. Let ‘\(\text{Res}\)’ be the predictive output of a sample ‘x’, to be generated using ensemble mechanism. Then, the average value is calculated as follows:

\[
\text{average} = \frac{\text{sum of outcomes of classifiers}}{\text{total number of classifiers}}
\]

\[
\text{average} = \frac{\sum_{i=1}^{n} O_i(x)}{n}
\]

In this study, there are only two classes that represent PD (=1) and HC (=0). 4 DCNN models were employed in the proposed study, that might generate the outcomes as either ‘0’ or ‘1’.

\[
\text{Res} = \begin{cases} 
1, & \text{If average} > 0.5, \text{indicating PD.} \\
0, & \text{If average} \leq 0.5, \text{indicating HC.}
\end{cases}
\]

3.7.2 Majority Voting method

In this scheme, the outcomes \([\mathbf{O}_1, \mathbf{O}_2, \mathbf{O}_3, \ldots, \mathbf{O}_n]\), belonging to multiple classes \([\mathbf{C}_1, \mathbf{C}_2, \mathbf{C}_3, \ldots, \mathbf{C}_j]\), generated from different classifiers (\(\text{DCNN}_1, \text{DCNN}_2, \ldots, \text{DCNN}_n\)) are counted and the class with the highest count is considered as the resultant prediction, for the subject being analyzed using ensemble method. In this study, there are only two classes that represent PD (=1) and HC (=0). Let ‘\(T_r\)’ and ‘\(T_s\)’ represent the total number of outcomes from different classifiers resulting in ‘1’ and ‘0’ respectively. Let ‘\(\text{Res}\)’ be the predictive output to be generated using ensemble mechanism and it can be calculated as:

\[
\text{Res} = \begin{cases} 
1, & \text{If} \ (T_p \geq T_N), \text{indicating PD} \\
0, & \text{If} \ (T_p < T_N), \text{indicating HC.}
\end{cases}
\]

3.7.3 OWPM method

OWPM method in an ensemble mechanism, which is an extension of weighted majority voting scheme, that introduces “priority” as an additional layer of information about the relative importance of classifiers. The proposed method uses classifier fusion ensemble mechanism [57] which involves a slightly different mechanism for combining classifier outputs. In this scheme, let (\(\text{DCNN}_1, \text{DCNN}_2, \ldots, \text{DCNN}_n\)) represent various classifiers generating different outcomes \([\mathbf{O}_1, \mathbf{O}_2, \mathbf{O}_3, \ldots, \mathbf{O}_n]\) belonging to multiple classes \([\mathbf{C}_1, \mathbf{C}_2, \mathbf{C}_3, \ldots, \mathbf{C}_n]\). These classifiers are assigned priorities \((P_1, P_2, \ldots, P_n)\) and Weights \((W_1, W_2, \ldots, W_n)\). The classification approach is as follows:

- Arrange the models in descending order of efficiency; The highest performance model is assigned with the highest priority \((P=n)\) and largest weight \((W=n)\); The model with the next highest performance is assigned with the next highest priority \((P=n-1)\) and the next largest weight \((W=n-1)\); This process is continued until the lowest performance model is assigned with the lowest priority \((P=1)\) and smallest weight \((W=1)\).
- Let ‘\(\text{Res}\)’ be the predictive output of a sample ‘x’, to be generated using ensemble mechanism.
- Calculate the Weighted Priority Mean as the ratio of sum of products of weights and priorities of models w.r.t sample ‘x’ along with their outcomes to the sum of their products of weights and priorities. i.e.,

\[
\text{Weighted Priority Mean} = \frac{\sum_{i=1}^{n} W_i P_i(x) \cdot O_i(x)}{\sum_{i=1}^{n} W_i P_i(x)}
\]

Where ‘i’ represents the classifier, and ‘n’ represents the total number of classifiers. In this study, there are only two classes that represent PD (=1) and HC (=0). 4 DCNN models were employed in the proposed study, that might generate the outcomes as either ‘0’ or ‘1’.
The pseudocode is presented in Algorithm-3. The final classification of a subject in a case where equal number of votes results for both the classes may need more data for the model to train on, or, an external intervention by an expert medical practitioner. The subject was assigned to either of the classes ‘HC’ or ‘PD’, in this study.

Algorithm-3: The OWPM Algorithm

1. Input: The trained and analyzed DCNN models \( \text{DCNN}_1, \text{DCNN}_2, \ldots, \text{DCNN}_n \), utilized in this study; the models efficiencies \( E_1, E_2, E_3, \ldots, E_n \); the models outcomes \( O_1, O_2, O_3, \ldots, O_n \); total number of models \( n \); the prediction for the subject with unknown label ‘Res’.
2. Output: Prediction for the subject with unknown label as PD (or) HC, using the OWPM algorithm.
3. Begin
   3.1 Arrange the models \( \text{DCNN}_1, \text{DCNN}_2, \ldots, \text{DCNN}_n \) in descending order of their efficiency and assign priorities \( P_1, P_2, \ldots, P_n \) to these models and Weights \( W_1, W_2, \ldots, W_n \) in such a way that the highest performance model \( \text{DCNN}_n \), will be assigned the highest priority \( P_1 = n \) and the highest weight \( W_1 = n \), the model with the next highest performance \( \text{DCNN}_{n-1} \), will be assigned priority \( P_2 = n-1 \) and highest weight of \( W_2 = n-1 \) and so on. This process is repeated until the model with the least performance will be assigned the lowest priority \( T \) and lowest weight \( T \).
   3.2 As a subsequent step, calculate the weighted priority mean as the ratio of sum of the product of outcomes \( O_1, O_2, O_3, \ldots, O_n \) and their associated weights \( W_1, W_2, \ldots, W_n \) and priorities \( P_1, P_2, \ldots, P_n \) to the sum of the products of weights and priorities, as follows:
   
   \[
   \text{Weighted Priority Mean} = \frac{\sum_{i=1}^{n} W_i P_i \cdot O_i}{\sum_{i=1}^{n} W_i P_i}
   \]

4. If the value obtained in the previous step is greater than 0.5, Res=1 (PD); otherwise, Res=0 (HC);
5. End

Advantages of OWPM model over Weighted Majority Voting Schema:

- Weights focus on the contribution of a classifier's prediction to the final decision, while priorities focus on the relative ranking of efficiency among classifiers. Weights alone might not be sufficient, if there is a need to explicitly consider the order of efficiency among classifiers. Priorities alone might not capture the reliability of classifiers. The combination of weights and priorities allows the adaptability to problem-specific characteristics, providing a more versatile and fine-grained ensemble approach.
- Weights control the influence of each classifier based on its individual performance, while priorities introduce a global ranking that affects the overall contribution of each classifier. The use of both priorities and weights provides a comprehensive approach to ensemble learning, enabling a system to consider both the accuracy and order of efficiency among classifiers.

3.8. Training And Fitting

It is vital to employ a pre-trained model, also known as transfer learning, to increase the performance of image classification. This is because the pre-trained model's weights had previously been designed with characteristics that were pertinent to the large majority of computer vision problems. To facilitate transfer learning, a data set consisting of training, verification, and test information was constructed. Because the dataset that was used had images that were comparable to those that were used in ImageNet, it is conceivable to use the transfer-learning technique in this investigation. It would have been essential to retrain some of the layers in the model in the event that the images are not comparable to those that were used to train the models that are now available. Fine tuning can be accomplished by making appropriate modifications to the topmost layer that is already fully connected. All the tasks are carried out on a laptop equipped with an Intel® Core (TM) i5-1135G7 microprocessor and 8GB of Random-Access Memory (RAM).
3.9. Output Layer Training

To enhance the model's performance, the original output layer was removed as the first step, followed by introducing a unique layer with two units in its place (PD and HC, as per the application) and a sigmoid activation function. During training, only the weights of this newly added output layer are adjusted, while the rest of the network remains frozen. This approach allows us to focus on fine-tuning specific aspects without disrupting the existing architecture. For optimization, Adam, a renowned optimizer in deep learning [38], was utilized. Selecting an appropriate initial learning rate for the DCNN model during fine-tuning can be challenging, as the learning rate may decrease throughout the process. To address this challenge, the Grid search optimization approach was employed, which proves substantially more efficient than random search. It enables us to methodically explore parameter combinations and select the most suitable configuration, ultimately improving the model's effectiveness [20]. Each classification problem is subjected to

3.10. Hyperparameter Tuning

Grid search method was employed for performing hyperparameter tuning. In this method, all of the potential combinations of the hyperparameters and their values are first laid out on a grid. Next, an algorithm is built for each combination of the grid's components. After making a note of the respective parameters' degrees of accuracy, an analysis of the validation data using those parameters was done. It took five steps to obtain the desired level of hyperparameter tuning. In the first stage of the process, the goal was to randomly assess how well each hyperparameter performed with reference to accuracy and loss, during both the training and validation phases. Next, 8 different experiments were conducted, out of which the best two models with reference to accuracy and loss function in the validation process are selected, for the classification of the test dataset. The initial weights assigned during training will be adjusted according to accuracy and loss function in the validation process are selected. The initial weights assigned during training will be adjusted accordingly, during the process. Next stage was to investigate this disparity, and the two studies that performed the best overall in terms of accuracy underwent two further training cycles. The batch size for the best sample, which was obtained in the fourth stage, varied between 32, 64, 128, and 256 in the following stage. When either the algorithm reached its maximum epochs or training phase's validation output started to decline over a set amount of time, the training procedure was stopped. Overfitting was avoided, as two different data sets were used for validation and testing, in this study. Additionally, Dropout method was employed during the training phase. The top level's goal is to lessen the average loss, while the bottom level's goal is to maximize the accuracy. Regular adjustments are made to hyper-parameters and the optimal ones are determined, based on the validation set. They can be observed from Table 6. The step-by-step classification procedure is outlined in Algorithm-4.

Algorithm-4: PD classification using pre-trained DCNN

Input: Pretrained DCNN model (L_k), Augmented training MRI (I_{train} = I_k + I_{g}), Validation Images \( I_{val}\), Validation data \( I_{val}\), validation accuracy \( \beta\), iterations per stage < Z = Z_1, Z_2, Z_3, ..., Z_2 >, Total no. of stages X, Training data per stage \( I_{train} = I_{train}^1, I_{train}^2, ..., I_{train}^Z >\), hyper parameters \( h_{1,k}\), Real MRI (I_k), Synthetic MRI (I_g)

Output: hyper parameters \( h^*\), Predicted labels (0-for HC, 1-for PD patients)

Begin

learning_rate = initial_learning_rate // Set the initial learning rate for Adam
beta1 = 0.9 // Adam parameter
beta2 = 0.999 // Adam parameter
epsilon = \frac{1}{10^6} // Adam parameter

for \gamma iterations

for each \( i \in I_{train}\)

\text{DCNN} \leftarrow L_{\alpha}(i) // Initialization of any one of the DCNN models (EfficientNetB1, ResNet-50, ResNet152V2, MobileNetV2)

\text{DCNN.fc.pop()} // Removal of last fully connected layer

\text{DCNN.dense(2, activation function)}

for stage = 1 to X do

for \( j = 1\) to k do

\( \beta_j \) = evaluate \( \beta(h_k, I_{train}^j, I_{val})\)

end

for \( i = 1\) to Z

\( g = \text{grid-search}(h, \beta_j)_{max} ; h_{max} = \text{args}_{max} a(h,g) ; \beta_j = \text{evaluate}(h, I_{train}^j, I_{val})\)

end for

end for

End
During the testing procedure. It can be finally concluded that over the proposed DCNN models performance along with the computational complexity can also be observed. The impact of data preprocessing can be depicted from the Fig. 12(a), 12(b), 12(c) and 12(d) respectively. Table 7 illustrates the ensemble model’s performance as well as transfer learned DCNN models performance. The impact of data preprocessing and augmentation over the proposed DCNN models performance along with the computational complexity can also be observed.

### 4. RESULTS

Appendices. As a first step, the classifier is trained on the pre-trained networks. Next, transfer learning was used to improve the weights further, starting with backpropagation based on the most recent training datasets. The final model was applied for a total of 100 epochs, following the selection of the pertinent hyper-parameters. Both SGD and Adam were studied as optimization measures and the training was carried out for a total of 100 epochs. The parameters that gave the best performance of the model are a learning rate of 0.001, weight decay of 0.05, momentum factor of 0.8 and batch size of 64, with Adam being the best optimizer. After running through all of the earlier programmes, the model is saved with learned weights and is put through its paces during the testing procedure. It can be finally concluded that Adam is the best optimizer, with a learning rate of 0.001, batch size of 64 and total epochs of 100, to classify HC and PD patients. The confusion matrices can be observed from Fig. 11. Loss and Accuracy corresponding to SGD and Adam can be depicted from the Fig. 12(a), 12(b), 12(c) and 12(d) respectively. Table 7 illustrates the ensemble model’s performance as well as transfer learned DCNN models performance. The impact of data preprocessing and augmentation over the proposed DCNN models performance along with the computational complexity can also be observed.
Fig. 11. Confusion matrices of Ensemble, DCNN models.

Fig. 12(a). Training / Validation loss using SGD

Fig. 12(b). Training / Validation loss using Adam
Matthews Correlation Coefficient (MCC) is a measure used to assess the quality of binary classifications. It is given by the formula:

\[
\text{MCC} = \frac{TP \times TN - FP \times FN}{\sqrt{(TP + FP)(FP + FN)(FN + TP)(TN + FP)}}
\]  

(11)

Where TP is the number of True Positives, TN is the number of True Negatives, FP is the number of False Positives and FN is the number of False Negatives. The MCC value ranges from -1 to +1, where ‘+1’ indicates a perfect prediction, ‘0’ indicates no better than random prediction, and ‘-1’ indicates total disagreement between prediction.

FPR stands for ‘False Positive Rate’. It is a metric that represents the proportion of actual negative instances that are incorrectly predicted as positive by the model. It indicates how well the model identifies negative instances. A lower FPR indicates better performance in terms of minimizing false positives. It is calculated using the following formula:

\[
\text{FPR} = \frac{FP}{FP + TN}
\]  

(12)

Where TN is the number of True Negatives and FP is the number of False Positives.

FNR stands for ‘False Negative Rate’. It is a metric that represents the proportion of actual positive instances that are incorrectly predicted as negative by the model. It indicates how well the model identifies positive instances. A lower FNR indicates better performance in terms of minimizing false negatives. It is calculated using the following formula:
Where $FN$ is the number of False Negatives and $TP$ is the number of True Positives.

**Sensitivity** is the proportion of actual positive instances that the model correctly predicts as positive. It quantifies the model's ability to "sensitively" detect or capture instances of the positive class. The values range from 0 to 1, where 1 indicates perfect sensitivity (all positive instances are correctly identified) and 0 indicates no sensitivity (no positive instances are correctly identified). It is calculated using the following formula:

$$\text{Sensitivity} = \frac{TP}{TP+FN}$$  \hspace{1cm} (14)

Where $FN$ is the number of False Negatives and $TP$ is the number of True Positives.

**Specificity** is the proportion of actual negative instances that the model correctly predicts as negative. It quantifies the model's ability to be "specific" in identifying instances of the negative class. The values range from 0 to 1, where 1 indicates perfect specificity (all negative instances are correctly identified as negative) and 0 indicates no specificity (no negative instances are correctly identified). It is calculated using the following formula:

$$\text{Specificity} = \frac{TN}{TN+FP}$$  \hspace{1cm} (15)

Where $FP$ is the number of False Positives and $TN$ is the number of True Negatives.

**Balanced accuracy** is a metric used to evaluate the performance of a classification model, especially when dealing with imbalanced datasets, where accuracy alone may be misleading. Balanced accuracy takes into account both sensitivity (true positive rate) and specificity (true negative rate) to provide a more comprehensive assessment. It ranges from 0 to 1, where 1 indicates perfect classification, and 0.5 indicates no better than random classification. The formula for balanced accuracy is:

$$\text{Balanced Accuracy} = \frac{\text{Sensitivity} + \text{Specificity}}{2}$$ \hspace{1cm} (16)

**Accuracy** is the proportion of all instances that the model correctly predicts. i.e., It measures the overall correctness of a model by considering both true positives and true negatives. The values range from 0 to 1, where 1 indicates perfect accuracy (all predictions are correct), and 0 indicates no accuracy (all predictions are incorrect). It is calculated using the formula:

$$\text{Accuracy} = \frac{TP + TN}{\text{total predictions}}$$ \hspace{1cm} (17)

Where $TN$ is the number of True Negatives and $TP$ is the number of True Positives.

**Precision** measures the accuracy of the positive predictions made by the model. A high precision indicates that the model is making positive predictions with a low rate of false positives. It is calculated as:

$$\text{Precision} = \frac{TP}{TP + FP}$$ \hspace{1cm} (18)

Where $PN$ is the number of True Positives and $FP$ is the number of False Positives.

**Recall** measures the ability of the model to capture all the positive instances. A high recall indicates that the model is effectively identifying a large proportion of the positive instances. It is calculated as:

$$\text{Recall} = \frac{TP}{TP + FN}$$ \hspace{1cm} (19)

Where $FN$ is the number of False Negatives and $TP$ is the number of True Positives.
**F-score** is a metric that combines precision and recall into a single value, by taking their harmonic mean. The formula for the F1 score is:

$$F - score = \frac{2(\text{precision} \cdot \text{recall})}{\text{precision} + \text{recall}}$$

(20)

To determine the possibility that an image will be classified as either HC or PD, each image was examined while incorporating the final ensemble model into account. The impact of data preprocessing and augmentation on the performance of the proposed models can be observed from Fig. 13. When assessing the network's effectiveness, the Receiver Operating Characteristic (ROC) is used. Fig. 14 shows that the AUC calculated from the ROC curve has an estimated value of 0.9861, 0.9524, 0.7986, 0.7361 and 0.7083 for Ensemble Method, EfficientNet B1, ResNet-50, ResNet152V2 and MobileNetV2 respectively. This shows that Ensemble method results in a satisfying result of the estimated value of 0.9861, 0.9524, 0.7986, 0.7361 and 0.7083 for performance of the proposed models incorporating the final ensemble model into account. The impact of data preprocessing and augmentation on the performance of the proposed models can be observed from Fig. 13. When assessing the network's effectiveness, the Receiver Operating Characteristic (ROC) is used. Fig. 14 shows that the AUC calculated from the ROC curve has an estimated value of 0.9861, 0.9524, 0.7986, 0.7361 and 0.7083 for Ensemble Method, EfficientNet B1, ResNet-50, ResNet152V2 and MobileNetV2 respectively. This shows that Ensemble method results in a satisfying result of the trained model. The training and validation accuracies as well as losses, can be depicted from Fig. 15 and Fig. 16 respectively. Table 8 illustrates the proposed model’s performance in comparison with state-of-the-art approaches. Clearly, the ensemble model outperforms all other published research works, confirming that the innovative deep-learning approach that was developed, has its advantages and is competitive. The purpose of this work is to suggest a system that uses ensemble learning using deep learning models to build imaging biomarkers that can predict early PD.

<table>
<thead>
<tr>
<th>Performance Measure</th>
<th>Ensemble Method</th>
<th>TL-EfficientNet B1</th>
<th>CC (minutes)</th>
<th>TL-ResNet-50</th>
<th>CC (minutes)</th>
<th>TL-ResNet152V2</th>
<th>CC (minutes)</th>
<th>TL-MobileNetV2</th>
<th>CC (minutes)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Accuracy (%)</td>
<td>83.33</td>
<td>80</td>
<td>76.77</td>
<td>80.00</td>
<td>86.67</td>
<td>70.00</td>
<td>66.67</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Specificity (%)</td>
<td>85.71</td>
<td>84.62</td>
<td>78.57</td>
<td>80.00</td>
<td>78.57</td>
<td>71.43</td>
<td>66.67</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sensitivity (%)</td>
<td>81.25</td>
<td>76.47</td>
<td>75.00</td>
<td>80.00</td>
<td>76.47</td>
<td>68.75</td>
<td>66.67</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Table 7. Preprocessing and augmentation effect on DCNN and Ensemble models Performance

*CC: Computational Complexity; TL: Transfer Learned*
Evaluating the efficiency of a deep learning model involves not only performance metrics but also the interpretability of its predictions, particularly in critical domains such as healthcare. The ability to explain the rationale behind specific predictions is predominant, in the context of implementing intelligent systems. For the interpretation of predictions generated by the developed DCNN model, A Gradient weighted Class Activation Mapping (Grad-CAM) was employed [60, 61]. It is an extended version of CAM that calculates the gradient directly using the backpropagation from each neuron in the last convolution layer feature map.

Fig. 13. Impact of data preprocessing and augmentation.
CAM shows the score of importance of each part of the input by resizing 2D map that predicts class ‘C’ to the original input image. Then, it adds up these gradients within the ith feature map, to generate the weight of each map and predict class ‘C’. The Grad-CAM of class ‘C’ can be generated using the eq (24).

\[ L_{\text{Grad-CAM}} = \text{ReLU} \left( \sum_i w_t^i \text{GAP}^i \right) \]  

(21)

Where \( \text{GAP}^i \) is the Global Average Pooling of ith feature map and ‘\( w_t^i \)’ is the weight of each map.

Table 8. state-of-the-art works comparison

<table>
<thead>
<tr>
<th>Reference</th>
<th>Methodology</th>
<th>Accuracy (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>[10]</td>
<td>SVM classifier</td>
<td>85.8</td>
</tr>
<tr>
<td>Proposed</td>
<td>TL MobileNetV2 + data augmentation</td>
<td>86</td>
</tr>
<tr>
<td>Proposed</td>
<td>TL ResNet152V2 + data augmentation</td>
<td>89</td>
</tr>
<tr>
<td>[20]</td>
<td>TL AlexNet + data augmentation</td>
<td>89.23</td>
</tr>
<tr>
<td>Proposed</td>
<td>TL ResNet-50 + data augmentation</td>
<td>92.6</td>
</tr>
<tr>
<td>[18]</td>
<td>CNN + Bayesian SMBO optimization</td>
<td>95.29</td>
</tr>
<tr>
<td>Proposed</td>
<td>TL EfficientNet B1 + data augmentation</td>
<td>98</td>
</tr>
<tr>
<td>Proposed</td>
<td>Ensemble Learning</td>
<td>99</td>
</tr>
</tbody>
</table>

*TL-Transfer Learned;
The ReLU function is employed to eliminate negative contribution scores. This is because Grad-CAM focuses on considering only the input features that positively impact the prediction score for class ‘C’. Grad-CAM can be applied to interpret any type of DCNN without making changes to the DCNN model, as it directly uses gradients from backpropagation. The flow of action can be observed from Fig. 17 [22]. From Fig. 18(b), it can be observed that the CAM highlights the regions of interest in MRI slices predicted as Parkinson’s Disease. Notably, the model demonstrates an enhanced focus on the Substantia Nigra pars compacta (SNc), a region significantly impacted by the loss of dopaminergic neurons. This interpretative approach, utilizing the Grad-CAM technique, enhances our understanding of the model’s decision-making process.

Fig. 18 (a). MR Image predicted as PD Fig. 18(b). Grad-CAM Heatmap predicted as PD.

5. DISCUSSION

In Results section, we demonstrated the performances of 4 DCNN models for binary classification based on the images obtained from PPMI and LSSH, and also compared them against the efficiency obtained using Ensemble learning in Table 6. According to this study, the training period for the network was on an average of five hours long and consisted of 100 epochs, which results in high computational complexity. The model achieved an accuracy of 99% and a balanced accuracy of 98.99%, which proves its high performance. Moreover, the interpretability model using Grad-CAM has proven its trustworthiness, by concentrating on the Substantia Nigra region to identify whether the image belongs to PD or HC category. Table 6 shows that the developed ensemble method using OWPM approach yields the best performance over all other models. Because the model achieved a greater level of classification accuracy when it was applied to the test set, it is clear that the model did not suffer from overfitting when it was applied to the training set. It is important to point out that the incorporation of synthetic images results in a significant reduction of the standard deviation of the outcomes in the validation of accuracy, sensitivity, specificity, as well as the Area Under the Curve (AUC). When compared to the results of classification carried out using only the original images, the ensemble model designed from preprocessed and augmented data, attains a greater accuracy of up to 15.67%. The early findings using various existing methodologies provided in Table 1 indicate an accuracy of 40%–90%, while the method developed offers an accuracy of 99%. MCC is used to assess the model’s quality and having achieved 98.64% MCC, we can claim that our model stands as one of the best proposed models, in diagnosing early PD. Despite the high computational complexity, our model recommended for clinical use with its outstanding features.

6. CONCLUSION

PD ranks as the second most prevalent neurological disorder, next to Alzheimer's disease. In this study, the challenge of accurately classifying brain MRI as either PD or HC, is investigated by developing a comprehensive framework that combines data augmentation, transfer learning, ensemble learning and DCNN models. Leveraging the custom dataset created from the data obtained from LSSH, a comparative analysis was conducted against the clinical standard, which involves visual interpretation and quantification by experts besides facilitating real-time clinical comparisons in the PD diagnosis process. Remarkably, the proposed ensemble approach using OWPM method achieved exceptional accuracy of 99%, average recall of 0.986, average precision of 0.993, average specificity of 0.9931, f1-score of 0.9899, showcasing its potential as a reliable tool for discriminating PD from HCs, surpassing all the prominent architectures.
including EfficientNet B1, ResNet-50, ResNet152V2, and MobileNetV2. Impressively, the ensemble model yielded substantial improvement of 9.77%, compared to the state-of-the-art models. This remarkable advancement highlights the process of the approach in diagnosing the PD accurately, based on whole brain MRI. Furthermore, the ensemble model exhibited high differential fluency, as evidenced by an impressive AUC value of 0.9867 according to the ROC curve. This demonstrates the capability of the model to effectively distinguish between PD and HC cases. Moreover, the interpretation of the model over the MRI scans was also evaluated using Grad-CAM approach, and it was found that the model paid maximum attention to the substantia nigra region for predicting a particular MRI scan as PD. Nevertheless, this study has certain limitations that must be acknowledged. Specifically, patient characteristics such as different age groups, locations, dietary habits, and gender were not taken into consideration, which could impact the classification performance. Another limitation is being unable to identify the disease stage of a PD patient, as per the H&Y scale. A single clinical metric may not be sufficient to accurately track PD progression. Therefore, a combination of metrics or global statistical tests must be utilized (Huang et al., 2009). The training data set considered in this study suffers from class imbalance problem (25 HC and 45 PD). Efficient results in limited time may be possible, if data balance can be maintained. It is highly recommended to perform the research by considering specific subcortical structures for the detection of PD. Future research endeavors should aim to incorporate the discussed factors, in addition to the development of imaging biomarkers for PD and other related conditions. This study serves as a significant step forward, providing valuable insights for researchers and clinicians alike in the realm of clinical image evaluation. The objective identification of PD will become increasingly accessible and reliable for clinicians in the near future, with the ongoing advancements in deep learning techniques.

7 DECLARATIONS

7.1 Ethical approval
This manuscript reports studies which does not involve human participants, human data or human tissue and animals. The authors declare that the present research work has fulfilled all relevant ethical guidelines required by COPE.

7.2 Conflict of interest
The authors have no conflict of interests to declare that are relevant to the content of this article.

7.3 Authors contributions
Anitha Rani Palakayala contributed to conceptualization, methodology, software, data curation, formal analysis, writing—original draft. Kuppusamy P contributed to conceptualization, methodology, supervision, result analysis, writing—review and editing final draft and validation.

7.4 Funding
This research did not receive any specific grant from funding agencies in the public, commercial, or not-for-profit sectors.

7.5 Availability of data and materials
Data used in this study is available from PPMI public database and can be downloaded from the web site: www.ppmi-info.org/data.

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APPENDIX

Algorithm-2: Generative Adversarial Network using ADAM optimizer

1. Input: Real MRI of brain (`Real_Imgs`), Random noise value (Z, 0.0 <= Z <= 1.0).
2. Output: GAN
3. Begin
4. Model=setup(hyp_param)
5. Dis_model=build_net(Dis_Layers);
6. Gen_model=build_net(Gen_Layers);
7. Train_img=random(num(Real_Imgs))
8. for i=1 to model.tot_batches
9.   tot_img=read(Real_Imgs, train_img);
10.  //Generator and Discriminator construction
11.  if for i=1 to model.tot_batches
12.    train_img=random(num(Real_Imgs))
13.    tot_img=read(Real_Imgs, train_img);
3.3.5  Gen_Model ← train(tot_imgs, Z)
3.3.6  Gen_Model ← adamopt (Gen_Model, Dis_Model, imitation_Imgs)
3.3.7  Dis_Model ← train(tot_imgs, Z, imitation_Imgs);
3.3.8  Dis_Model ← adamopt (Gen_Model, Dis_Model, imitation_Imgs)
3.3.9  End for
3.3.10 End for
3.4  GAN ← {Gen_Model, Dis_Model}

4. End

# Algorithm adamopt
Repeat:
update t', t:=t+1

$g_t = \nabla \text{grad}(\theta_t)$ // Obtain the gradients/derivatives 'g' w.r.t. 't'

$m_t = \beta_1 \cdot m_{t-1} + (1-\beta_1) \cdot g_t$ // Update the first moment 'm_t', second moment 'v_t'

$v_t = \beta_2 \cdot v_{t-1} + (1-\beta_2) \cdot g_t^2$ // Compute the bias corrected $m_t$ and bias corrected $v_t$

$\theta_t = \theta_{t-1} - \alpha \cdot \frac{m_t}{\sqrt{v_t} + \epsilon}$ // Update the parameter 'θ_t'
until $\theta_t$ do not converge.

# Algorithm compute_generator_loss
Function compute_generator_loss(Dis_Model, Fake_Imgs)
Fake_predictions = Dis_Model.predict(Fake_Imgs) // Compute the discriminator's prediction for the fake images
Generator_loss = compute_loss(Fake_predictions, target_labels='real') // Calculate generator loss using the predicted labels
return Generator_loss
End

# Algorithm compute_discriminator_loss
Function compute_discriminator_loss(Dis_Model, Real_Imgs, Fake_Imgs)
Real_predictions = Dis_Model.predict(Real_Imgs) // Compute discriminator's predictions for real and fake images
Fake_predictions = Dis_Model.predict(Fake_Imgs)
Real_loss = compute_loss(Real_predictions, target_labels='real') // Calculate the discriminator loss using the predicted labels
Fake_loss = compute_loss(Fake_predictions, target_labels='fake')
Discriminator_loss = Real_loss + Fake_loss // Combine the losses to compute the overall discriminator loss
return Discriminator_loss
End

4. End of the algorithm