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MASTER THESIS

A comparison of Time Series Classification Algorithms based on their ability to learn on diminishing time series

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A comparison of Time Series Classification Algorithms based on their ability to learn
in an Early Classification Context

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Abstract

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List of Abbreviations

ABR	Auditory Brainstem Response
BOSS	Bag of SFA Symbols
CNN	Convolutional Neural Networks
DFT	Discrete Fourier Transformation
DTW	Dynamic Time Warping
ECTS	Early Classification of Time Series
EE	Elastic Ensemble
ERP	Edit Distance with Real Penalty
eTSCA	Early Time Series Classification Algorithms
FS	Fast Shapelets
HM	Harmonic Mean
IG	Information Gain
i.i.d	Independent and Identically Distributed
KNN	K-Nearest Neighbor Algorithm
KNNED	K-Nearest Neighbor using Euclidean Distance
LCSS	Longest Common Subsequence
LOOCV	Leave One Out Cross Validation
LS	Learned Shapelets
MCB	Multiple Coefficient Binning
MPL	Minimum Prediction Length
MSM	Move-Split-Merge
MTSC	Multivariate Time Series Classification
PAA	Piecewise Aggregate Approximation
PF	Proximity Forest
SAX	Symbolic Aggregate Approximation
SFA	Symbolic Fourier Approximation
ST	Shapelet Transform
TSC	Time Series Classification
TSCA	Time Series Classification Algorithms
TSF	Time Series Forest
TWE	Time Warp Edit
WDTW	Weighted Dynamic Time Warping

Chapter 1

Introduction

Time Series Classification (TSC) is a research field which has grabbed a lot of attention in the data mining community during the last decade. This goes back to the fact that time series data exists, by nature, in numerous scenarios in our life; medical examination records of patients [GM01; GO12], consumption of electrical devices in smart homes and smart grids [Gao+14; JZ14] and astronomy [Pro+06; RK13] are only few examples of them.

The problem of time series data classification has been tackled in different ways and motivated by various objectives. When the main driving objective is to attain high classification accuracy, then this is referred to simply as Time Series Classification (TSC). TSC has also been extended to cover space and time complexity along with accuracy. On the other hand, if time plays a critical role in the problem being tackled beside accuracy; like early detection of malfunctions in industrial processes or diagnosis of a patient with a life threatening disease. In this case the classification problem is referred to as Early Time Series Classification (eTSC).

A lot of algorithms have been introduced to tackle the problem of TSC. According to the [Bag+17], these algorithms can be divided, based on their technique, into six groups. Distance-based time series algorithms compare instances of time series, usually using 1-NN classifiers and an elastic distance measure between the data points of both time series. Phase dependent interval algorithms operate by extracting informative features from intervals of time series, they are more suitable for long and noisy data than distance-based time series algorithms. Phase independent interval algorithms are used when a class can be identified using a single or multiple patterns regardless of when they occur during the time series. Dictionary based algorithms consider the number of repetitions of patterns as a factor of classification and not just simple occurrence of one. Ensembles combine the power of different algorithms, either of different or same core technique, then make the final classification decision based on voting. In addition to the previous algorithms, there are also deep learning time series algorithms which build classifiers using generative as well as discriminative neural networks models.

On the other hand, Early Time Series Classification algorithms are designed to deal with less data in order to achieve earliness of prediction, without sacrificing accuracy as much as possible. Since accuracy and earliness are contradicting by nature, eTSC's main goal is to find a tradeoff between both objectives [Gup+20]. Many of the techniques that have been applied in TSC have also been applied for eTSC. Using 1-

NN classifiers is a very popular TSC technique, which has been extended for eTSC by combining it with Minimum Prediction Length (MPL)[XPP12]. Phase independent intervals[GO12; He+15], generative and discriminative model based classifiers[LLF14; Che+18] as well as deep learning algorithms [HLT18; Ruß+19] have all been extended to tackle eTSC problems.

Both, Time Series Classification Algorithms (TSCA) and Early Time Series Classification Algorithms (eTSCA), have introduced well performing algorithms in terms of their respective performance measures. The existence of comprehensive benchmark data archives [Dau+18; Bag+18] has helped regulate and standardize the process of comparison between classifiers. Algorithms can be benchmarked on a diverse set of datasets with different characteristics to avoid biased results and cherry picking.

Like other fields, the *No free lunch theorem* exists in the TSC domain as well [Bag+17; Bos18]. This means that there is no one specific algorithm that can generalize to all problems and prevail over all other competing algorithms. With that being the case, this leaves an open door for continuous investigation of how the performance of algorithms would change on different data sets and in different contexts.

Inspired by the of the problem of eTSC and its application in diverse areas. We define a situation in which a classifier is put into test to learn on only a part of the available data; to see if it will be able to gracefully degrade. We consider a classifier to be graceful, or fault tolerant, if it is able to maintain statistically significant results by learning on less data in comparison to it's results when learning on full length data. We refer to this situation as the early classification context.

The motivation behind experimenting with such a context is driven by the medical field, where sometimes patients have to go through strenuous tests to be diagnosed. This is specially the case in audiological examinations; as patients are exposed to signals which pushes the limits of their hearing discomfort to be able to examine their problems. A classifier which can attain comparable results while only learning on a fraction of the whole time series data would shorten this process and thus relief patients from their discomfort.

Goal of this Thesis

The goals of this master thesis can be addressed as:

- Provide a solution for adapting existing time series workflows for an early classification context. The solution should run experiments on a group of data sets from the UCR/UEA archive and measure the performance of classifiers within the early classification context. By the end of the experiments the solution should collect the results and provide an analysis of the results, the analysis should:
 - Identify if classifiers can sustain good performance while learning on fewer data points in comparison to learning on the full length
 - Identify if there is a winning classifier across the executed experiments
 - Identify the aspects of the data set which affect the performance of classifiers within the context. Such as train size, number of classes, number of dimensions, length, etc.
- Evaluate the proposed solution
- Evaluate the performance of the different classifiers within the new context on an Auditory Brainstem Response (ABR) data set.

Structure of the Thesis

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Chapter 2

Related Work

The main contribution of this thesis is to provide a framework that compares TSCAs in an early classification context. The idea of creating frameworks to standardize the process of comparison between algorithms is not new to the time series data domain. It was actually needed due to the increasing interest in time series data and the vast amount of newly introduced algorithms. In this chapter we will briefly discuss 3 other frameworks that had been developed to compare TSCA on univariate and multivariate data sets.

2.1 The great time series classification bake off

One of the most famous studies in comparing TSCAs is [Bag+17] and which has inspired some of the work done in this thesis. At the time when this paper was published, the TSC domain had already witnessed hundreds of algorithms being proposed, each claiming superior performance. There was no clear structure, at the time, for experimenting and benchmarking algorithms' performance, which pushed for having a more concrete structure to be followed in succeeding research. The primary goal of the experiment was to assess the average accuracy of the different classifiers on a broad set of univariate data sets. The Scalability and Efficiency of the classifiers were considered as secondary goals.

2.1.1 Inclusion Criteria

The experiment collected 18 different classifiers to be compared with 2 baseline classifiers on the UCR data archive (discussed in 4.1.1). Algorithms were selected based on three criteria; an algorithm had to be published in a high impact conference or journal, used some data sets from the UCR archive to assess performance and the availability of the code.

In order to better describe the various techniques applied by the algorithms and what differences characterize each of them, they formulated a taxonomy which groups algorithms based on the features they try to find to discriminate between different classes. These groups were; whole series similarity (also referred to as distance-based algorithms), phase dependent intervals, phase independent intervals (shapelets), dic-

tionary based and ensemble algorithms. We follow the same taxonomy for our experiment as well and describe these groups in more details in section 3.3.

2.1.2 Experimental Design

The framework ran 100 resampling folds for each of the data sets for each classifier, totalling 8500 (85 data sets \times 100 resamples) experiments for each classifier. The first run used the default train/test split provided with the data set, while the remaining resamples were stratified to preserve the class distribution in the default split. Using the same parameters as the relevant published papers, the experiment limited the number of parameter values for each classifier to a maximum of 100. They used different number of cross validation folds based on the type of algorithm. LOOCV was used for algorithms utilizing distance based measures, while for the other classifiers a new model was created for each parameter set. Resulting in a total of 850,000 models per classifier.

2.2 Deep learning for time series classification: a review

Another paper that focused on comparison of deep learning algorithms for TSC was [Faw+19a]. Deep learning algorithms had already proved competency in many application fields including image classification, speech recognition and natural language processing [He+16; SK16; KSH12; Gua+19] and was gaining more popularity in TSC problems [Zhe+14; Zhe+16; Zha+17]. The main goals were to benchmark the performance of deep learning algorithms on TSC problems; as they were less studied than the other algorithms and to provide an open source framework for deep learning on TSC.

2.2.1 Inclusion Criteria

The framework included 9 discriminative model end-to-end deep learning algorithms developed to work on TSC problems. They started by grouping deep learning algorithms into two main groups; generative and discriminative approaches.

Generative models involve an unsupervised step which tries to learn a quality representation of the time series, then this new representation is fed to another classifier in a learning phase. But this family of algorithms was excluded due to its complexity and incompetence compared to the other group [LGI17; Bag+17].

As for the discriminative models, algorithms that needed preprocessing of features prior to learning were also excluded; to avoid the bias of hand crafted features. The remaining discriminative approaches were; Multi Layer Perceptron (MLP), Fully Convolutional Neural Network (FCN), Residual Network (ResNet), Encoder, Multi-scale Convolutional Neural Network (MCNN), Time Le-Net, Multi Channel Deep Convolutional Neural Network (MDCNN), Time Convolutional Neural Network (Time-CNN) and Time Warping Invariant Echo State Network (TWIESN).

These 9 algorithms were to be compared to each other and to 7 other classifiers. A group consisting of the best 4, out of the 18, classifiers that were included in the experiment by [Bag+17] which are; Elastic Ensemble (EE), Bag Of SFA Symbols (BOSS), Shapelet Transform (ST), Collective of Transformation-based Ensemble(COTE). They

have also included Hierarchical Vote Collective of Transformation-Based Ensembles (HIVE-COTE), which is an extension of COTE using hierarchical voting and two extra classifiers. In addition to Proximity Forest, an ensemble based on the same concept of Random Forests but using class exemplars instead of feature split values. Finally, the classic 1-NN classifier utilizing Dynamic Time warping (DTW) elastic distance measure with the warping window set through cross-validation on training data set. Many of these classifiers we cover later on in section 3.3.

2.2.2 Experimental Design

The experiment covered both univariate data sets and multivariate data sets. The univariate data sets were represented by the same 85 z-normalized data sets from the UCR archive as [Bag+17]. As for the multivariate data sets, 12 data sets from the archive by Mustafa Baydogan¹ were used. Due to the existence of instances with unequal lengths in the multivariate archive, they used linear interpolation, suggested by [RK05], such that all instances inside the same data set are adjusted to the length of the longest instance. Z-normalization was not applied to any of the Baydogan archive.

In order to avoid bias from the initial weights assigned to the classifiers, for each of the data sets for each classifier, the framework used 10 runs for training and then considered the mean accuracy of all runs together. The original train/test split was used in all 10 runs, but the initial weights were randomized. The framework applied optimization for the hyperparameters of the deep learning algorithms, but not for the other classifiers. With the exception of TWIESN, the number of epochs used during optimization ranged between 100 and 5000. A model checkpoint procedure was involved. This meant that after training a model for 1000 epochs, the model which attains the least error on the validation data set is chosen for evaluation. All the models were initialized randomly using the methodology from [GB10] and were optimized using variants of stochastic gradient descent; like Adam [KB14] and AdaDelta [Zei12]. For FCN, MLP, and ResNet if the training loss had not improved for 50 consecutive epochs, then learning rate was decreased by 0.5 to a minimum of 0.0001.

2.3 The great multivariate time series classification bake off

A recent study was carried out by [Rui+20] focusing on comparing multivariate TSCAs. With previous research paying more attention to univariate classifiers and univariate problems, multivariate time series classification (MTSC) recieved less attention. Which meant that the MTSC domain is now in the same position that univariate TSC had been in some years ago and there is a need for a benchmarking framework to guide upcoming research on how to compare multivariate TSCAs to already existing ones.

MTSC is simply the classification of time series data collected, where multiple features are collected at each time point and a single label is assigned. MTSC is more challenging than univariate TSC because the discriminative features can be in the interaction between the multiple dimensions and not only based on the interdependency of consecutive values in one dimension. The framework included two techniques that algorithms follow to deal with multivariate data; either dedicated multivariate

¹<http://www.mustafabaydogan.com/multivariate-time-series-discretization-forclassification.html>

TSCAs which are, by design, able to handle multiple dimensions, or adapted univariate classifiers which can handle multivariate time series instances. The adapted univariate classifiers technique is based on two pillars; assuming independence between the dimensions of the data and fitting separate univariate classifiers to each dimension then ensemble their results. We used this technique in our experiment for handling multivariate data sets.

2.3.1 Inclusion Criteria

Although univariate and multivariate TSC problems differ in their complexity, but the algorithms for both can be grouped, based on the technique they apply, in the same way. These are; distance based similarity, phase dependent intervals, phase independent intervals (shapelets), dictionary based and deep learning algorithms.

The framework collected 16 classifiers from these different groups and compared them on the 30 data sets from the UEA multivariate data archive (discussed in 4.1.2). The included classifiers had to meet 2 simple criteria; code availability and runnable code. The classifiers encompassed 3 variations of the classic DTW which represented the baseline; DTW using Independent Warping (DTW_I) which calculates DTW distance for each dimension separately then sums all distances, DTW using Dependent Warping (DTW_D) which simultaneously calculates the DTW distance across all dimensions for each time point and DTW using Adaptive Warping (DTW_A) which adapts to each instance by calculating both DTW_I and DTW_D and then chooses between them based on a threshold score. The multivariate classifiers were; Generalized random shapelet forest (gRFS), WEASEL+MUSE, Canonical interval forest (CIF), Random Convolutional Kernel Transform (ROCKET), Multiple Representation Sequence Learner (MrSEQL), Residual network (ResNet), InceptionTime, Time series attentional prototype network (TapNet). The adapted univariate classifiers were; the ensemble HIVE-COTE and each of its single components; Shapelet Transform (ST), Time Series Forest (TSF), Contractable Bag of Symbolic Fourier Approximation Symbols (CBOSS) and Random Interval Spectral Ensemble (RISE). Two specialized toolkits in time series machine learning were used for the experiment. One of them is python based, while the other is java based. Most of the algorithms are implemented in both, but few exist only in one. More information is available in [Rui+20].

2.3.2 Experimental Design

The framework ran 30 resamples for each data set for each classifier to calculate performance metrics, the first run using the default train/test split of the data set, while the remaining 29 runs using stratified sampling; to maintain the class distribution of the original split. In contrast to the previous experiments where only accuracy was measured, this experiment had other measures beside which are; area under the ROC, balanced accuracy, F1, negative log likelihood, Matthew's correlation coefficient, recall/sensitivity, precision and specificity. Still accuracy was the primary performance measure used in comparisons; due to its easier interpretation.

The classifiers were initialized using the default structure and hyperparameters in their original published papers. Apart from the classifiers that have internal tuning processes, no other external tunings were applied. Even for the window size of DTW, which is as was proved to have a small but significant improvement of performance for univariate data sets [RK05]. Using a side experiment, where an untuned DTW was compared to a naive implementation of DTW with window sizes between 0% and

100% on 21 of the data sets. The experiment had proved that the untuned DTW was better on 14 data sets, but with no significant difference between both. Which meant that tuning was not needed as far as the experiment is concerned.

The data sets of the UEA archive are not normalized and were presented to the classifiers without any preprocessing. This might have caused a disadvantage for WEASEL+MUSE, which doesn't have an internal normalization process, against other classifiers like, ROCKET; gRSF; TapNet; InceptionTime; ResNet; CBOSS; STC and CIF, which do. The decision of excluding data normalization was based on another side experiment that they carried out. A comparison was done using the three DTW variants, once on normalized data and once on non-normalized data. The experiment had proved that data normalization had no significant effect on the performance of the classifiers, but overall the performance of DTW declined with normalization. The same experiment was repeated for HIVE-COTE and its components and the same result was observed. This meant that data normalization was also not considered important for the scope of the experiment.

2.4 Evaluation Criteria

All three frameworks follow the same methodology, introduced by [Dem06], for comparing their classifiers over multiple data sets. When using multiple resamples over data sets, the score of a classifier on a given data set is calculated as the average of its accuracy over all resamples. The methodology recommends using a Friedman ranking test to refute the null hypothesis; that there is no difference between the ranks of the classifiers. After refuting the null hypothesis and applying ranks to each classifier, a pairwise post-hoc analysis is done according to the recommendation of [BCM16]. Which applies a Wilcoxon signed rank test using a Holm's alpha correction of value 5% to form cliques of classifiers. Each clique represents a group of classifiers where there is no significance between their pairwise comparisons. Results are described by a critical difference diagram, where the average rank of each classifier is noted and cliques are expressed by a thick line.

Chapter 3

Classification Algorithms

This Chapter discusses the definitions and background of the topics mentioned in this thesis. We discuss the nature of time series data, the two problems of time series classification (TSC) and early time series classification (eTSC) then present the different techniques encompassed by them. We will discuss TSCAs in more details as they are the main focus of our framework, but a review of the problem of early classification and a review of its algorithms is also included; as it motivates the context in which the framework operates.

3.1 Time Series Data

3.1.1 Definitions

A time series is a finite sequence of ordered observations, either based on time or another aspect [AML19; Bag+17]. The existence of the time component makes time series an abundant form of data that covers various domains like; medicine, finance, engineering and biology [LTB18].

A time series dataset is a collection of time series instances.

Definition 1

A set T of n time series instances, $T = \{T_1, T_2, \dots, T_n\}$.

Each of the time series instances T_i consists of a sequence of observations collected at consequent time points.

Definition 2

A time series T_i , of length L is represented as $T_i = [t_1, t_2, \dots, t_L]$.

An observation t_i represents the value of the collected measurement at some point of time i .

Time series data can come in different forms. It is important to comprehend what different forms the data can take and what implicit assumptions they convey; to be able to choose the suitable algorithms and tools to deal with it.

The first form of time series data is referred to as univariate time series. This form of data exists when the observations of instances capture a singular value across time.

Definition 3

*Univariate time series T_i , of length L is represented as $T_i = [t_1, t_2, \dots, t_L]$
With t_j as a real valued number.*

The other form is when multiple measurements are captured by the observations. According to [Lön+19], it is essential to differentiate between the two ways multiple time series can be generated; panel data and multivariate time series data. If more than one variable is being observed during a single experiment, with each variable representing a different measurement; this is called multivariate time series.

Definition 4

*Multivariate time series T_i , of length L is represented as $T_i = [t_1, t_2, \dots, t_L]$
With t_j having M dimensions, each one of them is a univariate time series.*

While panel data is when the same kind of measurements is collected from independent instances; like different patients or diverse industrial processes. For panel data, it is possible to assume that the different instances are independent and identically distributed (i.i.d), but this assumption doesn't hold for observation of a single instance. The same goes for multivariate time series, individual univariate observations are assumed to be statistically dependant.

3.1.2 Nature of Time Series Data

Having discussed the dependency assumptions in time the different forms of time series data. It is this dependency that makes time series data challenging for conventional machine learning algorithms, which are used for tabular and cross-sectional data. Tabular and cross-sectional data assume observations to be i.i.d [Lön+19].

If we were to tabularize time series data; convert it into a tabular form by considering each observation as an individual feature. Then it would be possible to apply conventional machine learning algorithms, under the implicit modelling assumption that observations are not ordered. This means that if the order of the features was changed, still the model result will not change. This assumption can work for some problems, but it doesn't have to work for all problems.

3.2 Time Series Classification

Time series classification is a subtype of the general classification problem, which considers the the unique property of dependency between adjacent features of instances [BB17b]. The main goal of time series classification is to learn a function f , which given a training dataset $T = \{T_1, T_2, \dots, T_n\}$ of time series instances along with their corresponding class labels $Y = \{y_1, y_2, \dots, y_n\}$ where $y_i \in \{1, 2, \dots, C\}$, can predict class labels for unseen instances [Den+13].

Time series classification has been studied with different objectives, some papers focused on attaining the highest accuracy of classification as the main goal [Kat16; JJO11; BB17a; LTB18; SL17b; Faw+20], while other papers focused on attaining lower time complexity [RK04; Bag+17; TPW20; Pet+16; SL17a].

In this master thesis, we are more interested in assessing the results in terms of accuracy than time complexity. We define accuracy like [SL20]; as the percentage of

correctly classified instances for a given dataset D , either being a training or testing dataset.

Definition 5

$$Accuracy = \frac{\text{number of correct classifications}}{|D|}$$

3.3 Time Series Classification Algorithms

This chapter will introduce different types of TSCAs and discuss the various techniques that each type apply. There are multiple ways to divide TSCAs in order to better understand them. In this thesis we follow the grouping defined by [Bag+17].

3.3.1 Whole Time Series Algorithms

Whole time series similarity algorithms, also called distance-based algorithms, are methods that compare pairs of time series instances. An unlabeled time series instance is given the class label of the nearest instance in a training data set [Kat16]. There are two main techniques for carrying out the comparison; either by comparing vector representations of the time series instances, or by combining a defined distance function with a classifier, KNN being the most common one [LTB18]. Whole time series algorithms are best suited for problems where the unique features can exist anywhere along the whole time series [Bag+17].

One of the simplest forms of whole time series is 1-NN with Euclidean Distance [FRM94], yet it can surprisingly attain high accuracy compared to other distance measures [XPK10]. But Nearest Neighbor with Euclidean Distance (KNNED) is an easy to beat baseline, due to its sensitivity for distortion and inability to handle time series of unequal lengths [XPK10; Kat16; LTB18]. This lead many of the researchers to focus on defining more advanced and elastic distance measures that can compensate for misalignment between time series [AML19]. The standard and most common baseline classifier utilizing elastic distance measures is 1-NN with Dynamic Time Warping (DTW) [Bag+17]. In contrast to the idea that more powerful machine learning algorithms will be able to defeat the simple KNN and an elastic measure, DTW has proved to be a very tough opponent to other algorithms and other elastic distance measures as well [Kat16; LB15; Wan+13]. But there were also other distance metrics that have been introduced in literature, these include extensions of DTW on one hand like; Weighted Dynamic Time Warping (WDTW) which penalizes large warpings based on distance [JJO11] and Derivative Dynamic Time Warping (DDTW) [KP01; GL13] which uses derivatives of sequences as features rather than raw data to avoid singularities. On the other hand, Edit Distance with Real Penalty (ERP) [CN04], Time Warp Edit (TWE) [Mar08], Longest Common Subsequence (LCSS) [DGM97] and Move-Split-Merge (MSM) [SAD12] are all alternatives for distance measures, yet multiple experiments have considered DTW to be relatively unbeatable [Bag+17; AML19; BB17a]. To the extend of our knowledge, the most powerful whole time series classifiers is Proximity Forest (PF) [Luc+19].

Nearest Neighbor with ED

The Euclidean distance is a remarkably simple technique to calculate the distance between time series instances. Given two instances $T_1 = [t1_1, t1_2, \dots, t1_n]$ and $T_2 =$

$[t2_1, t2_2, \dots, t2_n]$, the euclidean distance between them can be determined as:

$$ED(T_1, T_2) = \sqrt{\sum_{i=1}^n (t1_i - t2_i)^2} \quad (3.1)$$

Euclidean distance has been preferred to other classifiers due to its space and time efficiency, but it suffers from two main shortcomings [BRT13; JJO11; Kat16]. The first one is that it cannot handle comparisons between time series of different lengths. While the second one is its sensitivity to minor discrepancies between time series; it would calculate large distance values for small shiftings or misalignments. Although other metrics have been introduced to overcome the drawbacks of euclidean distance, experimental proof showed that this is only the case for small datasets, but for larger datasets the accuracy of other elastic measures converge with euclidean distance [Hil+14; Din+08; Bag+12].

Nearest Neighbor with DTW

Dynamic Time Warping was a very strong baseline for time series classification for a long time [AML19; Bag+17]. It was first introduced as a technique to recognize spoken words that can deal with misalignments between time series that Euclidean Distance couldn't handle [TPW20]. Figure 3.1 shows a comparison between ED and DTW while computing the distance between two time series instances.

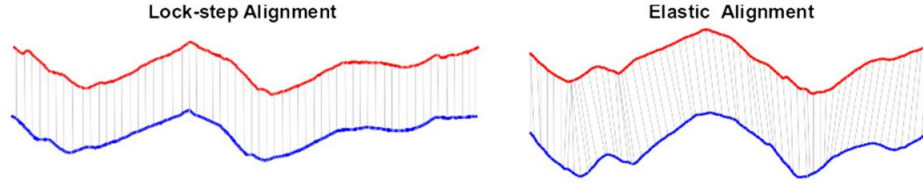


Figure 3.1: Mapping of Euclidean distance (lock-step measure) versus mapping of DTW distance (elastic measure) [AML19].

To calculate the distance between two time series instances $T_1 = [t1_1, t1_2, \dots, t1_n]$ and $T_2 = [t2_1, t2_2, \dots, t2_n]$; a distance matrix $M(T_1, T_2)$, of size $n \times n$, is calculated for T_1 and T_2 . With $M_{i,j}(t1_i, t2_j)$ representing the distance between $t1_i \in T_1$ and $t2_j \in T_2$. The goal of DTW is to find an optimal path that minimizes the cumulative distance between points of T_1 and T_2 .

A candidate path $P = [p_1, p_2, \dots, p_p]$ is to be found by traversing M . For a path to be valid it must conform to three conditions:

- $p_1 = (t1_1, t2_1)$
- $p_p = (t1_n, t2_n)$
- for all $i < m$:
 - $0 \leq t1_{i+1} - t1_i \leq 1$
 - $0 \leq t2_{i+1} - t2_i \leq 1$

Finding an optimal path under DTW can be computationally expensive with complexity of $O(n^2)$ for a time series of length n [SL17a; Pet+16]. Consequently it is usual to use a constraint with the path; to prevent comparison of points outside a certain window [TPW20]; like the famous Sakoe-Chiba Band [SC78], Itakura Parallelogram [Ita75] and Ratanamahatana-Keogh Band [RK04]. Typically DTW can make use of its warping window to handle distortion in time series, but still it is vulnerable to cases where the difference in length between instances length is larger than the warping window [Tan+19].

Nearest Neighbor with MSM Distance

Move-Split-Merge is distance measure that was first introduced in [SAD12]. The main purpose of introducing MSM was to combine certain characteristics within one distance measure. These are; robustness to misalignments between time series instances, being an actual metric unlike other distance measures like DTW and LCSS, assure translation invariance and achieve quadratic run-time [LB15].

The way MSM works is pretty much like other edit distance methods; it determines the similarity between two instances through the usage of a set of operations to transform one of them to the other. These operations, as the name indicates, are; move, split and merge [Bag+17].

Move is the substitution of one single time point of a series with another. Split divides a single time point of a series into two consecutive time points holding the same value as the original time point. Merge can be seen as the opposite of Split, it combines two consecutive time points holding the same value into one time point.

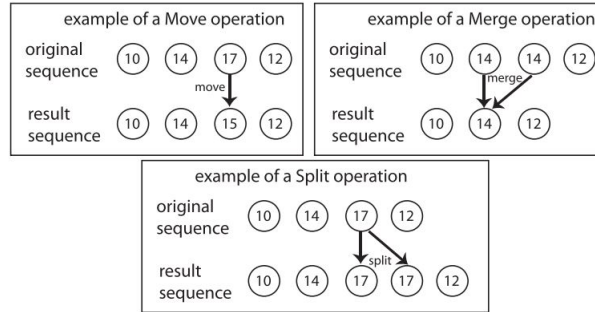


Figure 3.2: Examples of the Move, Split, Merge operations [SAD12].

Each of the previously mentioned operations is associated with a cost. The cost of a move is equal to the absolute difference between the old and the new value of the time point. The costs of split and merge are equal and they are set to a constant to satisfy the symmetry property of metricity [SAD12; TPW20].

Proximity Forest

Proximity forest was developed by [Luc+19]. It was introduced as an addition to scalable time series classification, offering a more scalable and accurate classifier than EE [TPW20]. On one side, EE was an accurate classifier being one the state of the

art algorithms and the best among distance based algorithms, as it combines 11 NN-algorithms each using a different elastic measure. But on the other hand, EE's training process was very slow as it scales quadratically with the training size of the data set [LB15; Bag+17]. This goes back to the leave-one-out-cross-validation (LOOCV) used to optimize the parameters for each used metric [Shi+20].

Proximity Forest wanted to achieve two main goals. The first was to offer an adaptable algorithm that can scale with huge data sets consisting of millions of time series instances. Beating EE, by orders of magnitude, and other state of the art algorithms in terms of training and testing run time complexity. While the other goal was to develop a competitive algorithm on the UCR data sets archive without the need to sacrifice accuracy for scalability as is the case with BOSS-VS [Luc+19].

Capitalizing on the previous research that has been put in developing specialized time series distance measures and inspired by the existing EE [Faw+20; Faw+19b]. Proximity forests combine the eleven elastic distances from EE along with a tree-based algorithms to form an ensemble of decision trees. The reason behind using tree-based algorithms lies in the divide-and-conquer strategy that they adopt, which makes them scalable for large data sets. Also a stochastic process is used for the selection of distance measures and their hyper-parameters, which usually hinders the performance of other algorithms, like KNN, that need to learn the hyper-parameters of the utilized distance measure for each data set before using it [Luc+19]. Proximity forests can scale sublinearly with training data set size, but quadratically with the length of the time series [Shi+20].

Proximity forests are based on a similar concept as Random Forests [Bre01], another tree-based ensemble, which learns only on a subset of the available features for building tree nodes. This process insinuates a factor of variability between the trees that form the ensemble but each with a low bias. The collective classification accuracy of the ensemble then tends to provide better results than any if it's single classifiers [Luc+19].

The building unit of a proximity forest is called the proximity tree. A proximity tree and a decision tree are similar on all aspects, but they differ in the tests they apply in internal nodes. A conventional decision tree builds its nodes using attributes. When an instance is being tested, it is compared to the value of the attribute and then follows the branch to which it conforms.

Unlike conventional decision trees, that use feature values for their nodes, proximity trees build their nodes using randomly selected exemplars. When an instance to be tested, an elastic distance measure is calculated and then it follows the branch of the nearest exemplar.

An internal node in the tree holds two attributes; *measure* and *branches*. As noted in [Luc+19], a measure is function $object \times object \rightarrow \mathbb{R}$. Proximity Forest uses the same 11 distance measures used by EE; Euclidean distance (ED) Dynamic time warping using the full window (DTW); Dynamic time warping with a restricted warping window (DTW-R); Weighted dynamic time warping (WDTW); Derivative dynamic time warping using the full window (DDTW); Derivative dynamic time warping with a restricted warping window (DDTW-R); Weighted derivative dynamic time warping (WDDTW); Longest common subsequence (LCSS); Edit distance with real penalty (ERP); Time warp edit distance (TWE); and, Move-Split-Merge (MSM). Proximity Forest saves a lot of the computational cost by replacing parameter searches with random sampling [Faw+20; Faw+19a]. While branches is a vector of the possible branches to follow, each branch holding two attributes; *exemplar* and *subtree*. *exemplar* is a time series instance to which a query instance is compared, and *subtree* refers to the

tree an instance should follow in case it is closest to a specific exemplar.

If all time series in a specific node share the same class, then a leaf node is created and the value of the class label is assigned to the *class* attribute of this node. During classification, if a query instance is to reach this node, it is directly labeled with the value of its *class* attribute.

When a query time series is to be classified, it starts at the root node of a proximity tree. The distance between the query and each of the randomly selected exemplars is calculated, using the randomly selected distance measure at the node. Then the query travels down the branch of the nearest exemplar. This process is repeated, passing through the internal nodes of the tree till the query reaches a leaf node, where it is assigned the class label of that node. This whole process is then repeated for all the trees constituting the forest. The final classification of the forest is made by majority voting between its trees.

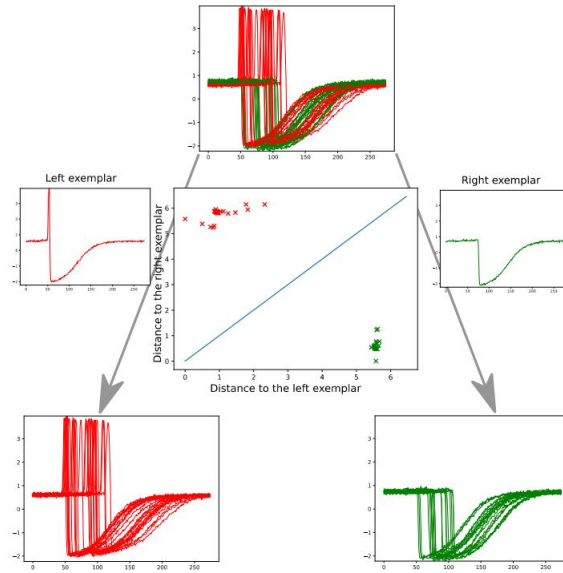


Figure 3.3: Visual depiction of the root node for the ‘Trace’ dataset (simplified to 2 classes). The top chart represents the data at the root node (one colour per class) while the data at the bottom left and right represent the data once split by the tree. The two time series in the middle left and right are the exemplars on which the tree is splitting. The scatter plot at the center represents the distance of each time series at the root node with respect to the left and right exemplars (resp. x- and y-axes) (Color figure online) [Luc+19].

3.3.2 Phase Dependent intervals Algorithms

Phase dependent algorithms is a group of algorithms that extract temporal features from intervals of time series. These temporal features help with the interpretability of the model; as they give insights about the temporal characteristics of the data [BR16],

unlike whole time series algorithms that base their decisions solely on the similarities between instances. Another advantage of phase dependent algorithms is that they can also handle distortions and misalignments of time series data [Den+13].

According to [Bag+17], phase dependent algorithms are best used for problems where discriminatory information from intervals exist, this would be the case with long timer series instances and which might include areas of noise that can easily deceive classifiers. Like the case with the SmallKitchenAppliances dataset, in which the usage of three classes; a kettle, a microwave and toaster is recorded every 2 minutes for one day. Not only the pattern of usage is discriminatory in such case, but also the time of usage.

Typically using interval features requires a two phase process; first by extracting the temporal features and then training a classifier using the extracted features [Den+13]. There are $n(n-1)/2$ possible intervals, for a time series of length n [Bag+17]. There is also a wide variety of features, also called literals, to extract for each interval. These cover simple statistical measures as well as local and global temporal features [SK16; RA04; Den+13]. This introduces one of the main challenges for phase dependent algorithms, that is which intervals to consider for the feature extraction step. Which [RA04] proposed a solution for by only considering intervals with lengths equal to powers of two [Bag+17].

Time Series Forest

Time Series Forest (TSF) is an algorithm that was introduced in 2013 by [Den+13]. They motivated their model with two main criteria; contributing to interpretable time series classification through the use of simple statistical temporal features and reaching this goal by creating an efficient and effective classifier.

TSF considers three types of interval features; mean, standard deviation and slope. If we were to consider an interval with starting point t_1 and with ending point t_2 . Let v_i be the value at a specific point t_i . Then the three features can be denoted as:

$$mean(t_1, t_2) = \frac{\sum_{i=t_1}^{t_2} v_i}{t_2 - t_1 + 1} \quad (3.2)$$

$$std(t_1, t_2) = \begin{cases} \frac{\sum_{i=t_1}^{t_2} (v_i - mean(t_1, t_2))^2}{t_2 - t_1} & \text{if } t_1 < t_2 \\ 0 & \text{if } t_1 = t_2 \end{cases} \quad (3.3)$$

$$slope(t_1, t_2) = \begin{cases} m & \text{if } t_1 < t_2 \\ 0 & \text{if } t_1 = t_2 \end{cases} \quad (3.4)$$

Where m denotes the slope of the least squares regression line for the training dataset.

For building the trees, TSF introduced a new splitting criteria at the tree nodes, which they called the Entrance. A combination of Entropy and distance; to break the ties between features of equal entropy gain by preferring splits that have the furthest distance to the nearest instance. They also use a specific number of evaluation points rather than checking all split points for the highest information gain. In their experiment [Bag+17] found these two criteria to have negative effect on accuracy.

As mentioned earlier the feature space for creating interval features is huge, TSF adopts the same random sampling technique that Random Forests use reducing the feature space from $O(M^2)$ to only $O(M)$, by considering only $O(\sqrt{M})$ random interval sizes and $O(\sqrt{M})$ random starting points at each tree node [Den+13]. The final

classification of a testing instance is done using majority voting of all time series trees created.

3.3.3 Phase Independent intervals Algorithms

Phase independent intervals, or just shapelets as less formally known, are subseries which are ultimately distinctive of classes regardless of their place on the time series [SL17a; Bag+17]. They were first introduced in [YK09] as an alternative for KNN approaches; to overcome their shortcomings.

Shapelets reduce the space and time complexity needed by KNN, because they are formed from subsequences which are shorter than the original time series. Needing only one shapelet at classification time, they form a compressed format of the classification problem [BB17a; YK09; MKY11]. While KNN classify based on comparison to other instances, shapelets provide insight about the unique features of classes and thus more interpretable results of how the classification was carried out. Finally, shapelets are best suited for problems where a certain pattern can differentiate instances which is harder to detect when comparing whole series [Bag+17; BB17b]. Figure 3.4 shows how shapelets can separate instances by calculating distances and provide interpretable explanation by projecting the data in a shapelet transform space.

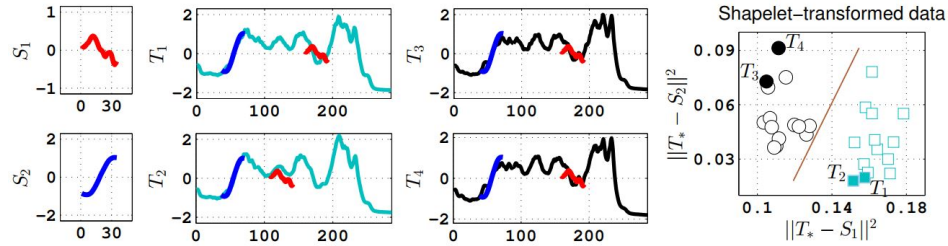


Figure 3.4: An illustration of two shapelets S_1 , S_2 (leftmost plots) learned on the Coffee dataset. Series' distances to shapelets can optimally project the series into a 2-dimensional space, called the shapelet-transformed representation [Lin+12] (rightmost plot). The middle plots show the closest matches of the shapelets on series of two classes having light-blue and black colors [Gra+14].

The original shapelet algorithm enumerated all possible shapelets and embedded the best ones, based on information gain assessment, in a decision tree. Together with a calculated distance threshold, the shapelets and the threshold are used together as splitting criteria [LTB18; Sch15]. There have been many attempts to speed up the process of shapelets discovery, by determining good shapelets in a faster manner. Two of them are; Fast Shapelets (FS) [RK13] and Learned Shapelets (LS) [Gra+14]. FS applied discretization through Symbolic Aggregate Approximation (SAX) to reduce the length of time series, while LS tried to learn the shapelets [Shi+20]. Later on the idea of transforming time series data to an alternative space was adopted in [Hil+14], the transformed data consists of distances to the best k shapelets, then classification is done using an ensemble of eight classifiers.

Learned Shapelets

Learned Shapelets (LS) was proposed in [Gra+14] as a new prospective for approaching time series shapelets. Instead of searching for shapelets through enumeration of all candidates, LS learns K near-to-optimal shapelets that can linearly separate instances through a stochastic gradient objective [LTB18; Bos18]. The found shapelets need not to be a subsequence of one of the training examples [Bag+17; SL17a].

LS follows a two steps technique. In the beginning LS looks for a set of shapelets from the training data set using two parameters; L controls the length of shapelets searched, while R controls the scaling of subsequences. Then these shapelets are clustered using a K-Means algorithm and instances are represented in a new K -dimensional format where the values of the features represent the minimum distance between the instance and one of the shapelets.

For the second step, using the new features representation, LS can start learning class probabilities for instances by considering a logistic regression model for each of the classes and optimizing a regularized logistic loss function. The regularized loss function updates the shapelets and the weights of features. This process keeps iteratively going until either the model converges or the maximum number of iterations is reached. [Bos18]. In summary, the main objective of the algorithm is to learn collectively the optimal shapelets and the weights linear hyper-plane that minimizes the objective function [Bag+17; Gra+14].

Shapelet Transform

The first Shapelet Transform (ST) was introduced in [Hil+14]. While the original algorithm embeded shapelets discovery in decision trees and assessed candidates through enumeration and the use Information Gain (IG) at each node, ST proposed a different way that saved repeating the brute force multiple times [Bos18]. ST segregated the shapelets discovery process from the classifier. This segregation opened the door for choosing classifiers freely and considering more accurate classifiers than decision trees [Bag+17; LB15]. Also [Hil+14] experimented with other shapelet assessment metrics like Kruskal-Wallis, F-stat and Mood's median to find out that F-stat attained higher accuracies than the other three and than IG [Bos18].

ST follows a three step procedure. In the beginning, a data transformation phase is carried out by utilizing a single-scan algorithm and extracting K best shapelets from the training data set where K represents a cutoff threshold for the maximum number of shapelets to extract without affecting the quality of the shapelets extracted. Then a reduction process is done by clustering the shapelets together until they reach a user defined number. Finally, The clustered shapelets are then used to transform the original dataset, by representing instances in terms of their distances to each one of the extracted shapelets. They experimented with different classifiers other than decision trees, these are; C4.5 tree, 1-NN, naive Bayes, Bayesian network, Random Forest, Rotation Forest, and support vector machine, for which decision trees proved to be the worst among all, while support vector machine proved to be the best [Hil+14].

ST was then extended again by [BB17b], the intuition behind it was that the previously used assessment technique couldn't handle multi-class problems [BB17b]. Instead of assessing shapelets that discriminate between all classes, they accommodated a one-vs-all technique so that shapelets are assessed on their ability to separate one class to all other classes. They also introduced a balancing technique to represent each of the classes with the same number of shapelets [Bag+17]. For the classification, a

combination of tree based, kernel based and probabilistic classifiers were used in an ensemble on the transformed data set [Shi+20; LTB18]. Each of the classifiers was given a weight based on its training accuracy and the final classification used weighted voting [BB17b]. Although ST has proved to be a competent accurate classifier, it suffers from high training-time complexity [Shi+20].

3.3.4 Dictionary Based Algorithms

As previously discussed, shapelet algorithms are very good at identifying unique sub-series which can be related to class labels either by their presence or absence. This technique although very helpful for some problems, but it can be deceived if the class labels are defined by the relative frequency of some patterns and not just the a single match [Lar+18; Bag+17]. Dictionary based algorithms is a family of algorithms that uses the frequency of words as the foundation of finding discriminatory features of classes [MVB19]. They can address this type of problems by tracking the frequency counts of repeating patterns, which makes same resilient to noisy data [Shi+20]. To achieve this, dictionary based algorithms transform time series data into words; to reduce their dimensionality and approximate their values. Then based on the distribution of the words they measure similarity between instances [LKL12; Sch15].

In general, the process of dictionary based classifiers involves using a sliding window, of some length w , over the data. Then for each window, an approximation process is carried out to reduce its length to from w to a shorter length l [Bag+20]. A quantization process then follows; to discretize the real values into words by assigning letters to each value. When all the windows are finished, the number of occurrences of words is calculated and each time series is represented by a histogram [Shi+20]. In the end, similarity between instances is calculated based on similarity between histograms.

Different dictionary based algorithms apply different approximation and discretization techniques. For example two of the older algorithms; Bag of Patterns (BoP) [LKL12] and Symbolic Aggregate Approximation-Vector Space Model (SAX-VSM) [SM13] use an approximation method called Symbolic Aggregate Approximation (SAX) [Lin+07]. SAX extends Piecewise Aggregate Approximation (PAA), a method that divides a time series into equal consecutive segments and concatenates their means [Shi+20]. Then to transform the series to words, it creates break points using quantiles from the normal distribution. In addition to using SAX, SAX-VSM merges it with the vector space model from the Information Retrieval domain. The major difference between BOP and SAX-VSM, is that SAX-VSM calculates its frequencies on the class level rather than on the series level. Then applies weighting using inverse document frequency [Bag+17].

On the other hand, the newer algorithms; BOSS [Sch15], BOSS-VS [Sch16] and WEASEL [SL17a] use another method which is called Symbolic Fourier Approximation (SFA) which approximates data using coefficients of Discrete Fourier Transformation (DFT). Then apply an adaptive technique called Multiple Coefficient Binning (MCB) to quantize the data. These two techniques; SFA and MCB, will be discussed in more details in the next parts.

BOSS

Bag of SFA Symbols (BOSS) is a dictionary based algorithm that was introduced by [Sch15] in 2015. Like the previous dictionary based classifiers, BOP and SAX-VSM, BOSS also used a windowing technique to extract patterns from the data and

transform them into words, but it had other significant differences to them [Bag+17]. BOSS was concerned with the issue of dealing with noisy data which, at that time, received little attention; due to the common practice of either using raw data directly or handling noise in a preprocessing stage. The goal was to introduce an algorithm faster than it's rivals of the same group, robust for existence of noise in the data and competitive with existing TSCA.

The BOSS model is divided into several steps. In the beginning, it passes a window of size w over the time series to extract substructures. Each of the extracted windows is then normalized to obtain amplitude invariance, while obtaining offset invariance by subtracting the mean value for each window is data set specific and can be decided upon based on a parameter. Then the substructures are quantized using SFA transformations which transforms the data into unordered words; this helps further reduce the noise in the data by making it phase shift invariant and makes it possible to apply string matching algorithms on the data. Since some data sets might happen to have long constant signals, this will lead to SFA transformations of their windows to produce the same words multiple times and cause higher weights to be assigned for them. BOSS applies a numerosity reduction technique adopted from [Lin+07; LKL12] that ignores multiple sequential occurrences of the same word. Finally time series instances can be compared for differences, using the noise reduced substructures using a customized distance measure inspired by Euclidean distance. We will, briefly, discuss the main components that are used for each of the previously mentioned steps.

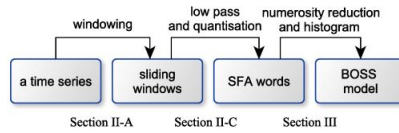


Figure 3.5: The BOSS workflow [Sch15]

To extract the substructures from a given time series instance $T = [t_1, t_2, \dots, t_n]$, a windowing function is used to split it into fixed sized windows $S_{i:w} = (t_i, \dots, t_{i+w-1})$, each of them is of a size w . The total number of windows that can be created for a time series of length n is $n - w + 1$, and each consecutive windows overlap on $w - 1$ points. In order to achieve offset and amplitude invariance, each window is z-normalized by subtracting the mean from it's original values and then dividing the difference by the standard deviation.

$$windows(T, w) = \{S_{1:w}, S_{2:w}, \dots, S_{n-w+1:w}\} \quad (3.5)$$

After running the windowing function, the real values of the time points inside the windows are transformed into words using SFA [SH12]. SFA is an alternative way of representing time series data, that instead of using real values, uses a sequence of symbols which are referred to as SFA words based on a predefined alphabet of specific length. SFA accomplishes two things; low pass filtering through removal of noisy components from the data and string representation which allows for the use of string matching algorithms.

For SFA to achieve it's goals, two main operations have to be carried out; approximation and quantization. Approximation is the process of representing a specific

signal of length n using another signal of lower length l . This is achieved using Discrete Fourier Transformation (DFT); a transformation technique which is applied to a signal represented by a sequence of equally spaced values into a sequence of coefficients of complex sinusoid waves ordered by their frequencies [LLY17]. The higher coefficients in a DFT refer to data with rapid changes, which can be considered as noise in the signal and thus ignored. So considering only the first $l \ll n$ coefficients acts as a low pass filter for noise producing a smoother signal.

Quantization also helps reduce noise by splitting the frequency domain into equi-depth bins, then maps each of the Fourier real and imaginary coefficients into a bin. BOSS utilizes Multiple Coefficient Binning (MCB), an adaptive technique that minimizes the loss of information which is a side effect of quantization. After the approximation step, a matrix is built from the Fourier transformations of the training data set using only the first $\frac{l}{2}$ coefficients, including both the real and imaginary values for each coefficient. Then using an alphabet Σ of size c , MCB creates for each column of the matrix $c+1$ break points using equi-depth binning. During classification, to acquire the SFA word for a Fourier transformed time series, a lookup is done on the precomputed MCB bins and a word is assigned if the value falls within its bin's boundaries.

After the transformation of instances to SFA words, BOSS uses a customized distance measure referred to as BOSS distance; to measure the similarity between the transformed instances. BOSS distance is a variation of Euclidean distance, which compares instances based on the histograms formed from their transformed forms. The appearance of the same SFA words in both instances, is considered to be a notion of similarity. While the absence of SFA words might be caused by one of two reasons; the absence of some substructures from either instances, or due to the presence of noise which disfigures the time series. Instances under BOSS distance are compared based on shared SFA words only, thus excluding words of frequencies equal to 0. BOSS distance as noted by [Sch15] is:

Given two time series instances T_1 and T_2 and their corresponding BOSS histograms $B_1 : \Sigma^l \rightarrow \mathbb{N}$ and $B_2 : \Sigma^l \rightarrow \mathbb{N}$, where l is the word length and Σ is an alphabet of size c . Their BOSS distance is:

$$D(T_1, T_2) = dist(B_1, B_2) \quad (3.6)$$

where

$$dist(B_1, B_2) = \sum_{a \in B_1; B_1(a) > 0} [B_1(a) - B_2(a)]^2 \quad (3.7)$$

Each single BOSS classifier utilizes 1-NN approach along with its BOSS model. The reason behind using 1-NN is that it is a simple technique that doesn't add to the parameters of the model, but rather proved to be a robust one. When classifying a query instance, BOSS searches for the nearest neighbor in a sample of candidates by choosing the closest instance based on the BOSS distance.

Finally, BOSS Ensemble is introduced as an ensemble of multiple BOSS classifiers. While a fixed window length is used over time series instances for BOSS classifier, BOSS Ensemble considers representing each time series instance by ensembling multiple BOSS classifiers, each of a different window length. When the training data is fitted, BOSS Ensemble acquires a group of scores for each of the different window lengths. To classify a query instance, BOSS Ensemble acquires the best accuracy score from the score sets returned during training. Then considers all window lengths that obtained accuracies within a factor of the best accuracy score. Each of the considered

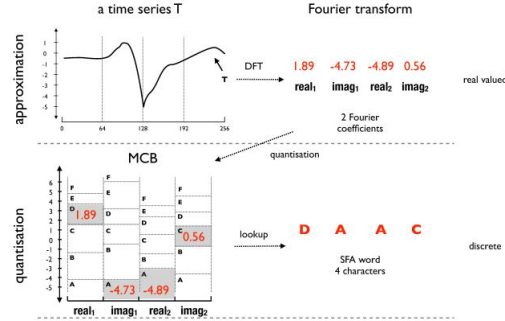


Figure 3.6: A time series is (a) approximated (low pass filtered) using DFT and (b) quantised using MCB resulting in the SFA word DAAC [Sch15]

windows predicts a class label for the query instance using 1-NN. Majority voting is applied and the most dominant class label is assigned.

In [MVB19], some enhancements were introduced to BOSS. These included randomising the parameter settings, subsampling training data for each classifier, adding a voting scheme to assign weights to classifiers based on the probability distribution from cross validation on training dataset, fixing the number of classifiers and adding a time contract.

The new enhanced version of BOSS is named the Contractable BOSS or CBoss. In the contractable BOSS, the ensemble is assigned to build as many individual BOSS classifiers as possible during a specified duration of time (the contracting time) despite the properties of the data set. In cases of small data sets, this might require huge memory resources; as the number of possible classifiers to create would increase. This was overcome by introducing a parameter to limit the number of maximum classifiers to create. On the other side, for huge data sets there is a chance of building problems to happen and the loss of progress. Which was overcome by adding checkpoints to periodically save the current state of the classifier.

Multiple experiments were carried out using different variations of the enhancements. None of the enhancements alone proved competence to the original BOSS performance, but a combination of them together made it possible to achieve significant results. More details about the experiments can be found in [MVB19]. In our framework, we use the implemented contractable version of BOSS as a replacement for the original BOSS, as suggested by [MVB19].

WEASEL

After the success that BOSS achieved, it became one of the baseline classifiers of TSC. Other research has either included it as a building component of their ensembles [LTB18; Bag+15], or have used it, as a baseline for comparison of their algorithm's performance [Faw+20; Shi+20; Luc+19]. Later on, the same team that developed BOSS introduced a newer algorithm, Word ExtrAction for time SERIES cLassification (WEASEL) [SL17a]. A dictionary based classifier which is very similar to BOSS and can be thought of as an extension to it, with more focus on scalability [MVB19]. WEASEL motivated their work with the absence of scalable classifiers, at the time,

that can deal with big data sets; as the existing were either not scalable enough or not accurate enough. Despite being a non-ensemble classifier, WEASEL can compete with powerful ensembles in terms of accuracy without the need for long prediction times, but this comes with the cost of resource expensive training [MVB19].

Like the other dictionary based approaches, WEASEL uses a sliding window over the data instances to extract substructures. These substructures are then discretized per window to extract features and finally a machine learning classifier is learned over the transformed features. But WEASEL differentiates itself from the other algorithms in the way it constructs and filters the features. We will discuss the new approaches that WEASEL uses for extracting discriminative features, building a model to deal with multiple occurrences of words and variable windows' lengths and selecting features to reduce runtime and exclude irrelevant features.

In the beginning, WEASEL uses multiple sliding windows of different lengths; to extract substructures. While keeping track of their order; in order to use them later on as bi-gram features. This replaces the process of building multiple models then choosing the best one, with building only one model which can learn from the concatenated high-dimensional feature vector. Then each of the substructures is normalized and a DFT is applied. Instead of filtering out the higher Fourier coefficients, WEASEL applies an ANalysis Of VAriance (ANOVA) F-test to keep real and imaginary Fourier values that best separate instances from different classes. Then the kept coefficients are discretized into words, based on alphabet of size c , using binning boundaries. Instead of using equi-depth binning, WEASEL applies an information gain based binning technique; which further more helps separating instances of classes. In the end, a histogram is built using all the windows lengths and the extracted features, uni-grams and bi-grams. Irrelevant features are then filtered out using a Chi-squared test. The final highly discriminative feature vector produced is then used to learn a logistic regression classifier.

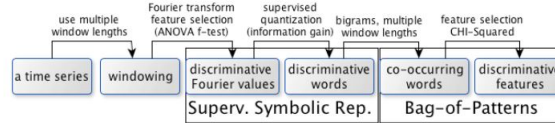


Figure 3.7: WEASEL Pipeline: Feature extraction using a novel supervised symbolic representation, and a novel bag-of-patterns model [SL17a]

WEASEL, like BOSS, transforms extracted windows from a time series into words using an alphabet of size c . They identify two main drawbacks for SFA, and introduce a new supervised symbolic representation technique which is based on SFA but overcomes the drawbacks. The first drawback of SFA is that it acts like a low pass filter and excludes the high frequency components from the Fourier transformation, which might discard important features in some scenarios. The other drawback, is that SFA defines the boundaries of bins during quantization independent of the class labels; which might cause SFA words of equal frequencies to appear unnecessarily in multiple classes. In order to overcome the drawbacks of SFA, WEASEL follows two steps; discriminative approximation using ANOVA F-test and discriminative quantization using information gain.

As mentioned earlier, approximation involves representing a time series of length

n with a shorter, yet informative, representation of length l by applying a Fourier transformation. WEASEL aims at keeping the best class separating l , real and imaginary, Fourier coefficients by applying a one-way ANOVA F-test. The test verifies the hypothesis that the means of two or more distributions/groups differ from each other [Low14]. This can be tested by comparing two variances; the variance between groups and the variance within the groups. Using the notations in [SL17a], *mean square between* (MS_B) and *mean square within* (MS_W) respectively. The F-values is then calculated as :

$$F = \frac{MS_B}{MS_W} \quad (3.8)$$

Then l coefficients with the highest F-values are kept; as these represent features which have big differences between the classes (high MS_B) and small differences within the same class (low MS_W).

After that, discretization is carried out to set the split thresholds for each of the extracted Fourier value. Discretization involves a binning process, where each Fourier value is divided into a number of bins. Each bin is represented by an upper and a lower boundary, and assigned a letter from an alphabet c . Previous quantization techniques used equi-depth or equi-width binning, but these techniques are solely based on values and ignore the distribution of classes. WEASEL applies a binning technique based on IG; assuring that for each partition the majority of the values would belong to the same class.

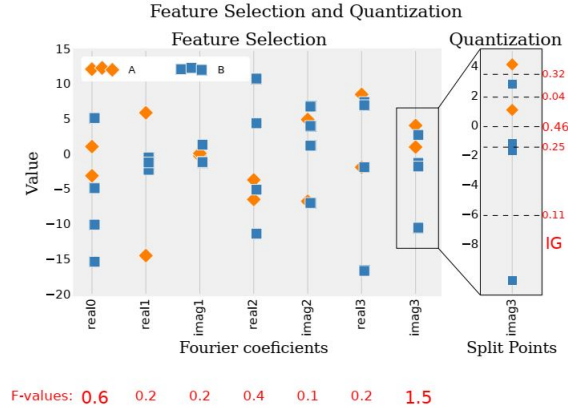


Figure 3.8: On the left: Distribution of Fourier coefficients for a sample dataset. The high F-values on imag3 and real0 (red text at bottom) should be selected to best separate the samples from class labels 'A' and 'B'. On the right: Zoom in on imag3. Information gain splitting is applied to find the best bins for the subsequent quantization. High information gain (IG) indicates pure (good) split points [SL17a]

The result of the feature extraction phase is a high dimensional feature vector with a dimensionality of $\mathcal{O}(\min(Nn^2, c^l))$, where c is an alphabet, l is the length of a word, n is the total number of instances and n is the length of the time series. Since WEASEL uses bi-grams and $\mathcal{O}(n)$ window lengths, the dimensionality of the feature space increases to $\mathcal{O}(\min(Nn^2, c^{2l} \cdot n))$. This enormous feature space is then reduced

using a Chi-squared (χ^2) test. Chi-squared tests is a statistical test used to determine if there is a significant difference between the recognised frequencies and the expected frequencies of a features within a group. This implies that features which have high (χ^2) values are statistically frequent in certain classes. By comparing the (χ^2) values of features to a threshold, all features with scores lower than the threshold can be excluded from the feature space. This usually reduced the feature space by 30% - 70% and helped train the logistic regression classifier in a timely manner.

3.3.5 Deep Learning Algorithms

Deep learning is a long established group of machine learning algorithms which has proved competence in many areas [LBH15] and which have encouraged the introduction of deep learning algorithms for time series classification [WYO17]. Deep learning is appealing for investigating time series data; due to the role that the time dimension play as a structure for the data and because deep learning algorithms can scale linearly with training size [Shi+20].

The general framework of deep learning neural networks is described by [Faw+19a] as an architecture of L layers, each one of them is considered an input domain representation. Each of the layers consists of small computing units referred to as neurons. Neurons compute elements for the output of each layer. A layer l_i applies an activation function to it's input then passes the output to the next layer l_{i+1} . The behavior of activation functions in each layer is controlled by a set of parameters θ_i and are referred to by the term weights. The weights are assigned to the links between the inputs and outputs of the network's layers. A neural network carries out a series of computations to predict the class label for a given input x , these calculation are noted as:

$$f_L(\theta_L, x) = f_{L-1}(\theta_{L-1}, f_{L-2}(\theta_{L-2}, \dots, f_1(\theta_1, x))) \quad (3.9)$$

Where f_i represents the activation function applied at layer l_i .

In the training phase of a neural network, the weights are randomly initialized. Then a forward pass of the input x is done through the model and an output vector is computed, in which every component of the vector represent a class probability. Using a cost function, the model's prediction loss is calculated and weights of the network are updated, in a backward pass process, using a gradient descent. By continuous iteration of forward passes and weight updates by backward passes, the neural network learns the best weights to minimize the cost function. For the prediction of unseen instances, or inference phase, the input data is passed through the network in a forward pass. Then using the class probabilities of the output vector, the class with the highest probability is assigned.

Many of the deep learning research on time series focused on the use of variants of Convolutional Neural Networks (CNN). [Faw+19b] CNN model is based on the idea of learning convolutional filters that can accurately represent the data. Using these filters, CNNs can learn the hierarchical structure of the data while incorporating translation invariance[LMT16]. Multi-Scale Neural Networks (MCNN) [CCC16] and LeNet [LMT16] were among the first experiments of using CNN in time series classification. MCNN was composed of three stages; transformation stage, local convolution stage and full convolution stage. The transformation stage aimed at applying a low pass filter to exclude noise and to capture different temporal patterns. Local convolution is a stage where different scale features are extracted and local features of the time series are learned. In the final stage, the full convolution stage, all the features are

concatenated and the final prediction is made. On the other hand, LeNet consists of 2 convolutional layers, after each one a max pooling is carried out to do sub-sampling. For the first layer, 5 filters are applied and the max pooling size is 2. While for the second there are 20 filters and max pooling size is 4. In [WYO17] MultiLayer Perceptrons (MLP), Fully Convolutional Networks (FCN) and the Residual Networks (ResNet) were tested on 44 datasets from the UCR time series archive, where FCN was not significantly different from state of the art and MCNN was not significantly different from COTE and BOSS [SL17a].

A recent more comprehensive review of deep learning algorithms was done by [Faw+19a]. In which an experiment between nine deep learning algorithms was done. These included the classical MCNN and LeNet, in addition to the three algorithms from the previously mentioned experiment; MLP, FCV and ResNet. The other algorithms included were Encoder [SPK18], Multi Channel Deep Convolutional Neural Network (MDCNN) [Zhe+14; Zhe+16], Time Convolutional Neural Network [Zha+17] and Time Warping Invariant Echo State Network (TWIESN) [TH16]. For more details about the algorithms and their structures, please refer to [Faw+19a].

Finally, the newest deep learning algorithm introduced for time series classification is InceptionTime [Faw+20], which is the newest state of the art and which is an ensemble of deep convolutional neural networks. It can attain high accuracy scores while maintaining scalability. We will discuss in more details the InceptionTime algorithm in the next section.

InceptionTime

InceptionTime [Faw+20] is a recent deep learning algorithm for TSC problems, which is able to achieve high accuracy score [Rui+20]. Motivated by the increasing interest in deep learning algorithms in the TSC domain along with the need for scalable algorithms that can deal with large data sets, either in number of instances or in length of the time series, and scale to them. InceptionTime is inspired by AlexNet [KSH12], an algorithm that was considered a breakthrough for deep learning algorithms [Alo+18], and wanted to achieve the same but for the domain of TSC.

InceptionTime is based on the idea that the combination of deep CNN with residual connections, like in ResNet, can attain higher classification performance [Faw+19a]. Since CNN has proved competency with image classification, there seemed a potential opportunity to be able to use deeper CNN for time series; since time series data is mainly structured on only one dimension which is time, while images have two spacial dimensions. This opened the door for using more complex models for TSC problems which would be computationally challenging to use for images. The building blocks of an InceptionTime network are called Inception modules, these are were introduced by [Sze+15] and evolved later on to Inceptionv4 [Sze+17]. The InceptionTime classifier is an ensemble of 5 InceptionTime networks each initialized with random weights and are assigned equal weights for the final prediction. We will discuss in more details the structure of an InceptionTime network using the notation mentioned in [Faw+20].

InceptionTime's structure is similar to that of a ResNet, but instead of using three residual blocks, InceptionTime uses only two. Each of the residual blocks is composed of three Inception modules instead of the traditional fully convolutional network. Like residual networks, a linear skip connection exists between the two residual blocks of the network, passing the input from one block to be concatenated with the input of the other; this helps passing information from earlier layers of the network to deeper layers and thus mitigating the vanishing gradient issue [He+16]. After the residual

blocks, a Global Average Pooling (GAP) layer exists where the multivariate time series output is averaged over the time dimension. The final component of the network is a conventional fully-connected softmax layer with a count of neurons similar to the count of output classes.

Inside the inception module, there are two main components; the bottleneck layer and the variable length filters. Assuming that the input in a multivariate time series data of M dimensions. The job of the bottleneck layer is to transform the data from having M dimensions, into a multivariate data set having m dimensions, where $m \ll M$. This is done by passing a group of sliding filters m with length 1 and a stride of size 1. Which will substantially reduce the dimensionality of the time series data and consequently will also decrease the model's complexity making it more robust for overfitting problems on data sets of small sizes. The other benefit of including the bottleneck layer, is that it allows utilizing longer filters on the data than the original ResNet using approximately the same number of parameters; due to the lower number of dimensions that the filters will have to deal with.

The output of the bottleneck layer is then passed for a set of variable length filters, the second component of the network, of length l where $l \in 10, 20, 40$. In addition to the filters, a parallel MaxPooling operation is carried out followed by a bottleneck layer; to make the model robust to small data noises. The final multivariate output is then formed by concatenating the output from each filter based convolution along with the output of the MaxPooling operation. This whole process is executed for each Inception module in the network.

In the end, an InceptionTime network is able to learn the underlying hierarchical structures of a time series by stacking multiple Inception modules and learning from the different filter sizes, which had been learned during training, inside them.

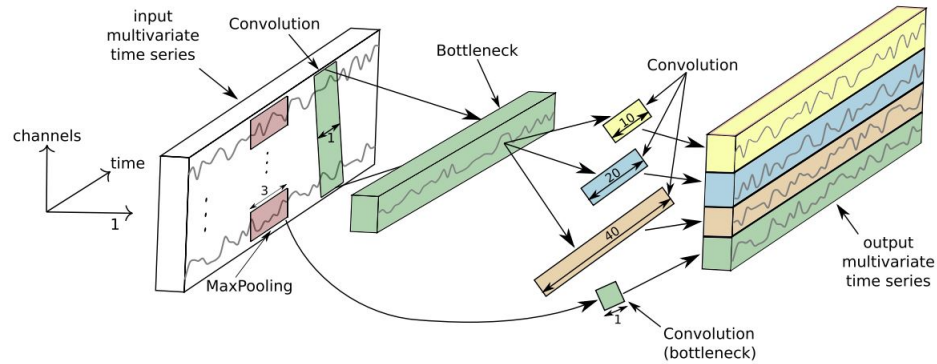


Figure 3.9: Inside Inception module for time series classification. For simplicity a bottleneck layer of size $m = 1$ is used [Faw+20]

The final InceptionTime classifier is an ensemble of 5 InceptionTime networks. InceptionTime networks accuracy scores showed high variances, a problem which have been discussed by [SW17] and was found in ResNet networks as well [Faw+19b]. This happens due to the random initialization of the networks and due to the stochastic approach used for optimization. InceptionTime classifier follows an ensembling technique for neural networks to handle TSC [Faw+19b] in order to leverage the high variability; adhering to the idea that combining multiple networks would yield better results than

one classifier. During the classification of an instance, InceptionTime combines the logistic output of the 5 networks and assigns an equal weight to each of them. This can be denoted by the equation

$$\hat{y}_{i,c} = \frac{1}{n} \sum_{j=1}^n \sigma_c(x_i, \theta_i) | \forall c \in [1, C] \quad (3.10)$$

Where $\hat{y}_{i,c}$ is the class probability for instance x_i as belonging to the class c , which is the averaged logistic output σ_c over the randomly initialized models n .

3.4 Early Time Series Classification

On another side, Early Time Series Classification (eTSC) is also a classification problem which considers the temporal nature of data, but with a focus on slightly different objectives. and had been used in multiple domains.

eTSC's main objective is to learn a model which can classify unseen instances as early as possible, while maintaining a competitive accuracy compared to a model that uses full length data or to a user defined threshold [XPP09]. Which is a very challenging objective; due to the, naturally, contradicting nature of earliness and accuracy. In general, the more data is made available for the model to learn the better accuracy it can attain [Mor+19; TM16; XPP12; Mor+17b]. This is why eTSC is considered as a problem of optimizing multiple objectives.

We have already defined accuracy for TSC in section 3.2. For earliness, we follow the definition mentioned by [SL20]; as the mean number of data points s after which a label is assigned.

Definition 6

$$Earliness = \frac{\sum_{T_i \in D} \frac{s}{len(T_i)}}{|D|}$$

As well as the objective measure, Harmonic mean (HM), mentioned by [GO12; SL20], which combines both accuracy and earliness in one equation. For the scope of our framework, we consider HM as a weighted average between accuracy and earliness.

Definition 7 $F_\beta = (1 + \beta^2) \frac{accuracy(1 - earliness)}{\beta^2(1 - accuracy) + earliness}$

The value of β can be used to give higher importance to one of the aspects over the other, but we use equal weights for both.

eTSC is needed in situations in which waiting for more data to arrive can be costly or when making late decisions can cause unfavorable results [Mor+17a; Par+13; Lin+15]. This is why eTSC has been applied in various domains like early medical diagnosis [GM01; GO12], avoiding issues in network traffic flow [Ber+06], human activity recognition [YD19; Gup+20] and early prediction of stock crisis [GRO14].

3.5 Early Time Series Classification Algorithms

3.5.1 ECTS

The idea of eTSC evolved over time. In the beginning, similar concepts to eTSC existed where the main goal was to classify prefixes of data, but these techniques lacked the reasoning of how to provide reliable classification results [Lv+19; SK16]. One of the

earliest papers that formally defined eTSC as a tradeoff between earliness and accuracy was the paper by [XPP09], proposing a technique which is named after the problem itself; Early Time Series Classification (ECTS). The main idea was to examine the stability of of Nearest Neighbor approaches, to keep correctly classifying data instances, in the space of full length series and the subspaces created from subsequences of the training data set [Lv+19; Mor+17a]. ECTS utilizes the concept of reverse nearest neighbors (RNN) to learn for each training instance the Minimum Prediction Length (MPL). MPL represents the earliest point in time for which a training instance can be used to classify a query instance without loss of accuracy. The definition mentioned by [XPP09] defines MPL as:

Definition 8 *In a training data set T with full-length L , for a time series teT , the minimum prediction length (MPL for short) of t , $MPL(t) = k$ if for any $l(k \leq l \leq L)$, (1) $RNN^l(t) = RNN^L(t) \neq \emptyset$ and (2) $RNN^{k-1}(t) \neq RNN^L(t)$. Specifically, if $RNN^L(t) = \emptyset$, then $MPL(t) = L$.*

During classification MPL is calculated for each instance (in the same study a relaxed variant is introduced using clusters of instances). As for the classification, as the data points of the testing instance start to arrive, the classification method tries to find the nearest instance which has reached it's MPL. If there was no such instance found, then more data points are needed till a trusted classification can be done. ECTS was criticized [He+15; GRO14; Xin+11] because it is based on nearest neighbor approaches; which only provide classification results based on distance without deriving patterns from the data. This means that data users cannot attain insights about the problem using the results.

3.5.2 Shapelet Based Algorithms

A group of methods [Xin+11; GO12; GRO14; He+15] focused on the interpretability of the results obtained by eTSCAs, motivated by domains where not only correct classification is important, but also getting insights and understanding what contributed to the final result. One of these domains would be the medical field, where understanding the reason for a patient's disease is very important than just linking the patient to previously seen patients [Xin+11]. Shapelets provided a good solution for these methods. Since shapelets, as previously mentioned, are defined as unique subsequences which can uniquely identify classes. Early Distinctive Shapelet Classification (EDSC) was a framework consisting of 2 steps; feature extraction and feature selection. In the feature extraction step, all local shapelets are extracted and a distance threshold is learned. EDSC developed a technique which they called the best-matching-distance (BMD), which considers a shapelet to be distinctive, by comparing it to all instances in the training data set and considering target classes distribution. If the majority of instances closer to the shapelet belong to a target class, it is considered to be distinctive of this class. In the feature selection step, EDSC selects a smaller set of features based on three main criteria; earliness, frequency and distinctiveness. EDSC was later extended in two different directions.

In [GO12] EDSC was extended in the aspect of data dimensionality; as the original EDSC only covered univariate data sets. The extension covered; adapting the feature extraction to discover multivariate shapelets along with calculating distance thresholds for each dimension and using a distance threshold based on information gain, adapting the feature selection step, by modifying the equations calculating the notions of earliness, frequency and distinctiveness to multiple dimensions. In [GRO14] EDSC was

extended in the aspect of data uncertainty. The main idea was that shapelets use a distance threshold when being compared to instances, if the distance between the shapelet and the instance is less than the threshold, the instance is given the label of the shapelet. On the other hand, since the threshold represents a range, a shapelet is more certain of the class label given to an instance which has a distance closer to 0 than the class label given to an instance falling on the limits of the threshold. Motivated by the interest of uncertainty in the medical field, the goal was to provide a notion of uncertainty along with the interpretability of shapelets. The proposed Modified EDSC with Uncertainty estimates (MEDSC-U) calculated uncertainty as $(1 - \text{confidence})$, where confidence is a numeric value representing two aspects; confidence that the distance between the shapelet and the instance is less than the threshold and confidence that the shapelet can correctly classify the instance. This confidence notion could then be generalized for classes represented by multiple shapelets. The issue of confidence for eTSC had been discussed in other papers like [Par+13] and [Lv+19].

3.5.3 ECDIRE

The paper by [Mor+17b] proposed a new framework; Early Classification of Time Series based on Discriminating Classes Over Time (ECDIRE). There were two goals that ECDIRE wanted to achieve [Mor+17a]. The main goal of the framework was to track the accuracy evolution for a group of probabilistic classifiers across time; to identify safe time points after which predictions can be made. This would allow for avoiding unnecessary checking at each time point. The other goal was to offer an outliers discarding mechanism using a reliability condition. ECDIRE consisted of three training steps. In the first step the framework analyzes the data set and tries to identify the safe points at which classes can be differentiated from each other. This is done by defining time points as percentages of the full length and training, at each time point, a probabilistic classifier. The early points are then defined as those which maintain an accuracy of a desired percentage from that obtained on the complete data [SK16]. In the second step, a reliability threshold is determined using the class probabilities of the assigned class and the second highest class; this helps avoid uncertain classifications or very soon time points. In the last step, the final probabilistic classifiers are trained using the early time points identified in the first step. As a backup plan, if the last early time point did not coincide with the full length of the time series, ECDIRE acts like other methods and builds classifiers on each time point.

3.5.4 TEASER

One of the latest classifiers introduced is the Two-tier Early and Accurate Series classifier (TEASER) [SL20]. TEASER's main focus was challenging the logic of previous classifiers on determining the point of time at which a classification can be assigned to a time series instance. Previous methods would try to learn from the training data set a fixed prefix needed to be seen before it can start classification; like MPL in ECTS and EDSC, or to learn a safe point like in ECDIRE. These techniques would require the classifiers to always wait till they reach these fixed points of time before they can assign a label. Whereas TEASER was designed to give a classification once confident of its decision, and not to depend on any fixed time points nor assume that all data instances were recorded at the same starting point. TEASER models the problem of eTSC with a two tier solution. In the first tier, a slave classifier is regularly checking

the incoming data and assigning class probabilities. These probabilities are then assessed in the second tier by a master classifier, which either accepts the classification from the slave, or rejects it and decides to wait for more data to arrive. The classifiers are paired through out the whole process, such that for each slave classifier there is a master classifier that reads its output and assesses its results. TEASER starts off by defining the number of classifier pairs that will be needed, this is calculated by dividing the length of the longest data instance in the training data set by the window length which is a user defined parameter. Which is noted as $S = \lceil n_{max}/w \rceil$, where n_{max} is the length of the longest instance and w is the window length. During training, slave classifiers are trained on subsequences s_i , or snapshots, of the time series where the length of $s_i = i.w$. These snapshots are z-normalized based on the values existing only in them and not on values from the full length data as recommended by [Mor+17a].

The slave classifiers would then calculate three features; the class label with the highest probability $c(s_i)$, the vector of class probabilities for all available classes $P(s_i)$, and the difference between the class probability of the highest two classes denoted as Δd . These three features are then passed to the paired master classifier, which trains a model over them using a one-class Support Vector Machine (oc-SVM). A parameter that plays an important role in this process is v , it is the number of consecutive times the same class label is given to an instance. Once the same label is given v times for consecutive snapshots, the master classifier is confident of this class label. Figure 3.10 shows an example of how TEASER components handle a misleading data instance from the traces of microwave and a digital receiver.

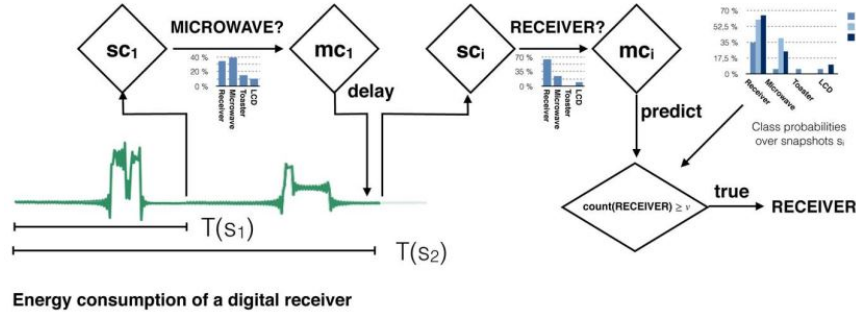


Figure 3.10: TEASER is given a snapshot of an energy consumption time series. After seeing the first s measurements, the first slave classifier sc_1 performs a prediction which the master classifier mc_1 rejects due to low class probabilities. After observing the i -th interval which includes a characteristic energy burst, the slave classifier sc_i (correctly) predicts RECEIVER, and the master classifier mc_i eventually accepts this prediction. When the prediction of RECEIVER has been consistently derived v times, it is output as final prediction [SL20]

Chapter 4

Data Sets

In order to address our research questions, this thesis considered a group of data sets from the UCR data archive [Dau+18] and the UEA data archive [Bag+18]. This Chapter will describe the data sets that were used to conduct this empirical experiment and their general properties.

4.1 UCR/UEA Archives

The University of California Riverside (UCR) and the University of East Anglia (UEA) data archives have been used as references in literature for many experiments on time series problems [AML19; Faw+20; Bag+17; YD19; Rui+20; Faw+19a] and specially for benchmarking the performance of algorithms.

4.1.1 UCR Archive

The UCR archive previously had 85 univariate data sets but in 2018 was then expanded to reach 128 data sets. Despite sharing the property of being univariate, the data sets cover a variety of values for other aspects like; training data size [16, 8926]; testing data size [20, 16800]; number of classes [2, 60]; length of time series [15, 2844] and for some data sets length of instances vary within the data set; and the nature of the data being collected. The archive offers a single predefined train/test split for each of the data sets to facilitate reproducibility of results. Finally, the data is z-normalized; to remove offset and scaling. Z-normalization is the transformation of the data to have a mean of zero and one unit of standard deviation.

4.1.2 UEA Archive

The other archive we used is the UEA archive for multivariate data sets. Like the UCR archive, was developed and expanded over time. It started as a small archive of 12 data sets collected by Mustafa Baydogan, then later on, in 2018, was expanded to reach 30 data sets as a collaboration between researchers of UEA and UCR. The data sets vary in their characteristics; training data size [12, 30000]; testing data size [15, 20000]; number of classes [2, 39]; length of time series [8, 17984]; number of dimensions [2, 1345] and six different different groups based on their application area. The archive

was processed so that the data instances have equal lengths, instances of missing data points were excluded and a single predefined train/test split is provided.

4.1.3 Used Data Sets

For the scope of our experiment, a total of 77 data sets were selected from both archives to be used within the framework. We have based our selection criteria for data sets on:

- The data belongs to a real time series structure behind it and not converted from another data format.
- All instances in the data sets has the same length of time series.
- The data data sets should not contain any missing values.

Based on the previously mentioned criteria, we excluded data sets that belonged to either of the types; Simulated, Image or Motion type; to comply with our first selection criteria. For the other two criteria; we avoided any further preprocessing than what is offered by the data archives. As mentioned before, the univariate data sets from the UCR are already Z-normalized. Z-normalization is a common and recommended practice for time series data as it has shown to have a positive impact on the performance of classifiers [Bag+17; Faw+19a] This form of preprocessing is not true for the multivariate data sets from the UEA archive.

We excluded data sets with missing data and avoided filling these missing values; as we believe that data imputation is a whole another topic. We also avoided any interpolation of data. A technique which has been previously used in [RK05; Faw+19a] to make all instances of the data set of the same length as the longest instance. Our goal in this experiment is to focus on comparing the performance of the classifiers in our proposed context based while ensuring, as much as possible, that the main contributing factor is the technique behind the classifiers. This doesn't mean that preprocessing of time series data is of a less importance, but we simply excluded it because we believe that it doesn't fit into our current scope.

The remaining collection of data sets still represented a wide variety in terms of data criteria. We present the statistics about the remaining data sets all together in figure 4.1. Since the smallest data set (StandWalkJump) and the largest data set (InsectWingbeat) are the same as the UEA archive; training and testing data sizes are the same as previously mentioned ranges in the original archive, [12, 30000] and [15, 20000] respectively. The data sets covers covers from binary classification problems up to problems with 52 classes, the range [2, 52]. The range of the time series length is [24, 3000], with Chinatown being the shortest and MotorImagery the longest data set. For the 22 multivariate data sets, the number of dimensions still covered the whole range [2, 1345] like the UEA archive. Finally after our exclusion criteria, the remaining 14 problem types are AUDIO, DEVICE, ECG, EEG, EOG, EPG, HAR, HEMODYNAMICS, MISC, OTHER, SENSOR, SOUND, SPECTRO and Traffic.

Although the data sets that we used covered a variety of properties, but within some properties there were huge imbalances in the distribution of values. Within grouping by training data set size, the group of data sets with sizes [501, 1000] was the smallest, with only 4 data sets which is less than half of the next smallest group. Grouping by series length was the best, among the different groupings, in terms of imbalance. Data sets of lengths [501, 1000] were the smallest group with a count of 6 compared to the largest group, length [101, 250], of 19 data sets. Number of classes was highly imbalanced; the binary classification group contained 29 data sets,

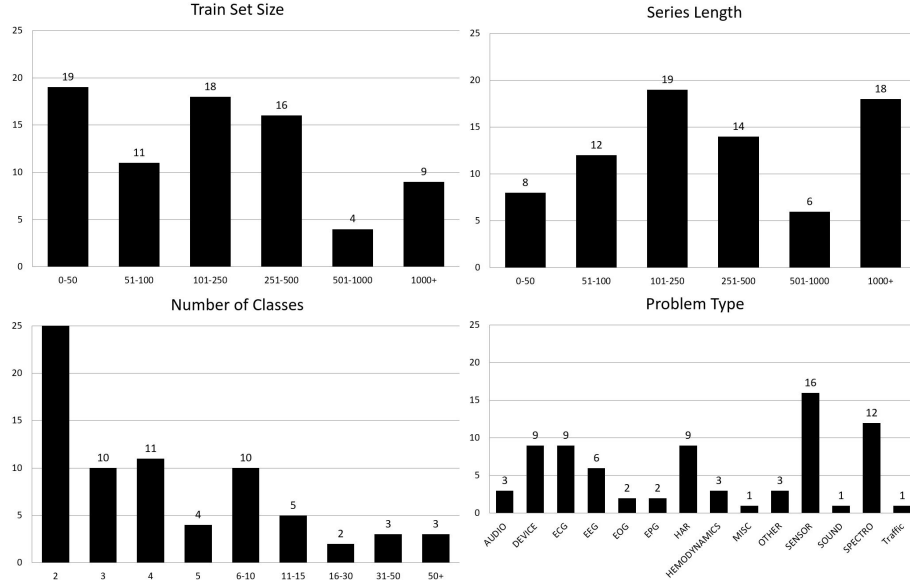


Figure 4.1: Metadata of the used data sets

which was more than double the second largest group [6, 10] of 10 data sets. This is expected because all the data sets from the UCR archive fall into the same bin, while the multivariate data sets, which are already less in number, are distributed among multiple bins. Finally, there was an over representation of type SENSOR than any other type with a total of 16 data sets. While multiple other groups were represented by very few data sets like MISC, SOUND and TRAFFIC which only had 1 data set; or like EOG and EPG which had 2 data sets.

Since the final goal of our work is to learn a recommender which will be able to suggest the best performing classifier to unseen data sets based on their properties. It is important to include all necessary and useful metadata about the data sets, which will help the recommender correctly give its recommendations. Imbalanced data sets can easily produce misleading results in terms of performance, specially if using a simple performance metrics like accuracy. In cases of extreme imbalance, one can attain very high accuracy scores, simply by always predicting the majority class. Handling the data imbalance is a two fold solution. The first step to help learn about the skewness is to have a definition for imbalance. Then based on this definition flag the data sets as either having balanced classes or not. The second step is to use a suitable performance metric which can give actual performance results and not be misled by the imbalance.

We discuss the first step here as it is a feature of the data set, the other step we consider in the design of our experiments in section 5.2. To be able to inform our recommender about the skewness property of a data set, we create a new feature, in addition to the previously discussed ones ; to indicate whether a data set is balanced in terms of class distribution or skewed. We define a data set to be skewed if there exists a class in this data set which is, at least, as twice as likely as any other class.

To better explain how this property works, consider the two data sets; DS_1 and DS_2 each of which contains 5 classes. DS_1 has a class distribution: $P(C_1) = 20\%$,

$P(C_2) = 15\%$, $P(C_3) = 15\%$, $P(C_4) = 25\%$ and $P(C_5) = 25\%$. While DS_2 has a class distribution: $P(C_1) = 20\%$, $P(C_2) = 15\%$, $P(C_3) = 15\%$, $P(C_4) = 30\%$ and $P(C_5) = 20\%$. In this case we would flag DS_1 as a balanced data set; because there is no prior class probability which is double any other. On the other hand, DS_2 would be flagged as skewed/imbalanced because class C_4 has a probability of 30% which is double the value of C_2 and C_3 . We denote this feature with the parameter *Balanced*, which is a boolean flag.

4.2 Karolinska data set

data set	Train	Test	Length	Classes	Type
ACSF1	100	100	1460	10	DEVICE
Beef	30	30	470	5	SPECTRO
Car	60	60	577	4	SENSOR
Chinatown	20	345	24	2	Traffic
CinCECGtorso	40	1380	1639	4	ECG
Coffee	28	28	286	2	SPECTRO
Computers	250	250	720	2	DEVICE
DodgerLoopDay	78	80	288	7	SENSOR
DodgerLoopGame	20	138	288	2	SENSOR
DodgerLoopWeekend	20	138	288	2	SENSOR
Earthquakes	322	139	512	2	SENSOR
ECG200	100	100	96	2	ECG
ECG5000	500	4500	140	5	ECG
ECGFiveDays	23	861	136	2	ECG
ElectricDevices	8926	7711	96	7	DEVICE
EOGHorizontalSignal	362	362	1250	12	EOG
EOGVerticalSignal	362	362	1250	12	EOG
EthanolLevel	504	500	1751	4	SPECTRO
FordA	3601	1320	500	2	SENSOR
FordB	3636	810	500	2	SENSOR
FreezerRegularTrain	150	2850	301	2	SENSOR
FreezerSmallTrain	28	2850	301	2	SENSOR
Fungi	18	186	201	18	OTHER
Ham	109	105	431	2	SPECTRO
HouseTwenty	34	101	3000	2	DEVICE
InsectEPGRegularTrain	62	249	601	3	EPG
InsectEPGSmallTrain	17	249	601	3	EPG
InsectWingbeatSound	25000	25000	600	10	AUDIO
ItalyPowerDemand	67	1029	24	2	SENSOR
LargeKitchenAppliances	375	375	720	3	DEVICE
Lightning2	60	61	637	2	SENSOR
Lightning7	70	73	319	7	SENSOR
Meat	60	60	448	3	SPECTRO
MelbournePedestrian	1194	2439	24	10	TRAFFIC
MoteStrain	20	1252	84	2	SENSOR
NonInvasiveFetalECGThorax1	1800	1965	750	42	ECG
NonInvasiveFetalECGThorax2	1800	1965	750	42	ECG
OliveOil	30	30	570	4	SPECTRO
Phoneme	214	1806	1024	20	SOUND

Chapter 5

Framework and Experimental Design

This chapter describes our proposed framework which compares TSCAs in a context that is inspired by the problem of eTSC and the design of our experiment. We begin by defining what we mean by early classification context and how does our framework simulate it. Then we discuss the structure of the framework, the software tools that were used to build it and how to evaluate it. In the second section of the chapter we represent our experimental setup.

5.1 The Framework

As motivated earlier, eTSC is a field which is concerned with classification of time series data with earliness and accuracy as the main objectives. Dedicated eTSCAs focus on building models that can learn class labels of the data as early as possible while maintaining accuracy [Mor+17a]. Our framework, on the other hand, investigates the adaptation of the context in which conventional TSCA operate to learn a specific classification problem and the effect it has on the classifiers' performance.

5.1.1 Early Classification Context

We define our early classification context as the problem in which a TSCA is trained on labeled but incomplete time series data, then used to classify data instances of full length.

To answer our first research question *How can we adapt existing ts classification workflow for the early TS scenario ?*

We propose a framework that simulates an early classification context and runs a series of experiments using different TSCAs. It divides whole timeseries data instances into smaller subseries using a chopping algorithm, then starts multiple learning processes. In each learning process, the data is exposed incrementally to the classifier by revealing the next sub-series. These processess simulate an incremental learning scenario, where the classifier is not exposed to all the data points collected from the beginning to the end at once. However in each learning process it gets to know more

information about the data instances than the previous process. In order to assess the performance of classifiers that were exposed to less data points, we use the performance of classifiers that were exposed to full length training data as a baseline. In the next subsection 5.1.2 we discuss the details and the structure of the framework.

5.1.2 Structure and Toolkits

We implemented our framework in python language. For the implementation of the non-deep learning algorithms we used the open source libraires *sktime*V.0.5.3 [Lön+19] and *pyts*V.0.11.0 [FJ20], while for the implementation of the deep learning algorithm InceptionTime we used the library *sktimedl*V.0.2.0 which provides an interface for the implementation provided in the study [Faw+20]. Finally, we used the library *Tslearn*V.0.5.0.5 [Tav+20] as a mediator between the different model implementations; due to it's capacity to process raw data into the needed input format by any of the libraries, as well as converting data format between the different libraires.

Our framework consists of two components; the comparison testbed and the recommender. The testbed is the engine responsible for comparing the implemented algorithms on the data sets provided. The testbed mainly collects user input about the experiment configuration, preprocesses the data in preparation for the learning process and finally calculates performance metrics using testing data. By the end of each performance assessment step, the testbed produces an analysis file and a logs file. The logs file provides information about the setup of the experiment, the execution status, the duration and reasons of failure. While the analysis file provides information about the cross validation scores, the final hyperparameter values used, processing time of training and testing steps and the performance measures.

The second step of the framework is learning a recommender which will be able to predict the best performing algorithm(s) for an unseen data set. Given the different analysis reports generated from the testbed, these reports are then consolidated and fed to the recommender to learn on. First, the logging files are processed and filtered to get the data of the successfully executed processes. Then the analysis files for these processes are consolidated into a unified structure and the included information is enriched by metadata about the data sets. In the end, the data is preprocessed again to be fed to a regression classifier which learns from the results of already carried out experiments. When given a new data set, the regression classifier tries to predict the performance of each implemented classifier on the data set. These scores are then used to rank the classifiers and the final recommendation is presented.

Our testbed component can be divided into several consecutive steps:

1. User input
2. Experiment preparation
3. Experiment execution
4. Analysis reports and logging

In the next subsections, we will discuss the details and implementation of each step involved in the construction of our 2 steps framework.

User Input

In the user input step, the user provides the necessary information required by the framework to start it's experiments. These information provide the guidelines of the

process which will be carried out and decide what steps should be done in the experiment preparation step. Table 5.2 shows a summary of each of the needed information, represented as a parameter for the framework.

Parameter	Data Type	Description
Dataset	String List	The name of the data set(s)
Splits	Integer	The number of splits to apply on the data set
Hyperparam	Boolean	Learn hyperparameters for the model or skip
NumIterations	Integer	Number of hyperparameters combinations to sample
ChunksToKeep	Integer List	The indexes of the chunks to use
FromBeg	Boolean	Reveal data chunks from beginning or end
ScoringFunction	String	The strategy for evaluating the model on test data

Table 5.2: Input parameters required for using the framework

We implemented the user input interface using *docopt*¹. A library which offers command-line interfaces through the use of simple arguments and elements. It also allows setting default values for parameters which can be overridden if a value is passed. In order to start running the framework, one provides the necessary parameter values through a simple terminal command. Listing 5.1 shows a sample command for running our master python file *run.py* on the Computers data set using hyperparameters optimization, while applying a split into 10 chunks.

```
python run.py --etsc --dataset=Computers --cv --split=10
```

Listing 5.1: Sample command for providing user input to the framework

Experiment Preparation

In the experiment preparation step, the framework starts preparing the different components of the experiment to be run. These components include data set management, models initialization, preparation of the early classification context and configuration of the training and testing processes.

Our framework's data sets handling module is configured to process data sets in either of two ways; automatic download from the UCR/UEA data archives, or manually provided data sets that comply with some specifications. Since the data loading

¹<https://github.com/docopt/>

module from *pyts* [FJ20] offers integrated tools with the UCR/UEA data archives for fetching and downloading data, we have configured our data module to extend *pyts* by transparently checking both data archives when provided with a data set name. The other option would be to provide a data set folder holding the name of the data set. Inside the folder there should exist one training data set file with the name “FolderName_TRAIN” and one testing data set file with the name “FolderName_TEST”. Which is the case for most of the data sets from both archives. For multivariate data sets, our framework assumes that the training data set file, as well as the testing data set file, contains data of all dimensions. For maximum compatibility, our framework extends the data processing utilities from *sktime* [Lön+19] which can read data sets that comply with the guidelines provided by the UCR and the UEA archives. We consider “arff” format as the primary expected data type.

As mentioned earlier in section 2.3, there are two ways to work on multivariate time series data sets [Rui+20]. The first way is by using bespoke multivariate classifiers which can deal with the multiple dimensions available in the data. While the other is by using ensembles of univariate classifiers, each fit to a separate dimension under the assumption of independence between the dimensions. Our framework applies the second technique for handling multivariate data sets.

In the experiment preparation step, the framework checks the data and determines the number of dimensions in consists of. If the data set is univariate then the model initializes the default classifier implementations provided by the libraries. In case the data set was found to be multivariate, we initialize a special classifier type which extends the usage of univariate classifiers for multivariate problems. This special classifier is referred to as *ColumnEnsembleClassifier* in the *sktime* library implementation and as the *MultivariateClassifier* in *pyts*.

There are two assumptions that we make at this step in our framework. First, we assume that to extend a specific univariate classifier to a problem, we should fit the same type of the classifier on all dimensions of the data. This is goes back to our goal, that we want to compare classifiers from different groups and not to create an ensemble which makes use of different classifiers applying different techniques, like HIVE-COTE does. Second, we assume that we always have to fit one classifier per dimension, which have proved to be not a feasible solution for high dimensional data sets. We will discuss this in more details in our results.

In preparation for the following steps in the framework, the experiment preparation step plays a role in configuring the training and the testing processes. For the training process, it is possible to either run using the default configuration of the classifiers from their libraries, or to learn it’s hyperparameters through cross validation. We use a randomized search over the hyperparameters space of classifiers. The framework parameter *NumIterations* represents the number of random samples to consider when optimizing the hyperparameters. If the space of the hyperparameters is less than *NumIterations*, then the space is exhausted by applying a grid search. We have considered 5-folds cross validation for all experiments.

For the testing process, the framework allows the usage of any chosen performance metric which is supported by *sklearn* library [Ped+11]. The chosen performance metric is utilized for the evaluation during both; the cross validation process and for calculating performance on the test data set. We consider “balanced accuracy” as our default performance metric; to account for problems where data sets have imbalanced classes distribution.

Based on the assumption that all data instances are of the same length, we implement our early classification context using a time series chopping algorithm. The

algorithm is provided with some time series data set T with instances of length L and three user defined parameters; s which decides the number of splits L should be split to, *FromBeg* a flag which indicates whether the data chopping should occur from the beginning or the end of the time series T , and *ChunksToKeep* which provides the user with the option to use only specific chunks of interest and exclude the processing for all other chunks. Algorithm 1 demonstrates how the chopping algorithm works.

The result of the algorithm is then used to create new copies of T ; $[T_1, T_2, \dots, T_s]$, each copy T_i containing all instances from T but revealing a subsequence of L which we call the chunk. A chunk is a sequence of p data points starting from the beginning or the end of the instances in T based on the value of the parameter *FromBeg*. Each data set copy T_{i+1} reveals one more chunk than the previous data set T_i and the data set T_s represents the set with instances of the full length L . It should be noted that our algorithm tries to have an equal value for p used for each chunk, but this is limited by the length of the time series and the provided number of splits. The resulting data sets are then passed to the training process, where each algorithm applies it's own technique to learn on the data set.

Algorithm 1 The Chopping Algorithm

```

1: function GetSplitIndexes( $T, s, ChunksToKeep$ )
2:    $L \leftarrow \text{ExtractLength}(T)$ 
3:    $ChunksSizes \leftarrow \text{GetChunkSizes}(L, s)$ 
4:    $SplitIndexes \leftarrow \text{CumSum}(ChunksSizes)$ 
5:   if KeepChunks is not Null then
6:      $FilteredSplitIndexes \leftarrow \text{KeepChunks}(SplitIndexes, ChunksToKeep)$ 
7:   else
8:      $FilteredSplitIndexes \leftarrow SplitIndexes$ 
9:   end if
10:  return  $FilteredSplitIndexes$ 
11: end function

```

To better understand how the chopping algorithm works, consider the next example. Let's assume we have a data set T with instances of length $L = 100$, we set the value of $s = 10$ and *FromBeg* = True. The algorithm starts by calculating the value of p for each of the required 10 chunks using the function 2.

The resulting list *ChunksSizes* from the algorithm will contain the value 10 for each of the chunks ($ChunksSizes = [10, 10, 10, 10, 10, 10, 10, 10, 10, 10]$); this is because the length of the time series is divisible by the number of splits provided, so it results in perfect splits of the data and thus chunks of equal sizes. If we assume the same scenario again but with a different s value like 8, the algorithm will try to provide as equal values of p as possible giving back a *ChunksSizes* of values $[13, 13, 13, 13, 12, 12, 12, 12]$.

After calculating the chunk sizes, the chopping algorithm translates *ChunksSizes* into a list of indexes called *SplitIndexes*. Which is then used to create the different copies T_i by selecting subsequences. The values of *SplitIndexes* are the indexes of the last time point that a specific chunk should read. We calculate these values by carrying out a cumulative sum over the values of *ChunksSizes*. For example, the list $ChunksSizes = [10, 10, 10, 10, 10, 10, 10, 10, 10, 10]$ is translated into the list $SplitIndexes = [10, 20, 30, 40, 50, 60, 70, 80, 90, 100]$. This means that the

Algorithm 2 Function to Get Chunks Sizes

```

1: function GetChunkSizes( $L, s$ )
2:    $ChunksSizes \leftarrow []$ 
3:   for  $i \leftarrow 0$  to  $s - 1$  do
4:     if  $i < L \pmod{s}$  then
5:        $p = (L \div s) + 1$ 
6:     else
7:        $p = L \div s$ 
8:     end if
9:      $ChunksSizes.append(p)$ 
10:  end for
11:  return  $ChunksSizes$ 
12: end function

```

corresponding data set T_1 will contain all its data instances represented by the subsequences from the 1st time point till the 10th time points, and the data set T_3 will contain subsequence from the 1st time point till the 30th time point. When we apply the same translation to the list $ChunksSizes = [13, 13, 13, 13, 12, 12, 12, 12]$, it gets translated into $SplitIndexes = [13, 26, 39, 52, 64, 76, 88, 100]$.

Finally, the algorithm filters out the list $SplitIndexes$, if the user provided a list of specific chunks to keep using the variable $ChunksToKeep$. The values passed through $ChunksToKeep$ corresponds to the indexes of the chunks the user is interested in, all the other chunks are excluded. Note that this process is different from reducing the number of splits; as the number of splits affects the sizes of the chunks being created. While $ChunksToKeep$ is used only to filter out the chunks after their sizes are already determined and translated into $SplitIndexes$. The space of possible values for $ChunksToKeep$ is between $[1, s]$. The logic of the function is demonstrated in function 3

Algorithm 3 Function to filter out Chunks

```

1: function KeepChunks( $SplitIndexes, ChunksToKeep$ )
2:    $FilteredSplitIndexes \leftarrow []$ 
3:   for  $i \in ChunksToKeep$  do
4:      $FilteredSplitIndexes.append(SplitIndexes[i])$ 
5:   end for
6:   return  $FilteredSplitIndexes$ 
7: end function

```

If we consider the list $ChunksToKeep = [1, 3, 5]$ for our example $SplitIndexes = [10, 20, 30, 40, 50, 60, 70, 80, 90, 100]$. The resulting filtered list will be $FilteredSplitIndexes = [10, 30, 50]$, which will result in only 3 copies of T ; T_1 , T_3 and T_5 , each represented by its respective subsequence length.

If we have a closer look at the parameter s , we can recognise that it contributes to the granularity of the earliness factor that we are calculating. The value of s decides the total number of chunks we will use, which subsequently decides also the ratio represented by each chunk to the total length of the time series T . That means the

greater the value of s , the smaller the chunk sizes and consequently the more granular results we can test for the algorithms. If we consider our previous example, when we set the value of $s = 10$ then the length of each subsequence would be close to 10% of the total length. If we decided to change the value of $s = 20$, then each chunk will roughly represents 5% of the total length and so on. Increasing the number of splits to be used comes with the cost of time and resources, each extra split adds one extra run per algorithm per data set.

Experiment Execution

The experiment execution step is where the actual processing happens. After the preparation of the chunks for the data sets in the previous step, learning processes start and each of the implemented classifiers starts training on the prepared data sets. Each of these training processes is then followed by a testing process, where performance scores are calculated using the function provided by the parameter *ScoringFunction* on testing data sets.

As mentioned in section 3.3, we follow the grouping criteria by [Bag+17] which arranges TSCAs into 5 main categories based on their techniques. We have included classifiers representing each of the 5 groups, in addition to a 6th group which represents the family of deep learning time series algorithms. We tried, as far as the implemented libraries allowed, to represent each group with 2 classifiers; one that is a non-ensemble and another which is an ensemble. A total of 10 classifiers are included in the framework covering all 6 groups, these can be broken down into the following:

- Distance based : 1NN using MSM distance and PForest
- Phase dependent algorithms : TSF
- Shapelets: LS and ST
- Dictionary based : WEASEL and BOSS
- Deep learning : InceptionTime
- Classical baselines: 1NN using euclidean distance and 1NN using DTW

Table 5.4 shows the configuration and hyperparameters for all the included classifiers. We used the same hyperparameter space like that of the original published papers as closely as possible. During training, the number of hyperparameter sampling is decided by the framework parameter *NumIterations* and we fix the number of cross validation folds to 5.

After the training processes are finished, the classifiers are then shown the testing data sets to evaluate their performance. If the experiment is configured to do hyperparameter optimization, then the classifier version that attained the highest validation score, from the 5 cross validation folds, is used for calculating the performance score on the testing data set.

Analysis Reports and Logging

The last step of our framework is the analysis report preparation. After the successful running of the experiments and calculating performance scores using the provided scoring function, we create a single report for each run experiment. We produce one report per classifier per data set. We refer here by data set to each data set that was finally produced by the chopping algorithm. These reports collect the essential data

Classifier	Parameters
MSM	$c = \{0.01, 0.1, 1, 10, 100\}$
PForest	$k = 100$ $r = 5$
TSF	$r = 500$ splitting = {entropy, gini} maxfeatures = {sqrt, log2}
LS	$\lambda_w = \{0.01, 0.1, 1\}$ $L_{min} = \{0.025, 0.075, 0.125, 0.175, 0.2\}$ $R = \{1, 2, 3\}$ $K = \{0.05, 0.15, 0.3\}$ $\eta = 0.01$ maxIter = {2000, 5000, 10000}
ST	$t = 60$ mins $n = \{500, 100\}$
CBOSS	$t = 60$ mins $k = \{50, 100, 250, 500\}$ $s = 250$ $p = [0.5 - 1]$
WEASEL	ANOVA = {True, False} bigrams = {True, False} binning = {equi-depth, equi-width, information-gain}
Inception	epochs = 1500 batch size = 64 $\eta = 0.001$ kernel sizes = {10, 20, 40}
DTW	full warping window

Table 5.4: Parameters and configuration for TSCAs

and statistics that form the basis of our results. Table 5.6 shows the structure of the output for the analysis report.

Most of these information are already available once the testing process is finished; because either they are metadata about the experiment or metrics calculated during training and testing. Metadata about the process is already known before the running of the process; like the *Classifier*, *Dataset* and *Revealed%*. While the processes metrics are *Traintime*, *Testtime*; which represent the total duration of the learning and testing processes, and the *Testscore* which represents performance of the classifier.

There is only one metric which is calculated during the analysis report phase after all the other metrics are calculated; the harmonic mean. We previously defined the HM in chapter 3.4, it has been used by [SL20] as the objective function for evaluating

Item	Description
Classifier	The classifier name
Train time	The total training time (CPU Time)
Test time	The total testing time (CPU Time)
Test score	Performance score on testing data set
Params	List of hyperparameters used and their values
Revealed %	Percent of data points used for training from the total length
Harmonic Mean	Harmonic mean between test score and revealed %
Data set	The name of the data set

Table 5.6: Analysis report created for each experiment

their algorithm *TEASER*. The HM is a popular choice for calculating a score that combines 2 objectives at the same time, which is in our case earliness and performance like the other eTSC problems. Earliness is represented by the value of the parameter *Revealed%* which is a fraction of the full series length, e.g. 0.5 means 50% of the total length. On the other hand, performance is represented by the value of the parameter *Testscore*. We use HM as the means of evaluating classifiers in our experiment; due to it's capacity to incorporate both objectives in one score. Which allows us to compare the trained classifiers in terms of a one score value without worrying about fixing the value of either the earliness or the performance.

5.2 Experiments

Our experiments were conducted on univariate data sets from the UCR archive as well as multivariate data sets from the UEA archive. The data sets were chosen based on the criteria discussed in the subsection 4.1.3. Each of the data archives offers a train and test split of the data which we have used unchanged for all the classifiers. All our experiments were run on a LINUX server with an AMD Ryzen 7 1700 Processor and 32GB RAM using Python 3.7.10.

We used the same experimental configuration through out all our experiments. We fixed the value of the parameter *Splits* = 10, that is for every data set the data would be revealed for the classifiers incrementally in batches of 10% from the total length. The chopping algorithm was set to always reveal data from the beginning of the time series. Due to time limitations, we restricted our experiments to run only on the 1st, 2nd, 3rd and 10th chunksof the data sets by setting *ChunksToKeep* = {1,2,3,10}. We included the 10th chunk to represent the baseline performance of each classifier if shown the full length data. While the first 3 chunks were used to represent the classifiers in the early context scenarios. Hyperparameters optimization was always carried out on all data sets with a *NumIterations* maximum sampling value of 50 iterations, unless proven unfeasible for any of the classifiers due to memory shortage

or time constraint. In this case the experiment is repeated for all classifiers without hyperparameters optimization. The selected performance metric for our experiments was *BalancedAccuracy*. Balanced accuracy is a performance metric which can handle data sets with skewed class distributions by avoiding inflated performance estimates. To calculate balanced accuracy, the *Recall* value is computed for each class then averaged over the total number of classes. We have set all our experiments to use the same configuration for all classifiers on the same data set.

5.2.1 Excluded Classifiers

Some of the included classifiers in the framework were excluded from our experiments; either because they couldn't operate in the early classification context or because they couldn't attain comparable results to the published performances by previously published frameworks.

For example, KNNED and InceptionTime were excluded due to the nature of their techniques which couldn't handle our created context. Both algorithms are clearly able to learn on chopped training data sets. However once the testing phase is reached, they would fail to classify instances of the testing data, showing errors related to mismatches between the expected length of instances and the provided length. Yet they would finish the last chunk where the data is provided in its original full length. The reason why KNNED fails such scenario, is that it uses ED which is a point-wise comparison distance measure that cannot compare time series of unequal lengths [Tan+19]. On the other hand, InceptionTime is a deep learning model whose input layer architecture, represented by number of nodes, depends on the length of the input time series [Faw+19a]. Since we use full length instances for testing, this caused an overflow of input data than what the model structure was expecting. There have been literature discussing adapting TSCAs to unequal time series, but this is out of the scope of our experiments. For more details refer to [CCP09; Tan+19; Faw+19a]

Although KNNDTW has been a competent time series classifier for decades; thanks to the exploitation of the elastic distance measure DTW. We couldn't get either implementation of KNNDTW from *sktime* and *pyts* to work on the chopped data; due to errors in the data representation needed by lower level internal libraries. KNNDTW would have been able to operate in the early classification context if used with a full warping window; which would have been successful in handling extreme classification cases like classifying full length data even when learning on the 10% chunk data set.

Two classifiers were excluded because they attained significantly inferior results compared to the published scores; these are KNNMSM and LS. Specially on the InsectWingbeatSound data set, for which we attained a difference in performance of -45.71% for KNNMSM and -20.71% for LS than the results published by [Bag+17]

5.2.2 Included Classifiers and Adjustments

Our experiments proceeded with the remaining 5 classifiers; PForest, TSF, ST, CBOSS and WEASEL. To focus only on the comparison of performance between classifiers and not preprocessing, we consider only data sets where all instances have the same length and no attributes have missing data. We also focus our interest on data which is originally a timeseries data. This means that it is a collection of specific measurements across a span of time. Which lead us to exclude some data set types from taking part in the experiment, more details are presented in subsection 4.1.3. We cover a total of

77 data sets from both the UR and the UEA archives, out of which 55 data sets are univariate and 22 data sets are multivariate.

Time and resources limitations played a big role shaping our experiments. Running classifiers is computationally expensive [SL20] and takes up to hundreds of processing days. This lead us to modify some algorithms to so that they can finish promptly; like in the case of PForest, or to use an enhanced version of the original classifier with an option to control the learning phase time; like BOSS and ST.

Due to time limitations; We used the contractable implementations of BOSS and ST offered by *sktime*. These implementations allows passing a time parameter which controls the sampling space for BOSS and the shapelets searching time for ST. We set the value of the time contract for both to 60 minutes.

We applied 2 adjustments to the PForest classifier. Firstly, we excluded ED from the pool of distance measures that PForest selects from; since the ED has previously proven not to be compatible with our early classification context when used with the KNN classifier. Secondly, we excluded TWE from the pool of distance measures when the size of the training data set exceeded 150 instances, which is the median of all data sets training sizes. TWE is a very slow distance measure. It has been reported by [Bag+17] to be the slowest among all elastic distance measures. When measured on classifying 10 test instances from the StarlightCurves data set, it performed 10,000 times slower than ED.

Chapter 6

New Framework

Chapter 7

Results

This chapter should discuss the Results

The baseline performance for each classifier is determined by it's performance on the full length data.

Chapter 8

Findings Discussion

This chapter should discuss the Findings

Chapter 9

Summary and Outlook

This chapter should discuss the Summary

Appendix

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9.1 Declaration of Authorship

I hereby declare that I have written this thesis "TITLE TITLE TITLE" without any help from others and without the use of documents and aids other than those stated above. Furthermore, I have mentioned all used sources and have cited them correctly according to the citation rules. Moreover, I confirm that the paper at hand was not submitted in this or similar form at another examination office, nor has it been published before.

Magdeburg, DATE, SIGNATURE