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Abstract

Clustering high-dimensional data has been a challenging problem in data mining and machining learning. Spectral clustering via sparse representation has been proposed for clustering high-dimensional data. A critical step in spectral clustering is to effectively construct a weight matrix by assessing the proximity between each pair of objects. While sparse representation proves its effectiveness for compressing high-dimensional signals, existing spectral clustering algorithms based on sparse representation use those sparse coefficients directly. We believe that the similarity measure exploiting more global information from the coefficient vectors will provide more truthful similarity among data objects. The intuition is that the sparse coefficient vectors corresponding to two similar objects are similar and those of two dissimilar objects are also dissimilar. In particular, we propose two approaches of weight matrix construction according to the similarity of the sparse coefficient vectors. Experimental results on several real-world high-dimensional data sets demonstrate that spectral clustering based on the proposed similarity matrices outperforms existing spectral clustering algorithms via sparse representation.

Keywords: spectral clustering, high-dimensional data, weight matrix, sparse representation

1. Introduction

As an important task in data mining cluster analysis aims at partitioning data objects into several meaningful subsets, called clusters, such that data objects are similar to those in the same cluster and dissimilar to those in different clusters. With advances in database technology and real-world need of informed decisions, data sets to be analyzed are getting bigger—with many more records and variables. Examples of high-dimensional data sets include document data [1], user ratings data [2], multimedia data [3], financial time series data [4], gene expression data [5] and so on. Due to the “curse of dimensionality” [6], clustering
high-dimensional data has been a challenging task and therefore, attracts much research in data mining and related research domains [7].

Many of the existing high-dimensional clustering approaches can be categorized into the following three types: dimension reduction [8], subspace clustering [9] and spectral clustering [10–12]. The first two types transform the original feature space to a lower-dimensional space and then apply an ordinary clustering algorithm (such as K-means). The focus is on how to extract more important features of data objects and avoid noises from the less important dimensions. Spectral clustering is based on the spectral graph model, which searches for clusters in the full feature space and is equivalent to graph min-cut problem based on a graph structure constructed from the original objects in vector space [12]. Spectral clustering is also considered superior to traditional clustering algorithms due to its deterministic and polynomial time solution. All these characteristics make spectral clustering suitable for high-dimensional data clustering [10].

The effectiveness of spectral clustering depends on the weights between each pair of data objects. Thus, it is vital to construct a weight matrix that faithfully reflects the similarity information among objects. Traditional simple weight construction, such as ε-ball neighborhood, k-nearest neighbors, inverse Euclidean distance [13, 14] and Gaussian radial basis function (RBF) [12], is based on the Euclidean distance in the original space, thus not suitable for high-dimensional data.

In this chapter, we focus on effective weight construction for spectral clustering, based on sparse representation theory. Sparse representation aims for representing each object approximately by a sparse linear combination of other objects, which comes from the theory of compressed sensing [15]. Coefficients of the sparse linear combination represent the closeness of each object to other objects. Traditional spectral clustering methods based on sparse representation [16] use these sparse coefficients directly to build the weight matrix, thus only local information is utilized. However, exploiting more global information of the whole coefficient vectors promises better performance, followed by an assumption that the sparse representation vectors corresponding to two similar objects should be similar, since they can be reconstructed in a similar fashion using other data objects. If two objects are contributing in a similar manner to the reconstruction of all other objects in the same data set, they are considered similar.

Therefore, this chapter presents a spectral clustering approach of high-dimensional data exploiting global information from sparse representation solution. More specifically, using sparse representation, we firstly convert each high-dimensional data object into a vector of sparse coefficients. Then, the proximity of two data objects is assessed according to the similarity between their sparse coefficient vectors. This construction considers the complete information of the solution coefficient vectors of two objects to analyze the similarity between these two objects rather than directly using a particular single sparse coefficient, which only considers local information. In particular, we propose two different weight matrix construction approaches: one of which is based on consistent sign set (CSS) and the other is based on the cosine similarity (COS) between the two vectors. Extensive experimental results on several image data sets show that similarly exploiting the global information from the solutions of sparse representation works better than using local information of the solutions under a variety of clustering performance metrics.
2. Related work

2.1. Techniques for high-dimensional data

There are many techniques for dealing with high-dimensional signals (or data), popular of which include non-negative matrix factorization (NMF), manifold learning, compressed sensing and some combinations between them.

Non-negative matrix factorization (NMF) is a powerful dimensionality reduction technique and has been widely applied to image processing and pattern recognition applications [17], by approximating a non-negative matrix $X$ by the product of two non-negative low-rank factor matrices $W$ and $H$. It has attracted much attention since it was first proposed by Paatero and Tapper [18] and has already been proven to be equivalent in terms of optimization process with K-means and spectral clustering under some constraints [19]. The research about NMF can be generally categorized into the following groups. The first group is focused on the distance measures between the original matrix and the approximate matrix, including Kullback–Leibler divergence (KLNMF) [17], Euclidean distance (EucNMF) [20], earth mover’s distance metric [21] and Manhattan distance-based NMF (MahNMF) [22]. Besides, there are researches about how to solve the optimization of NMF efficiently and the scalability of NMF algorithms for large-scale data sets, for example, fast Netwon-type methods (FNMA) [23], online NMF with robust stochastic approximation (OR-NMF) [24] and large-scale graph-regularized NMF [25]. Moreover, how to improve the performance of NMF using some constrains or exploiting more information of data is also popular, such as sparseness constrained NMF (NMFsc) [26], convex model for NMF using $l_1,\infty$ regularization [27], discriminant NMF (DNMF) [28], graph-regularized NMF (GNMF) [29], manifold regularized discriminative NMF (MD-NMF) [30] and constrained NMF (CNMF) [31] incorporating the label information.

Manifold learning is another theory to process high-dimensional data, assuming that the data distribution is supported on a low-dimensional sub-manifold [32]. The key idea of manifold learning is that the locality structure of high-dimensional data should be preserved in low-dimensional space after dimension reduction, which is exploited as a regularization term [33–35] or constraint [36, 37] to be added to the original problem. It has been widely used to machine learning and computer vision, such as image classification [38], semi-supervised multiview distance metric learning [39], human action recognition [40], complex object correspondence construction [41] and so on.

Besides the abovementioned two approaches for high-dimensional data, in recent years, sparse representation coming from compressed sensing has also attracted a great deal of attention and proves to be an extremely powerful tool for acquiring, representing and compressing high-dimensional data. The following section will briefly review of sparse representation.

2.2. Brief review of sparse representation

Given a sufficient high-dimensional training data set, $X = [x_1, x_2, \ldots, x_n] \in \mathbb{R}^{m \times n}$, where $x_i = [x_{i1}, x_{i2}, \ldots, x_{im}]^T \in \mathbb{R}^m$ is the column vector of the $i$-th object. Research on manifold learning [32] has
proved that any new test data \( y \) lie on a lower dimensional manifold, which can be approximately represented by a linear combination of the training objects:

\[
y = a_1 x_1 + \cdots + a_i x_i + \cdots + a_n x_n = Xa \in \mathbb{R}^m.
\] (1)

Obviously, if \( m \gg n \), Eq. (1) is overdetermined, and \( a \) can usually be found as its unique solution. Typically, the number of attributes is much less than that of training objects (i.e. \( m \ll n \)) and Eq. (1) is undetermined, so its solution is not unique.

However, if we add the constraint that the best solution of Eq. (1) should be as sparse as possible, which means that the number of non-zero elements is minimized, the solution becomes unique. Such a sparse representation can be obtained by solving the optimization problem:

\[
a^* = \arg \min_a \|a\|_0 \quad \text{subject to} \quad y = Xa,
\] (2)

where \( \| \cdot \|_0 \) denotes the \( l_0 \)-norm of a vector, counting the number of non-zero entries in the vector. Donoho [42] proves that if matrix \( X \) satisfies restricted isometry property (RIP) [43], Eq. (2) has a unique solution of \( a \).

However, it is NP-hard to find the sparsest solution of an underdetermined equation: that is, there is no known approach to find the sparsest solution that is significantly more efficient than exhausting all subsets of the entries for \( a \). Researchers in emerging theory of compressed sensing [44] reveal that the non-convex optimization in (2) is equal to the following convex \( l_1 \) optimization problem if the solution \( a \) is sparse enough:

\[
a^* = \arg \min_a \|a\|_1 \quad \text{subject to} \quad y = Xa,
\] (3)

where \( \| \cdot \|_1 \) denotes the \( l_1 \)-norm of a vector, summing the absolute value of each entry in the vector. This problem can be solved in polynomial time by standard linear programming methods [45].

Since the real data contains noise, it may not be possible to express the test sample exactly as a sparse representation of the training data. The sparse solution \( a \) can still be approximately obtained by solving the following stable \( l_1 \) optimization problem:

\[
a^* = \arg \min_a \|a\|_1 \quad \text{subject to} \quad \|y - Xa\|_2 \leq \varepsilon,
\] (4)

where \( \varepsilon \) is the maximum residual error; \( \| \cdot \|_2 \) denotes the \( l_2 \)-norm of a vector.

In many situations, we do not know the noise level \( \varepsilon \) beforehand. Then we can use the Lasso (least absolute shrinkage and selection operator) [46] optimization algorithm to recover the sparse solution from the following \( l_1 \) optimization:

\[
a^* = \arg \min_a \lambda \|a\|_1 + \|y - Xa\|_2.
\] (5)
where $\lambda$ is a scalar regularization parameter of the Lasso penalty, which directly determines how sparse $\alpha$ will be and balances the trade-off between reconstruction error and sparsity.

In addition to Lasso, other sparse learning models are also developed. It will be the elastic net model [47] if the $l_2$-norm of $\alpha$ is also added to Eq. (5) as another penalty term. Double shrinking algorithm (DSA) [48] compresses image data on both dimensionality and cardinality via building either sparse low-dimensional representations or a sparse projection matrix for dimension reduction. Go decomposition (GoDec) [49] tried to efficiently and robustly decompose a matrix with the low-rank part $L$ and the sparse part $S$. Locality structure of manifold can also be combined with sparse representation, such as manifold elastic net (MEN) [50] and graph-regularized sparse coding (GraphSC) [51], laplacian sparse coding (LSc) [35] and Hypergraph laplacian sparse coding (HLSc) [35].

Learning tasks such as classification and clustering usually perform better and cost less (time and space) on compressed representations than on the original data [48]. Therefore, supervised learning and pattern recognition based on the sparse representation coefficients using these sparse learning models are proposed, such as sparse representation-based classification (SRC) [52], Local_SRC [53], Kernel_SRC [54] and the methods outperform traditional classifier, such as SVM, nearest neighbor (NN) and nearest subspace (NS).

2.3. Sparse representation for clustering

Inspired by the successful application of sparse representation in the above-supervised learning approaches, researchers have also exploited sparse representation in unsupervised [55–57] and semi-supervised clustering [58, 59]. The main idea of clustering via sparse representation is to build weight matrix directly from normalized and symmetrized coefficients of sparse representation coefficients, called sparsity-induced similarity (SIS) measure [59]. To a certain extent, weight measure approaches derived from sparse representation can reveal the neighborhood structure without calculating Euclidean distance, which means a great potential to clustering high-dimensional data.

Some significant work applying SIS to spectral clustering is reviewed as follows. Sparse subspace clustering [55] directly uses the sparse representation of vectors lying in a single low-dimensional linear subspace to cluster the data into separate subspaces, followed by applying spectral clustering. It is also extended to clustering data contaminated by noise, missing entries or outliers. Experiments show that its performance for clustering motion trajectories outperforms state-of-the-art methods, such as power factorization and principal component analysis. Image clustering via sparse representation [56] characterizes the graph adjacency structure and graph weights by sparse linear coefficients, which is more effective than Gaussian RBF [12] to cluster an image data set. In semi-supervised learning by sparse representation [18], the graph adjacency structure as well as the graph weights of the directed graph construction is derived simultaneously and in a parameter-free manner to utilize both labeled and unlabeled data. Experiments on semi-supervised face recognition and image classification demonstrate the superiority over the counterparts based on traditional graphs (e.g. $\epsilon$-ball neighborhood, k-nearest neighbors). Compared to approaches using SIS of real
numbers, non-negative SIS measure [57] exploits the symmetric coefficients of non-negative sparse representation as weight matrix, which outperforms similarity measures, such as SIS and Euclidean (with Gaussian RBF baseline [12]), in cluster analysis of spam images.

However, all the above-existing approaches based on sparse representation treat directly the coefficients or just normalized coefficients of sparse representation as the weight matrix. These cannot exactly reflect the similarity between objects because the coefficients of sparse representation are somehow local similarity and sensitive to outliers. Our approach is expected to provide more effective weight matrix construction using more global content from the solution coefficients of sparse representation.

2.4. Graph construction with sparse representation

In clustering analysis, given a high-dimensional object data set \( X = [x_1, x_2, \ldots, x_n] \in \mathbb{R}^m \times n \), \( x_i = [x_{i1}, x_{i2}, \ldots, x_{im}]^T \in \mathbb{R}^m \), we can use Eq. (5) to represent each objects \( x_i \) as a linear combination of other objects. The coefficients vector \( \alpha_i \) of \( x_i \) can be calculated by solving the following Lasso optimization:

\[
\alpha_i^* = \arg \min_{\alpha_i} \lambda \| \alpha_i \|_1 + \| x_i - X \alpha_i \|_2^2, \tag{6}
\]

where \( X_i = X \setminus x_i = [x_1, \ldots, x_i \setminus 1, x_i + 1, \ldots, x_n]^T; \alpha_i = [\alpha_{i1}, \ldots, \alpha_{i(i-1)}, \alpha_{i(i+1)}, \ldots, \alpha_{in}]^T \).

Once we get the coefficient vector \( \alpha_i \) for each object \( x_i \) \( (i = 1, 2, \ldots, n) \) as a sparse representation of all other data objects by solving the \( l_1 \) optimization Eq. (6), we can construct the weight matrix using different approaches.

Existing weight matrix constructions via sparse representation are based on the assumption that coefficients in the sparse representation reflect the closeness or similarity between two data objects. For example, the SIS measure [20] is computed as:

\[
w_{ij} = \max \left\{ \alpha_{ij}, 0 \right\}; \quad \text{SIS}_{ij} = \frac{w_{ij} + w_{ji}}{2}. \tag{7}
\]

The \( l_1 \) Directed Graph Construction (DGC) measure [19] is computed as:

\[
\text{DGC}_{ij} = \frac{|\alpha_{ij}| + |\alpha_{ji}|}{2}. \tag{8}
\]

Obviously, the similarity calculation using the absolute coefficients in Eq. (8) will mistake the big negative coefficient as high similarity, resulting in a cluster of two objects with apparent opposite attributes value.

The non-negative SIS measure [22] adds a non-negative constraint in \( l_1 \) optimization Eq. (6):

\[
\alpha_i^* = \arg \min_{\alpha_i} \lambda \| \alpha_i \|_1 + \| x_i - X \alpha_i \|_2 \text{ s.t. } \alpha_{ij} \geq 0. \tag{9}
\]
Then the non-negative SIS measure is computed as:

$$NN_{ij} = \frac{\alpha_{ij}}{\sum_{k=1, k \neq i}^{n} \alpha_{i,k}}.$$  \hspace{1cm} (10)

3. Sparse representation for spectral clustering

Our proposed clustering algorithm consists of three steps: (1) solving $l_1$ optimization of sparse representation to obtain the coefficients of each object; (2) constructing weight matrix between objects on the basis of coefficients using more global content forms the solution coefficients of sparse representation; and (3) exploiting the spectral clustering algorithm with the weight matrix to get partition of the graph.

Compared to the direct construction methods using the independent solutions of Eq. (6), we have the assumption that for any two objects $x_i$ and $x_j$, the more similar they are, the more similar the corresponding coefficient vectors (e.g., $\alpha_i$ and $\alpha_j$) are not only a particular coefficient ($\alpha_{i,j}$ or $\alpha_{j,i}$). According to this assumption, we propose the following two graph adjacency structure and weight matrix constructions, which are expected to use the global information of the solution coefficients.

3.1. Proximity based on a consistent sign set

To get the similarity of two objects clearly and logically, we firstly find an object set for each of the two different objects $x_i$ and $x_j$ from the object data set $X$, called CSS. This definition is based on the assumption that the more objects of which a pair of objects both positively contribute to the reconstruction, the more similar the pair of objects are. In particular, the sparse reconstruction coefficients corresponding to $x_i$ and $x_j$ for every object in this set are both positive, defined as follows:

$$CSS(x_i, x_j) = \{X_k | \alpha_{k,i} > 0 \land \alpha_{k,j} > 0, k \neq i, k \neq j \} \ \forall i \neq j.$$  \hspace{1cm} (11)

Furthermore, we can construct graph adjacency structure and weight matrix as follows. A directed edge is placed between objects $x_i$ and $x_j$ if $CSS(x_i, x_j) \neq \Phi$ and the weight between object $x_i$ and $x_j$ is defined as the ratio of the $CSS(x_i, x_j)$’s cardinal to the total number of objects:

$$w_{ij} = \begin{cases} \frac{|CSS(x_i, x_j)|}{n} & i \neq j \\ 0 & i = j \end{cases},$$  \hspace{1cm} (12)

where $n$ is the total number of objects in $X$. Obviously, the weight is between 0 and 1.

3.2. Proximity based on cosine similarity of coefficient vector

We can construct coefficient matrix $A$ of data set $X$, to which transforming solution coefficients of Eq. (6) are:
A directed edge is placed from object $x_i$ and $x_j$ if angle cosine of the two corresponding vectors is greater than 0, that is:

$$\alpha_0^i \cdot \alpha_0^j > 0, \quad (14)$$

where $\alpha_0^i$ denotes the $i$-th row vector of $A$.

The weight between object $x_i$ and $x_j$ is defined as the cosine similarity of $\alpha_i'$ and $\alpha_j'$:

$$w_{ij} = \max \left\{ 0, \frac{\alpha_i' \cdot \alpha_j'}{\|\alpha_i'\|_2 \times \|\alpha_j'\|_2} \right\} \quad i \neq j$$

$$w_{ij} = \begin{cases} 1 & A(i,j) > 0 \\ 0 & \text{else} \end{cases}; \quad (15)$$

From the above similarity calculation formula, two objects have large similarity in condition that the corresponding solution coefficients of Eq. (6) are much similar, which is expected to use the whole solution coefficients.

3.3. The relationship between consistent sign set (CSS) and cosine similarity of coefficient vector (COS)

Since proposed proximity based on both CSS and COS are trying to exploit more information from the solution coefficient of sparse representation, the relationship between each other is following:

1. Both of them assess the weight between two objects according to the similarity between the corresponding coefficient vectors of the two objects. However, the difference is that proximity based on CSS uses the column vectors of the coefficient matrix $A$ while proximity based on COS calculates the similarity between row vectors, which means two understandings of the coefficient matrix. The reason for defining these two approaches like this is just experimental.

2. Proximity based on COS is to calculate the similarity of the original coefficient vector, while CSS can be considered as the discretization of the original coefficient vector with threshold zero. Therefore, proximity based on COS can be seen as the generalization of that based on CSS.

3. Specifically, another equivalent way to understand proximity based on CSS is as follows:
   - Transform the coefficients matrix $A$ to $DA$: $DA(i,j) = \begin{cases} 1 & A(i,j) > 0 \\ 0 & \text{else} \end{cases}$
   - The weight between $x_i$ and $x_j$ is:
\[ w_{ij} = \begin{cases} \frac{DA^i \cdot DA^j}{n} & i \neq j, \\ 0 & i = j \end{cases} \]

Obviously, the inner product \((DA^i \cdot DA^j)\) between \(DA^i\) and \(DA^j\) is equal to CSS \((x_i, x_j)\)'s cardinal \(|CSS(x_i, x_j)|\).

To illustrate the differences between our approaches for weight construction and others also using sparse representation, an example is given as follows. Assume that the coefficient matrix \(A\) of a data set with five objects obtained from solution coefficients of Eq. (6) is as the following 5 \times 5 matrix:

\[
A = \alpha_{ij} = \begin{bmatrix}
0 & 0.3 & 0.6 & 0.6 & -0.7 \\
0.4 & 0 & 0.5 & 0.6 & -0.6 \\
0.4 & 0.4 & 0 & -0.1 & -0.2 \\
-0.6 & -0.3 & 0.2 & 0 & 0.7 \\
-0.5 & 0.3 & 0.2 & 0.4 & 0
\end{bmatrix}
\]

According to the above introduction of different weight constructions:

1. \(SIS_{13} = 0.4, SIS_{12} = 0.2, DGC_{13} = 0.5\) and \(DGC_{12} = 0.35\), and these numbers show that the similarity between \(x_1\) and \(x_3\) is larger than that between \(x_1\) and \(x_2\). However, in our approaches using more entries in \(A\), \(CSS_{13} = 1/5 = 0.2, CSS_{12} = 2/5 = 0.4, COS_{13} = 0.24\) and \(COS_{12} = 0.98\) and these numbers show the different weights compared to the first group, where CSS and COS are the abbreviation of the above two proximity approaches, respectively.

2. \(DGC_{25} = 0.45, CSS_{25} = 0, COS_{25} = 0.16\), thus DGC mistakes the big negative coefficient \((\alpha_{25})\) as high similarity while CSS and COS both give lower similarity.

### 3.4. Algorithm description

Algorithm 1 describes the general procedure for spectral clustering of high-dimensional data, using sparse representation. The basic idea is to extract coefficients of sparse representation (Lines 1–4); construct a weight matrix using the coefficients (Line 5); and feed the weight matrix into a spectral clustering algorithm (Line 6) to find the best partitioning efficiently.

### Algorithm 1. General procedure for spectral clustering of high-dimensional data.

**Input:** high-dimensional training data set \(X = [x_1, x_2, \ldots, x_n] \in \mathbb{R}^m \times n\), where \(x_i = [x_{i1}, x_{i2}, \ldots, x_{im}]^T \in \mathbb{R}^m\) represents the \(i\)-th data object; the number of clusters \(K\).

**Parameter:** penalty coefficient \(\lambda\) for Lasso optimization

**Output:** cluster labels corresponding to each data object: \(c = [c_1, c_2, \ldots, c_n]\)

//standardize the input data for Lasso optimization
for each data object \(x_i \in X\) do

\[
// \text{Solve Eq. (6) with Lasso optimization}
\]

1. Set \(X_i = X_{(i)} = [x_{i1}, \ldots, x_{i-1}, x_i, x_{i+1}, \ldots, x_n];\)
2. \(\alpha_i^* = \arg \min \lambda ||\alpha_i||_1 + ||x_i - X_{(i)}\alpha_i||_2;\)
3. end

\[W = \text{ConstructWeightMatrix}(\alpha);\]
4. \(c = \text{SpectralClustering}(W);\)
5. return \(c.\)

The construct weight matrix () sub-routine can exploit any weight matrix construction method, such as those mentioned in Section 4. In particular, we describe the algorithm for computing the two newly proposed weight matrices, one based on the CSS (see Section 4.1) and the other based on COS of sparse coefficient vectors (see Section 4.2).

Algorithm 2 describes the procedure to construct the weight matrix according to the concept of CSS. To find the CSS of each pair of data objects (the two outermost loops), there is the need of checking the sparse coefficients of each remaining object to these two objects, so the time complexity of weight matrix constructions based on CSS is \(O(n^3).\)

**Algorithm 2. Construct weight matrix based on consistent sign set.**

**Input:** Coefficients for sparse representation \(\alpha\)

**Output:** Weight matrix \(W\)

1. for \(i = 1\) to \(n\) do
2. 
3. 
4. 
5. \(\text{if } j = i \text{ then } w_{ij} = 0;\)
6. 
7. 
8. 
9. 
10. 
11. 
12. 
13. 
14. end

Algorithm 3 describes the procedure to construct the weight matrix according to the COS of the sparse coefficients between each pair of items. The computation complexity for calculating the
COS of two vectors of length n is $O(n)$, and there are $O(n^2)$ pairs of data objects whose COS needs to be computed. Thus the complexity for COS-based weight matrix construction is $O(n^3)$.

**Algorithm 3.** Construct weight matrix based on similarity of coefficient vector.

**Input:** Coefficients for sparse representation $\alpha$

**Output:** Weight matrix $W$

1. for $i = 1$ to $n$ do
2. for $j = 1$ to $n$ do
3. if $j = i$ then $w_{ij} \leftarrow 0$;
4. else
5. $\text{cosine} = \frac{\alpha_i \cdot \alpha_j}{||\alpha_i||_2 ||\alpha_j||_2}$
6. if $\text{cosine} > 0$ then $w_{ij} \leftarrow \text{cosine}$;
7. else $w_{ij} \leftarrow 0$;
8. end
9. end
10. end

As shown in line 6 of Algorithm 1, after constructing the weight matrix $W$, we can use the classical spectral clustering algorithm [10] to discover the cluster structure of high-dimensional data.

The main characteristics of our proposed algorithm include the following: (1) compared to traditional graph construction induced from the Euclidean distance or other measures in the original high-dimensional space, the weight matrix is constructed after transforming the high dimensional data space into another space via sparse representation, which is expected to have better performance resulting from the superiority of compressed sensing [58] for high-dimensional data; (2) our graph construction based on consistent sign set or similarity of coefficient vector can simultaneously complete both the graph adjacency and weight matrix, while traditional graph constructions (such as $\epsilon$-ball neighborhood or k-nearest neighbors) complete the two tasks separately, which are interrelated and should not be separated [19]; (3) rather than existing graph constructions via sparse representation directly and independently applying the solution of $l_1$ optimization for each object in Eq. (6) to determine a row of the weight matrix, our approach considers the global information from the coefficients of the whole object set to calculate one element in the weight matrix.

### 4. Experimental results

In this section, we use experimental results to demonstrate the performance of our proposed approaches on real-world data sets using several effectiveness evaluations.
We select three data sets from the UCI machine-learning repository [60] and three face recognition data sets [61–63], which are well known in the machine learning and data mining research community. Table 1 lists a summary of these data sets.

In Yale and ORL face data sets, each image is transformed into a $32 \times 32$ pixel configuration using Matlab Image Processing Toolbox. In Yale B data set, which has 10 clusters, and each cluster has 585 image data, since the original size (5850) is too big, which leads to too much time consumption in clustering, we randomly select 60 images from the totally 585 images in each cluster. In all the data sets, each image is normalized to have unit norm.

We use interior-point method-based l1_ls_matlab tool [64] to solve Eq. (6) for each data object and then implement different weight matrices for algorithm 1 in Matlab to cluster each data set. Table 2 shows a summary of the proposed and baseline algorithms.

Since the true class labels of each data set are known, five commonly used external cluster validation metrics [66–68] are employed to evaluate the clustering results, namely clustering accuracy (CA) and normalized mutual information (NMI).

<table>
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<tr>
<th>Data set</th>
<th># Instance</th>
<th># Attributes</th>
<th># Classes</th>
<th>Source</th>
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<td>Heart</td>
<td>270</td>
<td>13</td>
<td>2</td>
<td>UCI</td>
</tr>
<tr>
<td>Image (Image segmentation)</td>
<td>2310</td>
<td>18</td>
<td>7</td>
<td>UCI</td>
</tr>
<tr>
<td>Yale</td>
<td>165</td>
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<td>15</td>
<td>[61]</td>
</tr>
<tr>
<td>Yale B</td>
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<td>1200</td>
<td>10</td>
<td>[62]</td>
</tr>
<tr>
<td>ORL Face</td>
<td>400</td>
<td>1024</td>
<td>40</td>
<td>[63]</td>
</tr>
<tr>
<td>Movement</td>
<td>360</td>
<td>90</td>
<td>15</td>
<td>UCI</td>
</tr>
</tbody>
</table>

Table 1. Summary of data sets.

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<th>Name</th>
<th>Description</th>
<th>Source</th>
<th>Role</th>
</tr>
</thead>
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<tr>
<td>CSS</td>
<td>Spectral clustering with weight matrix from consistent sign set</td>
<td>Section 3.1</td>
<td>Solution proposed</td>
</tr>
<tr>
<td>COS</td>
<td>Spectral clustering with weight matrix from cosine similarity of sparse coefficients</td>
<td>Section 3.2</td>
<td>Solution proposed</td>
</tr>
<tr>
<td>RBF</td>
<td>Spectral clustering with weight matrix from Gaussian RBF</td>
<td>[12]</td>
<td>Baseline</td>
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<tr>
<td>SIS</td>
<td>Spectral clustering with weight matrix from sparsity induced similarity measure</td>
<td>[59]</td>
<td>Baseline</td>
</tr>
<tr>
<td>DGC</td>
<td>Spectral clustering with weight matrix from $l_1$, Directed Graph Construction</td>
<td>[58]</td>
<td>Baseline</td>
</tr>
<tr>
<td>NN</td>
<td>Spectral clustering with weight matrix from non-negative sparsity induced similarity measure</td>
<td>[57]</td>
<td>Baseline</td>
</tr>
<tr>
<td>KM</td>
<td>$k$-means clustering</td>
<td>[65]</td>
<td>Baseline</td>
</tr>
</tbody>
</table>

Table 2. Summary of algorithms to be compared.

---

We use the matlab toolbox from: http://www.cad.zju.edu.cn/home/dengcai/Data/data.html
To illustrate the weight matrices from different approaches, we demonstrate the visual property of the proposed graph weight matrices in comparison with traditional ones in Figure 1, taking the Yale data set as an example. In Figure 1, each subfigure is a weight matrix with $N \times N$ (entries larger than the threshold is shown in white, otherwise black) and images from the same cluster are arranged together. These sparse representation-based graphs include consistent sign set (CSS), cosine similarity of coefficient vectors (COS), induced similarity measure (SIS), $l_1$ directed graph construction (DGC), nonnegative sparsity induced similarity measure (NN) and Gaussian RBF (RBF).

Since none of the original weight matrices constructed by the five approaches is sparse, we set threshold values (0.2 for COS; 0.388 for CSS; 0 for RBF ($\sigma$ is set 4 in RBF); 0.02 for the other three matrices) to get the best sparse matrices of different threshold values in Figure 1. A value larger than the threshold is shown in white, otherwise black. Normally, the clustering performance will be good if the weights between two objects from different clusters are little while weights from the same cluster are large. This comment can be equalized in the matrix with above arrangement that good matrix should be compact in diagonal position and sparse in other positions.

From Figure 1, we have the following observations: (1) matrices in all subfigures are compact in diagonal position; (2) the matrix of COS is sparser than others in lower left or upper right positions.
parts. This means that there are less inter-cluster adjacency connections in the COS than other graphs, so COS can encode more discriminating information and hence is more effective in spectral clustering than other traditional graphs; (3) CSS has a similar performance to SIS, DGC and NN Graph in Yale data set.

The clustering results obtained from the seven clustering algorithms with different evaluation metrics are reported in Tables 3 and 4, each of which corresponds to one evaluation metric. For each data set, the best results are in bold. All the numbers, except the last two rows in each table, represent the best clustering results using different lasso parameters ($\lambda$). The last two rows in each table present the average performance of each algorithm over all six data sets. Since the k-means clustering within spectral clustering is sensitive to initial centroids, we run spectral clustering 50 times for each case and report the mean and standard deviation (std).

From Tables 3 and 4, we can clearly see that, generally, CSS or COS algorithm gets the best clustering performance with all the two evaluation metrics. However, there are also some particular cases where CSS or COS does not get best result. For example, though NN gets the best CA on Yale B data sets, COS gets almost the same CA result as NN, that is, from 0.8937 to 0.8940; though NN also gets the best NMI for movement data set, COS gets best result in other metric on this data set. In particular, COS performs better than CSS with mean value of evaluation metrics, and the average standard deviation between 50 random tests of CSS is lowest for all metrics except CA.

Overall, for most data sets, CSS and COS show better performance than those baselines, which are robust across various external validation metrics. However, it is noticed that COS outperforms CSS in terms of all average mean metrics except CA, and CSS outperforms COS in

<table>
<thead>
<tr>
<th>Data set</th>
<th>CSS</th>
<th>COS</th>
<th>DGC</th>
<th>SIS</th>
<th>NN</th>
<th>BRF</th>
<th>KM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart</td>
<td>Mean</td>
<td>0.7704</td>
<td>0.8174</td>
<td>0.5852</td>
<td>0.7889</td>
<td>0.7519</td>
<td>0.7963</td>
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<td></td>
<td>(Std)</td>
<td>(0.0000)</td>
<td>(0.0000)</td>
<td>(0.0000)</td>
<td>(0.0000)</td>
<td>(0.0000)</td>
<td>(0.0882)</td>
</tr>
<tr>
<td>Image</td>
<td>Mean</td>
<td>0.7631</td>
<td>0.7921</td>
<td>0.7020</td>
<td>0.7820</td>
<td>0.7360</td>
<td>0.5335</td>
</tr>
<tr>
<td></td>
<td>(Std)</td>
<td>(0.0148)</td>
<td>(0.0323)</td>
<td>(0.0339)</td>
<td>(0.0341)</td>
<td>(0.0379)</td>
<td>(0.0305)</td>
</tr>
<tr>
<td>Yale</td>
<td>Mean</td>
<td>0.6823</td>
<td>0.7408</td>
<td>0.7178</td>
<td>0.7023</td>
<td>0.6417</td>
<td>0.6635</td>
</tr>
<tr>
<td></td>
<td>(Std)</td>
<td>(0.0432)</td>
<td>(0.0345)</td>
<td>(0.0414)</td>
<td>(0.0314)</td>
<td>(0.0412)</td>
<td>(0.0395)</td>
</tr>
<tr>
<td>Yale B</td>
<td>Mean</td>
<td>0.8572</td>
<td>0.8997</td>
<td>0.8320</td>
<td>0.8620</td>
<td>0.8940</td>
<td>0.6918</td>
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<tr>
<td></td>
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<td>(0.0713)</td>
<td>(0.0768)</td>
<td>(0.0635)</td>
<td>(0.0767)</td>
<td>(0.0226)</td>
</tr>
<tr>
<td>ORL face</td>
<td>Mean</td>
<td>0.7315</td>
<td>0.7570</td>
<td>0.7225</td>
<td>0.7243</td>
<td>0.6903</td>
<td>0.7314</td>
</tr>
<tr>
<td></td>
<td>(Std)</td>
<td>(0.0247)</td>
<td>(0.0218)</td>
<td>(0.0222)</td>
<td>(0.0244)</td>
<td>(0.0207)</td>
<td>(0.0252)</td>
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<tr>
<td>Movement</td>
<td>mean</td>
<td>0.5241</td>
<td>0.5472</td>
<td>0.5009</td>
<td>0.5183</td>
<td>0.5304</td>
<td>0.4874</td>
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<tr>
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<td>(Std)</td>
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<td>(0.0248)</td>
<td>(0.0193)</td>
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<td>(0.0232)</td>
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<tr>
<td>Average</td>
<td>Mean</td>
<td>0.7214</td>
<td>0.7580</td>
<td>0.6767</td>
<td>0.7296</td>
<td>0.7074</td>
<td>0.6506</td>
</tr>
<tr>
<td></td>
<td>(Std)</td>
<td>(0.0307)</td>
<td>(0.0298)</td>
<td>(0.0332)</td>
<td>(0.0288)</td>
<td>(0.0339)</td>
<td>(0.0235)</td>
</tr>
</tbody>
</table>

Table 3. Evaluation of all algorithms with CA as metric.
terms of all average standard metrics. It can be explained that CSS is more stable because its discretization may lower the variance of the pairwise of similarity, while COS get more generalized information of the pairwise of similarity leading to better average metrics but higher variance. Therefore, the choice between stability and quality should be taken into account when it is facing the clustering problem, in practice, using this kind of approach.

Finally, we plot the averages of the mean value and standard deviation (from the last two rows of the five tables), for comparing clustering algorithms, as shown in Figure 2.

<table>
<thead>
<tr>
<th>Data set</th>
<th>CSS</th>
<th>COS</th>
<th>DGC</th>
<th>SIS</th>
<th>NN</th>
<th>BRF</th>
<th>KM</th>
</tr>
</thead>
<tbody>
<tr>
<td>Heart Mean</td>
<td>0.2208</td>
<td>0.3149</td>
<td>0.0511</td>
<td>0.1791</td>
<td>0.0331</td>
<td>0.2712</td>
<td>0.2028</td>
</tr>
<tr>
<td>(Std)</td>
<td>(0.0000)</td>
<td>(0.0000)</td>
<td>(0.0000)</td>
<td>(0.0312)</td>
<td>(0.0000)</td>
<td>(0.1013)</td>
<td></td>
</tr>
<tr>
<td>Image Mean</td>
<td>0.7088</td>
<td>0.7451</td>
<td>0.5921</td>
<td>0.7319</td>
<td>0.6637</td>
<td>0.7451</td>
<td>0.6122</td>
</tr>
<tr>
<td>(Std)</td>
<td>(0.0071)</td>
<td>(0.0184)</td>
<td>(0.0171)</td>
<td>(0.0176)</td>
<td>(0.0357)</td>
<td>(0.0284)</td>
<td>(0.0437)</td>
</tr>
<tr>
<td>Yale Mean</td>
<td>0.7137</td>
<td>0.7815</td>
<td>0.7513</td>
<td>0.7641</td>
<td>0.6926</td>
<td>0.6989</td>
<td>0.6484</td>
</tr>
<tr>
<td>(Std)</td>
<td>(0.0211)</td>
<td>(0.0183)</td>
<td>(0.0203)</td>
<td>(0.0188)</td>
<td>(0.0198)</td>
<td>(0.0271)</td>
<td>(0.0342)</td>
</tr>
<tr>
<td>Yale B Mean</td>
<td>0.9008</td>
<td>0.9526</td>
<td>0.9202</td>
<td>0.9379</td>
<td>0.9510</td>
<td>0.7768</td>
<td>0.7858</td>
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<tr>
<td>(Std)</td>
<td>(0.0302)</td>
<td>(0.0360)</td>
<td>(0.0422)</td>
<td>(0.0356)</td>
<td>(0.0398)</td>
<td>(0.0224)</td>
<td>(0.0621)</td>
</tr>
<tr>
<td>ORL face Mean</td>
<td>0.8477</td>
<td>0.8688</td>
<td>0.8492</td>
<td>0.8547</td>
<td>0.8426</td>
<td>0.8512</td>
<td>0.8620</td>
</tr>
<tr>
<td>(Std)</td>
<td>(0.0116)</td>
<td>(0.0108)</td>
<td>(0.0118)</td>
<td>(0.0118)</td>
<td>(0.0109)</td>
<td>(0.0140)</td>
<td>(0.0157)</td>
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<tr>
<td>Movement Mean</td>
<td>0.5891</td>
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<td>0.5933</td>
<td>0.6000</td>
<td>0.6306</td>
<td>0.5741</td>
<td>0.5818</td>
</tr>
<tr>
<td>(Std)</td>
<td>(0.0128)</td>
<td>(0.0124)</td>
<td>(0.0140)</td>
<td>(0.0130)</td>
<td>(0.0173)</td>
<td>(0.0169)</td>
<td>(0.0180)</td>
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<tr>
<td>Average Mean</td>
<td>0.6635</td>
<td>0.7090</td>
<td>0.6262</td>
<td>0.6779</td>
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<td>0.6155</td>
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<tr>
<td>(Std)</td>
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<td>(0.0258)</td>
<td>(0.0181)</td>
<td>(0.0458)</td>
</tr>
</tbody>
</table>

Table 4. Evaluation of all algorithms with NMI as metric.

Figure 2. Error bar of different algorithms (a) CA, (b) NMI.
5. Conclusion

In this chapter, we present a study of spectral clustering based on sparse representation, using two novel weight matrix construction approaches to assess the consistency of two sparse vectors. This construction considers the global information of the solution coefficient vectors of two objects to analyze the similarity between these two objects rather than directly using the sparse coefficients, which only considers local information. Evaluation experiments on real-world data sets show that spectral clustering for high-dimensional data using our novel weight matrix construction exploiting global information outperforms direct k-means and spectral clustering approaches using Gaussian RBF, SIS, $l_1$-directed graph construction and non-negative SIS in five evaluation metrics (CA and NMI).

These results demonstrate a reliable performance of our algorithm and therefore promise wide applicability in practice. The findings also shed light on developing global solutions theories in the future work.

Figure 2 clearly demonstrates that COS and CSS algorithms outperform other algorithms, and COS is better than CSS on average. CSS obtains the least average value of standard deviation among all seven algorithms. The KM and DGC algorithms have comparable performance, which is usually worse than the other algorithms.

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