A New Methodology for Classifying QRS Morphology in ECG Signals

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Abstract—The electrocardiogram (ECG) is a non-invasive method to detect cardiovascular diseases (CVD), the most common cause of death in the world. The recognition of heartbeat morphologies present in the ECG signal is an effective way to detect CVDs prematurely. Many approaches were developed for this purpose, such as the use of Wavelets, High Order Statistics (HOS), Local Binary Patterns (LBP), Random Projection, Fiducial points, and Hermite Polynomials. Unfortunately, most parts of these approaches suffer from the high variability of ECG signal features and conditions. Also, it is common to use more than one of them simultaneously, which makes it hard to infer the contributions of each one. This work presents a new robust methodology to extract features for heartbeat morphology classification. Moreover, we introduce new labels for a small set of morphologies present in MIT-BIH Arrhythmia database, taking into account only the QRS complex (the 3 more representative waves of a heartbeat) instead of the whole heartbeat. We evaluate each approach in isolation and the results show that our method outperforms other well-known strategies.

Index Terms—ECG signal, QRS morphology, feature extraction, prediagnosis.

I. INTRODUCTION

Cardiovascular diseases (CVD) are the most common cause of death globally, producing immense health and economic burdens. According to the World Health Organization (WHO): “People with cardiovascular disease or who are at high cardiovascular risk need early detection and management using counseling and medicines, as appropriate”. Despite the continuous advance of medicine, the ECG continues to be a crucial non-invasive tool to detect CVD. As a result, the ECG data analysis has become a significant field of study. Most of the useful information within ECG data is present on the intervals and amplitudes which characterize the significant regions (wave peaks and boundaries) that compose the standard cardiac cycle (heartbeat) [14]. These characteristic waves occur due to electrical changes caused by depolarization/repolarization cycles inside the heart. Depolarizations result in atrial contractions, which are associated with P waves, and ventricular contractions, which are associated to the QRS complex. Polarization results in the return of the ventricular mass to the relaxation state, producing the T waves. The junction of all these waves forms the PQRST cycle.

During, PQRST complex analysis (shape, duration, interval patterns and etc.) has received much attention from the research community, resulting in significant advances in many areas like disease diagnosis [12, 17], heartbeat segmentation [14], and heartbeat delineation [1, 14, 25]. Incidentally, the QRS complex is the most characteristic waveform of the PQRST cycle, presenting, in general, a higher amplitude than the other waves.

The morphological structure (curve shape) of the QRS complex is one of the main features of a heartbeat and may indicate many types of diseases or disorders [2]. In [7], the authors have demonstrated that morphologies serve to robustly predict long-term mortality in Left Ventricular Pacing (LVP) patients. Another important task solved by recognizing morphologies is detecting arrhythmias [22, 23]. Furthermore, the correct identification of the heartbeat shape helps on prognostics: patients that present non-LBBB (Left Bundle Branch Block) morphology commonly do not respond well for specifics treatments, like Cardiac Resynchronization Therapy (CRT) [20]. Finally, the QRS morphology is a better indicator than the QRS duration for long-term survival, since people with LBBB morphology present less ischaemic cardiomyopathy and atrial fibrillation [9].

The importance of the QRS morphology for diagnostics/prognostics motivated many approaches to recognize such shapes, such as the use of Wavelets [13, 14], Higher-Order Statistics (HOS) [3, 18, 19], Random Projection [6], Fiducial points of the heartbeat [17], and Hermite Polynomial [10]. Unfortunately, most of these features do not perform well when mismatch data is present, i.e., they need the test instances to be similar to the training set. Besides, since most parts of these methodologies are evaluated in conjunction with other approaches, it is hard to determine the real efficiency of each one.

This work introduces a new approach to extract features from the QRS complex based on the so-called mathematical models [5] and presents a comparative study with all methodologies previously mentioned. We named our approach as feature extraction via residuals from modeling with composition of mathematical functions (RCMF). From five mathematical models built upon three functions (Rayleigh, Gaussian, and Mexican-hat), we create a morphological descriptor related only to the shape of each beat, instead of the amplitude and duration of its waves.
To validate our approach, we consider the well-known public dataset MIT-BIH Arrhythmia database. Typically, this dataset contains five classes of morphologies representing arrhythmias: N corresponding to any heartbeat which does not pertain to other categories, S, supraventricular ectopic beat, V, ventricular ectopic beat, F, fusion beat, Q, unknown beat. Different from the previous works, we do not focus on classifying the entire heartbeat morphology, but only on the QRS complexes morphology. This classification is already made implicitly in the other approaches, which generally extract the QRS complex, apply its descriptor and then classify the complete cardiac cycle. However, to ascertain the describing ability of the QRS complex provided by each method, we will only classify its morphologies. For this purpose, we use a synthetic ECG simulator to produce artificial ECG data for the four typical morphologies qRs, RS, rRs and QS to constitute our training set. After that, we select a small portion of the MIT-BIH dataset (14 subects) whose QRS complexes contain the four types of morphologies qRs, RS, rRs, QS and apply an algorithm to detect QRS complex. The full set of beats is labeled by a specialist concerning each morphology and serves as our validation set. The performed experimental results show that our methodology outperforms other well-know feature extraction methodologies available in the literature.

The remaining of this paper is organized as follows. Section II presents the methodology applied for generating artificial signals and the common features applied for heartbeat classification. Section III shows the proposed approach based on mathematical models. Section IV details the corresponding related work within the literature. Section V describes the training/test datasets applied in this work. Description of the experiments and results using the proposed labels for MIT-BIH can be seen in Section VI. Finally, Section VII presents our conclusions and comments for future works.

II. ECG DATA SIMULATION

The QRS complex presents a high variability, since both the shape and the amplitude of the waves are governed by multiple individual factors, like the shape/position of the heart and the presence and nature of pathologies, among others [11]. As a result, the QRS complex can be categorized depending on the shape of its waves. In this study, we use the 4 common variations(qRs, RS, QS and rRs), presented in Fig. 1. Table I presents the average of corresponding wavelengths and amplitudes.

<table>
<thead>
<tr>
<th>Type of Wave</th>
<th>ECG normal values</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Amplitude(mV)</td>
</tr>
<tr>
<td>Q</td>
<td>0.00 - 0.30</td>
</tr>
<tr>
<td>R</td>
<td>0.60 - 2.10</td>
</tr>
<tr>
<td>S</td>
<td>0.00 - 0.60</td>
</tr>
</tbody>
</table>

To construct a pattern recognition model based on different morphologies, the initial knowledge base must be composed by specific ECG signals of each morphology. Unfortunately, labeling the ECG signals involves time and the involvement of a specialist, besides patients’ permission. To mitigate this problem, a heartbeat simulator has been developed which can accurately approximate a real beat [15].

The simulator is characterized by a dynamic system based on three differential equations capable of generating realistic synthetic ECG signals. It is possible to specify the mean frequency and the standard deviation of the heart rate, the power spectrum of the RR time-series, the QRS complex morphology, Low Frequency (LF) and High Frequency (HF) bandwidth for heart rate variability along with the LF/HF ratio. Finally, the amplitudes and modulations of the PQRST cycle can also be specified, allowing a considerable variability for generated synthetic ECG signals.

Fig. 2 illustrates an example of a synthetic PQRST complex containing a QRS-complex with a qRs morphology.

III. ECG FEATURE EXTRACTION METHODOLOGY

Typical ECG signals are composed by many heartbeats. Each beat represents one ECG complex, or in other words, a PQRST cycle. Also, every ECG complex contains a QRS complex formed by the waves Q, R, and S. For our proposal, we need to find and extract the QRS-complex held in each ECG complex. The task of separating each QRS complex of an ECG signal is named QRS detection.

Among all approaches, Pan Tompkins[4] is a well-known technique to detect the QRS-complex. The Pan Tompkins algorithm identifies QRS complexes using digital analysis of amplitude, width, and slope of the ECG wave. Also, it uses a
patient-specific threshold for QRS peak detection, periodically adjusted to adapt to the changes in QRS morphology and heart rate.

After using Pan Tompkins algorithm for beat detection within the ECG signal, we obtain a set of QRS complexes, from which the features will be extracted. Our goal is to classify the morphology present in each QRS complex. Due to the significant morphological variability of the heartbeats, the process for feature extraction needs to be accurate and capture the correct information for each morphology, independent of amplitude levels, frequency sampling, and intrinsic noise of the ECG signal.

In our approach, we adopt three different functions that present a strong correlation with the structures of typical ECG morphologies presented in Fig. 1. The functions are Gaussian, Mexican Hat and Rayleigh, respectively defined below:

\[
N(x|\mu, \sigma^2) = \frac{1}{\sqrt{2\pi\sigma^2}} e^{-\frac{(x-\mu)^2}{2\sigma^2}},
\]

\[
M(x|\lambda) = \frac{2}{\sqrt{3\lambda\pi}} \left(1 - \frac{x^2}{\lambda^2}\right) e^{-x^2/2\lambda^2},
\]

\[
R(x|\lambda) = \frac{x}{\lambda^2} e^{-x^2/2\lambda^2},
\]

where \(\mu\) and \(\lambda\) are location parameters and \(\sigma^2\) is the variance/scale parameter of the functions. Each of these functions has similarities when compared with some ECG morphologies and, by combining them, it is possible to find patterns that identify a high number of morphologies. Fig. 3 shows the distribution of the cited functions.

Fig. 3: Illustration of the functions Rayleigh, Gaussian and Mexican Hat.

We build five different mathematical models by combining the individual functions: Gaussian, Mexican Hat, and Rayleigh. Each model is composed by two of these functions, according to Table II, the choice of functions was made in order to represent the largest number of morphologies, with the least number of models possible. For simplicity, we will call A/B, the first and second/functions used to build each model.

To compose models I and II, first we use each composition function to generate an individual signal model in which the number of points on the output window is linearly proportional to the number of points of the QRS complex window. We do this process as follows: given the size of the QRS complex window \(w\), we build two models, A and B, with length \(s_a\) and \(s_b\), respectively. The length of the models (i.e., the number of points) is chosen from the range \([0.5w, 0.7w, 0.9w, 1.1w, \ldots, 1.9w, 2w]\). Then, we remove the first/last \(k\) points of the models A and B, where \(k\) is chosen from \([0.1s, 0.2s, \ldots, 0.5s, 0.6s]\). The process for selecting the parameters will be discussed in detail later. Besides that, we extend the models A and B by repeating their first/last value \(w\) times. After that, we select the peak with most significant absolute value for models A and B, and normalize them, forcing their peaks to have the same value. Finally, we merge the models overlapping both peaks. More specifically, the left part of the final model has the same distribution of the left part of the composition function A centered on their most significant peak, while the right part of the model has the same distribution of the right part of the function B, also centered in their most significant peak. Fig. 4 shows the final model merged from models A and B. From this point, for the sake of simplicity, we will omit that the left and right parts of a signal are always related to its most significant peak.

Fig. 4: Fitting the original signal above extented version of model II. In green line: the heartbeat; Black pointed line: RCMF model; Cut A: cut point for function A. Cut B: cut point for function B. Start/end beat represent the limits of heartbeat window.

As illustrated by Figs. 5a, 5b, 5c and 5d, for the models III, IV, and V, the intersection of the two composition functions is the first point with zero value after/before respectively from the corresponding most significant peak of the functions A and B. The highest peaks from both functions are preserved.

**TABLE II: Composition of each proposed mathematical model.**

<table>
<thead>
<tr>
<th>Model</th>
<th>Mathematical Models</th>
</tr>
</thead>
<tbody>
<tr>
<td>I</td>
<td>Gaussian</td>
</tr>
<tr>
<td>II</td>
<td>Mexican Hat</td>
</tr>
<tr>
<td>III</td>
<td>Rayleigh (-1)Rayleigh</td>
</tr>
<tr>
<td>IV</td>
<td>Rayleigh Rayleigh</td>
</tr>
<tr>
<td>V</td>
<td>Rayleigh</td>
</tr>
</tbody>
</table>
is a task very sensitive to noise presence. Another common
versions of the ECG signal [14]. However, detecting peaks
by thresholds, at scales that work, the morphology of QRS complexes depends on
baseline wandering, and changes in the QRS morphology. In
Martínez et al. [14], a Wavelet-based ECG delineator
techniques proposed for identifying heartbeat morphologies.
Wavelet-based approaches have been one of the first family of
results when applied to identify a variety of heart disorders.

Different works to detect/prevent such diseases. Among all
in medicine. The advances of computer-aided diagnosis propel
assumptions are needed, like a zero mean of the heartbeat
formation for nonlinear behavior. On the other hand, additional
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The best feature set for defining a range of parameters to experiment and construct
the current model using the composition functions A and B, usually \( \mu = 0, \sigma^2 \) and \( \lambda \in \{0.1, 0.5, 1, 1.5, \ldots 9.5, 10\} \). After
building the model, it is necessary to find the R peak (for
inverse R peaks like on QS morphology, we multiply the signal
by -1) of a given heartbeat and the most significant peak of
the model to normalize them in such a way that both maxima
are at the same point. The construction of the proposed model
ensures that the size of the model is greater than or equal
to the size of the QRS complex window. To correct this, we
cut the remaining portion of the model based on a window of
size equal to the QRS complex window, centered on the most
significant value of the model.

At this point, both the signals, the heartbeat and the model,
have the same length. To evaluate the selected parameters, we
calculate the Mean Square Error (MSE) for the left/right part
of the heartbeat/model. There is no difference in calculating
the MSE separately for segmented parts of the ECG heartbeat,
but since their morphology is not symmetric, it is possible
to acquire some information concerning how differently the
model fits different portions of the signal.

After the previous step, we have ten features composed by
the evaluation errors of the mathematical models for each
heartbeat piece. A normalization procedure is then applied
for the feature set, to force the interval range \([0, 1]\). This procedure
ensures that the features do not represent absolute errors, but
relative indicators concerning the performance of the different
models for fitting a given heartbeat.

Finally, we add a variable indicator concerning the sign of
the R peak to the feature set. This indicator will help to decide
between morphologies with opposite signs but with very
similar relative errors. Algorithm 1 summarize the proposed
method.

\begin{algorithm}
\caption{Feature extraction via residuals from modeling
with composition of mathematical functions (RCMF)}
\begin{algorithmic}
\State {\bf Input:} QRS complex.
\State {\bf Output:} the best found MSE (left and right parts) for each
mathematical model. Original sing of R peak.
\For {each function in Eq. (1), Eq. (2) and Eq. (3)}
\For {each parameter set \((w, s, k, \lambda, \mu, \sigma^2)\)}
\State Build the models \(A\) and \(B\) from composition functions
\(A\) and \(B\) and QRS window of size \(w\).
\State Remove the first/last \(k\) points on the models \(A\) and
\(B\).
\State Repeat the first/last \(w\) points on the models \(A\) and
\(B\).
\State Normalize the models \(A\) and \(B\).
\State Combine the models by overlapping their most signi-
ficant peak peaks for models I and II and preserving
both peaks for models III, IV and V.
\State Match the most significant value of the final model
and the inputted QRS complex.
\State Cut the remaining points on the final model to build
a model with window of size \(w\).
\State Find the most significant peak of the final model.
\State Calculate the MSE from left/right part of the
model/heartbeat matching the most significant peak
of the final model and the R peak of the QRS
complex.
\EndFor
\State Select the best MSE (sum of the MSE of left and right
parts) among all tested possibilities.
\EndFor
\State Calculate the sign of the R peak.
\end{algorithmic}
\end{algorithm}

\end{algorithmic}

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the R peak to the feature set. This indicator will help to decide
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IV. RELATED WORK

Prevention of death by heart disorders is an essential field
in medicine. The advances of computer-aided diagnosis propel
different works to detect/prevent such diseases. Among all
approaches, heartbeat shape recognition has shown promising
results when applied to identify a variety of heart disorders.
Wavelet-based approaches have been one of the first family of
techniques proposed for identifying heartbeat morphologies.
In Martínez et al. [14], a Wavelet-based ECG delineator
was proposed to deal with motion artifacts, muscular noise,
baseline wandering, and changes in the QRS morphology. In
that work, the morphology of QRS complexes depends on
the number of negative/positive peaks, which are governed
by thresholds, at scales \(2^1, 2^2, 2^3, 2^4, 2^5\) related to filtered
versions of the ECG signal [14]. However, detecting peaks
is a task very sensitive to noise presence. Another common
wavelet approach is to consider Daubechies wavelet (db1) to
extract features according to Mar et al. [13]. As in [17], we
have opted to use a 3-level decomposition to build a 23-feature
array.

Later, statistical ECG descriptors were proposed in [3, 19],
both based on the second, third, and fourth-order cumulants,
with robustness to additive noise and capable of extracting in-
formation for nonlinear behavior. On the other hand, additional
assumptions are needed, like a zero mean of the heartbeat
signal. As in [17], each heartbeat was split into five bins and
had their kurtosis and skewness extracted, resulting in a 10-
feature array.

Concomitantly, [16] presented a curve fitting methodology
for on-line heartbeat type recognition. After the QRS detec-
tion, the heartbeat was fitted using Hermite basis functions,
which have the coefficients extracted and applied as features
to characterize the shape of the signal. The main idea of
this work is based on [10], which considers the similarity of
the forms between fitted Hermite polynomials and QRS
complexes. Unfortunately, it is possible to insert unusable
features, since not all coefficients need to represent QRS
morphologies well.
Moreover, many other approaches were proposed to classify arrhythmia. The first one is the 1D-local binary patterns (a variant of 2D-local binary pattern [8]) where each point of the heartbeat is compared with the neighbors producing a binary code. The histogram of the frequency of each binary pattern is used as a feature. The original work proposes an 8-neighbor 1D-LBP built 59-dimensional descriptor for a 180-dimensional heartbeat [17]. Secondly, in [6], an ensemble of SVM’s with multiple random projections was applied to heartbeat classification. The features can be understood as multiple independent groups of random features. In our work, we reproduce the author’s experiment using 15 classifiers with 50 features for each projection and using the majority vote to predict the classes. Finally, Mondéjar-Guerra et al. [17] recently proposed a morphological descriptor of the PQRST complex based on four windows. For a heartbeat with a window size of 180 samples (500ms) (centered on the R peak), the morphological descriptor is calculated using the Euclidean distance between the R peak and the points below:

1) \( \max(\text{beat}[0:40]) \)
2) \( \min(\text{beat}[75:85]) \)
3) \( \min(\text{beat}[95:105]) \)
4) \( \max(\text{beat}[150:180]) \)

After that, all features are normalized between 0 and 1. Nevertheless, all of these approaches are severely affected by noise incidence.

Finally, many deep neural networks were proposed showing promising results [21, 24]. In this work, we reproduce the CNN 1D network at [21], with seven blocks of convolutional layers (128 filters, width 5), max-pooling, and dropout (rate=0.5). In the end, a global average pooling, 3 Full connected layers with (256/128/64), and softmax.

V. DATASET AND PRE-PROCESSING

We apply a pre-processing step before extracting features from the available data. Following most of the previous works, for each signal we remove the baseline. We use a high order (10) polynomial to calculate our baseline. Then, we subtract it from the original signal. Similarly to [17], we have opted not to add any other noise filter or modification to preserve most of the original signal.

After extracting the heartbeats, we select the QRS segment as 71 data points around the R peak (31 points before and 35 ones after). Since the data sample rate is 360 Hz, a window of 71 points gives us approximately 200 ms, which is long enough to cover all the QRS complexes experimented in this work.

A. Artificial dataset

Using the previously described ECG data simulator, 10 synthetic signals were created for the morphologies qRs, RS,
QS, and rRs, composing a total of 40 signals with varying amplitudes and durations, according to Table I. The frequency of the signals was set at 360 Hz, with the heart rate ranging from 60 to 100 beats per minute and LF/HF ratio of 0.5 and approximately 2 hours in duration. For each signal, we apply the Pan Tompkins method to extract 25 heartbeats, which had the P and Q waves removed, leaving only the QRS complex. This results in a total of 1000 heartbeats, 250 for each morphology. For each beat, a window of size 35 samples centered around its R peak was used to extract the QRS complex.

B. MIT-BIH Dataset

The MIT-BIH Arrhythmia dataset contains 48 half-hour excerpts of two-channel ambulatory ECG recordings obtained from 47 subjects studied by the BIH Arrhythmia Laboratory between 1975 and 1979. To build our test dataset, we select fourteen subjects (100-lead 1, 100-lead 2, 101-lead 1, 103-lead 1, 105-lead 2, 106-lead 1, 108-lead 2, 111-lead 2, 215-lead 1, 215-lead 2, 121-lead 2, 223-lead 2, 123-lead 2) that contain the morphologies qRs, RS, QS, and rRs. Then, we apply the Pan Tompkins method to extract a set of heartbeats for each subject/morphology class. After that, the extracted QRS complex windows with 71 samples, or approximately 200ms, were manually labeled by a specialist, forming a dataset with 240 QRS complexes (60 for each morphology).

VI. EXPERIMENTAL RESULTS

To investigate the efficiency of the proposed features, we conducted two experiment designs that aim to compare our proposed work and other standard feature extraction techniques used on ECG signal classification.

In the first one, we assessed the efficiency of all feature extraction methods using only the simulated data provided by the synthetic generator. We randomly selected 2/3 of the instances for training and 1/3 for testing, ensuring that QRS complexes of the same time-series must be present only on training or testing set but not on both at the same time. We chose Support Vector Machine (SVM) as the learning model, with the hyperparameters tuned following a 5-fold cross-validation in a grid-search procedure. For the hyperparameter search space, we chose \( C \in \{2^{-5}, 2^{-1}, 2^{-1}, \ldots, 2^{12}, 2^{15}\} \) and \( \gamma \in \{2^{-15}, 2^{-13}, 2^{-11}, \ldots, 2^{1}, 2^{13}\} \) for the Gaussian kernel, and \( C \in \{2^{-1}, 2^{-1}, 2^{-1}, \ldots, 2^{12}, 2^{15}\} \) for the linear kernel. The experiment was repeated 20 times.

To promote a fair comparison, we propose some adaptations for each approach when necessary. At first, since the morphological descriptor by Mondéjar-Guerra et al. [17] was designated for a heartbeat with length 180 samples (500ms) instead of 71 (~200ms), a rescale interpolation was applied to ensure the correct length. Besides that, the original methodology was designed for the whole heartbeat (including the P and T waves) choosing 4 regions to extract their features. In this work, we set the first and four regions on the extremities, and move the second and third regions, both with window size of 10 samples along the time axis in opposite directions, skipping the same time of their window. Finally, we repeat this procedure increasing their window(second and third fiducial regions) size for 20, 30 and 40.

The original approach of Hermite Polynomials extends the QRS complex by adding 45 zero values to each of their extremities. Then, all QRS complexes are linearly normalized in the range \([-1, 1]\), and finally, the mean level of the first and the last data points is subtracted. We choose not apply this preprocessing step, since, in our preliminary experiments, it achieved poor results.

For the wavelet approach by Martín et al. [14], the hyperparameter \( \alpha \) was tuned within the values \([0.4, 0.5, 0.55, 0.6]\). This hyperparameter represents a threshold to identify real peaks instead of noise. The remaining methods were used without any modifications. We emphasize that only the artificially labeled data were used to select the hyperparameters.

It is interesting to note that according to Table III, all feature extraction methods achieve a good performance. In fact, only the 1D-LBP method had an accuracy lower than 90%. This finding confirms our initial hypothesis that artificial data can be learned by well-known heartbeat feature extraction methods.

<table>
<thead>
<tr>
<th>Methods</th>
<th>Accuracy</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mondéjar-Guerra et al. [17]</td>
<td>0.980 ± 0.015</td>
</tr>
<tr>
<td>Wavelets Martinez et al. [14]</td>
<td>0.930 ± 0.030</td>
</tr>
<tr>
<td>Wavelets Mar et al. [13]</td>
<td>0.981 ± 0.054</td>
</tr>
<tr>
<td>Random Projection</td>
<td>1.000 ± 0.000</td>
</tr>
<tr>
<td>RCMP</td>
<td>0.997 ± 0.003</td>
</tr>
<tr>
<td>1D-IBP</td>
<td>0.898 ± 0.035</td>
</tr>
<tr>
<td>Hermite Polynomial</td>
<td>0.961 ± 0.051</td>
</tr>
<tr>
<td>1D CNN</td>
<td>0.982 ± 0.031</td>
</tr>
<tr>
<td>High Order Statistics</td>
<td>0.991 ± 0.018</td>
</tr>
</tbody>
</table>

The second experiment consists of using the artificial data to train the models and then evaluate them using the real data. We tuned all parameters using a 5 k-fold cross-validation with the artificial data set. Besides that, we repeat the experiment 20 times. To make the artificial data as more realistic as possible, for each extracted heartbeat, we repeat the experiment.
adding a Gaussian noise with mean 0 and variance within the values $[0, 0.2\text{rms}(s), 0.3\text{rms}(s), 0.4\text{rms}(s)]$, where $\text{rms}(s)$ represent the root mean square level of the heartbeat. Fig. 6 illustrates the noise levels considered in this experiment.

Initially, we expected that all methods would have similar performance on the first and second experiments. However, as shown in Table IV, the feature sets obtained by wavelets and 1D-LBP achieved a poor performance in the second experiment when compared to the first experiment. Thus, we can hypothesize that these methods cannot handle slight differences between training and test sets. For instance, the waveform features are based on peak detection, being therefore very sensitive to noise addition.

Apart from this slight discordance, the other methods performed well with accuracy greater or equal than 70%, including the other waveform approaches. As expected, the artificial noise on the training dataset provided a significant impact on the performance of the methods. It can be noticed that in most cases there were improvements in the accuracy for the 1° and 2° noise levels, possibly indicating that those noisy artificial signals are better approximations of real heartbeats with muscle noise.

It is important to note that Hermite polynomial and RCMF (our approach) acquired the best results among all the techniques. Both methodologies are based on fitting curves, which explain why they can better recognize the shape of the heartbeats. Our approach is the best in 3 of 4 scenarios, and it has obtained equal statistical results in one. This finding reinforces the efficiency of the proposed method. To support this, we have used the Kolmogorov-Smirnov test to verify whether two empirical data distributions are the same. In our experiments, it was used to compare pairs of classifiers and assess if the performance of any classifier is significantly different. Table V shows the results for our approach and Hermite Polynomials, for all noise levels.

According to the performed Kolmogorov-Smirnov test, we cannot reject the null hypothesis (two samples are drawn from the same distribution) if the p-value is greater than 0.05, which means that the performance of two models is significantly different. Considering this definition, we can verify that the RCMF approach significantly outperformed Hermite polynomial in the first 3 of 4 scenarios and it has statistically equal results on the other one. In summary, RCMF constitutes a valid alternative to other standard approaches, since they present a better or equal accuracy.

VII. CONCLUSION

Early detection of cardiovascular diseases (CVD) is a relevant field in medicine. The current approaches have many drawbacks, such as being noise sensitive or presenting poor generalization on unseen data. To overcome those issues, we proposed a new method, named feature extraction via residuals from modeling with composition of mathematical functions (RCMF), for the classification of the QRS morphology in ECG signals.

Based on the results obtained from computational experiments, we have verified that the RCMF approach achieved better accuracies when compared to other well-known feature extraction techniques for heartbeat classification. Additionally, we proposed new labels for some records of the MIT-BIH Arrhythmia database.

Future work may include new compositions of functions, like beta, triangular and bimodal distributions, aiming to extract information for new morphologies.

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REFERENCES


TABLE IV: Performance comparison for all the evaluated feature extraction methods.

<table>
<thead>
<tr>
<th>Method</th>
<th>Without Noise ±</th>
<th>0.1rms(s) ±</th>
<th>0.2rms(s) ±</th>
<th>0.3rms(s) ±</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mondéjar-Guerra et al. [17]</td>
<td>0.777±0.048</td>
<td>0.728±0.019</td>
<td>0.691±0.087</td>
<td>0.676±0.029</td>
</tr>
<tr>
<td>Wavelet/Martínez et al. [14]</td>
<td>0.641±0.009</td>
<td>0.474±0.014</td>
<td>0.443±0.071</td>
<td>0.587±0.003</td>
</tr>
<tr>
<td>Wavelets Mar et al. [13]</td>
<td>0.867±0.002</td>
<td>0.765±0.001</td>
<td>0.867±0.124</td>
<td>0.867±0.100</td>
</tr>
<tr>
<td>1D-LBP</td>
<td>0.217±0.091</td>
<td>0.546±0.026</td>
<td>0.360±0.015</td>
<td>272±0.037</td>
</tr>
<tr>
<td>Random Projection</td>
<td>0.873±0.006</td>
<td>0.870±0.049</td>
<td>0.862±0.015</td>
<td>900±0.005</td>
</tr>
<tr>
<td>RCMF</td>
<td>0.991±0.013</td>
<td>0.995±0.001</td>
<td>0.989±0.003</td>
<td>0.950±0.048</td>
</tr>
<tr>
<td>Hermite Polynomial</td>
<td>0.961±0.013</td>
<td>0.971±0.011</td>
<td>0.975±0.028</td>
<td>0.949±0.032</td>
</tr>
<tr>
<td>1D CNN</td>
<td>0.931±0.014</td>
<td>0.922±0.022</td>
<td>0.922±0.026</td>
<td>0.919±0.017</td>
</tr>
<tr>
<td>High Order Statistics</td>
<td>0.707±0.017</td>
<td>0.707±0.019</td>
<td>0.786±0.008</td>
<td>0.711±0.005</td>
</tr>
</tbody>
</table>

TABLE V: Kolmogorov-Smirnov test for Hermite Polynomials and RCMF.

<table>
<thead>
<tr>
<th>Noise level variance</th>
<th>KS statistic</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>9.00000000</td>
<td>0.00000036</td>
</tr>
<tr>
<td>0.1rms</td>
<td>0.94999999</td>
<td>0.000000004</td>
</tr>
<tr>
<td>0.2rms</td>
<td>0.69999999</td>
<td>0.000041504</td>
</tr>
<tr>
<td>0.3rms</td>
<td>0.25000000</td>
<td>0.497342335</td>
</tr>
</tbody>
</table>


