
PNEUMONIACXR: AI-ENABLED PNEUMONIA DETECTION

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ABSTRACT

Chest X-ray (CXR) interpretation for pneumonia diagnosis can be challenging, especially in resource-limited settings. We developed machine learning models to classify COVID-19 pneumonia, non-COVID pneumonia, and normal CXR images. Employing a multi-modal feature engineering approach (HOG, radiomics, ResNet) with dimensionality reduction, we explored Logistic Regression, Support Vector Machines (SVM), Gradient Boosting, and Random Forest classifiers. Logistic Regression and SVM achieved the highest accuracies (>0.89), with Logistic Regression demonstrating exceptional efficiency. Our findings suggest the potential of machine learning to improve pneumonia diagnosis, particularly in resource-limited settings. Future work should prioritize model explainability, probabilistic outputs, and rigorous validation across diverse datasets.

1 Introduction

The global COVID-19 outbreak has emphasized the need for fast, accurate diagnostic methods. Chest X-ray (CXR) imaging is a widely accessible tool for diagnosing respiratory diseases. However, traditional reliance on radiologist expertise can lead to variability and create diagnostic bottlenecks. Automated systems could offer greater diagnostic consistency and alleviate pressure on medical professionals.

Pneumonia, a common respiratory condition manifests in chest X-rays (CXR) as airspace opacification, where normally dark areas turn white/grey due to alveoli filling with infectious materials. These changes, ranging from patchy to confluent, are key indicators for pneumonia, alongside air bronchograms [5]. For COVID-19, specific patterns include patchy and/or confluent asymmetric airspace opacities, bilateral ground-glass opacities or consolidations with a peripheral and mid-to-lower lung distribution, and ill-defined margins of lesions [3] [8] [9]. However, it's important to note that while these visual characteristics are suggestive, there may be some overlap in the appearance of different respiratory diseases on chest X-rays. This diagnostic complexity highlights the potential value of automated diagnostic tools.

We propose a machine learning-based approach to classify chest X-ray images into COVID-19 pneumonia, Non-COVID pneumonia, and normal cases. This approach leverages diverse features, including edge gradients (HOG), texture patterns (radiomics), and deep learning representations (ResNet). Following feature extraction, we employ Principal Component Analysis (PCA) for dimensionality reduction and explore various classification models (e.g., SVM, Logistic Regression, Gradient Boost, Random Forest). Our goal is to identify a model that balances strong performance with computational efficiency, enabling rapid training and diagnosis. This system has the potential to assist clinicians in making more informed and timely diagnosis, ultimately improving patient outcomes.

2 Data and Preprocessing

Our study leverages the COVID-QU-Ex Dataset, a publicly available collection of chest-X ray images compiled from 10 separate studies [11]. The dataset comprises 32,103 chest X-ray images categorized as COVID-19 positive, non-COVID infections, and normal cases. To ensure a balanced representation of classes, we performed random undersampling, resulting in an even 33% split for each. Table 1 summarizes the class distribution before and after applying random undersampling.

Data Set	COVID-19		Non-COVID		Normal	
	Before	After	Before	After	Before	After
Training	7,658	6,849	7,208	6,849	6,849	6,849
Validation	1,903	1,712	1,802	1,712	1,712	1,712
Test	2,395	2,140	2,253	2,140	2,140	2,140
Total	11,956	10,701	11,263	10,701	10,701	10,701

Table 1: Number of Samples by Diagnostic Class and Split, Before and After Resampling

Each record included a chest X-ray image and a corresponding lung mask delineating the lung regions. The images retained their original grayscale intensities (0 to 255), while lung masks were binarized for effective segmentation. Fig. 1 highlights inconsistencies in the original dataset, including variations in contrast, the proportions of each image occupied by the lungs (with some images exhibiting black borders), and inconsistent image orientation.

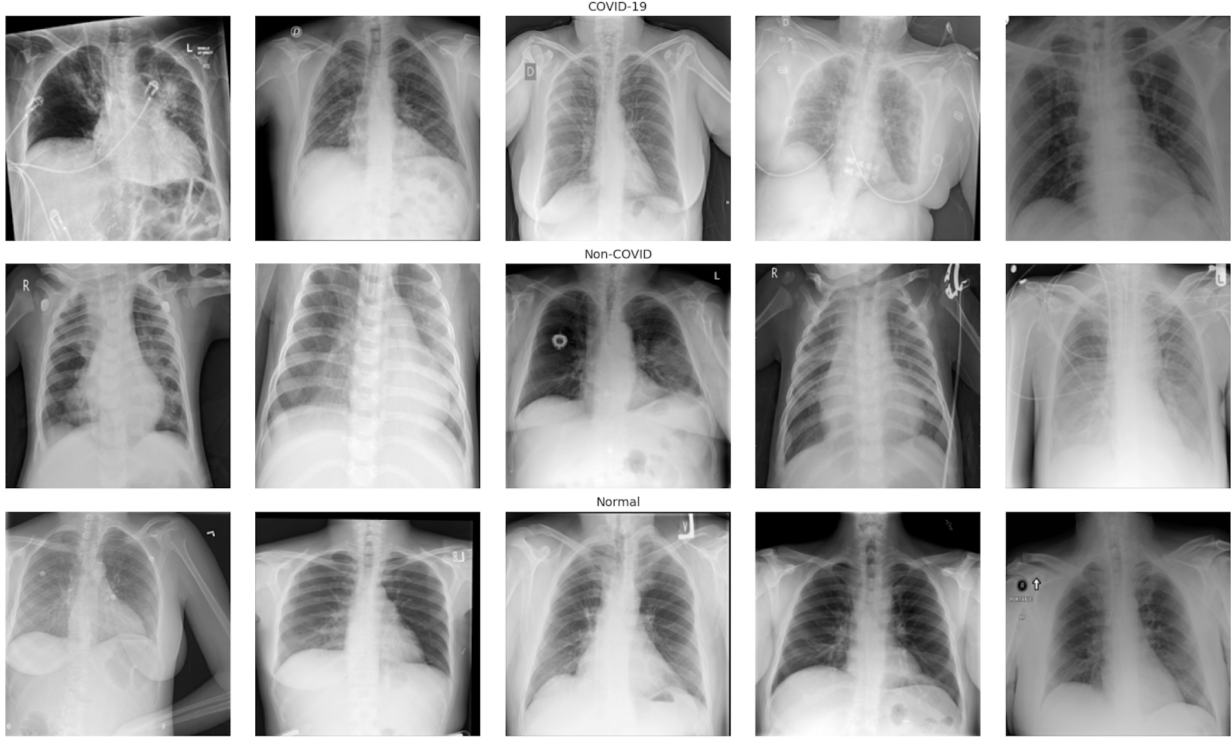


Figure 1: Unprocessed Sample Images

To address these inconsistencies, we performed several preprocessing steps. First, we applied z-score scaling to standardize the contrast range within each image. Next, we cropped images and their corresponding masks, focusing on lung regions identified by the masks. We added padding during cropping to preserve valuable anatomical features outside the lungs (e.g., the diaphragm). Finally, we used OpenCV to rotate images, achieving a standard perpendicular orientation based on spinal cord alignment.

3 Feature Engineering

To effectively analyze the complex visual patterns of pneumonia, we employed a multi-modal feature engineering approach combining Histogram of Oriented Gradients (HOG), Radiomics, and features extracted via a pre-trained ResNet model. This approach was designed to capture diverse aspects of chest X-ray images, with HOG emphasizing edge-based patterns of opacity, Radiomics quantifying subtle textural variations, and ResNet extracting deeper patterns through its learned representations.

For HOG and Radiomics features, Hyperparameter tuning was performed on a 10% subset of the training data using Optuna, a well-established optimization library. The objective function was set to maximize the sum of mutual information between the extracted HOG features and the disease labels (pneumonia or normal). Mutual information quantifies the amount of information one variable (features) provides about another (disease labels). In feature engineering, higher mutual information between features and the class labels suggests the features' potential usefulness for classification.

Following individual feature extraction, we adopted Principal Component Analysis (PCA) for dimensionality reduction, ensuring the most informative aspects of each feature set are retained while mitigating redundancy and noise, and reducing computational requirements during model training. The culmination of this process involved concatenating the principal components of the feature sets, thus creating a comprehensive representation of each image in the dataset.

3.1 Histogram of Oriented Gradients (HOG)

HOG is a feature descriptor well-suited for analyzing variations in texture within areas of opacity in chest X-rays. By detecting the distribution of gradient orientations, HOG effectively highlights the distinct edge patterns associated with consolidations, ground-glass opacities, and their relative locations within the lung field. These characteristics can aid in distinguishing between different pneumonia types.

Parameters	Initial Configuration	Search Values	Parameter Importance	Best Configuration
Image Size	(128, 255)	(128,255), (255,128), (255,255)	30.9%	(255, 255)
Orientations	9	7, 8, 9	4.3%	9
Pixels per Cell	(16, 6)	(8,8), (12,12), (16,16)	58.7%	(16, 16)
Cells per Block	(2, 2)	(2,2), (3,3)	6.2%	(2, 2)

Table 2: HOG Hyperparameter Tuning Results

Key parameters for HOG include image size, pixels per cell, and cells per block. These parameters control the level of detail captured, the granularity of image partitioning, and the extent of local gradient patterns considered. Table 2 summarizes our hyperparameter tuning process for HOG, initially using configurations aligned with a similar study on pneumonia detection [7]. Hyperparameter tuning was conducted using 40 trials, in which trials leading to vector sizes exceeding 15,000 were discarded. L2 normalization was applied across all tested configurations.

Outcomes of the HOG hyperparameter tuning study (Table 2) reveal that pixels per cell is the most important parameter, followed by image size. This is likely because pixels per cell directly controls the fineness of edge detection. A smaller pixels-per-cell leads to the detection of more subtle variations in texture, crucial for distinguishing different types of opacities. Image size is also important, as it determines the overall visual field considered by the HOG descriptor. Interestingly, cells per block and orientations had a relatively minor impact on mutual information.

Based on these findings, the configuration for HOG feature extraction was set with (255, 255) image size (maintaining the original image size), (16,16) pixels per cell, and (2,2) cells per block, with L2 normalization applied. This improved total mutual information in the validation set by 230%, demonstrating the value of higher-resolution detail for pneumonia classification.

Examination of sample HOG images 2 helps elucidate how this descriptor aids in pneumonia classification. The emphasis on edges allows HOG to delineate areas of consolidation. However, in cases of diffuse opacities, the edges become less distinct, reflecting HOG's primary focus on localized gradient patterns.

3.2 Radiomics

Radiomics, the process of extracting quantitative features from medical images, offers valuable insights beyond traditional visual assessment. It analyzes subtle textural patterns correlated with underlying disease characteristics. Applied to CXR images, radiomics can differentiate between normal tissue, non-Covid pneumonia and COVID pneumonia based on variations in texture. This technique complements HOG, which primarily focuses on shape and edge patterns. Radiomics' potential in pneumonia classification is supported by recent studies. One study demonstrated its potential in differentiating COVID-19 pneumonia from influenza virus pneumonia based on CT scan images [1]. Another successfully used a radiomics-based model to detect pneumonia in CXR images based on textural features [6].

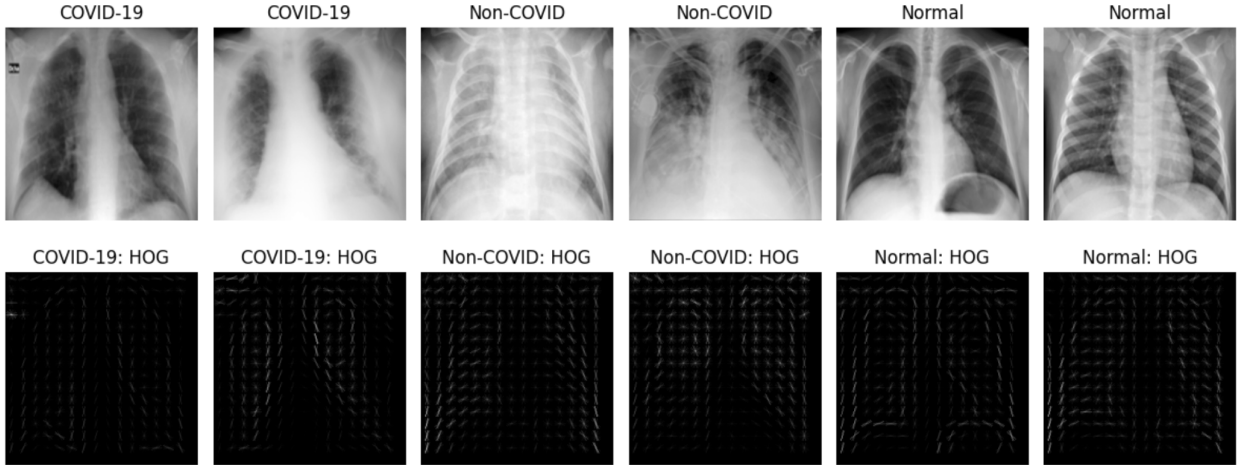


Figure 2: HOG Features for Example Images by Diagnosis Class with (255, 255) Image Size, (16, 16) Pixels per Cell, (2,2) Cells per Block, and L2 Normalization

We extracted radiomics features from the segmented lung regions (defined by the provided masks) using the well-established PyRadiomics package [12]. Images were z-score normalized, but cropping and rotation adjustments were not used due to the invariance of the selected features to orientation and the use of lung segmentation.

Radiomics feature engineering followed a four-step process. First, we explored a wide range of potential textural features on a 10% subset of the training data, using multiple image transformations and feature classes. Next, we optimized hyperparameters for the most promising transformations and feature classes based on mutual information analysis, again utilizing the 10% subset. We then performed feature selection on the complete training dataset, leveraging the insights from hyperparameter optimization to identify the most informative features while minimizing redundancy. Finally, we extracted the selected features with their optimal hyperparameter configurations across the full training, validation, and test data splits for use in downstream classification models.

3.2.1 Initial Feature Extraction

To thoroughly analyze textural changes relevant to pneumonia, we employed multiple image transformations. The original image provided a baseline, while the gradient image emphasized edges and transitions in intensity to highlight potential abnormalities by calculating the rate of change in pixel values. Local Binary Patterns (LBP2D), which compare a central pixel to its neighbors to create a binary code, captured microtextural variations that may indicate early signs of pneumonia. Finally, wavelet transform decomposed the image into various frequency scales, uncovering textural details potentially obscured at the original resolution.

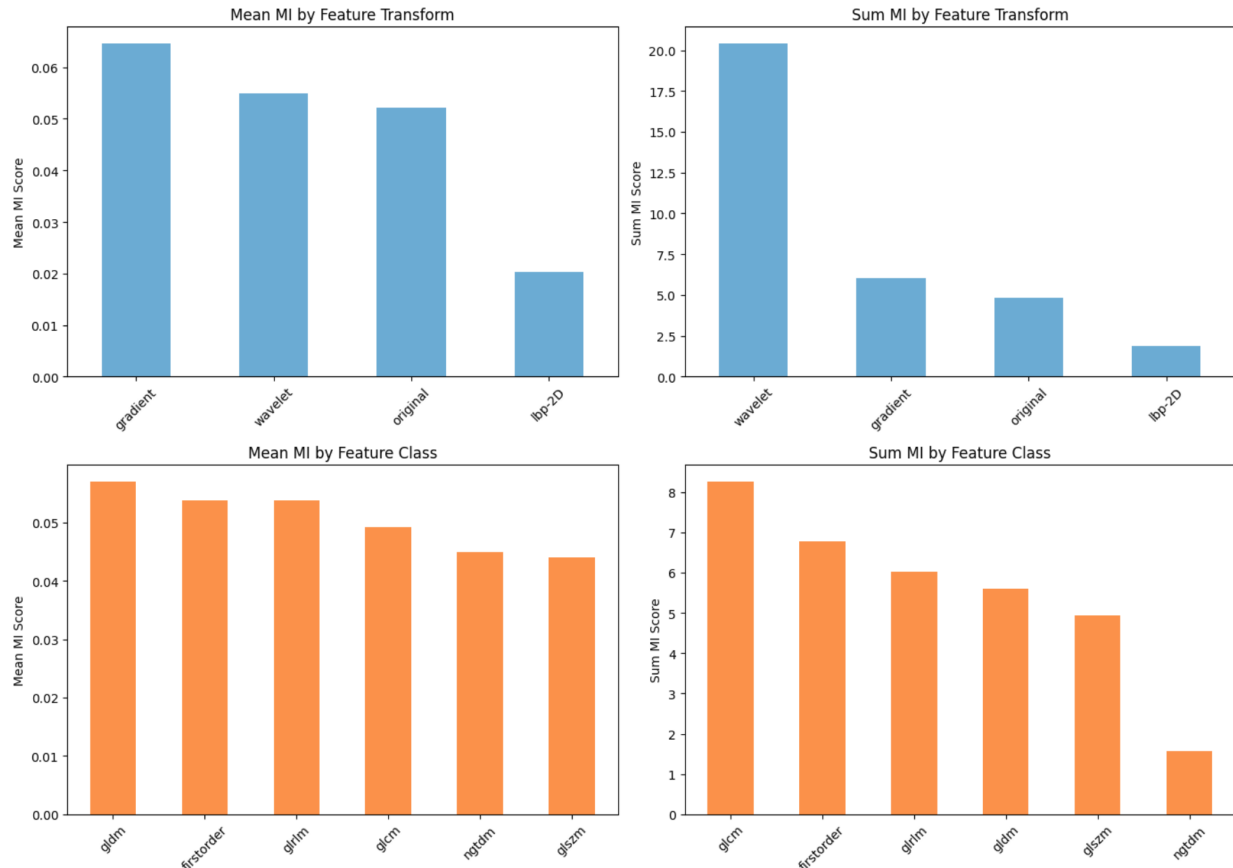
Radiomics feature classes, including First Order Statistics, Gray Level Dependence Matrix (GLDM), Gray Level Co-occurrence Matrix (GLCM), Gray Level Size Zone Matrix (GLSZM), Gray Level Run Length Matrix (GLRLM), and Neighboring Gray Tone Difference Matrix (NGTDM), play crucial roles in characterizing the complexity of lung textures in pneumonia. These features capture various aspects of pixel intensity distribution, texture patterns, and spatial relationships, collectively differentiating between normal tissue, non-Covid pneumonia, and COVID-19 pneumonia. Further details of these features and their relevance to pneumonia are presented in Table 3.

The combination of image transformations and feature classes yielded a total of 651 radiomics features initially extracted from a 10% subset of the training data. Analysis of the mean and sum of mutual information scores by feature transformation (Fig. 3) revealed several insights. Gradient and wavelet transformations generally enhanced mutual information compared to original images. Gradient-transformed features demonstrated the highest average mutual information, followed by wavelet-transformed features. Notably, while the wavelet transform generated many features (resulting in the highest sum of mutual information), linear binary patterns (LBP2D) displayed lower mean and sum mutual information, suggesting they may be less informative for this specific application. These findings informed our decision to prioritize gradient and wavelet transformations in hyperparameter tuning.

Analysis of mutual information scores also highlighted insights regarding feature classes (Fig. 3). The feature classes GLCM, GLDM, first-order statistics, and GLRLM emerged as the most promising based on their overall mutual

Feature Class	# Features	Description	Relevance to Pneumonia
First Order Statistics	19	Measures pixel intensity distribution (e.g., mean, variance).	Detects variations in lung tissue density, useful for distinguishing between normal and pathological states.
Gray Level Dependence Matrix (GLDM)	14	Captures dependencies of pixel pairs, considering distance.	Highlights textural changes in the lung indicative of pneumonia, such as areas of consolidation.
Gray Level Co-occurrence Matrix (GLCM)	24	Analyzes spatial relationships of pixel intensities (contrast, homogeneity).	Useful for differentiating types of pneumonia by assessing changes in lung texture homogeneity.
Gray Level Size Zone Matrix (GLSZM)	16	Quantifies homogeneous zones by size and intensity distribution.	Identifies regions of lung consolidation, contributing to distinguishing pneumonia in lung images.
Gray Level Run Length Matrix (GLRLM)	16	Measures consecutive pixels of the same intensity in various directions.	Effective in analyzing vascularity by identifying continuous lines (blood vessels). Lesions may disrupt these patterns, indicating pathological changes.
Neighboring Gray Tone Difference Matrix (NGTDM)	5	Analyzes intensity differences between a pixel and its neighborhood.	Highlights local heterogeneity in lung texture, useful for identifying early signs of pneumonia.

Table 3: Description of Radiomics Feature Classes


 Figure 3: Mean and Sum Mutual Information, by Feature Transformation and Feature Class
 (10% Subset of Training Data)

information scores. Consequently, subsequent feature engineering efforts focused primarily on features derived from gradient, wavelet, and original transformations in combination with the GLCM, GLDM, first-order statistics, and GLRLM feature classes.

3.2.2 Hyperparameter Tuning

To maximize the informativeness of our radiomics features and enhance their potential for accurate pneumonia classification, we employed a sequential hyperparameter tuning strategy. Mutual information (MI) gain with the target class labels served as our primary optimization metric. We adopted the Optuna framework to efficiently explore the parameter space.

Prioritizing computational efficiency, we tuned hyperparameters associated with image sampling, transformations, and feature classes sequentially. We began by optimizing resampling and binning settings, considering that regions of interest (ROI) of images had varying image resolutions. Subsequent tuning focused on gradient and wavelet transformations, inheriting the optimized sampling settings. Finally, we optimized hyperparameters for the GLRLM and GLDM feature classes, which also inherited settings from prior stages. Initial iterations suggested that tuning GLCM parameters was unlikely to substantially improve MI. Due to the computationally intensive nature of radiomics feature extraction, we performed tuning on a 1% subset of the original training data (76 Normal, 68 COVID-19, and 62 Non-COVID images).

Transformation / Class	Parameters	Initial Configuration	Search Values	Best Configuration
Sampling	do_resample	False	True, False	False
	binWidth	None	None, 5, 10, 15, 20	None
Wavelet Transformation	Wavelet Type	coif1	coif1, db1, sym2	coif1
	Start Level	0	0, 1	1
Gradient Transformation	Gradient Use Spacing	False	True, False	False
	GLRLM	None	Manhattan, Euclidian, None	Manhattan
GLDM	Distance	1	1, 2, 3	1
	Alpha	0	0, 1, 2, 3, 4, 5	1

Table 4: Radiomics Hyperparameter Tuning Results

Table 4 summarizes the hyperparameter tuning process and outcomes. Importantly, this optimization led to a 146% increase in mutual information within our validation dataset. Our analysis of the tuning process reveals that several key parameters remained optimal at their initial configuration, including sampling settings (no resampling or binning). This suggests the robustness of our chosen radiomics features to variations in ROI resolution. On the other hand, optimizing the wavelet transform and feature class parameters yielded the most significant MI gains. Interestingly, the optimal wavelet start level falls at the edge of our tested range, indicating the potential for further improvements by exploring even higher start levels in future investigations.

It is crucial to note that while maximizing the sum of MI led to aggregate feature set improvements, this approach has limitations. As each level of the wavelet transform generates a larger number of features, the sum of MI naturally increases. In future work, using mean MI or a composite metric might provide a more balanced assessment of feature informativeness, independent of sheer feature quantity.

3.2.3 Feature Selection and Final Extraction

Feature selection aimed to reduce redundancy and improve computational efficiency. Fig. 4a demonstrates the high correlation of the initial radiomics features. To address this, we applied a two-step selection process. First, we removed features with mutual information scores in the bottom 5th percentile, reducing the feature set from 864 to 820. Next, we iteratively selected the feature with the highest mutual information score and removed all other features with a correlation above 0.95. This process further reduced the feature set from 820 to 241 features, as visually demonstrated by the decrease in correlation in Fig. 4b.

Visual inspection of the top 50 radiomics features (ranked by mutual information) highlights the importance of employing multiple image transformations and feature classes (Fig. 5). Wavelet-transformed features dominate the top ranks, with various levels of decomposition represented (Fig. 5a), demonstrating their effectiveness in extracting valuable information for pneumonia classification. Furthermore, while first-order and GLCM features exhibit the

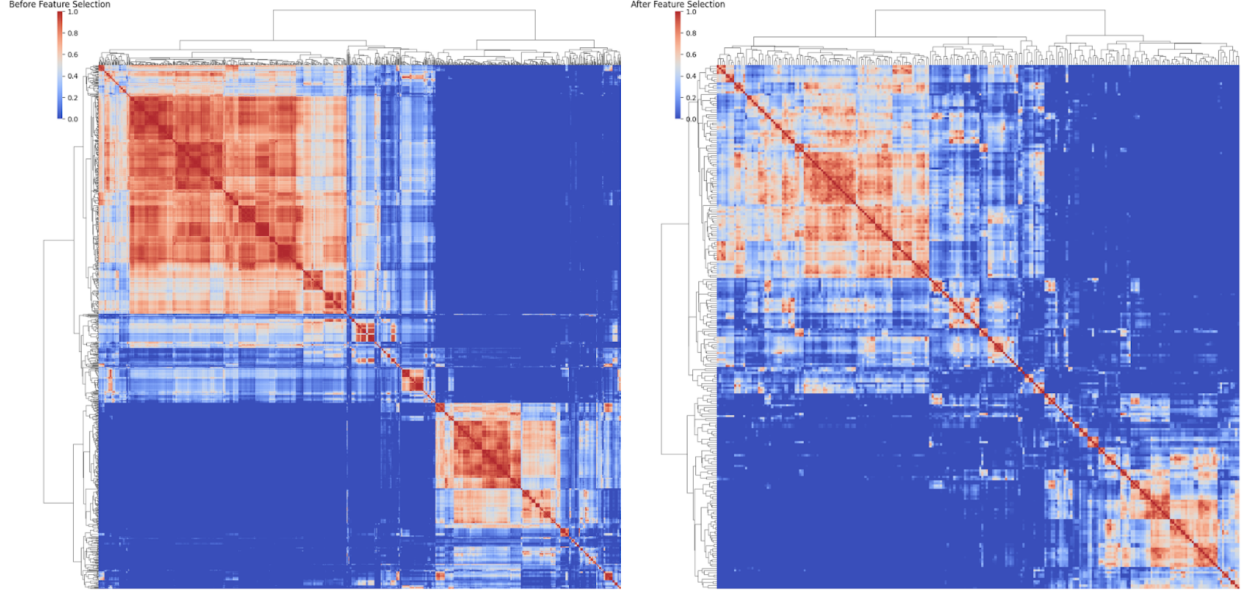


Figure 4: Radiomics Feature Correlations a) Before and b) After Feature Selection

highest mutual information, GLDM and GLRLM features are also well-represented (Fig. 5b). This diversity in feature classes ensures the radiomics features captures a broad range of image characteristics.

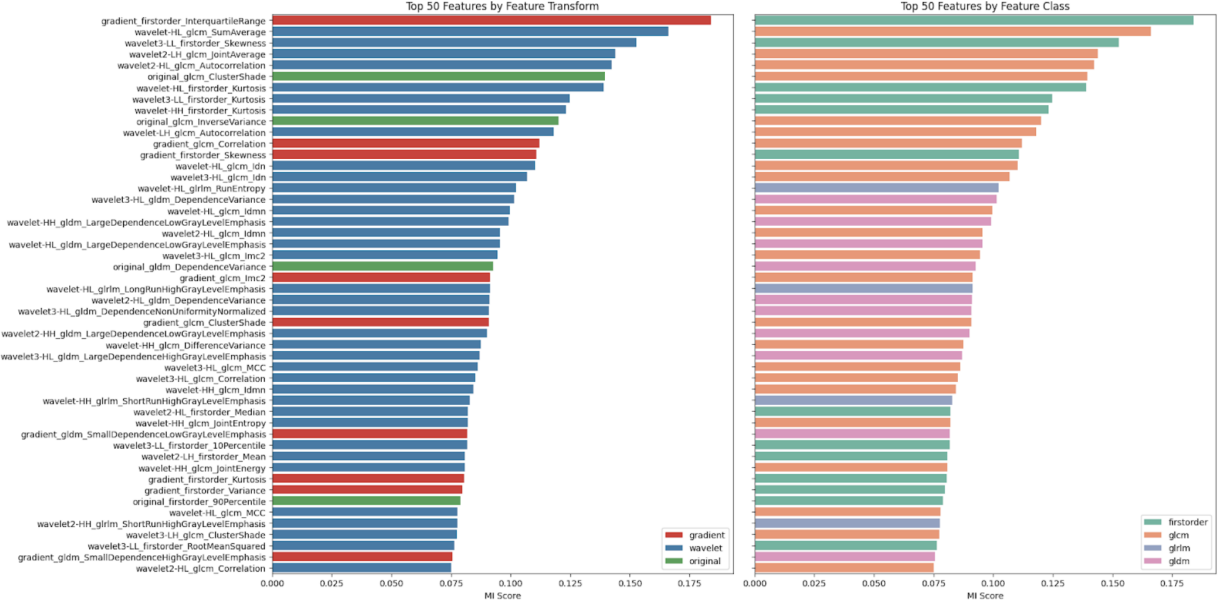


Figure 5: Top 50 Radiomics Features a) Image Transformation Applied and b) Feature Class

3.3 RESNET

Residual Network (ResNet), a convolutional neural network architecture, addresses the problem of vanishing gradients encountered in deeper networks [2]. Its core innovation, the residual block, introduces skip connections that allow for more efficient learning of deeper representations crucial for complex image analysis tasks. This makes ResNet well-suited for medical image applications.

The effectiveness of ResNet for pneumonia classification is supported by recent research. For instance, in a study on viral, bacterial, and normal case classification using chest X-ray images, ResNet50 achieved an accuracy of 93.01%, while ResNet-101 demonstrated an accuracy of 97.78% in distinguishing between COVID-19 and other viral pneumonia [4]. We chose ResNet50 over other variants for its balance of depth and computational efficiency.

To extract meaningful features from CXRs, we employed a pre-trained ResNet50 model. Images were resized to (224, 224), converted to grayscale, and normalized using standard ImageNet statistics. We opted for direct resizing to preserve detail around the periphery (since images were pre-cropped). The extracted embeddings were then flattened into a 1D vector, providing us with the raw ResNet50 embeddings.

3.4 T-distributed Stochastic Neighbor Embeddings

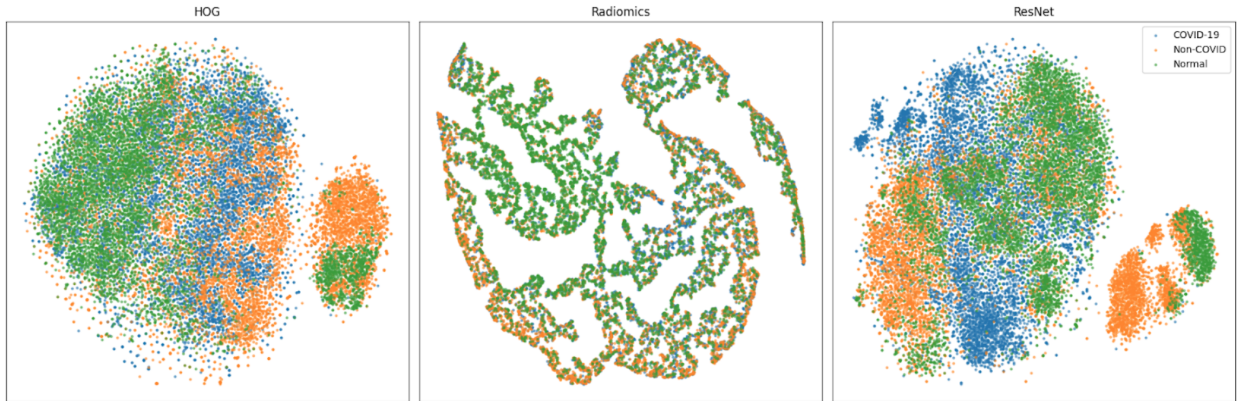


Figure 6: t-SNE Visualization of HOG, Radiomics and ResNet Features (Training Set)

T-distributed Stochastic Neighbor Embeddings (t-SNE) is a dimensionality reduction technique that projects how high-dimensional data points onto a lower-dimensional space (typically 2D or 3D) for easier visualization. In the context of machine learning and feature extraction, t-SNE helps us analyze how well the extracted features separate data points belonging to different classes.

As shown in Fig. 6, our t-SNE visualizations reveal that while perfect class separation is not achieved, certain methods demonstrate tendencies toward clustering. HOG and ResNet features exhibit some clustering behavior, suggesting that these feature extraction techniques might inherently capture characteristics that allow for partial differentiation between COVID-19, non-COVID, and normal cases. Conversely, the lack of clustering observed with the radiomics feature set could stem from the diverse nature of the extracted features. This highlights a limitation of t-SNE when visualization relationships between heterogenous features, where preserving global structure in a lower-dimensional space can be challenging, particularly if features are uncorrelated or orthogonal.

3.5 Principal Component Analysis

Principal Component Analysis (PCA) was employed to reduce the high dimensionality of our feature sets, specifically HOG (13,689 dimensions), radiomics (241 dimensions), and ResNet (2,048 dimensions). This technique mitigates the risk of overfitting during model training and improves computational efficiency by creating a more manageable feature space.

Fig. 7 illustrates the cumulative explained variance captured by PCA for each feature set as the number of principal components increases. As expected, the inherent complexity of HOG features is reflected in the plot, with a relatively high number of components (607 for 80% variance, 1,065 for 90%) required to capture most of the variation. Conversely, radiomics features exhibit a much steeper curve, requiring only 11 and 22 components to reach 80% and 90% explained variance, respectively. This suggests some redundancy within the radiomics features, indicating potential benefit from further feature selection techniques. ResNet features show a similar trend to HOG, also requiring a substantial number of components (84 for 80% variance, 193 for 90%) to capture important variations.

By applying PCA to each feature set and then concatenating the resulting principal components, we obtain a consolidated feature vector that is significantly lower in dimensionality (702 and 1280 features for 80% and 90% cumulative explained

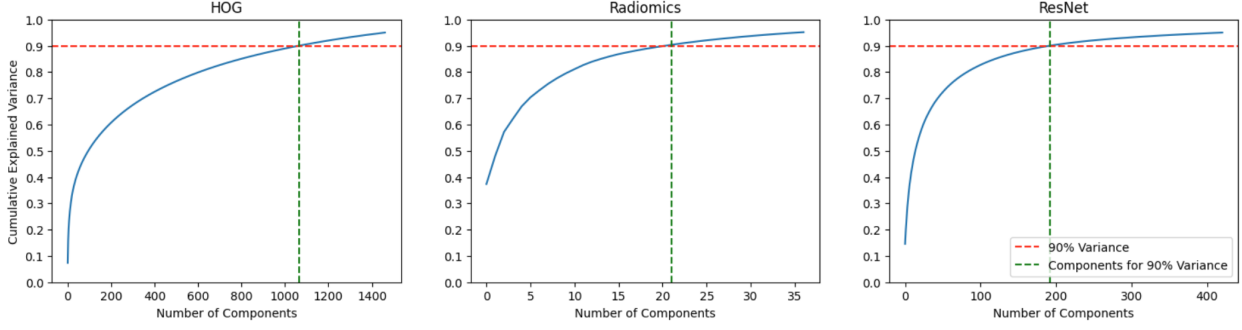


Figure 7: Cumulative Explained Variance vs. Number of PCA Components for HOG, Radiomics and ResNet Features (Training Set)

variance, respectively) while still retaining a high percentage of the original information. This transformation allowed us to train more complex models on this reduced feature space while minimizing overfitting issues.

4 Results

Four models are introduced for classification: Logistic Regression, a linear classifier known for its efficiency and interpretability; Support Vector Machine (SVM), which excels at finding optimal boundaries between classes even in high-dimensional spaces; Gradient Boosting, an ensemble method combining weak learners for improved accuracy; and Random Forest, another ensemble method that leverages multiple decision trees to mitigate overfitting and provide insights into feature importance.

4.1 Hyperparameter Optimization

As with feature engineering, we employed the Optuna framework to perform a Bayesian search over the hyperparameter space for each of the four models, aiming to identify the configurations that yielded the best performance metrics. We used accuracy as the objective function and conducted the hyperparameter search using the validation set, to avoid overfitting while maximizing generalizability. The parameters investigated, their search values, and the best parameter values for each model are shown in Table 5 below.

Model	Number of Trials	Search Time Per Trial (s)	Parameter	Search Values	Best Value
Logistic Regression	25	63.360	C	[0.01, 0.1, 1.0, 10]	0.1
			penalty	['l1', 'l2']	l1
SVM	25	63.080	C	[1e-4, 1e-3, 1e-2, 1e-1, 1, 1e1, 1e2]	1
			kernel	'linear', 'sigmoid', 'rbf'	rbf
Gradient Boost	15	2027.400	learning_rate	[0.01, 0.05, 0.1]	0.1
			n_estimators	[100, 200]	100
			subsample	[0.5, 1.0]	1.0
Random Forest	25	53.440	n_estimators	[50, 100, 200]	200
			max_depth	[5, 10]	10
			criterion	["gini", "entropy"]	"entropy"
			min_samples_split	[2, 5]	5

Table 5: Model Hyperparameter Tuning Results

4.2 Classification Performance

As shown in Fig. 6, our models demonstrated robust classification performance, with Logistic Regression and SVM achieving exceptional accuracy and F1-scores exceeding 0.89. This, combined with their balanced precision and

recall scores 10, highlights their potential for reliable pneumonia diagnosis in settings where minimizing both false positives and false negatives is critical. While all models performed well on the training set (accuracy and F1-scores above 0.85), a slight performance drop was observed on the validation and test sets (ranging from 0.81 to 0.88). This decrease, particularly for SVM and Random Forest, suggests a degree of overfitting, which will be further explored in the "Generalizability" section.

Model	Accuracy			F1-Score		
	Training Set	Validation Set	Test Set	Training Set	Validation Set	Test Set
Logistic Regression	0.91	0.88	0.90	0.91	0.88	0.90
SVM	0.97	0.87	0.89	0.97	0.87	0.89
Gradient Boost	0.86	0.85	0.85	0.86	0.85	0.85
Random Forest	0.92	0.81	0.82	0.92	0.81	0.82

Table 6: Tuned Model Accuracy and F1 Scores on Test Set (PCA: 90% Variance)

Confusion matrix analysis (Fig. 8) revealed a significant challenge: for logistic regression, 6.4% of COVID-19 and 6.7% of non-COVID pneumonia cases were misclassified as Normal. This highlights the inherent difficulty of accurately diagnosing pneumonia from chest X-rays, even for trained classifiers. Interestingly, further examination of misclassified images (Fig. 9) showed many cases where even untrained observers might struggle to discern the correct label. This aligns with clinical findings that chest X-ray interpretation can be subjective, with subtle or atypical presentations leading to diagnostic uncertainty [10].

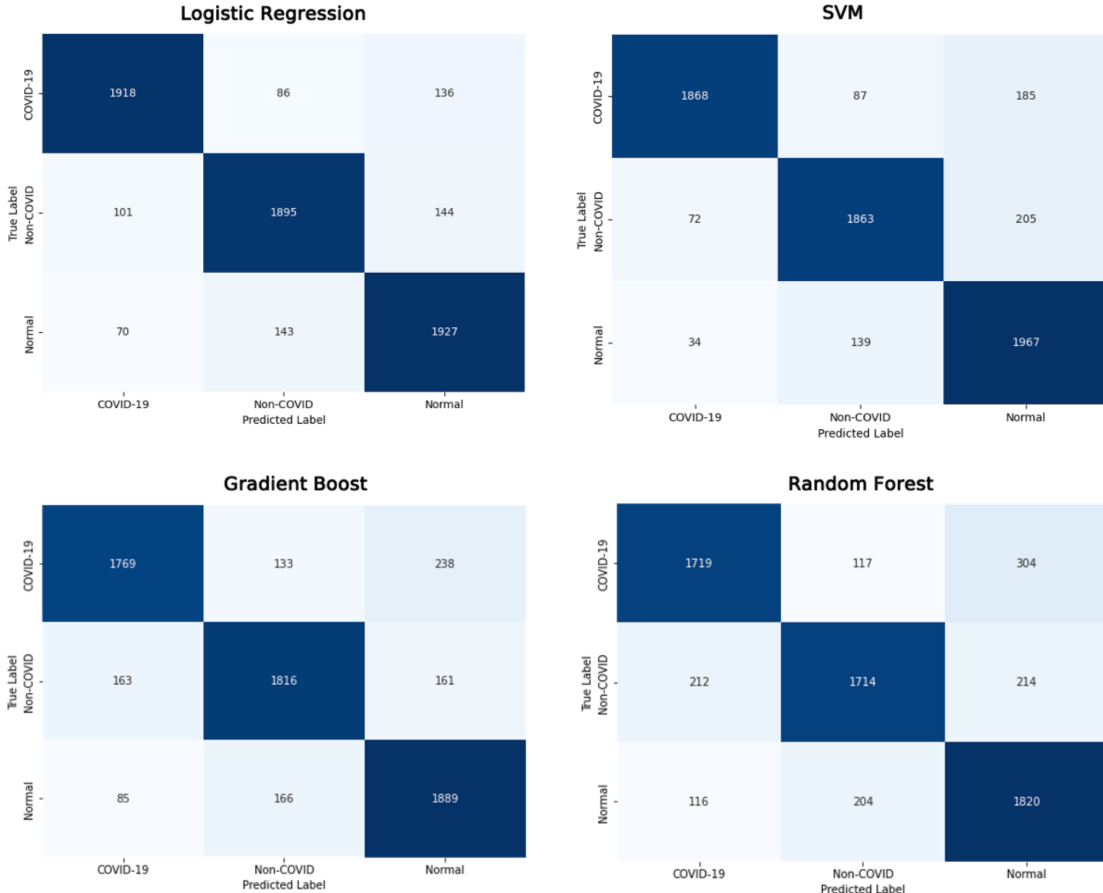


Figure 8: Confusion Matrices for a) Logistic Regression, b) SVM, c) Gradient Boost, and d) Random Forest on Test Set ((PCA: 90% Variance))

True Label	COVID-19	Normal	Non-COVID	Normal
Predicted Label	Normal	COVID-19	Normal	Non-COVID

Figure 9: Example Misclassified Images for Logistic Regression

To examine factors influencing our models’ decisions, we conducted Random Forest feature importance analysis to understand factors influencing our models’ decisions. Random Forest feature importance analysis (Table 7, Table 8) underscores the value of our multi-modal feature approach. All three feature classes (HOG, ResNet, and radiomics) contributed significantly to predictive power. While HOG features accounted for the largest overall importance (50.14%), ResNet (26.59%) and radiomics (23.27%) features demonstrated higher average importance per feature. This highlights the distinct and complementary insights captured by each modality. The prominence of HOG features highlights the importance of edge and gradient patterns for identifying consolidations and opacities, which are hallmarks of pneumonia in chest X-rays. Radiomics features complement this by quantifying nuanced textural variations which may be especially useful for distinguishing pneumonia types. ResNet likely extracts additional complex patterns, encompassing both shape and texture, that are not fully captured by HOG or radiomics alone. These insights, while derived from Random Forest, offer a valuable starting point for understanding the significance of our diverse feature set across all models.

	HOG	Radiomic	Resnet
Number of Features	1065	22	193
Sum of Feature Importance	50.14%	23.27%	26.59%
Average Importance per Feature	0.05%	1.06%	0.14%
Maximum Feature Importance	6.01%	2.91%	4.34%

Table 7: Summary of Random Forest Feature Importance

Rank of Feature	Feature Class	Feature Importance
1	HOG	6.0%
2	ResNet	4.3%
3	HOG	4.2%
4	ResNet	3.4%
5	Radiomics	2.9%
6	Radiomics	2.9%
7	Radiomics	2.2%
8	ResNet	2.2%
9	HOG	2.1%
10	HOG	2.1%

Table 8: Top 10 Important Random Forest Features

The models’ difficulty with cases where opacities have indistinct boundaries or atypical presentations, despite the presence of ResNet and radiomics features, highlights the complexity of pneumonia diagnosis. This suggests that even with multi-modal features, there may be subtle visual cues that our current models are not fully capturing. Investigating

the specific nature of these challenging cases could inspire the development of more refined features or the integration of additional modalities like clinical information. Exploring these avenues offers a promising direction for future research to enhance pneumonia classification models.

4.3 Generalizability

In image classification, a generalizable model can correctly identify diseases or objects even when those images were acquired using different equipment, within different healthcare settings, or contain variations not present in the training dataset. In our study, we evaluated generalizability primarily by assessing our models' performance on a held-out test dataset containing unseen images and measuring differences in performance across the training, validation, and test sets.

Our models demonstrated promising generalization capabilities, evidenced by their strong performance on the held-out test set. Logistic Regression and Gradient Boosting were particularly robust, maintaining stable accuracy across training, validation, and testing (with training-to-validation accuracy drops of 0.01). This suggests an ability to effectively classify new pneumonia cases, even with potential variations in image characteristics. While SVM and Random Forest exhibited greater differences between training and testing scores (with training-to-validation accuracy drops of 0.08 and 0.1, respectively) their overall performance on unseen data remains strong. These more significant drops suggest a greater degree of overfitting for SVM and Random Forest, highlighting a potential area for improvement.

Despite these promising results, there is room to further improve generalizability, particularly for mitigating the overfitting observed in SVM and Random Forest. To further improve these models, future studies should prioritize several strategies. Data augmentation, by introducing artificial variations that mimic real-world image differences, can expand the training dataset and reduce overfitting. Additionally, hyperparameter tuning could benefit from incorporating an objective function that explicitly balances accuracy with minimizing the difference between training and validation scores. Nonetheless, even with these refinements, systematic validation of the models' generalizability across diverse settings is crucial.

The dataset's origin from 10 different studies, encompassing images from various platforms and healthcare systems (Qatar, Europe, etc.), offers promising potential for generalizability across diverse settings. Manual investigation suggests representation of both genders. To validate generalizability and uncover potential biases, future studies should employ stratified cross-validation using metadata such as country, platform used, and patient demographics (gender, age, ethnicity). If, for instance, discrepancies arise between platforms, developing platform-specific models or integrating platform metadata into non-linear models could address such variations and improve performance.

4.4 Efficiency

Rapid and accurate diagnosis is essential in medical image analysis, especially for conditions like pneumonia where timely treatment influences outcomes. Efficiency is therefore crucial alongside accuracy for image classification models. This study, conducted in a standard Colab environment with CPU, highlights the importance of efficient models suitable for real-world deployment where computational resources might be limited.

Table 9 reveal significant insights into model efficiency across our dataset. Logistic Regression stands out, demonstrating the fastest tuning, training, and prediction times, while also achieving the highest accuracy and F1 scores. While Gradient Boost had considerably longer tuning and training times, its prediction speed was second only to Logistic Regression. Interestingly, PCA played a crucial role – without dimensionality reduction, Logistic Regression was unable to process the full feature set. It's worth noting that in the context of pneumonia detection, all prediction times across both tables fall within an acceptable range.

Model	Per Trial Tuning Time (s)	Training Time (s)	Prediction Time (s)			Accuracy	F1-Score
			Training Set	Validation Set	Test Set		
Logistic Regression	63.36	16.97	0.02	0.00	0.01	0.90	0.90
SVM	63.08	20.89	50.10	11.79	16.20	0.89	0.89
Gradient Boost	2027.44	2089.13	0.30	0.06	0.08	0.85	0.85
Random Forest	53.44	111.38	0.60	0.14	0.17	0.82	0.85

Table 9: Tuned Model Efficiency and Accuracy (PCA: 90% Variance).
Times shown are for 6,849 training images, 1,712 validation images, and 2,140 test images.

Table 9 shows the efficiency gains achieved through PCA. Moving from 90% to 80% explained variance resulted in faster training times for all models. Logistic Regression and Gradient Boosting exhibited the most significant reductions in training time, with Logistic Regression also demonstrating a slight improvement in prediction speed. SVM and Random Forest also benefited from PCA, with notable decreases in training and prediction times.

Interestingly, the impact of lowering PCA variance on accuracy varied. While Logistic Regression experienced a minor decrease (from 0.90 to 0.88), SVM and Gradient Boosting maintained their accuracy levels. Random Forest saw a slight increase in accuracy (0.82 to 0.83), likely due to reduced overfitting with the lower-dimensional feature set.

Model	Test Set Metrics	PCA Variance	
		90% (1280 Dimensions)	80% (702 Dimensions)
Logistic Regression	Training Time (s)	16.97	9.75
	Prediction Time (s)	0.01	0.01
	Accuracy	0.90	0.88
SVM	Training Time (s)	20.89	14.39
	Prediction Time (s)	16.20	10.45
	Accuracy	0.89	0.89
Gradient Boost	Training Time (s)	2089.13	1116.67
	Prediction Time (s)	0.08	0.05
	Accuracy	0.85	0.85
Random Forest	Training Time (s)	111.38	82.21
	Prediction Time (s)	0.17	0.15
	Accuracy	0.82	0.83

Despite efficiency gains from lowering PCA variance, Logistic Regression at 90% variance emerges as the optimal choice. Its fast prediction time (0.01s for 2,140 test images) makes the minor efficiency trade-off worthwhile. Given these promising results, future studies should explore Logistic Regression with even higher PCA variance thresholds to further optimize the balance between efficiency and accuracy.

5 Discussion and Conclusions

Our study demonstrates the potential of machine learning for accurate pneumonia classification using chest X-rays. Logistic Regression and SVM achieved particularly impressive results, with accuracies exceeding 0.89 across COVID-19, non-COVID pneumonia, and normal classes. However, the observed overfitting in SVM highlights the need to explore data augmentation and regularization techniques for improved generalizability if SVM were to be pursued.

Feature importance analysis underscores the value of our multi-modal approach. HOG effectively captured edge-based opacity patterns, while radiomics quantified subtle textural variations, and ResNet likely extracted more complex patterns. This synergy suggests that diverse feature sets are crucial for robust pneumonia detection.

For real-world applications, three key areas require further development. First, explainability is crucial for clinical adoption. Techniques like SHAP could provide insights into the features driving predictions, fostering trust in the model’s output. Second, instead of binary classifications, our models could provide probabilities or confidence levels for each class, empowering healthcare workers to make informed triage and referral decisions. Finally, despite encouraging initial findings regarding generalizability, rigorous validation across diverse settings remains crucial. Future studies should prioritize data augmentation, objective functions explicitly balancing accuracy and training-validation consistency, and stratified cross-validation using metadata (country, platform, demographics) to uncover biases. Addressing potential performance discrepancies across settings may necessitate platform-specific adaptations or metadata integration within the models.

This work advances the field of AI-assisted pneumonia diagnosis, demonstrating its potential for improving patient outcomes, particularly in resource-limited regions. By addressing explainability, providing probabilistic outputs, and validating generalizability, future research can pave the way for the integration of such models into real-world clinical practice.

6 Appendix

Model	Precision			Recall		
	Training Set	Validation Set	Test Set	Training Set	Validation Set	Test Set
Logistic Regression	0.91	0.88	0.90	0.91	0.88	0.90
Random Forest	0.92	0.81	0.83	0.92	0.81	0.82
SVM	0.97	0.87	0.89	0.97	0.87	0.89
Gradient Boost	0.87	0.85	0.86	0.86	0.85	0.85

Table 10: Tuned Model Precision and Recall on Test Set (PCA: 90% Variance)

Model (80% Variance)	Training Time (s)	Prediction Time (s)			Accuracy	F1-Score
		Training Set	Validation Set	Test Set		
Logistic Regression	9.75	0.01	0.00	0.01	0.88	0.88
SVM	14.39	29.53	7.98	10.45	0.89	0.89
Gradient Boost	1116.67	0.15	0.03	0.05	0.85	0.85
Random Forest	82.21	0.50	0.11	0.15	0.83	0.83

Table 11: Untuned Model Efficiency and Accuracy (PCA: 80% Variance).

Times shown are for 6,849 training images, 1,712 validation images, and 2,140 test images.

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