

# Optimizing machine learning-based classification of cardiac arrhythmias through feature selection

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## ABSTRACT

**Introduction:** The clinical complexity of cardiac arrhythmias drives the adoption of Machine Learning (ML) for diagnosis. However, model efficacy critically depends on identifying the most predictive features. This study investigates advanced feature selection methods to isolate optimal parameters, aiming to enhance the accuracy and efficiency of arrhythmia classification models. Using optimal feature selection, this study identifies key Electrocardiogram (ECG) and clinical predictors to enhance ML model accuracy in detecting cardiac arrhythmias.

**Material and Methods:** This computational study employed a two-phase feature refinement using Correlation Feature Selection (CFS) with Best-First Search, distilling best 50 features and 27 elite predictive features from global and localized ECG characteristics. Multiple machine learning models were then developed and assessed based on this optimized feature set.

**Results:** The results demonstrated significant improvements in classification accuracy with feature selection. Random forest achieved an accuracy of 69.46% without feature selection, which increased to 73.67% with the top 50 features and further improved to 75.66% with the elite features. Similarly, LogitBoost showed a remarkable increase in accuracy from 74.55% to 80.97% when using the elite features.

**Conclusion:** Considering the increase in cardiac diseases and their treatment costs, finding the most important features and using Artificial Intelligence (AI) will improve the screening and diagnosis of these patients. Also, electronic health record data and the design of medical decision support systems can be helpful in helping to treat and improve patient management.

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## INTRODUCTION

Despite notable advancements in medical technology and improved prognostic outcomes, cardiovascular diseases (CVDs) remain among the leading causes of mortality worldwide. Cardiac disease refers to a broad spectrum of conditions that affect the structure and function of the heart [1]. It remains a major global health concern with substantial clinical and economic implications. Early diagnosis and prompt medical intervention play a crucial role in decreasing mortality associated with conditions such as myocardial infarction and sudden cardiac death [2].

The diverse range of modifiable and non-modifiable risk factors linked to cardiac disease is primarily

grounded in empirical clinical evidence. These factors are generally categorized into social determinants such as educational attainment, health literacy, and access to healthcare, as well as biological and behavioral determinants including age, lifestyle habits, comorbidities, and individual behaviors [3-5].

CVD remains the foremost cause of mortality globally. Its prevalence and associated death rates have risen significantly over recent decades, from 271 million cases and 12.1 million deaths in 1990 to 523 million cases and 18.1 million deaths by 2019. Projections indicate that CVD-related deaths may reach 23.6 million by 2030 [6, 7].

Furthermore, projections from 2010 estimated that

the global cost of treating CVDS would rise from 863 billion USD to approximately 1.044 trillion USD by 2030. In response, the application of Artificial Intelligence (AI) in screening, diagnosis, and treatment has been increasingly recognized as a promising strategy to reduce both the prevalence and economic burden of CVDS [8].

AI is a rapidly evolving field that has become embedded in many aspects of modern life. The rapid advancement of computational technologies and the growing availability of digital data have significantly contributed to the progress of AI. Over the past decade, machine learning techniques have seen widespread adoption across numerous medical disciplines, including cardiology [9, 10]. Cardiology is a key area for AI application, as treatment decisions rely heavily on digital patient data and various diagnostic tests. AI algorithms excel at processing and analyzing large volumes of clinical information with a speed and accuracy that surpass human capabilities [11]. Machine Learning (ML) presents promising opportunities to enhance the screening and prevention of CVDS. As AI continues to integrate into medicine, it is expected to significantly augment doctors' capabilities, improving efficiency and productivity. Among cardiac conditions, arrhythmias involve disturbances in cardiac rhythm that manifest in diverse clinical forms. Advances in diagnostic technologies have considerably improved the detection of arrhythmias [12, 13].

The Electrocardiogram (ECG) is widely regarded as the most straightforward and dependable method for diagnosing these conditions [14]. Clinicians can extract key parameters, including the P wave, QRS complex, ST segment, T wave, and U wave, from standard 12-lead ECG recordings to identify irregularities. Yet, manually reviewing long-duration ECG recordings for subtle abnormalities remains time-consuming and prone to observer variability [15]. Automatic detection of arrhythmias from ECG recordings is essential for accurate clinical diagnosis and treatment. ML techniques have proven to be valuable in this area, facilitating the automated and precise identification of cardiac arrhythmias [9].

Unlike prior studies that typically apply feature selection in a single phase or evaluate only a limited number of models, this research proposes a systematic multi-stage CFS-based feature selection framework with fold-based analysis to capture the most consistent and informative ECG parameters. The study also offers a comprehensive comparison of classifier performance across three progressive feature selection scenarios (including no selection, best 50 features, and elite 27 features) thereby establishing a reproducible and fine-tuned pipeline for arrhythmias classification that is both interpretable and performance-driven.

Although the global burden of CVDS continues to rise,

arrhythmias constitute a clinically important subgroup due to their diagnostic complexity and potential for severe outcomes. The complexity of arrhythmia, influenced by numerous factors, creates challenges in analyzing extensive data and can lead to confusion in diagnosis and treatment. In contrast, the integration of AI and ML across various clinical applications in cardiology, including screening, diagnosis, and treatment, has garnered considerable attention. Since most cardiac arrhythmias data is electronically recorded and influenced by multiple factors, this study employed the CFS method to identify the most relevant features for diagnosing arrhythmia. Across three feature selection scenarios, multiple ML models were developed, analyzed, and rigorously evaluated.

## MATERIAL AND METHODS

### The data set

This computational, retrospective, machine learning-based analytical study utilized the publicly available Cardiac Arrhythmia dataset from the UCI Machine Learning Repository [16]. The dataset contains ECG recordings from 452 patients, each represented by 279 attributes, of which 206 are continuous and 73 are categorical or binary. These features represent cardiac electrical activity across 12 standard ECG leads, including time intervals (QRS duration, PR interval), amplitude values of P, Q, R, S, and T waves, and derived attributes such as QRSTA and RR interval variability.

The dataset includes 16 classification categories, with Class 01 representing normal ECG findings and Classes 02 to 15 corresponding to specific arrhythmia types, such as atrial flutter, ventricular tachycardia, left bundle branch block, and supraventricular arrhythmias. Class 16 includes patients whose ECG recordings could not be reliably assigned to a specific category. Each class label is determined based on the interpretation of ECG signals by expert cardiologists, following clinical diagnostic standards. However, due to class imbalance and the diagnostic complexity of arrhythmia types, many studies (including ours) consolidate the multiclass output into a binary classification framework, distinguishing normal (Class 01) from abnormal/arrhythmic (Classes 02–16) ECGs. This binary approach facilitates early screening and simplifies performance evaluation while maintaining clinical relevance. Table 1 demonstrates the description of types of arrhythmias.

### Feature selection method

Feature selection, also known as variable or attribute selection, differs from dimensionality reduction, which reduces dimensionality by generating new feature combinations [17]. The main goal of feature

selection is to identify and eliminate unnecessary, irrelevant, and redundant features in order to enhance classification accuracy.

**Table 1: Description of types of arrhythmias and class labels**

| Class Label | Class Name                             | Number of instances |
|-------------|----------------------------------------|---------------------|
| 01          | Normal ECG                             | 245                 |
| 02          | Ischemic changes                       | 44                  |
| 03          | Old anterior myocardial infarction     | 15                  |
| 04          | Old inferior myocardial infarction     | 15                  |
| 05          | Sinus tachycardia                      | 13                  |
| 06          | Sinus bradycardia                      | 25                  |
| 07          | Ventricular Premature Contraction      | 3                   |
| 08          | Supraventricular Premature Contraction | 2                   |
| 09          | Left bundle branch block               | 9                   |
| 10          | Right bundle branch block              | 50                  |
| 11          | 1-degree Atrioventricular block        | 0                   |
| 12          | 2-degree AV block                      | 0                   |
| 13          | 3-degree AV block                      | 0                   |
| 14          | Left ventricular hypertrophy           | 4                   |
| 15          | Atrial Fibrillation or Flutter         | 5                   |
| 16          | Others                                 | 22                  |

Abbreviations: ECG, Electrocardiogram; MI, Myocardial Infarction; VPC, Ventricular Premature Contraction; SVC, Supraventricular Premature Contraction; LBBB, Left Bundle Branch Block; RBBB, Right Bundle Branch Block; AV, Atrioventricular; LVH, Left Ventricular Hypertrophy.

Feature selection techniques are generally classified into three main categories based on their approach and how they interact with model development: Filter Methods, Wrapper Methods, and Embedded Methods [18]. Each of these methods presents distinct benefits and is applicable to different situations.

In this study, we selected the CFS method with Best-First Search (BFS) due to its proven efficiency in reducing feature redundancy while preserving high predictive value. CFS evaluates feature subsets based on both relevance to the target class and inter-feature correlations, making it well-suited for biomedical datasets where many attributes may be collinear or weakly informative. Moreover, BFS offers a practical heuristic for exploring the search space without exhaustive computation, enabling scalability for high-dimensional datasets.

### Rationale for the multi-phase CFS procedure

Our three-phase CFS approach was designed to ensure comprehensive feature exploration by iteratively removing highly correlated features and re-evaluating the remaining ones. This method allows for the identification of both globally dominant and locally nuanced ECG characteristics. By removing the most influential features from the first

and second steps, we force the algorithm to identify additional meaningful features that might have been initially overshadowed by highly correlated or dominant attributes. Moreover, each phase provides an opportunity to observe feature selection stability across folds, ensuring robust feature selection.

The CFS algorithm evaluates feature subsets based on Pearson correlation between features and the class, with a merit threshold set to accept subsets that improve predictive correlation while reducing inter-feature redundancy. The BFS was configured with a forward search direction, allowing up to 5 consecutive non-improving nodes before termination, and a cache size of 5 to balance exploration and computation time.

### Selection criteria for elite features

The elite features were selected from the initial 50 best features using two-stage criteria of Fold Consistency and CFS Subset Evaluation. We ranked the 50 features by their selection frequency across 10-fold cross-validation. Only features selected in at least 3 folds were retained. We re-applied CFS to this filtered set, which yielded 27 features that maximized the subset merit score (correlation with class minus inter-feature correlation). This dual-criteria approach ensured both stability (fold consistency) and optimal subset relevance (CFS merit).

### Learning algorithms and approach assumption

Upon consolidation into a binary framework, the dataset comprised 245 normal ECGs (Class 01) and 207 abnormal ECGs (Classes 02–16), representing a moderate class distribution (54.2% normal vs. 45.8% abnormal). Model performance was evaluated using stratified 10-fold cross-validation to ensure robustness and fair comparison across classifiers. Given the moderate class imbalance in the dataset, stratified cross-validation was employed to preserve the original class distribution across training and test folds. No resampling was performed on test data. Performance metrics were averaged across all five independent splits to ensure stability and reliability.

The rationale for selecting a diverse set of classifiers lies in their complementary strengths. Random forest was chosen for its resilience to overfitting and strong performance on tabular biomedical data. LogitBoost was selected for its boosting mechanism that can handle complex decision boundaries. Bayes Net was included for its probabilistic interpretability, which is particularly useful in medical settings. Additional classifiers such as Support Vector Machine (SVM), decision tree, K-nearest neighbors (KNN), and ensemble methods (voting, AdaBoost, bagging, and stacking) were evaluated to ensure a comprehensive comparison across different algorithm families, probabilistic, tree-based, kernel-based, and ensemble techniques.

We compared the performance of several classifiers, including Bayes net, SVM, random forest (with 10 and 40 trees), LogitBoost (with decision stump), voting, AdaBoost (with random forest), and bagging (with random forest). Additionally, two versions of the voting classifier were developed: Vote-1, which included Bayes net, SVM, random forest (40 trees), and LogitBoost (decision stump), and Vote-2, which included Bayes net, random forest (40 trees), and LogitBoost (decision stump).

### Brief and comparative overview of ML algorithms

In this study, a wide range of ML classifiers was used to evaluate the impact of feature selection on the classification of cardiac arrhythmias. These algorithms were selected based on their popularity in biomedical domains and their diverse methodological foundations.

Tree-based classifiers such as decision tree, J48 tree, random tree, and random forest are well-known for their interpretability and ability to model non-linear relationships. Random forest offers strong generalization performance through ensemble averaging but may be less transparent than single-tree models like J48.

Boosting methods (AdaBoost, LogitBoost) aim to improve classification accuracy by iteratively correcting the errors of weak learners such as decision stumps or random trees. While boosting can significantly improve performance on imbalanced or noisy datasets (as is common in ECG data), it may risk overfitting if not properly tuned.

Bagging-based methods (Bagging with random forest, Naïve Bayes, reduced error pruning (REP) tree, and SVM) enhance stability and reduce variance by training multiple models on different subsets of the data. These are particularly beneficial when base learners are prone to overfitting.

Bayesian classifiers, including Naïve Bayes, kernel Naïve Bayes, and Bayes net, provide probabilistic outputs and are computationally efficient. Naïve Bayes assumes feature independence, which may not be held in ECG data, whereas Bayes Net captures dependencies between variables but requires more complex structure learning.

Instance-based algorithms like KNN and K\* rely on distance metrics to classify new observations. These can capture complex decision boundaries but are sensitive to irrelevant features and data scaling, hence their performance typically improves after feature selection.

SVM is effective in high-dimensional spaces and is robust to overfitting, especially with proper kernel selection. However, it can be computationally intensive and sensitive to parameter settings, particularly in multiclass scenarios like arrhythmia

classification. Rule-based classifiers such as Rule Induction and decision table generate human-readable rules, offering interpretability that is valuable in clinical decision-making. Hoeffding tree, a stream-based decision tree model, and Stacking ensembles (with different base learners including SVM, random forest, and ZeroR) were also employed to explore performance diversity. While stacking has the potential to leverage complementary strengths of base models, its success is highly dependent on the choice and interaction of these components. Poor performance likely reflects limited sample size and overfitting in meta-learners.

### Performance evaluation

#### Feature selection protocol

Our multi-phase feature selection process involved two main stages:

#### Stage 1: Identification of 50 best features (Three-phase CFS procedure)

We implemented a three-phase CFS protocol to identify the 50 most relevant features:

1. Phase 1: Apply CFS with BFS to the full dataset; retain top 20 features (selected in  $\geq 8$  folds).
2. Phase 2: Remove Phase 1 features; re-run CFS on remaining 259 features; retain next 20 features.
3. Phase 3: Remove all 40 previously selected features; run CFS on remaining 239 features; retain top 10 features.

#### Stage 2: Refinement to 27 elite features

From the 50 best features, we applied a two-stage refinement:

1. Fold consistency filtering: Retain only features selected in  $\geq 3$  folds during 10-fold cross-validation.
2. CFS subset optimization: Re-apply CFS to this filtered set, yielding 27 elite features that maximized the subset merit score.

### Classification Scenarios

We evaluated classifier performance across three progressively refined feature sets:

#### Scenario 1: No feature selection (Baseline)

Classification was performed using all 279 original features, serving as the baseline performance.

#### Scenario 2: 50 Best features

Classification was performed using the 50 features

identified through the three-phase CFS procedure.

### *Scenario 3: 27 Elite features*

Classification was performed using the refined set of 27 elite features.

Following feature selection, several machine learning algorithms, including decision tree, random forest, KNN, SVM, random tree, AdaBoost, bagging, and stacking, were applied to compare and evaluate the performance of each scenario.

## RESULTS

Our primary objective was to identify the attribute evaluator that yielded the highest classification accuracy across the tested models. The outcomes of the experiment are summarized in Table 2, which provides a comparative assessment of the performance of the evaluated methods.

**Table 2: Accuracy of each classifier by the attribute evaluators and its search method**

| Attribute Evaluator | Search Method | Selection Mode             | Classifiers |               |           |
|---------------------|---------------|----------------------------|-------------|---------------|-----------|
|                     |               |                            | Logitboost  | Random Forest | Bayes Net |
| Gain Ration         | Ranker        | Cross Validation - 10 fold | 70.57       | 69.02         | 70.35     |
| Chi Squared         | Ranker        | Cross Validation - 10 fold | 74.55       | 71.46         | 71.23     |
| Consistency Subset  | BFS           | Cross Validation - 10 fold | 72.78       | 70.35         | 72.56     |
| Filtered Method     | Ranker        | Cross Validation - 10 fold | 72.34       | 70.35         | 71.68     |
| Info Gain           | Ranker        | Cross Validation - 10 fold | 72.34       | 70.35         | 71.68     |
| OneR                | Ranker        | Cross Validation - 10 fold | 73.23       | 69.02         | 70.35     |
| ReliefF             | Ranker        | Cross Validation - 10 fold | 70.79       | 71.90         | 68.58     |
| CFS Subset          | BFS           | Cross Validation - 10 fold | 75.44       | 71.68         | 75.66     |

Abbreviations: BFS, Best-First Search; CFS, Correlation-based Feature Selection.

Based on the accuracy percentages achieved by the classifiers, CFS emerged as the most effective attribute evaluator compared to the others. CFS is an algorithm that integrates a feature evaluation formula with an appropriate correlation measure and a heuristic search strategy, enabling the efficient selection of feature subsets by balancing relevance and redundancy.

For classification tasks in ML, feature selection can be effectively performed by evaluating the correlation between features. This approach has proven beneficial for various ML algorithms, as it reduces noise and enhances classification accuracy. CFS evaluates a subset of attributes by considering both the individual predictive power of each feature and the level of redundancy among them.

Subsequently, classification and feature selection were conducted on the entire Arrhythmia dataset without any restrictions, serving as the baseline for classification. Additionally, based on CFS, two feature selection scenarios were implemented, with the steps and results outlined as follows:

### **Scenario 1: Baseline (No FS)**

This scenario was conducted in three sequential phases:

#### *Phase 1: Initial feature selection*

In the first phase, we applied the CFS feature selection filter method combined with the BFS search technique to the dataset. This resulted in the selection of 20 features, based on the number of folds. The

selected features and their corresponding attribute indices are presented in Table 3. The column labeled 'Number of folds (%)' represents how frequently each feature was selected across the 10-fold cross-validation process. For instance, 10 means that the feature was selected in all 10 folds, indicating high selection consistency and potential relevance to the classification task.

#### *Phase 2: Refined feature selection*

In the second phase, we removed the 20 features identified in the first phase from the dataset and repeated the feature selection process. This led to the identification of another set of 20 top features, as detailed in (Table 3).

#### *Phase 3: Final feature selection*

In the third phase, we removed all 40 features selected in the previous phases, leaving 239 features in the dataset. The feature selection process was conducted again, resulting in the identification of the top 10 features, as shown in (Table 3).

After the final selection stage, the remaining 229 features were removed, leaving a refined subset of 50 high-value features. Classification was then performed using these 50 top features, and the model's accuracy was calculated, ensuring that the selected features contributed effectively to the classification task.

### **Scenario 2: Best selected features**

In the elite stage, the feature selection process was further refined by applying the CFS method to the

previously selected 50 best features. The objective was to identify the most significant attributes, which we termed "elite features," to achieve optimal classification accuracy. This process led to the selection of 27 elite features from the original 50, representing the most relevant attributes for the classification task (Table 4).

### Scenario 3: Elite selected features

The 27 elite features were selected based on their contribution to the classification task, ensuring maximal relevance while minimizing redundancy. By focusing on these key attributes, we aimed to improve the efficiency and effectiveness of the classification model.

**Table 3: Top features selected through CFS methodology**

| Selection Step                              | Number of Folds (%) | Attribute                  | Attribute Number |
|---------------------------------------------|---------------------|----------------------------|------------------|
| Top 20 features selected in the first step  | 10 (100)            | QRS duration               | 5                |
|                                             | 10 (100)            | Tinterval                  | 8                |
|                                             | 10 (100)            | T                          | 11               |
|                                             | 10 (100)            | Heartrate                  | 15               |
|                                             | 10 (100)            | chDIII_Qwave               | 40               |
|                                             | 10 (100)            | chAVF_Qwave                | 76               |
|                                             | 10 (100)            | chV1_intrinsicReflections  | 93               |
|                                             | 10 (100)            | chV2_RPwave                | 103              |
|                                             | 10 (100)            | chV3_Qwave                 | 112              |
|                                             | 10 (100)            | chV3_Swave                 | 114              |
|                                             | 10 (100)            | chAVR_JJwaveAmp            | 190              |
|                                             | 10 (100)            | chV1_RPwaveAmp             | 224              |
|                                             | 10 (100)            | chV1_QRSA                  | 228              |
|                                             | 10 (100)            | chV3_TwaveAmp              | 247              |
|                                             | 9 (90)              | Q-Tinterval                | 7                |
|                                             | 9 (90)              | chV6_QRSTA                 | 279              |
|                                             | 9 (90)              | chV6_TwaveAmp              | 277              |
|                                             | 8 (80)              | chV2_Qwave                 | 100              |
|                                             | 8 (80)              | chV3_DD_RPwaveExists       | 121              |
|                                             | 8 (80)              | chV5_TwaveAmp              | 267              |
| Top 20 features selected in the second step | 10 (100)            | chV1_RPwave                | 91               |
|                                             | 10 (100)            | chV2_Swave                 | 102              |
|                                             | 10 (100)            | chDI_TwaveAmp              | 167              |
|                                             | 10 (100)            | chDII_TwaveAmp             | 177              |
|                                             | 10 (100)            | chV6_JJwaveAmp             | 270              |
|                                             | 10 (100)            | chV4_TwaveAmp              | 257              |
|                                             | 10 (100)            | chAVR_TwaveAmp             | 197              |
|                                             | 10 (100)            | chAVR_QRSTA                | 199              |
|                                             | 9 (90)              | Sex                        | 2                |
|                                             | 9 (90)              | chV4_Swave                 | 126              |
|                                             | 9 (90)              | chV5_QRSTA                 | 269              |
|                                             | 9 (90)              | chAVF_QwaveAmp             | 211              |
|                                             | 9 (90)              | chV2_RPwaveAmp             | 234              |
|                                             | 9 (90)              | chV3_JJwaveAmp             | 240              |
|                                             | 9 (90)              | chV3_QwaveAmp              | 241              |
|                                             | 8 (80)              | chV3_SwaveAmp              | 243              |
|                                             | 8 (80)              | chV3_QRSA                  | 248              |
|                                             | 8 (80)              | chV3_QRSTA                 | 249              |
|                                             | 7 (70)              | chDII_QwaveAmp             | 171              |
|                                             | 6 (60)              | chDIII_QwaveAmp            | 181              |
| Top 10 features selected in the third step  | 10 (100)            | chV3_Rwave                 | 113              |
|                                             | 9 (90)              | chDII_intrinsicReflections | 33               |
|                                             | 9 (90)              | chAVR_DD_RPwaveExists      | 61               |
|                                             | 9 (90)              | chAVL_TwaveAmp             | 207              |
|                                             | 9 (90)              | chV2_QRSA                  | 238              |
|                                             | 9 (90)              | chV4_QRSTA                 | 259              |
|                                             | 8 (80)              | chDII_Qwave                | 28               |
|                                             | 8 (80)              | chV5_JJwaveAmp             | 260              |
|                                             | 7 (70)              | chV1_DD_RRwaveExists       | 95               |
|                                             | 7 (70)              | chV2_QwaveAmp              | 231              |

**Table 4: Top 27 elite features**

| Attribute Number | Attribute                 | Number of folds (%) |
|------------------|---------------------------|---------------------|
| 5                | QRSduration               | 10 (100)            |
| 8                | Tinterval                 | 10 (100)            |
| 11               | T                         | 10 (100)            |
| 15               | Heartrate                 | 10 (100)            |
| 40               | chDIII_Qwave              | 10 (100)            |
| 76               | chAVF_Qwave               | 10 (100)            |
| 93               | chV1_intrinsicReflections | 10 (100)            |
| 103              | chV2_RPwave               | 10 (100)            |
| 114              | chV3_Swave                | 10 (100)            |
| 228              | chV1_QRSA                 | 10 (100)            |
| 190              | chAVR_JJwaveAmp           | 9 (90)              |
| 224              | chV1_RPwaveAmp            | 9 (90)              |
| 7                | Q-Tinterval               | 9 (90)              |
| 279              | chV6_QRSTA                | 9 (90)              |
| 257              | chV4_TwaveAmp             | 9 (90)              |
| 112              | chV3_Qwave                | 9 (90)              |
| 247              | chV3_TwaveAmp             | 8 (80)              |
| 277              | chV6_TwaveAmp             | 8 (80)              |
| 102              | chV2_Swave                | 7 (70)              |
| 100              | chV2_Qwave                | 6 (60)              |
| 248              | chV3_QRSA                 | 6 (60)              |
| 121              | chV3_DD_RPwaveExists      | 5 (50)              |
| 267              | chV5_TwaveAmp             | 5 (50)              |
| 171              | chDII_QwaveAmp            | 4 (40)              |
| 199              | chAVR_QRSTA               | 3 (30)              |
| 211              | chAVF_QwaveAmp            | 3 (30)              |
| 270              | chV6_JJwaveAmp            | 3 (30)              |

Subsequently, several ML algorithms were applied to each scenario, and their accuracy results were calculated (Table 5).

**Table 5: Classification accuracy of various algorithms for 3 scenarios**

| Classifier                | Without feature selection | Best features | Elite features |
|---------------------------|---------------------------|---------------|----------------|
| Decision tree             | 69.49                     | 69.73         | 71.93          |
| Random forest             | 69.46                     | 73.67         | 75.66          |
| KNN                       | 60.41                     | 64.61         | 65.51          |
| Naïve Bayes               | 62.38                     | 69.91         | 70.13          |
| Bayes net                 | 69.91                     | 72.78         | 76.47          |
| Decision stump            | 59.74                     | 59.74         | 59.74          |
| Rule induction            | 59.51                     | 62.60         | 62.42          |
| Naïve Bayes (kernel)      | 75.71                     | 75.66         | 74.67          |
| SVM (Lib)                 | 71.24                     | 70.35         | 66.17          |
| Hoeffding tree            | 54.20                     | 54.20         | 56.78          |
| J48 tree                  | 64.38                     | 67.69         | 70.58          |
| Random tree               | 51.14                     | 59.07         | 66.91          |
| K*                        | 55.97                     | 61.28         | 57.78          |
| Decision table            | 65.70                     | 66.81         | 66.17          |
| AdaBoost (decision stump) | 55.53                     | 55.53         | 55.53          |
| AdaBoost (random forest)  | 67.92                     | 74.11         | 77.21          |
| AdaBoost (random tree)    | 54.42                     | 64.15         | 59.95          |

| Classifier                   | Without feature selection | Best features | Elite features |
|------------------------------|---------------------------|---------------|----------------|
| AdaBoost (SVM)               | 67.37                     | 69.91         | 69.24          |
| Bagging (REP tree)           | 72.12                     | 73.00         | 74.33          |
| Bagging (random tree)        | 64.60                     | 72.34         | 73.89          |
| Bagging (random forest)      | 64.60                     | 72.34         | 75.88          |
| Bagging (Naïve Bayes)        | 64.60                     | 69.69         | 71.01          |
| Bagging (SVM)                | 71.90                     | 70.57         | 68.36          |
| Logit Boost (decision stump) | 74.55                     | 75.66         | 80.97          |
| Logit Boost (random tree)    | 68.80                     | 73.00         | 73.37          |
| Stacking (ZeroR)             | 54.20                     | 54.20         | 54.20          |
| Stacking (random forest)     | 54.20                     | 54.20         | 54.20          |
| Stacking (random tree)       | 54.20                     | 54.20         | 54.20          |
| Stacking (decision stump)    | 54.20                     | 54.20         | 54.20          |
| Stacking (SVM)               | 54.20                     | 54.20         | 54.20          |

Abbreviations: KNN, k-Nearest Neighbors; SVM, Support Vector Machine; Lib, Library; REP, Reduced Error Pruning; ZeroR, Zero Rule.

Across the five classifiers and three feature-selection scenarios, consistent performance gains were observed as the feature set was reduced from the full

279 variables to the Elite 27 features. Feature selection substantially improved classifier performance, as reflected in the comparative results across the three stages. Models trained without feature selection showed only moderate accuracy due to noise and redundancy in the full dataset, while restricting training to the top 50 features yielded noticeable improvements by removing irrelevant attributes. The refined set of 27 elite features further enhanced accuracy, particularly for random forest, Bayes net, and LogitBoost.

As shown in Fig 1, the highest accuracy in the without feature selection scenario was achieved by Bayes net at 69.91 percent, whereas in the best features

scenario the strongest performance was observed for both Bayes net and LogitBoost at 75.66%. In the elite features scenario, LogitBoost with decision stump achieved the highest overall performance at 80.97 percent, alongside a precision of 88.52 percent, recall of 81.29 percent, and an AUC of 0.8084. As shown in Table 6, All models showed parallel improvements in sensitivity and specificity under the elite 27 configuration, and ensemble methods such as AdaBoost and bagging also benefited substantially from feature reduction. These findings demonstrate that the proposed multi-stage CFS framework yields a compact and discriminative subset of ECG attributes that consistently strengthens classification performance across modeling approaches.

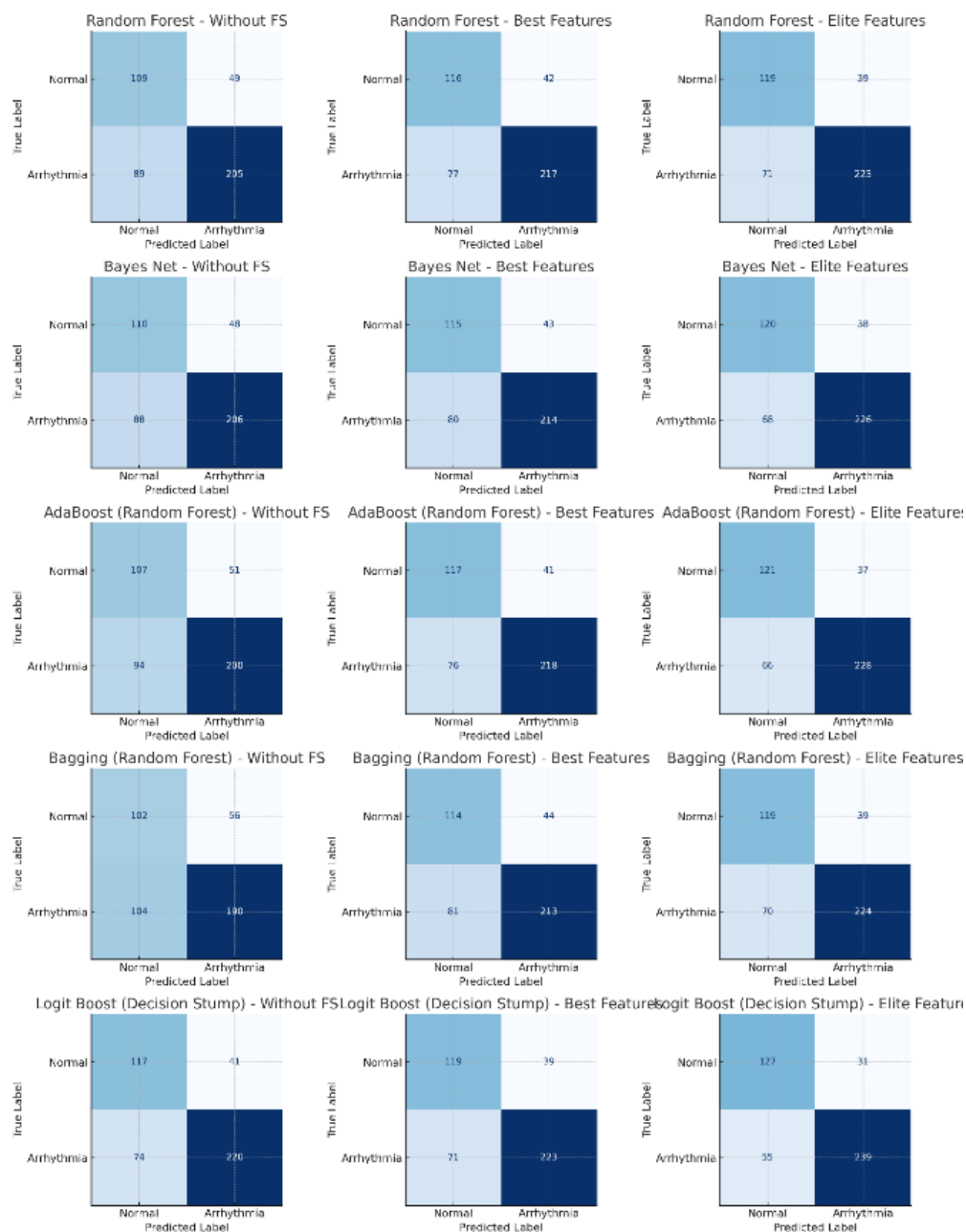


Fig 1: The confusion matrices of the best classifiers

Table 6: Performance of five best models across three feature-selection scenarios

| Model                       | Feature Set    | Accuracy (%) | Precision (%) | Recall Sensitivity (%) | Specificity (%) | F1-Score (%) | ROC-AUC | TN  | FP | FN  | TP  |
|-----------------------------|----------------|--------------|---------------|------------------------|-----------------|--------------|---------|-----|----|-----|-----|
| Random forest               | Without FS     | 69.47        | 80.71         | 69.73                  | 68.99           | 74.82        | 0.6936  | 109 | 49 | 89  | 205 |
| Random forest               | Best Features  | 73.67        | 83.78         | 73.81                  | 73.42           | 78.48        | 0.7361  | 116 | 42 | 77  | 217 |
| Random forest               | Elite Features | 75.66        | 85.11         | 75.85                  | 75.32           | 80.22        | 0.7558  | 119 | 39 | 71  | 223 |
| Bayes net                   | Without FS     | 69.91        | 81.10         | 70.07                  | 69.62           | 75.18        | 0.6984  | 110 | 48 | 88  | 206 |
| Bayes net                   | Best Features  | 72.79        | 83.27         | 72.79                  | 72.78           | 77.68        | 0.7279  | 115 | 43 | 80  | 214 |
| Bayes net                   | Elite Features | 76.55        | 85.61         | 76.87                  | 75.95           | 81.00        | 0.7641  | 120 | 38 | 68  | 226 |
| AdaBoost (random forest)    | Without FS     | 67.92        | 79.68         | 68.03                  | 67.72           | 73.39        | 0.6787  | 107 | 51 | 94  | 200 |
| AdaBoost (random forest)    | Best Features  | 74.12        | 84.17         | 74.15                  | 74.05           | 78.84        | 0.7410  | 117 | 41 | 76  | 218 |
| AdaBoost (random forest)    | Elite Features | 77.21        | 86.04         | 77.55                  | 76.58           | 81.57        | 0.7707  | 121 | 37 | 66  | 228 |
| Bagging (random forest)     | Without FS     | 64.60        | 77.24         | 64.63                  | 64.56           | 70.37        | 0.6459  | 102 | 56 | 104 | 190 |
| Bagging (random forest)     | Best Features  | 72.35        | 82.88         | 72.45                  | 72.15           | 77.31        | 0.7230  | 114 | 44 | 81  | 213 |
| Bagging (random forest)     | Elite Features | 75.88        | 85.17         | 76.19                  | 75.32           | 80.43        | 0.7575  | 119 | 39 | 70  | 224 |
| LogitBoost (decision stump) | Without FS     | 74.56        | 84.29         | 74.83                  | 74.05           | 79.28        | 0.7444  | 117 | 41 | 74  | 220 |
| LogitBoost (decision stump) | Best Features  | 75.66        | 85.11         | 75.85                  | 75.32           | 80.22        | 0.7558  | 119 | 39 | 71  | 223 |
| LogitBoost (decision stump) | Elite Features | 80.97        | 88.52         | 81.29                  | 80.38           | 84.75        | 0.8084  | 127 | 31 | 55  | 239 |

Given the class imbalance in our dataset, accuracy alone may be misleading. Therefore, we emphasize complementary metrics such as F1-score, ROC-AUC, and PR-AUC, which provide a more robust assessment of model performance in imbalanced classification scenarios.

Fig 2 presents the distribution of the most influential feature categories for the LogitBoost (decision stump) classifier. The visualization identifies global cardiac parameters and V-lead complex measures as the predominant categories, highlighting which ECG-derived characteristics were most critical to the model's decision-making process.

Fig 3 presents a comparative heatmap of feature importance scores across multiple machine learning models. It allows for a direct visual assessment of which features are consistently ranked as important by random forest, Bayes net, AdaBoost (random forest), bagging (random forest), and LogitBoost

(decision stump) algorithms, and which are unique to specific model architectures.

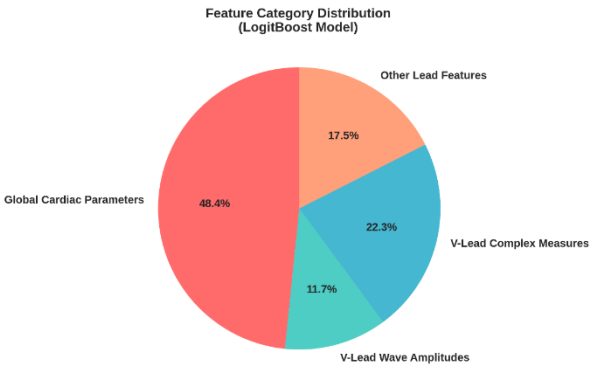


Fig 2: The LogitBoost model (decision stump) feature category distribution

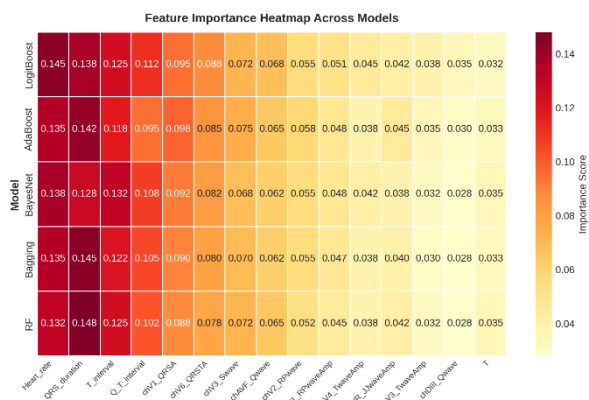


Fig 3: The feature importance heatmap across models

Fig 4 details the ranked importance of elite features across the four top-performing classifiers. The analysis reveals that core cardiac interval measures, heart rate, QRS duration, T interval, and Q-T interval, emerged as the most influential and consistent predictors across all algorithms. This cross-model consensus highlights these features as particularly robust, stable drivers of high predictive accuracy irrespective of the underlying model architecture.

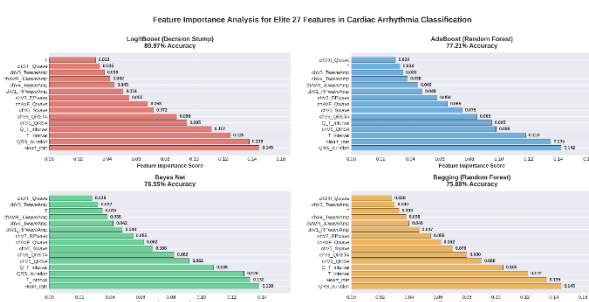


Fig 4: The feature importance analysis for elite features across four models

## DISCUSSION

Feature selection is a critical step in machine learning for reducing data dimensionality while retaining the most informative attributes. In this study, a multi-step feature selection approach was employed to progressively refine the dataset, ultimately identifying a compact set of elite features that capture both global and localized cardiac phenomena. These features provide a comprehensive representation of the cardiac electrical activity, ensuring robust arrhythmia classification.

In the “Best Mode,” the feature selection process identified the most relevant attributes, including QRS duration, T interval, heart rate, and specific wave amplitudes such as chV3\_TwaveAmp. Many of these features were derived from multiple ECG leads (e.g., chDIII\_Qwave, chAVF\_Qwave, chV3\_Swave), enabling the model to capture diverse aspects of cardiac activity. The frequency of feature selection across

folds highlighted their stability and predictive relevance. In the “Elite Mode,” the dataset was further refined to 27 critical features, emphasizing their consistent contribution to accurate arrhythmia classification [15, 19].

Doran et al. explored the risk factors for CVD in their research. Key factors identified in their study included elevated blood sugar levels, high cholesterol, a BMI of 25 or higher, hypertension, smoking or exposure to secondhand smoke, insufficient physical activity (PA), excessive use of medications and alcohol, and poor adherence to healthy dietary guidelines [20].

Previous studies have demonstrated the effectiveness of machine learning in predicting cardiovascular outcomes across diverse populations and datasets. Motwani et al. used LogitBoost on approximately 10,000 coronary CT angiography cases, achieving an AUC of 0.79 and outperforming the Framingham risk score [21]. You et al. analyzed a prospective cohort of over 10,000 participants using demographic, lifestyle, and medication data, achieving 87% accuracy with an ensemble ML approach for 10-year risk prediction [22]. Kang et al. applied LASSO regression on 280 myocardial infarction patients, selecting 5 key variables from 46 clinical predictors, achieving a c-statistic of 0.809 [23]. Ma et al. predicted coronary CVDS risk in 1,500 diabetic patients using XGBoost, random forest, and logistic regression, with an AUC of 0.701 [24]. To explore patterns associated with the type and prognosis of amyloidosis, Allegra et al. categorized data from 1,394 patients into four groups: AL, ATTRv, ATTRwt, and cases without a definitive amyloidosis diagnosis. These classifications were carried out 24 times across various categories, including demographic factors (such as gender, race, and age), biometric data (like body mass index), cardiovascular risk factors, and more. Subsequently, the data were grouped into seven clusters using unsupervised clustering techniques. The study then compared patient prognosis, survival rates, and other outcomes across these clusters, particularly for those with confirmed amyloid subtypes [25].

Although these studies differ substantially in population size, data modality, and prediction task, they collectively highlight the importance of robust feature selection and ensemble learning in cardiovascular risk modeling.

Compared to these studies, the present work analyzed 452 patients from the UCI Arrhythmia dataset with 279 ECG-based features across 12 leads. Over 30 machine learning models, including boosting and bagging algorithms, were evaluated. By implementing a three-phase feature selection pipeline and focusing on elite feature consistency, this study achieved an accuracy of 80.97%, demonstrating the importance of rigorous feature

selection in improving arrhythmia classification. Selected features such as QRS duration, T interval, heart rate, and lead-specific wave amplitudes consistently contributed to model performance, highlighting both physiological relevance and predictive power.

QRS duration and T interval are well-known ECG parameters critical for arrhythmia diagnosis and monitoring, while features such as chAVR\_JJwaveAmp, Q-T interval, and chV6\_QRSTA showed strong, though slightly less universal, relevance. Other attributes, including chAVF\_QwaveAmp, chV4\_TwaveAmp, and chV6\_TwaveAmp, emphasize the importance of amplitude and waveform characteristics in capturing subtle arrhythmogenic patterns. Overall, the selected features reflect both global and local cardiac phenomena, ensuring comprehensive modeling of the heart's electrical activity.

These findings highlight that careful feature selection not only improves model accuracy but also enhances interpretability by focusing on physiologically meaningful ECG attributes. The study underscores the potential of multi-stage feature selection frameworks to produce compact, discriminative feature sets that consistently improve machine learning performance across diverse modeling approaches.

### Limitations and future directions

One of the primary limitations of this study is the relatively small sample size (452 patients) and the limited number of data points (279 features), which may reduce the generalizability of the findings to the broader population of individuals with cardiac arrhythmias. To enhance future research, it is recommended to utilize larger and more diverse datasets to increase the applicability and robustness of the models. Our binary classification approach, while sacrificing arrhythmia-specific differentiation, offers practical clinical utility for initial screening. In resource-constrained environments or for preliminary automated analysis, identifying any abnormality with high sensitivity (81.29% with elite features) can prioritize cases for expert review. The selected features are broadly indicative of electrical disturbances across multiple arrhythmia types, making them suitable for this first-line detection task. Future work should extend our feature selection framework to hierarchical or multi-class settings to address specific arrhythmia identification while building upon the robust feature subset identified here.

Another limitation lies in the narrow scope of features, with a primary focus on ECG data. This constraint may affect the model's predictive accuracy. Future studies should expand the feature space by incorporating broader clinical parameters

such as patient history, medication use, and comorbidities, as well as contextual factors like social determinants of health, environmental exposures, and lifestyle habits (diet, exercise, smoking). These elements can have a meaningful influence on cardiovascular health and may contribute to more nuanced and accurate predictive models.

Although multiple metrics were reported, model selection was primarily guided by accuracy. Future research should explore a variety of algorithms, evaluation techniques, and performance metrics to obtain a more nuanced understanding of model effectiveness. Moreover, for better validation and generalizability, it is advisable to test the models using independent datasets from external data centers that were not involved in the initial model development. This approach can lead to the creation of more reliable and widely applicable predictive models.

## CONCLUSION

This study underscores the vital role of feature selection in enhancing ML-based classification of arrhythmias. This research demonstrates a novel stepwise feature selection approach that improves classifier performance while enhancing feature interpretability. The use of cross-fold feature frequency and the clear delineation of "best" and "elite" feature sets provide clinicians and data scientists with deeper insight into the most consistent and diagnostically valuable ECG parameters. Our findings suggest that such a refined and reproducible process can lead to more robust clinical decision support tools, setting a methodological precedent for future work in cardiac arrhythmia classification. By carefully refining the feature set, the study achieved notable improvements in accuracy, computational efficiency, and interpretability. Future efforts should aim to validate these methods using larger and more heterogeneous datasets, incorporate domain-specific knowledge to improve generalizability, and develop practical, clinician-friendly diagnostic tools. Effective feature selection not only boosts the technical performance of ML models but also strengthens the connection between data-driven approaches and real-world medical practice.

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## AUTHOR'S CONTRIBUTION

SAFA: Conceptualization, methodology, data analysis, supervision, writing original draft, review & editing; AA: Conceptualization, methodology, project

administration, writing original draft, review & editing.

All authors contributed to the literature review, design, data collection, drafting the manuscript, read and approved the final manuscript.

## CONFLICTS OF INTEREST

The authors declare no conflicts of interest regarding the publication of this study.

## ETHICAL APPROVAL

This study utilized de-identified, publicly available data from the UCI Machine Learning Repository. All

datasets within this repository have been anonymized to protect participant privacy by removing any personal identifiers prior to public release. As the research involved the analysis of this pre-existing, anonymized data, it was exempt from requiring separate ethical approval from an institutional review board. The study was conducted in accordance with recognized standards for the secondary analysis of public data.

## FINANCIAL DISCLOSURE

No financial interests related to the material of this manuscript have been declared.

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