Extending the SACOC algorithm through the Nyström method for Dense Manifold Data Analysis

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Abstract: Data analysis has become an important field over the last decades. The growing amount of data demands new analytical methodologies in order to extract relevant knowledge. Clustering is one of the most competitive techniques in this context. Using a dataset as a starting point, clustering techniques aim to blindly group the data by similarity. Among the different areas, manifold identification is currently gaining importance. Spectral-based methods, which are one of the main used methodologies in this area, are sensitive to metric parameters and noise. In order to solve these problems, new bio-inspired techniques have been combined with different heuristics to perform the cluster selection, in particular for dense datasets. Dense datasets are featured by areas of higher density, where there are significantly more data instances than in the rest of the search space. This paper presents an extension of a previous algorithm named Spectral-based Ant Colony Optimization Clustering (SACOC), a spectral-based clustering methodology used for manifold identification. This work focuses on improving the SACOC algorithm through the Nyström extension in order to deal with dense data problems. We evaluated the performance of the proposed approach, called SACON, comparing it against online clustering algorithms and the Nyström extension of the Spectral Clustering algorithm using several benchmark datasets.

Keywords: Ant Colony Optimization, Clustering, Data Mining, Machine Learning, Spectral, Nyström, SACON, SACOC

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1 Introduction

The importance of analysing large quantities of data is growing as a consequence of the rapid generation of data from social interactions, smart devices, WiFi and networks, among other sources [1]. Several methodologies have been extended in order to deal with the scale of the data. Recently, there has been an increased interest in methodologies based on MapReduce [1] and online analysis [2]. The former is an approach designed to parallelize the execution of an algorithm by using several nodes to distribute computation in two steps: map (takes an input pair and produces a set of intermediate values) and reduce (merges intermediate values to form a smaller set of values). Online analysis is a methodology where data instances are only processed once and the system keeps no information about old instances, reducing the memory use to allow an algorithm to work with datasets where there are strong density variation.

There are several data mining approaches that deal with dense data. The most relevant ones are those based on supervised and unsupervised learning [3]. Since supervised techniques usually need human assistant in a manual labelling process, which in many cases is costly, unsupervised techniques are gaining importance in this area [2]. One of the most relevant unsupervised techniques is clustering. Clustering is defined as the process of grouping data blindly, using a similarity criterion. Clustering algorithms have been extensively used in several heterogeneous fields [3].

Clustering can be divided into several sub-areas [3], such as centroid, medoid, hierarchical and continuity-based. This work is based on the latter. Continuity-based clustering is a methodology focused on the identification of manifolds within the data. Manifolds are structures defined by the data points inside the search space [4]—i.e., groups of data instances that define a specific form, such as a sphere or a cube. This work aims to extend the Spectral-based ACO Clustering (SACOC) algorithm by incorporating the Nyström extension [5]. SACOC is a bio-inspired clustering algorithm based on Ant Colony Optimisation (ACO) [6].

ACO algorithms are based on the foraging behaviour of the ants: many ants species are able to find the shortest path between their nest and a food source without any visual aid. All communication is performed indirectly by means of pheromone. Ants use pheromones to indicate the path that they have followed, and since ants following shorter paths are faster, more pheromone is deposited—the greater the pheromone concentration, the more attractive a path becomes for other ants. Eventually, all ants will follow the same path, which in most cases corresponds to the shortest path between the nest and a food source. ACO extends this idea to optimization problems and it has been successfully applied in several fields, such as data mining [7, 8].

The SACOC extension proposed in this paper, called SACON, applies a space approximation using a sub-sample of the dataset. This reduces the computational time of the algorithm since it does not need to calculate the similarity metric value for each data pair, allowing the algorithm to deal efficiently with larger datasets. It guarantees accurate solutions even when the reduction has been applied. It is evaluated using several continuity-based dense datasets and the results are compared against online clustering algorithms and the Nyström extension of spectral clustering.

The paper is structured as follows: Section 2 presents the related work, Section 3 introduces the SACOC algorithm, which is extended in Section 4 to create the new SACON algorithm. Section 5 shows the experimental setup used in the experiments presented in Section 6. Finally, Section 7 presents the conclusions.

2 Related Work

Clustering has been widely used in several heterogeneous areas, where the vast amount of data make clustering a promising technique given that it can deal with unlabelled data, while classification usually needs a previous (manual) labelling process. Current challenges are focused on dense and stream data analysis, where online clustering techniques are popular. Another area receiving increasing attention is continuity-based clustering for manifold identification, since it can be used to deal with dense data problems. In this section we review related work on both online and continuity-based clustering, and previous ACO approaches for clustering.

2.1 Online Clustering

The idea behind online clustering algorithm is to analyse this data using real-time techniques. These techniques usually need to deal with large quantities of data, making them suitable for dense data problems. One of the main challenges of real-time analysis is that they need a complete representation of the data space to update the information, limiting their applicability to design new clustering algorithms—e.g., there are clustering algorithms that might not be easily adapted to this kind of analysis [2].

One of the main tools used for online clustering analysis is the Massive On-line Analysis (MOA) tool. This framework provides the following online clustering algorithms:

- Online K-means [2]: This online algorithm updates the centroid position when a new instance arrives. Only one centroid is updated per iteration. It is similar to classical K-means algorithm.
- ClusTree [9]: This online algorithm iteratively updates the information of the clusters. It is able to consider the speed of the data stream generating the concept of the age of an object. It also maintains stream summaries.
2.2 Spectral Clustering

This section presents an overview of the Spectral Clustering (SC) algorithm, which is one of the most relevant continuity-based and manifold identification clustering algorithm. For each data pair, SC starts generating a similarity graph among the data instances. There are three main methodologies that are used [11]:

1. $\epsilon$-neighbourhood graph: all the components whose pairwise distance is smaller than $\epsilon$ are connected.

2. $k$-nearest neighbour graphs: the vertex $v_i$ is connected with vertex $v_j$ if $v_j$ is among the $k$-nearest neighbours of $v_i$.

3. fully connected graph: all points with positive similarity are connected with each other.

This work focuses in the fully connected graph. One of the most important metrics used in SC is the RBF kernel [12] defined by:

$$s(x_i, x_j) = e^{-\sigma \|x_i - x_j\|^2},$$

(1)

where the metric calculates the inverse exponent of the Euclidean distance between points $x_i$ and $x_j$ using a normalization factor $\sigma$.

The second step is related to the study of the eigenvectors of the Laplacian matrix of the similarity graph. Depending on the Laplacian matrix calculation, there are three different techniques related to SC [11]:

1. unnormalized Spectral Clustering: it defines the Laplacian matrix as: $L = D - W$

2. normalized Spectral Clustering: it defines the Laplacian matrix as: $L_{sym} = D^{-1/2}LD^{-1/2} = I - D^{-1/2}WD^{-1/2}$

3. random walks-based normalized Spectral Clustering: it defines the Laplacian matrix as: $L_{rw} = D^{-1}L = I - D^{-1}W$

In these equations, $I$ is the identity matrix, $D$ represents the diagonal matrix whose $(i, i)$-element is the sum of the similarity matrix $i$-th row and $W$ represents the similarity graph. Once the Laplacian is calculated, its eigenvectors are extracted. One of the main problems of SC is related to the consistency of the two classical methods used in the analysis: normalized and unnormalized. A deep analysis about the theoretical effectiveness of normalized clustering over unnormalized can be found in [13].

The last step is the application of a clustering algorithm to the projective space formed by the normalized eigenvectors, considering each row of the matrix as a point. The most frequently applied algorithm is K-means. There are several versions of SC according to the algorithm that is used in this step—e.g., the SACOC [14] algorithm is a Spectral-based algorithm which applies an ACO clustering algorithm instead of K-means.

The main challenge of SC lies in how to compute the eigenvectors and the eigenvalues of the Laplacian matrix of the similarity graph, avoiding a huge memory consumption. For example, when large datasets are analysed, the similarity graph of the SC algorithm requires a high memory storage that makes extremely hard the eigenvalues and eigenvectors computation.

2.3 Ant Colony Optimization in Clustering

ACO algorithms have been extensively applied to supervise learning—e.g., classification rules [15, 16], decision tree induction [7], neural networks [17] and naïve-bayes model [18]. ACO has also been applied to clustering with promising results. Kao and Cheng designed a centroid-based ACO clustering algorithm (ACOC) [19], where ants assign each data instance to one of the available clusters and clusters centroids are adjusted based on this assignment; following a similar direction, we presented the SACOC algorithm [14], which is a spectral extension of ACOC; a similar approach is also used to design a medoid-based ACO clustering (MACOC) algorithm [8], with the difference that ants assign data instances to medoids; Ashok and Messinger focused their work on graph-based clustering of spectral imagery [20], where the data is represented as a graph and an ACO procedure is used to find long paths through the data; several other approaches are discussed in [21].

It is interesting to remark that both MACOC and SACOC are based on ACOC algorithm in terms of the search, where ants visit all data instances to decide the cluster assignation. The main difference between them is the cluster/search space representation used: in ACOC, clusters are represented by centroids in an euclidean space; MACOC represents clusters as medoids (i.e., data instances are used to define the clusters); SACOC uses a spectral transformation to project the original space to a representation where similarities are more apparent.

3 SACOC: Spectral-based ACO Clustering Algorithm

This section presents SACOC [14], as it is the base for the proposed SACON algorithm. SACOC is similar to Spectral Clustering, where the goal of the algorithm is to discriminate the data using the information of a similarity graph and partition it into different clusters.
Figure 1 The construction graph of SACOC. The arrows illustrate a potential trail of an ant.

Algorithm 1 High-level pseudocode for the ACO-based clustering procedure.

Require: \( X = x_1, \ldots, x_N \) data instances and \( k \) number of clusters

Ensure: \( c_1, \ldots, c_k \) best \( k \) centroids

1. Initialize the pheromone matrix \( \tau_0 \).
2. for iteration \( t = 0 \) to maxIterations do
   3. Initialize ants: \( C^a = \{k \text{ randomly chosen centroids}\} \), \( W_a = 0 \) and \( tb^a = 0 \).
   4. for all ant \( a \in A \) (the ants set) do
      5. while \( |tb^a| < N \) do
         6. Select the next instance \( i \).
         7. Choose exploration or exploitation.
         8. Select \( c_u \in C^a \) as the closest centroid.
         9. Set \( w^a_{i,u} = 1 \).
         10. add instance \( i \) to \( tb^a \).
      end while
   11. Calculate the objective function for each ant: \( J^a = \sum_{i=1}^{N} \sum_{j=1}^{k} w^a_{i,j} \cdot d(i,j^a) \).
   12. end for
   13. Rank the ants according to \( J^a \).
   14. Choose the best ant \( a^* \) (iteration-best solution).
   15. Compare it with the best-so-far solution (\( a^{**} \)) and update this value with the maximum between them.
   16. Update pheromone values: \( \tau_{ij}(t+1) = (1 - \rho) \cdot \tau_{ij}(t) + \sum_{h=1}^{r} w^h_{i,j} \cdot \Delta \tau^h_{ij} \).
   17. end for
   18. Re-centralize instances based on \( a^{**} \).
   19. end

3.1 Clustering data

SACOC performs the clustering of the data by using a strategy based on the ACOC algorithm [19], as illustrated in Algorithm 1. The search space is defined in terms of instances and centroids, which can be represented as a graph with an associated \( N \times k \) matrix (where \( N \) is the number of instances and \( k \) is the number of centroids or clusters).

The algorithm uses several ants looking for the best path in the graph, illustrated in Figure 1. Each ant \( a \) has the following features: a list of visited objects (\( tb^a \)), a set of chosen centroids \( C^a \) and a weighted matrix \( W^a \) (related to the assignment of instances to clusters). To create a solution, an ant uses two different strategies: exploration and exploitation. It chooses the strategy according to:

\[
  j = \begin{cases} 
    \arg\max_{u \in N} \{[\tau(i,u)]^{\gamma}(i,u)\}^\beta, & \text{if } q \leq q_0 \\
    S, & \text{otherwise}
  \end{cases}
\]

where \( N \) is the set of nodes associated to instance \( i \), \( [\tau(i,u)]^{\gamma}(i,u) \) is the pheromone value between instance \( i \) and centroid \( u \), \( q_0 \) is a user-defined exploitation probability, \( q \) is a random number for strategy selection, \( \beta \) is an ACO parameter that controls the influence of the heuristic \( j \) in the chosen cluster; \( \gamma(i,u) \) is the heuristic value between \( i \) and \( u \) for \( a^* \), defined as \( 1/d(i,u^a) \), where \( i \) is a data instance and \( u^a \) is the \( a \)-th centroid from the ant \( a \) centroid list. When \( q \) is greater than \( q_0 \), an ant uses the probabilistic exploration strategy \( S \), defined by:

\[
  S = P^a(i,u) = \frac{[\tau(i,u)]^{\gamma}(i,u)\beta}{\sum_{j=1}^{N}[\tau(i,j)]^{\gamma}(i,j)\beta},
\]

where \( P^a(i,u) \) is the probability of assigning instance \( i \) to cluster \( u \) and \( k \) is the total number of clusters.

The algorithm steps presented in Algorithm 1 can be divided in:

1. Initialize pheromone matrix (line 1).
2. Initialize ants (line 3): \((tb^a, C^a, W^a)\), for each ant \( a \) in the colony; then, each ant repeats until \( tb^a \) is full:
   a. Select randomly an instance \( i \) satisfying \( i \notin tb^a \) (line 6).
   b. Select a cluster \( j \); first the ant chooses a strategy; then, it calculates the transition probability and visits a node (line 7).
   c. Update \( tb^a \) and \( W^a \) (lines 9 and 10).
3. Calculate the objective function for each ant (line 12):

\[
  J^a = \sum_{i=1}^{N} \sum_{j=1}^{k} w^a_{i,j} \cdot d(i,j^a),
\]

where \( w^a_{i,j} \) is a weight value of the assignment matrix \( W^a \).
4. Select the best solution (lines 14 to 16). First, rank ants solutions, select the iteration-best solution, apply local search (for more details of local search see [22]) to improve the solution and, finally, compare it with the best-so-far solution and update this value with their maximum.
5. Update pheromone trails (line 17): only the best \( r \) ants are able to add pheromones. Let \( \rho \) be the pheromone evaporation rate \((0 < \rho < 1)\), \( t \) the iteration number, \( r \) the number of elitism ants and \( \Delta \tau^h_{ij} = 1/J^h \) (line 17):

\[
  \tau_{ij}(t+1) = (1 - \rho) \cdot \tau_{ij}(t) + \sum_{h=1}^{r} w^h_{i,j} \cdot \Delta \tau^h_{ij}.
\]
is the sum of the similarity matrix $v$ in columns. Finally, we form the matrix $V$ in the case of repeated eigenvalues—and form the matrix $I$—renormalizing each row of clustering procedure (line 6) to the representation of each $Y$. SACOC. Algorithm 2 presents the high-level pseudocode in a similar way that K-means is modified to generate space. However, it can be modified to consider any kernel originally designed to use Euclidean space as a search. The clustering procedure presented in Algorithm 1 was algorithm [14].

Algorithm 2 High-level pseudocode of SACOC

Require: $X = x_1, \ldots, x_N$ data instances and $k$ number of clusters

Ensure: $c_1, \ldots, c_k$ best $k$ clusters

1: Form the similarity graph $W \in \mathbb{R}^{N \times N}$ defined by $W_{ij} = e^{-\|x_i - x_j\|^2/2\sigma^2}$ if $i \neq j$, and $W_{ii} = 0$.
2: Define $D$ to be the diagonal matrix whose $(i, i)$-element is the sum of the $i$-th row of $W$.
3: Construct the matrix $L = D^{-1/2}WD^{-1/2}$.
4: Find $v_1, \ldots, v_k$, the $k$ largest eigenvectors of $L$ (chosen to be orthogonal to each other in the case of repeated eigenvalues) and form the matrix $V = [v_1 v_2 \ldots v_k] \in \mathbb{R}^{N \times k}$ by stacking the eigenvectors in columns.
5: Form the matrix $Y$ from $V$ by renormalizing each row of $V$ to have unit length (i.e. $Y_{ij} = V_{ij}/(\sum_j V_{ij}^2)^{1/2}$).
6: ClusterData($Y,k$).

6. Check the termination condition: if the number of iterations is greater than the maximum limit, finish; otherwise, go to step 2.

3.2 The spectral hybridisation

The clustering procedure presented in Algorithm 1 was originally designed to use Euclidean space as a search space. However, it can be modified to consider any kernel in a similar way that K-means is modified to generate the SC algorithm—a similar strategy is employed in SACOC. Algorithm 2 presents the high-level pseudocode of SACOC. Consider a graph $G$ and its associated weighted matrix $W$, which is a pairwise similarity graph among the data. The similarity is calculated using a similarity function defined by a kernel $k(x_i, x_j)$, where $x_i$ and $x_j$ are the $i$-th and $j$-th data instances, respectively (line 1). The spectrum of the graph is calculated in a similar fashion as proposed by Ng et al. [23] (lines 2 and 3) to create the SC algorithm. First, we calculate the Laplacian matrix defined by: $L_{sym} = I - D^{-1/2}WD^{-1/2}$, where $I$ is the identity matrix and $D$ represents the diagonal matrix whose $(i, i)$-element is the sum of the similarity matrix $i$-th row. After the creation of the Laplacian matrix, we extract the $v_1, \ldots, v_z$ (line 4), which corresponds with the $z$ largest eigenvectors of $L$—chosen to be orthogonal to each other in the case of repeated eigenvalues—and form the matrix $V = [v_1 v_2 \ldots v_z] \in \mathbb{R}^{N \times z}$ by stacking the eigenvectors in columns. Finally, we form the matrix $Y$ from $V$ by renormalizing each row of $V$ to have unit length (i.e., $Y_{ij} = V_{ij}/(\sum_j V_{ij}^2)^{1/2}$) (line 5). Then, we can consider $Y$ as a projection of the original space and apply the clustering procedure (line 6) to the representation of each point.

4 SACON: Improving the SACOC algorithm through the Nyström extension

The goal of combining SACOC with the Nyström extension is to reduce the dimensions sampling of the similarity matrix. The reduction is achieved by choosing a subset of points $S = \{s_1, \ldots, s_n\} \in X = \{x_1, \ldots, x_N\}$ (where $n < N$)—the high-level pseudocode of SACON is presented in Algorithm 3. Given a similarity matrix $W$ (related to the similarity graph), we need to extract the eigenvectors of its spectrum in order to project the data. In this work, the spectrum is defined by $L_{sym}$. In order to improve the computational time through the Nyström method, we need to use the sub-sample $S$ and reformulate the whole process to describe how to extract approximate eigenvectors of the spectrum using less information related to the original similarity matrix.

The approximation is obtained by applying the Nyström extension [5], defined as follows:

**Nyström Extension:** Let $k(x_i, x_j)$ be a kernel whose Gram matrix $K$ is symmetric and positive semi-definite satisfying $K_{ij} = k(x_i, x_j)$. We assume that the eigendecomposition is $K = U \Lambda U^T$ where $U$ is the orthogonal matrix of eigenvectors and $\Lambda$ is the diagonal matrix composed by the eigenvalues. Then, the Nyström extension for a new instance $x$ is the eigenvector approximation $\hat{u}^k(x)$ to the real $u^k(x)$ given by:

$$
\hat{u}^k(x) = \frac{1}{N} \sum_{j=1}^N k(x, x_j) \hat{u}_j^k,
$$

where $\hat{u}_j^k$ is the extension of the eigenvector $u_j^k$ calculated from the sub-sample set of the matrix $K$, $u_j^k$ is the coordinate $j$ of the eigenvector and $\lambda_j^k$ is the eigenvalue associated to the eigenvector $u_j^k$.

In order to apply the extension to calculate the eigenvectors of $L_{sym}$, we simplify the process as follows. First, the eigenvectors of $L_{sym} = I - D^{-1/2}WD^{-1/2}$ are equivalent to the eigenvectors of $P = D^{-1/2}WD^{-1/2}$ since the identity matrix can be omitted.

The eigenvectors approximation follows the scheme specified in Algorithm 3. In this case, we use $W$ and take the sub-samples set $S$ to define a sub-matrix of $W$ called $A$. The matrices satisfy:

$$
W = \begin{pmatrix}
A & B \\
B^t & C
\end{pmatrix},
$$

where $A \in \mathbb{R}^{n \times n}$, $B \in \mathbb{R}^{n \times (N-n)}$ and $C \in \mathbb{R}^{(N-n) \times (N-n)}$ (line 3). The Nyström extension will only need the augmented matrix $(A|B)$ formed by $A$ and $B$. The matrix $C$ is the part that we want to approximate and satisfies that has more elements than $A$, due to $n < N$. Using matrix $C$, it will be able to approximate eigenvectors of $U$ calculating the eigenvectors of $A$, denoted by $\hat{U}$ (line 4). The eigenvectors $\hat{U}$ are extended to form the approximation...
$\mathcal{U}$, which is close to the original $U$. Once the eigenvectors $\tilde{U}$ have been calculated, we have the eigendecomposition of $A$, denoted as $A = \tilde{U} \check{\Lambda} \tilde{U}^T$ (where $\tilde{U}$ is orthonormal). The approximation $\mathcal{U}$ of the eigenvectors of $W$ is calculated as (line 5):

$$\mathcal{U} = \left( \begin{array}{c} \tilde{U} \\ B^T \check{\Lambda} \end{array} \right).$$

Now, we need to calculate the Laplacian eigenvectors, which are approximated by the matrix (line 7): $L' = D^{-1/2} W' D^{-1/2}$, where $W' = \tilde{U} \check{\Lambda} \tilde{U}^T$, $\check{\Lambda} = \check{\Lambda}^T$. Setting $D_{i,i} = \sum_{j=1}^N w'_{ij}$, we can redefine $D$ as:

$$D = \begin{pmatrix} D_1 & 0 \\ 0 & D_2 \end{pmatrix},$$

where $D_1 \in \mathbb{R}^{n \times n}$, $D_2 \in \mathbb{R}^{(N-n) \times (N-n)}$ and $L$ can be approximated by (lines 7 and 8):

$$L' = \left( \begin{array}{c} A' \\ (B')^T \\ B' \\ (A')^{-1} B' \end{array} \right),$$

where $A' = D_1 A$ and $B' = D_2 B$. In order to approximate the eigenvectors of $L'$ we will use the same methodology that we have to approximate $W$. Assuming that the eigenvectors of $A'$ and eigenvalues are $\lambda^o$ and $\lambda^n$, and using $A'$ as the base, the approximation is given by:

$$\lambda' = \frac{N}{n} \lambda^n, \quad \mathcal{V} = \sqrt{\frac{n}{N}} \left[ \begin{array}{c} A' \\ (B')^T \end{array} \right] \mathcal{V}^o \lambda^o,$$

where $\mathcal{V}$ and $\lambda'$ are the extended eigenvectors and eigenvalues of $L'$. Since $\mathcal{V}$ is not orthogonal, which is required by the spectral clustering algorithm, we will use the Fowlkes et al. [5] transformation and define the matrix $R$ as (line 9):

$$R = A' + (A')^{-1/2} B'(B')^T (A')^{-1/2}.$$

This matrix can be decomposed as $R = U_R \check{\Lambda}_R U_R^T$, since $A'$ is positive define. Then, $\mathcal{V}$ is defined as:

$$\mathcal{V} = \left[ \begin{array}{c} A' \\ (B')^T \end{array} \right] (A')^{-1/2} U_R \check{\Lambda}_R^{-1/2}$$

and finally, $P$ is defined as:

$$P = D^{-1/2} W' D^{-1/2} = \mathcal{V} \check{\Lambda}_R \mathcal{V}^T.$$

At the end of this process, $\mathcal{V}$ are the eigenvectors used cluster the data (i.e., the projection of the data based on the Spectrum). This whole process helps to reduce the computational time consumed by the pairwise calculation, which is usually performed to create the whole similarity graph, and also reduces the memory consumption.

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### Algorithm 3: High-level pseudocode of SACON algorithm.

**Require:** $X = x_1, \ldots, x_N$ data instances and $k$ number of clusters

**Ensure:** $c_1, \ldots, c_k$ best $k$ clusters

1. Select a subsample of the data instances: $S = \{s_1, \ldots, s_n\} \subset X = \{x_1, \ldots, x_N\}$ where, $n < N$.
2. Form the similarity graph $A \in R^{n \times n}$ defined by $A_{ij} = e^{-\sigma |s_i - s_j|^2}$ if $i \neq j$, and $A_{ii} = 0$.
3. Calculate the matrix $B$ formed by the similarities among the elements of $A$ and the rest of data instances.
4. Calculate the eigenvectors of $A$, named $\tilde{U}$ and the eigenvalues, named $\check{\Lambda}$.
5. Calculate the approximate eigenvectors $\mathcal{U}$ of $W$ which try to approximate the real eigenvectors, named $U$, as:

$$\mathcal{U} = \left( \begin{array}{c} \tilde{U} \\ B^T \check{\Lambda} \end{array} \right).$$

6. Define $D$ to be the diagonal matrix whose $(i,i)$-element is the sum of the $i$-th row of $W$.
7. Construct the matrix $L' = D^{-1/2} W' D^{-1/2}$.
8. Separate $L' = \left( \begin{array}{c} A' \\ (B')^T \end{array} \right) (A')^{-1} B' (A')^{-1/2}$, and calculate its eigenvector decomposition $R = U_R \check{\Lambda}_R U_R^T$.
9. Set $\mathcal{V} = \left[ \begin{array}{c} A' \\ (B')^T \end{array} \right] (A')^{-1/2} U_R \check{\Lambda}_R^{-1/2}$.
10. Find $v_1, \ldots, v_k$, the $k$ largest eigenvectors of $L'$ and form $V = [v_1, v_2, \ldots, v_k] \in R^{n \times k}$ by stacking the eigenvectors in columns.
11. Form the matrix $Y$ from $V$ by renormalizing each row of $V$ to have unit length (i.e. $Y_{ij} = V_{ij} / (\sum_j V_{ij}^2)^{1/2}$).
12. ClusterData$(Y,k)$.

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### 5 Experimental Setup

The evaluation of a clustering algorithm is a sensitive process, due to clustering being a blind process. There are no universal methods to evaluate it, in particular when the algorithm deals with large datasets. In this work we have focused the evaluation on comparing the proposed SACON algorithm against state-of-the-art algorithms. Experiments have been carried out using the Nyström extension of Spectral Clustering (SC+Nyström) [5], CluStream, Online K-means and ClusTree.

The parameters of SACON are: the sub-sample size fixed to 500 and the sigma value was calculated using the methodology described by Ng et al. [23], the same value used for SC+Nyström); the number of ants is 10, elitism is 1, exploitation probability is 0.0001, initial pheromone values have been set to $1/k$ (where $k$ is the number of clusters), $\beta = 2.0$, $\rho = 0.1$, local search probability is
0.001 and the maximum number of iterations is 1000. We have made no attempt to tune the parameters to individual datasets.

The metrics used are the Euclidean Distance for online algorithms (CluStream, Online K-means and ClusTree) and the RBF for the spectral-based algorithms (SACOC and SC+Nyström). The evaluation is performed comparing the results with the real labels to measure the accuracy, defined by:

$$\text{sim}(C_i, C_j) = \frac{\sum_{q=1}^{n} \delta_{C_i}(x_q) \delta_{C_j}(x_q)}{2 \left( \frac{1}{|C_i|} + \frac{1}{|C_j|} \right)} ,$$

where $|C_i|$ is the number of elements of cluster $C_i$ and $\delta_{C_i}(x_q)$ is the Kronecker $\delta$ defined by:

$$\delta_{C_i}(x_q) = \begin{cases} 0 & \text{if } x_q \notin C_i \\ 1 & \text{if } x_q \in C_i \end{cases} .$$

All algorithms have been executed 100 times. The statistical test analysis was performed using the non-parametric Wilcoxon test, which does not require a normal distribution. The aim is to compare SACON against SC+Nyström, since both algorithms use the Nyström extension but differ in the way they cluster the data. We have considered that there is statistically difference when the $p$-value of the test is lesser than 0.05 (5% significance level).

### 5.1 Dataset Description

The datasets have been generated using the R package mlbench [24], a collection of standard benchmark problems. The datasets—all composed by 50,000 instances—that have been generated are the following (see Figure 2 for some examples):

- **Cassini**: 2-dimensional problem with 3 clusters (2 external banana-shaped clusters with a circular cluster between them).
- **Cuboids**: 3-dimensional problem with 3 cuboids and a small cube in the middle of them.
- **Hypercube**: 8 spherical Gaussians distributed at the corners of an 8-dimensional cube.
- **Shapes**: a Gaussian, square, triangle and wave in 2 dimensions.
- **Simplex**: 4 spherical Gaussians distributed at the corners of a 4-dimensional simplex.
- **Smiley**: 2 Gaussian eyes, a trapezoid nose and a parabola mouth (with vertical Gaussian noise).
- **Spirals-1**: 2 entangled spirals without noise.
- **Spirals-2**: 2 entangled spirals with noise.

### 6 Computational Results

Table 1 presents the results achieved by the algorithms in each dataset. As we can observe, the proposed SACON algorithm obtains very good results overall.

In the case of **Cassini** dataset, SACON obtains the best results—the statistical test shows that SACON is significantly better than SC+Nyström. It is important to remark that the standard deviation (SD) of SACON is 0, which means that these results are stable. SC+Nyström also obtains good results, while the remaining algorithms obtain poor results. The Cassini dataset has two sections that can be easily defined using a Gaussian distribution, but when the number of instances is high the boundaries between these distributions are harder for the discrimination process; SC+Nyström is also sensitive to the noise produced in this situation.

For **Cuboids** dataset, Clustream obtains the best median results (100.0%) followed by SC+Nyström (99.74%), with SACON achieving close results (99.33%). SACON is the most stable of the algorithms in this dataset (0.0066 of SD). The Wilcoxon test shows that there is not statistical difference between the SACON and SC+Nyström.

**Hypercube** results show that SACON is able to discriminate the clusters perfectly. Clustream also obtains good results during the discrimination process. SC+Nyström, Clustree and Online K-means have problems to discriminate the cluster distribution. This dataset is challenging due to the large quantities of data, which introduces noise during the cluster separation (Figure 2(b)). The algorithms are also less stable than SACON (the SD of SACON is 0). Again, SACON is statistically significantly better than SC+Nyström.

In the case of **Shapes**, the best results according to the median are achieved by Clustream, Clustree and Online K-means. SACON obtains good results (99.98%) and it is also very stable (the SD is 0.0002). SC+Nyström obtains the worst results (68.71%). According to the statistical test, SACON is statistically significantly better than SC+Nyström again.

[Figure 2](#) Illustration of the synthetic datasets used in the experiments.
Table 1 Median and standard deviation accuracy results obtained by each algorithm, measured over 100 executions. Values in bold show the best results. The ▲ symbol indicates the datasets where the results of SACON are statistically significantly better than the results of the benchmark SC+Nystrom algorithm.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>SACON</th>
<th>SC+Nystrom</th>
<th>Online K-means</th>
<th>Clustree</th>
<th>Clustream</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cassini</td>
<td>99.98% ± 0.0000 ▲</td>
<td>98.61% ± 0.1681</td>
<td>65.47% ± 0.0003</td>
<td>66.97% ± 0.1301</td>
<td>67.39% ± 0.0478</td>
</tr>
<tr>
<td>Cuboids</td>
<td>99.33% ± 0.0066</td>
<td>99.74% ± 0.1736</td>
<td>88.84% ± 0.1219</td>
<td>72.74% ± 0.1108</td>
<td>100.0% ± 0.1157</td>
</tr>
<tr>
<td>Hypercube</td>
<td>100.0% ± 0.0000 ▲</td>
<td>76.71% ± 0.1565</td>
<td>81.36% ± 0.1122</td>
<td>82.29% ± 0.1126</td>
<td>100.0% ± 0.0541</td>
</tr>
<tr>
<td>Shapes</td>
<td>99.98% ± 0.0002 ▲</td>
<td>68.71% ± 0.4425</td>
<td>100.0% ± 0.1389</td>
<td>100.0% ± 0.1765</td>
<td>100.0% ± 0.0001</td>
</tr>
<tr>
<td>Simplex</td>
<td>100.0% ± 0.0000</td>
<td>100.0% ± 0.0000</td>
<td>100.0% ± 0.1508</td>
<td>100.0% ± 0.1703</td>
<td>100.0% ± 0.0000</td>
</tr>
<tr>
<td>Smiley</td>
<td>99.56% ± 0.0003 ▲</td>
<td>74.99% ± 0.1627</td>
<td>62.54% ± 0.1729</td>
<td>64.41% ± 0.1608</td>
<td>91.60% ± 0.1451</td>
</tr>
<tr>
<td>Spirals-1</td>
<td>100.0% ± 0.0000</td>
<td>100.0% ± 0.0991</td>
<td>50.00% ± 0.0000</td>
<td>50.01% ± 0.0002</td>
<td>50.01% ± 0.0006</td>
</tr>
<tr>
<td>Spirals-2</td>
<td>63.03% ± 0.0001 ▲</td>
<td>59.59% ± 0.0392</td>
<td>59.36% ± 0.0000</td>
<td>58.84% ± 0.0295</td>
<td>59.59% ± 0.0392</td>
</tr>
</tbody>
</table>

6.1 Discussion

SACON shows competitive results identifying manifolds when is applied to continuity-based data—such as Cassini, Shapes, Smiley and Spirals—while the remaining algorithms have problems (e.g., in Spirals, which is a pure continuity-based problem). It also performs better than the rest when we add noise to the dataset. The algorithm also obtains more stable results than the others according to the standard deviation. The spectral transformation in combination with the ACO clustering procedure is probably the main reason for the algorithm performance—the transformation makes the regularities in the data space more apparent and the ACO global search procedure is able to find the (near-)optimal cluster assignment. The performance is correlated to the type of clustering that this algorithm can face, i.e., continuity-based clustering.

In order to compare the memory usage of the proposed algorithm against SACOC (its predecessor), it is important to remark that SACON uses a matrix of 50,000 × 500 instances, whereas SACOC uses 50,000 × 50,000 instances. The memory usage of the former is around 0.18 GB while the latter consumes around 18.63 GB. The memory consumption of SACON grows linearly, whereas SACOC grows exponentially. The remaining algorithms are not spectral algorithm, with the exception of SC+Nystrom—i.e., they do not use a similarity matrix. Regarding the memory consumption, the online algorithms only keep in memory the information about the centroids they are using (similar to K-means), i.e., the memory grows according to the number of centroids.

7 Conclusions

This paper proposed an improvement of the SACOC algorithm [14] using the a Nystrom extension [5]. The new algorithm, called SACON, applies the spectral transformations and the Nystrom extension to the original search space in order to cluster the data in the projective space. The projection consists on transforming the original data into a graph-based representation (through a similarity graph) and calculate its Laplacian matrix. During this calculation, the Nystrom extension guarantees that the Laplacian calculation is accurate enough using less information. Once the Laplacian has been obtained, the eigenvectors are extracted and normalized to generate the projective space—this extraction also applies the Nystrom extension in order to reduce the memory usage.

The proposed SACON algorithm shows good results in the studied datasets. It is able to discriminate manifolds and continuity-based clusters with more stable results in dense datasets, when compared to Spectral Clustering using the Nystrom extension and modern online clustering algorithms. The future work will be mainly focused on running experiments with real-world datasets.
large datasets, in particular large datasets with cluster intersections.

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References