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# Study on density peaks clustering based on k-nearest neighbors and principal component analysis



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# ABSTRACT

Density peaks clustering (DPC) algorithm published in the US journal Science in 2014 is a novel clustering algorithm based on density. It needs neither iterative process nor more parameters. However, original algorithm only has taken into account the global structure of data, which leads to missing many clusters. In addition, DPC does not perform well when data sets have relatively high dimension. Especially, DPC generates wrong number of clusters of real-world data sets. In order to overcome the first problem, we propose a density peaks clustering based on k nearest neighbors (DPC-KNN) which introduces the idea of k nearest neighbors (KNN) into DPC and has another option for the local density computation. In order to overcome the second problem, we introduce principal component analysis (PCA) into the model of DPC-KNN and further bring forward a method based on PCA (DPC-KNN-PCA), which preprocesses high-dimensional data. By experiments on synthetic data sets, we demonstrate the feasibility of our algorithms and spectral clustering (SC) algorithm in accuracy. Experimental results show that our algorithms are feasible and effective.

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# 1. Introduction

Clustering, used mostly as an unsupervised learning method, is a major technique for data mining. The main aim of cluster analysis is to divide a given population into groups or clusters with common characteristics, since similar objects are grouped together, while dissimilar objects belong to different clusters. Clustering is useful in exploratory pattern-analysis, grouping, decision-making, and machine-learning situations, including data mining, document retrieval, image segmentation, and pattern classification [1]. Clustering methods are generally divided into five groups: hierarchical clustering, partitioning clustering, density-based clustering, gridbased clustering and model-based clustering [2]. Each method has its own strengths and weaknesses.

Density-based clustering [3–7] is represented by DBSCAN [3]. In density-based clustering, clusters are defined as areas of higher density than the remainder of the data set. Density-based clusters can have an arbitrary shape in the feature space. In addition, DB-SCAN does not require one to specify the number of clusters in the data a priori. However, it is very sensible to the user-defined

\* Corresponding author at: School of Computer Science and Technology, China University of Mining and Technology, Xuzhou 221116, China. Tel.: +86 15852494396. *E-mail address:* dingsf@cumt.edu.cn (S. Ding). parameter values, often producing very different clustering results in the data set even for slightly different parameter settings [2].

Like DBSCAN and the mean-shift method [8], density peaks clustering (DPC) algorithm [9] proposed by Rodriguez and Laio is able to detect non-spherical clusters and does not require one to specify the number of clusters. This method is robust with respect to the choice of  $d_c$  as the only parameter. DPC is based on the idea that cluster centers are characterized by a higher density than their neighbors and by a relatively large distance from points with higher densities. Several researches [10–15] have been going on around this method.

But DPC still has some defects. The local structure of data has not been taken into account in DPC when it calculates the local density. For example, DPC does not perform well when clusters have different densities. Having clusters of different densities is very common in data sets. The local density of DPC will lead to missing many clusters. Fig. 1 presents that clusters cannot be all detected with the local density of DPC. If p is small, two clusters in the lower-left corner are detected as a single cluster. However, if p is high, two clusters near the bottom are detected as a single cluster. In this case, DPC is not able to find clusters.

In order to overcome this problem, we propose a novel DPC based on k nearest neighbors (DPC-KNN). The proposed method makes use of the ideas of the k nearest neighbors for the local density computation.

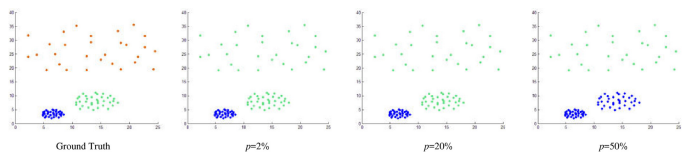


Fig. 1. DPC on these clusters of different densities.

In addition, it does a poor job of finding the clusters of highdimensional data. It may generate wrong number of clusters of real-world data sets. This is because many of the dimensions in high dimensional data are often irrelevant. These irrelevant dimensions can confuse DPC by hiding clusters in noisy data. Another reason is its overwhelming dependence on the distances between points. Two main quantities of DPC are both relevant to the distances. And for this reason, the problem, "the curse of dimensionality", is exacerbated. As the number of dimensions in a dataset increases, distance measures become increasingly meaningless [16]. Additional dimensions spread out the points until, in very high dimensions, they are almost equidistant from each other. More specific details are shown in Section 4. On the basis of the former, we further bring forward a method based on principal component analysis (DPC-KNN-PCA).

We test our algorithms on synthetic data sets to demonstrate their feasibility. In order to assess the performance of proposed algorithms, we compare proposed algorithms with other algorithms on some UCI data sets. Our algorithms have achieved satisfactory results in most data sets. The rest of this paper is organized as follows. In Section 2, we describe the principle of the DPC method, and introduce the k nearest neighbors and principal component analysis. In Section 3, we make a detailed description of DPC-KNN and DPC-KNN-PCA. In Section 4, we present experimental results in synthetic data sets and UCI data sets, then we analyze the performance of proposed algorithms. Finally, some conclusions and the intending work are given in the last section.

# 2. Related works

The proposed DPC-KNN is based on DPC and KNN. The proposed DPC-KNN-PCA is based on former theories and PCA. This section provides brief reviews of DPC, KNN, and PCA.

# 2.1. Density peaks clustering

Rodriguez and Laio proposed an algorithm published in the US journal Science. Its idea is that cluster centers are characterized by a higher density than their neighbors and by a relatively large distance from points with higher densities [9]. This method utilizes two important quantities: One is the local density  $\rho_i$  of each point *i*, and the other is its distance  $\delta_i$  from points of higher density. The two quantities correspond to two assumptions with respect to the cluster centers. One is that the cluster centers are surrounded by neighbors with a lower local density. The other is that they have relatively larger distance to the points of higher density. In the following, we will describe the computation of  $\rho_i$  and  $\delta_i$  in much more detail.

Assume that the data set is  $\mathbf{X}_{N \times M} = [\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_N]^T$ , where  $\mathbf{x}_i = [\mathbf{x}_{1i}, \mathbf{x}_{2i}, \mathbf{x}_{Mi}]$  is the vector with M attributes and N is the number of points. The distance matrix of the data set needs to be computed first. Let  $d(\mathbf{x}_i, \mathbf{x}_i)$  denote the Euclidean distance between the point

 $\boldsymbol{x}_i$  and the point  $\boldsymbol{x}_i$ , as follows:

$$d(\boldsymbol{x}_i, \boldsymbol{x}_j) = \|\boldsymbol{x}_i - \boldsymbol{x}_{j_2}\|$$
(1)

The local density of a point  $\mathbf{x}_i$ , denoted by  $\rho_i$ , is defined as:

$$\rho_{i} = \sum_{j} \chi \left( \mathbf{d} \left( \mathbf{x}_{i}, \mathbf{x}_{j} \right) - d_{c} \right)$$

$$\chi \left( x \right) = \begin{cases} 1, & x < 0 \\ 0, & x \ge 0 \end{cases}$$
(2)

where  $d_c$  is a cutoff distance.  $\rho_i$  is defined as the number of points that are adjacent to point  $\mathbf{x}_i$ . There is another local density computation in the code presented by Rodriguez and Laio. If the former is called a hard threshold, the latter will be called a soft threshold. Specifically,  $\rho_i$  is defined as a Gaussian kernel function, as follows:

$$\rho_i = \sum_j exp\left(-\frac{\mathrm{d}(\boldsymbol{x}_i, \boldsymbol{x}_j)^2}{d_c^2}\right)$$
(3)

where  $d_c$  is an adjustable parameter, controlling the weight degradation rate.

 $d_c$  is the only variable in Formulas (2) and (3). The process for selecting  $d_c$  is actually that for selecting the average number of neighbors of all points in data set. In the code,  $d_c$  is define as:

$$d_c = d_{N_d \times \frac{p}{100}} \tag{4}$$

where  $N_d = \binom{N}{2}$  and  $d_{N_d \times \frac{p}{100}} \in D = [d_1, d_1, \dots, d_{N_d}]$ . *D* is a set of all the distances between every two points in data set, which are sorted in ascending order. *N* denotes the number of points in data set.  $N_d \times \frac{p}{100}$  is the subscript of  $d_{N_d \times \frac{p}{100}}$ , where  $[\cdot]$  is the ceiling function and *p* is a percentage.

The computation of  $\delta_i$  is quite simple. The minimum distance between the point of  $\mathbf{x}_i$  and any other points with higher density, denoted by  $\delta_i$ , is defined as:

$$\delta_{i} = \begin{cases} \min_{j:\rho_{i}>\rho_{j}} (\mathbf{d}(\mathbf{x}_{i}, \mathbf{x}_{j})), & \text{if } \exists j \text{ s.t.} \rho_{i} > \rho_{j} \\ \max_{j} (\mathbf{d}(\mathbf{x}_{i}, \mathbf{x}_{j})), & \text{otherwise} \end{cases}$$
(5)

Only those points with relative high  $\rho_i$  and high  $\delta_i$  are considered as cluster centers. The points with high  $\rho_i$  and  $\delta_i$  value are also called as peaks that have higher densities than other points. A point is assigned to the same cluster as its nearest neighbor peak.

After cluster centers have been found, DPC assigns each remaining points to the same cluster as its nearest neighbors with higher density. A representation named as decision graph is introduced to help one to make a decision. This representation is the plot of  $\delta_i$  as a function of  $\rho_i$  for each point.

The following algorithm is a summary of DPC.

 Algorithm 1. DPC algorithm.

 Inputs:

 The samples  $X \in \mathbb{R}_{N \times M}$  

 The parameter  $d_c$  

 Outputs:

 The label vector of cluster index:  $y \in \mathbb{R}_{N \times 1}$  

 Method:

 Step 1: Calculate distance matrix according to Formula (1)

 Step 2: Calculate  $\rho_i$  for point *i* according to Formula (2) or (3)

 Step 3: Calculate  $\delta_i$  for point *i* according to Formula (5)

 Step 4: Plot decision graph and select cluster centers

 Step 5: Assign each remaining point to the nearest cluster center

Step 6: Return y

## 2.2. k Nearest neighbors

k Nearest neighbors (KNN) has already been exploited for classification [17–21]. This approach has been shown to be a powerful technique for density estimation [22], clustering [23–25] and other fields. As the name implies, the goal of this approach is to find the k-nearest neighbors of a sample among N samples. In general, the distances between points are achieved by calculating the Euclidean distance.

To compute the local density preferably, we use an idea based on k nearest neighbors (KNN).

We always assume that we are given *N* data points  $\mathbf{X} = [\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_N]^T$  which have been drawn in *M* dimensional space. As distance function between points we use the Euclidean distance, which is denoted by  $d(\cdot, \cdot)$ . It is required to compute the k nearest neighbors of a sample  $\mathbf{x}_i$  among  $\{\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_{i-1}, \mathbf{x}_{i+1}, \dots, \mathbf{x}_N\}$ . Sorting these distances in ascending order is to find the first K distances. The *k*th distance corresponds to the *k*th nearest neighbors of  $\mathbf{x}_i$  among  $\{\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_{i-1}, \mathbf{x}_{i+1}, \dots, \mathbf{x}_N\}$ . By kNN( $\mathbf{x}_i$ ) we denote the set of the k nearest neighbors of  $\mathbf{x}_i$  among  $\{\mathbf{x}_1, \mathbf{x}_2, \dots, \mathbf{x}_{i-1}, \mathbf{x}_{i+1}, \dots, \mathbf{x}_N\}$ . More specifics of the idea are discussed in Section 3.

The nearest neighbor step is simple to implement, though it scales in the worst case as  $O(N^2)$ , if performed without any optimizations. Some efficient methods such as K-D tree [26] or ball trees [27] can be used to compute the neighbors in  $O(N\log N)$  time.

#### 2.3. Principal components analysis

Principal components analysis (PCA) is a dimensionality reduction algorithm that can be used to significantly speed up unsupervised feature learning algorithm. The basic idea of PCA is to project the original data onto a lower-dimensional subspace, which highlights the principal directions of variation of the data.

The following steps describe this algorithm procedure.

- (1) Make each of the features have the same mean (zero) and variance.
- (2) Calculate the covariance matrix  $\Sigma$ .
- (3) Calculate the eigenvectors  $\boldsymbol{u}_i$  and the eigenvalues  $\lambda_i$  of  $\Sigma$ .
- (4) Sort these eigenvalues in decreasing order and stack the eigenvectors  $u_i$  corresponding to the eigenvalue  $\lambda_i$  in columns to form the matrix **U**.

# **3.** Density peaks clustering based on KNN and density peaks clustering based on KNN and PCA

There are still some defects in DPC. To solve these problems, we propose the following solutions.

#### 3.1. Density peaks clustering based on k nearest neighbors

Firstly, the local structure of data is not assessed by the local density in DPC. For this, we propose a novel DPC based on k nearest neighbors (DPC-KNN).

A shortcoming of the local density is that it is not sensitive to the local geometric of the data. Especially, when there is a great difference between the clusters in the density, there is a great difference between cluster centers on the local density. If  $d_c$  is so low that the distances between cluster centers are distinguished on decision graph, it is hard to select cluster centers. In our work, we introduce the idea of KNN into the calculation of the local density.

Let  $\mathbf{x}_i \in \mathbf{X}$ , d( $\cdot$ ,  $\cdot$ ) the Euclidean distance function, and the NN<sub>k</sub>( $\mathbf{x}_i$ ) be the *k*th nearest point to  $\mathbf{x}_i$  according to *d*, the *k*-nearest neighbors (kNN( $\cdot$ )) of  $\mathbf{x}_i$  is defined as:

$$kNN(\boldsymbol{x}_i) = \left\{ j \in X | d(\boldsymbol{x}_i, \boldsymbol{x}_j) \le d(\boldsymbol{x}_i, NN_k(\boldsymbol{x}_i)) \right\}$$
(6)

We can use  $kNN(x_i)$  to calculate the local density. This new local density is calculating the mean distance to k nearest neighbors, as follows:

$$\rho_{i} = exp\left(-\left(\frac{1}{k}\sum_{x_{j}\in kNN(\boldsymbol{x}_{i})}d(\boldsymbol{x}_{i},\boldsymbol{x}_{j})^{2}\right)\right)$$
(7)

where *k* is computed as a percentage (p) of the number of points *N*, so  $k = p \times N$ . Because a higher value of the local density means a higher density, so Formula (7) is a Gaussian kernel function as an inverse measure of the distance.

Then we can adopt the idea of KNN to the local density in DPC. The following algorithm is a summary of the proposed DPC-KNN.

Inputs:	
The s	amples $\boldsymbol{X} \in \mathbb{R}_{N \times M}$
The p	arameter p
Outputs	:
The l	abel vector of cluster index: $\boldsymbol{y} \in \mathbb{R}_{N \times 1}$
Method	
Step 1: 0	Calculate distance matrix according to Formula (1)
Step 2: 0	Calculate $\rho_i$ for point i according to Formula (7)
Step 3: 0	Calculate $\delta_i$ for point i according to Formula (5)
Step 4: I	Plot decision graph and select cluster centers
Step 5: A	Assign each remaining point to the nearest cluster center
Step 6: I	Return y

Complexity Analysis: Suppose N is the total number of points in data set. The complexity in calculating the similarity matrix is  $O(N^2)$ . DPC-KNN also needs  $O(N^2)$  to compute the local density. In addition, we cost O(Nlog N) in the sorting process with quick sort. For the progress to determine the cluster centers, we take no account of the time. As the complexity in assignment procedure is O(N), the total time complexity of our DPC-KNN method is  $O(N^2) + O(N^2) + O(N \log N) + O(N) + O(N) \sim O(N^2)$ .

# 3.2. Density peaks clustering based on k nearest neighbors and principal component analysis

In addition, DPC does not perform well when data has relatively high dimension. Especially, in real-world data sets, DPC generates wrong number of clusters of real-world data sets. On the basis of DPC-KNN, we further bring forward a method based on principal component analysis (DPC-KNN-PCA).

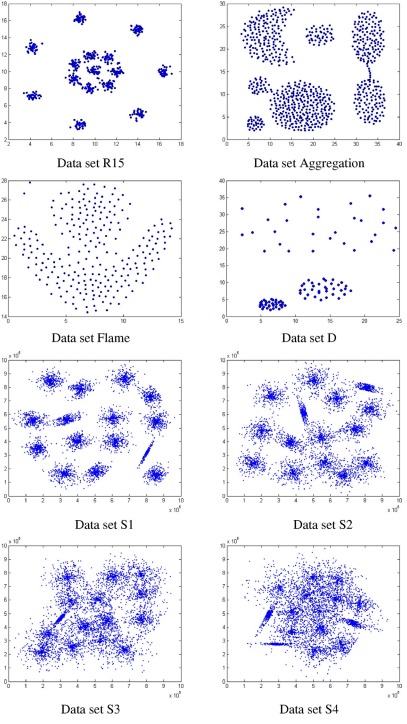


Fig. 2. Visualization of two-dimensional data sets.

Firstly, we make each of the features have the same mean (zero). Then, compute the matrix  $\Sigma$  as follows:

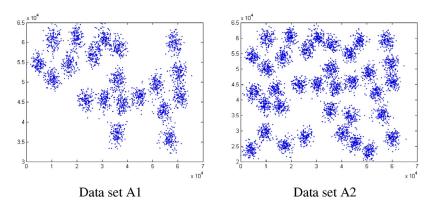
$$\Sigma = \frac{1}{N} \sum_{i=1}^{N} x_i x_i^T \tag{8}$$

If **x** has zero mean, then  $\Sigma$  is exactly the covariance matrix of **x**. We can compute the eigenvectors of  $\Sigma$ , and stack the eigenvectors

$$\mathbf{U} = \begin{bmatrix} | & | & | & | \\ \boldsymbol{u}_1 & \boldsymbol{u}_2 & \cdots & \boldsymbol{u}_M \\ | & | & | & | \end{bmatrix}$$
(9)

in columns to form the matrix **U**:

where  $\boldsymbol{u}_1$  is the principal eigenvector (corresponding to the largest eigenvalue),  $\boldsymbol{u}_2$  is the second eigenvector, and so on. Also, let  $\lambda_1, \lambda_1, \ldots, \lambda_M$  be the corresponding eigenvalues.



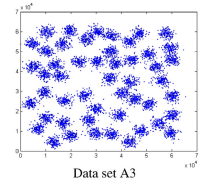


Fig. 2. (Continued).

We can represent  $\mathbf{x}$  in the  $(\boldsymbol{u}_1, \boldsymbol{u}_1, \ldots, \boldsymbol{u}_M)$ -basis by computing

$$\mathbf{x}_{rot} = \mathbf{U}^T \mathbf{x} = \begin{bmatrix} \mathbf{u}_1^T \mathbf{x} \\ \mathbf{u}_2^T \mathbf{x} \\ \vdots \\ \mathbf{u}_1^T \mathbf{x} \end{bmatrix}$$
(10)

The subscript "rot" comes from the observation that this corresponds to a reflection of the original data. We can compute  $\mathbf{x}_{rot}^{(i)} = \mathbf{U}^T \mathbf{x}^{(i)}$  for every point *i*. If we want to reduce this data to one dimension (the principal direction of variation of the data), we can set

$$\tilde{\boldsymbol{x}}^{(i)} = \boldsymbol{x}_{\text{rot},1}^{(i)} = \boldsymbol{u}_1^T \boldsymbol{x}^{(i)} \in \mathbb{R}$$
(11)

If **x** in  $\mathbb{R}^M$  and we want to reduce it to a *k* dimensional representation  $\mathbf{\tilde{x}} \in \mathbb{R}^k$  (where k < M), we would take the first *k* components of  $\mathbf{x}_{rot}$ , which correspond to the top *k* directions of variation. In other words, our definition of  $\mathbf{\tilde{x}}$  can also be arrived at by using an approximation to  $\mathbf{x}_{rot}$  where all but the first *k* components are zero, as follows:

$$\tilde{\mathbf{x}} = \begin{bmatrix} \mathbf{x}_{rot,1} \\ \vdots \\ \mathbf{x}_{rot,k} \\ 0 \\ \vdots \\ 0 \end{bmatrix} \approx \begin{bmatrix} \mathbf{x}_{rot,1} \\ \vdots \\ \mathbf{x}_{rot,k+1} \\ \vdots \\ \mathbf{x}_{rot,M} \end{bmatrix} = \mathbf{x}_{rot}$$
(12)

The reason that it drops the later components of  $\mathbf{x}_{rot}$  is that the first few components are considerably larger than the later components.

To decide how to set *k*, we will usually look at the percentage of variance retained for different values of *k*. Generally, let  $\lambda_1, \lambda_1, \cdots$ ,

 $\lambda_M$  be the eigenvalues of  $\Sigma$  (sorted in decreasing order), so that  $\lambda_i$  is the eigenvalue corresponding to the eigenvector  $\boldsymbol{u}_i$ . Then if we retain k principal components, the percentage of variance retained is given by:

$$\frac{\sum_{i=1}^{k} \lambda_i}{\sum_{i=1}^{M} \lambda_i} \tag{13}$$

In this paper, we pick the smallest value of *k* that satisfies

$$\frac{\sum_{i=1}^{K} \lambda_i}{\sum_{i=1}^{M} \lambda_i} \ge 0.99 \tag{14}$$

The following algorithm is a summary of the proposed DPC-KNN-PCA.

Algorithm 3. DPC-KNN-PCA algorithm.					
Inputs:					
The samples $X \in \mathbb{R}_{N \times M}$					
The parameter <i>p</i>					
Outputs:					
The label vector of cluster index: $\boldsymbol{y} \in \mathbb{R}_{N \times 1}$					
Method:					
Step 1: Make all of the features have the same mean (zero) and variance					
Step 2: Compute the covariance matrix $\Sigma$ according to Formula (8)					
Step 3: Compute the eigenvectors $\boldsymbol{u}_i$ and the eigenvalues $\lambda_i$ of $\Sigma$					
Step 4: Reduce the data and keep 99 % principal component to Formula (14)					
Step 5: Calculate distance matrix according to Formula (1)					
Step 6: Calculate $\rho_i$ for point <i>i</i> according to Formula (7)					
Step 7: Calculate $\delta_i$ for point <i>i</i> according to Formula (5)					
Step 8: Plot decision graph and select cluster centers					
Step 9: Assign each remaining point to the nearest cluster center					
Step 10: <b>Return y</b>					

Complexity Analysis: Suppose *N* is the total number of points in data set and *M* is the number of features of each point. Covariance matrix computation is  $O(M^2N)$ . Its eigenvalues decomposition is. So,  $O(M^3)$  the complexity of PCA is  $O(M^3 + M^2N)$ . The total time complexity of our DPC-KNN-PCA method is  $O(M^3 + M^2N + N^2)$ .

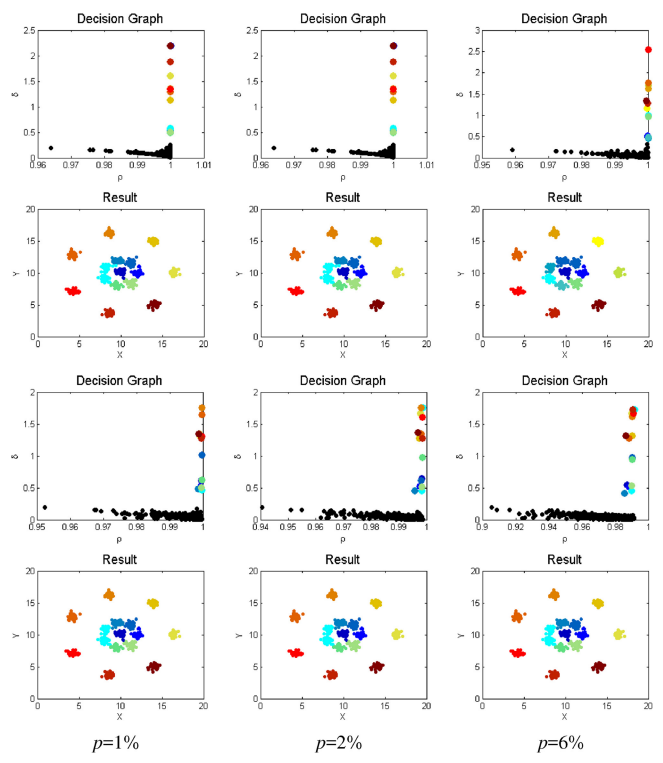


Fig. 3. DPC-KNN-PCA on R15 set with different values of *p*.

# 4. Experiments and results

In this section, we will test the performance of DPC-KNN-PCA through two types of the experiments. By experiments on synthetic data sets, we demonstrate the feasibility of the algorithm. By experiments on real-world data sets, we compared this algorithm with k-means algorithm, spectral clustering (SC) algorithm in accuracy.

We do experiments in a work station with a core i7 DMI2-Intel 3.6 GHz processor and 18 GB RAM running MATLAB 2012B. We run k-means algorithm, SC algorithm, 10 times in real-world data sets. This paper measures the similarity between data points with the famous Euclidean distance, which is used widely to measure the similarity of spatial data, as shown Formula (1). In DPC-KNN and DPC-KNN-PCA, we select the parameter *p* from [0.1% 0.2% 0.5% 1% 2% 6%]. In DPC, the parameter *d<sub>c</sub>* is also selected from [0.1% 0.2%

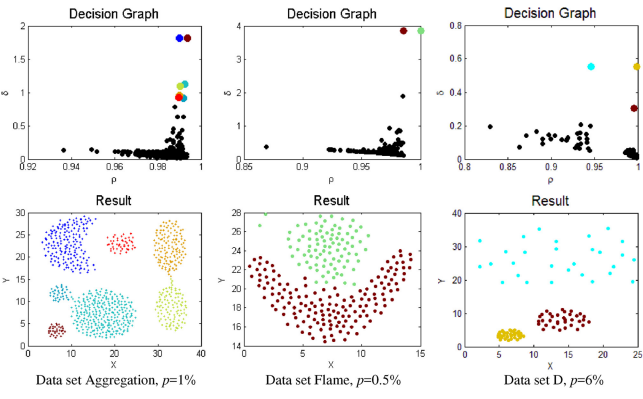


Fig. 4. DPC-KNN-PCA on Aggregation and Flame sets.

0.5% 1% 2% 6%]. The kernel of spectral clustering (SC) algorithm is the Gaussian kernel. In SC, we select the parameter  $\delta$  from [0.5 1 2 3 4].

# 4.1. Experiments on synthetic data sets

We test the performance of our algorithms on synthetic data sets. The synthetic data sets are two dimensional, which makes things easy from the visualization point of view. Because the performances of DPC-KNN and DPC-KNN-PCA are similar in 2dimensional data sets, we only test the performance of DPC-KNN-PCA.

#### 4.1.1. Synthetic data sets

Our algorithm is tested by 11 data sets whose geometric shapes are shown in Fig. 2. The first data set, R15 [28] is generated as 15 similar 2-D Gaussian distributions that are positioned in rings. The second data set, aggregation [29] consists of the seven perceptually distinct groups of points where there are non-Gaussian clusters. The third data set, Flame [30], is of different size and shape. The fourth data set, D, is of clusters of different densities. S1, S2, S3 and S4 [31] are four 2-D data sets with varying complexity in terms of spatial data distributions: i.e., in S1 the overlap is the smallest, whereas in S4 the overlap is the greatest. A1, A2 and A3 [32] are two-dimensional sets with varying number of circular clusters (M = 20, 35, 50). We demonstrate the power of these algorithms on these test cases.

# 4.1.2. The evaluation of clustering results in synthetic data sets

There are 15 clusters, 600 points in R15. In this case, our approaches are robust respect to parameter *p*. Fig. 3 presents a lot of clusters found by our algorithms with different *p* and selections of cluster centers. As shown, DPC-KNN-PCA gets perfect results, except when p = 0.5%.

There are 2 clusters, 240 points in Aggregation set and are 7 clusters, 788 points in Flame set. The two data sets consist of some

clusters that are of different size and shape. D set has 2 clusters and 97 points. Its clusters are of different densities. Clustering results proposed by DPC-KNN-PCA have been given in Fig. 4.

The data sets S1 to S4 are two-dimensional sets with varying complexity in terms of spatial data distributions. The data sets have 5000 points around 15 clusters with a varying degree of overlap. As shown in Fig. 5, the performance of DPC-KNN-PCA is perfect for data sets with varying complexity.

A1, A2 and A3 sets are large data sets with varying number of clusters. A1 has 3000 points around 20 clusters. A2 has 5250 points around 35 clusters. There are 7500 points and 50 clusters on A3. We demonstrate the robustness of the DPC-KNN-PCA for the quantity, as shown in Fig. 6.

As these experiments illustrate our algorithms are very effective in finding clusters of arbitrary shape, density, distribution and number.

#### 4.2. Experiments on real-world data sets

The performances of our algorithms are compared with classical methods (k-means algorithm and spectral clustering algorithm).

#### 4.2.1. Real-world data sets

N

The data sets used in the experiments are all from the UCI Machine Learning Repository, which include Iris, LED digits, Seeds, Heart, Pen-based digits, Waveform, Sonar. The details of those data sets are given in Table 1.

# 4.2.2. Quality of the clustering results

This paper uses clustering accuracy (ACC) [33] to measure the quality of the clustering results. For *N* distinct samples  $\mathbf{x}_i \in \mathbb{R}^j$ ,  $y_i$  and  $c_i$  are the inherent category label and the predicted cluster label of  $\mathbf{x}_i$ , the calculation formula of ACC is

$$ACC = \sum_{i=1}^{N} \delta(\mathbf{y}_i, map(\mathbf{c}_i)) / N$$
(15)

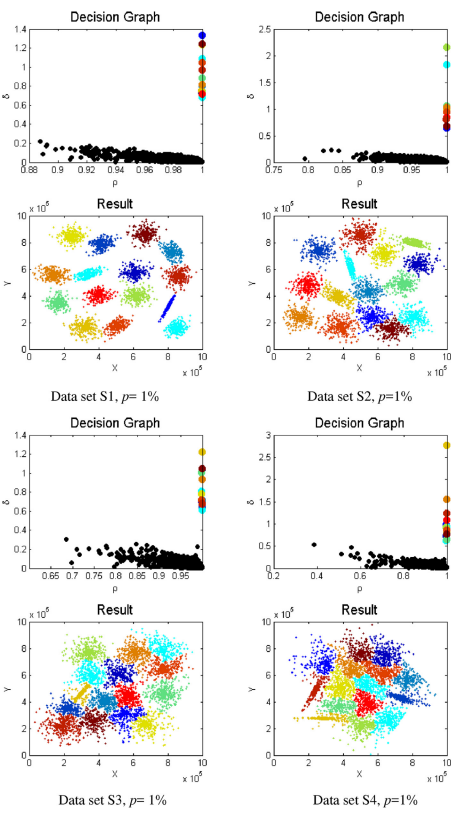


Fig. 5. DPC-KNN-PCA on the S1, S2, S3, and S4 sets.

where  $map(\cdot)$  maps each cluster label to a category label by the Hungarian algorithm [34] and this mapping is optimal, let  $\delta(y_i, c_i)$  equal to 1 if  $y_i = c_i$  or equals to 0 otherwise. The higher the values of the ACC are, the better the clustering performance will be.

The comparison of these algorithms is shown in Table 2. In Table 2, the symbol – means that the algorithm cannot work in the data set.

There are less than ten features in the first three data sets. And the last two data sets have more than 10 features. As shown

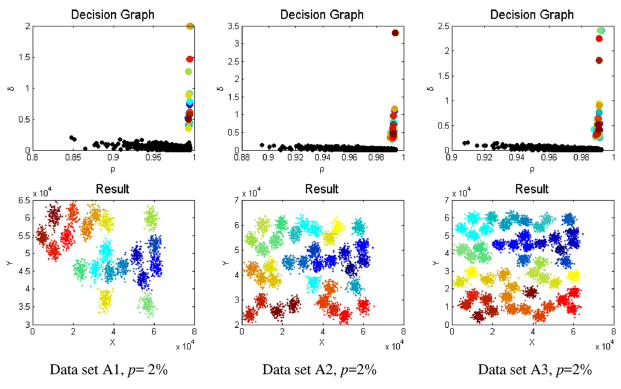


Fig. 6. DPC-KNN-PCA on the A1, A2, and A3 sets.

Table 1The details of UCI data sets.

Data sets	Cluster	Dimension	Ν
Iris	3	4	150
LED digits	10	7	500
Seeds	3	7	210
Heart	2	13	270
Pen-based digits	10	16	109,962
Waveform	3	21	5000
Sonar	2	60	208

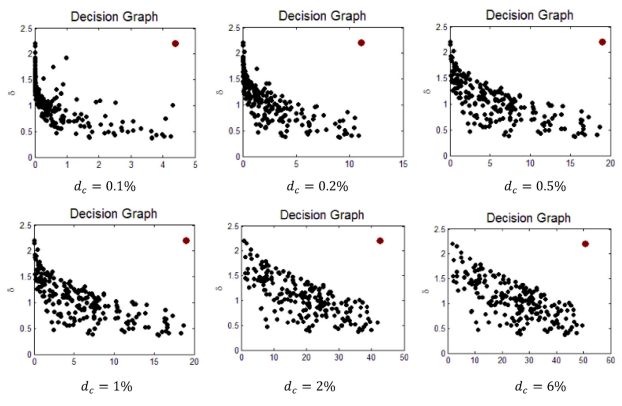
in Table 2, DPC-KNN outperforms others in low-dimensional data sets such as Iris, LED digits and Seeds. However, DPC-KNN-PCA has a better performance compared to DPC-KNN in relatively high-dimensional data sets. To be specific, the higher the features of the data set are, the greater advantage DPC-KNN-PCA has over

Table 2			
	The performance comparison of proposed algorithms.		

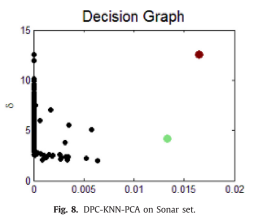
DPC-KNN. It is obvious that DPC-KNN-PCA algorithm has achieved gratifying results in most data sets.

Then, the reason that produced above results is discussed. DPC-KNN, without missing any information of features, performed better than DPC-KNN-PCA in low-dimensional data set. Due to the phenomenon of the so called curse of dimensionality, the similarity between samples becomes meaningless in high dimensional data spaces. Although similarity between high dimensional samples is not meaningful, similarity according to subsets of attributes is still meaningful. PCA not only reduces the dimensionality of the data, but also maintains as much information as possible. When some data sets have relatively high dimensions, DPC does a poor job of finding the clusters, which we need to pay extra attention to. For example, Sonar data set has relatively high dimensions compared to the number of samples. Fig. 7 shows only one cluster center found by DPC with different  $d_c$  on decision graph. In this case, it is unacceptable that we were incapable of making the right choices. DPC-KNN-PCA has a favorable performance compared to the

Data sets	Accuracy	DPC	DPC-KNN	DPC-KNN-PCA	SC	k-means
Iris	Mean	0.94	0.96	0.88	$0.8867~\pm~0$	0.8560 ± 0.097
	Parameter	$d_{\rm c} = 0.1\%$	p = 1%	p = 4%	$\delta = 0.5$	
LED digits	Mean	_	0.7460	0.6700	$0.6360~\pm~0.0552$	$0.5208~\pm~0.0713$
	Parameter		p = 6%	p = 6%	$\delta = 0.5$	
Seeds	Mean	0.8952	0.9143	0.9143	$0.9048~\pm~0$	$0.8905~\pm~0$
	Parameter	$d_c = 1\%$	p = 2%	p = 2%	$\delta = 1$	
Heart	Mean	-	0.8111	0.8259	$0.7963~\pm~0$	$0.7166~\pm~0.0640$
	Parameter		p = 1%	p = 6%	$\delta = 4$	
Pen-based digits	Mean	-	0.7618	0.7623	$0.7177 ~\pm~ 0.0276$	$0.7004~\pm~0.0466$
	Parameter		p = 1%	p = 0.2%	$\delta = 3$	
Waveform	Mean	0.5676	0.5840	0.6452	$0.5054~\pm~0$	$0.5012~\pm~0$
	Parameter	$d_c = 0.5\%$	p = 0.2%	p = 0.1%	$\delta = 0.5$	
Sonar	Mean	-	-	0.6442	$0.5433~\pm~0$	$0.5433 ~\pm~ 0.0124$
	Parameter			p = 1%	$\delta = 2$	



**Fig. 7.** DPC on Sonar set with different values of  $d_c$ .



original algorithm, as shown Fig. 8. In consequence, DPC-KNN-PCA outperforms DPC-KNN in high-dimensional data sets.

# 5. Conclusions

DPC-KNN has another option based on k nearest neighbors (KNN) for the local density computation. On the basis of DPC-KNN, a method based on principal component analysis (DPC-KNN-PCA) is presented to improve the performance of the former on real-world data sets. In this paper, presented algorithms show the power in some synthetic data sets. Besides the good feasibility, proposed algorithms get better clustering performances compared to classical methods (k-means algorithm and spectral clustering algorithm) and DPC in UCI data sets. In low-dimensional data sets, DPC-KNN outperforms others. And DPC-KNN-PCA has achieved gratifying results in relatively high-dimensional data sets.

However, the proposed algorithm does not perform well when there is a collection of points forming vertical streaks in data set. Future research is to improve the performance of DPC-KNN-PCA algorithm on manifold data set.

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