Pediatric pneumonia diagnosis using stacked ensemble learning on multi-model deep CNN architectures

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Abstract
Pediatric pneumonia has drawn immense awareness due to the high mortality rates over recent years. The acute respiratory infection caused by bacteria, viruses, or fungi infects the lung region and hinders oxygen transport, making breathing difficult due to inflamed or pus and fluid-filled alveoli. Being non-invasive and painless, chest X-rays are the most common modality for pediatric pneumonia diagnosis. However, the low radiation levels for diagnosis in children make accurate detection challenging. This challenge initiates the need for an unerring computer-aided diagnosis model. Our work proposes Contrast Limited Adaptive Histogram Equalization for image enhancement and a stacking classifier based on the fusion of deep learning-based features for pediatric pneumonia diagnosis. The extracted features from the global average pooling layers of the fine-tuned MobileNet, DenseNet121, DenseNet169, and DenseNet201 are concatenated for the final classification using a stacked ensemble classifier. The stacking classifier uses Support Vector Classifier, Nu-SVC, Logistic Regression, K-Nearest Neighbor, Random Forest Classifier, Gaussian Naïve Bayes, AdaBoost classifier, Bagging Classifier, and Extra-trees Classifier for the first stage, and Nu-SVC as the meta-classifier. The stacking classifier validated using Stratified K-Fold cross-validation achieves an accuracy of
98.62%, precision of 98.99%, recall of 99.53%, F1 score of 99.26%, and an AUC score of 93.17% on the publicly available pediatric pneumonia dataset. We expect this model to greatly help the real-time diagnosis of pediatric pneumonia.

**Keywords**  Pediatric pneumonia · Chest X-rays · Computer-aided diagnosis · Contrast limited adaptive histogram equalization · Deep learning · Transfer learning · Stacking classifier · Stratified K-fold

1 Introduction

Acute diseases have become a reason for concern over the past few years. The exponential rise in such ailments results in high mortality rates across various countries and long-term economic losses [25, 46]. Pneumonia is one such disease that causes respiratory infection; as a result, harming the normal functioning of the human body. Some of the known pathogen that causes pneumonia include viruses and bacteria. A notable agent for the pervasive transmission of viruses and bacteria is the degradation of air quality [35]. The human lungs constitute tiny bags or sacs called the alveoli. These air sacs are responsible for exchanging essential gases, namely oxygen and carbon-di-oxide. When an individual is affected by pneumonia, these sacs fill up with pus and fluid, decreasing the gas exchange between the blood and the lungs. Pneumonia causes difficulty in breathing and other complications like chest pain, cough, vomiting, diarrhoea, and fatigue.

Pneumonia generally affects children under five years, the geriatric population, and people with co-morbidities like diabetes, cardiovascular disorders, and auto-immune disorders [35]. In 2017, over 850,000 lives perished due to pneumonia-induced disorders. The mortality rate is specifically high in South Asian and Sub-Saharan countries [17]. Pediatric pneumonia has accounted for the high mortality in children in the past five years [32]. In 2011, nearly 1.2 million children under the age of five died due to pneumonia [1]. In 2016, more than 800,000 children died of pneumonia, most of whom were no more than two years old, and the death toll was more than the totality of malaria, AIDS, and measles [20]. In 2019, pneumonia killed 740,180 children under the age of five, accounting for 14% of all deaths of children under five. Pediatric pneumonia accounts for 19% of the overall mortality rate in children below five years of age.

Studies indicate that the other causes of pediatric pneumonia are malnutrition, air pollution, and lack of immunity. The ailments of pediatric pneumonia are easily preventable when diagnosed at a very early stage. Different approaches for diagnosing pediatric pneumonia include chest X-rays to find inflammation in the lungs, blood sample test for arterial blood gas analysis, and sputum test. Other diagnostic measures include pulse oximetry, Complete Blood Cell (CBC), chest CT scan, bronchoscopy, and thoracentesis. Even though highly advanced diagnostic measures exist, the trend in mortality does not seem to decrease due to high prices and inaccessibility. The factors mentioned are vital, especially in underdeveloped countries where people have limited access to such resources. Chest X-ray-based diagnosis, being comparatively inexpensive, was adopted as a standardized test for pediatric pneumonia.

The chest X-ray is the commonly used test for diagnosing various lung diseases as it is a painless and non-invasive method. It is fast, easy, and inexpensive when compared to other techniques. Though chest X-ray diagnosis is time and cost-effective, it heavily relies on the diagnostic conditions and the expertise of doctors and radiologists. Sometimes the chest X-
rays may appear different due to outdated equipment and tools in clinical practice. In other cases, getting a proper X-ray in an ideal posture might be difficult. Hence, these factors will affect the quality of the image captured and diagnosis. In addition, using lower radiation levels for chest X-rays in children makes pneumonia diagnosis a cumbersome task.

Detecting the cause of pneumonia in children is tedious due to the lack of rapid commercially available accurate lab tests for most of the existing pathogens which cause pediatric pneumonia [19]. According to World Health Organization (WHO), radiologists face difficulty distinguishing between pus-filled lesions and the lesions caused by external diagnosis conditions. Computer-aided systems are, therefore, necessary to help radiologists and doctors diagnose pediatric pneumonia and standardize chest X-ray-based diagnosis. The advancement in artificial intelligence, specifically machine learning, has made drastic improvements in the automated diagnosis of diseases. Computer-aided diagnosis (CAD) systems make the diagnosis of diseases and prediction tasks easier with minor error margins. Several CAD approaches using machine learning have been employed to diagnose major respiratory problems such as pneumonia [26, 63, 67], tuberculosis [22], Parkinson’s disease classification [52], and cardiac diseases [64]. A research team used a machine learning approach to detect diabetic retinopathy, and the results showed that their algorithms performed tantamount to ophthalmologists [8]. The conventional supervised machine learning approach requires handcrafted filters to extract the input data based on domain knowledge. Feature engineering is a cumbersome task requiring a tremendous amount of time and expertise to develop custom filters. Thus, the scalability of handcrafted filters is limited. In unsupervised learning, the features extracted from input data are not labelled, and machines have to discover patterns among the samples, which generally requires a more extensive training dataset. Large datasets, especially in the medical domain, are limited due to privacy concerns. These existing methods are highly improbable in medical imaging for an immediate and accurate CAD model.

Therefore, deep learning emerged as a state-of-art method [39], where it learns the features on its own, making it more robust compared to machine learning and thereby adding advantages to computer-aided diagnostic systems. Deep learning started with multiple hidden layers and gradually extended to multiple CNN layers resulting in deep CNN architectures. These architectures are the most commonly used technique for pattern recognition and image classification tasks [24, 40]. CheXNet, which uses a convolutional neural network that contains 121 layers, was authored by researchers at Stanford University to diagnose pneumonia [7]. A significant challenge while developing an accurate and generalized deep learning model is that the dataset should be large and well-curated for training. It has to cover variability in patients spread all over the globe, tools used for imaging, and other metrics. Collecting such a vast dataset is generally impossible for all medical diseases, especially labelling and annotating, which requires time. A modern breakthrough in artificial intelligence called transfer learning serves to overcome this impediment [45]. Based on the literature survey, it was noticed that the existing deep learning architectures perform well; however, their performance is limited. A large number of neurons in deep architectures lead to the problem of overfitting, which limits the generalizability property of models. The literature survey concludes that the existing models are not guaranteed to perform well on unseen data. Our proposal of a simple and easily replicable solution for the pediatric pneumonia diagnosis is summarized as follows:

- A feature concatenation-based stacking ensemble leveraging the strengths of various machine learning classifiers.
The contributions made in the field of pediatric pneumonia diagnosis that motivated us to advance with the idea of feature fusion-based stacked ensemble learning are as follows:

- The initial research on pediatric pneumonia diagnosis with transfer learning and the dataset’s open-source availability [38].
- A transfer learning approach using dilated convolutions and residual structures to solve the limited performance of deep convolution layers [20].
- The effects of different spatial domain image enhancement techniques on COVID-19 detection using chest X-ray images [21].
- A fusion technique involving a deep CNN model with PCA and logistic regression [41].
- A weighted average ensemble of deep CNN models incorporating deep transfer learning [9].
- A majority voting ensemble of the predictions from deep CNN models [65].
- CheXNet [7], a DenseNet121 model trained on the ChestX-ray14 dataset whose performance exceeded that of the average radiologist.

The rest of this article is organized as follows: Section 2 describes the literature survey and discusses the existing gap in the literature and how our approach completes it. The proposed approach is discussed in Section 3. Section 4 contains the description of the dataset. The performance metrics used in this study are detailed in Section 5. The experimental results are analysed and discussed with plots in Section 6. Finally, in Section 7, we conclude our work, summarizing the problem and the limitations of our approach, along with the possible future works.

2 Literature survey

Pattern classification, which earlier required specific filters for feature extraction, switched to automatic feature extraction processes using deep learning architectures. Initial studies for pattern classification were mainly based on MLP approaches. The downside to this approach was its inability to capture local information. Convolutional Neural Networks (CNNs) were introduced to solve this issue. CNN works based on the convolution operation done using the convolution layer, followed by max pooling for reduced computations. The beginning of pediatric pneumonia-based classification research started with the availability of the dataset in Kermany et al. [38]. Earlier studies on MLP and simple CNN architectures were introduced in [5]. Saraiva et al. [5] extended the findings using the cross-validation technique for extensive learning on the existing simple CNN architectures in [50]. Stanford’s research radiologists
validated the authenticity of the highly sensitive pediatric pneumonia dataset, and CheXNet was introduced to the research community [7]. The CheXNet is a deep CNN architecture based on DenseNet121 with a performance better than the average radiologists. Several custom deep CNN models were developed; however, their performance was limited. The limitation was due to the increasing number of layers in deep CNN models. The impact of the loss of spatial information in chest X-rays when passing through increasing convolutional layers was studied by Gaobo Liang et al. [20]. This spatial information is of utmost importance in detecting pediatric pneumonia. Gaobo Liang et al. [20] introduced a transfer learning-based dilated convolution CNN model to increase the receptive field, thereby reducing the risk of spatial information degradation. A purely depthwise separable convolution approach followed this variation in CNNs [51]. Siddiqi et al. [55] proposed a deep sequential CNN for pediatric pneumonia detection. Nahid et al. [54] suggested a novel two-channel CNN architecture for pneumonia diagnosis. Yu et al. [34] introduced a graph-based feature reconstruction for pediatric pneumonia detection called CGNet. The new class of deep learning architectures, Capsule Networks for pediatric pneumonia detection, was introduced by Mittal et al. [68]. Wu et al. [31] proposed a hybrid system for diagnosing pneumonia from chest X-Ray images consisting of an adaptive median filter CNN recognition model based on random forest.

Several research studies introduce architectures with competing performances. Rahman et al. [66] studied the performance of AlexNet, ResNet18, DenseNet201, and SqueezeNet utilizing transfer learning. El Asnaoui et al. [42] compared the results of fine-tuned deep learning architectures for binary classification in chest X-rays. Predefined weights are essential in determining a model’s performance in transfer learning. Mahajan et al. [3] analyzed the performance differences between CheXNet weights, ImageNet weights, and random weights for the task at hand. The problem of class imbalance in machine learning is a necessity that must be addressed. For unbiased training, machine learning relies mainly on a balanced dataset. Sampling serves as a solution to this class imbalance problem. Using the Xception network, Luján-Garca et al. [30] investigated random undersampling (RUS) for unbiased training and applied a cost-sensitive learning strategy.

Researchers began to analyze the importance of using spatial domain preprocessing techniques like Histogram Equalization, Adaptive Histogram Equalization (AHE), Contrast Limited Adaptive Histogram Equalization (CLAHE), and Gamma correction. Rubini et al. [37] examined two popular spatial processing approaches for improving MRI images: AHE and CLAHE. In [29], the authors propose using CLAHE as the preprocessing technique and custom CNN architectures for prediction. Nneji et al. [33] propose using dynamic histogram enhancement techniques for pediatric pneumonia detection. Tawsifur et al. [66] detail the effects of HE, CLAHE, image complement, gamma correction, and balance contrast enhancement approaches.

In recent studies, ensemble methods proved to show improved performances. Chouhan et al. [65] analyzed the performance of a majority voting ensemble composed of AlexNet, DenseNet121, Inception V3, GoogLeNet, and ResNet18. Sagar Kora Venu [9] proposed combining these deep CNN models - MobileNetV2, Xception, DenseNet201, ResNet152V2, and InceptionResNet - into a weighted average ensemble. Nahida et al. [41] proposed combining a deep convolutional neural network for feature extraction, principal component analysis for dimensionality reduction, and logistic regression for classification. On ensembled features from VGG-19 and CheXNet, Nahida et al. [47] proposed using Random Under Sampling, Random Over Sampling and SMOTE for classification. Islam et al. [10] illustrate using feature concatenations from SqueezeNet and InceptionV3 combined with ANNs to
make predictions. El Asnaoui et al. [18] proposed average fusion of prediction made by ResNet50, InceptionResNetV2, and MobileNetV2. Ensemble of RetinaNet and Mask R-CNN for pneumonia detection and localization was proposed by Sirazitdinov et al. [2]. A weighted average ensemble of GoogLeNet, ResNet18, and DenseNet121 with a novel assignment weights criteria was proposed in [57]. Rajaraman et al. [23] compared the performance of iterative model pruning of deep learning architectures with majority voting, weighted average ensemble averaging, and stacking on the COVID-19 dataset.

The lack of transparency is the most significant impediment to a complete transition to artificial intelligence (AI). Explainable AI (XAI), a promising research topic, has recently gained much attention. Nguyen Hai et al. [44] introduced a novel technique incorporating explainability for pneumonia detection. They proposed a mixture of custom CNN architecture with Grad-CAM for pneumonia detection. Liz et al. [36] used the XAI framework and an average ensemble of five custom CNN predictions.

An abundance of research has been done in this field. However, there exist limitations are discussed below:

1. Studies emphasize the use of CheXNet weights for custom CNN training which is a challenging task.
2. Lots of research proposes using custom complex architectures that are not easily replicable and hampers the reproducibility of the work.
3. The absence of exploring ensemble approaches pertinent to pediatric pneumonia diagnosis was observed. The same was witnessed concerning the use of machine learning classifiers.
4. Data sampling methods like RUS, ROS, and SMOTE lead to longer training times and over-fitting.
5. Most of the studies mentioned above failed to cover the aspect of feature visualization.
   Feature visualization is crucial to ensure the learned features are meaningful for predictions.
6. Most studies fail to prove the generalizability of their proposed models and approaches.

The distinguishing factor in our work compared to the rest of the works is detailed in Table 1. Our work proposes a detection pipeline to bridge the gap in the existing literature. The dataset has been redistributed for unbiased training instead of data sampling methods. We propose the use of a contrast enhancement technique for image preprocessing. The proposed methodology is based on the feature fusion of DenseNet121, DenseNet169, DenseNet201, and MobileNet architectures pre-trained on the commonly available ImageNet weights for feature extraction. The extracted feature maps from the global average pooling layer are passed to the t-SNE and CAM for visualization. Stacking ensemble classifier approach with KNN, SVC, Random Forest Classifiers, Nu-SVC, Logistic Regression, Extra-trees Classifier, Bagging Classifier, Gaussian Naïve Bayes, and AdaBoost classifier was used along with Stratified K-Fold cross-validation to overcome overfitting. All additional details are discussed in the forthcoming sections.

3 Proposed approach

This section details the workflow of the proposed pediatric pneumonia detection architecture, from the enhancement of images to the final classification, as illustrated in Fig. 1. The input
chest X-ray is preprocessed and sent to pre-trained deep CNN architectures with transfer learning to extract the features from the global average pooling layer. The next step is the column-wise concatenation of the extracted features. The concatenated features are passed through the stacking classifier with level one classifiers as SVC, KNN, Logistic Regression, NuSVC, Random Forest Classifier, AdaBoost Classifier, GaussianNB, Bagging Classifier, and Extra-trees Classifier. The binary predictions from the ML classifiers are sent to the NuSVC meta classifier for the final predictions.

The dataset contains images of varying sizes. In this study, we reshape the images according to the requirement of different deep CNN models. Each image is normalized to bring the pixel values between the range 0–1. Image augmentations were introduced on the fly using the Keras image generator as a necessary part of modeling to prevent over-fitting. The

| Reference            | Dataset            | Pretrained model                          | Feature visualization | Feature fusion | Ensemble method               |
|----------------------|--------------------|-------------------------------------------|-----------------------|----------------|------------------------------|
| Gaobo Liang et al.  | Kermany et al. [38] | Custom CNN based on dilated convolutions  | Yes                   | No             | No                           |
| Nahida Habib et al. | Kermany et al. [38] | CheXNet (DenseNet121)                     | No                    | No             | No                           |
| Mahajan et al.      | Kermany et al. [38] | InceptionV3, DenseNet121                  | No                    | No             | No                           |
| Yu et al. [34]      | Kermany et al. [38], CT-Pneumonia dataset | CGNet                                   | No                    | No             | No                           |
| Siddiqi et al.      | Kermany et al. [38] | Custom deep sequential CNN                | No                    | No             | No                           |
| Nahida Habib et al. | Kermany et al. [38], COVID-19 database | CheXNet, VGG-19                       | No                    | Yes            | Ensemble of CNN models       |
| Siddiqi et al.      | Kermany et al. [38] | PneumoniaNet                              | No                    | No             | No                           |
| Chouhan et al.      | Kermany et al. [38] | AlexNet, DenseNet121, ResNet18, InceptionV3, GoogLeNet, DenseNet201 | Yes                   | No             | Majority Voting              |
| Rahman et al.       | Kermany et al. [38] | CapsNet                                   | No                    | Yes            | Ensemble of convolution capsules |
| Mittal et al.       | Kermany et al. [38] | SqueezeNet, InceptionV3                   | No                    | Yes            | No                           |
| Islam et al.        | Kermany et al. [38] | DenseNet201, Xception, InceptionResNet, ResNet152V2, MobileNetV2 | No                    | No             | Weighted average ensemble    |
| Kora Venu Sagar     | Kermany et al. [38] | Adaptive median filter CNN                | No                    | No             | Ensemble of trees             |
| Wu et al. [31]      | Kermany et al. [38] | MobileNet, DenseNet121, DenseNet169, DenseNet201 | Yes                   | Yes            | Stacking ensemble            |
| Current work        | Kermany et al. [38] |                                           |                       |                |                              |
augmentations used include zoom, shear, and flip. Table 2 describes the corresponding augmentation values.

### 3.1 Image enhancement

Spatial image enhancement techniques are widely used to enhance specific features in an image. Contrast enhancement adjusts the relative difference between darkness and brightness to improve visibility. Contrast enhancement becomes very important in medical image diagnosis, which deals with chest X-rays and CT scans. Several studies strongly recommend using contrast enhancement techniques for better results [21, 28, 49, 53]. Adaptive Histogram Equalization (AHE) is an image processing technique for enhancing contrast.

| Methods       | Corresponding Parameters |
|---------------|--------------------------|
| Rescale       | 255                      |
| Shear         | 0.2                      |
| Zoom          | 0.2                      |
| Horizontal Flip | True                   |

![Table 2](image)
It generates histograms for each image part and uses them to redistribute its values. As a result, this is well suited for improving the local contrast and edge regions in images. However, in relatively homogeneous areas of an image, AHE tends to overamplify noise. Contrast Limited Adaptive Histogram Equalization (CLAHE), a type of adaptive histogram equalization, is thus used to restrict the amplification. CLAHE works on small regions (pixels) in the image known as tiles, where the adjacent tiles are combined using bilinear interpolation. The slope of the transformation function determines the contrast amplification in the neighborhood of a specific pixel value in CLAHE. The Blind/Reference less Image Spatial Quality Evaluator (BRISQUE) metric assessed the image quality after enhancement. The lower the BRISQUE score, the more perceptual quality it retains.

3.2 Transfer learning

The performance of any deep learning model depends on the available training data. Massive datasets are proven to increase the performance metrics of the deep learning models. However, this is not usually the case in the real world due to privacy concerns, especially in medical image data. We use transfer learning to overcome this challenge of limited data. Transfer learning uses the weights pre-trained on a similar dataset; it is then adapted to the new target task by fine-tuning it for the task at hand. We fine-tuned the already pre-trained models on ImageNet in our proposed method.

3.3 Deep learning models

The literature survey discussed earlier ascertained that using pre-trained deep CNN models provided an overall better performance. Existing pre-trained deep CNN models such as VGG16 [48], VGG19 [48], MobileNet [56], MobileNetV2 [56], InceptionResNetV2 [14], DenseNet121 [60], DenseNet169 [60], DenseNet201 [60], InceptionV3 [16], ResNet50 [59], ResNet101 [59], ResNet152 [59], ResNet50V2 [12], ResNet101V2 [12], ResNet152V2 [12], EfficientNetB0 [13] and Xception [61] are trained on the Contrast Limited Adaptive Histogram Equalization (CLAHE) enhanced dataset to select the top four best performing models. The features of the selected architectures are extracted from the global average pooling layer to retrieve a feature map for each image. These extracted features are concatenated for stacked ensemble learning.

3.3.1 MobileNet

MobileNet is a deep convolutional neural network model open-sourced by Google [56], designed for mobile and embedded vision applications. It is a streamlined architecture that uses depthwise separable convolution for lightweight computations. Depthwise separable convolution comprises two factorized convolutions: standard depthwise convolution and pointwise convolution, as shown in Fig. 2. The first phase is the depthwise spatial convolution, where the convolution is done depthwise for each input channel with a single filter. The following phase uses pointwise convolution, which applies $1 \times 1$ convolution to combine the output of depthwise convolution, thereby reducing the
computational cost. This method of factorizing the convolution process into two phases reduces the model’s size. MobileNet uses 28 convolutional layers with $3 \times 3$ depthwise separable convolutions. All layers in the architecture are followed by batch normalization and ReLU nonlinear activation. Figure 3 describes the entire architecture of MobileNet.

### 3.3.2 DenseNet

Convolutional Neural Networks rely on the gradients in the image for feature retrieval. Increasing convolution layers introduces the vanishing gradient problem, hence explaining the staggering performance with the increasing number of convolutional layers. A unique network architecture called the DenseNet was proposed by Huang et al. [60] as a solution to this vanishing gradient problem. In DenseNet, each layer is interconnected in a feed-forward manner. Features maps of all preceding layers are used as input to each layer, and their feature maps are used as inputs into all subsequent layers. This collective knowledge retained several advantages: an improved flow of information in the network alleviated the problem of relearning redundant features and decreased the number of learnable parameters.

Furthermore, since each layer has direct access to the gradients of any preceding layer, training deep network architectures become much more manageable. In addition, the regularizing effect introduced by the deep connections reduces overfitting when trained on small datasets. Our proposed approach uses DenseNet121, DenseNet169 and DenseNet201. The DenseNet121 was the first model released from the DenseNet family with 121 convolutional layers. After that, researchers began experimenting with added convolutional layers and eventually released DenseNet169 with 169 convolutional layers. The DenseNet201 with 201 layers is the most recent advancement in the DenseNet models and outperforms all other deep CNN models. Figure 4 illustrates the architecture of the DenseNet model.

### 3.3.3 Hyperparameters

Hyperparameter tuning plays a significant role in increasing the performance of the model. Thus, after extracting the feature maps from global average pooling, we added a dropout rate of 0.4 to avoid overfitting. Next, the feature maps are sent to the softmax layer, which uses the sigmoid activation function for the binary - NORMAL and PNEUMONIA classification.
Finally, we fine-tuned the models based on different optimizers to find the best hyperparameters, as shown in Table 3.

The Binary Cross Entropy calculates the difference between the expected and actual output. The value for the loss function ranges between 0 and 1 and is given by Eq. 1.

$$\text{Loss} = \sum_{i=0}^{\text{outputsize}} y_i \log(\hat{y}_i) + (1-y_i) \log(1-\hat{y}_i)$$  \hspace{1cm} (1)

3.4 T-SNE feature representation

t-Distributed Stochastic Neighbour Embedding (t-SNE) is a non-linear dimensionality reduction technique primarily used for visualizing high-dimensional data. This algorithm calculates...
the resemblance between pairs of instances in high and low dimensional space and then optimizes them using the cost function. The feature maps of the test data are visualized using the t-SNE plot. Unlike PCA, this representation allowed us to visualize the binary classes (normal and pneumonia) as clusters. It gives an insight into how well the predictions are made. This visualization gives an insight into the required classifiers for the task.

Table 3 List of hyperparameters and their values used in our study to finalize the perfect combination for the task at hand

| Hyper Parameter     | Corresponding Values                      |
|---------------------|-------------------------------------------|
| Optimizer           | Adam, SGD, Nadam, RMSprop, Adamax, Adagrad |
| Learning rate       | 0.001                                     |
| Batch size          | 32                                         |
3.5 Stacking classifier

Ensemble learning is the most commonly used technique to increase the performance of a model, thereby making it much more robust. Stacking classifiers is one of the most widely used ensemble techniques for classification or regression. This type of classifier combines different heterogeneous machine learning algorithms such as logistic regression, support vector classifier, and k-nearest classifier, unlike other techniques such as bagging and boosting. The stacking classifier consists of two stages. The first stage consists of several machine learning classifiers stacked together whose predictions are trained on the second stage classifier. The second stage classifier, called the meta classifier, provides the final classification.

3.5.1 SVC

The support vector classifier generates decision boundary in such a way that maximizes the margin between the support vectors belonging to different classes. This operation is preceded by mapping to high dimensional spaces with different kernel functions.

3.5.2 KNN

The K-Nearest Neighbour algorithm uses the proximity between k neighbours to classify the data points. The distance between the data points is calculated using different methods namely: hamming, Manhattan and Euclidean. The majority class of the first k distances in the ascending order assign the categorical class value to a given data point.

3.5.3 Logistic Regression

The Logistic Regression is the simplest solution to any binary classification task. It uses a nonlinear log transformation to map and classify points without a linear relationship. If the value of the logistic function is greater than 0.5, it is said to belong to class 1 or else class 0.

3.5.4 NuSVC

The NuSVC is very similar to SVC, except that instead of parameter C, which penalizes the wrong predictions in the optimization algorithm, the NuSVC uses the regularization parameter nu. While C has no direct interpretation, the parameter nu sets the upper bound and lower bound on the fraction of margin errors and the fraction of support vectors, respectively.

3.5.5 Random Forest Classifier

The Random Forest algorithm, used for classification and regression, is one of the most commonly used machine learning algorithms. Decision Trees are the main components of Random Forests. The algorithm becomes more advanced as the number of trees increases. It chooses the best result from the votes gathered by the trees, making it robust. The higher the number of trees in the forest, the more accurate it is, and the problem of overfitting is avoided. It comprises two phases: the first is to combine N decision trees with building a random forest, and the second is to generate predictions for each tree created in the first phase.
3.5.6 Gaussian Naïve Bayes Classifier

The Gaussian Naïve Bayes model works based on the Bayes’ theorem, assuming that the features are independent of each other. The model learns the conditional probability distribution of the input features such that the likelihood of obtaining the correct class is maximized, as shown in Eq. 2.

\[
\hat{y} = \arg\max_y P(y) \prod_{i=1}^{n} P(x_i|y)
\]  

(2)

3.5.7 AdaBoost Classifier

AdaBoost, also known as Adaptive Boosting, is a machine learning approach utilized as a part of an ensemble method. Decision trees with one level, or decision trees with only one split, are the most popular algorithm used with AdaBoost. This approach creates a model by assigning equal weights to all data points. It then gives points that are incorrectly categorized a higher weight. In subsequent models, points with greater weights are given more importance. This iterative process will continue to train models until a lower error is received.

3.5.8 Bagging Classifier

A bagging classifier is an ensemble meta-estimator technique that fits base classifiers on random subsets of the original dataset and then combines their predictions by voting or average to generate the final prediction.

3.5.9 Extra Trees Classifier

The Extra Trees Classifier uses a meta estimator to get the final prediction based on predictions made by randomized decision trees that fit on various sub-samples of the data.

3.6 Stratified K-fold cross-validation

Cross-validation is the most widely employed technique to estimate the model’s performance on unseen data. The performance on unseen data is of utmost importance for real-world deployment. The Stratified K-fold Cross-Validation technique is used when we have an imbalanced dataset. It is an extension of normal K-fold cross-validation. But here, rather than splits being completely random, the ratio of target classes is the same in each fold. Hence this method is used to preserve the class ratio for our target classes and when we have relatively fewer training examples.

4 Dataset description

All the experiments in this study were conducted on the Kermany et al. [38] dataset. The dataset contains 5856 chest X-ray images, which belong to two classes Normal (1583 X-rays) and Pneumonia (4273 X-rays). The existing dataset was imbalanced and was therefore redistributed for unbiased training. Table 4 illustrates the new data distribution. The chest X-ray images are from children aged between 1 to 5 years from the Guangzhou Women and
Children’s Medical Centre. The second row with white patches in the alveolar region of the lungs in the X-ray image shown in Fig. 5 represents the residence of pus and fluid.

### 4.1 Performance metrics

The critical aspect of evaluation metrics is distinguishing between models’ performances. The primary metrics used in this study to evaluate our models are accuracy, precision, recall, F1-score, and the AUC value. The confusion matrix shown in Fig. 6 illustrates the predictions made by the model. The rows and columns of the confusion matrix represent the actual values and the predicted values of the target variable, respectively.

| Category   | Train | Validation |
|------------|-------|------------|
| Normal     | 1400  | 183        |
| Pneumonia  | 1700  | 2573       |
| Total      | 3100  | 2756       |

True Positive (TP) - number of pneumonia X-rays correctly predicted as pneumonia

False Negative (FN) - number of pneumonia X-rays wrongly predicted as normal

True Negative (TN) - number of normal X-rays correctly predicted as normal

False Positive (FP) - number of normal X-rays predicted wrongly as pneumonia

The accuracy of a model is calculated as the ratio of correct predictions to the total number of predictions, as shown in Eq. 3. Precision is the ratio of the true positives and the number of predicted positives, as shown in Eq. 4. The recall for a class label is calculated as the ratio between the true positive and the total number of actual positives, as shown in Eq. 5. It measures the

![Fig. 5 Samples of Normal x-rays and Pneumonia x-rays from the dataset in the first row and second row respectively](image-url)
model's ability to detect positive samples. If the model correctly classifies all the positive samples, then the recall will be 1. F1 score is the harmonic mean of precision and recall, as shown in Eq. 6. AUC is a measure to distinguish between classes, i.e., when the AUC value is equal to 1, the classifier will be able to differentiate between children with and without pneumonia.

\[
\text{Accuracy} = \frac{TP + TN}{TP + TN + FP + FN}
\]  

(3)

\[
\text{Precision} = \frac{TP}{TP + FP}
\]  

(4)

\[
\text{Recall} = \frac{TP}{TP + FN}
\]  

(5)

\[
\text{F1 score} = 2 \left( \frac{\text{Precision} \times \text{Recall}}{\text{Precision} + \text{Recall}} \right)
\]  

(6)

**Table 5** Parameter used for CLAHE image enhancement with the Brisque average calculated over 20 pneumonia samples

| Image enhancement used | Clip limit | Tile grid size | Mean Brisque score |
|------------------------|------------|----------------|-------------------|
| CLAHE                  | 0.03       | 8.8            | 129.47            |
|                        | 0.02       | 8.8            | 129.22            |
|                        | 0.01       | 4.4            | 129.26            |
|                        | 0.01       | 8.8            | 129.16            |
|                        | 0.01       | 16.16          | 133.40            |
|                        | 0.01       | 32.32          | 139.53            |
|                        | 0.01       | 64.64          | 142.89            |
5 Results and discussion

Several deep CNN architectures were trained and validated on the CLAHE enhanced dataset to find the best-performing models suited for the task at hand. The parameters for CLAHE enhancement were selected meticulously to ensure that the crucial features in these chest X-rays do not disappear in the process. Table 5 describes parameters and their BRISQUE scores for the image enhancement techniques used. These scores are calculated as the mean of twenty pneumonia X-rays. The parameters resulting in the least BRISQUE scores are selected for Table 6 Fine-tuning information and the number of trainable parameters associated with each model used in our study.

| Deep CNN model      | Fine-tuned from | Total number of trainable parameters |
|---------------------|----------------|-------------------------------------|
| MobileNet V2        | 77             | 2.064,769                           |
| MobileNet           | 50             | 2.665,473                           |
| EfficientNet B0     | 118            | 3.700,169                           |
| DenseNet121         | 213            | 4.632,897                           |
| DenseNet169         | 297            | 8.544,833                           |
| DenseNet201         | 353            | 12.741,185                          |
| VGG16               | 9              | 13.569,793                          |
| Xception            | 66             | 14.860,313                          |
| InceptionV3         | 155            | 16.791,489                          |
| VGG19               | 11             | 17.699,329                          |
| ResNet50V2          | 95             | 21.352,449                          |
| ResNet50            | 87             | 21.364,225                          |
| ResNet101V2         | 188            | 30.625,793                          |
| ResNet101           | 172            | 30.640,129                          |
| ResNet152V2         | 282            | 39.836,673                          |
| ResNet152           | 257            | 39.855,617                          |
| InceptionResNet V2  | 390            | 41.922,529                          |

Table 7 Performance chart of deep learning models with values rounded off to the nearest two decimal positions on the CLAHE enhanced images.

| MODEL            | ACCURACY | PRECISION | RECALL | F1-SCORE | AUC  |
|------------------|----------|-----------|--------|----------|------|
| RESNET152        | 0.47     | 1.00      | 0.43   | 0.60     | 0.71 |
| MOBILENET V2     | 0.61     | 1.00      | 0.59   | 0.74     | 0.79 |
| RESNET101        | 0.63     | 1.00      | 0.66   | 0.80     | 0.82 |
| VGG16            | 0.85     | 1.00      | 0.84   | 0.91     | 0.90 |
| RESNET152 V2     | 0.86     | 1.00      | 0.85   | 0.92     | 0.92 |
| INCEPTION RESNET V2 | 0.89   | 0.89      | 0.88   | 0.94     | 0.94 |
| RESNET101 V2     | 0.91     | 0.96      | 0.94   | 0.95     | 0.71 |
| XCEPTION         | 0.91     | 1.00      | 0.90   | 0.95     | 0.97 |
| RESNET50 V2      | 0.92     | 1.00      | 0.91   | 0.95     | 0.95 |
| VGG19            | 0.93     | 0.93      | 1.00   | 0.97     | 0.50 |
| RESNET50         | 0.93     | 0.93      | 1.00   | 0.97     | 0.50 |
| EFFICIENT NET B0 | 0.93     | 0.93      | 1.00   | 0.97     | 0.50 |
| INCEPTION V3     | 0.94     | 1.00      | 0.93   | 0.97     | 0.96 |
| DENSENET 121     | 0.94     | 1.00      | 0.94   | 0.97     | 0.97 |
| MOBILENET        | 0.97     | 0.99      | 0.97   | 0.98     | 0.95 |
| DENSENET 169     | 0.98     | 1.00      | 0.98   | 0.99     | 0.96 |
| DENSENET 201     | 0.98     | 0.99      | 0.98   | 0.99     | 0.94 |
| PROPOSED METHOD  | 0.99     | 1.00      | 0.99   | 0.99     | 0.93 |
enhancement. Looking at the mean BRISQUE values, it is noticed that increasing the tile grid size leads to loss of perceptual quality. The parameters clip limit = 0.01, and tile grid size = 8,8 were selected for this study.

Literature survey concludes that several deep CNN architectures achieve competing accuracies for pediatric pneumonia diagnosis. To find the best architectures for the task at hand, the following models are analyzed and compared: VGG16, VGG19, MobileNet, MobileNetV2, InceptionResNetV2, DenseNet121, DenseNet169, DenseNet201, InceptionV3, ResNet50, ResNet101, ResNet152, ResNet50V2, ResNet101V2, ResNet152V2, Xception and EfficientNetB0. Each of the models above were pre-trained on ImageNet weights with the corresponding input image of size 224 × 224, 224 × 224, 224 × 224, 224 × 224, 224 × 224, 224 × 224, 224 × 224, 224 × 224, 299 × 299, 224 × 224, 224 × 224, 224 × 224, 224 × 224, 299 × 299, 299 × 299, 299 × 299, and 224 × 224 respectively. All the architectures were trained for 30 epochs with Adam as the optimization function and the learning rate as 0.001 in Google Colab resourced with K80 GPU and 12 GB RAM. Tensorflow2 and Keras2 were used to build and evaluate the models. The models were fine-tuned from their specified layers, as shown in Table 6. Table 6 also includes the trainable parameters for all the models used in this study.

Fig. 7 Confusion matrix for (a) MobileNet predictions on the test data (b) DenseNet121 predictions on the test data (c) DenseNet169 predictions on the test data (d) DenseNet201 predictions on the test data
The results in Table 7 show that DenseNet169 and DenseNet201 are the best performing models. The family of DenseNet models performs consistently well compared to other architectures. DenseNet169 and DenseNet201 achieve the highest accuracy of 97.79%. MobileNet precedes them with an accuracy of 97%, followed by DenseNet121 with an accuracy of 94%. One common attribute in the best-performing models is the residual connection. The residual connections are a key factor that has suppressed over-fitting and thus enabled the above models to perform well on the test data.

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Fig. 8 ROC curve for (a) MobileNet predictions on the test data (b) DenseNet121 predictions on the test data (c) DenseNet169 predictions on the test data (d) DenseNet201 predictions on the test data

Fig. 9 MobileNet model performance on the validation set using different optimizers
knowledge and residual connections in Densenets has enabled it to achieve the highest accuracy. The confusion matrix for the test data predictions from MobileNet, DenseNet121, DenseNet169, and DenseNet201 are shown in Fig. 7. MobileNet predicts 67 false negatives and 13 false positives. DenseNet121, DenseNet169 and DenseNet201 predicts 9, 49, 43 false negatives and 66, 12, 18 false positives respectively. DenseNet121 has the lowest false-negative
predictions in the Densenet family. The confusion matrices conclude that the architectures are unable to deal with false predictions. The ROC curves for test data predictions using MobileNet, DenseNet121, DenseNet169, and DenseNet201 are shown in Fig. 8. DenseNet201 has the lowest AUC value of 0.94 compared to the rest. DenseNet121 and DenseNet169 share the highest AUC score of 0.96. Our proposed method achieves the highest accuracy, precision, recall, and f1-score but shows a reduced AUC score.

Extensive experiments were conducted for hyperparameter tuning of each of the best-performing models. The models were fine-tuned on different optimizers to find the best fit for
the task at hand. The Adam optimizer performs best for all models, as seen in Figs. 9, 10, 11 and 12. Based on Figs. 9, 10, 11, and 12, the optimal hyperparameters with the Adam optimizer and a constant learning rate of 0.001 were chosen for feature extraction. The validation loss and accuracy plots of MobileNet, DenseNet121, DenseNet169, and DenseNet201 are shown in Figs. 13, 14, 15 and 16. All the architectures exhibit initial oscillatory behavior. This oscillation is persistent in MobileNet and DenseNet121 whereas, in DenseNet169 and DenseNet201, it gradually reduces with the increasing number of epochs. For the MobileNet model, the validation loss and accuracy are constrained to 1–0 and 0.75–1, respectively. For the DenseNet121 model, the validation loss and accuracy are constrained to 1.2–0 and 0.8–1, respectively. For the DenseNet169 model, the validation loss and accuracy are constrained to 3.7–0 and 0.4–1, respectively. For the DenseNet201 model, the validation loss and accuracy are constrained to 2.8–0 and 0.6–1, respectively.

With the best performing models selected from Table 7, we propose using MobileNet, DenseNet121, DenseNet169, and DenseNet201 for feature extraction. The extracted features
are concatenated for the final classification. The extracted features are visualized using the t-SNE representation for the layman’s interpretability of the features predicted by the models. The parameter values used for visualization are $n\_\text{components} = 2$, perplexity = 40, and $n\_\text{iter} = 300$. Figs. 17, 18, 19, and 20 illustrate the t-SNE plot of the extracted feature maps from each selected architecture.

The t-SNE feature representations conclude that all the models have minor overlapping cluster formations and are non-linearly separable. In the cluster formation of the concatenated

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**Fig. 18** t-SNE feature representation of the test data extracted from the DenseNet121 model

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**Fig. 19** t-SNE feature representation of the test data extracted from the DenseNet169 model
features, Fig. 21, we notice fewer overlaps than their individual counterparts. The feature representations indicate a possible classifier that can deal with such complexity. This study proposes using the stacking classifier to deal with the non-linearly separable classification.

Class Activation Maps (CAM) are employed to understand the region of interest proposed by these deep CNN architectures. An overview of the regions of interest in the predictions is crucial before real-time deployment as a life-saving resource. The first set of class activation
maps for each model misclassified normal as pneumonia, normal as pneumonia, and pneumonia as normal, as shown in Figs. 22, 24, 26, and 28, respectively. The second set of the class activation maps for each model correctly classifies normal and pneumonia X-rays, as shown in Figs. 23, 25, 27, and 29, respectively. From the CAMs of each model, we conclude that the misclassified samples are possibly due to over-fitting. The plots conclude that the accurateness of class activation maps of the concatenated features relies purely on the strength of the individual models. The CAMs of the extracted features from MobileNet, DenseNet121, DenseNet169, and DenseNet201 and their concatenated features are shown in Figs. 30 and 31 for pediatric pneumonia diagnosis for misclassified and correctly classified samples, respectively.

Several machine learning classifiers were trained on the concatenated features and validated against the stacking classifier. Table 8 concludes that the stacking classifier outperforms all machine learning classifiers by leveraging the strength of individual estimators. The first stage in the stacking classifier leverages the RandomForests, Support Vector Classifier, KNeighborsClassifier, GNBClassifier, LogisticRegression, Nu-Support Vector Classifier, ExtraTreesClassifier, AdaBoost Classifier, and a Bagging Classifier. The hyperparameters for
each of these classifiers were selected using the Bayesian optimization strategy in optune and are
detailed in Table 9. Individual predictions from each of the five classifiers are sent to the meta-
classifier for the final classification. The meta classifier uses Nu-SVC with nu = 0.5, kernel =
"rbf", degree = 3. Stratified K-Fold cross-validation with n_splits = 30 was employed to help the
model learn the most from the existing limited dataset and prevent over-fitting.

The confusion matrix for Stratified K-Fold cross-validation stacking classifier predictions
on the test set is shown in Fig. 32. The false predictions are lower in number compared to the
raw predictions from the deep CNN architectures due to the collective strength of individual
CNN models and the wide range of machine learning classifiers. The perfect combination of
base estimators is of utmost importance. Though the accuracy increased by 0.83%, the AUC
value has not increased. The number of false-positive predictions is minimal and equal to the
number of false positives in DenseNet169. Though the number of false negatives is not as
minimal as that of DenseNet121, the proposed approach has overall reduced false predictions.

Looking at the confusion matrix, the loss of 1.38% in the model’s accuracy might favorably be
due to the imbalanced dataset or insufficient training samples. Another possible reason is the
use of a simple feature concatenation technique that solely relies on their individual predictors. The proposed method achieves an accuracy of 98.62%.

Table 10 compares our proposed approach’s performance, technique, and classification classes with other recent works. The proposed work exhibits competing performances with other literary works. All the works mentioned in the Table 1 validated their results tested on the Kermany et al. [38] dataset. The advantage of the proposed method compared to other works in Table 10 is in the visualization of features learned by the model using both CAMs and t-SNE plots. Since the feature concatenation of MobileNet, DenseNet121, DenseNet169, and DenseNet201 are used as the feature extractor is based on the commonly available ImageNet weights, reproducibility is easier. In addition to stacking various machine learning classifiers for rich predictions, the proposed models were tested on unseen similar lung disease datasets for generalization and robustness, which was previously absent in recent works.

The proposed model’s limitation is its heavy reliance on the individual deep CNN architectures and the correct combination of base classifiers for accurate classification. It was noticed that the meta classifier played a crucial role in determining the performance of
the proposed architecture. While a few experiments resulted in AUC scores reaching 98%, it reduced other metrics by a negligible percentage, and the same was observed vice-versa. The dependence on the fit deep CNN models is a drawback as any one of the models being overfit can affect the final predictions. Though ensemble learning was used to prevent overfitting, the performances indicate the need for cost-sensitive learning-based approaches to learn from the small-scale imbalanced dataset. Other limitations include the inability to subclassify into different stages of pediatric pneumonia like early, latent and severe.

5.1 Robustness and generalization of the proposed approach for lung disease classification

Generalization is an important criterion to be considered before real-time deployment. The proposed deep CNN models trained on the Kermany et al. [38] pediatric pneumonia dataset are tested on similar lung diseases like COVID-19, Tuberculosis, and Pneumonia [6, 15, 27, 43, 62]. The extracted features from each of MobileNet, DenseNet121, DenseNet169, and

![Fig. 25 Class activation maps of correctly classified X-rays from DenseNet121 (row 1: normal classified as normal, row 2: pneumonia classified as pneumonia, row 3: pneumonia classified as pneumonia)
DenseNet201 are concatenated and classified using the k-means clustering algorithm. The k-means algorithm was employed to infer the ability to correctly form clusters in the extracted features for reliable classification of similar lung diseases. The results for each dataset are analyzed as classification reports and confusion matrices.

In the COVID-19 vs. normal vs. pneumonia classification dataset [4], the proposed feature fusion achieves an accuracy of 45% with good classification margins for the normal category, as shown in Table 11. The models achieve 1643 correct classifications for normal images. It shows misclassification for the viral adult pneumonia, and COVID-19 predictions as the model are unable to extract the required features for COVID-19 and adult pneumonia detection. The same can be observed in the normal vs. pneumonia vs. tuberculosis classification dataset [4, 58], where the proposed feature fusion achieves an accuracy of 48% with good classification margins for the normal category, as shown in Table 12. The models achieve 1646 correct classifications for normal images. It shows misclassification for the viral adult pneumonia and tuberculosis predictions as the models are unable to extract the required features for the same. The proposed method achieves an accuracy of 60% in the normal vs. pneumonia dataset [43].
with 80 false negative predictions, as shown in Table 13 and Fig. 33. The results show the proposed models’ capability to classify normal chest X-rays from chest X-rays with infections correctly. The results also conclude that though the challenges of pediatric pneumonia diagnosis are characteristically different from adult pneumonia and other similar lung diseases, the proposed method can be extended to aid with diagnosing other lung diseases.

6 Conclusion and future work

This work proposes a computer-aided diagnosis model for detecting pediatric pneumonia using chest X-rays. The use of low radiation levels in chest X-rays for children makes detection a difficult task. Highly trained physicians then diagnose these chest X-rays meticulously to confirm the presence of the acute infection. This process requires loads of time and heavily relies on the availability of experts, which is not always feasible. Other works in the same field include using novel architectures and an ensemble of deep CNN models with the
added advantage of using an augmented dataset to increase the number of samples in each category. For predictions, the proposed approach uses Contrast Limited Adaptive Histogram Equalization (CLAHE) enhanced chest X-ray images. Our work uses the existing deep CNN models for feature extraction; visualized using t-SNE feature representations and class activation maps. The best-performing models’ features are concatenated and sent to the stacking classifier for the final - Normal, Pneumonia classification. Redistribution of the dataset instead of added augmentations to ensure unbiased training was the initial dominant factor for reliable performance. Our work uses transfer learning on pre-trained models to compensate for the availability of a limited dataset and introduces data augmentations to prevent overfitting. The features from MobileNet, DenseNet121, DenseNet169 and DenseNet201 are concatenated for stacked ensemble learning. The advantage of the proposed models for this task in specific has been studied in detail. A stacking classifier covering nearly all machine learning models was employed. Stacking classifier with Stratified K-Fold cross-validation results in an accuracy of 98.62%. The proposed models were tested on other lung disease datasets to validate the performance across unseen data for inference.

Fig. 28 Class activation maps of misclassified X-rays from DenseNet201 (row 1: normal classified as pneumonia, row 2: normal classified as pneumonia, row 3: pneumonia classified as normal)
As future work, we would like to explore more about other histogram equalization techniques such as Gamma Correction (GC), Multipeak Histogram Equalization (MPHE), and Multipurpose Beta Optimized Bihistogram Equalization (MBOBHE) for pediatric pneumonia prediction. We notice minor outliers and feature overlap between normal and pneumonia chest X-rays in the t-SNE plots. The t-SNE plots conclude the need for a better architecture to capture many more intrinsic patterns. Introducing attention-based networks and transformers is another possible future direction. Attention-based concatenation instead of simple feature concatenations can be utilized. Augmentations for training might help the model perform much better and reduce the current misclassification rate of 1.38%. With our work performing better CheXNet, which has surpassed normal radiologist level categorization, it will be of immense help to all physicians and radiologists for accurate diagnoses in a matter of seconds. This early detection will help reduce the mortality rate of children suffering from pneumonia.

Fig. 29 Class activation maps of correctly classified X-rays from DenseNet201 (row 1: normal classified as normal, row 2: pneumonia classified as pneumonia, row 3: pneumonia classified as pneumonia)
Fig. 30 Class activation maps of an X-ray with pneumonia misclassified as normal from each of the selected architectures. As seen the CAM of the final concatenated feature map is spread over the entire image due to the drawback of using simple feature concatenation techniques.
Fig. 31 Class activation maps of an X-ray without pneumonia correctly classified as normal from each of the selected architectures. As seen the CAM of the final concatenated feature map is concentrated to a specific part of the entire X-ray.
### Table 8: Performance comparison of different machine learning classifiers with the stacking classifier with values rounded off to the nearest two decimal positions

| Classifier                  | Accuracy | Precision | Recall | F1-score | AUC  |
|-----------------------------|----------|-----------|--------|----------|------|
| Logistic regression         | 97.42    | 99.96     | 97.28  | 98.60    | 98.37|
| Support vector classifier   | 97.75    | 99.92     | 97.67  | 98.78    | 98.29|
| Nu-Support vector classifier| 98.55    | 99.92     | 97.32  | 98.60    | 98.11|
| K-Nearest classifier       | 97.24    | 99.96     | 97.09  | 98.50    | 98.27|
| MLP classifier              | 97.24    | 99.96     | 97.09  | 98.50    | 98.27|
| Gaussian naïve bayes        | 97.57    | 99.60     | 97.78  | 98.69    | 96.16|
| BernoulliNB                 | 96.84    | 99.60     | 97.00  | 98.29    | 95.77|
| Gradient boosting classifier| 96.99    | 99.76     | 97.28  | 98.52    | 97.27|
| XGB classifier              | 96.52    | 99.84     | 96.42  | 98.10    | 97.12|
| DecisionTree classifier     | 97.61    | 99.88     | 97.55  | 98.70    | 97.96|
| RandomForest classifier     | 97.71    | 99.88     | 97.67  | 98.76    | 98.01|
| ExtraTrees classifier       | 96.77    | 99.76     | 96.77  | 98.24    | 96.75|
| Bagging classifier          | 97.50    | 99.80     | 97.51  | 98.64    | 97.39|
| AdaBoost classifier         | 97.35    | 99.84     | 97.32  | 98.56    | 97.57|
| LGB classifier              | 97.53    | 99.84     | 97.51  | 98.66    | 97.66|
| CatBoost classifier         | 97.17    | 99.80     | 97.16  | 98.46    | 97.22|
| Proposed method             | 98.62    | 98.99     | 99.53  | 99.26    | 93.17|

### Table 9: Fine-tuning information and the number of trainable parameters associated with each model used in our study

| Classifier                  | Hyperparameters                                                                                                                                 |
|-----------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------|
| Support Vector classifier   | C=2, kernel=’poly’, degree=3, gamma=’scale’, coef0=0.0, tol=0.0003                                                                                                                                 |
| K-Nearest Neighbors classifier| n_neighbors=175, weights=’uniform’, algorithm=’brute’, p=2, leaf_size=42                                                                                                                                  |
| Logistic Regression         | tol=3.89253667505987e-05, C=1, penalty=’l2’, max_iter=102, nu=0.38, kernel=’rbf’, gamma=’scale’, tol=2.522320748167019e-05, probability=False, n_estimators=100, criterion=’gini’, min_samples_split=2, min_samples_leaf=1, min_weight_fraction_leaf=0.0, min_impurity_decrease=0.0, ccp_alpha=0.0 |
| NuSVC                       | n_estimators=100, criterion=’gini’, min_samples_split=2, min_samples_leaf=1, min_weight_fraction_leaf=0.0, min_impurity_decrease=0.0, ccp_alpha=0.0                                                                 |
| GaussianNB                  | var_smoothing=1e-09                                                                                                                                                                                         |
| AdaBoostClassifier          | base_estimator=DecisionTreeClassifier, n_estimators=50, learning_rate=1.0, algorithm=’SAMME.R’                                                                                                             |
| BaggingClassifier           | base_estimator=DecisionTreeClassifier, n_estimators=10, max_samples=1, max_features=1                                                                                                                      |
| ExtraTreesClassifier        | n_estimators=10, criterion=’gini’, min_samples_split=2, min_samples_leaf=1, max_features=’auto’                                                                                                        |
Table 10 Performance of other recent works on the Kermany et al. [38] dataset with values rounded off to the nearest two decimal positions

| Author                  | Classes              | Technique                                      | Accuracy (%) | Precision (%) | Recall (%) | AUC (%) | Feature Extraction Time |
|-------------------------|----------------------|-----------------------------------------------|--------------|---------------|------------|---------|-------------------------|
| Kermany et al. [38]     | Normal and Pneumonia | Inception V3 pretrained CNN model             | 92.8         | 90.1          | 93.2       | –       | –                       |
| Nahida et al. [54]      | Normal and Pneumonia | Two-Channel CNN model                         | 97.92        | 98.38         | 97.47      | 97.97   |                         |
| Stephen et al. [4]      | Normal and Pneumonia | Custom CNN model without Transfer Learning    | 93.73        | –             | –          | –       |                         |
| Chouhan et al. [65]     | Normal and Pneumonia | Majority voting ensemble model                 | 96.39        | 93.28         | 99.62      | 99.34   | –                       |
| Rajaraman et al. [58]   | Normal and Pneumonia | Custom VGG-16 model                           | 96.2         | 97.0          | 99.5       | 99.0    | –                       |
| Siddiqi et al. [55]     | Normal and Pneumonia | Deep sequential CNN model                     | 94.39        | 92.0          | 99.0       | –       | –                       |
| Hashmi et al. [43]      | Normal and Pneumonia | Weighted classifier                            | 98.43        | –             | –          | 99.76   |                         |
| Yu Xiang et al. [34]    | Normal and Pneumonia | CGNET                                         | 98.72        | 97.48         | 99.15      | –       | –                       |
| El Asnaoui et al. [42]  | Normal and Pneumonia | Deep CNN model                                | 96.27        | 98.06         | 94.61      | –       | –                       |
| Mittal et al. [68]      | Normal and Pneumonia | CapsNet architecture                          | 96.36        | –             | –          | –       | –                       |
| Rahman et al. [66]      | Normal and Pneumonia | Deep CNN model                                | 98.0         | 97.0          | 99.0       | 98.0    | –                       |
| Sagar Kora Venu et al. [9]| Normal and Pneumonia | Weighted average ensemble model                | 98.46        | 98.38         | 99.53      | 99.60   | –                       |

Fig. 32 Confusion matrix for predictions made on the test dataset using the stacked classifier with feature concatenations
Table 10 (continued)

| Author | Classes                  | Technique                                                                 | Accuracy (%) | Precision (%) | Recall (%) | AUC (%) | Feature Extraction Time |
|--------|--------------------------|----------------------------------------------------------------------------|---------------|---------------|------------|---------|------------------------|
| Toğar et al. [11] | Normal and Pneumonia | Deep CNN model                                                             | 96.84         | 96.88         | 96.83      | 96.80   | –                      |
| Nahida et al. [47] | Normal and Pneumonia | SMOTE on ensembled features from VGG-19 and CheXNet                      | 98.90         | –             | –          | 99.00   | –                      |
| Islam et al. [10] | Normal and Pneumonia | Feature concatenations with ANN                                            | 98.99         | 99.18         | 98.90      | –       | 00:04:16               |
| Proposed Work | Normal and Pneumonia | Stacking Classifier based feature concatenations from MobileNet, DenseNet121, DenseNet169 and DenseNet201 | 98.62         | 98.99         | 99.53      | 93.17   | 00:01:30               |

Table 11 Classification report for predictions made on the COVID-19 vs normal vs pneumonia classification dataset [4] using feature fusion from MobileNet, DenseNet121, DenseNet169 and DenseNet201

|                  | Precision | Recall | F1-score | Support |
|------------------|-----------|--------|----------|---------|
| COVID-19         | 0.24      | 0.29   | 0.26     | 1086    |
| NORMAL           | 0.67      | 0.54   | 0.60     | 3053    |
| VIRAL PNEUMONIA  | 0.09      | 0.18   | 0.12     | 407     |
| accuracy         |           |        | 0.45     | 4546    |
| macro avg        | 0.34      | 0.34   | 0.33     | 4546    |
| weighted avg     | 0.52      | 0.45   | 0.48     | 4546    |

Table 12 Classification report for predictions made on the normal vs pneumonia vs tuberculosis classification dataset [4, 27, 58] using feature fusion from MobileNet, DenseNet121, DenseNet169 and DenseNet201

|                  | Precision | Recall | F1-score | Support |
|------------------|-----------|--------|----------|---------|
| NORMAL           | 0.83      | 0.53   | 0.65     | 3090    |
| VIRAL PNEUMONIA  | 0.10      | 0.17   | 0.13     | 381     |
| TUBERCULOSIS     | 0.06      | 0.27   | 0.10     | 231     |
| accuracy         |           | 0.48   | 0.29     | 3702    |
| macro avg        | 0.33      | 0.33   | 0.29     | 3702    |
| weighted avg     | 0.71      | 0.48   | 0.56     | 3702    |

Table 13 Classification report for predictions made on the normal vs pneumonia classification dataset [15, 43] using feature fusion from MobileNet, DenseNet121, DenseNet169 and DenseNet201

|                  | Precision | Recall | F1-score | Support |
|------------------|-----------|--------|----------|---------|
| NORMAL           | 0.56      | 1.00   | 0.71     | 100     |
| PNEUMONIA        | 1.00      | 0.20   | 0.33     | 100     |
| accuracy         |           | 0.60   | 0.60     | 200     |
| macro avg        | 0.78      | 0.60   | 0.52     | 200     |
| weighted avg     | 0.78      | 0.60   | 0.52     | 200     |
Informed consent

None.

Data Availability

The data that support the findings of this study are available from the first author upon reasonable request.

Declarations

Compliance with ethical standards

None.

Conflicts of interest/competing interests

The authors declare no conflict of interest.
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