Master Thesis

Combining Multi-scale Kernels and Transformer Encoder for multi-label ECG classification

Kilian Laurin Kramer

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Thesis Committee:

Prof Dr. Pietro Bonizzi Prof Dr. Stef Zeemering Prof. Dr. Joël Karel

Maastricht University Faculty of Science and Engineering Department of Advanced Computing Sciences

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Abstract

The accurate classification of electrocardiogram (ECG) signals is crucial for diagnosing various cardiovascular diseases, such as Atrial Fibrillation and to provide cardiologists with reliable predictions. With recent advances in deep learning, Transformer models [1] have emerged as powerful tools for the accurate single and multi-lead ECG classification of Atrial Fibrillation [3] [5] [9] [19] [22] [26] [45] [48]. This thesis investigates the potential of applying Transformer-based models to single-lead ECG classification and provides a comparison with non deep learning (based on extracted features) and other deep learning based models, such as Convolutional Networks.

Chapter 1

Introduction

With recent advances in deep learning models, Transformer-based architectures have emerged as powerful tools for accurate single-lead and multi-lead ECG classification [3] [5] [9] [22] [19] [26] [45] [48]. Multi-lead ECGs provide a view of the heart's electrical activity from different angles and can localise abnormalities more reliable. It is a standard in clinical monitoring. Accurate single-lead ECG classification is an attractive area of research when it comes to continuous processing of single-lead ECGs by remote monitoring systems, such as wearable devices like Apple Watches [19], for continuous monitoring of an individual's condition to provide early suggestions for a doctor's visit. This thesis investigates the potential of applying Transformer-based models to ECG classification and provides a comparison with non deep learning (based on extracted features) and other deep learning based models, such as Convolutional Networks. The experiments focus in particular on the classification of Sinus Rhythm, Atrial Fibrillation, Atrial Flutter, Premature Atrial Contractions and Premature Ventricular Contractions.

1.1 Problem statement

1.1.1 Electrocardiograms

Electrocardiograms (ECGs) monitor the condition of a patient's heart. An ECG measures the electricity flowing through the heart in repeated cardiac cycles. ECGs are an essential tool for cardiologists to diagnose heart diseases. To measure the heart's activity, electrodes (called leads within the ECG) are placed on the skin of the upper chest and back. The number of leads affects the quality of the measurements, i.e. single-lead ECGs are based on two electrodes, while multilead ECGs use more than two electrodes. Multi-lead ECGs can contain up to 12 leads, defined in the literature as I, II, III, aVR, aVL, aVF, V1, V2, V3, V4, V5 and V6 [2] [23]. Multi-lead ECGs are used in preference to single-lead ECGs because they provide a view of the heart from different angles and can therefore localise abnormalities more accurate through spatiality. Typically, these recordings last from a few seconds to a few minutes and are called short-term ECGs. However, sometimes long-term ECGs are performed. Long-term ECGs monitor the heart's activity over a longer period of time, e.g. 24 hours, and provide a more complete picture. This is necessary to detect abnormalities that are less common and harder to detect. The scope of this thesis focuses on short-term single-lead ECGs. An electrocardiogram can be recorded with different devices, e.g. in clinical settings 12-lead recordings are standard, while for remote monitoring often portable devices are used, such as the Holter monitoring system. The Holter monitoring system can be adapted to different numbers of leads as required, e.g. three leads or up to 12 leads. Special watches [19] can now also be used to record the heart's activity. These watches use a technology called Photoplethysmography (PPG), where LED lights under the watch emit light into the skin [30]. The light is absorbed and reflected by blood vessels and the changes in light absorption are measured to determine heart rate. However, PPG is a measure of the blood pressure from which the heart rate can be derived, but which is not direct an ECG recording of the electrical activity of the heart. Usually, the derived quality of the sensor is not as accurate as a standard ECG, making it a less reliable method. A watch is still an interesting device because it can monitor a patient's heart condition on a regular basis. It also has the advantage of not requiring the involvement of a cardiologist and can provide possible advice for a more transparent in-house check in a doctor's surgery. Accuracy in arrhythmia detection is important for several reasons, including arrhythmia risk and correct treatment. For example, increased accuracy in automated ECG processing can reduce false positives and false negatives. This can avoid unnecessary doctoral visits and lead to more efficient and effective clinical monitoring and treatment. In addition to developing accurate models, a goal of this thesis will be to transfer and apply the pre-trained models from the Physionet 2021 challenge data [31] to the database provided by the University of Maastricht, here referred to as the MyDiagnostick database. Both datasets contain recordings from different monitoring systems. Analysis and pre-processing steps need to take into account the type of monitoring system and electrodes, number of leads, recording duration, sampling rate and post-processed filters applied, which will be discussed in section 4.1.3.

Figures 1.1 and 1.2 illustrate the activity of a heart from a physiological perspective and a single normal heartbeat from an ECG, defined as a Sinus Rhythm. In the literature [2] [23] a heartbeat is described by the P, Q, R, S, T waves, segments and their intervals. Each wave and segment corresponds to a specific event in the electrical cycle of the heart. Below is a brief description of the entire contraction and depolarisation process of a single normal Sinus Rhythm heartbeat.

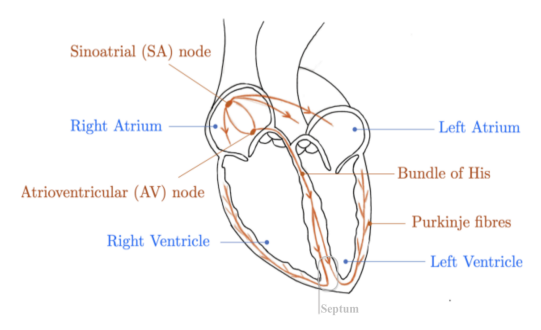


Figure 1.1: Human heart anatomy Source: Collimator



Figure 1.2: Heartbeat rhythm composition Source: Wikipedia

The two graphs show a single heart contraction and depolarisation recorded on an ECG, starting with the electrical impulse from the Sinoatrial (SA) node and the atrial contraction, followed by the depolarisation phase and the pumping of blood from the ventricles into the aorta and pulmonary artery and the final repolarisation and relaxation of the heart. The SA node, located in the right atrium, is the heart's natural pacemaker. It initiates all heartbeats and determines the heart rate. The P wave in an ECG shows the electrical activation of the SA node, which causes the atria of the heart to contract. It is the first wave in a heartbeat and is usually small and positive. When the atria of the heart are filled with blood, the SA node fires and the electrical impulse from the SA node spreads to the two upper atria, causing them to contract. Atrial depolarisation, which fills the blood from the atria into the ventricles, begins about 100 milliseconds after the SA node fires. The Atrioventricular (AV) node, located opposite the SA node in the right atrium, receives the electrical signal from the SA node and marks the start of ventricular depolarisation. The electrical depolarisation is slowed down before propagating to the ventricles, here part of the PR interval. The PR segment represents the time it takes for the signal to travel from the SA node to the AV node. The AV node acts as an electrical gateway to the ventricles and delays the electrical conduction to the ventricles. This delay ensures that the atria contract all the blood into the ventricles before the ventricles depolarise. The AV node conducts the electrical impulse to the AV bundle (also called the Bundle of His). The bundle is divided into right and left branches, through which the electrical impulse is conducted to the Purkinje Fibres and causes the main depolarisation effect of the ventricles. This ventricular depolarisation is seen on the ECG as the QRS complex. The Q wave corresponds to the depolarisation of the interventricular septum. The R wave is produced by the depolarisation of the main mass of the ventricles. The S wave represents the final phase of ventricular depolarisation. Atrial repolarisation also occurs during this time, but the signal is obscured by the large QRS

complex. The ST segment, which follows the QRS complex, is the initial phase of the ventricular repolarisation. The ST segment reflects when the ventricles contract, pumping oxygen rich blood and oxygen poor blood into the aorta and pulmonary artery. The final T wave represents ventricular repolarisation before the heart relaxes. Repolarisation continues until the end of the T wave. The QT interval represents the total ventricular contraction activity (depolarisation and repolarisation). At the end of this process, the atria are filled with blood again and the SA node fires to repeat the cardiac cycle. This process represents a normal Sinus Rhythm.

1.1.2 Arrhythmia diseases

In the literature [2] [23], normal Sinus Rhythm is described as being within normal limits and ranges. For Sinus Rhythm, this is usually between 60 and 100 beats per minute. A rate below 60 is generally defined as Sinus Bradycardia and a rate above 100 beats per minute as Sinus Tachycardia. The two classes group several subtypes of arrhythmia. Arrhythmia subtypes are characterised by different origins and causes in the heart and need to be treated individually. In the Physionet 2021 challenge data, more than 100 arrhythmia subtypes are present 4.1. For the evaluation of the models transferred from the pre-training on the Physionet 2021 challenge data to the MyDiagnostick data, this thesis focuses the classification on Sinus Rhythm (SR), Atrial Fibrillation (AF), Atrial Flutter (AFL), Premature Atrial Contractions (PAC) and Premature Ventricular Contractions (PVC). This is due to the limited class annotation of the MyDiagnostick dataset, which only consists of these class labels. However, a broader performance evaluation of the models is performed on 26 different classes of the annotated Physionet 2021 challenge data. Most of the more than 100 available classes in the Physionet 2021 challenge data contain few examples, so the challenge uses a subset of 26 classes for the official scored metrics. Much research has been done to develop accurate models for binary classification of non-abnormality vs. abnormality, i.e. Sinus Rhythm vs. Atrial Fibrillation. However, fewer research has been done on distinguishing specific arrhythmia subtypes, such as Atrial Fibrillation and Atrial Flutter. The limited research in this area tends to show weaker outcomes [33] [32]. Atrial Fibrillation and Atrial Flutter are often confused by algorithms because they have similar characteristics. PAC and PVC are also examined, because the MyDiagnostick dataset provided contains these annotations. In the following is a brief summary of each rhythm and arrhythmia type studied:

- Sinus Rhythm (SR): Regular P waves followed by a narrow QRS complex, typically lasting between 80 and 100 milliseconds. The PR interval remains constant throughout. Heartbeats are regular, between 60 and 100 beats per minute.
- Atrial Fibrillation (AF): Atrial fibrillation is abnormal electrical activity that causes the atrial muscle fibres to contract at different times. Atrial Fibrillation is characterised by a rapid and irregular heartbeat. These uncoordinated contractions produce a quivering or fibrillating activity. Atrial Fibrillation does not have constant P waves preceding the QRS complexes, although the fibrillation effect may resemble a P wave at times when it is not expected. Only some of the electrical signals are conducted down into the ventricles, resulting in ventricular depolarisation. However, there is no real pattern to which impulses are conducted. In the literature [2] [23], AF is also described as an irregular heart rhythm, which explains the variable lengths of the RR intervals.
- Atrial Flutter (AFL): Atrial Flutter is often confused with Atrial Fibrillation because of its similar characteristics. The main difference is that Atrial Flutter is characterised by coordinated electrical activity in the atria due to a re-entry pathway, resulting in rapid contraction of the atria. This is usually around 250 and 300 beats per minute with a regular

atrial rate and a narrow QRS complex. The AV conducts the signal slower, resulting into a slowed ventricular depolarization. Atrial Flutter is therefore characterised by the number of P waves compared to the number of ventricular contractions, which shows the ratio of non conducted to conducted beats, e.g. a 3:1 conduction means that every third atrial impulse is conducted to the ventricles. Atrial Flutter is often characterised by a "sawtooth" pattern of atrial activity, known as flutter waves, caused by the rapid atrial depolarisation. A conduction ratio of one to one is also possible and is associated with instability and progression to ventricular fibrillation. These ratios can be variable in the same patient, making the ventricular rate irregular, which is why it can often be mistaken for Atrial Fibrillation. Atrial Flutter is less dangerous than Atrial Fibrillation but can progress if left untreated.

- Premature Atrial Contractions (PAC): Premature Atrial Contractions are heartbeats that originate in the atria. PACs occur as early and extra beats that disrupt the regular heart rhythm. On an ECG, PAC shows a premature and often abnormal P wave followed by a QRS complex. Premature Atrial Contractions are characterised by a different P wave morphology compared to the normal sinus P wave, which follows a narrow QRS complex. The beat following the PAC may resemble a pause. However, if the locations of the P waves are followed, they should occur at expected times approximately. Normally, PACs are a fairly common finding on ECGs and usually do not require further investigation.
- Premature Ventricular Contractions (PVC): Premature Ventricular Contractions are early heartbeats that originate in the Purkinje Fibre region of the ventricles. They are extra, abnormal beats. On an ECG, PVCs are seen as wide and bizarre QRS complexes that are not preceded by a P wave. The QRS complex in PVCs is longer than 120 milliseconds and there is a compensatory pause before the next beat. PVCs are rarely dangerous on their own, unless they are frequent, i.e. if they occur more than 10 to 30 times per hour, or if they occur every beat in a row, they can be diagnosed as ventricular tachycardia, which is critical for stroke.

Figure 1.3 shows for normal Sinus Rhythm and each of the investigated arrhythmias an example from the Physionet 2021 challenge database.

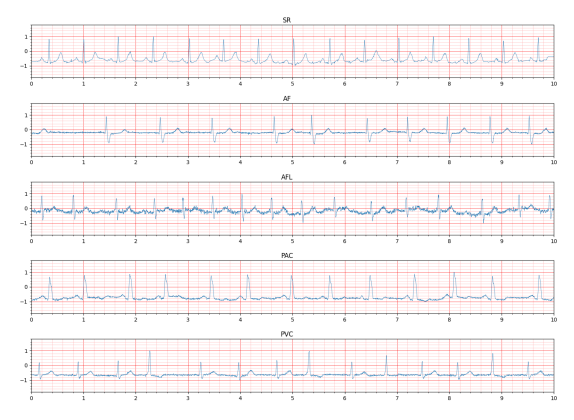


Figure 1.3: Examples for Sinus Rhythm (1), Atrial Fibrillation (2), Atrial Flutter (3), Premature Atrial Contractions (4) and Premature Ventricular Contractions (5) from the Physionet 2021 database

1.2 Goal

The main goal of this thesis is to investigate and compare different transformer architectures with non deep learning (based on extracted features) and other deep learning approaches, e.g. Convolutional Networks, for the specific classification of Sinus Rhythm, Atrial Fibrillation, Atrial Flutter, Premature Atrial Contractions and Premature Ventricular Contractions. The main advantage of the Transformer models is the attention mechanism which is designed to learn relationships within data. This could be useful for tasks such as classifying Atrial Fibrillation in ECGs, where relationships within the data could lead to a better understanding of underlying patterns that depend on specific events. A Convolutional Network focuses on local patterns and is less suited to understanding relationships within extracted features. This work explores this mechanism to extract the harder to detect abnormalities, such as distinguishing between Atrial Fibrillation and Atrial Flutter, or capturing Premature Atrial Contractions and Premature Ventricular Contractions, which relate to different events within the repolarisation and depolarisation cycle within the ECG. Premature Atrial and Ventricular Contractions often occur only once on an ECG. Compared to convolutional filters, Transformer models use the weight-based attention mechanism that can reinforce the model to attend to specific parts of the input. The research question is whether the Transformer model with its attention mechanism can capture these patterns better than a traditional Convolutional Network. In addition, different Transformer

architectures are investigated and analysed, i.e. input preparation, applied positional encoding, number of blocks and heads, attention dimension and further hyperparameter tuning. For the evaluation of the models the Physionet 2021 challenge data [31] and the provided MyDiagnostick databases will be used. The work will first evaluate the models on the Physionet 2021 challenge data itself and then attempt to transfer the pre-trained models from the Physionet 2021 challenge data to the MyDiagnostick database. Necessary pre-processing steps and considerations are discussed. Although the Physionet 2021 challenge data provides 12-lead ECGs, the thesis will limit the experiments to single-lead ECGs to narrow the problem statement and because the provided MyDiagnostik dataset contains only single-lead ECGs. Four research questions are formulated, which will be addressed in the experiments and answered by the end of this thesis:

1.2.1 Research questions

- 1. How well does a Transformer-based model perform on the Physionet 2021 challenge data compared to a feature-based model or a Convolutional Network?
- 2. Can an ensemble Transformer model and Convolutional Network effectively capture spatiotemporal information and improve accuracy?
- 3. How is the performance of the most promising model at discriminating SR, AF, AFL, PAC and PVC on both datasets?
- 4. What are the challenges in transferring the pre-trained models from the Physionet 2021 challenge data to the MyDiagnostick database? Do the models generalise well, even though different ECG devices were used?

1.3 Chapter overview

Chapter 1 provided an introduction to the problem statement and research objective of this thesis. Chapter 2 discusses related work in this area. Chapter 3 explains the relevant mathematical background and presents the own approaches. Chapter 4 discusses the experiments and evaluates the models. Chapter 5 summarises the results and gives an outlook for further research.

Chapter 2

Related work

This chapter discusses related work. It is divided into three sections. The first section discusses related work on Transformer-based models for the general classification of various abnormalities in ECGs. The second section discusses related work on models developed for the Physionet 2021 challenge, which provides an annotated dataset for the classification of 26 different arrhythmia types. The final section discusses related work in the specific area of classifying Sinus Rhythm, Atrial Fibrillation, Atrial Flutter, Premature Atrial Contraction and detection of Premature Ventricular Contraction based on deep-learning models.

2.1 Related work on the developed models for the CinC Physionet 2021 challenge

Several paper in the Physionet 2021 challenge incorporate an attention-based mechanism. To give an overview, the four highest ranked papers with the best performing models are discussed, from which the first, second and fourth ranked model utilize an attention-based mechanism. The fourth ranked paper highlights several different attention-based methods and is discussed in more detail. The first ranked model of the Physionet 2021 challenge [27] by Nejedly et al. proposes a deep residual Convolutional Network combined with a single Multi-head Attention layer. As all approaches in the Physionet 2021 challenge, their model is designed for 12-lead ECG classification, while for fewer lead configurations the unused leads are padded with zeros. Their approach achieves about 58% on all 2, 3, 4, 6 and 12-lead tasks within the challenge metric and is able to achieve on all lead tasks the 1st ranked performances. Key features of their model are that the model is based on the ResNet architecture [16] proposedvby Zhang et al., which introduced the deep Convolutional Network based on residual connections. Nejedly et al. use large convolutional filters, i.e. 15x15 on the first layer and 9x9 in subsequent residual blocks. The model is composed of five residual blocks, each consisting of two convolutional layers, a batch normalisation layer and a feed-forward layer with leaky relu as activation function. In the final bottom layer the model integrates a single Multi-head Attention block, as the authors justify to also capture temporal relations [27]. However, notable is a follow-up study [28] on the same experiments, where the authors find that the Multi-head Attention layer slightly worsened the classification performance of the same model. Based on the experiments the authors find that without the Multi-head Attention block the model achieves an overall performance of 58% and with the Multi-head attention block only 57% on the official Physionet 2021 challenge data. A question that arises from this, is whether the features that are passed into the attention

block are not appropriate prepared. In their paper the authors do not explain this in detail. Furthermore, the authors do not state, whether positional encoding have been applied, that potentially could help the attention mechanism with extracting relational patterns. Another key feature of their approach is that the authors propose a loss function that incorporates the Physionet 2021 challenge evaluation metrics to optimize the model on the challenge metric. The loss function is an altered version from Vicar et al. [39] [40]. This work will adopt the cost function for fine-tuning the models on the Physionet 2021 data, which is discussed in section fine-tuning 3.3.7. The second ranked paper by Han et al. [15] achieves between 55%-58% in the Physionet 2021 challenge. In their approach the authors use a deep Convolutional Network combined with demographic features, i.e. age and gender. The features are directly incorparated into the final output dense layer of the network. Within the model a Squeeze and Excitation Network is integrated. A Squeeze and Excitation Network is based on a mechanism called channel-attention proposed by Hu et al. in [21]. This work benchmarks a Squeeze and Excitation Network as it follows a related attention mechanism approach compared to the idea of Transformer models. The main ideas of the Squeeze and Excitation Network will be discussed in 3.3.5. The authors of [15] highlight that the model achieves high generalisation performance by using a "leaveone-dataset-out-cross-validation" strategy. In addition, their approach use a data augmentation method called "Mixup" [46], which smoothes the decision boundaries of the model through a regularisation technique that mixes two input samples with their features and labels based on a coefficient. The third ranked paper by Wickramasinghe and Athif [43] proposes two deep Convolutional Networks with four residual blocks, which achieves an accuracy of 51%-55% on the final test set. The two networks work in parallel, while one Convolutional Network receives the raw ECG signal with temporal information as input and the second Convolutional Network applies a Fast Fourier Transformation (FFT) on the ECGs to capture features from the frequency domain. Both models are combined by a dense and pooling layer. Previously the authors apply standard pre-processing steps such as normalisation, resampling and zero padding. The fourth ranked paper by Srivastava et al. [37] applies a residual Convolutional Network, which incorporates several convolution- and attention-based methods. The methods include "inception" proposed in the GoogLeNet by Szegedy et al. in [38], channel-attention based on the squeeze and excitation network architecture [21] and attention-pooling proposed by Ilse et al. in [24]. This work will also experiment with the Inception Network in more detail. The main ideas of the Inception Network will be discussed in 3.3.4. Although, the original paper apply 1x1 convolutions in the inception module, the authors from the challenge paper [37] do not apply these 1x1 convolutions, instead they use 1D convolutions with dimensions 1x3, 1x4 and 1x5. In addition, the authors alter the method of channel-attention to channel self-attention. Instead of using global average pooling the authors extract 32 features from each channel and calculate attention instead between within each channel based on the pooled feature vector. Interrelation among channels is captured through sharing weights across channels. Furthermore, the authors extend the model output with a technique called attention-pooling [24] and alter it as described to Multi-head Attention, which pools the final feature vectors into a feature space of size NxL, where N is the number of predicted classes and L are feature aggregations. The inputted feature vectors are subdivided into N segments. The attention block uses two trainable weights matrices U and W that linearly project the feature vectors and apply softmax to scale the weights. The scaling values are multiplied with the inputted feature vector. However, the paper [37] does not state in detail the shape and size of the inputted feature maps from previous channel-attention layer. Based on the above approaches about residual Convolutional Networks, Multi-head Attention, Squeeze and Excitation Network and Inception Network it is hard to reason which of the above presented approaches are the most improving building blocks that lead to a good performance for capturing arrhythmias. Therefore, this work analysis each of these approaches, especially

the Transformer encoder model, for multi-label classification of 26 cardiac arrhythmias in more depth by investigating each approach independently and discusses modifications, which will be described in the own approach section 3.3. Objective will also be to reduce the model size, while aiming for high performance.

2.2 Related work for the classification of ECGs based on Transformer models

In recent years, Transformer models have gained considerable popularity because their model and training design can be much more easily accelerated on a graphic processing unit (GPU), due to the capability of parallel weight updates during backpropagation through the shared attention (query, key and value) matrices. This leads to a significant reduction in training time and hardware costs, while maintaining comparable or even better performance, compared to the previous Long-Short-Term-Memory (LSTM) [17] or Recurrent Neural Network (RNN) [20], which were widely used until Transformer models for sequential data. LSTMs and RNNs need to process data sequentially through their units and do not scale well for large training applications leading to computational slowness and costs. In addition, Transformer models have the ability to process long-range sequential data much better, because LSTMs and RNNs can suffer from long-range dependencies caused by vanishing gradient problems during backpropagation. Choi et al. [6] propose a more complex encoder-based Transformer model and training methodology for ECG arrhythmia classification, called ECGBERT, that closely models the architecture and training procedure of BERT [8]. BERT, based on the Transformer encoder architecture, was one of the first successful and popular language models fine-tuned on various tasks. Following the architecture design of BERT, ECGBERT is fine-tuned on multiple down-stream tasks, e.g. atrial fibrillation classification (binary), heart-beat classification (normal, unknown, supraventricular ectopic, ventricular ectopic and fusion heartbeats), user verification by using the ECGs as biometric authentication to predict patient IDs and sleep apnea detection. Similarly to BERT, the authors create an own vocabular, here a wave segment vocabular, to tokenize each ECG and apply then BERT's training procedure Masked Language Modeling (MLM) to pre-train the model. In a preparation step, the ECG signals are pre-processed (normalised, filtered etc.) and splitted by fiducial points into wave segments. Time-frequency analysis and Hamilton's algorithm [29] are applied to clean the ECG signals and extract onset and offset points. The obtained wave segments are used to cluster the waves using K-mean and Dynamic Time Warping. For this, the authors train four classifiers: P, QRS, T and background waves. 70 clusters are obtained with 12 P, 19 QRS, 14 T and 25 background (e.g. PR or ST intervals) wave clusters. Next, each segment is classified by the corresponding classifier and assigned to a wave cluster. The classified wave segments are then mapped to pre-defined tokens. The authors do not describe how the tokens are encoded, an assumption could be that these are random initialized embeddings. In addition, CNN features are extracted from the raw ECG using an U-Net architecture [34]. The authors reason that the model can only learn high-level wave context relations from the ECG tokens, but CNN features provide refined pattern information and therefore are crucial for extracting fine-granular features. The tokens are combined with the CNN embeddings and positional information is added, likewise in the original Transformer paper [1]. Based on this, the authors create training samples from the ECGs that form sentences of wave segments, where each sentence contains 1-8 consecutive heartbeats and a heartbeat is composed of several wave segments. The constructed ECG wave sentences are then inputted to the Transformer-encoder module, where several sequences can be inputted, which are splitted by a separation "[SEP]" token. The model is pre-trained using 15% masking (similar to Masked Language Modeling in LLMs). For pre-training of the Transformer

the authors use the MIT-BIH database (AFIB/arrhythmia) [13] and the Apnea-ECG database from Physionet. Based on the pre-trained model weights the authors use the model to fine-tune it on several down-stream tasks by adapting the output layer to the corresponding tasks. The model achieves for binary classification of Atrial Fibrillation an accuracy score of 97% and for beat classification about 86%. Another work from Zhao [48] lists in a review from 2023 several Transformer-based models for ECG classification. Most of the Transformer-based ECG models discussed in the review combine the Transformer encoder module with a Convolutional Network [3]. [5] [9] [22] [26] [48]. The author reason that the Convolutional Network have a limited receptive field, which prevents them from learning distant dependencies and extract local patterns. On the other hand, Transformers learn long-range dependencies through their attention mechanism. The combination of both capture spatio-temporal information from the ECG signal [48]. Hu et al. [22] propose a encoder- and decoder-based Transformer architecture. The model first extracts features from a single-lead ECG signal using multiple convolutional layers. Instead of applying a global pooling layer, the authors treat each feature map as an input token to a Transformer encoder and decoder block, similar to the original Transformer architecture [1]. In addition, the authors add positional encoding. The decoder block unmasks each token by token (here the positional encoded feature maps), through which the decoder block sequentially classifies the next ECG segment. The paper uses the MIT-BIH arrhythmia database [13], which contains annotations for each heartbeat (Normal, Ventricular, Supraventricular, etc.). Bing et al. [3] propose an encoder-based Transformer architecture that combines a Vision Transformer with a Convolutional Neural Network, called ConVit. Vision Transformer (ViT) [10] was one of the first papers that effectively applied Transformer models to computer vision tasks. In Vision Transformers an embedding is obtained by a subpart/pixel group from an image and the model extract global relations within the image for classification task by considering relations within image subspaces.

2.3 Related work for the specific classification of SR, AF, AFL, PAC and PVC based on deep learning models

Wang et al. [42] analyse in their paper an approach, called PVCNet, for the particular detection of PVC. The authors apply a deep Convolutional Network in combination with several feed-forward layers. The input of the model is the full sequence of a single-lead ECG. The authors use the publicly available MIT-BIH database [13] for pre-training and test it on the St. Petersburg INCART database. Their model is able to achieve an accuracy of 98% on the test set. In their work, the authors highlight the importance of several pre-processing steps, including data augmentation to overcome class imbalance, splitting the training and validation sets by patients, ECG resampling and filtering, i.e. a butterworth band-pass filter, as the authors justify that this filter can retain the main frequency components of the signal. Li et al. highlight in their paper [18] the importance of improving the accuracy of assistive ECG devices. In their work they develop a deep learning approach that specifically classifies Atrial Fibrillation, Premature Atrial Contractions and Premature Ventricular Contractions from the first two leads of an ECG. In the study the authors find that their assistive ECG device makes about 18%-24% false positives after revisiting the labels. In their approach the authors use a combination of a Convolutional Network and a Long-Short-Term-Memory (LSTM). In the first stage, the CNN model extracts features with the P, QRS and T intervals. The extracted features are then fed into an LSTM model, which classifies the successively extracted features. The authors justify that the interaction between the two models ensures that spatial-temporal features are extracted. With this approach the authors are able to increase the accuracy compared to the device from 0.77 to 0.86 (AF), 0.76 to 0.84 (PVC) and 0.82 to 0.87 (PAC), after manually revisiting labels. Another study by Marco et al. [7] focuses on the classification of non PVC and PVC using only the QRS complex from an ECG. For this the authors utilize the MIT-BIH database [13] by extracting all QRS complexes from long-term ECGs in the MIT-BIH database. In their study they compare several models, e.g. random forest, LSTM, bidirectional LSTM, ResNet-18, MobileNetv2 and ShuffleNet. MobileNetv2 [36] and ShuffleNet [47] CNN networks designed for efficiency and to work on mobile devices. Ribero et al. highlight in their work [32] and [33] that the classification of Atrial Fibrillation and Atrial Flutter is not a trivial task. In their paper, the authors propose a 1D and 2D Convolutional Network trained on 1D ECG signals and 2D ECG images. The underlying datasets are a combination of the Physionet 2021 challenge database and private datasets collected from various cardiologists and hospitals. However, in their study [33] the authors state that the performance of the model trained on only the Physionet 2021 challenge data is significantly lower than that trained on their own dataset, as the authors justify due to quality of the Physionet 2021 data. Wang [41] investigates the classification of Atrial Fibrillation and Atrial Flutter and proposes a combination of a Convolutional Network with an improved version of the Elman Neural Network (IENN) [12], a specific form of the Recurrent Neural Network (RNN) architecture, which uses a context node for improved contextual memory. The combined model is able to achieve around 99% accuracy on the MIT-BIH [13] database for the binary classification problem.

Chapter 3

Methods

This chapter discusses the utilized and modified approaches developed for this thesis. It is divided into three sections. The first section discusses the mathematical background of Transformer models. It highlights the architectural differences between a Transformer model trained on language and applied to ECG signals. The second section gives a brief overview from the main components of a Convolutional Network. The third section presents the modified approaches that are used to address the problem of multi-label ECG classification into 26 different arrhythmia classes, here a feature-based random forest classifier, a residual Convolutional Network with and without dilation, a Transformer encoder model, an Inception Network, a Squeeze and Excitation Network and the own approach, a modified ensemble model of an Inception Network combined with the Multi-head Attention mechanism. Finally, section 3.3.7 describes how the models are fine-tuned on the Physionet 2021 challenge data.

3.1 Mathematical background about Transformer models

The Transformer model was introduced in the paper "Attention Is All You Need" by Vaswani et al. [1]. Figure 3.1 shows the model architecture.

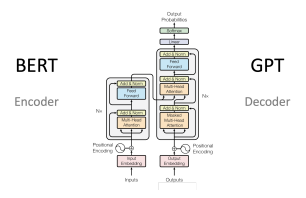


Figure 3.1: Transformer architecture Source: "Attention Is All You Need" [1]

The model consists of two parts, the encoder block, left half in 3.1, and the decoder block, right half in 3.1, connected by the arrow in the middle. Initially, this architecture was designed for

language translation. The encoder block receives the input sequence and encodes the words (tokens). The encoder block maps then the input sequence to the translated output sequence via the decoder block, which predicts the next most likely output token. The main strength of the Transformer model is the "attention mechanism" implemented in the "Multi-Head Attention" block 3.1. The attention mechanism might be compared to a convolutional filter, although it has a different objective and uses a different approach. Similarly, the attention block applies a "filter" that projects and merges each feature with surrounding features from the input sequence. However, the objective of the Transformer model is to learn relations within the input based on weighted attention scores, unlike the Convolutional Network, which aims to extract invariant features from local inputs and feature groups. That means, depending on the kernel size, a convolutional filter applies only to local input and feature groups, while the attention block applies to the entire input sequence as whole. The attention block computes query, key and value pairs by considering each input element with each other input element individually. During training, the attention block adapts its attention weights to different arrangements in the training data, i.e. as it sees different variations in the training data it can extract relationships and learn meaningful patters from various data arrangements, by forwarding the most important relations and features. The attention mechanism is in more detail discussed in section 3.1.2. Many researchers have adopted the original Transformer architecture and use parts of it for various downstream tasks, such as (text) classification tasks, e.g. BERT (Bidirectional Encoder Representation from Transformers) [8], or for (text completion) regression tasks, such as GPT (Generative Pre-trained Transformer) [4]. BERT uses only the encoder block, while GPT uses only the decoder block. This thesis utilises and implements only the encoder block of the Transformer model. The reason is that the encoder block considers all inputted embeddings (bidirectional), whereas the decoder is designed for prediction tasks and focuses its attention on the output of the encoder block, also referred as cross-attention, and on its own past states (unidirectional) by masking future states. This is achieved by adding masking to the embeddings in the decoder block. This makes the decoder less suitable for classification tasks and could prevent a classification model from learning a more meaningful representation from the entire ECG because valuable information would be masked. Therefore, this works investigates an encoder-based Transformer model. In general, a Transformer model divides the input sequence into subspaces, called embeddings. Embeddings are vectors that represent specific input features. In language-based Transformers, each embedding represents an assigned sub-word or character in the known vocabulary of the model, also called tokens. The input preparation process for language-based Transformers work different as in this work and is usually handled by a tokenizer, a separate sequence mapping and vocabulary indexing tool of the model. Tokenizers are one of the key differences between Transformers trained on language tasks and Transformers trained on ECG signals. In this work, the Transformer model does not utilize a tokenizer to map the ECG signal. One reason is that the signals are already present as processable numbers and do not need to be encoded. The other reason is that words in language are repetitive, whereas heartbeats or segments of ECG signals have different shapes and are unique, making it superfluous to index these or train an input embedding matrix that can only represent a finite feature space of arrhythmia features. In this work, both raw ECGs and features obtained from convolutional filters are utilized for inputting features to the Transformer encoder block. This includes either splitting the signal into fixed-size segments or extracting features maps / channels by a Convolutional Network. This will be analysed and discussed in the evaluation 4.2 and discussion 4.2.2. The input size of a Transformer model needs to be fixed-size, also known as the context window of the model. In this work, this is either a 10 or 60 seconds long single-lead ECG signal, depending on the investigated experiments (either Physionet 2021 or MyDiagnostick data). The ECG embedding features are fed directly into the encoder block, whereas in language-based Transformer models an additional

trainable embedding matrix represents the first layer of the model before the encoder block.

3.1.1 Positional encoding

The original Transformer model adds positional information to each embedding before these are passed into the attention block. The positional encoding layer is connected with the output of the attention block via a residual connection. This ensures that the temporal information about the order of the embedding sequence is preserved and propagated throughout the network. The original paper proposes a positional encoding technique using sine and cosine functions of different frequencies:

$$PE(pos, 2i) = \sin\left(\frac{pos}{10000^{\frac{2i}{d_{\text{model}}}}}\right)$$
(3.1)

$$PE(pos, 2i + 1) = \cos\left(\frac{pos}{10000^{\frac{2i}{d_{\text{model}}}}}\right)$$
(3.2)

"Pos" represents the index position of each embedding in the sequence and i is the embedding dimension, which is equal sized for all input embeddings. Each dimension in the embedding corresponds to a single sinusoid and the wavelengths form a geometric progression from 2π to $10000 \cdot 2\pi$ that projects the embedding values into a positional order. The authors chose this function, because they hypothesised that it would allow the model more easily to learn and attend to relative positions, since for any position k, PE_{pos+k} can be represented as a linear function of PE_{pos} . [1]. This work investigates the effect of adding positional information to ECG segments for ECG classification into arrhythmias and discusses this in the evaluation 4 and discussion 4.2.2 sections.

3.1.2 Attention mechanism

The input to the attention layer (Transforemr encoder block) is a sequence of length N, consisting of N positional encoded embeddings (features) with dimension d, e.g. a tensor of the shape $N_{sequence\ length} \cdot d_{embedding\ dimension}$. The attention layer uses three trainable query, key and value matrices for projecting the input sequence into three new feature spaces, denoted by the matrices WQ, WK and WV. The multiplication of the input with each weight matrices WQ, W^{K} and W^{V} , yields the matrices Q, K and V that go into the attention layer, which are represented by the three arrows in 3.1. An important advantage is that the weights WQ, WK and WV can be computed in parallel during the forward and backward pass and do not depend on sequential states as in LSTMs or RNNs. The next steps form the core of the Transformer model, which is the "scaled dot-product attention", shown in figure 3.2 and formula 3.3. The first step in this process is to compute the dot-product between Q and K by multiplying Q with K^{T} , which gives an unscaled attention weight matrix A. Since every projected embedding in Q is multiplied with every transposed projected embedding in K^{T} , the shape of the attention weight matrix is $N_{sequence\ length} \cdot N_{sequence\ length}$ (the quadratic number of the inputted sequence). A disadvantage is that the trainable projection weights WQ, WK and WV scale quadratic with the length of the input sequence. The obtained attention matrix A represents how much each embedding relate to each embedding in the same sequence (self-attention), which is learned through WQ, WK and WV and which is utilized to scale the embeddings in the projected value matrix V by taking into account relations among the embeddings. The attention scores in Aare divided by a normalising constant $\sqrt{d_k}$ (the row dimension of WK, which is of shape d_k · $N_{sequence\ length}$, more precisely h_{heads} · d_k · $N_{sequence\ length}$ - but which is discussed in more detail in the next section about Multi-head Attention 3.1.3), which helps the model to reduce

the variance of the computed relational values from Q and K and stables training. Next, the attention weight matrix A is scaled by a softmax function to obtain softmax-normalised and interpretable weights. The softmax-normalised weights are the final attention weights of A that will be used to scale the value matrix v by matrix multiplication from the obtained attention matrix A. This operation forces the model to attend and propagate specific embeddings more than others in the network. A linear matrix 3.3 is followed after the calculation with V. The linear matrix, which should not be interchanged with the additional feed-forward layer as part of the Transformer encoder block, is a trainable linear weight matrix that is used to weightbased concatenate the heads, which will be discussed in the next section 3.1.3. Usually, the outputted shape of the sequence and embeddings from the Transformer encoder block should have the same shape as the inputted sequence and embeddings to the Transformer encoder block, e.g. $N_{sequence\ length} \cdot d_{embedding\ dimension}$. Depending on the dimension configuration of WQ, WK and WV, the initial shape might be restored by this linear matrix operation, which is also discussed in more detail in the next section 3.1.3. The masking operation (optional) 3.2 is skipped here, because it is only used in the decoder block and rather suitable for temporal forecasting tasks. In short, the masking operation adds a triangular matrix masking to the attention matrix A so that the processed attention scores depend only on unmasked and past embeddings. The output of the encoder block is added with the input embeddings by a residual connection, which is an important step. This ensures that the initial feature and it's temporal information is propagated throughout the entire network, when stacking multiple Transformer encoder blocks. In addition, layer normalisation is applied to stabilise the weights during training. Next, a feed-forward layer follows the attention block 3.1. The feed-forward layer allows the model to learn non-linearity, since the operations inside the attention block remain linear matrix multiplications. In the original paper a single hidden-layer with a ReLU activation function is used [1]. For comparison, GPT-3 uses a feed-forward layer with a hidden-layer size of 24. The output of the feed-forward layer is then reconnected by a second residual connection and again normalised. Since the output of the entire encoder block should have the same shape as the input $N_{sequence\ length} \cdot d_{embedding\ dimension}$, it allows to easily stack multiple encoder blocks without further modifications, thus each block can learn and attend to different relations among the inputted embedding.

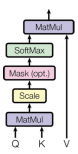


Figure 3.2: Scaled Dot-Product Attention Source: "Attention Is All You Need" [1]

Attention
$$(Q, K, V) = \operatorname{softmax}\left(\frac{QK^T}{\sqrt{d_k}}\right)V$$
 (3.3)

3.1.3 Multi-Head Attention

In addition, the original Transformer model introduces a mechanism called "Multi-Head Attention" [1]. The idea of Multi-Head Attention is to train multiple attention blocks on different inputted embedding sub-spaces simultaneously in the same attention layer. This allows to capture a wider range of relationships and to encapsulate partial information from the embeddings into partial contexts. Multi-head Attention splits each embedding into h sub-embeddings before these are passed in parallel to h query WQ, key WK and value matrices WV. In the original paper 8 heads were used [1]. Following the example from the original paper, an attention layer, which receives $N_{sequence\ length} \cdot 512_{embedding\ dimension}$ as input, splits each embedding into 8 sub-embeddings, yielding $N_{sequence\ length} \cdot 8_{heads} \cdot 64_{sub-embedding\ dimension}$, as input for the attention layer. This can be thought by slicing horizontally the input sequence with vertical feature embedding vectors in 8 pieces to obtain 8 parallel partial embeddings sequences as input for the Transformer encoder, where each sequence is separately passed to another (smaller) attention block that work in parallel, also referred to as "heads". The dimension of the sliced subembeddings is here described as $qkv_{dimension}$, where $h_{heads} \cdot qkv_{dimension}$ should normally yield the initial inputted embedding dimension $d_{embedding dimension}$. Moreover, $d_{embedding dimension}$ should be divisible by h_{heads} to obtain equal-sized attention blocks. The computation then proceeds as described in the previous section about the scaled dot-product attention, but here in parallel on different embedding sub-spaces and for multiple WQ, WK and WV matrices. After the computation within the attention block the outputted sub-embeddings are then concatenated again and multiplied by a trainable linear layer, shown in figure 3.3 as "Linear" at the top. The multiple "Linear" blocks at the bottom simply represent the reshaping procedure of Q, K and V into h heads. However, the "Linear" operation at the top represent trainable weights that serve as a linear projection operation, without the use of an activation function, which weighted-based concatenate the sub-embeddings from the heads. In addition, it can ensure that the initial input shape of $N_{sequence\ length} \cdot d_{embedding\ dimension}$ is restored, even if the shape after "Concat" does not match the initial shape. Generally, the input and the output dimension of the standard attention block has an equivalent shape of $N_{sequence\ length} \cdot d_{embedding\ dimension}$. This means that $h_{heads} \cdot qkv_{dimension}$ does not necessarily be equivalent to the embedding shape $d_{embedding\ dimension}$, thus the attention block can be of variational size and learn attention weights on a densed embedding feature space. In addition, it serves as a trainable linear projection operation, that weighted-based concatenate the sub-embeddings received from the heads and the concatenation operation. Figure 3.3 illustrates the entire Multi-Head Attention process.

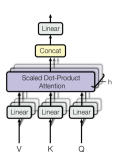


Figure 3.3: Multi-head Attention Source: "Attention Is All You Need" [1]

3.2 Mathematical background about Convolutional Networks

Convolutional Networks are often applied deep-learning models for the classification of ECG signals [18] [7] [42], including the first three ranked models and many other models within the Physionet 2021 challenge [27] [15] [43]. Traditionally used for computer vision tasks, a convolutional network can similarly be applied to one dimensional data. The main component in a convolutional network is the convolutional filter (also called kernel), which is a fixed-sized window that stripes over the input data, where each data point and it's surrounding data points within the window are multiplied with the kernel weights and added to project the data points into a new single outputted data point. The kernel then stripes over the entire input data. The outputted projection is called feature map or channel. At the beginning the kernel weights are randomly initialized, then the kernel later optimizes the kernel weights to the training data by learning meaningful patterns, such as corners or edges to more abstract concepts like faces in computer vision tasks. Commonly, several convolutional filters work in parallel within the same convolutional layer, yielding several outputted features maps or channels that are stacked in depth. Therefore, a convolutional layer is typically configured by the number of convolutional filters, kernel, stride and padding size. The number of filters in a layer determines how many distinctive patterns the convolutional layer can learn within a specific network level from the data, while each feature map contain different features from this layer. Usually, this number starts small, e.g. 32 and increases from one layer to the next deeper layer, e.g. 64, 128 and so on. The reason for this is that in deeper layers more abstract concepts need to be learned by combining simpler corner and edge features from previous layers. This process requires a larger number of filters to capture the increasing complexity and diversity of features. The filter size describes the size of the stripe window and mainly influences the number of trainable parameters. Smaller filters can focus on details, while larger filters can receipt distant associated patterns. Smaller windows may also prevent overfitting and enable deeper and more complex network hierarchies. Stride determines the number of data points the kernel window is shifted when sliding over the input data. Therefore, some data points can be left out by skipping. Padding adds margin to the borders of the input and can reduce the input size. Stride that is larger than the kernel size and padding can miss relevant input information by ignoring or skipping features, but reduce the feature space and number of trainable parameters. Beside standard convolutional filters, stride and padding, dilated convolutions can be applied. Dilated convolutions employ a kernel that incorporates strides. These kernels extent the receptive field and extract features with less resolution. Figure 3.4 illustrates a dilated convolution applied on one dimensional ECG data.

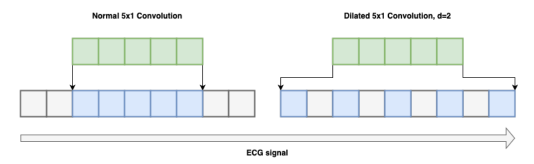


Figure 3.4: Dilated convolution

This work investigates the effect of dilated convolutions for the classification of ECGs. Typically, a convolutional layer is followed by a pooling layer that aims to reduce dimensionality. Other than the convolutional layer, the pooling layer uses a single kernel across all channels that (most commonly used) applies "min", "max" or "average" pooling. This means, the pooling kernel forwards either the minimum, the maximum value or averages the values within the window. For example, in a 1D signal a pooling window size of two will reduce the input dimensionality by half, thus for two data points a single data point is outputted and forwarded to the next layer. This process enables the convolutional network to extent the receptive field and learn from detailed features to more widely scattered features. As an alternative to pooling layers strides can be used for downsampling the input. Other common components in convolutional networks include feed-forward layers using non-linear activations. These layers are utilized to learn non-linear features, since convolutional kernels only apply linear multiplication operations. The final layers of a convolutional network Commonly apply global pooling, that extract from all values within a channel a single value, e.g. min, max or average, from which then a feed-forward or softmax layer follows as the final model output, depending on the task.

3.3 Approaches

3.3.1 Feature-based Random Forest

As baseline comparison, a feature-based random forest is trained based on the Biobss Python package. The Biobss library provides a set of toolkits for extracting features from ECG signals. Another library to work with ECG data is the Neurokit2 package, which is a more maintained library, but which is due to similar methods not further discussed here. Table 3.3.1 shows 39 extracted features using the Biobss library and a short description for each feature. In the first step, the Biobss library calculates all R peaks and other fiducial points in the ECG, e.g. P, Q, S and T wave onset and offset points. For the point localisation the package has one of these three methods implemented: "pantompkins" [29], "hamilton" [14] or "elgendi" [11]. For the experiments the "pantompkins" method is used, because according to the research work by Khan and Imtiaz [25] that compare different fiducial points detection methods, it is the most widely used approach for RR peak detection because it offers high accuracy. These methods are also the default implemented algorithms for fiducial points localisation in the Neurokit2 package. In the next step, inbuilt functions in the Biobss library calculate 39 morphological features from the extracted ECG locations (peaks and wave onset/offset points). The Biobss library splits the ECG into segments using the detected fiducial points, where for each segment peak amplitudes, intervals and ratios are calculated as features. The library provides two functions: biobss.ecgtools.ecg_features.from_Rpeaks and biobss.ecgtools.ecg_features.from_waves. "from_Rpeaks" takes as reference four consecutive R peaks and calculates the corresponding (current) R peak amplitude, RR intervals (the RR interval before the current R peak and two RR intervals after the current R peak), ratios and the mean of the intervals (3.3.1, row 1-8). "from_waves" takes as reference two consecutive R peaks and calculates the corresponding P, Q, R, S and T amplitudes, their intervals and ratios (3.3.1, row 9-39). Each feature is an average from all segments in the ECG signal. After feature extraction some feature vectors had "nan", "-inf" or "inf" entries, which have been manually imputated by means. The features are then used to train a random forest classifier. For this, the work utilises the Sklearn Python framework. The classifier results are discussed in section 4 and 4.2.2.

Biobss features						
Features (averaged)	Description					
a_R	Amplitude of R peak					
RR0	Previous RR interval					
RR1	Current RR interval					
RR2	Subsequent RR interval					
RRm	Mean of RR0, RR1, and RR2					
RR_0_1	Ratio of RR0 to RR1					
RR_2_1	Ratio of RR2 to RR1					
RR_m_1	Ratio of RRm to RR1					
t_PR	Time between P and R peak locations					
$t_{-}QR$	Time between Q and R peak locations					
t_SR	Time between S and R peak locations					
$t_{-}TR$	Time between T and R peak locations					
t_PQ	Time between P and Q peak locations					
t_PS	Time between P and S peak locations					
$t_{-}PT$	Time between P and T peak locations					
$t_{-}QS$	Time between Q and S peak locations					
$t_{-}QT$	Time between Q and T peak locations					
t_ST	Time between S and T peak locations					
t_PT_QS	Ratio of t_PT to t_QS					
$t_{-}QT_{-}QS$	Ratio of t_QT to t_QS					
a_PQ	Difference of P wave and Q wave amplitudes					
a_QR	Difference of Q wave and R wave amplitudes					
a_RS	Difference of R wave and S wave amplitudes					
a_ST	Difference of S wave and T wave amplitudes					
a_PS	Difference of P wave and S wave amplitudes					
a_PT	Difference of P wave and T wave amplitudes					
a_QS	Difference of Q wave and S wave amplitudes					
a_QT	Difference of Q wave and T wave amplitudes					
a_ST_QS	Ratio of a_ST to a_QS					
a_RS_QR	Ratio of a_RS to a_QR					
a_PQ_QS	Ratio of a_PQ to a_QS					
a_PQ_QT	Ratio of a_PQ to a_QT					
a_PQ_PS	Ratio of a_PQ to a_PS					
a_PQ_QR	Ratio of a_PQ to a_QR					
a_PQ_RS	Ratio of a_PQ to a_RS					
a_RS_QS	Ratio of a_RS to a_QS					
a_RS_QT	Ratio of a_RS to a_QT					
a_ST_PQ	Ratio of a_ST to a_PQ					
a_ST_QT	Ratio of a_ST to a_QT					

3.3.2 Encoder-based Transformer

As second investigated model a standard Transformer encoder model is utilized. Two types of the Transformer model are investigated within this work. First a "standalone" Transformer encoder-based model and secondly an own modified version of the Transformer model is proposed that uses a Convolutional Network based on several residual convolution-inception blocks as

input features and which will be described in section 3.3.6. The main parts of the Transformer encoder block applied in this work are similarly to the standard Transformer encoder block from the original paper [1]. As described in section 3.1 and shown figure 3.1, the Transformer encoder block is composed of a positional encoding layer, a Multi-head Attention layer, a layer-normalization layer, a feed-forward layer and again a layer-normalization layer with residual connections in between. Since the Transformer encoder model is intended to learn relational features from embeddings, this version of the Transformer model segments each ECG into fix-sized embeddings. For example, an ECG with 2000 datapoints is reshaped into 40 segments each with 50 datapoints. This operation refers to the inputted embedding layer in 3.1. The positional encoding layer that adds temporal information to the embeddings is an optional layer, which is conducted in the experiments 4.2. In the experiments different parameter configurations of the Multi-head Attention layer are analysed, e.g. the performance of stacking several Transformer encoder blocks.

3.3.3 Residual CNN with and without dilation

As third model two residual Convolutional Networks are investigated, based on the idea of ResNet [16]. Both networks consist of 6 residual blocks. The residual blocks are composed of two 1D convolutional layers with kernel size 5x1, strides 1 and no padding, where the channels from the first convolution layer are connected via skip connection to the last layer of each block. The second convolutional layer is followed by a batch normalization layer, a feed-forward layer using leaky relu for extracting non-linear features and a dropout layer with rate 0.1. The final output of the dropout layer is connected by the skip connection from the first convolution. The channel maps increase per residual block with 16, 32, 64, 128, 256 and 512. Furthermore, between each residual block a 1D average pooling layer with stride 2 is applied, which halves the signal length and features after each residual block. Finally, the model is concluded by a global average pooling layer and a dense layer to the number of outputted classes using sigmoid activations. The corresponding model graph is given in figure graph 3.5. The second residual Convolutional Network experiments with dilated convolutions using similarly a kernel size of 5x1, although with a dilation rate of 2.

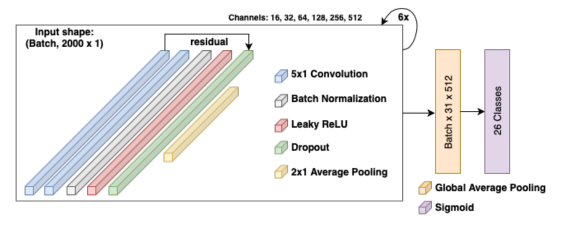


Figure 3.5: Residual CNN architecture

3.3.4 Multi-scale Convolutional Network

The fourth investigated method adopts the idea from the Inception Network. "Inception" proposed by Szegedy et al. in "Going deeper with convolutions" [38] set new state-of-the-art performance on the ImageNet classification challenge [35] in 2014, while it is a 20 layers deep network it uses much fewer trainable parameters. The idea of inception is to use several convolutional filters in parallel in the same convolutional layer, e.g. in the original paper 1x1, 3x3 and 5x5 have been used. The outputted channels are then concatenated into one feature map block for further processing. The idea behind 1x1 convolution is that it reduces dimensionality. Unlike pooling, which takes for example the minimum, maximum or average values within a channel, 1x1 convolutions apply dimension reduction across channels on channel depth by learning a set of weights equal to the number of channels. The expression 1x1 convolution might be a bit misleading, because more precisely it applies a $n_{number\ of\ channels}$ x1 convolution in which each channel value is weighted-based summed into one channel. The overall strength in inception is extracting and combining features from different kernel sizes, while keeping the dimensional overhead low using 1x1 kernels. Although, the term "attention" is not directly used in the paper [38], the 1x1 convolution might be compared to an attention operation, since it learns a set of linear weights that weight channel information, while channel-attention, which is described in the next section 3.3.5, incorporates non-linear functions, e.g. relu and sigmoid, to weight channels base on a set of learnable attention weights. This work makes a few modifications to the proposed Inception Network [38]. The modified inception block in this work reverses a little bit the idea, as it does not apply a max-pooling operation or 1x1 convolutions within the inception block [38]. Instead it is intended to use multi-scale kernels based on the idea of combining many small and large sized kernels, i.e. 3x1, 5x1, 9x1 and 15x1 in parallel. The modified version consists of 4 blocks and is shown in figure 3.6.

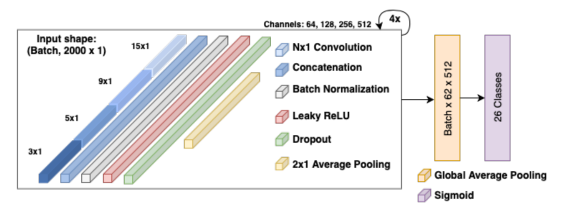


Figure 3.6: Multi-scale Convolutional Network

3.3.5 Squeeze and Excitation Network

The fifth investigated method is channel-attention implemented through a network architecture, called Squeeze and Excitation Network (SENet) proposed by Hu et al. [21]. The model set new state-of-the-art performance on the ImageNet classification challenge [35] in 2017. The application of this model is motivated, as it uses a related variant of an attention mechanisms and is incorporated within the residual Convolutional Network of the second and fourth ranked

model from the Physionet 2021 challenge [15] [37]. The idea of channel-attention is to recalibrate the channels of a convolutional layer. This is achieved by learning in a parallel "Squeeze and Excitation Network" a set of attention weights. In a first step, the Squeeze and Excitation Network "squeezes" all channels from the previous convolutional layer by applying global average pooling. The channel values are then fed into a two hidden layer feed-forward network using relu activations in the first layer and sigmoid activations in the second layer. The network is intended to learn inter-dependency between the channels. In the original paper [15] the authors experiment with the size of the hidden layer. They use a dense ratio to the number of channels, to evaluate the performance of the network and which has been emperically determined, e.g. 2-32, while smaller ratios increase the block sizes. The authors find that are ratio of 16 offers a good balance between accuracy and model complexity. The dimensionality is then increased in the second hidden layer to the number of channels using sigmoid activations. The outputted thresholds are used as channel-attention weights to scale the channels from the inital convolutional layer, which are multiplied by a residual connection parallel to the Squeeze and Excitation Network, also described as "excitation operation". An improved version of the Squeeze and Excitation network is the Convolutional Block Attention Module (CBAM) proposed by Woo et al. [44]. CBAM uses two inputs for the attention maps, channel-attention and spatial-attention, which are obtained from both, a max and average pooling operation. As comparison model this work experiments with the CBAM approach. It uses 3 CBAM blocks with 64, 128 and 256 channels. The motivation is to directly evaluate the performance of channel-attention in the own experiments, while considering the model independently, compared to [15] and [37], which use the model in combination with other approaches.

3.3.6 Own approach

The own approach is motivated by the idea of applying Multi-head Attention on feature maps obtained from multi-scale convolutions. Similarly as illustrated in 3.3.4 it uses 1x3, 1x5, 1x9 and 1x15 kernels to capture small and large sized features. 256 channels, from which each 64 correspond to a different kernel size, are then forwarded into 3 consecutive Transformer encoder blocks to capture spatial relations among the channels. Each Transformer encoder block is composed of the same Transformer encoder layers as described in 3.3.2. The Multi-head Attention layer uses 8 heads with $qkv_{dimension}$ 32. The feed-forward hidden layer is 24. The corresponding model is illustrated in 3.7. The Transformer encoder block does not incorporate

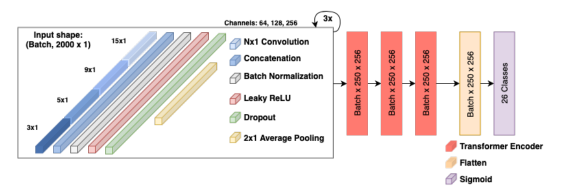


Figure 3.7: Multi-scale channels combined with multiple Transformer encoder

positional encoding. The dimensions within each Transformer encoder remain equal. The output

of the Transformer blocks are feeded into a global average pooling layer and a final dense layer times the 26 classes using sigmoid activations. Previously a flattening layer has been conducted, which did let the model overfit to quickly.

3.3.7 Fine-tuning

The Physionet 2021 challenge organisiers add partial rewards to misclassifications that lead to similar cardiac treatments. Therefore the challenge score is calculated by incorporating class weights for each correct and incorrect classification. To fine-tune the performance of the models on the Physionet 2021 challenge metrics the presented approaches adopt an altered custom lost function used by the first ranked model of the Physionet 2021 challenge [27], which was initially proposed by Vicar et al. [39] [40]. The custom loss function incorporates the evaluation weights of the challenge and decreases the total loss as follows [27]:

$$Loss = \sum_{hom} \left(BCE(T, P) - CL(T, P) + SL(P) \right)$$
 (3.4)

$$CL(T, P) = \sum_{i,j=1}^{i,j=class n} \omega_{ij} a_{ij}$$
(3.5)

$$SL(P) = average(-4P(P-1))$$
(3.6)

The calculated total loss of the mini-batch is composed of three functions, first the standard binary cross-entropy loss for multi-label classification (BCE), second a custom loss function (CL) and third a sparsity loss function (SL). T and P are 26-dimensional vectors that contain the true labels as binary values and predicted output classes as continuous values. The sparsity loss (SL) function adds a penalty value ranging between 0 to 1 for each outputted logit that is close to 0.5. It is a down concaved parabolic curve with vertex at 0.5 and roots 0 and 1. The sparsity loss is intended to improve the final threshold optimization, as it forces the model to output values close to 0 or 1. The custom loss function incorporates the evaluation weights w_{ij} , where a_{ij} represent a multi-label confusion matrix A, which use a continuous version of the logical OR. This confusion matrix is calculated by equation A and N, where N is a normalizing constant obtained from the continuous version of the logical OR [39]:

$$A = T^T \left(\frac{P}{N}\right) \tag{3.7}$$

$$N = ((T + P - T \odot P) \mathbf{1}_{c \times 1}) \mathbf{1}_{1 \times c}$$

$$(3.8)$$

 1_{cx1} and 1_{1xc} are 26-dimensional vectors consisting of only ones, $\frac{P}{N}$ and \odot are point-wise division and multiplication operators.

The proposed deep-learning approaches in this work are fine-tuned based on this compound cost function to maximize the challenge score. The fine-tuning is performed on a small top network, which is concatenated to each base model after the pre-training. The small network consists of one hidden-layer and an output layer, both with dimension 26 and sigmoid activations. Therefore, the deep-learning models are pre-trained and treated as base models with standard binary cross-entropy loss for multi-label classification (BCE) for at most 100 epochs trained to maximize the validation accuracy and then concatenated with the small top layer network and fine-tuned for 15 epochs to maximize the challenge score. This idea is adopted from [27], which

utilize the cost function in combination with a small fine-tuned network that do not propagate the fine-tuned gradients to the base model. Additionally, a grid search threshold optimization, iterating 1000 decimal places, is performed on the fine-tuned model that optimizes the F1 score for each class independently.

Chapter 4

Evaluation

4.1 Datasets

4.1.1 Physionet 2021 challenge database

This section summarises the publicly available Physionet 2021 challenge database that is used to train all models within this work. In section 4.1.2 the MyDiagnostick data provided by University of Maastricht is briefly discussed. The Physionet 2021 challenge database provides 12-lead ECGs annotated by cardiologist experts and is a diverse dataset collection from seven sources in four countries and three continents (see Physionet 2021)[31]. The sources of the challenge data include the Chapman-Shaoxing database and Ningbo database, PTB and PTB-XL database, Georgia 12-lead challenge data, CPSC and CPSC-extra database and the INCART database. In total the challenge data contains 88,253 publicly shared 12-lead ECG recordings. The annotations contain labels from more than 100 known arrhythmia types (see all Physionet 2021 database arrhythmias). Each arrhythmia class has a 'Snomed CT code' assigned, an unique ID representing each arrhythmia type. The ECGs are multi-label annotated, meaning that multiple arrhythmia types can occur within the same ECG. Table 4.1 shows an overview of the Physionet 2021 challenge database composition, the sources, recording length, sampling frequency and data proportion.

Dataset source	Recording length in seconds	Sampling frequency	ECG samples	
Chapman-Shaoxing database			_	
& Ningbo database	10s	500 Hz	45,152	
PTB database	10-120s	either 500	22,353	
& PTB-XL database	10-1208	or 1000 Hz	22,333	
Georgia 12-lead	5-10s	500 Hz	10,344	
challenge database	9-108	500 Hz	10,344	
CPSC database	6-144s	500 Hz	10.330	
& CPSC-extra database	0-1448	500 Hz	10,330	
INCART 12-lead database	1800s (30 min)	257 Hz	74	

Table 4.1: Physionet 2021 challenge data composition

This work only uses the published training data, since the test data for the official challenge and evaluation scoring metrics are withhold by the organisers and are not publicly available. Figure 4.1 shows the class distribution for the entire publicly available Physionet 2021 database (top), as well as the distribution of the scored challenge classes (bottom). The official Physionet 2021 challenge uses a subset of 30 classes that are scored within the challenge and the evaluation metrics (see scored Physionet 2021 database arrhythmias). Due to limited samples, eight arrhythmia types are treated as pairs and same class within the challenge metrics, namely Premature Atrial Contractions (PAC) and Supraventricular Premature Beats (SVPB), Premature Ventricular Contractions (PVC) and Ventricular Premature Beats (VPB), Complete Left Bundle Branch Block (CLBBB) and Left Bundle Branch Block (LBBB), Complete Right Bundle Branch Block (CRBBB) and Right Bundle Branch Block (RBBB). Therefore, to benchmark the proposed approaches, the developed models follow these requirements and annotations are similar grouped to output 26 classes instead of 30, while other class annotations and ECGs that are not part of the scored metrics are discarded. Both graphs in 4.1 show that the data is quite unbalanced. Considering the bottom graph that show the scored challenge data distribution, 28,971 (22.40%) Sinus Rhythm samples are provided, while many classes with less than 2,000 (1% <)samples are present. For an unbiased model training, the dataset needs to be pre-processed and properly balanced to avoid overfitting on some major classes, which is described in section 4.1.3.



Figure 4.1: Physionet 2021 classes distribution

4.1.2 MyDiagnostick data

The MyDiagnostick data contains single-lead ECGs (first lead I) with a sampling frequency of 200 Hz and 60 seconds in recording length. For the evaluation of the transferred models on the MyDiagnostick data, part of the experiments will focus only on a subset of 5 classes, namely Sinus Rhythm (SR), Atrial Fibrillation (AF), Atrial Flutter (AFL), Premature Atrial Contractions (PAC) and Premature Ventricular Contractions (PVC). This is due to the limited class annotations in the MyDiagnostick data, which are limited to these class annotations. ECG preprocessing steps to transfer the models are discussed in the next section 4.1.3. Figure 4.2 shows the class distribution of the corresponding classes for both, the Physionet 2021 data and the My-Diagnostick data. The most underrepresented class in both datasets is Premature Ventricular Contraction with 1279 samples, respectively with 160 samples. In many MyDiagnostick samples both classes, Atrial Ventricular Contractions and Premature Ventricular Contractions, are present within the same ECG. Although, the MyDiagnostick provides 10,587 manually annotations, the dataset consists of more than 40,000 samples. The remaining samples are automatically labelled as either Sinus Rhythm or Atrial Fibrillation by an assistive ECG device. As part of this thesis the aim of the experiments will be to evaluate the best performing pre-trained model from the Physionet 2021 challenge data on the manually annotated MyDiagnostick data and to provide annotations for the remaining 30,000 samples, which will be discussed in section 4.2.2.

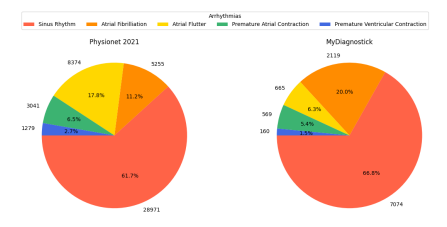


Figure 4.2: Physionet 2021 and MyDiagnostick class distribution - SR, AF, AFL, PAC and PVC

4.1.3 Pre-processing

This section summarises the pre-processing steps that have been applied on both datasets to address data inequalities. Pre-processing steps include train-test split by patient, class balancing, zero padding/truncation, resampling, normalisation, filtering. The train-test splits are: 50% training, 25% validation and 25% test set. The Physionet 2021 challenge data was previously split by patient to ensure that the models learn general features rather than specific features of individual patients. The MyDiagnostick data is only utilized for transferring and testing the pre-trained models from the Physionet 2021 challenge data, meaning the MyDiagnostick data is not used for fine-tuning. As in figure ?? shown, the training data is highly unbalanced. To

address this issue, standard class down- or upsampling is applied on the Physionet data using the Python package imbalanced-learn. In the case of the 26-classes models up- and downsampling is set to 2000 samples by randomly discarding or duplicating some ECGs. In the case of the 5-classes models down- and upsampling is set to 6000 samples. The class upsampling was applied after splitting the training data by patients into train, validation and test sets to avoid evaluating on trained ECGs. However, since the data is multi-label annotated the class balance is only to some extent achieved, because imbalanced-learn does not support class balancing for multi-label classification. The workaround solution was to transform the multi-label annotations into multi-classification annotations and balance it from there, while during the recovering process of the annotations the labels have been ascending sorted from minor to major classes and duplicates are ignored for the major classes. The class balancing results are shown in figure 4.1.3. The second graph on the left for the 26 balanced train set classes shows still quite unbalanced data, which is probably due that many minor arrhythmias contain more than one label with one of the frequent classes, e.g. Sinus Rhythm.

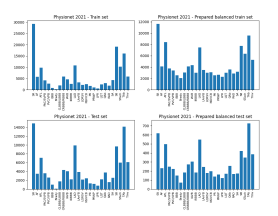


Figure 4.3: Prepared Physionet 2021 data (All scored classes)

Figure 4.4: Prepared Physionet 2021 data (5 classes)

Since the Physionet 2021 challenge data consists of 12-lead ECGs and the MyDiagnostick data consists of single-lead ECGs (first lead I), all models in the experiments are pre-trained on the first lead from the Physionet data. Most of the ECGs in the Physionet data are recorded at a sampling frequency of 500 Hz, while the MyDiagnostick ECGs are recorded at a sampling frequency of 200 Hz. Most of the Physionet 2021 challenge ECGs contain 10 second long recordings (5000 data points for 10 seconds - see 4.1). In comparison the MyDiagnostick data contains 60 seconds long recorded ECGs with 12000 data points (2000 data points are equal to 10 seconds). To address the data inequality, first each ECG in the Physionet 2021 challenge is truncated or zero padded to 5000 data points. Next, each Physionet ECG is downsampled to 200 Hz, yielding 2000 data points. This makes both datasets uniform in terms of sample density and temporal resolution. In addition, each ECG is normalised within the range of -1 and 1 using min-max scaling. This is particularly helpful in reducing signal amplitude variation due to differences in electrode placement, body size and individual heart activity. In addition, normalisation helps to stable training with gradient descent and avoids gradient explosion problems during backpropagation. By keeping the model weights in an uniform range prevents the model from being biased towards certain input features (e.g. R peaks). In the next step, a standard Butterworth bandpass filter is applied with a bandwidth of 0.3 Hz to 21 Hz, which reduces small noise fluctuations in the ECG signal and facilitates the model to learn key features from arrhythmias, such as RR intervals. Finally, all ECGs in the Physionet data are again zero padded to 12000 datapoints. This is necessary, because a model applied on both datasets need to receive as input an equal sized feature vector. Figure 4.5 visualizes step by step the ECG pre-processing procedure. The last step after the filtering is only utilized for the model that is transferred and tested on the MyDiagnostick data. The last is an ECG example from the MyDiagnostick data to show the mismatch between the lengths of the ECGs, which only needed to be normalised and filtered as it is already present with a sample frequency of 200 Hz.

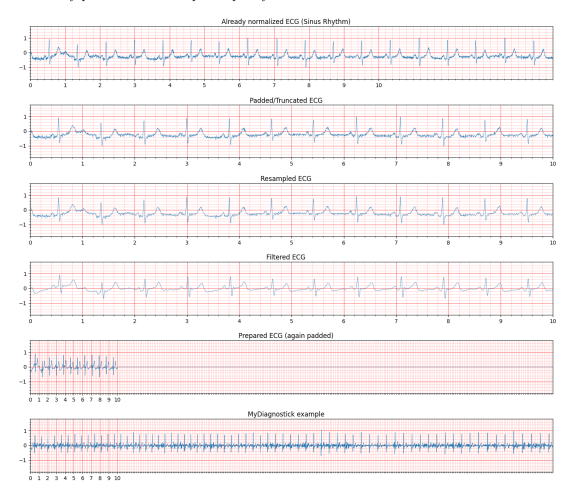


Figure 4.5: ECG pre-processing steps applied to the Physionet 2021 data

4.2 Experimental setup and results

This section describes the experimental setup and results of the developed approaches. The section is divided into two parts. The first part discusses the experimental setup and results for the models trained and tested on the Physionet 2021 database. The second part discusses the experimental setup and results of the pre-trained models on the Physionet 2021 database transferred and evaluated on the MyDiagnostick data. All models have been trained in Google

Colab using an Nvidia T4 GPU with 16GB VRAM for at most 100 epochs and early stopping enabled with regard to not decreasing validation loss (patience set to 5). Adam optimizer is used with an initial learning rate of 0.001, which decreases by 1/10 with regard to not decreasing validation loss (patience set to 5). As loss function binary-crossentropy for multi-label classification is used, because the Physionet 2021 and MyDiagnostick data are both multi-label annotated. All models use sigmoid activations in the final output layer times the number of classes. For simplicity, all models are trained with a fixed mini-batch size of 32, although which could be determined empirically. Due to minimize computational costs and required training time, as some developed model took up to one hour for training, the main part of these experiments focus on a fixed train-test split, meaning that the train-test samples within these experiments are identically for each tested model as described in 4.1.3. Using scikit-learn the weighted average of accuracy, precision, recall and f1 is computed for each model. In addition, the official Physionet 2021 evaluation metrics are calculated to benchmark the own developed approaches with other approaches within the challenge, for other model results see challenge results. The official Physionet challenge metrics use for evaluation AUROC, AUPRC, Accuracy, F1 measure and the own calculated challenge score.

4.2.1 Physionet 2021 challenge

This section will answer the first two research questions of this work based on the Physionet 2021 challenge data:

- 1. How well does a Transformer-based model perform on the Physionet 2021 challenge data compared to a feature-based model or an Convolutional Network?
- 2. Can an ensemble model of Transformer and Convolutional Network effectively capture spatio-temporal information and improve accuracy?

As part of the experiments to address the above research questions several variations of a standard Transformer encoder block without any feature extraction, e.g. convolution layers, are trained on the raw Physionet ECGs for the classification into 26 classes. Gridsearch is applied on various parameters. Conducted parameters include input shape, positional encoding, number of stacked encoder blocks, number of heads, dimensions of the projection matrices q, k or v, feed-forward layer dimension and dropout rate. The input shape describes in how many embeddings the inputted ECG sequence is splitted before feeded into the Transformer encoder block. (40, 50) means that the pre-processed ECG with 2000 datapoints in length is splitted into 40 temporal consecutive segments (embeddings), where each embedding contains 50 datapoints. Number of heads and q, k and v dimension describe the amount of trainable parameters of the encoder block, i.e. whether several heads are used in each encoder block and the size of the q, k and v matrices that project the embeddings into the feature space. Feed-forward dimension determines the hidden-layer dimension of the feed-forward layer that is part of each encoder block. Dropout is applied to all layers, either set 0.1 or 0.4. The corresponding results are below in table 4.2 shown.

Input shape	Pos. enc.	Enc. blocks	Heads	qkv dim	ff dim	Drop- out	Train. param.	Acc.	Prec.	Rec.	F1
(40, 50)	True	1	1	25	24	0.1	155.375	0.096	0.514	0.120	0.194
(40, 50)	True	1	1	25	24	0.4	155.375	0.065	0.418	0.083	0.139
(40, 50)	True	8	1	25	24	0.1	878.818	0.061	0.579	0.079	0.139
(40, 50)	True	8	1	25	24	0.4	878.818	0.098	0.592	0.122	0.203
(40, 50)	False	1	1	25	24	0.1	155.375	0.227	0.747	0.25	0.374
(40, 50)	False	1	1	25	24	0.4	155.375	0.224	0.742	0.253	0.378
(40, 50)	False	8	1	25	24	0.1	878.818	0.228	0.765	0.261	0.389
(40, 50)	False	8	1	25	24	0.4	878.818	0.226	0.737	0.255	0.379
(10, 200)	False	8	1	25	24	0.1	1.004.818	0.160	0.744	0.174	0.283
(10, 200)	False	8	8	25	24	0.1	2.129.018	0.177	0.709	0.197	0.308
(40, 50)	False	8	8	400	24	0.1	6.035.018	0.223	0.762	0.247	0.374
(40, 50)	False	8	8	25	2048	0.1	65.947.210	0.219	0.740	0.257	0.382
(40, 50)	False	8	8	400	2048	0.1	70.819.210	0.226	0.751	0.257	0.383
(40, 50)	False	8	8	400	2048	0.4	70.819.210	0.245	0.739	0.286	0.413

Table 4.2: Standard Transformer encoder evaluated on the Physionet 2021 challenge data

Based on the experimental results it can be seen that a standard Transformer encoder-based model is able to extract relational features from the ECGs for the classification of 26 different arrhythmias. However, in terms of accuracy there is still room for improvements, while precision is higher. Using smaller ECG segments as input sequence, e.g. (40, 50) instead of (10, 200), offers slightly improved performance. Surprising is that positional encoding negatively affects the performance. A reason for this might be that the variance of the inputted ECG sequence segments is to high, for example the wave segments are not equally aligned within the each segments across the samples. By adding positional information it might on top confuse the Multi-head Attention network to extract meaningful relations among the segments. Based on the experiments, the size of the model and amount of trainable parameters, e.g. number of encoder blocks, q, k and v dimensions and feed-forward layer size, improves the model performance, but which should be critically seen in terms of required training costs, especially considering a model with 70,000,000 parameters. The effect of a higher dropout rate varies among the experiments, although in the last model it slightly improves the generalization ability of the model.

The next experiments on the Physonet 2021 compare all previously discussed models on the multi-label classification of 26 arrhythmias. 4.3 and 4.4 show the evaluation scores after the models have been fine-tuned and optimized by thresholds. 4.3 has two values for accuracy, because the first accuracy correspond to the accuracy score before the models have been fine-tuned and threshold optimized to the F1 score. All other values correspond to the fine-tuned versions. Based on the experiments it can be seen that the feature-based random forest yields the highest accuracy. Although most models offer comparable results, the Transformer encoder performs a bit lower than other models. Even the own proposed approach is not able to show good results, also the results partial better than only the Transformer encoder. Based on the challenge metric all models, except the random forest, are able to achieve high performance, which is because of the fine-tuning with the adopted cost function [39] [40].

Model	Parameters	Accuracy	Precision	Recall	F-measure	
Random Forest	,	0.390	0.714	0.382	0.406	
(Biobss features)	/	0.590	0.714	0.362		
Residual CNN	3.702.282	0.387-0.310	0.593	0.692	0.625	
(without dilation)	3.702.202	0.367-0.310	0.595	0.092		
Residual CNN	3.702.282	0.386-0.311	0.590	0.695	0.624	
(with dilation)	3.102.202	0.300-0.311	0.590	0.090	0.024	
Transformer Encoder	919,418	0.225-0.100	0.380	0.619	0.432	
(8 heads, 8 blocks)	919,418	0.225-0.100	0.300	0.019	0.432	
Multi-scale	53.202.522	0.358-0.299	0.581	0.673	0.609	
Convolutional Network	03.202.322	0.556-0.299	0.561	0.073	0.009	
Modified Squeeze and	152.210	0.257-0.184	0.463	0.622	0.501	
Excitation Network	192.210	0.201-0.104	0.405	0.022	0.501	
Multi-scale Kernels and	11,105,058	0.3310.314	0.503	0.618	0.540	
Transformer Encoder	11,100,000	0.5510.514	0.505	0.016	0.540	

Table 4.3: Physionet 2021 train/test split evaluation

Model	AUROC	AUPRC	Accuracy	F-measure	Challenge metric
Random Forest (Biobss features)	0.560	0.142	0.390	0.153	0.147
Residual CNN (without dilation)	0.874	0.439	0.310	0.482	0.556
Residual CNN (with dilation)	0.842	0.404	0.311	0.442	0.554
Transformer Encoder (8 heads, 8 blocks)	0.701	0.211	0.100	0.258	0.402
Multi-scale Convolutional Network	0.832	0.382	0.299	0.414	0.526
Modified Squeeze and Excitation Network	0.768	0.290	0.184	0.322	0.445
Multi-scale Kernels and Transformer Encoder	0.822	0.358	0.265	0.402	0.514

Table 4.4: Physionet 2021 challenge metric scores

4.2.2 MyDiagnostick

This section will answer the last two research questions of this work based on the Physionet 2021 challenge data and MyDiagnostick:

- 1. How is the performance of the most promising model at discriminating SR, AF, AFL, PAC and PVC on both datasets?
- 2. What are the challenges in transferring the pre-trained models from the Physionet 2021 challenge data to the MyDiagnostick database? Do the models generalise well, even though different ECG devices were used?

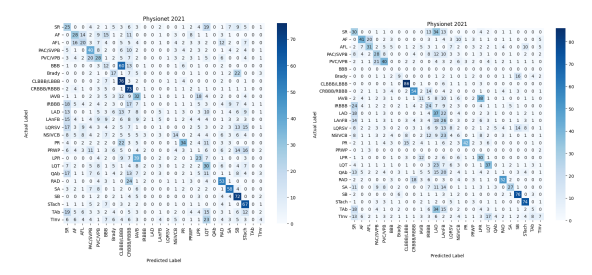


Figure 4.6: Results of several runs on the Physionet 2021 data

Model	Accuracy	Precision	Recall	F-measure
Residual CNN				
(without dilation)	0.402	0.744	0.414	0.432
5 classes				
Residual CNN				
(without dilation)	0.111	0.899	0.210	0.240
26 classes				

Table 4.5: MyDiagnostick model evaluation

Chapter 5

Conclusion

5.1 Discussion

5.2 Summary and Outlook

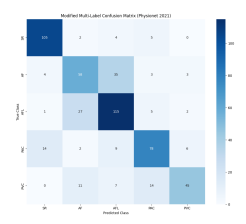


Figure 5.1: Results of the 5-classes residual CNN model tested on the Physionet 2021 data

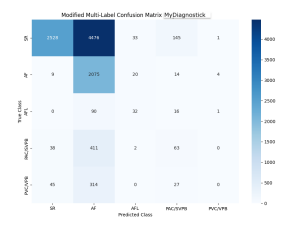


Figure 5.2: Results of the 5-classes residual CNN model transferred and tested on the MyDiagnostick data

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Appendix A

Implementation

```
# Transformer Encoder single-lead (TensorFlow)
def get_positional_encoding(seq_length, d_model):
   position = np.arange(seq_length)[:, np.newaxis]
   div_term = np.exp(np.arange(0, d_model, 2) * -(np.log(10000.0) / d_model))
   pe = np.zeros((seq_length, d_model))
   pe[:, 0::2] = np.sin(position * div_term)
   pe[:, 1::2] = np.cos(position * div_term)
   pe = pe[np.newaxis, :]
   return tf.constant(pe, dtype=tf.float32)
def transformer_encoder_block(input, input_shape, num_heads, key_dim, ff_dim, dropout):
   # Multi-Head Attention
   x = keras.layers.MultiHeadAttention(num_heads=num_heads,
   key_dim=key_dim, dropout=dropout, kernel_regularizer=regularizers.12(0.001))(input, input)
   # Add & Normalize
   res = x + input
   x = keras.layers.LayerNormalization(epsilon=1e-6)(res)
   # Feed-Forward Layer
   x = keras.layers.Flatten(input_shape=input_shape)(x)
   x = keras.layers.Dense(units=ff_dim, activation='relu', kernel_regularizer=regularizers.12(0.
   x = keras.layers.Dense(input_shape[0] * input_shape[1], kernel_regularizer=regularizers.12(0.
   x = keras.layers.Reshape(input_shape)(x)
   x = keras.layers.Dropout(rate=dropout)(x)
   # Add & Normalize
   x = x + res
   x = keras.layers.LayerNormalization(epsilon=1e-6)(x)
   return x
def build_encoder(num_classes, input_shape, positional_encoding, num_encoder_blocks, num_heads, k
    inputs = keras.Input(shape=input_shape)
   x = inputs
    if positional_encoding:
        positional_encoding_values = get_positional_encoding(input_shape[0], input_shape[1])
```

```
x = x + positional_encoding_values
for _ in range(num_encoder_blocks):
    x = transformer_encoder_block(x, input_shape, key_dim, num_heads, ff_dim, dropout)
x = keras.layers.Flatten(input_shape=input_shape)(x)
outputs = keras.layers.Dense(num_classes, activation='sigmoid')(x)
return keras.Model(inputs, outputs)
```

Appendix B

Graphics

t.b.c.