Remote ECG Monitoring Kit to Predict Patient-Specific Heart Abnormalities
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ABSTRACT
Electrocardiogram (ECG) signals are widely used to examine heart rhythms and general health conditions. However, the majority of commercial ECG kits are generic and their normal ranges are set based on the averages obtained from a large population of people with normal heart conditions. This averaging ignores the extreme inherent variability of normal heart signals. As such, many false alarms are generated if the thresholds are selected too strict and true alarms are missed if the thresholds are set too loose. Furthermore, false alarms may arise due to the high physical activity of the test person. In this paper, we developed a prototype for patient-specific heart monitoring kit, which learns the properties of a patient’s normal ECG signal over time and reports significant deviations from this normal behavior, in addition to presenting significant violations from the global norms. Further, false alarms due to high physical activity levels are eliminated through processing the utilized accelerometer signal. This personalized remote heart monitoring kit with the proposed signal processing and self-tuning capabilities and wireless connectivity provides more detailed information and insightful interpretations of ECG signals compared to generic devices, therefore can be used for remote heart monitoring of high-risk people.

Keywords: Personalized Diagnosis, Wearable Devices, ECG Signal Analysis, Predictive Modeling, Remote Heart Monitoring.

1. INTRODUCTION
Healthcare service in today’s society has been significantly improved by using different technological advances such as fully automated diagnosis tools, data-driven predictive modeling and remote monitoring systems. In particular, cardiovascular disease, one of the top ranked killers in the world over the past decades, has been heavily investigated from different perspectives by the biomedical research community [1-3].

A majority of this high mortality rate is due to late diagnosis and delayed therapeutic interventions [4]. Therefore, a constant heart condition monitoring and timely prediction of potential heart abnormalities can have a significant impact on saving lives, especially for high-risk patients.

In order to diagnose heart abnormalities, Electrocardiogram (ECG) is widely used in healthcare industry, since it can be implemented using low-cost and affordable circuitry with a relatively high accuracy. Electrocardiograms involve the measurement of electrical activities of different parts of a hearts [3, 5, 6].

Many practical implementations of ECG-based heart monitoring kits in terms of wearable devices (e.g. fitbit [7]), mobile apps (e.g. Alivecor’s heart monitor app [8]) as well as more sophisticated commercial cardiac monitoring equipment are developed to assess heart conditions based on ECG signals. However, a majority of these devices suffer from the following drawbacks. Firstly, these devices lack advanced signal processing and information extraction capabilities; hence, they require results interpretation by an expert. Secondly, the predefined normal ranges for signal parameters (e.g. heart rate) are set based on a typical ECG signal for a normal heart. However, there exists extreme inherent variations among normal heart signals collected from people, influenced by many factors such as age, gender, race, genetic patterns, as well as environmental conditions (e.g. temperature, elevation from sea level) [9,10]. Therefore, a generic equipment with a pre-defined set of thresholds is not well suited for precise monitoring of different patients’ heart functionality. For tightly selected thresholds, the device generates many false alarms. On the other hand, the device may miss significant true alarms for loosely selected thresholds. Thirdly, physical activity of a person may impose misleading changes on the ECG signal morphology [11], which should not be confused with the actual hearth abnormalities. This fact is overlooked in the current off the shelf kits.

In this project, we developed a prototype of a personalized remote heart monitoring kit that probes a patient’s heart functionality as well as his physical activity and transmits the collected information to a remote processing unit through wireless communication for further analysis and interpretation.

The design of this system allows a more flexible and continuous cardiac monitoring. The developed learning algorithm builds a patient-specific model to analyze and interpret the ECG signal by detecting deviations from the patient’s normal ECG trends. The results of this analysis in terms of minor and major alarms are displayed on the kit’s display module in order to assist the patient to take proper actions accordingly.

Further, the device is equipped with an accelerometer in addition to ECG measurement probes to incorporate the patient’s physical activity into the learning algorithm and avoid false alarm generation due to the patient’s physical activities. The results provided in section 4, shows that the developed kit outperforms generic methods based on a global classifier by providing more detailed information about heart functionality and predicting potential upcoming heart abnormalities before their occurrence. This device can be used to remotely monitor heart behavior and general health conditions of high-risk people (such as seniors and patients with heart attack history) and assist patients to take necessary therapeutic actions such as taking rest, calling their doctors and taking blood thinning medications.

The following of this paper is organized as follows. An overview of the entire system is presented in Section 2 with elaborating on the details of the designed wearable ECG monitoring module. The proposed signal processing and prediction processing is presented in Section 3. The results are presented in Section 4, followed by concluding remarks in Section 5.
2. SYSTEM MODEL

The proposed patient-specific remote heart monitoring kit comprises two modules: a Wearable Sensing Module (WSM) and a computer-based Personalized Processing Unit (PPU), as shown in Fig. 1a. The two modules communicate using Wi-Fi technology with our proprietary half-duplex overlay communication protocol. The WSM module collects analog ECG signal and 3-dimensional accelerometer signal and transmits them to PPU for further analysis. PPU processes the received signals and classifies the received signals into normal and abnormal classes using a global classifier trained using a public ECG dataset. The normal signal segments are used by an online predictive learning algorithm for constant tuning of the patient-specific model parameters. Simultaneously, significant deviations from the normal trend are recognized to trigger a minor alarm. The results of processing are sent back to WSM via a feedback channel to be displayed in terms of user-friendly text messages. Fig. 1b shows the block-diagram of the designed prototype. The following sections elaborate on the details of the kit and the developed software.

Wireless Sensing Module

WSM is designed based on Arduino Uno microcontroller and includes a single-lead ECG sensing module with three electrodes, an accelerometer and a display module. The ECG electrodes are attached to the left arm, the right arm, and the lower abdominal cavity of the patient. The kit is attached to the arm to maximize sensitivity to physical activities. The analog signals from the two sensors are sampled evenly with sampling rate of 215 samples/sec. WSM receives the feedback messages and displays respective user-friendly text messages on a build-in LCD. Minor and major alarms are shown by two yellow and red LEDs. The displayed messages include the result of signal processing, the patient’s physical activity level, the functionality of the sensors, and the communication channel status indicators.

Communication Module

The sensing module constantly collects ECG and 3-dimensional accelerometer signals and bundles them into transmit packets. Each packet includes a start flag for flow control, a checksum code for error detection, and the measurement samples, as depicted in Fig. 2. The PPU analyzes the received hyperpacket including 1498 information packets and two control packets and sends back the result of the analysis in a feedback packet, which includes start and end flags, user ID, a set of message codes as shown in Fig. 2.

3. SIGNAL PROCESSING AND PREDICTIVE MODELING

The proposed Software is developed in MATLAB Environment with Graphic User Interface (GUI) to facilitate easy data entry (e.g. personal and clinical information), signal processing and result display. In addition to the received signal, a set of representative features as well as the result interpretations are presented. The core part of the software is the proposed learning-based prediction algorithm, which includes the following stages:

1) Data preprocessing and de-noising
2) Segmentation
3) Feature extraction
4) Global classifier
5) Personalized local deviation analysis

De-noising and Baseline removal

Here, we use the popular method of Wavelet decomposition for denoising purpose. We use the 8-level Daubechies wavelet transform (db8), which has been proven to be efficient [12]. Since we use the public MIT-BIH arrhythmia database with sampling frequency of 360Hz [13] to train the global classifier, we use the relation described in Eq. (1) in order to identify the levels of wavelet transform. The high –frequency details are considered noisy and subject to discard according to the recommendations of the Association for the Advancement of Medical Instrumentation (AAMI) [14].

\[ L = \left[1 + \log_2 \frac{F_s}{360}\right] \]  

(1)

Here, \([x]\) is the floor operator and represents the maximum integer number lower than \(x\). To de-noise the ECG signal with sampling rate of \(F_s = 215 \text{ samples/sec}\), the high frequency details at the first layer is discarded. The baseline wander is another common artifact in ECG signal. In this study, a five ordered fitted polynomial is subtracted from the ECG signal to eliminate the wandering trend. Finally, the resulting signal is resampled to 360Hz to match the training dataset.
Segmentation
We first identify the cardiac cycles using wavelet transform. A typical cardiac circle includes five fiducial peaks, namely P, Q, R, S and T. The QRS complex, which composed of Q, R and S peaks, is the most representative wave of one cycle. The detection of other peaks often depends on the location of QRS complexes. Both normal beats and abnormal beats exhibit QRS complexes in the frequency range of 5 to 22Hz [1]. With a sampling frequency of 360Hz, the detail coefficients of level 4 and level 5 covers the information of QRS complexes. The algorithm described in [13] and [15] uses Maximal Overlap Discrete Wavelet Transform (MODWT) to detect R peaks. We use db4 as basis function instead of sym4, since it exhibits more similarity with a typical QRS complex [1, 12]. The rest of fiducial peaks are then detected within a certain segment around R peaks; resulting in the P, QRS onset, Q, S, QRS offset and T waves within each cardiac circle [1].

If \( p_t \) is the location of R peak for the \( i \)th cycle, the range of this cycle is defined as \( \left[ p_t + \frac{i - 1}{3}, p_t + \frac{i + 1}{3} \right] \), noting that R peak approximately splits a cycle of length \( r \) into two parts of length \( r/3 \) and \( 2r/3 \) [9]. Further, we combine \( m \) consecutive cycles to obtain one analysis segment, in order to eliminate transient artifacts and obtain local averages of signal parameters. In the subsequent learning algorithm, each segment considered as a data-point to be classified. The next segment is obtained by sliding the segment \( k \) cycles forward, where \( 1 \leq k \leq m \). Fig. 3 shows the segmentation concept for \( k = 1 \) and \( m = 3 \). Accelerometer signals are also segmented accordingly.

Feature Extraction
In order to develop an accurate learning algorithm and avoid the well-known overfitting issue by using all time-points, here we extract a set of informative features that capture the main properties of the signals as follows.

For the ECG signal, the morphological details can be used to distinguish between the normal and abnormal cardiac cycles. In Fig. 4, the ventricular cardiac cycle can be easily recognized as it has a larger maximum negative peak and longer R-R duration compared to the adjacent normal cardiac cycle.

As depicted in Table 1, we extract three categories of features including temporal features, morphological features and frequency domain features to make a comprehensive list of features that are reported in the literatures [1, 3, 5, 12]. For each category, we include two sets of features. Set 1 includes 8 features that are calculated per cycle for all \( m \) cycles and then we include their mean and coefficient of variations as representative features of the morphological properties of the signal. Set 2 contains 6 features, which includes general properties of the signal and are calculated once per segment. Therefore, we have a total of \( 2 \times 8 + 6 = 22 \) features per segment. The 22 dimensional feature space is mapped into 8 dimensional space by Principal Component Analysis (PCA). Finally, these 8 features are normalized to yield zero mean unit variance features.
Table 1: Features extracted from ECG signal.

<table>
<thead>
<tr>
<th>Feature Type</th>
<th>SET 1</th>
<th>SET 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Temporal Features</td>
<td>1) QRS duration, 2) QT duration, 3) PR duration</td>
<td>1) segment mean ( RR ), 2) segment mean(( R_1 - R_{avg} ))</td>
</tr>
<tr>
<td>Morphological Features</td>
<td>3) max positive peak to second peak ratio</td>
<td>3) signal average energy, 4) max positive peak, 5) max negative peak, 6) peak to energy ratio</td>
</tr>
<tr>
<td>Freq- Domain Features</td>
<td>5-8) signal power level at 7.5Hz, 10Hz, 12.5Hz, 15Hz</td>
<td></td>
</tr>
</tbody>
</table>

For the accelerometer signal, we extract representative features for each segment to build a binary classifier that distinguishes between the active and test modes as presented in Table 2 following suggestions provided in [16]. For the sake of simplicity, we first combine the three axes of data using the following equation:

\[
Ac(i) = \sqrt{AcX(i)^2 + AcY(i)^2 + AcZ(i)^2} \tag{2}
\]

If the reading of the accelerometer is zero across all axes, then an accelerometer signal loss alarm is displayed.

**Proposed Personalized Predictive Modeling**

In this section, we elaborate on the proposed personalized predictive modeling, which includes a global classifier and a local deviation analysis module under a hierarchical structure as depicted in Fig. 5. In order to verify the global classifier of the proposed algorithm, we apply it to well-annotated MIT-BIH database, which includes two subsets DS1 and DS2, each of which contains ECG signals for 22 patients [14, 17]. Here, we use DS1 as training dataset and DS2 as test dataset. Each ECG signal includes cardiac cycles, which are annotated by experts and mapped into four classes including Normal, Supraventricular, Ventricular and Fusion [5]. The number of classes in the two database subsets DS1 and DS2 is presented in Table 3.

We first label the ECG segments. A segment is labeled as Normal, if all \( m = 3 \) member cycles are labeled as \( N \), otherwise it is labeled as the only abnormal class (\( V, S \) or \( F \)) within the segment. If a segment includes two or more different abnormal classes, the segment is discarded.

Table 2: Features extracted from accelerometer signal.

<table>
<thead>
<tr>
<th>Metrics Name</th>
<th>Formula</th>
</tr>
</thead>
<tbody>
<tr>
<td>Differences</td>
<td>[ \max[Ac(i)] - \min[Ac(i)] ]</td>
</tr>
<tr>
<td>Stand Deviation</td>
<td>[ \frac{1}{N} \sum_{i=1}^{N} (Ac(i) - \frac{1}{N} \sum_{j=1}^{N} Ac(j))^2 ]</td>
</tr>
<tr>
<td>Minimum</td>
<td>[ \min[Ac(i)] ]</td>
</tr>
<tr>
<td>Range</td>
<td>[ \max[Ac(i)] - \min[Ac(j)] ]</td>
</tr>
<tr>
<td>Energy</td>
<td>[ \frac{\sum_{i=1}^{N}</td>
</tr>
</tbody>
</table>

Table 3: The utilized public dataset overview

<table>
<thead>
<tr>
<th>Dataset</th>
<th># of N segments</th>
<th># of V segments</th>
<th># of S segments</th>
<th># of F segments</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td>DS1</td>
<td>12633</td>
<td>2053</td>
<td>550</td>
<td>121</td>
<td>15357</td>
</tr>
<tr>
<td>DS2</td>
<td>11721</td>
<td>2356</td>
<td>862</td>
<td>256</td>
<td>15195</td>
</tr>
<tr>
<td>Total</td>
<td>24354</td>
<td>4409</td>
<td>1412</td>
<td>377</td>
<td>30552</td>
</tr>
</tbody>
</table>

The following are the details of the global classifier and local deviation analysis module.

**Global Classifier**

We first build a global classifier using all segments of the signals for all data-points in database DS1. Note that the data after feature extraction and preprocessing is not guaranteed to be independently drawn and identically distributed (iid), a set of candidate classification algorithm are tested using data from DS1. Finally, we chose k-Nearest Neighbors method for this stage and Weighted k-Nearest Neighbor for the following stage with \( k = 10 \) to build the classifier, since they demonstrate the best performance as shown in Table 4. However, the method is general and not sensitive to the choice of classifiers.

During the test phase, we process the received signal segments and classify them to one of the four major classes \( N, V, S, \) or \( F \). The segment at time \( t \) is denoted by \( x_t \) and the predicted label by the global classifier is denoted by \( y_t \). A segment which is classified as abnormal \((y_t \in \{V, S, F\})\) triggers a red alarm in the system, while the normal samples \( y_t = N \) are used to develop an online local model for patient-specific signal properties as presented in Fig. 5.

Table 4: Accuracy of different classification methods for the global classifier.

<table>
<thead>
<tr>
<th>Classifier Name</th>
<th>Accuracy (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Complex Tree</td>
<td>88.9</td>
</tr>
<tr>
<td>Medium Tree</td>
<td>88.8</td>
</tr>
<tr>
<td>Simple Tree</td>
<td>87.2</td>
</tr>
<tr>
<td>Fine KNN</td>
<td>88.4</td>
</tr>
<tr>
<td>Medium KNN</td>
<td>90.8</td>
</tr>
<tr>
<td>Coarse KNN</td>
<td>90.5</td>
</tr>
<tr>
<td>Cubic KNN</td>
<td>90.6</td>
</tr>
<tr>
<td>Weighted KNN</td>
<td>90.9</td>
</tr>
<tr>
<td>Subspace KNN</td>
<td>89.5</td>
</tr>
<tr>
<td>Boosted Trees</td>
<td>89.5</td>
</tr>
</tbody>
</table>

The Personalized Deviation Analysis

Here, we propose a novel deviation analysis method through a two stages process including a deviation detection stage followed by a re-classification stage to obtain final class labels denoted by \( p_t \). The first stage is to verify whether a normally classified sample by a global classifier is a firm normal \((p_t = N)\) or leaning towards one of the major alarm classes \( \{V, S, F\} \). In the later case, a yellow alarm is triggered based on re-classifying the sample into one of the three abnormality classes \( p_t \in \{V, S, F\} \). Therefore, the set of the outcomes of the whole system is \( \{N, V_t, S_t, F_t\} \), where subscripts \( g \) and \( p \) represent global and personal alarms, respectively.
In order to perform the local analysis, we first partition the abnormal samples in the training dataset DS1 (with true labels(V, S, F) into three disjoint subsets denoted by $\Omega_V, \Omega_S, \Omega_F$). Then, for a new sample $x_t$, we calculate the following distance metrics:

$$R_{max} = \max_{x_t \in \Omega(t)} \left\{ \frac{1}{\alpha} \left( x_t - x_n \right)^2 \right\}$$

(2)

$$D_N(x_t) = \text{median}_{x_n \in \Omega(t)} \left\{ \frac{1}{\alpha} \left( x_t - x_n \right)^2 \right\}$$

(3)

$$D_V(x_t) = \text{median}_{x_n \in \Omega(t)} \left\{ \frac{1}{\alpha} \left( x_t - x_n \right)^2 \right\}$$

(4)

$$D_S(x_t) = \text{median}_{x_n \in \Omega(t)} \left\{ \frac{1}{\alpha} \left( x_t - x_n \right)^2 \right\}$$

(5)

$$D_F(x_t) = \text{median}_{x_n \in \Omega(t)} \left\{ \frac{1}{\alpha} \left( x_t - x_n \right)^2 \right\}$$

(6)

In the equations above, $D_N(x_t), D_V(x_t), D_S(x_t), D_F(x_t)$ are the median of the distances of the sample $x_t$ from the members of the sets $\Omega_N(t), \Omega_V, \Omega_S$ and $\Omega_F$, respectively. $R_{max}$ represents the largest pairwise distance for the aggregated normal samples. The sets of abnormal samples $\Omega_N, \Omega_V, \Omega_S$ and $\Omega_F$ are static and developed using the training dataset DS1, whereas $\Omega_N(t)$ is a dynamic set including aggregated normal samples until time $t$ through collecting samples that are labelled Normal and do not trigger a yellow alarm, as elaborated below in Algorithm 1. Therefore, $\Omega_N(t)$ reflects the personal normal history.

In order to improve the accuracy of the local deviation analysis and avoid biasing to the very few first samples, we initialize the set $\Omega_N(0)$ with $n_0 = 300$ normal samples collected from training dataset DS1, which exhibit minimum distance with the first test sample $x_0$ ($n_0$ nearest neighbors of $x_0$ in set DS1). $n_0$ is chosen based on the average number of normal samples for each patient in dataset DS1. A sample $x_t$ is deemed normal ($p_t = N$) if we have:

$$D_{max}(x_t) < \alpha R_{max}$$

(7)

and

$$D_N(x_t) < \min(D_V(x_t), D_S(x_t), D_F(x_t))$$

(8)

where $\alpha$ is a regularizing parameter, chosen by maximizing accuracy over training dataset.

Eq. (7) monitors the relationship between the new sample and the personal normal cluster ($\Omega_N(t)$) by comparing it with the clusters ‘diameter’ ($R_{max}$), while Eq. (8) guarantees that the new sample is more likely to be a member of personal normal cluster ($\Omega_N(t)$) than other abnormal clusters ($\Omega_V, \Omega_S$ and $\Omega_F$). The illustration of this idea in two-dimensional space is shown in Fig. 6.

If Eqs. (7) and (8) do not hold at the same time, the sample is mapped to a minor abnormality of type $p_t = S$. We use a k-Nearest Neighbor classifier with $k = 10$ and a square inversion kernel trained using datasets $\Omega_V, \Omega_S, \Omega_F$ [16]. The classification steps can be summarized as:

$$k = 10, \ K(d) = \frac{1}{d^2}$$

(9)

$$d_\theta(x_t) = \sum_{x_n \in \Omega(t)} K \left( \frac{1}{\alpha} \left( x_t - x_n \right)^2 \right)$$

for $\theta \in \Omega_V, \Omega_S, \Omega_F$ (10)

$$\theta = \arg\min_{\theta \in \{V,S,F\}} d_\theta(x_t)$$

(11)

For subsequent normal samples ($x_t, t > 1$), if the new sample $x_t$ is classified as normal ( $p_t = N$), then it joins the set of aggregated normal samples $\Omega_N(t)$ and one member of $\Omega_N(t)$ with a maximum pairwise Mahalanobis distance to the other members of this set ($x_1^{(t)} = \arg\max_{x \in \Omega_N(t)} \sum_{x \in \Omega_N(t)} (x - y)^2$) is excluded.

Therefore, the set $\Omega_N(t)$ accumulates the confirmed normal samples for the patient over time and the less-representative initially chosen samples leave the set gradually in order to reinforce the personal history over time. This set can be used as
Algorithm 1: Personal Classifier
Input: Samples Labelled Normal by Global Classifier (\(g_z = N_g\))
Output: Labels by Personal Classifier (\(p_t \in \{N_p, V_p, S_p, F_p\}\))

Initialization:
Partition \(\Omega^{(0)}\) into \(\Omega_N, \Omega_S, \Omega_V, \Omega_F\);
Set \(\Omega_N^{(0)} = \emptyset\);
\(n_0 = 300\)
for \(i = 1 : n_0\) do
\(\Omega_N^{(i)} = \Omega_N^{(i-1)} \cup \text{argmin}_{x \in \Omega_N} (x_0 - x)^2;\)
end for
for \(t = 1 : \infty\) do
Calculate \(R_{\text{max}}, D_N(x^t), D_V(x^t), D_S(x^t), D_F(x^t)\);
Set \(\Omega_N^{(t)} = \Omega_N^{(t-1)}\);
If \((D_{\text{max}}(x^t) < aR_{\text{max}})\) and \(D_N(x^t) < \min(D_V(x^t), D_S(x^t), D_F(x^t))\) then
\(p_t = N;\)
\(x_e^t = \text{argmax}_{x \in \Omega_N} \sum_{y \in \Omega_N}(x - y)^2;\)
\(\Omega_N^{(t)} = \Omega_N^{(t)} \setminus x_e^t;\)
\(\Omega_N^{(t)} = \Omega_N^{(t)} \cup x^t;\)
else
\(\theta_y = \text{argmin}_{\theta \in \{V, S, F\}} d_p(x^t);\)
p\(t = \theta_y;\)
end if
end for

a reference to decide the subsequent samples. After running the algorithm for a couple of minutes, this set includes only the most recent normal samples and hence is an indicative of the personal normal history of the current patient. The summary of this algorithm is shown in Algorithm Personal Classifier.

4. RESULTS

In this section, the performance of the proposed method in terms of Classification Accuracy (AC), Specificity (SP) and Sensitivity (SE) is investigated. We first, obtain the classification accuracy by comparing the true labels \(c_t\) (expert’s annotation) with the outcome of the proposed two-step algorithm \(p_t\). At this step, we combine all abnormalities into one set defined as \(\Omega_p = \Omega_V \cup \Omega_S \cup \Omega_F\). Then, we calculate the true positive, true negative, false positive and false negative rates of class \(N\) for all test samples. In order to avoid bias to training dataset DS1, we use the 22 unseen samples in DS2 for test purpose. We also, calculate the median, interquartile range (IQR), mean and standard deviation values for all performance metrics (AC, SE, SP) in order to examine the robustness of the proposed method and performance variations among 22 test samples. We use robust statistics (median and IQR) to assess the performance of the proposed algorithm. Table 5 summarizes the overall results and Fig. 7a), shows the histogram of accuracy for class \(N\), where interquartile range (IQR) is used to measure the stability of performance.

Table 5: Classification accuracy and sensitivity of the proposed method for Class N

<table>
<thead>
<tr>
<th>Class N</th>
<th>median (%)</th>
<th>IQR (%)</th>
<th>mean(%)</th>
<th>std(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC</td>
<td>97.6</td>
<td>13.2</td>
<td>88.3</td>
<td>19.4</td>
</tr>
<tr>
<td>SE</td>
<td>98.3</td>
<td>9.9</td>
<td>86.2</td>
<td>26.5</td>
</tr>
</tbody>
</table>

Figure 7: Accuracy histogram among all patients for classes: a) N, b) V, c) S and d) F.
The results are promising and suggest that the probability of missing a true alarm is negligible using the proposed two-step method with minor alarm recognition capability. The cost paid is more complexity and reporting minor alarms. There is a tradeoff between the rate of minor alarms and the sensitivity of the method, that can be regularized by tuning regularizing parameter \( \alpha \) in Eq. (7). Here, we used \( \alpha = 1 \).

In order to analyze the specificity of the method, we consider all classes individually and count the number of calling false alarm types per class. The results are summarized in Table 6, which demonstrate an excellent specificity per class. Likewise, Figs. (6b), (6c) and (6d) present the histogram for accuracy across samples for classes V, S and F, respectively. In the results presented in Table 5 and Table 6, both minor (yellow) and major (red) alarms are included.

<table>
<thead>
<tr>
<th>Class</th>
<th>Median (%)</th>
<th>IQR (%)</th>
<th>Mean (%)</th>
<th>Std (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC</td>
<td>82.8</td>
<td>26.1</td>
<td>73.9</td>
<td>25.0</td>
</tr>
<tr>
<td>SP</td>
<td>99.8</td>
<td>9.9</td>
<td>89.6</td>
<td>20.2</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Class</th>
<th>Median (%)</th>
<th>IQR (%)</th>
<th>Mean (%)</th>
<th>Std (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AC</td>
<td>68.6</td>
<td>22.2</td>
<td>65.9</td>
<td>20.2</td>
</tr>
<tr>
<td>SP</td>
<td>99.5</td>
<td>2.0</td>
<td>93.6</td>
<td>21.2</td>
</tr>
</tbody>
</table>

From the result presented in Table 6, it is easy to notice that the performance for V and F classes are better, while the performance for class F is more stable compared to class V. This result also is consistent with the fact that class F is a Fusion class considered as an intermediate state between classes N and V [14].

A unique property of the proposed method is the utility of the minor (yellow) alarms in predicting upcoming major (red) alarms. In order to investigate this property, for each patient, we count the number of subsequent red alarm types that come after a yellow alarm. To evaluate the global predictive capacity of the method across the test data, we firstly assess the segments collected from all samples in the test dataset. The general results are presented in Table 7. As shown, there are in total 439 type V samples, 97 type S samples and 24 type F samples in DS2 according to the ground truth. For instance, the Table 6 suggests that we detect 20 yellow alarms of type S that are followed by a red alarm of type F. Also, the probability of observing major alarms are simply calculated as

\[
\text{Pr(alarm of type } \theta) = \frac{n_\theta}{n_S + n_V + n_F}, \quad (\theta \in \{S, V, F\})
\]  

(12)

where \( n_\theta \) is the number of red alarms of type \( \theta \). The results are interesting and in general show that

\[
\text{Pr(red alarm } \theta \text{ after a yellow alarm } \theta) > \text{Pr(red alarm } \theta)
\]  

(13)

For instance, comparing the subsequent major alarms after a yellow alarm \( V_p \) reveals that there is 84% chance of having an upcoming red alarm of type V. However, the probability of having an alarm of type V, without considering the preceding yellow alarms of type V is only 78%. The same trends holds for alarm types S as well.

Additionally, noticing the records in DS2 usually have either type V or type S dominant over other type of alarms. It’s also necessary to consider the predictive capacity of individual record. For example, in Table 8, record number 219 has type V dominant over other abnormal sample types. The conclusion drawn from Table 7 is also valid for this patient. For the patients whose dominant alarm type is S, an example from record number 100 is shown in Table 9. All yellow alarms of type S are followed by a red alarm of type S. The predictive accuracy is 100% for this record.

At the same time, it is worthwhile to compare the performance of proposed method for different alarm types and the consistency of the performance as well. For the 22 records in DS2, there are 19 records which have at least one major alarm. Among these records, 10 records showing the same trend for alarm type S and 7 records showing a good predictive capacity for type V. However, the prediction of alarm type F is not as promising as it is for other alarm types, reflecting the fact that alarm type F is a fusion type of class V and class N [17]. In general, the algorithm show a better performance over alarm type S than type V.

This demonstrates an important utility of the yellow alarms, since it means that by detecting yellow alarms, we can predict the upcoming red alarms with a higher certainty. A high-risk patient may take into account these yellow alarms and take cautionary actions (e.g. avoiding physical activity).

<table>
<thead>
<tr>
<th>Yellow alarm tag</th>
<th>Number of next abnormalities</th>
<th>Probability of next abnormality (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( V_p )</td>
<td>( S_p )</td>
</tr>
<tr>
<td>True V</td>
<td>179</td>
<td>256</td>
</tr>
<tr>
<td>True S</td>
<td>30</td>
<td>67</td>
</tr>
<tr>
<td>True F</td>
<td>4</td>
<td>20</td>
</tr>
</tbody>
</table>

Table 8: Predictive power of yellow alarms, result for record number 219.

<table>
<thead>
<tr>
<th>Yellow alarm tag</th>
<th>Number of next abnormalities</th>
<th>Probability of next abnormality (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( V_p )</td>
<td>( S_p )</td>
</tr>
<tr>
<td>True V</td>
<td>21</td>
<td>7</td>
</tr>
<tr>
<td>True S</td>
<td>2</td>
<td>3</td>
</tr>
<tr>
<td>True F</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>
Table 9: Predictive power of yellow alarms, result for record number 100.

<table>
<thead>
<tr>
<th>Yellow alarm tag</th>
<th>Number of next abnormalities</th>
<th>Probability of next abnormality (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$V_p$</td>
<td>$S_p$</td>
</tr>
<tr>
<td>True V</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>True S</td>
<td>0</td>
<td>20</td>
</tr>
<tr>
<td>True F</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

5. CONCLUSIONS

A prototype of a remote heart monitoring system with wireless connectivity is implemented. A novel two-step predictive modeling is designed by adding an additional analysis layer to the common approach of using global classifiers. With the proposed method, a normal trend of ECG signal for each patient is learned from the received samples over time. This trend is subsequently used to trigger minor (yellow) alarms in case of deviating from the patient-specific normal history. The results suggest that these yellow alarms are informative in the sense that they provide insight about the upcoming major (red) alarms. Since the reaction time in cardiac disease is very important and can significantly reduce catastrophic consequences, the proposed device can be used to notify the users about potential upcoming major alarms in order to take necessary preventive actions. This kit also can be used to assist the physicians and healthcare staff to interpret the patient’s ECG signal more accurately. Finally, by incorporating the accelerometer signal into the predictive modeling, we avoid calling fake alarms due to the patient’s physical activities.

6. REFERENCES