Barycentre Averaging for the Move-Split-Merge Time Series Distance Measure

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Keywords: Time Series Distances, Time Series Clustering, Move Split Merge, Barycentre Averaging, Dynamic Barycentre Averaging, DBA.

Abstract: Distance functions play a core role in many time series machine learning algorithms for tasks such as clustering, classification and regression. Time series often require bespoke distance functions because small offsets in time can lead to large distances between series that are conceptually similar. Elastic distances compensate for misalignment by creating a path through a cost matrix by warping and/or editing time series. Time series are most commonly clustered with partitional algorithms such as $k$-means and $k$-medoids using elastic distance measures such as Dynamic Time Warping (DTW). The distance is used to assign cases to the closest cluster representative. $k$-means requires the averaging of time series to find these representative centroids. If DTW is used to assign membership, but the arithmetic mean is used to find centroids, $k$-means performance degrades significantly. An averaging technique specific to DTW, called DTW Barycentre Averaging (DBA), overcomes the averaging problem however, can only be used with DTW. As such alternative distance functions such as Move-Split-Merge (MSM) are forced to use the arithmetic mean to compute new centroids and suffer similar degraded performance as $k$-means-DTW without DBA. To address this we propose a averaging method for MSM distance, MSM Barycentre Averaging (MBA) and show that when used to find centroids it significantly improves MSM based $k$-means and is better than commonly used alternatives.

1 INTRODUCTION

Machine learning applied to time series data is an extensively studied area within the literature. One of the key factors contributing to its advancement is the improved ability to collect temporal data using cost-effective sensor technology across various scientific disciplines. Specifically, time series data can be subjected to different tasks, each with its own objective. Among these tasks, Time Series Classification (TSC) (Bagnall et al., 2017; Middlehurst et al., 2023) has gained widespread attention, involving the prediction of a discrete output value for a given time series. Furthermore, if the aim is to predict a continuous output value instead of a discrete one, the task is referred to as Time Series Extrinsic Regression (TSER) (Tan et al., 2021; Guijo-Rubio et al., 2023). Concerning unsupervised tasks, Time Series Clustering (TSCL) (Liao, 2005; Aghabozorgi et al., 2015) is particularly well-known, aiming to group time series without the need of labelled data. TSCL is the primary focus of this work.

TSCL is used across multiple different disciplines. It has been used to identify anomalous amplitude and shape patterns in disease outbreak data (Li et al., 2021) Climate researchers used TSCL to discover common patterns preceding important paleoclimate events (Nikolaou et al., 2015). TSCL has been used extensively in bioinformatics to group gene expression patterns (McDowell et al., 2018). These applications of TSCL demonstrate the versatility of this task across several disciplines.

With respect the TSCL taxonomy, it can be approached from two main points of view: 1) extracting features from the time series; and 2) employing a standard clustering approach along with a time series distance measure. The majority of existing TSCL techniques fall into the second category. These ap-
approaches critically depend on the choice of an appropriate distance measure, combined with the clustering approach used.

Measuring distance between time series is a primitive operation that can be used for a range of tasks such as classification, clustering, regression and query. Time series require bespoke distance functions because small offsets between series can lead to large distances between series that are conceptually similar. Elastic distances compensate for misalignment by creating a path through a cost matrix through either warping or editing time series. The most common elastic distance is Dynamic Time Warping (DTW) (Ratanamahatana and Keogh, 2005), however, there have been numerous others proposed.

A recent comparison study that evaluated 9 commonly used elastic distances found that the MSM distance measure was the best performing distance function for the k-means clustering algorithm (Holder et al., 2022). Unlike DTW, MSM satisfies the mathematical conditions of a metric, meaning it can, for example, exploit the triangle inequality for fast distance calculations. Figure 1 shows the method for finding the MSM distance between two series.

In addition to computing similarity, many TSCL approaches must also choose or synthesise exemplars to represent clusters. One of the most common techniques to do this is averaging. k-means (Lloyd, 1982) is one of the most commonly used TSCL approaches in the literature and requires both to compute the distance between time series and the creation of synthetic time series (by averaging) that represent a cluster. Various methods have been proposed to compute the average of a collection of time series (Brill et al., 2019; Holznigenkemper and Seeger, 2023).

Our contribution is to propose a method for averaging time series using MSM. Our method, MSM Barycentre Averaging (MBA) is based on the approximate barycentre averaging method which was made popular for DTW using DTW Barycentre Averaging (DBA) (Petitjean et al., 2011). We show that, when integrated with k-means clustering, MBA significantly outperforms clusters created using either DTW or MSM with arithmetic mean averaging and DBA.

We have conducted all experiments with the aeon time series machine learning toolkit1 and we demonstrate how to reproduce all our experiments with the associated experimental code and notebook2.

The rest of this paper is structured as follows. In Section 2 we provide background into elastic distance functions and time series averaging. Section 2.3 describes our approach to averaging using MSM, known as MSM Barycentre Averaging (MBA). In Section 3.1, we present related works in TSCL literature. Our results are presented in Section 4 before we conclude in Section 5.

2 BACKGROUND

Distance-based time series machine learning has been a popular theme in time series classification and clustering research. There have been numerous experimental evaluations of distance-based classification, such as (Ding et al., 2008; Lines and Bagnall, 2014). For many years, the received wisdom was that DTW was the best choice. For example, the first sentence of (Petitjean et al., 2016) is: “The last decade has seen increasing acceptance that the nearest neighbour algorithm with dynamic time warping as the distance measure is the technique of choice for most time series classification problems”. However, recent experimental papers (Lines et al., 2018; Paparrizos et al., 2020; Holder et al., 2022) have identified that MSM is more effective for both classification and clustering. Nevertheless, DTW is still by far the most widely used elastic distance measure. Hence, we limit our focus to DTW and MSM, as well as the standard Euclidean Distance (ED), and direct the interested reader to (Holder et al., 2022; Shifaz et al., 2023) for more detailed background on elastic distances.

2.1 Time Series Elastic Distance Functions

Suppose we want to measure the distance between two time series (assumed to be equal lengths and univariate), \( \mathbf{a} = \{a_1, a_2, \ldots, a_m\} \) and \( \mathbf{b} = \{b_1, b_2, \ldots, b_m\} \). The ED, \( d_{ED} \) is the L2 norm between series,

\[
d_{ED}(\mathbf{a}, \mathbf{b}) = \sqrt{\sum_{i=1}^{m} (a_i - b_i)^2}.
\]

\( d_{ED} \) puts no priority on the ordering of the series. Elastic distance measures allow for possible misalignment by attempting to optimally align two series. This is done by either distorting indices or by editing the series to add or remove values.

2.1.1 Dynamic Time Warping (DTW)

DTW mitigates distortions in the time axis by realigning (also known as warping) the series to best
match each other. Let \( M(a, b) \) be the \( m \times m \) pointwise distance matrix between series \( a \) and \( b \), where \( M_{i,j} = (a_i - b_j)^2 \). A warping path
\[
P = (e_1, f_1)(e_2, f_2), \ldots, (e_s, f_s)
\]
is a set of pairs of indices that define a traversal of matrix \( M \). A valid warping path must start at location \((1, 1)\), end at point \((m, m)\) and not backtrack, i.e. \(0 \leq e_{i+1} - e_i \leq 1\) and \(0 \leq f_{i+1} - f_i \leq 1\) for all \(1 < i < m\). The DTW distance between series is the path through \( M \) that minimises the total distance. The distance for any path \( P \) of length \( s \) is
\[
D_P(a, b, M) = \sum_{i=1}^{s} M_{e_i, f_i}. 
\]
(2)

If \( \mathcal{P} \) is the space of all possible paths, the DTW path \( P^* \) is the path that has the minimum distance, hence the DTW distance between series is
\[
d_{DTW}(a, b) = D_{P^*}(a, b, M). 
\]
(3)

The optimal warping path \( P^* \) can be found exactly through the dynamic programming formulation described in Algorithm 1. This can be a time-consuming operation, and it is common to put a restriction on the amount of warping allowed.

2.1.2 Move-Split-Merge (MSM)

At any step, elastic distances can use one of three costs: diagonal, horizontal or vertical, in forming an alignment. The alignment path is a series of moves across the cost matrix. DTW assigns no explicit penalty for moving off the diagonal. Instead, it uses an implicit penalty (long paths have longer total distance) and a hard cut off on window size to stop large warpings. An alternative family of distance functions are based on the concept of edit distance (e.g. Edit distance with Real Penalty (ERP) (Chen and Ng, 2004)). An edit distance, such as MSM, considers a diagonal move as a match, a vertical move as an insertion and an horizontal move as a deletion. MSM (Algorithm 2) follows this structure, where move is a match (diagonal), split is an insertion (vertical) and merge is deletion (horizontal).

Algorithm 1: DTW \((a, b)\) (both series of length \(m\), \(w\) (window proportion), default value \(w \leftarrow 1\), \(M\) (pointwise distance matrix)).

1. Let \( C \) be an \((m + 1) \times (m + 1)\) matrix initialised to zero, indexed from zero.
2. for \( i \leftarrow 1 \) to \( m \) do
3.     for \( j \leftarrow 1 \) to \( m \) do
4.         if \(|i - j| < w \cdot m\) then
5.             \( C_{i,j} \leftarrow M_{i,j} + \min(C_{i-1,j-1}, C_{i-1,j}, C_{i,j-1}) \)
6.     return \( C_{m,m} \)

Figure 1 shows the method for finding the MSM distance between two time series. As can be ob-
served, the three operations (match/insert/deletion) are identified with different colours (blue/green/red) and using the specific terminology of MSM distance (move/split/merge).

The move operation in MSM uses the absolute difference rather than the squared euclidean distance for matching in DTW. The cost of the split operation is given by cost function $C$ (Equation 4) with a call to $C(a_i, a_{i-1}, b_i, c)$. If the value being inserted, $b_i$, is between the two values $a_i$ and $a_{i-1}$ being split, the cost is a constant value $c$. If not, the cost is $c$ plus the minimum deviation from the furthest point $a_i$ and the previous point $a_{i-1}$ or $b_i$. The delete/merge is given by $C(b_l, b_{l-1}, a_i, c)$, which is simply the same operation as split but applied to the second series. Thus, the cost of splitting and merging values depends on the value itself and adjacent values.

$$C(a_i, a_{i-1}, b_j, c) = \begin{cases} c & \text{if } a_{i-1} \leq a_i \leq b_j \\ c & \text{if } a_{i-1} \geq a_i \geq b_j \\ c + \min(|a_i - a_{i-1}|, |a_i - b_j|) & \text{otherwise.} \end{cases}$$

(4)

Algorithm 2 describes how to calculate the MSM distance between two time series $a$ and $b$. MSM satisfies triangular inequality and is a metric. In Algorithm 2 the first return value is the MSM distance between $a$ and $b$, the second is the cost matrix used to compute the MSM distance (this is used in Algorithm 5).

**Algorithm 2: MSM(a (of length m), b (of length m), c (minimum cost)).**

1. Let $CM$ be an $m \times m$ matrix initialised to zero.
2. $CM_{1,1} = |a_1 - b_1|$
3. for $i \leftarrow 2$ to $m$ do
   4. $CM_{i,1} = CM_{i-1,1} + C(a_i, a_{i-1}, b_1, c)$
5. for $i \leftarrow 2$ to $m$ do
   6. $CM_{1,j} = CM_{1,j-1} + C(b_j, a_1, b + i - 1, c)$
7. for $i \leftarrow 2$ to $m$ do
   8. for $j \leftarrow 2$ to $m$ do
      9. move $\leftarrow CM_{i-1,j-1} + |a_i - b_j|$
     10. split $\leftarrow CM_{i-1,j} + C(a_i, a_{i-1}, b_j, c)$
     11. merge $\leftarrow CM_{i,j-1} + C(b_j, b_{j-1}, a_i, c)$
     12. $CM_{i,j} \leftarrow \min($move$,$split$,$merge$)
9. return $CM_{m,m}, CM$

**2.2 Time Series Averaging Methods**

Finding a consensus (average) representation of a set of sequences has been described as the Holy Grail by (Gusfield, 1997). One of the main issues with averaging sequences is that it is subjective. As such to develop an algorithm to solve this problem it is normally formulated as an optimisation problem (Brill et al., 2019; Holznigenkemper and Seeger, 2023). If we are not concerned with offset, then the simple arithmetic average over time points will minimise the ED between time series. For algorithms such as $k$-means this is the default approach. However, when using this approach with elastic distances (i.e. cluster membership is assigned based on an elastic distance measure), the arithmetic mean centroid may misrepresent the elements of a cluster. Simple averaging will tend to blur the underlying series and result in worse $k$-means clustering performance (Petitjean et al., 2016). Similarly, if series of the same class are condensed through arithmetic averaging, it is unlikely these exemplars will be useful for classification or regression.

As such many methods to average time series using DTW have been proposed. Initially NonLinear Alignment and Averaging Filters (NLAAF) (Gupta et al., 1996) was proposed. This method applies a tournament scheme whereby sequences are paired and averaged together step by step until only one final sequence remains. When two sequences are averaged, the DTW alignment path between the two series is computed and the point to point average is taken using this alignment path. The main drawback of this approach is it leads to a large growth in the average sequence produced (as every use of the averaging method can lead to the length of the sequence almost doubling) (Petitjean et al., 2011). Another issue with NLAAF is error propagation. Niennattrakul and Ratanamahatana (2009) proposed Prioritised Shape Averaging (PSA) which employs a hierarchical averaging method reducing error propagation but also leading to the same large sequence length growth as NLAAF. Due to this growth of sequence, both are impractical for time series data (given complexities of algorithms employed, such as DTW). Petitjean et al. (2011) addresses both of these problems by proposing DTW Barycentre Averaging (DBA) that uses the optimal warping path to compute a series of the same length while accounting for alignment of time series. This will be explained further in Section 2.3.

**2.3 Barycentre Averaging**

DBA (Petitjean et al., 2011) was proposed to overcome the limitations of other averaging methods for time series outlined in Section 2.2. DBA uses a heuristic strategy to compute a new series that minimises the DTW distance to cluster members rather than the ED. The DBA process begins with an initial centre, which is typically the medoids of the time series collection to be averaged. For each time series
in the collection, the optimal DTW warping path is computed to the centre. Using these warping paths a new centre is computed by determining which points warped onto each element of the initial centre. The mean is then calculated for each time point warped to each other time point, taking into account how often each time point was warped to. This process continues iteratively until there is no significant change in the sum of squared DTW distances to the centre, indicating that the optimum centre has been achieved.

While the original proposal for DBA was only for DTW, assuming an optimal alignment path can be obtained from an elastic distance (and it aims to minimise the dissimilarity between two series), any distance could be used with (Petitjean et al., 2011) barycentre averaging approach. Holder et al. (2022) reviewed 9 different elastic distance measures for TSCL and found the MSM distance significantly outperformed DTW across multiple clustering metrics. As such this paper seeks to expand on these finding by also using the best performing elastic distance in the averaging stage of computation.

3 MSM BARYCENTRE AVERAGE (MBA)

To change the distance measure used in the original DBA, the optimal warping path needs to be retrieved from the distance computation, necessitating the MSM algorithm to construct and return a cost matrix, as seen in Algorithm 2. This optimal path is identified by backtracking through the cost matrix and following the trajectory that minimises the total distance, as outlined in Algorithm 3. Given these traits, MSM can feasibly be paired with barycentre averaging.

With this prerequisites the DBA algorithm has been adapted and outlined in Algorithm 4 and 5. This new algorithm computes the MSM Barycentre Average (MBA). Each step in the MBA algorithm will now be outlined.

\[
MSM_{medoids} = \arg \min_{x_m \in C} \sum_{x_c \in C} MSM(x_c, x_m, \text{cost}).
\]

(5)

Algorithm 4 takes a collection of time series, a number of max iterations and a minimum cost for MSM as parameters and computes the MBA. Firstly an initial centre is computed (line 1). This is done using the MSM medoids given in Eq 5. Using this initial centre, a number of iterations are executed (max_iters) to refine the centre (line 3). This is done using Algorithm 5. Algorithm 5 begins by defining two values: num_warsps_\text{to}, which is an array that tracks how many times a point is warped to (due to how the optimal warping path is computed one point can be warped to multiple times in a single path), and

<table>
<thead>
<tr>
<th>Algorithm 3: Compute_path(CM (of size (n \times m)).)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1 Let alignment be a list.</td>
</tr>
<tr>
<td>2 Let (i = n - 1) and (j = m - 1).</td>
</tr>
<tr>
<td>3 while (i &gt; 0) or (j &gt; 0) do</td>
</tr>
<tr>
<td>4 Append ((i, j)) to alignment.</td>
</tr>
<tr>
<td>5 if (i == 0) then</td>
</tr>
<tr>
<td>6 (j \leftarrow j - 1).</td>
</tr>
<tr>
<td>7 else if (j == 0) then</td>
</tr>
<tr>
<td>8 (i \leftarrow i - 1).</td>
</tr>
<tr>
<td>9 else</td>
</tr>
<tr>
<td>10 (\min_{\text{index}} = \arg \min((CM_{i-1,j-1}, CM_{i-1,j}, CM_{i,j-1}))</td>
</tr>
<tr>
<td>11 if (\min_{\text{index}} == 0) then</td>
</tr>
<tr>
<td>12 (i \leftarrow i - 1, j \leftarrow j - 1).</td>
</tr>
<tr>
<td>13 else if (\min_{\text{index}} == 1) then</td>
</tr>
<tr>
<td>14 (i \leftarrow i - 1).</td>
</tr>
<tr>
<td>15 else</td>
</tr>
<tr>
<td>16 (j \leftarrow j - 1).</td>
</tr>
<tr>
<td>17 Append ((0, 0)) to alignment.</td>
</tr>
<tr>
<td>18 return alignment reversed.</td>
</tr>
</tbody>
</table>

(continued)
alignment, which is an array of length $m$ which will store the values of the new centre (line 1 and 2). Next, for each value in the collection of time series $X$, it computes the MSM Cost Matrix ($CM$) (Algorithm 2) between $X_i$ and the centre (line 4). Using the computed $CM$, the optimal alignment path can be found in the form (line 5):

$$curr\_alignment = < (e_1, f_1), (e_2, f_2), \ldots, (e_s, f_s) > .$$

Each tuple in the optimal alignment path is then looped over (line 6) and the $j^{th}$ value in $X_j$ is added to the value in $alignment_k$ (i.e., value $j$ is warped to the index $k$) (line 7). Furthermore $num\_warps_j$ is incremented (line 8). Once each time series in the collection $X$ has been warped to, the new centre can be computed by taking the mean of the alignments using $num\_warps_j$, which tracked the number of time each index was warped to (line 9). The new computed centre is then returned (line 10).

The result of this adaptation, while subtle, leads to a much different resulting average as shown in Figure 2. This figure shows the average time series of the whole set of time series with class 1 for the GunPoint dataset. The discriminatory features are the small peaks before and after the main peak. These represent the moment when the actor is drawing and replacing the gun. As can be observed, MSM is able to clearly identify these discriminatory features, whereas both arithmetic averaging or DTW do not.

### 3.1 Alternative TSCL Algorithms

To put the results of MBA into context we also explore alternative TSCL approaches. Most of the developments in TSCL are domain-specific, not being focused on TSCL as a whole. Nevertheless, there are a few approaches tackling TSCL as a whole. The first one is the U-shapelets technique (Zakaria et al., 2012), which, instead of computing pairwise distances, considers only relevant subsequences of time series. This technique shares similarities with its classification counterpart (Hills et al., 2014). First of all, subsequences are extracted from the data and ranked based on their utility, which reflects their discriminative power. This value attempts to maximise the separation gap between two subsets of time series: one subset comprises time series with subsequences similar to the shapelet being evaluated, while the other subset consists of the remaining time series. The subsequences with high utility, referred to as shapelets, are retained. Once the final set of shapelets is achieved, a transformation matrix is built with cells representing distances between U-shapelets and time series. Finally, the standard k-means method is applied to the transformation matrix. The main advantages of this approach are that U-shapelets mitigates the sensitivity to noise and other irrelevant data, and their ability to provide additional insights into the data.

Another well-known approach is the Two-step Time series Clustering (TTC) (Aghabozorgi et al., 2014). This method firstly reduces the size of the dataset using the concept of affinity. For this, time series are grouped according to similarity in time and then applying an affinity search technique. Subsequently, for each cluster a prototype is defined according to the affinity of the time series belonging to it. The second step of this approach involves computing the DTW distances between the subclusters prototypes. This distance measurement aims to represent the dissimilarity between the subgroups, in such a way that the complexity is reduced as much as possible. Finally, similar subclusters are merged by means of the $k$-medoids standard clustering method.

In addition, one of the approaches in the state-of-the-art of TSCL is the well-known $k$-shapes (Paparrizos and Gravano, 2015). It is a partitioning clustering algorithm that aims to create homogeneous and well-separated clusters through an iterative process. In a similar way to $k$-means, $k$-shapes also performs two main processes: 1) the assignment step; and 2) the refinement step. For the first one, $k$-shapes employs an efficient adaptation of the cross-correlation measure known as Shape-Based Distance (SBD). In the refinement step, the centroids of the clusters are recomputed by solving an optimisation problem that minimises (or maximises) the sum of squared distances (or squared similarities) to all the time series, found significantly better than computing the average time series. A key advantage of $k$-shapes is that it groups time series based on their shape similarity, regardless of differences in amplitude and phase. Thus, it preserves the shapes of the time series while measuring the distance between them.

Finally, a range of deep learning approaches have been recently compared and analysed in (Lafabregue et al., 2022). This study represents the first exploration of deep learning techniques in TSCL. Hence, three components have been studied: architecture, clustering loss, and pretext loss. Through separate assessments of each component it has been determined that a simple autoencoder architecture using a reconstruction-based pretext loss is the best combination. Interestingly, the results also indicated that the incorporation of clustering losses did not lead to a performance increase. Therefore, its addition is not justified. Finally, authors discussed that more research is required for improving the performance of these ap-
4 COMPARISON OF OUR METHOD

This section details the experimental settings used as well as the results obtained, highlighting some of the most important aspects to be analysed.

4.1 Experimental Settings

We compare MBA to alternative approaches using the 112 univariate, equal-length time series in the UCR archive (Dau et al., 2019) for clustering. We use the provided default train/test splits for all experiments. We z-normalise all datasets prior to clustering. In addition each model takes a value of $k$ as a parameter. We set the value of $k$ equal to the number of unique class labels for each dataset. To measure performance, we use three measures of the cluster labels: 

- **CLustering ACCuracy (CL-ACC)**, is calculated by dividing the number of correct predictions by the total number of cases, similar to classification accuracy. To do this, once each value has been assigned to a cluster the maximum accuracy from every permutation of cluster and class value is taken. The cluster that achieved the highest accuracy for each label is then assigned that ground truth label (this is done using the Hungarian algorithm).

- The Rand Index (RI) works by measuring the similarity between two sets of labels such as the predicted and actual class values. The RI is the number of pairs that agree on a label divided by the total number of pairs. One of the limiting factors of RI is that the score is inflated, especially when the number of clusters is high. The Adjusted Rand Index (ARI) compensates for this by adjusting the RI based on the expected scores on a purely random model.

- The Mutual Information (MI) score, is a function that measures the agreement of the two clusterings or a clustering and a true labelling, based on entropy. Normalised Mutual Information (NMI) rescales MI onto $[0, 1]$.

We compare the performance of the following 10 clustering algorithms:

1. **$k$-means-DBA** and **$k$-means-MBA**: $k$-means clustering with DTW/MSM barycentre averaging (i.e. DBA/MBA respectively), and DTW/MSM distance assignment.

2. **$k$-means-ED**, **$k$-means-DTW** and **$k$-means-MSM**: $k$-means clustering with arithmetic mean for centres, and ED, DTW and MSM distance assignment, respectively.

3. **$k$-shapes** (Paparrizos and Gravano, 2015).

4. **TTC**: Two-step Time series Clustering (TTC) (Aghabozorgi and Wah, 2014)

5. **$k$-medoids-ED**, **$k$-medoids-DTW** and **$k$-medoids-MSM**: $k$-medoids clustering with
ED, DTW and MSM distance assignment, respectively. 
k-means, k-medoids and k-shapes at their core are variations of Lloyds algorithm (Lloyd, 1982) and so share many of the same parameters. As such this means by keeping many of the same parameters constant better insight can be gained when comparing metric performance. The values used in our experiment for Lloyds based models are given in table 1.

Max_iters is a maximum number of iterations the algorithm can run before forceful termination. This limit will only ever be reached if the algorithm does not converge (i.e. an iterations cluster values do not change compared to the previous iteration). Our experiments found many of the algorithm converged (the cluster values did not change between iterations) in under 20 iterations and so the limit of 300 is set for redundancy. The init_algo is the initialisation algorithm used to select the initial centres. Our experiment uses the most common initialisation technique: random initialisation (MacQueen et al., 1967). This technique consists of choosing the initial centres randomly from the dataset. The rationale behind this is that random selection is likely to pick points from dense regions. Rerunning the model multiple times with random initialisation and taking the best clustering (as measured by the sum of distances to their closest cluster centres) is the most common way of initialising k-means (Bradley and Fayyad, 1998). The number of reruns is defined by n_init which for our experiment is set to 10 as this is the most common value used in other similar experiments and is the default value for scikit-learn’s k-means clusterer.

The metric parameter is our first independent variable. Different metrics have been outlined in Sections 2.1 and 2.3. Finally the centroid computation defines the technique used to compute a new cluster centre from a collection of time series. MBA, DBA and arithmetic mean have been defined in Section 2.3. MSM medoids is given in Eq 5 and DTW medoids is similar but instead of using the MSM distance in Eq 5 the DTW distance is employed. Finally shape extraction is a shape based averaging technique using SBD.

Results are expressed using an adaptation of the critical difference diagram (Demšar, 2006), replacing the post-hoc Nemenyi test with a comparison of all classifiers using pairwise Wilcoxon signed-rank tests, and cliques formed using the Holm correction (García and Herrera, 2008; Benavoli et al., 2016).

Table 1: Lloyds based algorithm variation parameters. For all models max_iters = 300, n_init=10 and init_algo="random".

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Metric</th>
<th>Centroid Computation</th>
</tr>
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<tbody>
<tr>
<td>k-means-MBA</td>
<td>MSM</td>
<td>MBA</td>
</tr>
<tr>
<td>k-means-MSM</td>
<td>MSM</td>
<td>arithmetic mean</td>
</tr>
<tr>
<td>k-means-DBA</td>
<td>DTW</td>
<td>DBA</td>
</tr>
<tr>
<td>k-means-DTW</td>
<td>DTW</td>
<td>arithmetic mean</td>
</tr>
<tr>
<td>k-means-ED</td>
<td>ED</td>
<td>arithmetic mean</td>
</tr>
<tr>
<td>k-shapes</td>
<td>SBD</td>
<td>SE</td>
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<tr>
<td>k-medoids-MSM</td>
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<td>MSM medoid</td>
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<tr>
<td>k-medoids-DTW</td>
<td>DTW</td>
<td>DTW medoid</td>
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</table>

4.2 k-means Variants

Figures 3, 4 and 5 show the average ranks of the five k-means clusters against our two benchmark algorithms, k-shapes and TTC, for CL-ACC, ARI and NMI. We had to exclude the HandOutlines dataset and hence reduce the number of datasets in our study to 111, given the computational time required by this dataset. The pattern of performance is the same for the three measures: k-means-MBA is significantly better than the other six algorithms. This is our primary support for using k-means-MBA. There is some consistency in the clique membership. k-means-MSM and k-means-DBA are always in the same clique and TTC is significantly better than k-shapes. k-means-MSM (k-means with MSM using arithmetic averaging to find centroids) is not significantly different to k-means-DBA (k-means with DTW and DTW barycentre averaging). Moreover, k-means-DTW (k-means with DTW and arithmetic averaging) is the worst performing algorithm and is significantly worse than k-means-ED (k-means with ED and arithmetic averaging) (confirming results presented in (Holder et al., 2022)).

Figure 6 shows the scatter plot in terms of CL-ACC of k-means-MBA against k-means-MSM. It demonstrates the improvement provided by using MBA.

Table 2 quantifies the summary performance statistics of these clusterers. Using MBA increase MSM based k-means by approximately 2% for all three metrics. The improvement does come at a cost: both DBA and MBA take much longer than arithmetic averaging. We run our experiments on a shared computing cluster in parallel, but we can say that whilst k-means-MSM and k-means-DTW take on average a few minutes per problem, k-means-DBA and MBA take on average over an hour.

Deep learning results presented in (Lafabregue et al., 2022) are available from the associated web-
Table 2: Summary performance measures for k-means based clustering.

<table>
<thead>
<tr>
<th></th>
<th>CL-ACC</th>
<th>ARI</th>
<th>NMI</th>
</tr>
</thead>
<tbody>
<tr>
<td>k-means-MBA</td>
<td>55.97%</td>
<td>23.91%</td>
<td>32.36%</td>
</tr>
<tr>
<td>k-means-MSM</td>
<td>54.07%</td>
<td>21.75%</td>
<td>29.87%</td>
</tr>
<tr>
<td>TTC</td>
<td>53.70%</td>
<td>22.38%</td>
<td>30.87%</td>
</tr>
<tr>
<td>k-means-DBA</td>
<td>53.96%</td>
<td>20.63%</td>
<td>29.86%</td>
</tr>
<tr>
<td>k-means-ED</td>
<td>51.59%</td>
<td>18.64%</td>
<td>27.31%</td>
</tr>
<tr>
<td>k-shapes</td>
<td>47.81%</td>
<td>11.24%</td>
<td>21.57%</td>
</tr>
<tr>
<td>k-means-DTW</td>
<td>48.95%</td>
<td>16.22%</td>
<td>23.30%</td>
</tr>
</tbody>
</table>

3.1081 k-means-MBA
3.6036 k-means-MSM
3.7568 TTC
4.0586 k-means-DBA
4.1667 k-means-ED
4.3063 k-shapes
5k-means-DTW

The alternative to using an elastic averaging technique with k-means is to use k-medoids, which trade extra memory to require fewer averaging operations. This is because they require a precomputed pairwise distance matrix. This is not needed in k-means as only distances to the generated centres each iteration are needed. Holder et al. (2022) found that k-medoids based on Lloyds algorithm was significantly better than k-means for seven elastic distance measures. If we use k-means-MBA, then the performance difference is removed and on NMI and ARI measures, k-means-MBA significantly outperforms k-medoids-MSM. Figures 7, 8 and 9 show the ranks by NMI, accuracy and ARI of five k-means variants against three k-medoids clusterers. The scatter plot of k-means-MBA against k-medoids-MSM in terms of ARI is shown in Figure 10.

As illustrated in Figures 7, 8, and 9, the choice of elastic distance measure plays a critical role in the performance of both k-means and k-medoids algorithms. However, when we hold the distance metric constant and solely vary the method for computing the cluster center - comparing k-medoids-MSM against k-means-MBA we find that MBA significantly enhances the quality of clustering across all evaluation metrics. In addition this is true against the arithmetic mean used in k-means- msm.

5 CONCLUSION

Distance functions play a crucial role in time series machine learning, particularly with clustering where it is commonly still used. A recent study found k-medoids more effective than standard k-means on the UCR data (Holder et al., 2022). However, k-medoids has the problem of always requiring a complete distance matrix, and is usually slower than k-means. The

Aghabozorgi, S. and Wah, T. Y. (2014). Clustering of

Figure 7: Average ranks and cliques for CL-ACC for five k-means and three k-medoids methods.

Figure 8: Average ranks and cliques for NMI for five k-means and three k-medoids methods.

Figure 9: Average ranks and cliques for ARI for five k-means and three k-medoids methods.

Figure 10: ARI scatter plot of k-means-MBA against k-medoids-MSM. The yellow points show k-means-MBA was better and the blue points show where k-medoids-MSM was better.

problem with standard k-means is that arithmetic averaging in the step to find centroids loses the elastic information. A solution for k-means-DTW based on barycentre averaging was proposed in (Petitjean et al., 2011), known as DTW Barycentre Averaging (DBA). However, it was also shown in (Holder et al., 2022) that DTW is less effective at finding good clusterings than alternative elastic distances. We have adapted the Move-Split-Merge (MSM) distance (Stefan et al., 2013) to be used with k-means barycentre averaging. We have shown (see Table 2, Figures 7, 9, 8) that using MSM with Barycentre Averaging (MBA) significantly improves the results of k-means. k-means-MBA is also significantly better than the popular k-shapes and Two-step Time series Clustering (TTC) algorithms, and similar to the best deep learning approach found through experimenting with over 300 models (see end of Section 4.2). We believe k-means-MBA offers a good alternative to k-medoids-MSM for Time Series Clustering (TSCL).

In future work we would seek to improve the time complexity of MBA by employing a techniques such as a bounding windows for MBA. Additionally we have set out a framework to adapt other elastic distances for barycentre averaging. This could lead to experimentation using other distances such as Time Warp Edit (TWE) (Marteau, 2009) which achieved similar performance to MSM for k-means (Holder et al., 2022). Finally we would like to investigate the possibility of using different elastic distances with our framework to create an ensemble elastic distance k-means clusterer similar to the elastic ensemble classifier proposed in (Lines and Bagnall, 2015).

ACKNOWLEDGEMENTS

This work has been supported by EPSRC (grant reference EP/W030756/1) and partially subsidised by “Agencia Española de Investigación (España)” (grant reference: PID2020-115454GB-C22/AEI/10.13039/501100011033). David Guijo-Rubio’s research has been subsidised by the University of Còrdoba through grants to Public Universities for the requalification of the Spanish university system of the Ministry of Universities, financed by the European Union - NextGenerationEU (grant reference: UCOR01MS). Some of the experiments were carried out on the High Performance Computing Cluster supported by the Research and Specialist Computing Support service at the University of East Anglia. We would like to thank all those responsible for helping maintain the time series classification archives and those contributing to open source implementations of the algorithms.

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