# Extracting Statistical Graph Features for Accurate and Efficient Time Series Classification

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## ABSTRACT

This paper presents a multiscale visibility graph representation for time series as well as feature extraction methods for time series classification (TSC). Unlike traditional TSC approaches that seek to find global similarities in time series databases (e.g., Nearest Neighbor with Dynamic Time Warping distance) or methods specializing in locating local patterns/subsequences (e.g., shapelets), we extract solely statistical features from graphs that are generated from time series. Specifically, we augment time series by means of their multiscale approximations, which are further transformed into a set of visibility graphs. After extracting probability distributions of small motifs, density, assortativity, etc., these features are used for building highly accurate classification models using generic classifiers (e.g., Support Vector Machine and eXtreme Gradient Boosting). Thanks to the way how we transform time series into graphs and extract features from them, we are able to capture both global and local features from time series. Based on extensive experiments on a large number of open datasets and comparison with five state-of-the-art TSC algorithms, our approach is shown to be both accurate and efficient: it is more accurate than Learning Shapelets and at the same time faster than Fast Shapelets.

# **1** INTRODUCTION

Time series data refer to sequences of data that are ordered either temporally, spatially or in another defined order. They can be frequently found in a variety of domains, including financial data analysis, medical and health monitoring and industrial automation applications [27, 28]. Recently, it turns out to be feasible to model software systems as time series in order to conduct malware detection and classification [43]. Due to their wide application scenarios and abundance, there has been an increasing need for efficient knowledge discovery methods to extract useful information from time series databases. One of the major tasks in time series mining is time series classification (TSC), which consists of applying a learning algorithm on labeled data to train a model that will then be used to predict the classes of samples from an unlabeled data set. Due to the sequential characteristic of time series data, state-of-the-art classification algorithms (such as SVM and Random Forest [13]) that perform well for generic data are generally not suitable for TSC. It is thus important and beneficial to have a feature extraction mechanism that transforms

the sequential characteristics of time series data into unordered feature vectors, so that any modern classification algorithm can be taken advantage of. After all, one of the most challenging aspects of TSC lies in the sequentiality property.

Traditionally, researchers often rely on one of the simplest classifiers for TSC: the *k* Nearest Neighbor (*k*NN) algorithm. As stated in [6], "all of the current empirical evidence suggests that simple nearest neighbor classification is very difficult to beat". To perform well, *k*NN classifiers leverage the Dynamic Time Warping (DTW) [7] distance which mitigates problems caused by distortion in the time axis. One intrinsic issue with DTW, however, is that it focuses on finding *global* similarities, *i.e.*, the overall curve *shape* of time series. It also requires applications to specify a proper warping window size or to properly align data samples. As a result, DTW can be sensitive to data alignment/segmentation and it performs better when data are properly curated. In practice, however, well-aligned time series data are difficult or expensive to come by [17].

To address the phase issue of DTW- and other distancebased 1NN approaches, the research community has proposed approaches that focuses on finding defining local features/subsequences in order to be invariant to data alignment and rotation. Popular methods that fall into this category include Bag-of-Patterns [31], SAX-VSM [39] and shapelets-based algorithms [47], such as Fast Shapelets (FS) [35] and Learning Shapelets (LS) [15]. The majority of these techniques have taken advantage of text feature extraction approaches - e.g., TF-IDF after converting time series into alphabetical strings. Such conversion is often done via Symbolic Aggregate approXimation (SAX) [30], which requires two parameters (i.e., cardinality and PAA window size) to be set and it may not always be trivial to find the best pair. Besides, many approaches attempt to find time series subsequences that are representative of each class, e.g., shapelets by definition are defining time series subsequences that are calculated by exhaustive or optimized search. Overall, many of these methods have suffered from high computation complexity or suboptimal classification accuracy [42].

Graph representations for TSC, on the other hand, have not been investigated extensively by the time series mining research community possibly due to their high computation complexity. Nevertheless, thanks to recent development of graph mining algorithms [5, 33], some of the formerly complex problems can be solved extremely efficiently with optimization and parallelization techniques [2]. Such advances give us the opportunity to re-evaluate the possibility of taking advantage of graph representations and extracting graph features for building an efficient and accurate TSC algorithm.

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This paper proposes a novel approach for TSC that considers time series as complex graphs/networks and extracts from these networks important statistical features, which are fed to modern generic classifiers to learn structural knowledge from the original time series. After evaluating the classification performance with a large open dataset, we find out that our approach is capable of making efficient and accurate classification predictions. The main contributions of this paper are listed as follows:

- We present a multiscale graph representation for time series, so that both global and local features from time series can be captured, making this approach agnostic to time series alignment and outperform major distance-based TSC algorithms.
- Since we transform time series that are intrinsically sequential into unordered feature vectors, it is then suitable for taking advantage of modern generic classifiers (*e.g.*, RF, SVM and XGBoost) for efficient feature selection and classification. This clear separation of feature extraction and actual classification can help researcher focus on finding insightful characteristics in time series data without the need for reinventing the wheel and designing a classifier from scratch specifically for time series.
- We propose a novel feature extraction and classification method for time series based on calculating probability distributions of small motifs (*i.e.*, repeated patterns) in visibility graphs and other statistical features such as density, degree statistics, assortativity and coreness. This feature extraction mechanism is parameter-free, so that it can be easy to use and help yield reproducible results.
- We have intentionally chosen a collection of statistical features that are computationally efficient to extract from graphs and validated their effectiveness in controlled experiments. Moreover, since our feature extraction and classification process is inherently parallel, it is suitable for and capable of large scale data explorations.
- After extensively evaluating our approach with a large number of open datasets and comparing with related research efforts, experiment results indeed suggest that accurate and efficient classification can be obtained following this paradigm.

The remainder of this paper is structured as follows. Section 2 lays down the necessary background. Next, we present our approach in section 3 and detail TSC accuracy and efficiency evaluation results along with a case study in section 4. For interested readers, section 5 introduces research work related to ours. Finally, we conclude the paper with future research directions in section 6.

## 2 BACKGROUND

Traditionally, time series refer to a sequence of numbers that are chronologically ordered. However, in the research community time series have a much broader scope and do not associate strictly with timestamps:

Definition 2.1 (Time series). A time series instance T is an ordered sequence of n real-valued variables, *i.e.*,  $T = (v_1, ..., v_n), v_i \in \mathbb{R}$ .

If we consider each point in a time series as a single feature in a vector, then time series data usually have a huge number of features. When considering these features as a vector in an *n*-dimensional space, time series data are often high dimensional. Due to difficulties to conduct knowledge discovery tasks on high dimensional data, it is frequently required to reduce the dimensionality of time series in order to improve computation efficiency:

Definition 2.2 (Dimensionality reduction). The dimensionality of a time series sample *T* is the length of *T*, denoted by |T|. If *T'* is an approximated representation of *T* and  $|T'| \ll |T|$ , then *T'* is a dimension-reduced representation of *T*.

The research community has proposed a number of dimensionality reduction techniques for time series, including sampling, Wavelet transform [25, 26], Piecewise Aggregate Approximation (PAA) [19], etc.. Among them PAA is perhaps one of the simplest and most widely applied approaches. PAA reduces time series  $T = (v_1, ..., v_n)$  from *n* dimensions to *s* dimensions by firstly dividing the data into *s* segments of equal size, then the approximation is a vector of the mean values of the data readings per segment [30]. Let  $T' = (v'_1, ..., v'_s)$  be this vector where  $v'_i$  is computed by equation 1. For the sake of simplicity,  $\frac{n}{s}$  is often chosen to be an integer or rounded to the nearest one.

$$v'_{i} = \frac{s}{n} \sum_{k=\frac{n}{s}(i-1)+1}^{\frac{n}{s}i} v_{k}$$
(1)

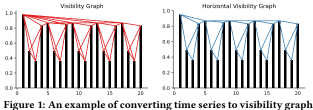
One of the core routines in distance-based classification algorithms involves evaluating the dissimilarity (or similarity) of two time series. There are a number of dissimilarity measures for time series, two of the most frequently used measures in the research community are Euclidean distance and DTW distance. The Euclidean distance maintains a one-to-one mapping of all the points in two series. On the other hand, the DTW distance tries to find the best mapping of points in two series using the dynamic programming paradigm, so that the minimum distance between these two series is achieved. The paradigm is called "time warping" since the time axis of series can be expanded or compressed in order to ensure the minimum distance, *i.e.*, an *i*<sup>th</sup> point in X can be mapped to a *j*<sup>th</sup> point (it is possible that  $i \neq j$ ), or one point in X may even be mapped to multiple points in Y.

### 2.1 Visibility Graph

Aiming for taking advantage of graph theories as a way of characterizing time series, an algorithm named visibility graph (VG) [22, 23] is proposed to transforms time series into a network structure. The creation of VGs relies on an extremely simple idea: each point in a time series is treated as a vertical bar, whose height is the corresponding numerical value. When considering these bars on a landscape, it is then straightforward that the top of a bar may be visible from the top of other bars. Assume each time-step as a vertex in a graph, then two vertices are connected if the top of the vertical bars are visible to each other, *i.e.*, there exists a straight line from the top of the two bars without intersecting with other bars. More formally,

Definition 2.3 (Visibility graph). Given a time series  $T = (v_1, ..., v_n)$ , its VG representation G = (V, E) has *n* vertices: V = (1, ..., n). An edge  $e = (i, j) \in E$  iff  $\forall k$  such that i < k < j  $(1 \le i, j \le n)$  inequality  $v_k < v_j + (v_i - v_j) \frac{j-k}{j-i}$  is satisfied.

It is obvious that VGs are undirected, although it is possible to create a directed version by limiting the direction of viewpoints from the vertical bars. Besides, VGs are always connected since each node will always be visible to its neighbors. Finally,



and horizontal visibility graph.

VGs are invariant when time series undergo affine transformations, *i.e.*, the visibility criterion remains fulfilled when rescaling the time series either horizontally or vertically. However, VGs are not suitable for non-stationary time series, *i.e.*, those having monotonically increasing/decreasing trends in time. Such trends should be removed before applying VG generation. The goal of VGs is to characterize structural properties [23] of time series, such as periodicity, fractality, *etc.*, although it is shown that it can be extended to weighted VGs in order to quantitatively distinguish generic time series [41].

Creation of VGs from time series without optimization generally has  $O(n^2)$  computation complexity, where *n* is the dimensionality of time series data. However, a more efficient VG generation algorithm [1] can have a sub-quadratic computation complexity. Specifically, this algorithm can reduce the complexity of time series VG generation down to  $O(n \log^2(n))$ . When further taking advantage of parallelization,  $O(n \log^2(n))$  work can be effectively solved within  $O(\log^2(n))$  time. Another simplified variant of VG, horizontal visibility graph (HVG) [32], only connects nodes *i* and *j* if a horizontal line can be drawn between these nodes. Creation of HVGs without any optimization generally has a computation complexity of O(n).

Definition 2.4 (Horizontal visibility graph). Given a time series  $T = (v_1, ..., v_n)$ , its HVG representation  $\overline{G} = (V, E)$  has *n* vertices: V = (1, ..., n). An edge  $e = (i, j) \in E$  iff  $\forall k$  such that i < k < j ( $1 \le i, j \le n$ ) inequality  $v_i, v_j > v_k$  is satisfied.

VG and its variants have been shown to be able to differentiate certain time series. For instance, [18] have extracted motifs from HVGs and claimed the motif statistics can be used for differentiating various types of time series data, including white Gaussian noises, fully chaotic logistic maps and noisy fully chaotic logistic maps. The authors further claimed that HVG motif profiles from heart rate time series can be used to cluster different types of meditative activities.

Intuitively, VGs and HVGs are very similar concepts and HVG is a subgraph of VG. However, when extracting statistics from them, VGs can be more capable of capturing global features, while HVGs are often more sensitive to local variations. As a result, VGs and HVGs can be joined to provide more accurate representations of time series data. We will discuss more about such heuristics in section 4.2. For simplicity, in the remainder of this paper the term VG indicates the combination of VG and HVG if HVG is not explicitly specified.

### 2.2 Graph Features

Extracting features from graphs has become a popular research topic thanks to the recent applications in social network analysis, physics as well as bio-informatics. There are a number of research avenues in graph mining, the most popular ones are about finding communities or clusters within large graphs and characterizing graphs by means of finding and counting recurrent patterns. Since time series VGs are always connected, it is thus not immediately helpful to extract clusters, since clusters in these graphs will always correspond to subsequences of original time series. Graph motifs or graphlets, however, are especially interesting since they are sub-graph structures or patterns in a larger graph that are recurrent and statistically significant. Finding motifs in graphs is generally a complex problem, but there have been a number of research work for efficiently extracting motifs from graph data, such as GTrieScanner [37] and Parallel Parameterized Graphlet Decomposition (PGD) [2]. Due to the fact that the number of motifs increases exponentially with motif size, the complexity of finding motifs is also exponentially correlated with motif size. Researchers thus generally focus on optimizing computation efficiency of locating small motifs (of size up to four). PGD is the state-of-the-art approach for efficiently finding and counting small motifs in graphs and can be several magnitudes faster than other approaches. Table 1 shows all possible small graph motifs up to size four. Note that PGD works only for undirected graphs, while GTrieScanner works on both directed and undirected graphs. However, motifs extracted by GTrieScanner are only connected motifs and the running time is significantly slower than PGD. As a result, in this paper we take advantage of PGD for small motif counting.

 Table 1: All graph motifs up to size 4. Note that connected graphs may contain disconnected motifs.

	#	Motif	Name		#	Motif	Name
	$M2_1$	1	2-edge		$M2_2$	•	2-node-independent
	$M3_1$	$\checkmark$	3-triangle		$M3_3$	•••	3-node-1-edge
_	$M3_2$		3-path	DISCONNECTED	$M3_4$	•••	3-node-independent
CONNECTED	$M4_1$	$\bowtie$	4-clique		$M4_7$	arphi	4-node-triangle
	$M4_2$	$\square$	4-chordal-cycle	Disco	$M4_8$		4-node-star
	$M4_3$	Ν	4-tailed-triangle		$M4_9$	•••	4-node-2-edges
	$M4_4$		4-cycle		$M4_{10}$	•••	4-node-1-edge
	$M4_5$	$\mathbf{\nabla}$	4-star		$M4_{11}$	•••	4-node-independent
	$M4_6$	$\square$	4-path				

Researchers argue that motif distribution extracted from VGs can be extremely helpful for identifying different types of artificially generated time series. However, in practice motif distributions may not be as distinguishable as those from artificial data. For example, Figure 2 illustrates the motif distributions of three different classes of time series from the ArrowHead dataset. As shown, it can be very difficult to tell one class apart from another given only these distributions since the probability distributions of different classes tend to overlap, especially in this case for instances from class 2 and 3. Besides small motifs, other graph features are also easy to obtain, *e.g.*, vertex and edge statistics as well as structural metrics. In this paper we consider some additional graph features listed as follows:

• Density: the ratio of the number of edges to the number of all possible edges. Graph density is computed following equation 2 and has a computation complexity of *O*(1).

$$p = \frac{2|E|}{|V|(|V|-1)}$$
(2)

• *K*-core: the *K*-core of a graph G = (V, E) is a maximal subgraph H = (V', E') in which each vertex has a degree of at least *K*, *i.e.*,  $\forall v \in V'$ :  $\deg_H(v) \ge K$ . Consequently, it is a cohesiveness measurement of interlinked subgraphs

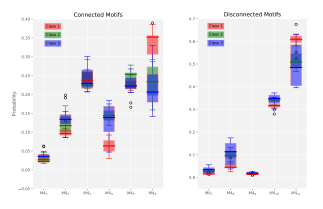


Figure 2: Boxplots of motif probability distribution of different classes from the ArrowHead Dataset's training set.

within a network and *K* is computed by equation 3. Computing *K* has an O(|E|) time complexity [5].

$$K = \operatorname*{argmax}_{H \in \mathcal{O}} \operatorname{Core}_{H} \ge K \tag{3}$$

 Assortativity coefficient: a metric to measure the correlation of vertices in a graph through calculating the Pearson correlation coefficient of degree between pairs of connected vertices. Equation 4 shows how assortativity coefficient is computed,

$$r = \frac{\sum_{xy} xy(e_{xy} - a_x b_y)}{\sigma_a \sigma_b} \tag{4}$$

where  $a_x$  and  $b_y$  represents respectively the fraction of edges that start and end at vertices with values x and y, and  $e_{xy}$  is the measure of assortativity, such that  $\sum_{xy} e_{xy} = 1$ ,  $\sum_y e_{xy} = a_x$  and  $\sum_x e_{xy} = b_y$ . Finally,  $\sigma_a$  and  $\sigma_b$  are the standard deviations of the distributions  $a_x$  and  $b_y$ . Computing r has an O(|E|) time complexity [33].

 Degree statistics including max, min and mean degree per vertex. Naturally, computing such statistics has an O(|V|) time complexity.

Note that there are a plethora of features that can be extracted from graphs [11], and such features are not necessarily equally easy to obtain. For instance, the diameter of a graph – which is the shortest distance between the two most distant vertices in the graph – can be computationally expensive to calculate since it demands O(|V|(|V| + |E|)) computation. Nonetheless, this paper does not intend to find the exclusive set of features that are both efficient and descriptive for time series VGs. Rather, we try to investigate if these aforementioned statistical features are indeed helpful for TSC.

## **3 MULTISCALE VISIBILITY GRAPH**

Time series data differ greatly in characteristics depending on how they are captured, sampled and their underlying applications: in one domain global features may be helpful, while in another domain local features (*i.e.*, defining subsequences) can become more important for classification. After transforming real-valued time series into VGs, specific features from such VGs – such as probability distributions of small motifs – are more reflective of local features than global ones. We refer to this type of representation as Uniscale Visibility Graph (UVG) in the remainder of this paper. In this case, extracting more global features (*i.e.*, probability distributions of large motifs) becomes exponentially expensive for computation. To mitigate this problem, we propose the Multiscale Visibility Graph (MVG) representation such that each time series sequence is transformed into a set of dimensionality-reduced approximations: these downscaled approximations are then converted into a set of VGs. Consequently, features are extracted from each graph in MVGs, thus approximating the process of extracting features of different scales. This approach is inspired by computationally efficient wavelet transform [10], with the exception that time series in each scale are represented with graph structures instead of numerical values. We first define the multiscale representation of time series as follows:

Definition 3.1 (Multiscale Approximation). Given a time series  $T_0 = (v_1, ..., v_n)$ , its approximated multiscale representation is a set of time series  $\hat{\mathbb{T}} = (T_1, T_2, ..., T_m)$ , where  $T_i$   $(1 \le i \le m)$  is a downscaled approximation of  $T_0$ , such that  $|T_i| = |T_0|/2^i = n/2^i$ . In addition, to avoid tiny and meaningless representations we enforce a constant threshold  $\tau$  for the downscaled approximations, *i.e.*,  $|T_m| > \tau$ .

The downscaling of  $T_0$  can be achieved with widely used dimensionality reduction techniques such as PAA (*cf.* equation 1). Since the size of downscaled time series representations follows an exponential decay, time series multiscale representations  $\hat{\mathbb{T}}$  often consists of a small number downscaled series. Besides, considering  $\sum_{i=1}^{\infty} n * 2^{-i} = n$ , theoretically the maximal dimension of  $\hat{\mathbb{T}}$  when fully expanded is n.

Definition 3.2 (Multiscale representation). Given a time series  $T_0$  and its approximated multiscale representation  $\hat{\mathbb{T}} = (T_1, T_2, ..., T_m)$ , its multiscale representation is the union of  $\hat{\mathbb{T}}$  and  $T_0$ , *i.e.*,  $\mathbb{T} = (T_0, T_1, T_2, ..., T_m)$ .

Approximated multiscale representations can help smoothing time series and reducing noises, while full multiscale representations consist of both the original time series and augmented versions. Each series in multiscale representations can be transformed into VGs, thus we have a natural definition for multiscale visibility graphs, which are supersets of approximated multiscale visibility graphs (AMVGs):

Definition 3.3 (Multiscale visibility graph). Given a time series T and its multiscale representation  $\mathbb{T} = (T_0, T_1, T_2, ..., T_m)$ , its multiscale visibility graph is a set of graphs  $\mathbb{G} = (G_0, G_1, G_2, ..., G_m)$ , where  $G_i \ (1 \le i \le m)$  is the corresponding visibility graph created from  $T_i$ .

Since the number of vertices in  $G_i$  equals the dimensionality of  $T_i$ , to avoid meaningless trivial graphs it is natural to set  $\tau$ to a small integer (*e.g.*,  $\tau = 15$ ), such that the smallest graph in  $\mathbb{G}$  contains more than  $\tau$  vertices. Note that  $\tau$  should not be considered as a parameter for the feature extraction process. Rather, it is more an optimization trick and bearing a default value of 0 will not cause any issues, since feature selection is done during classification.

### 3.1 Feature Extraction

As shown in Algorithm 1, feature extraction in MVGs follow the same paradigm as extracting features from individual graphs in an MVG and concatenating all features together, since graph features extracted in this paper are solely statistical and do not pertain orders. Consequently, these features can be fed into any generic classification algorithms [13] well studied in the machine learning and data mining community. It is utterly important that this feature extraction transforms the sequential characteristics of time series data into unordered feature vectors, so that any modern classification algorithm can be taken advantage of.

**Algorithm 1** Building time series MVGs and extracting features from them.

1: pr	ocedure ExtractFeatures(T)	
2:	$\mathbb{F} \leftarrow \emptyset$	▹ Feature set
3:	$\mathbb{G} \leftarrow \emptyset$	► MVGs
4:	$T' \leftarrow T$	
5:	while $ T'  > \tau$ do	▹ Ignore trivial graphs
6:	$\mathbb{G} \leftarrow \mathbb{G} \cup BuildVGAndHVG(T')$	
7:	$T' \leftarrow DimensionalityReduction(T)$	$\triangleright  T'  \le \frac{ T }{2}$
8:	for $G \in \mathbb{G}$ do	
9:	$\mathbb{M} \leftarrow MotifProbabilityDistribution($	G)
10:	$\mathbb{M} \leftarrow Normalize(\mathbb{M})$	
11:	$\mathbb{F} \leftarrow \mathbb{F} \cup \{\mathbb{M}, OtherStatistics(G)\}$	
12:	return ℝ	

Although features have become unordered, it is nevertheless important to carefully curate them in order to achieve better classification results. Note that PGD is very efficient in counting motifs in graphs, and the dominant features from time series MVGs will be the probability distribution of motifs of different sizes:

Definition 3.4 (Motif probability distribution). Given a time series visibility graph G and the set of motifs  $\mathbb{M}$ , the motif probability distribution  $\mathbb{P}_G$  is the set of probabilities corresponding to each motif in  $\mathbb{M}$ .

Empirically, the distribution of connected and disconnected motifs vary greatly in graphs. It is thus desirable to calculate separately motif probability distributions depending upon motif connectivity. To that end, the motif probability distributions (MPDs) are calculated per motif size and connectivity. This essentially normalizes extracted motif features. Specifically, MPDs are normalized according to the following five groups (*cf.* Table 1): { $M2_1, M2_2$ }, { $M3_1, M3_2$ }, { $M3_3, M3_4$ }, { $M4_1, ..., M4_6$ }, { $M4_7, ..., M4_{11}$ }. Finally, it is obvious that other graph features – *e.g.*, graph density and assortativity coefficient – are independent of MPDs. As a result, no further curation for such features is needed.

## 3.2 Classification

When features are extracted, we can feed them into a generic classifier to learn from labeled samples and predict the class of unlabeled ones. We are in favor of most well-known and widely accepted and well optimized algorithms such as SVM, Random Forest and eXtreme Gradient Boosting (XGBoost) [8] for this task. Especially, as a highly optimized distributed gradient boosting library, XGBoost runs on major distributed environment. It has gained remarkable adoption since its inception and is well known for its performance in machine learning and data mining competitions.

Typical classification tasks involve learning models and selecting a performant estimator through the process of validation. Although in this study we have proposed a parameter-free method for feature extraction, it is still required to tune the hyperparameters for generic classifiers. Note that hyper-parameters in machine learning refers to parameters that are external to the model and such values cannot be estimated from the training data. As a result, they are often set using heuristics. For instance, the *k* in *k*NN classifier can be considered as a hyper-parameter, since "there is no analytical formula available to calculate an appropriate value" [21]. In order to tune hyper-parameters in our classifiers, we conduct cross-validation to evaluate the performance of estimators and apply grid search to find the most satisfactory estimator based on the cross entropy scores:

$$-\log P(\hat{y}|y) = -(y\log(\hat{y}) + (1-y)\log(1-\hat{y}))$$
(5)

where y is the ground truth, and  $\hat{y}$  is the probabilistic predictions by an estimator. Since datasets may be highly imbalanced, which will lead to degraded classification results, we can apply random oversampling techniques over the minority class and use stratified cross validation to preserve class balance when validating models.

#### 4 EVALUATION

To evaluate our approach, we conduct experiments using a large number of publicly accessible datasets. We first introduce the datasets. Then we list and validate our heuristics, followed by creating an accurate classifier with stacked generalization. Next, we compare the accuracy and efficiency of our method with stateof-the-art approaches. Finally, we close this section with a case study and discussions.

## 4.1 Datasets

When validating our heuristics, we consider the most popular and largest open dataset for TSC: the UCR Time Series Classification Archive [9]. Specifically, we use a subset from the UCR archive that are more recent (added after the summer of 2015), which include datasets from various fields ranging from electrocardiograms (ECG) to intra-species image recognition data. These datasets have a uniform file format and consistent internal data structures, making it possible to conduct batch processing in a content-agnostic manner. Furthermore, datasets in this subset are generally larger in size, making it more reliable to evaluate the scalability of TSC algorithms.

When comparing our approach with state-of-the-art approaches, we take advantage of the UEA & UCR Time Series Classification Repository since it contains all datasets from the UCR archive. In addition, this repository also provides open source implementation for more than 18 TSC algorithms [3] and benchmarking results are publicly available from the repository website<sup>1</sup>. We compare our results with relevant algorithms using the default train and test split from this repository.

## 4.2 Validating Heuristics

Before diving directly into MVG representations, it would be a prerequisite to validate our heuristics step by step. We summarize these heuristics below:

- Motif statistics from VGs and HVGs can serve better during classification when combined with other graph features such as degree statistics.
- (2) Features from HVGs are more capable of capturing *local* characteristics while those from VGs are capable of capturing *global* characteristics. Combining features from both HVGs and VGs can help yield more accurate classification results.
- (3) Multiscale representations are able to reveal time series features at different scales, and a generic classifier is able to conduct feature selection and find important features to perform more accurate classification than using features without multiscale augmentation.

<sup>&</sup>lt;sup>1</sup>http://timeseriesclassification.com/

When validating these heuristics, we feed the features corresponding to each heuristic to an XGBoost classifier for 3-fold cross-validation and model selection. A set of hyper-parameters have been set for grid search, including the learning rate (three choices from 0.01 to 0.3), number of estimators (10 choices from 10 to 100), max tree depth (10 or 20). In order to prevent overfitting, the subsampling and column sampling hyper-parameters are both set to 0.5 to randomly collect half of the data instances and features to grow trees. To reduce the impact of random over sampling on minority classes and floating-point summation issues in parallel processing, all experiments have been repeated five times and average accuracy scores are calculated. Finally, all experiments are conducted on a Linux server with two Intel Xeon E5-2430 CPUs with a clock rate of 2.20GHz. With this setup, we illustrate step by step how experiments are setup and present the final classification results in Table 2.

4.2.1 Choosing Graph Features. We first investigate which graph features are helpful for extracting distinguishable information from time series. Specifically, we transform time series into UVGs consisting of both VG and HVG representations and try to extract MPDs as well as other features (density, assortativity and degree statistics) from these graphs. We are interested in finding out whether MPDs are sufficient for TSC and if extracting other statistic features from graphs are necessary. Next, we take advantage of XGBoost classifier to train on different feature sets and classify the test datasets. Columns A, B, C and D in Table 2 presents the classification error rates using HVG with only MPDs, HVG with MPDs and other graph features, VG with only MPDs and VG with all features. Comparison on the bottom of the table shows that including features such as density and assortativity coefficient indeed helps improving classification accuracy. For HVGs, including features other than MPDs increases classification accuracy in 32 datasets, while for VGs 29 datasets saw accuracy improvements. A Wilcoxon signed rank test with p-values 9.48e-7 and 9.56e-5 suggests that such improvement is indeed significant (both *p*-values < 0.05). Figure 3 shows the classification results in the form of scatter plots, where each point represent a dataset. Such results along with the Wilcoxon test indeed suggest that our Heuristic (1) is valid.

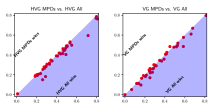


Figure 3: Comparison of classification error rates: using MPDs with or without other graph features.

4.2.2 VG and HVG. Next, it is necessary to show that graph features extracted from both VGs and HVGs are important. Recall that, intuitively, VGs are helpful for capturing global features while HVG can help locating local features. We separately test the distinguishing power of VGs and HVGs for time series in order to make sure that both can lead to satisfactory classification results. Furthermore, we conduct experiments to investigate if combining VGs and HVGs can better capture both global and local features for time series and thus lead to more accurate classification.

Figure 4 illustrates the comparison of classification accuracy with VG and HVG features as well as combining both VG and

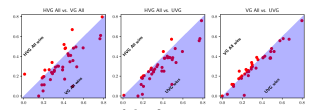


Figure 4: Comparison of classification error rates: using HVGs, VGs or combining two together (denoted as UVG here).

HVG. The first scatter plot shows that for majority datasets, VG features can yield more accurate classification performance. However, it is still worth mentioning that HVG features can also outperform VG features in some specific datasets, possibly due to the reason that local features are more influential in such datasets. Moreover, it is obvious from the other two scatter plots in Figure 4 that combining both VG features and HVG features can greatly boost the classification accuracy. This is probably due to the excellent feature selection capability of XGBoost, so that the classifier is able to find out which features are more important during the training process. In addition, as shown in Table 2, using VGs outperforms HVGs in 30 datasets with a pvalue of 3.09e-3, suggesting VGs are capable of capturing more characteristics in time series. Finally, combining features from both VGs and HVGs yields more accurate results in 27 datasets with a *p*-value of 5.01e-3, indicating significant improvement when combining two different types of graphs. As a result, the validity of our Heuristic (2) can be confirmed.

4.2.3 UVG, AMVG and MVG. Since we have demonstrated that taking advantage of both VG and HVG features can improve classification accuracy for UVG representations, now we can investigate whether Heuristic (3) holds, *i.e.*, if multiscale representations can help achieving even more accurate results. To visually inspect which representation suites best for TSC, we further draw scatter plots of the accuracy results in Figure 5. It is then obvious that AMVG and UVG (scatter plot on top) lead to similar classification accuracy, which suggests that AMVG can be good approximations for original time series data. Furthermore, the two scatter plots in the bottom of Figure 5 indeed confirm that MVG representations result in better classification performance, since almost all dots representing results from different datasets fall on the side of MVG.

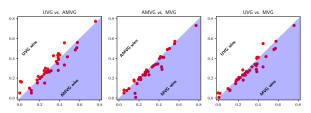


Figure 5: Comparison of UVG, AMVG and MVG's error rates.

From Table 2, we can see that multiscale approximations outperforms uniscale time series in 19 datasets, with a Wilcoxon *p*-value of 0.8623 > 0.05. As a result, AMVG representations are not statistically significantly different than UVGs. On the other hand, MVGs outperform AMVG in 29 datasets with a *p*-value of 1.72e-4 and MVGs are more accurate than UVGs in 30 datasets with a *p*-value of 8.74e-4, suggesting that MVG representations indeed contribute significantly towards a more accurate feature extraction and TSC process. As a result, our Heuristic (3) is valid.

						$Scales \rightarrow$		AMVG MVG					
					Туре	of Graphs $\rightarrow$	HVG		V	'G		VG+HVG	
						$Features \rightarrow$	MPDs	All	MPDs	All		All	
Dataset	#Cls.	#Train	#Test	Dim.	1NN-ED	1NN-DTW	A	В	C	D	E	F	G
ArrowHead	3	36	175	251	0.200	0.200	0.482	0.449	0.406	0.407	0.385	0.405	0.398
BeetleFly	2	20	20	512	0.250	0.300	0.440	0.410	0.250	0.170	0.250	0.150	0.180
BirdChicken	2	20	20	512	0.450	0.300	0.260	0.150	0.000	0.000	0.000	0.050	0.050
Computers	2	250	250	720	0.424	0.380	0.294	0.284	0.367	0.338	0.281	0.292	0.266
DistalPhalanxOutlineAgeGroup	3	139	400	80	0.218	0.228	0.204	0.202	0.214	0.196	0.202	0.196	0.188
DistalPhalanxOutlineCorrect	2	276	600	80	0.248	0.232	0.409	0.389	0.251	0.263	0.251	0.264	0.23
DistalPhalanxTW	6	139	400	80	0.273	0.272	0.342	0.348	0.298	0.300	0.315	0.275	0.279
ECG5000	5	500	4500	140	0.075	0.075	0.289	0.182	0.137	0.112	0.116	0.076	0.075
Earthquakes	2	139	322	512	0.326	0.258	0.262	0.255	0.286	0.265	0.245	0.276	0.283
ElectricDevices	7	8926	7711	96	0.450	0.376	0.503	0.493	0.392	0.357	0.366	0.368	0.338
FordA	2	1320	3601	500	0.341	0.341	0.009	0.009	0.254	0.220	0.007	0.167	0.000
FordB	2	810	3636	500	0.442	0.414	0.328	0.318	0.313	0.220	0.271	0.257	0.23
Ham	2	109	105	431	0.442	0.414	0.328	0.313	0.313	0.290	0.271	0.389	0.343
HandOutlines	2	370	1000	2709	0.199	0.197	0.293	0.288	0.275	0.225	0.221	0.215	0.206
Herring	2	64	64	512	0.484	0.469	0.425	0.419	0.450	0.484	0.381	0.431	0.28
InsectWingbeatSound	11	220	1980	256	0.438	0.407	0.425	0.763	0.586	0.404	0.557	0.484	0.488
LargeKitchenAppliances	3	375	375	720	0.438	0.422	0.303	0.703	0.380	0.478	0.380	0.346	0.4325
Meat	3	575 60	575 60	448	0.067	0.203	0.401	0.414	0.490	0.478	0.380	0.340 0.050	0.323
	3				0.067	0.067	0.497	0.490	0.160	0.097 0.238	0.117		0.080
MiddlePhalanxOutlineAgeGroup		154	400	80								0.266	
MiddlePhalanxOutlineCorrect	2	291	600	80	0.247	0.318	0.534	0.531	0.305	0.294	0.337	0.426	0.314
MiddlePhalanxTW	6	154	399	80	0.439	0.419	0.471	0.443	0.419	0.426	0.401	0.434	0.410
PhalangesOutlinesCorrect	2	1800	858	80	0.239	0.239	0.397	0.391	0.302	0.291	0.288	0.272	0.264
Phoneme	39	214	1896	1024	0.891	0.773	0.798	0.786	0.812	0.797	0.759	0.772	0.730
ProximalPhalanxOutlineAgeGroup	3	400	205	80	0.215	0.215	0.207	0.194	0.185	0.192	0.178	0.152	0.170
ProximalPhalanxOutlineCorrect	2	600	291	80	0.192	0.210	0.280	0.267	0.167	0.165	0.174	0.181	0.14
ProximalPhalanxTW	6	205	400	80	0.292	0.263	0.303	0.307	0.257	0.259	0.257	0.300	0.233
RefrigerationDevices	3	375	375	720	0.605	0.560	0.533	0.523	0.526	0.494	0.478	0.415	0.417
ScreenType	3	375	375	720	0.640	0.589	0.506	0.486	0.678	0.669	0.572	0.506	0.499
ShapeletSim	2	20	180	500	0.461	0.300	0.189	0.194	0.067	0.051	0.017	0.161	0.047
ShapesAll	60	600	600	512	0.248	0.198	0.715	0.595	0.585	0.485	0.448	0.332	0.313
SmallKitchenAppliances	3	375	375	720	0.659	0.328	0.239	0.212	0.282	0.261	0.205	0.205	0.206
Strawberry	2	370	613	235	0.062	0.062	0.217	0.205	0.117	0.113	0.097	0.099	0.094
ToeSegmentation1	2	40	228	277	0.320	0.250	0.354	0.339	0.289	0.301	0.296	0.261	0.259
ToeSegmentation2	2	36	130	343	0.192	0.092	0.185	0.185	0.205	0.182	0.185	0.218	0.185
UWaveGestureLibraryAll	8	896	3582	945	0.052	0.034	0.551	0.498	0.516	0.482	0.386	0.278	0.265
Wine	2	57	54	234	0.389	0.389	0.493	0.404	0.530	0.519	0.448	0.556	0.548
WordSynonyms	25	267	638	270	0.382	0.252	0.794	0.770	0.662	0.610	0.578	0.559	0.571
Worms	5	77	181	900	0.635	0.586	0.492	0.470	0.414	0.409	0.387	0.448	0.409
WormsTwoClass	2	77	181	900	0.414	0.414	0.328	0.306	0.242	0.248	0.243	0.239	0.23
		Com	parison	versus	G	G	В	D	D	Е	F	G	Е
Nu	mber of	f more acc			26	23	32	30	29	27	19	29	30
		Wilcoxe	on test p	-value	0.01	0.1638	9.48e-7	3.09e-3	9.56e-5	5.01e-3	0.8623	1.72e-4	8.74e

 Table 2: Error rates of classifying 39 UCR datasets compared with 1NN-Euclidean and 1NN-DTW. Different heuristic combinations are taken into account. Bold-faced values indicate lowest error rates (including ties) for specific datasets in all experiments.

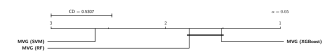
4.2.4 Summary of Heuristic Validation. Overall, all our heuristics are supported by experiment results and each heuristic contributes significantly to a more accurate TSC process. Table 2 also shows the comparison between our approach and Euclidean distance- as well as DTW-based nearest neighbor classification. Our approach outperforms 1NN-Euclidean in 26 datasets with a Wilcoxon *p*-value of 0.01, suggesting MVG features with XGBoost is significantly more accurate than 1NN-Euclidean. Moreover, MVG appears to be in par with 1NN-DTW in terms of classification accuracy since 23 datasets are in favor of MVG with a *p*-value of 0.1638. Next, we will try to build a more robust and accurate classifier using stacked generalization.

## 4.3 Stacked Generalization

Previously we have solely taken advantage of XGBoost for classifying time series with features extracted from graphs. However, it can also be interesting to investigate if other classifiers can achieve similar results. Besides, although modern classifiers such as XGBoost and RF are capable of efficiently conducting feature selection during training, we can still conduct feature selection processes and feed such *a priori* information to classifiers in order to achieve better classification accuracy. To that end, we propose feeding different sets of features to a collection of classifiers and then create a meta-classifier using stacked generalization (*a.k.a* stacking or blending) [44]. This meta-classifier will then hopefully generate better final predictions. In fact, a variant of this technique has led to winning the Netflix Prize with a reward of one million dollars [40].

Before building a meta-classifier with stacked generalization, we first make sure that features we previously extracted are suitable inputs for different classifiers such as RF and SVM. Generally, tree-based classifiers are not so sensitive to monotonic transformations of individual features. As a result, it is often not required to have features scaled to similar magnitudes for RF and XGBoost. However, since SVM's kernel functions (in an Euclidean space) are usually sensitive to different feature magnitudes, we use Min-Max scaling to transform each feature into range of zero and one. After that, we compare the classification performance of these three classifiers.

In order to evaluate the significance of the differences a generic classifier can incur, we take advantage of the Nemenyi test [12], which is a post-hoc test aiming for finding whether groups of data differ after a statistical test of multiple comparisons. This test can be illustrated by means of a critical difference diagram, where average ranks of all approaches are presented. Specifically, the location of vertical lines indicate the average ranking of an approach and bold lines (insignificance lines) indicate groups of approaches that are not significantly different. In this case, for one approach to be considered significantly better than another, its overall ranking has to be at least 0.5307 higher than its competitor. As shown in Figure 6, XGBoost performs slightly better than RF in general, and both are significantly more accurate than SVM. Such results are not surprising, since [13] have empirically tested hundreds of classifiers and concluded that RF produces the most accurate classification results. This study was conducted before the initial release of XGBoost, and the recent adoption momentum of XGBoost indeed suggests that XGBoost is great for yielding accurate classification results.



# Figure 6: Critical difference diagram comparison of RF, SVM and XGBoost.

Now that generic classifiers can be used for graph features extracted from time series, we then set to investigate whether stacking can help further increase classification accuracy. We first stack top performing classifiers in each family before blending classifiers from different families. Specifically, we first select the top five most accurate classifiers from RF, SVM and XGBoost through cross validation. Then these five classifiers are stacked to produce a meta-classifier, which is used for producing final predictions. Finally, when stacking classifiers of different families, five classifiers from each family have been selected, thus the metaclassifier has been trained with 15 classifiers. Our algorithm is described in Algorithm 2.

**Algorithm 2** Algorithm for creating an ensemble classifiers using stacked generalization.

	<b>Input:</b> Training dataset $\mathbb{D} = \{\mathbb{X}_i, y_i\}_{i=1}^m (\mathbb{X}_i \in \mathbb{R}^n, y_i \in \mathbb{Z})$ Base classifiers $\mathbb{H}$ (with different hyper-parameters) <b>Output:</b> An stacked ensemble <i>E</i>								
1:	procedure BuildStackingEnsemble(D, H)								
2:	$\mathbb{S} \leftarrow \text{CreateStratifiedKFolds}(\mathbb{D}, \text{cv=3})$	► 3-fold CV							
3:	$\mathbb{E} \leftarrow \emptyset$ >	Best performing base estimators							
4:	for all $h \in \mathbb{H}$ do								
5:	$H \leftarrow \emptyset$								
6:	for all $S = \{\mathbb{D}_{train}, \mathbb{D}_{validation}\}$	$e \in \mathbb{S}$ do							
7:	TrainClassifier( $h$ , $\mathbb{D}_{train}$ )								
8:	$\hat{y} \leftarrow \text{Predict}(h, \mathbb{X}_{validation})$								
9:	score $\leftarrow -\log P(\hat{y} y_{validation})$	)							
10:	$H \leftarrow H \cup \{h, score\}$								
11:	$H \leftarrow \arg \operatorname{sort}(H)$	▹ Sort estimators by score							
12:	$\mathbb{E} \leftarrow \mathbb{E} \cup \text{slice}(H, k)$	▹ Select top-k estimators							
13:	$W \leftarrow \text{ComputeEstimatorWeights}(\mathbb{E})$	▶ with logistic regression							
14:	$E \leftarrow \sum_{i=1}^{ \mathbb{E} } W_i \mathbb{E}_i$								
15:	return E								

Our experiments show that, for RF and SVM, stacking their top performing classifiers indeed increases final classification. In the case of XGBoost, however, stacking its most accurate classifiers does not seem to significantly increase classification accuracy, since the classification results of stacked generalization is on par with those with a single most accurate classification during cross validation. It possibly indicates that XGBoost has already very good generalization capabilities. Finally, stacking most accurate classifiers from three different families can help achieving better classification accuracy than single best XGBoost classifier in most datasets. As a result, stacked generalization can indeed be helpful for further improving the performance of MVG.

Figure 7 demonstrates how stacked generalization can help boosting classification accuracy. It is straightforward that stacking XGBoost and SVM produces similar classification accuracy, while stacking top performers from all three families can be significantly more accurate than using a single family. As a result, we are confident that staked generalization is favorable for more accurate TSC.

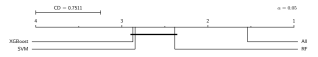


Figure 7: Critical difference diagram comparison of stacking single family of classifiers versus all families of classifiers.

### 4.4 Accuracy Benchmarking

Finally, we compare our results with the state-of-the-art and relevant approaches. Table 3 presents the classification error rates as well as the running time statistics. Specifically, datasets in this experiment are from the UEA & UCR Time Series Classification Repository thanks to the many benchmarking results that are publicly available. Note that although names of the datasets used in this repository are exactly the same compared to our previous experiments, the similarity of their contents is not guaranteed. In fact, the training and testing datasets may have been swapped for a number of datasets. An obvious example is the FordA dataset, where in this experiment the training dataset and testing set are of size 1320 and 3601 respectively, while in our previous experiments the training set has 3601 samples and the testing test has 1320.

We compare our method MVG with five state-of-the-art approaches: two distance-based global similarity matching algorithms including Euclidean- and DTW-based nearest neighbor classification (1NN-ED and 1NN-DTW), and three local pattern matching algorithms including Learning Shapelets (LS) [15], Fast Shapelets (FS) [35] and SAX-VSM [39]. Among all five approaches, LS is recognized as the most accurate classifier [42] by the research community. However, LS is also known for its high computation complexity. We first compare the classification accuracy of different approaches in this section and then investigate MVG's efficiency later in section 4.5.

Viewing the error rate comparison columns in Table 3, it is obvious that MVG is the most accurate classifier with 16 winning datasets. LS then follows MVG with 12 winning datasets. A Wilcoxon test between MVG and LS yields a *p*-value of 0.3421 > 0.05, suggesting that MVG should not be considered significantly better than LS. SAX-VSM is ranked third with 10 winning cases. The Wilcoxon test again suggest that such difference is not statistically significant. However, MVG is indeed significantly more accurate than FS, 1NN-DTW and 1NN-ED. Moreover, scatter plots in Figure 8 shows that most of the points in each comparison are located away from the diagonal line, suggesting that MVG is indeed a very different approach when compared to the state-of-the-art. Having confirmed MVG's excellent accuracy, we continue to investigate its runtime efficiency.

						C		MVG Runtime			FS			
Dataset	# Cls.	#Train	#Test	Dim.	1NN- ED	1NN- DTW	LS	FS	SAX- VSM	MVG	FE	Clf.	Σ	Runtim
ArrowHead	3	36	175	251	0.200	0.297	0.154	0.406	0.211	0.371	8	29	37	30
BeetleFly	2	20	20	512	0.250	0.300	0.200	0.300	0.100	0.050	2	21	23	54
BirdChicken	2	20	20	512	0.450	0.250	0.200	0.250	0.000	0.000	4	22	26	38
Computers	2	250	250	720	0.424	0.300	0.416	0.500	0.380	0.252	22	51	73	1293
DistalPhalanxOutlineAgeGroup	3	400	139	80	0.374	0.230	0.281	0.345	0.158	0.254	11	86	97	45
DistalPhalanxOutlineCorrect	2	600	276	80	0.283	0.283	0.221	0.250	0.272	0.281	19	93	112	101
DistalPhalanxTW	6	400	139	80	0.367	0.410	0.374	0.374	0.396	0.381	12	204	216	59
ECG5000	5	500	4500	140	0.075	0.076	0.068	0.077	0.090	0.069	162	190	352	170
Earthquakes	2	322	139	512	0.288	0.281	0.259	0.295	0.252	0.252	22	78	100	3689
ElectricDevices	7	8926	7711	96	0.448	0.398	0.413	0.421	0.295	0.332	406	6344	6750	3558
FordA	2	3601	1320	500	0.335	0.445	0.043	0.213	0.173	0.014	207	410	617	44832
FordB	2	3636	810	500	0.394	0.380	0.083	0.272	0.249	0.333	183	534	717	45874
Ham	2	109	105	431	0.400	0.533	0.333	0.352	0.190	0.343	8	32	40	670
HandOutlines	2	1000	370	2709	0.138	0.119	0.519	0.189	0.092	0.200	4431	182	4613	179745
Herring	2	64	64	512	0.484	0.469	0.375	0.469	0.375	0.344	19	30	49	286
InsectWingbeatSound	11	220	1980	256	0.438	0.645	0.394	0.511	0.453	0.459	85	130	215	705
LargeKitchenAppliances	3	375	375	720	0.507	0.205	0.299	0.440	0.123	0.288	32	89	121	7301
Meat	3	60	60	448	0.150	0.067	0.100	0.067	0.067	0.050	10	31	41	120
MiddlePhalanxOutlineAgeGroup	3	400	154	80	0.481	0.500	0.429	0.455	0.455	0.435	9	88	97	50
MiddlePhalanxOutlineCorrect	2	600	291	80	0.234	0.302	0.220	0.271	0.323	0.289	15	87	102	80
MiddlePhalanxTW	6	399	154	80	0.487	0.494	0.494	0.468	0.513	0.481	9	177	186	62
PhalangesOutlinesCorrect	2	1800	858	80	0.239	0.272	0.235	0.256	0.290	0.248	48	209	257	332
Phoneme	39	214	1896	1024	0.891	0.772	0.782	0.826	0.895	0.692	134	1419	1553	25604
ProximalPhalanxOutlineAgeGroup	3	400	205	80	0.215	0.195	0.166	0.220	0.176	0.166	11	70	81	41
ProximalPhalanxOutlineCorrect	2	600	291	80	0.192	0.216	0.151	0.196	0.172	0.144	18	80	98	80
ProximalPhalanxTW	6	400	205	80	0.293	0.239	0.224	0.298	0.390	0.220	12	160	172	49
RefrigerationDevices	3	375	375	720	0.605	0.536	0.485	0.667	0.347	0.421	37	77	114	12798
ScreenType	3	375	375	720	0.640	0.603	0.571	0.587	0.488	0.480	35	89	124	8473
ShapeletSim	2	20	180	500	0.461	0.350	0.050	0.000	0.283	0.000	6	22	28	76
ShapesAll	60	600	600	512	0.248	0.232	0.232	0.420	0.302	0.255	168	1833	2001	7939
SmallKitchenAppliances	3	375	375	720	0.656	0.357	0.336	0.667	0.421	0.208	30	75	105	5667
Strawberry	2	613	370	235	0.054	0.059	0.089	0.097	0.043	0.076	29	75	104	314
ToeSegmentation1	2	40	228	277	0.320	0.228	0.066	0.044	0.070	0.197	8	26	34	30
ToeSegmentation2	2	36	130	343	0.192	0.162	0.085	0.308	0.138	0.185	6	22	28	38
UWaveGestureLibraryAll	8	896	3582	945	0.052	0.108	0.047	0.211	0.201	0.258	470	343	813	32047
Wine	2	57	54	234	0.389	0.426	0.500	0.241	0.037	0.519	4	27	31	22
WordSynonyms	25	267	638	270	0.382	0.351	0.393	0.569	0.509	0.522	39	1017	1056	907
Worms	5	181	77	900	0.545	0.416	0.390	0.351	0.442	0.182	26	98	124	6852
WormsTwoClass	2	181	77	900	0.390	0.377	0.273	0.273	0.286	0.130	27	45	72	5044
											1	10	-	1
	Numbe	er of best (		<i>v</i> ,	1	2	12	3	10	16			24	15
		Wilcox	on test p	-value	0.0023	0.0044	0.3421	0.0005	0.5767	-	Total	time $\rightarrow$	21379	395075

Table 3: Classification error rates compared with five benchmark approaches and running time statistics (in seconds).

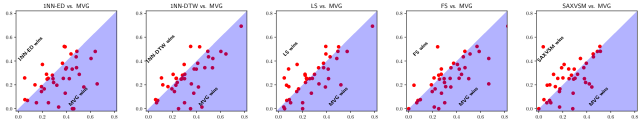


Figure 8: Comparison of classification accuracy with five state-of-the-art approaches in the form of scatter plots.

# 4.5 Efficiency

The pipeline of MVG consists of a feature extraction phase and a train-validate-test process. During the feature extraction process, time series are firstly transformed in to VGs and then features are extracted from such graphs. With optimization from [1], transforming time series of length *n* into VGs has a computation complexity of  $O(n \log^2(n))$  and it can be effectively solved within  $O(\log^2(n))$  time when taking full advantage of parallelization. Extracting motif features from VGs may be potentially expensive, fortunately PGD provides a fully parallel way of counting small motifs. For instance, counting graph cliques of size 4 – one of the most time consuming tasks in PGD – has a computation complexity of  $O(m \cdot \Delta \cdot T_{max})$ , where *m* is the number of 4-motifs

(*i.e.*, 11),  $\Delta$  is the maximum degree and  $T_{max}$  is the maximum number of triangles incident to an edge and  $T_{max} \ll \Delta$ . PGD is hundreds of times faster than other motif counting algorithms: counting motifs from a graph with more than 26,000 vertices can take only 0.01 seconds on a twelve-core commodity CPU [2]. Since other statistic features such as density, *k*-core, assortativity and degree statistics extracted from time series VGs are intentionally chosen to be simple metrics, collecting these features has a time complexity of  $O(\max(|V|, |E|))$ . Since we use a multiscale graph representation, ultimately graph generation has a monothread complexity of  $O(\max(|V|, |E|))$  and in parallel an  $O(\log^3(n))$  time complexity. As a result, feature extraction per graph has a computation complexity of  $O(\max(|V|, |E|) + m \cdot \Delta \cdot T_{max})$ . The

classification process leverages state-of-the-art classifiers that are widely used and well-optimized. Overall, the efficiency of MVG may be best illustrated when we compare our method against our benchmarking approaches.

Previous research [42] has shown that LS is extremely time consuming, and FS as an approximated approach can be 100X faster than LS. As a result, FS will be a good and strong baseline to which the running time of our approach can be compared. Thus we record the running time of FS and MVG and compare their efficiency. These experiments are conducted on a computer with Intel i7-4980HQ quad-core CPU clocked at 2.80GHz, 16GB memory and solid-state drive suitable for fast I/O. We use the FS implementation by its original authors [35] with default parameters. Columns located on the right side of Table 3 shows the running time per dataset for MVG and FS. For MVG, running time for feature extraction and training are recorded separately and then summed. Overall, MVG completes faster than FS for 24 datasets. The total running time of FS for all 39 datasets amounts to 395075 seconds, which is more than 18 times that of MVG. A scatter plot of the running time is provided in Figure 9, obviously MVG can be up to 100X faster than FS, suggesting that it is indeed an efficient TSC approach. In addition, we note that FS appears to be time consuming with large datasets with time series of high dimensionality, while the running time of MVG remains reasonable as datasets grow larger. Furthermore, since we have experimented only on a quad-core commodity computer that is not really powerful in terms of parallel processing, the efficiency of MVG can be easily boosted by adding more computing cores. Overall, MVG can indeed be considered an efficient TSC approach.

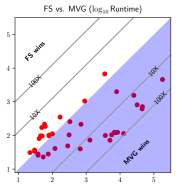


Figure 9: Runtime comparison between FS and MVG.

#### 4.6 Case Study

Although accuracy and efficiency are very important measurements for TSC approaches, it is also desirable for TSC methods to have comprehensible classification processes, so that users may gain extra insights into his/her time series data. Popular approaches based on shapelets and subsequences may have a natural advantage in classification comprehensibility. However, since features extracted in MVG are solely statistical, it can be difficult to locate exactly where a distinguishing subsequence lies in the original time series. However, for MVG it is still possible to understand which features have contributed most to correct classification. For instance, when feeding features to train an XGBoost classifier, it will assign weights to all the features. After ranking these features, we can also gain insights into the data. For instance, Figure 10 illustrates a scatter matrix plot for the test dataset of FordA showing ten most important features learned by the classifier<sup>2</sup>. Out of ten features, six are features from HVGs, which are created from the original series ( $T_0$ ). VGs features from dimensionality-reduced approximations ( $T_2$  and  $T_3$ ) are also present. Moreover, it appears that most highly ranked graph features for this dataset are MPDs and assortativity, suggesting that statistics other than MPDs are indeed helpful. Finally, visually from the kernel density estimation it is obvious that some features alone – *e.g.*,  $T_0$  HVG  $P(M4_4)$  – can already provide good classification guidelines, suggesting that it is feasible for MVG to present visually comprehensible cues regarding its classification decisions.

#### 4.7 Discussions

Due to specific characteristics of VGs, obviously MVG may not be suitable for every TSC scenarios. For instance, VGs are agnostic of affine transformations in time series. That is, in applications where the absolute oscillation is more important, MVG is less likely to detect such characteristics. Furthermore, when dealing with non-stationary time series data where trends are very frequently found, MVG works best when trends are not the deciding factor since VGs may not be able to capture long-term monolithic trends. Since many graph features have been taken into account in MVG, we believe it can be robust in practical applications.

In addition to limitations inherited from VGs, our approach is more suitable for applications where the size of training datasets is larger, so that classification models can generalize better. In fact, when we review the classification accuracy of MVG, it seems that a majority of its winning datasets have either long time series or large training datasets. This is perhaps innate to the nature of statistics: sample size has to be sufficiently large to make accurate estimations. Based on running time comparison, MVG also scales well with large datasets, which can be an important advantage in the era of big data.

### 5 RELATED WORK

TSC is a major task in time series mining thanks to its wide application scenarios. As a consequence, there are a plethora of classification algorithms for TSC. Classical TSC approaches involve utilizing 1NN classification together with similarity measures specific to time series data, *e.g.*, DTW [7] and its variants with lower bounding [36] and early abandoning techniques [34].

Another line of research transforms time series into texts and resort to text classification algorithms. For example, inspired by the well known bag-of-words approach, [31] proposes the Bag-of-Patterns approach for TSC. SAX-VSM [39] also takes advantage of bag-of-words approach and builds term frequency-inverse document frequency (TF-IDF) vectors in its training phase. It defines a similarity measure of two vectors (that are constructed from original series) based on their inner product. Both Bag-of-Patterns and SAX-VSM rely on SAX [30] for transforming time series into texts. A more recent work, Representative Pattern Mining (RPM) [42], also tries to classify time series by means of finding the most representative SAX-symbolized subsequences. Our previous work Domain Series Corpus (DSCo) [24, 29] also transforms time series into texts and approximates TSC to a pseudo language detection problem. [38] invents another symbolic representation based on Fourier transform and proposes a bag-of-patterns approach named BOSS ensemble based on this symbolic representation method.

 $<sup>^2\</sup>text{To}$  save space, we show more illustrations comparing original time series and important MVG features on project website. URL: http://daoyuan.li/mvg/

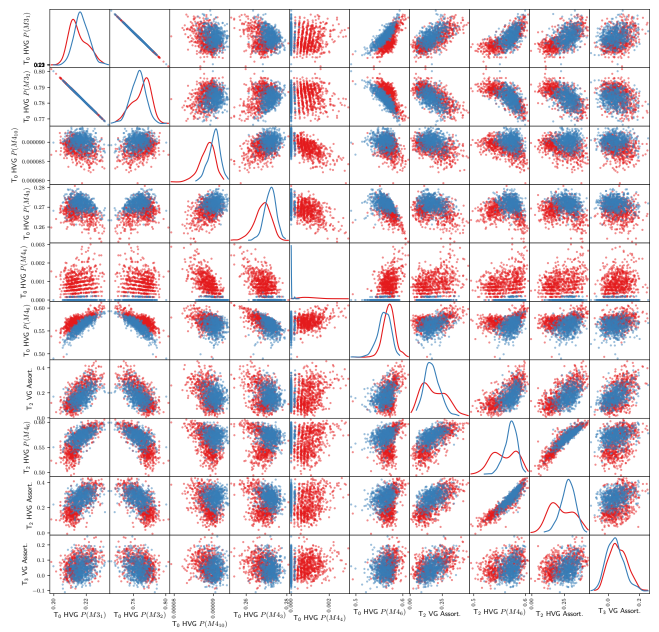


Figure 10: Scatter matrix of ten most important features for FordA's test dataset. Different point colors indicate different classes and the diagonal shows the Gaussian kernel density estimation for each feature.

Recently, shapelet-based approaches are gaining popularity among the research community. A shapelet [47] is a single subsequence in time series that is representative of its class. However, these approaches [15, 16] are known for their high computation complexity. As we have demonstrated in this paper, FS [35] as an approximated approach can also take long time to run. High computation complexity is also present for TSC approaches using deep neural networks [46]. Finally, [4] introduce an ensemble algorithm named Collective of Transformation-based Ensembles (COTE) – an ensemble of 35 different classifiers – that has shown to be very accurate but has a time complexity of  $O(m^4n^2)$ , where *m* is the dimensionality of time series and *n* is the size of training dataset.

Although the concept of time series VGs has appeared for almost a decade, using it for TSC has just started picking up. Machine learning approaches taking advantage of VG and its variants are generally using artificially generated data [45] or EEG signals. Finally, graph kernel methods [20] can be used for evaluating graph similarity, which may potentially be used for TSC as well. Our approach transforms TSC into a graph classification [14] problem. Since we convert time series into a set of graphs at different scales, we have not experimented generic graph classification algorithms. However, they are indeed interesting for future experiments.

# 6 CONCLUSIONS AND FUTURE WORK

This paper proposes a graph-based multiscale time series representation named MVG and a feature extraction method for TSC. MVG transforms time series into a collection of visibility graphs of various scales and feature extraction are conducted by investigating statistical graph features such as probability distributions of small motifs, assortativity as well as degree statistics. After a large scale evaluation with open datasets and comparison with a number of state-of-the-art TSC algorithms, our proposed approach appears to be both accurate and efficient: it can be more accurate than LS and up to  $\sim$ 100X faster than FS.

In the future, we plan to further investigate other useful and efficient graph features – such as degree distribution entropy, centrality, bipartivity, *etc.* [11] – for MVG in order to further improve its accuracy. Currently we have only evaluated MVG with univariate time series data, we are also excited to investigate the possibility of adopting MVG for multivariate TSC.

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