Atrial Fibrillation Classification in Electrocardiography using Deep Learning

Christoph Reich[‡]

Department of Electrical Engineering and Information Technology,
Technische Universität Darmstadt

†christoph.reich@bcs.tu-darmstadt.de

Abstract—The evaluation of electrocardiogram recordings is the most common approach to diagnose and monitor cardiac arrhythmia such as atrial fibrillation. The electrocardiogram evaluation typically requires expert knowledge, which is not always available. We present a novel approach for the automated classification of atrial fibrillation in electrocardiogram recordings with variable length. Our deep learning approach utilizes both input data in the time and frequency domain. The performance of the proposed dual network for ECG classification (ECG-DualNet) is showcased on the 2017 PhysioNet/CinC Challenge dataset. ECG-DualNet outperforms recent Convolutional Neural Network approaches in terms of classification accuracy. Code and trained models are available at https://github.com/ChristophReich1996/ECG_Classification.

Index Terms—deep learning, attention, arrhythmia classification, atrial fibrillation classification, electrocardiography.

I. INTRODUCTION

Electrocardiography (ECG) is the most important tool for the diagnosis and the monitoring of cardiac arrhythmia [1]–[3]. The first recorded human heartbeat dates back to the late 19th century [3]. Today 12-lead ECGs are the common standard [3]. The analysis of ECG recordings, especially the detection of cardiac arrhythmia [1]. However, in recent years edge devices like smartwatches became popular. These devices most often include a single-led ECG sensor. Analyzing such signals require expert knowledge, which is typically not available in an edge device settings. Since a fast and accurate diagnosis of cardiac arrhythmia in single-led ECG signals can positively affect a patient's chance of survival, an increasing interest in the automated detection of cardiac arrhythmia occurs [1], [4]–[6].

The most common human cardiac arrhythmia is atrial fibrillation (AF) (Fig. 1) [7]. AF mostly affects patients at an advanced age and can lead to an increased risk for dementia, stroke, and even heart failure. High blood pressure and obesity are common factors for an increased risk for AF. During AF undirected electoral impulses (Fig. 1) occur in the atrium of the heart. This leads to a fast movement of the heart atrium resulting in an impaired blood flow. Additionally, blood clots can occur which potentially lead to thrombosis. [1], [7]

In this study, we propose a novel deep learning approach for classifying AF in single-lead ECG recordings with variable lengths. Our novel deep learning approach ECG-DualNet utilized both input data for the time and frequency domain, enabling the network to learn features in both domains. The

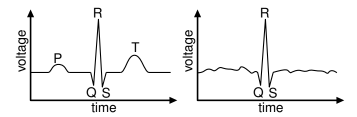


Fig. 1. Regular heart beat with QRS complex, P wave, and T wave on the left and atrial fibrillation on the right. AF can be detected by the noisy ECG at the P and T wave regions. Please note that this is only a theoretical illustration. Real ECGs of AF can be seen in the appendix.

proposed ECG-DualNet surpasses the classification accuracy of recent convolutional neural network (CNN) approaches which only utilize input data from the frequency domain [5].

II. RELATED WORK

Recent approaches for the task of AF classification in ECG recordings can be clustered into two groups. First classical machine learning approaches that typically extract features, first and classify them in a section learnable step [4], [8]. Hoog Antink *et al.* [4] proposed an approach that first extracts ECG timing features, robust interval features, and waveform features [4]. All extracted features are fed into a learned random forest for classification [4]. Other approaches perform similar feature extractions but utilize different learnable classification methods, such as support vector machines [8]. These approaches, however, require a lot of domain knowledge to extract relevant features. Additionally, hand-crafted feature extraction approaches are often complicated and error-prone to implement.

Recently deep learning-based approaches have been applied to AF classification in ECG recordings [5], [9]–[12]. These approaches typically produce a spectrogram of the ECG recording and feed it into a deep neural network for classification [5], [10]. Deep learning AF classification approaches tend to outperform more classical machine learning approaches [9], [11]. Using deep neural networks, however, require a reasonable amount of data, or extensive data augmentation [13], and has higher computational requirements [14], [15]. Since a deep neural network is able to learn the extraction of relevant features, heavy preprocessing is typically not required [5], [14], [15].

III. METHOD

In this section, we introduce our ECG-DualNet (Fig. 2). We take inspiration from the model by Mousavi *et al.* which utilizes two separate network paths to encode both time and frequency domain data [9]. Our data augmentation pipeline builds on the work of Nonaka *et al.* which presented multiple augmentations for ECG data [12].

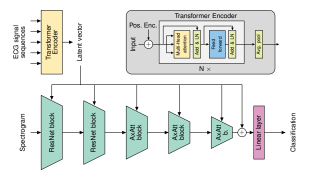


Fig. 2. ECG-DualNet++ architecture with spectrogram and ECG signal as inputs. The ECG signal sequence gets encoded by a Transformer encoder to a single latent vector. The spectrogram is encoded by multiple 2D blocks. The first two blocks are standard ResNet like blocks and the following three blocks are Axial-Attention blocks. All blocks of the spectrogram encoder utilize the latent vector by conditional batch normalization. Transformer encoder [16] architecture shown in the top right.

A. ECG-DualNet Architecture

Our ECG-DualNet (Fig. 2) utilizes two separate encoders. The first encoder takes the ECG signal as the input. We will refer to this encoder as the signal encoder. The second encoder is fed with a spectrogram. We refer to this encoder as the spectrogram encoder. The final part of the network builds a simple softmax classifier to produce the final classification. We utilize two network settings, ECG-DualNet and ECG-DualNet++ in which we use different building blocks for both encoders.

In the standard-setting (ECG-DualNet), the signal encoder consists of a standard long-short-term-memory (LSTM) [17] module. The LSTM module encodes temporal agnostic feautres into a latent vector. This latent vector is incorporated into the spectrogram encoder and the final linear layer. In the ECG-DualNet++ we replace the LSTM module with a Transformer encoder [16], [18] (Fig. 2 top right), using learnable encodings [16], [19], Layer Normalization [20], and Gaussian Error Linear Units [21]. To encounter overfitting dropout [22] is used in both the LSTM and the Transformer.

The spectrogram encoder is, in the standard setting, comprised of five ResNet-like [23] blocks. Each block consists of two 2D convolutions, two Padé Activation Units [24] two Conditional Batch Normalization layers (CBN) [25], an average pooling layer, and a skip connection. CBN is utilized to conditionalize the spectrogram encoder on the latent vector of the signal encoder. For ECG-DualNet++ the three highest blocks of the spectrogram encoder are replaced with Axial-Attention blocks [26]. Still, CBN and Padé Activation Units are employed. Similar to the signal encoder, dropout [22] is

also applied in each spectrogram encoder block.

To investigate the effect of the network size we employ different network sizes for both the ECG-DualNet and ECG-DualNet++. We vary the width and the depth of the signal encoder. For the spectrogram encoder, we diversify the width.

B. Training Approach

We train all networks on a weighted version of the crossentropy loss [15]

$$\mathcal{L} = -\frac{1}{N} \sum_{j=1}^{N} \sum_{i=1}^{4} \alpha_i \, y_{ji} \, \log(\hat{y}_{ji}). \tag{1}$$

Where $\mathbf{y}_j \in \mathbb{R}^4$ is the ground truth one-hot label, $\hat{\mathbf{y}}_j \in \mathbb{R}^4$ the network softmax prediction, and $\alpha \in \mathbb{R}^4$ the class weighting. The cross-entropy loss is averaged over a minibatch of the size N. The loss function (Eq. 1) is minimized by using the RAdam optimizer [27].

C. Validation Approach

To validate the performance of our networks we utilize the accuracy and the F1 score. The accuracy is computed over all classes by

$$ACC = \frac{1}{n} \sum_{j=1}^{n} \delta\left(\arg\max(\mathbf{y}_j), \arg\max(\hat{\mathbf{y}}_j)\right).$$
 (2)

Where $\delta(\cdot,\cdot)$ is the Kronecker delta, and $\arg\max(\cdot)$ estimates the position of the maximum value present in the input vector. n corresponds to the dataset size. The F1 score is computed as

$$F1 = \frac{1}{4} \sum_{i=1}^{4} \frac{2TP_i}{2TP_i + FP_i + FN_i}.$$
 (3)

 TP_i represents the true positive predictions of a class over the whole dataset, FP_i the false positive predictions, and FN_i the false-negative predictions.

D. Preprocessing

We utilize a simple preprocessing composed of four steps. In the first steps, the ECG signal gets standardization to a mean of zero and unit variance. In the second step, various augmentations (Sec. III-E) are applied to the ECG signal. In the following step, a log spectrogram of the ECG signal is produced. The log spectrogram is computed with a window length of 64, a hop size of 32, and 64 bins. Recent work showed, using the logarithmic spectrogram improves the classification accuracy of CNN's [5]. Finally, both the ECG signal and the spectrogram are zero-padded to a fixed length.

Our augmentation pipeline applies randomly multiple different data augmentations to the ECG signal. This improved the generalization of the trained network and prevents overfitting [12], [13], [28]. The following augmentations are used: dropping, cut-out, resampling, random resampling, scaling, shifting, sine addition, and bandpass filtering. The dropping augmentation sets random samples of the ECG signal to zero, the cut-out augmentation sets a random sequence to zero. In the resampling augmentation, the signal gets resampled to a different heartbeat rate. Random resampling is inspired by the random elastic deformation [29]-[31] used for image augmentation. The ECG signal gets resampled by smooth random offset, resulting in a changing heartbeat rate. In the scaling augmentation, the signal gets scaled by a random factor. The sine addition augmentation adds a sinusoidal signal with a random magnitude and phase to the ECG signal. The shift augmentation shifts the ECG signal by a random length. Finally, in the band pass filter augmentation, the ECG signal is filtered by a band-pass.

F. Implementation Details

ECG-DualNet is implemented in PyTorch [32]. For implementing the preprocessing SciPy [33], NumPy [34], and Torchaudio were used, in addition. For the Padé Activation Unit, we used the implementation by the authors [24].

Different network sizes (S, M, L, and XL) for both ECG-DualNet and ECG-DualNet++ (Tab. I) were employed. A detailed overview of the network configurations can be found in the appendix or in the provided implementation.

100 training epochs with a fixed batch size of 24 were performed. All models except ECG-DualNet++ 130M were trained on a single Nvidia 2080 Ti. Training took between 30min (ECG-DualNet S) and 3h (ECG-DualNet++ XL). Our biggest model ECG-DualNet++ 130M was trained on four Nvidia Tesla V100 (16GB) which took approximately 6h. The weights of the loss function (Eq. 1) were set to $\alpha = [0.4, 0.7, 0.9, 0.9]$, for counteracting the dataset class imbalance (Sec. IV-A). The initial learning rate of the RAdam optimizer was set to 10^{-3} . The learning rate were decreased after 25, 50, and 75 epochs by 0.1. The first and second-order momentum factors were set to 0.9 and 0.999, respectively. Each augmentation described in Section III-Ewas applied with a probability of 0.2. An overview of all hyperparameters is presented in the appendix.

We also consider pre-training on the Icentia11k dataset [35]. Similar to pre-training in computer vision [36]–[38], we first train on the very large Icentia11k dataset. Afterward, the pre-trained weights were used as the initial weights for training on the target PhysioNet dataset. For pre-training, we perform 20 epochs with a batch size of 100 on a single Nvidia Tesla V100 (32GB). Training took approximately 24h. During pre-training, the same learning rate schedule as described earlier with steps after 5, 10, and 15 epochs was employed. The training on the target dataset was performed as a normal training run on the PhysioNet dataset, described earlier.

IV. EXPERIMENTS

We conduct experiments on the 2017 PhysioNet/CinC Challenge dataset [6] (Sec. IV-A). We performed three training runs (if not stated differently) for each model with different random seeds, where the best run, based on the validation accuracy, was reported. Additional results can be found in the appendix.

A. 2017 PhysioNet/CinC Challenge Dataset

We utilize the 2017 PhysioNet/CinC Challenge dataset [6] for training. The dataset includes 8529 publicly available labeled single-lead ECG sequences. Each ECG sequence includes between 2714 and 18286 samples, recorded with a sampling frequency of 300. For each sequence, a ground truth classification label is provided. Four classes are given, namely, normal, indicating a normal cardiac rhythm, AF, indicating atrial fibrillation, other, for different rhythms, and noisy, for a noisy measurement. Each class includes 5050 (normal), 2456 (AF), 738 (other), and 284 (noisy) samples, respectively. Visualizations of multiple sequences and corresponding labels are provided in the appendix.

We split the data once randomly into a training and validation set. The training set includes 7000 samples and the validation set 1528 samples. We omit the use of k-fold cross-validation due to the computational expensiveness of each training run [15], [39]. All sequences are zero-padded, respectively, cropped to a fixed length of 18000 samples. For a detailed description of the preprocessing see Section III-D.

B. Icentia11k dataset

The Icentia11k dataset [35], used for pre-training, consists of single-lead ECG recordings of 11k patients. The dataset is partly labeled with six different heart rhythms including atrial fibrillation. Each of the 550k dataset sample is a sequences of approximately 1h with a sampling frequency of 250. We resample the ECG signal to match the sampling frequency of the PhysioNet dataset. Finally, we randomly crop each sequence to a random length of 9000 to 18000 samples. For unlabeled crops, we utilize a dummy class. For validation, we split the dataset patient-wise. The resulting training set includes 10k patients and the validation set 1k patients.

C. Results

Our archived classification results are presented in Table I, in which we present both the accuracy (Eq. 2) and the F1 score (Eq. 3). Table I also includes classification results of Zihlmann *et al.* [5]. The models trained by Zihlmann *et al.* are, however, trained on the publicly available 2017 PhysioNet/CinC Challenge dataset and tested on the private test dataset [5]. Our models are trained on a custom split of the publically available samples (Sec. IV-A). Additionally, the reported F1 scores by [5] are computed over three classes (excluding the noise class) and thus are not directly comparable to our F1 scores.

Although training was performed with fewer data samples, all ECG-DualNet architectures outperformed the baselines from the literature in classification accuracy. We observe that

TABLE I
CLASSIFICATION RESULTS OF OUR PROPOSED APPROACHES AND
BASELINES ON THE 2017 PHYSIONET VALIDATION SET.

Model	ACC ↑	F1 ↑	# Parameters
CNN baseline* [5]	0.812	0.790^{\dagger}	$\sim 3.5 \mathrm{M}$
CRNN baseline* [5]	0.823	0.792^{\dagger}	$\sim 3.5 \mathrm{M}$
ECG-DualNet S	0.8527	0.8049	1.8M
ECG-DualNet M	0.8560	0.7938	4.3M
ECG-DualNet L	0.8514	0.8038	6.2M
ECG-DualNet XL	0.8612	0.8164	20.7M
ECG-DualNet++ S	0.8174	0.7291	1.8M
ECG-DualNet++ M	0.8259	0.7730	2.6M
ECG-DualNet++ L	0.8449	0.7859	3.7M
ECG-DualNet++ XL	0.8593	0.8051	8.2M
ECG-DualNet++ 130M	0.8534	0.7963	128M

^{*} Reported literature values (private PhysioNet test set utilized).

larger network configurations tend to perform stronger compared to smaller configurations. Especially, ECG-DualNet++ profits from a larger network capacity.

Training on the Icentia11k dataset [35] yield the results presented in Table II. Only two models have trained on the Icentia11k dataset due to the immense computational requirements.

TABLE II

CLASSIFICATION RESULTS OF OUR PROPOSED APPROACHES ON THE ICENTIA11K VALIDATION SET. ONLY A SINGLE TRAINING FOR EACH MODEL RUN WAS CONDUCTED.

Model	ACC ↑	F1 ↑
ECG-DualNet XL	0.8989	0.5135
ECG-DualNet++ XL	0.8899	0.5017

Using the model weights trained on the Icentia11k dataset (Tab. II) for training on the PhysioNet dataset leads to the results presented in Table VII. Compared to the results, using no pre-trained weights (Tab. VI), the results, with pre-trained weights, lead to similar or slightly worse results.

TABLE III

CLASSIFICATION RESULTS OF OUR PROPOSED APPROACHES ON THE 2017
PHYSIONET VALIDATION SET AND PRE-TRAINED ON THE ICENTIA11K
DATASET. DIFFERENCES TO VALUES IN TAB. I IN RED.

Model	ACC ↑	F1 ↑
ECG-DualNet XL	0.8468 (↓ 0.0144)	0.8014 (\psi 0.0150)
ECG-DualNet++ XL	$0.8481 (\downarrow 0.0112)$	0.7817 (\psi 0.0234)

Additional experimental results of each training run are presented in the appendix.

D. Ablation Study

We conducted multiple ablation training runs to investigate the effectiveness of each of the proposed components. In the presented results in table IV, we omit one core component of our approach in each training run. We only perform ablations with one network configuration, namely ECG-DualNet M. This is due to the large computational requirements for each training run.

TABLE IV
CLASSIFICATION RESULTS ON THE 2017 PHYSIONET VALIDATION FOR DIFFERENT ABLATIONS. ECG-DUALNET L CONFIGURATION UTILIZED.

Data aug. & dropout	Signal encoder	Spectrogram encoder	ACC ↑	F1 ↑
Х	✓	✓	0.8272	0.7493
✓	×	✓	0.8440	0.7855
✓	✓	X	0.7264	0.5813
✓	✓	✓	0.8560	0.7938

When utilizing no data augmentation (Sec. III-E) and dropout the training loss is minimized to approximately zero. This indicates signs of overfitting, even though the achieved accuracy is 0.8272.

From the results in table 1, it can be observed that each main component is required to reach the best classification performance. The spectrogram encoder has the largest impact on the classification accuracy. The use of data augmentation and a signal encoder has a lesser impact on the performance.

V. DISCUSSION

The conducted experiments showcased that classical overfitting does not occur. Data augmentation and dropout lead to a better generalization. Omitting the respective methods results in a weaker classification performance but not in strong overfitting. This holds for all ECG-DualNet variants, even for the larges ECG-DualNet++ 130M, which includes significantly more parameters as data samples are in the 2017 PhysioNet dataset. This observation may description by the double decent phenomenon [40], [41]. A large-scale study with more network sizes may help to understand this behavior.

Vice versa to the current findings in computer vision [36]—[38] and natural language processing [42], [43], large-scale pre-training on the Icentia11k dataset does not lead to better classification results after fine-tuning on the target 2017 PhysioNet dataset. A large difference in the dataset domains can presumably be one reason for this behavior. However, a full explanation of this behavior can not be provided.

In the conducted ablation studies we showcased that all core components of the proposed pipeline are important to achieve a competitive classification performance. The most significant core component for a strong classification performance is the spectrum encoder. This indicates that the spectrum is a very suitable representation to learn a deep network from.

VI. CONCLUSION

In this study we presented a novel deep neural network architecture for classifying arrhythmia's in single-lead ECG signals with variable lengths. The proposed ECG-DualNet(++) employs input data from both the time and frequency domain. In the task of atrial fibrillation classification, on the 2017 PhysioNet dataset, ECG-DualNet outperforms recent CNN approaches. Additionally, extensive pre-training on the Icentia11k dataset was performed. This pre-training does not lead to improvements in classification performance on the target dataset.

[†] F1 score computed over three classes, thus not directly comparable.

REFERENCES

- [1] D. E. Becker, "Fundamentals of electrocardiography interpretation," *Anesthesia progress*, vol. 53, no. 2, pp. 53–64, 2006.
- [2] R. H. Anderson, E. J. Baker, A. Redington, M. L. Rigby, D. Penny, and G. Wernovsky, *Paediatric Cardiology*. Elsevier Health Sciences, 2009.
- [3] M. AlGhatrif and J. Lindsay, "A brief review: history to understand fundamentals of electrocardiography," *Journal of community hospital* internal medicine perspectives, vol. 2, no. 1, p. 14383, 2012.
- [4] C. Hoog Antink, S. Leonhardt, and M. Walter, "Fusing QRS Detection and Robust Interval Estimation with a Random Forest to Classify Atrial Fibrillation," in 2017 Computing in Cardiology (CinC). IEEE, 2017, pp. 1–4.
- [5] M. Zihlmann, D. Perekrestenko, and M. Tschannen, "Convolutional Recurrent Neural Networks for Electrocardiogram Classification," in 2017 Computing in Cardiology (CinC). IEEE, 2017, pp. 1–4.
- [6] G. D. Clifford, C. Liu, B. Moody, H. L. Li-wei, I. Silva, Q. Li, A. Johnson, and R. G. Mark, "AF Classification from a Short Single Lead ECG Recording: the PhysioNet/Computing in Cardiology Challenge 2017," in 2017 Computing in Cardiology (CinC). IEEE, 2017, pp. 1–4.
- [7] G. Herold, Innere Medizin. Walter de Gruyter GmbH & Co KG, 2019.
- [8] R. Smíšek, J. Hejč, M. Ronzhina, A. Němcová, L. Maršánová, J. Chmelík, J. Kolářová, I. Provazník, L. Smital, and M. Vítek, "Svm based ecg classification using rhythm and morphology features, cluster analysis and multilevel noise estimation," in 2017 Computing in Cardiology (CinC), 2017, pp. 1–4.
- [9] S. Mousavi, F. Afghah, A. Razi, and U. R. Acharya, "ECGNET: Learning Where to Attend for Detection of Atrial Fibrillation with Deep Visual Attention," in 2019 IEEE EMBS International Conference on Biomedical & Health Informatics (BHI). IEEE, 2019, pp. 1–4.
- [10] F. R. Mashrur, A. D. Roy, and D. K. Saha, "Automatic identification of arrhythmia from ecg using alexnet convolutional neural network," in 2019 4th International Conference on Electrical Information and Communication Technology (EICT). IEEE, 2019, pp. 1–5.
- [11] L. Khriji, M. Fradi, M. Machhout, and A. Hossen, "Deep Learning-based Approach for Atrial Fibrillation Detection," in *International Conference* on Smart Homes and Health Telematics. Springer, 2020, pp. 100–113.
- [12] N. Nonaka and J. Seita, "Data Augmentation for Electrocardiogram Classification with Deep Neural Network," preprint arXiv:2009.04398, 2020
- [13] L. Perez and J. Wang, "The Effectiveness of Data Augmentation in Image Classification using Deep Learning," preprint arXiv:1712.04621, 2017
- [14] Y. LeCun, Y. Bengio, and G. Hinton, "Deep learning," nature, vol. 521, no. 7553, pp. 436–444, 2015.
- [15] I. Goodfellow, Y. Bengio, A. Courville, and Y. Bengio, *Deep Learning*. MIT press Cambridge, 2016, vol. 1, no. 2.
- [16] A. Vaswani, N. Shazeer, N. Parmar, J. Uszkoreit, L. Jones, A. N. Gomez, L. u. Kaiser, and I. Polosukhin, "Attention is all you need," in *Advances in Neural Information Processing Systems*, vol. 30, 2017.
- [17] S. Hochreiter and J. Schmidhuber, "Long Short-Term Memory," *Neural Comput.*, vol. 9, no. 8, p. 1735–1780, Nov. 1997.
- [18] A. Dosovitskiy, L. Beyer, A. Kolesnikov, D. Weissenborn, X. Zhai, T. Unterthiner, M. Dehghani, M. Minderer, G. Heigold, S. Gelly et al., "An image is worth 16x16 words: Transformers for image recognition at scale," preprint arXiv:2010.11929, 2020.
- [19] T. Prangemeier, C. Reich, and H. Koeppl, "Attention-Based Transformers for Instance Segmentation of Cells in Microstructures," in 2020 IEEE International Conference on Bioinformatics and Biomedicine (BIBM), 2020, pp. 700–707.
- [20] J. L. Ba, J. R. Kiros, and G. E. Hinton, "Layer Normalization," preprint arXiv:1607.06450, 2016.
- [21] D. Hendrycks and K. Gimpel, "Gaussian Error Linear Units (GELUs)," preprint arXiv:1606.08415, 2016.
- [22] N. Srivastava, G. Hinton, A. Krizhevsky, I. Sutskever, and R. Salakhutdinov, "Dropout: A Simple Way to Prevent Neural Networks from Overfitting," *Journal of Machine Learning Research*, vol. 15, no. 56, pp. 1929–1958, 2014.
- [23] K. He, X. Zhang, S. Ren, and J. Sun, "Deep Residual Learning for Image Recognition," in *Proceedings of the IEEE conference on computer vision* and pattern recognition, 2016, pp. 770–778.
- [24] A. Molina, P. Schramowski, and K. Kersting, "Padé Activation Units: End-to-end Learning of Flexible Activation Functions in Deep Networks," in *International Conference on Learning Representations*, 2020.

- [25] H. de Vries, F. Strub, J. Mary, H. Larochelle, O. Pietquin, and A. Courville, "Modulating early visual processing by language," in Advances in Neural Information Processing Systems, 2017, pp. 6597– 6607
- [26] H. Wang, Y. Zhu, B. Green, H. Adam, A. Yuille, and L.-C. Chen, "Axial-DeepLab: Stand-Alone Axial-Attention for Panoptic Segmentation," in European Conference on Computer Vision, 2020, pp. 108–126.
- [27] L. Liu, H. Jiang, P. He, W. Chen, X. Liu, J. Gao, and J. Han, "On the Variance of the Adaptive Learning Rate and Beyond," in *International Conference on Learning Representations*, 2020.
- [28] F. N. Hatamian, N. Ravikumar, S. Vesal, F. P. Kemeth, M. Struck, and A. Maier, "The Effect of Data Augmentation on Classification of Atrial Fibrillation in Short Single-Lead ECG Signals Using Deep Neural Networks," in ICASSP 2020 2020 IEEE International Conference on Acoustics, Speech and Signal Processing (ICASSP), 2020, pp. 1264–1268.
- [29] P. Y. Simard, D. Steinkraus, J. C. Platt *et al.*, "Best Practices for Convolutional Neural Networks Applied to Visual Document Analysis," in *Icdar*, vol. 3, no. 2003, 2003.
- [30] O. Ronneberger, P. Fischer, and T. Brox, "U-Net: Convolutional Networks for Biomedical Image Segmentation," in *International Conference on Medical image computing and computer-assisted intervention*. Springer, 2015, pp. 234–241.
- [31] T. Prangemeier, C. Wildner, A. O. Françani, C. Reich, and H. Koeppl, "Multiclass Yeast Segmentation in Microstructured Environments with Deep Learning," in 2020 IEEE Conference on Computational Intelligence in Bioinformatics and Computational Biology (CIBCB), 2020, pp. 1–8.
- [32] A. Paszke, S. Gross, F. Massa, A. Lerer, J. Bradbury, G. Chanan, T. Killeen, Z. Lin, N. Gimelshein, L. Antiga, A. Desmaison, A. Kopf, E. Yang, Z. DeVito, M. Raison, A. Tejani, S. Chilamkurthy, B. Steiner, L. Fang, J. Bai, and S. Chintala, "PyTorch: An Imperative Style, High-Performance Deep Learning Library," in Advances in Neural Information Processing Systems, vol. 32, 2019.
- [33] P. Virtanen, R. Gommers, T. E. Oliphant, M. Haberland, T. Reddy, D. Cournapeau, E. Burovski, P. Peterson, W. Weckesser, J. Bright et al., "SciPy 1.0: fundamental algorithms for scientific computing in Python," Nature methods, vol. 17, no. 3, pp. 261–272, 2020.
- [34] C. R. Harris, K. J. Millman, S. J. van der Walt, R. Gommers, P. Virtanen, D. Cournapeau, E. Wieser, J. Taylor, S. Berg, N. J. Smith *et al.*, "Array programming with NumPy," *Nature*, vol. 585, no. 7825, pp. 357–362, 2020.
- [35] S. Tan, G. Androz, A. Chamseddine, P. Fecteau, A. Courville, Y. Bengio, and J. P. Cohen, "Icentia11k: An unsupervised representation learning dataset for arrhythmia subtype discovery," preprint arXiv:1910.09570, 2019.
- [36] M. D. Zeiler and R. Fergus, "Visualizing and Understanding Convolutional Networks," in *European conference on computer vision*, 2014, pp. 818–833.
- [37] R. Girshick, J. Donahue, T. Darrell, and J. Malik, "Rich Feature Hierarchies for Accurate Object Detection and Semantic Segmentation," in 2014 IEEE Conference on Computer Vision and Pattern Recognition, 2014, pp. 580–587.
- [38] K. He, R. Girshick, and P. Dollar, "Rethinking imagenet pre-training," in *Proceedings of the IEEE/CVF International Conference on Computer Vision (ICCV)*, October 2019.
- [39] C. M. Bishop, Pattern Recognition and Machine Learning. Springer, 2006
- [40] M. Belkin, D. Hsu, S. Ma, and S. Mandal, "Reconciling modern machine-learning practice and the classical bias-variance trade-off," *Proceedings of the National Academy of Sciences*, vol. 116, no. 32, pp. 15849–15854, 2019.
- [41] P. Nakkiran, G. Kaplun, Y. Bansal, T. Yang, B. Barak, and I. Sutskever, "Deep double descent: Where bigger models and more data hurt," in International Conference on Learning Representations, 2020.
- [42] J. Devlin, M.-W. Chang, K. Lee, and K. Toutanova, "BERT: Pre-training of Deep Bidirectional Transformers for Language Understanding," in Proceedings of the 2019 Conference of the North American Chapter of the Association for Computational Linguistics: Human Language Technologies, Volume 1 (Long and Short Papers), 2019, pp. 4171–4186.
- [43] T. Brown, B. Mann, N. Ryder, M. Subbiah, J. D. Kaplan, P. Dhariwal, A. Neelakantan, P. Shyam, G. Sastry, A. Askell, S. Agarwal, A. Herbert-Voss, G. Krueger, T. Henighan, R. Child, A. Ramesh, D. Ziegler, J. Wu, C. Winter, C. Hesse, M. Chen, E. Sigler, M. Litwin, S. Gray, B. Chess,

J. Clark, C. Berner, S. McCandlish, A. Radford, I. Sutskever, and D. Amodei, "Language Models are Few-Shot Learners," in *Advances in Neural Information Processing Systems*, vol. 33, 2020, pp. 1877–1901.

VII. APPENDIX

This section includes additional information and results of this study work.

A. Dataset

This subsection includes visualizations and additional information of the 2017 PhysioNet/CinC Challenge dataset [6].

TABLE V CLASS DISTRIBUTION OF THE 2017 PHYSIONET/CINC CHALLENGE DATASET [6].

Normal	Other rhythm	AF	Noisy
5050	2456	738	284

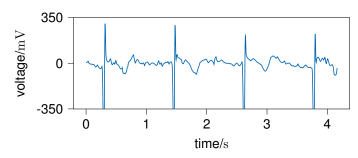


Fig. 3. ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] labeled as normal.

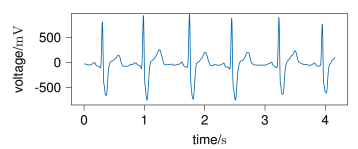


Fig. 4. ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] labeled as other rhythm.

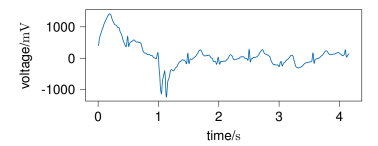


Fig. 5. ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] labeled as noisy.

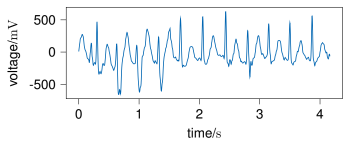


Fig. 6. ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] labeled as AF.

B. Augmentation Pipeline

This section includes visualization of the each augmentation included in our augmentation pipeline.

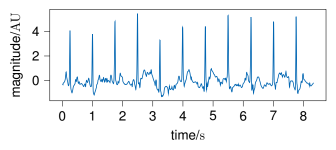


Fig. 7. Standardized ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] processed by scaling augmentation.

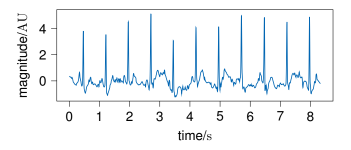


Fig. 8. Standardized ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] processed by shift augmentation.

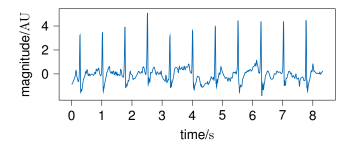


Fig. 9. Standardized ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] processed by band pass filter augmentation.

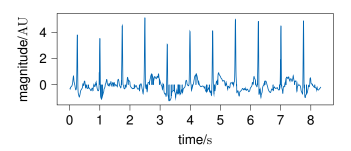


Fig. 10. Standardized ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] processed by dropping augmentation.

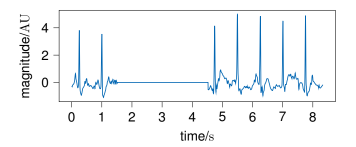


Fig. 11. Standardized ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] processed by cutout augmentation.

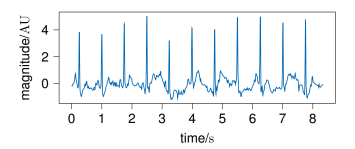


Fig. 12. Standardized ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] processed by sine augmentation.

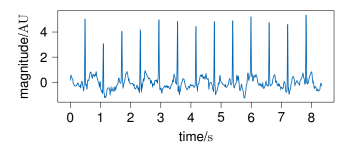


Fig. 13. Standardized ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] processed by resample augmentation.

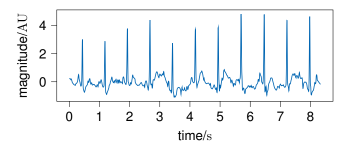


Fig. 14. Standardized ECG signal of the 2017 PhysioNet/CinC Challenge dataset [6] processed by random resample augmentation.

C. Failed Experiments

We experimented also with a full transformer-based architecture [16]. In which the ECG signal builds the input to the transformer encoder. The transformer decoder was feed with patches of the spectrogram. Unfortunately, this architecture achieved a weak classification accuracy (< 0.72) during early tests. Because of this, a full transformer-based architecture was not considered further.

D. Experimental Results

TABLE VI
CLASSIFICATION RESULTS OF ECG-DUALNET(++) ON THE PHYSIONET 2017 VALIDATION SET FOR THREE DIFFERENT TRAINING RUNS WITH DIFFERENT SEEDS.

	Rur	n 1	Rur	n 2	Rur	1 3
Model	ACC	F1	ACC	F1	ACC	F1
ECG-DualNet S	0.8527	0.8049	0.8410	0.7923	0.8455	0.7799
ECG-DualNet M	0.8560	0.7938	0.8442	0.7955	0.8495	0.7928
ECG-DualNet L	0.8508	0.8097	0.8213	0.7515	0.8514	0.8038
ECG-DualNet XL	0.7702	0.6899	0.8612	0.8164	0.7866	0.7162
ECG-DualNet++ S	0.7323	0.6239	0.8174	0.7291	0.7912	0.7127
ECG-DualNet++ M	0.8226	0.7544	0.8259	0.7730	0.7938	0.6947
ECG-DualNet++ L	0.8449	0.7859	0.8442	0.7750	0.8396	0.7671
ECG-DualNet++ XL	0.8593	0.8051	0.8051	0.7799	0.8501	0.7851
ECG-DualNet++ 130M	0.8475	0.7878	0.8534	0.7963	0.8462	0.7740

E. Challenge Submission

Our final challenge submission includes an ECG-DualNet XL. This model was pre-trained on the Icentia11k dataset [35] and fine-tuned on the 2017 PhysioNet/CinC Challenge

TABLE VII

CLASSIFICATION RESULTS OF ECG-DUALNET(++) ON THE PHYSIONET 2017 VALIDATION SET FOR THREE DIFFERENT TRAINING RUNS WITH PRE-TRAINED WEIGHTS (ICENTIA11K DATASET) AND DIFFERENT SEEDS.

	Rur	Run 1		Run 2		Run 3	
Model	ACC	F1	ACC	F1	ACC	F1	
ECG-DualNet XL	0.8534	0.8024	0.8167	0.7385	0.7663	0.5880	
ECG-DualNet++ XL	0.8534	0.8004	0.8534	0.7844	0.8508	0.7952	

dataset [6]. However, instead of the random training and validation split used in the reported results, an optimized split is used. Since it is known that a fraction of the final test set is taken from the publicly available samples of the 2017 PhysioNet/CinC Challenge dataset we detected these samples with the provided samples. Samples that are not included in the provided dataset but are included in the 2017 PhysioNet/CinC Challenge dataset are put into the training set. The final split includes 8000 training samples (including potential test data) and 528 samples to validate the network's performance during training. The achieved validation results are presented in Table VIII.

TABLE VIII

CLASSIFICATION RESULTS (4 CLASS) OF ECG-DUALNET XL PRE-TRAINED ON THE ICENTIA11K DATASET AND FINE-TUNED ON THE PHYSIONET DATASET WITH OPTIMIZED SUBMISSION SPLIT. METRIC COMPUTED ON THE SMALL VALIDATION SET.

Model	ACC ↑	F1 ↑
ECG-DualNet XL (4 class)	0.8840	0.8549

We also consider the two class case where the network only has to distinguish between a normal rhythm and AF. We set samples labeled as other rhythm and noisy to normal rhythm. The same split as in the four-class case is used. The following results were achieved (Tabl. IX).

TABLE IX

CLASSIFICATION RESULTS (2 CLASS) OF ECG-DUALNET XL PRE-TRAINED ON THE ICENTIA11K DATASET AND FINE-TUNED ON THE PHYSIONET DATASET WITH OPTIMIZED SUBMISSION SPLIT. METRIC COMPUTED ON THE SMALL VALIDATION SET.

 Model
 ACC ↑
 F1 ↑

 ECG-DualNet XL (2 class)
 0.9800
 0.9288

F. Hyperparameters

All important network and augmentation hyperparameters can be seen in the provided config file at: https://github.com/ChristophReich1996/ECG_Classification/blob/main/ecg_classification/config.py.

Additional hyperparameters can be found in the training script at: https://github.com/ChristophReich1996/ECG_Classification/blob/main/train.py