Arrhythmia Classification of Reduced-Lead Electrocardiograms by Scattering-Recurrent Networks

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Abstract

We describe an automatic classifier of arrhythmias based on 12-lead and reduced-lead electrocardiograms. Our classifier composes the scattering transform (ST) and a long short-term memory (LSTM) network. It is trained on PhysioNet/Computing in Cardiology Challenge 2021 data. The ST captures short-term temporal ECG modulations while reducing its sampling rate to a few samples per typical heart beat. We pass the output of the ST to a depthwise-separable convolution layer which combines lead responses separately for each ST coefficient and then combines resulting values across ST coefficients. At a deeper level, 2 LSTM layers integrate local variations of the input over long time scales. We train in an end-to-end fashion as a multilabel classification problem with a normal and 25 arrhythmia classes. We used canonical correlation analysis (CCA) for transfer learning from 12-lead ST representations to reduced-lead ones. For 12-, 6-, 4-, 3- and 2-leads, team “BitScattered” Challenge metrics on the hidden validation set were 0.46, 0.44, 0.45, 0.46 and 0.43; and on the hidden test set were 0.10, 0.11, 0.10, 0.10 and 0.10, respectively, ranking 34\(^{1}\)th on the hidden test set.

1. Introduction

The World Health Organization estimates that cardiovascular diseases (CVDs) caused 17.9 million deaths worldwide in 2016, and may reach 23.6 million in the year 2030. In this context, electrocardiography (ECG) plays a vital role in CVD prevention, diagnosis, and treatment. This is because each electrode in an ECG can reveal cardiac abnormalities, which are risk factors for CVDs.

The main advantage of ECG is that its acquisition is inexpensive and non-invasive. However, the visual interpretation of ECG is tedious, time-consuming, and requires expert knowledge. To address this, the PhysioNet/Computing in Cardiology Challenge 2021 offers a benchmark for automatic classification of cardiac abnormalities from 12-lead and reduced-lead ECGs.

Prior literature on ECG classification exhibits a methodological divide: signal processing versus machine learning. On one hand, digital signal processing methods include low-pass filters, fast Fourier Transform, and wavelet transform. On the other hand, machine learning methods include random forests, support vector machines, convolutional neural networks and long short-term memory (LSTM) networks. While feature engineering lacks flexibility to represent fine-grain class boundaries, a purely learned pipeline may lead to uninterpretable overfitting.

Our contribution to the Challenge aims to overcome the divide by combining insights from signal processing and machine learning. At a first stage, we extract time scattering transform (ST) coefficients for each ECG channel. Although this stage is not trainable, it offers numerical guarantees of stability to time warps. At a second stage, we train a depthwise separable convolution (DSC) network, followed by a bidirectional LSTM network. While DSC combines scattering coefficients from multiple leads simultaneously, the BiLSTM can also capture longer-term trends in cardiac activity. We also investigated transfer learning to the reduced-lead models using canonical correlation analysis (CCA). Our system extends previous Challenge work\(^1\) and is inspired from previous publications, which aimed at detecting sleep arousals from polysomnographic recordings \(^2\).

2. Methods

Figure \(^1\) summarizes our proposed system; this section explains the role of each system component in isolation.
2.1. Scattering transform

The scattering transform is a deep convolutional network whose filters are defined a priori instead of being learned from data. We refer to [3] for a mathematical introduction and to [2] for a recent review of the state of the art. Specifically, every layer contains filters of the form

$$\psi_j : t \mapsto 2^{-j/\xi} \psi(2^{-j/\xi} t),$$

where $\psi$ is a wavelet, $Q$ is a constant number of filters per octave, and the scale variable $j$ is an integer ranging between 0 and $J$. Hereafter, we take the “mother wavelet” $\psi$ to be a Morlet wavelet with a quality factor of $Q = 1$ and a center frequency of $\xi = 200$ Hz. The Morlet wavelet is a complex-valued function with a Gaussian envelope while being approximately analytic, i.e., with negligible Fourier coefficients outside of the half-line of positive frequencies ($\omega > 0$). Furthermore, we set the maximum wavelet scale to $J = 11$ after a process of trial and error.

Let $\phi_T$ be a Gaussian filter of cutoff frequency equal to $1/T$. The first two orders of the scattering transform are

$$S_1 x(t, j_1) = |x \ast \psi_{j_1} \ast \phi_T(t)$$

and

$$S_2 x(t, j_1, j_2) = |x \ast \psi_{j_1} \ast \psi_{j_2} \ast \phi_T(t),$$

where the vertical bars and the asterisk denote complex modulus and convolution product respectively.

For every discretized value of time $t$, we concatenate first-order coefficients $S_1 x(t, j_1)$ and second-order coefficients $S_1 x(t, j_1, j_2)$ to produce a multidimensional time series $S x(t, p)$; where the multindex $p$, known as scattering path, either denotes a singleton $(j_1)$ or a pair $(j_1, j_2)$. With $J = 11$, this results in 12 first-order and 63 second-order paths for a total number of $P = 75$ paths.

To control the degree of time invariance, we modified the Python scattering package Kymatio\(^1\) to set the time scale of Gaussian averaging to $T = 62.5$ ms. Note that this $T$ is less than the customary $2^J/\xi$. Rather, the filterbank $\{\psi_j\}_j$ covers the frequency range $[2^{-J}\xi; \xi] = [0.1 \text{Hz}; 200 \text{Hz}]$ whereas the scattering transform is discretized at a Nyquist rate of $2/T = 32$ Hz. This rate is chosen to be higher than typical patient heart rates yet considerably lower than the ECG acquisition rate (500 Hz).

We apply a pointwise compressive nonlinearity to the output of the ST, namely an offset log function: $\log(x + \epsilon)$ where $\epsilon = 10^{-4}$. Then per-path normalization subtracted the mean and divided by the standard deviation. Figure 2 illustrates the scattering transform of normal and atrial fibrillation ECG recordings, for the first two orders.

2.2. Depthwise separable convolution

A depthwise separable convolution (DSC) splits the computation into two operations: depthwise convolution $X$ linearly combines the leads for each ST path while the pointwise convolution $Y$ linearly combines these transformed paths, as in equations (3) and (4)

$$X[p] = \sum_{l=1}^{L} S[l, p] F[p, l]$$

$$Y[n] = \beta \left[ B[n] + \sum_{p=1}^{P} X[p] G[p, n] \right]$$

where $L \in \{12, 6, 4, 3, 2\}$ and $P$ represent the number of leads and paths, respectively. $F$ and $G$ refer to the filter maps, $N$ is the number of pointwise mixes, $B$ is the bias and $\beta$ represents the activation function. The total number of convolution coefficients including the bias weights is therefore $P \times L + (P + 1) \times N$. This is often a reduction

\(^1\)Official website of Kymatio: https://www.kymat.io
in parameters compared to regular convolution. We used a
DSC layer with $N = P = 66$ (chosen to be on the order
of the number of paths) and ReLU activation.

### 2.3. Transfer learning for reduced-lead models

For reduced-lead models, we apply transfer learning
from the 12-lead data using canonical correlation analy-
sis (CCA). CCA finds a pair of linear transformations for
two sets of multidimensional variables (views $S_i$), such
that the linear projections of the two views, $(S_1 w_1, S_2 w_2)$
are maximally correlated \(^4\). In our case, view $S_i$ is the
scattering of lead sets: $S_1$ corresponds to the lead set used
for prediction (2, 3, 4 or 6 leads) and $S_2$ corresponds to
the respective complements (10, 9, 8 and 6 leads). This is
done by maximizing the following equation:

$$
\rho = \max_{w_1, w_2} \text{corr} (S_1 w_1, S_2 w_2)
= \max_{w_1, w_2} \frac{w_1^T \Sigma_{12} w_2}{\sqrt{w_1^T \Sigma_{11} w_1} \sqrt{w_2^T \Sigma_{22} w_2}} \tag{5}
$$

where $\Sigma_{11}$, $\Sigma_{22}$ and $\Sigma_{12}$ are the covariances and cross-
covariance of $S_1$ and $S_2$; and $w_1$ and $w_2$ are determined
by singular-value decomposition.

We calculate $w_1$ and $w_2$ from fold training data prior to
network training. CCA uses $S_1$ and $S_2$ to find the projec-
tion vectors corresponding to the $k$ highest left- and right-
singular values, and $k = P \times L$ was chosen to include all
the singular values.

During training and prediction, $S_1$ is projected with
fixed $w_1$. This projection is intended to transfer informa-
tion from (possibly unavailable) $S_2$, correlated with
the complementary lead set, such that classification of
reduced-lead ECG records is improved.

### 2.4. Data

The PhysioNet/CinC Challenge 2021 data \(^5\) includes
88,000 public and 26,000 private ECG records. Each
record is assigned one or more diagnosis by experts. We
excluded subsets St. Petersburg having long durations (30
min) and PTB having non-uniform acquisition rates and
low class coverage (5/26).

### 2.5. Implementation

Although the ECG recording lengths in the training
set were as long as 120 s, the vast majority (78,181 of
87,663=89%) were 10 s or less. Therefore to reduce com-
putational requirements, we reduced the time span of the
learning batches to 10 s. Longer recordings were truncated
at 10 s or split into multiple training sub-sequences of 10 s.
We applied a padding target for sub-sequences of duration
less than 10 s to remove their unused samples from partic-
ipation in the loss function. 24 Georgia and 388 Ningbo
records were omitted from training because NaN values in
the ECG recordings prevented convergence.

We used two BiLSTM layers of 100 hidden units. The
dense layer used binary cross-entropy loss to support mul-
tiple classes. Predictions were averaged over time, and
over sub-sequences if present. Our decision rule chose any
class that exceeded probability threshold $p = 0.5$; other-
wise the maximum probability class was chosen.

The 10-fold cross-validation data partitions were 90% 
training and 10% testing for each fold. The validation set,
10% of training, was used for early stopping (60 epochs).
Table 1. Challenge metric (5; 6) for baseline and CCA lead models on cross-validation and hidden validation and test sets (ranking out of 40 teams in parentheses). * indicate two-sided t-test $p < 0.01$ compared to previous row.

<table>
<thead>
<tr>
<th>Model</th>
<th>Cross-validation</th>
<th>Validation</th>
<th>Test</th>
</tr>
</thead>
<tbody>
<tr>
<td>Base$_{12}$</td>
<td>0.601 ± 0.015</td>
<td>0.46 (23)</td>
<td>0.10 (34)</td>
</tr>
<tr>
<td>Base$_{6}$</td>
<td>0.584 ± 0.007*</td>
<td>0.44 (23)</td>
<td>0.11 (33)</td>
</tr>
<tr>
<td>CCA$_{6}$</td>
<td>0.573 ± 0.010</td>
<td>0.45 (21)</td>
<td>0.10 (34)</td>
</tr>
<tr>
<td>Base$_{4}$</td>
<td>0.582 ± 0.015</td>
<td>0.45 (21)</td>
<td>0.10 (34)</td>
</tr>
<tr>
<td>CCA$_{4}$</td>
<td>0.581 ± 0.009</td>
<td>0.45 (21)</td>
<td>0.10 (34)</td>
</tr>
<tr>
<td>Base$_{3}$</td>
<td>0.583 ± 0.006</td>
<td>0.46 (21)</td>
<td>0.10 (34)</td>
</tr>
<tr>
<td>CCA$_{3}$</td>
<td>0.576 ± 0.009</td>
<td>0.46 (21)</td>
<td>0.10 (34)</td>
</tr>
<tr>
<td>Base$_{2}$</td>
<td>0.570 ± 0.008</td>
<td>0.43 (22)</td>
<td>0.10 (34)</td>
</tr>
<tr>
<td>CCA$_{2}$</td>
<td>0.564 ± 0.010</td>
<td>0.43 (22)</td>
<td>0.10 (34)</td>
</tr>
</tbody>
</table>

3. Results

Table 1 shows cross-validation and hidden validation results for our baseline and CCA models using 10 s truncation. Our submitted entry completed training of the baseline models in just over 18 h and prediction of the hidden validation set in 18 min, within the maximum allowable times of 48 h and 24 h, respectively. Fig 3 shows the class incidence in the training data and cross-validation performance for the 12-lead model Base$_{12}$.

4. Discussion

Our approach achieved experimental success without need for feature engineering and with few parameters to select. We observe slight performance degradation for models with decreasing numbers of leads, suggesting that the correlation between leads is considerable. CCA did not significantly improve results but warrants further analysis.

Performance was only somewhat reduced on the hidden validation set compared to cross-validation, indicating good generalization; however hidden test set performance was drastically low in comparison, suggesting that the test set was quite different from the training and validation sets or that there was a systemic error in our processing of the test set. We anticipate that the organizers will provide further information to better assess these results.

Splitting rather than truncating records did not affect results, although benefits may have been masked by the predominance of 10 s recordings.

We note that higher incidence classes tended to perform better, especially normal sinus rhythm (28,891 of 87,663 training records $\approx$ 33%). Considering this imbalance could improve results for low incidence classes.

Extensions to our approach to explore include: improving the decision rule; exploring alternate loss functions; searching hyperparameters; and using age and sex demographic data, recognized risk factors for cardiac pathology.

Acknowledgements

We acknowledge resources provided by the Swedish National Infrastructure for Computing and PeriGen Inc.

References


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