The SpectACl of Nonconvex Clustering: A Spectral Approach to Density-Based Clustering

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Abstract

When it comes to clustering nonconvex shapes, two paradigms are used to find the most suitable clustering: minimum cut and maximum density. The most popular algorithms incorporating these paradigms are Spectral Clustering and DBSCAN. Both paradigms have their pros and cons. While minimum cut clusterings are sensitive to noise, density-based clusterings have trouble handling clusters with varying densities. In this paper, we propose SpectACl: a method combining the advantages of both approaches, while solving the two mentioned drawbacks. Our method is easy to implement, such as Spectral Clustering, and theoretically founded to optimize a proposed density criterion of clusterings. Through experiments on synthetic and real-world data, we demonstrate that our approach provides robust and reliable clusterings.

1 Introduction

Despite being one of the core tasks of data mining, and despite having been around since the 1930s (Driver and Kroeber 1932; Klimek 1935; Tryon 1939), the question of clustering has not yet been answered in a manner that doesn’t come with innate disadvantages. The paper that you are currently reading will also not provide such an answer. Several advanced solutions to the clustering problem have become quite famous, and justly so, for delivering insight in data where the clusters do not offer themselves up easily.

Spectral Clustering (Dhillon, Guan, and Kulis 2004) provides an answer to the curse of dimensionality inherent in the clustering task formulation, by reducing dimensionality through the spectrum of the similarity matrix of the data. DBSCAN (Ester et al. 1996) is a density-based clustering algorithm which has won the SIGKDD test of time award in 2014. Both Spectral Clustering and DBSCAN can find non-linearly separable clusters, which trips up naive clustering approaches; these algorithms deliver good results. In this paper, we propose a new clustering model which encompasses the strengths of both Spectral Clustering and DBSCAN; the combination can overcome some of the innate disadvantages of both individual methods.

For all their strengths, even the most advanced clustering methods nowadays still can be tripped up by some pathological cases: datasets where the human observer immediately sees what is going on, but which prove to remain tricky for all state-of-the-art clustering algorithms. One such example is the dataset illustrated in Figure 1: we will refer to it as the two circles dataset. It consists of two noisy separated concentric circles, each encompassing the same number of observations. As the leftmost plot in Figure 1 shows, Spectral Clustering does not at all uncover the innate structure of the data: two clusters are discovered, but both incorporate about half of each circle. It is well-known that Spectral Clustering is highly sensitive to noise (Bojchevski, Matkovic, and Günnemann 2017, Figure 1); if two densely connected communities are additionally connected to each other via a narrow bridge of only a few observations, Spectral Clustering runs the risk of reporting these communities plus the bridge as a single cluster, whereas two clusters plus a few noise observations (outliers) would be the desired outcome. The middle plots in Figure 1 shows how DBSCAN (with minpts set to 25 and 26, respectively) fails to realize that there are

\[\text{minpts}\]  

The scikit-learn clustering documentation (cf. https://scikit-learn.org/stable/modules/clustering.html) shows how SC and DBSCAN can succeed on this dataset, but that version contains barely any noise (scikit’s noise parameter set to 0.05, instead of our still benign 0.1).
two clusters. This is hardly surprising, since DBSCAN is known to struggle with several clusters of varying density, and that is exactly what we are dealing with here: since both circles consist of the same number of observations, the inner circle is substantially more dense than the outer circle. The rightmost plot in Figure 1 displays the result of the new clustering method that we introduce in this paper, SPECTACL (Spectral Averagely-dense Clustering): it accurately delivers the clustering that represents the underlying phenomena in the dataset.

1.1 Main Contributions

In this paper, we provide SPECTACL, a new clustering method which combines the benefits of Spectral Clustering and DBSCAN, while alleviating some of the innate disadvantages of each individual method. Our method finds clusters having a large average density, where the appropriate density for each cluster is automatically determined through the spectrum of the weighted adjacency matrix. Hence, SPECTACL does not suffer from sensitivity to the minPts parameter as DBSCAN does, and unlike DBSCAN it can natively handle several clusters with varying densities. As in the Spectral Clustering pipeline, the final step of SPECTACL is an embedding postprocessing step using $k$-means. However, unlike Spectral Clustering, we demonstrate the fundamental soundness of applying $k$-means to the embedding step in SPECTACL: from SPECTACL’s objective function we derive an upper bound by means of the eigenvector decomposition; we derive that the optimization of our upper bound is equal to $k$-means on the eigenvectors. Our Python implementation, and the data generating and evaluation script, are publicly available.

2 A Short Story of Clustering

A formal discussion of the state of clustering requires some notation. We assume that our data is given by the matrix $D \in \mathbb{R}^{n \times r}$. The data represents $m$ points $D_j$, for $1 \leq j \leq m$ in the $n$-dimensional feature space. For every point, we denote with $N_\epsilon(j) = \{l \mid \|D_j - D_l\| < \epsilon\}$ its $\epsilon$-neighborhood.

Given an arbitrary matrix $M$, we denote with $\|M\|$ its Frobenius norm. If $M$ is diagonalizable, then we denote its truncated eigendecomposition of rank $d$ as $M \approx V^{(d)} \Lambda^{(d)} V^{(d)\top}$. The matrix $\Lambda$ is a $d \times d$ diagonal matrix, having the eigenvalues of $M$ on its diagonal, which have the $d$-largest absolute values: $|\Lambda_1| \geq \ldots \geq |\Lambda_d|$. The matrix $V^{(d)}$ concatenates the corresponding eigenvectors.

We write 1 to represent a matrix having all elements equal to 1. The dimension of such a matrix can always be inferred from the context. We also use the shorthand $\mathbb{I}^m$ for the set of all binary partition matrices. That is, $M \in \mathbb{I}^m$ if and only if $M \in \{0, 1\}^{m \times l}$ and $|M_{j\cdot}| = 1$ for all $j$.

2.1 $k$-means

If there is one algorithm which comes to mind when thinking about clustering, it is likely the $k$-means algorithm (Lloyd 1982). The popularity of $k$-means is due to its simple and effective yet theoretically founded optimization by alternating minimization, and the intuitively easily graspable notion of clusters by the within-cluster point scatter. The underlying assumption for this kind of clustering is that points in one cluster are similar to each other on average. As fallout, this imposes the undesired constraint that clusters are separated by a Voronoi tessellation and thus, have convex shapes.

We denote the objective of $k$-means in its matrix factorization variant:

$$\min_{Y, X} \|D - Y X^\top\|^2 \text{ s.t. } Y \in \mathbb{I}^{m \times r}, X \in \mathbb{R}^{n \times r}.$$  

Here, the matrix $Y$ indicates the cluster memberships, i.e., $Y_{js} = 1$ if point $j$ is assigned to cluster $s$ and $Y_{js} = 0$ otherwise. The matrix $X$ represents the cluster centroids.

The objective is nonconvex, but convex if one of the matrices is fixed. The optimum with respect to the matrix $X$ for fixed cluster assignments $Y$ is attained at $X = D^\top Y(Y^\top Y)^{-1}$. This is a matrix formulation of the simple notion that each cluster centroid is equal to the average of points in the cluster (Bauckhage 2015).

The objective of Equation (1) is equivalent to a trace maximization problem (Ding et al. 2006), which derives from the definition of the Frobenius norm by means of the trace

$$\max_Y \text{tr}(Y^\top D D^\top Y (Y^\top Y)^{-1}) \text{ s.t. } Y \in \mathbb{I}^{m \times r}.$$  

Note that this objective formulation of $k$-means depends only on the similarities of data points, expressed by the inner product of points. This is a stepping stone to the application of kernel methods and the derivation of nonconvex clusters.

2.2 Graph Cuts

The restriction to convex clusters can be circumvented by a transformation of the data points into a potentially higher-dimensional Hilbert space, reflected by a kernel matrix (Schölkopf, Smola, and Müller 1998). Such a representation based only on the similarity of points introduces an interpretation of the data as a graph, where each data point corresponds to a node, and the denoted similarities are the weights of the corresponding edges. In this view, a cluster is identifiable as a densely connected component which has only weak connections to points outside of the cluster. In other words, if we cut the edges connecting the cluster to its outside, we strive to cut as few edges as possible. This is known as the minimum cut problem. Given a possibly normalized edge weight matrix $W$, the cut is:

$$\text{cut}(Y; W) = \sum_{s=1}^r Y_s^\top W (1 - Y_s).$$  

However, minimizing the cut often returns unfavorable, unbalanced clusterings, where some clusters encompass only a few or single points. Therefore, normalizing constraints have been introduced. One of the most popular objective functions incorporating this normalization is the ratio cut (Hagen and Kahng 1992). The objective of ratio cut is given as:

$$\text{RCut}(Y; W) = \sum_{s=1}^r \frac{Y_s^\top W (1 - Y_s)}{|Y_s|}.$$  

Minimizing the ratio cut is equivalent to a trace maximization problem (Dhillon, Guan, and Kulis 2004; Ding, He, and Simon 2005), given as:

$$\max_Y \text{tr}(Z^T (-L)Z)$$  \hspace{1cm} (5)

$$\text{s.t. } Z = Y (Y^T Y)^{-1/2}, Y \in \mathbb{R}^{m \times r},$$

where $L = \text{diag}(W1) - W$ is known as the difference Laplacian (Chung 1997). Note that a normalization of $W$ would change the matrix $L$ to another famous graph Laplacian: the symmetric or random walk Laplacian (Chung 1997).

The matrix $-L$ is negative semi-definite, having eigenvalues $0 = \lambda_1 \geq \ldots \geq \lambda_m$. We observe that substituting the matrix $-L$ from Equation (5) with the positive semi-definite matrix $|\lambda_m|I - L$ does not change the objective. Therefore, ratio cut is equivalent to kernel $k$-means, employing the kernel matrix $|\lambda_m|I - L$.

### 2.3 Spectral Clustering

According to the Ky Fan theorem (Fan 1949), the maximum of Equation (5) with respect to the matrix $Z \in \mathbb{R}^{m \times r}$, having orthogonal columns, is attained at the eigenvectors of the $r$ largest eigenvalues of $f |\lambda_m|I - L$, i.e., $Z = V^{(r)}$. These relaxed results then need to be discretized in order to obtain a binary cluster indicator matrix $Y$. Most often, $k$-means clustering is applied for that purpose.

The eigenvectors of eigenvalue zero from the Laplacian indicate connected components, which supports a theoretic justification of the embedding given by the eigenvectors $V^{(r)}$. Hence, Spectral Clustering is backed up by the thoroughly studied theory of graph Laplacians. Spectral Clustering is in general described by the following three steps:

1. choose the representation of similarities by the weight matrix $W$, calculate the Laplacian $L = \text{diag}(W1) - W$;
2. compute the truncated eigendecomposition of the translated Laplacian $|\lambda_m|I - L \approx V^{(r+1)} \Lambda^{(r+1)} V^{(r+1)^T}$;
3. compute a $k$-means clustering on the eigenvectors $V^{(r+1)}$ (excluding the first eigenvector) returning $r$ clusters.

The weighted adjacency matrix is usually generated by a $k$-nearest neighbor graph. Given $k$, one way to define the weighted adjacency matrix is $W = 1/2(A + A^T)$, where $A$ is the adjacency matrix of the $k$-nearest neighbor graph. The other variant, which is less often used, is the determination of $W$ by the $\epsilon$-neighbor graph. In this variant, we have $W_{ij} = 1$ if and only if $i \in N_\epsilon(j)$ and $W_{jj} = 0$.

It is advised to employ a normalization of the weighted adjacency matrix, e.g., the symmetric normalization $\tilde{W} = \text{diag}(W1)^{-1/2} W \text{diag}(W1)^{-1/2}$. The corresponding Laplacian is the symmetrically normalized Laplacian $L_{sym} = I - \tilde{W}$. For a more detailed discussion of Spectral Clustering methods, see (Von Luxburg 2007).

Spectral Clustering is a popular technique, since the truncated eigendecomposition is efficiently computable for large-scale sparse matrices (Saad 2011), and efficient to approximate for general matrices by randomized methods (Halko, Martinsson, and Tropp 2011). However, Spectral Clustering also has multiple shortcomings. One of the drawbacks of this pipeline of methods is the uncertainty about the optimized objective. Although there is a relation to the rational cut objective, as outlined, there are examples where the optimal solution of rational cut and the solution of Spectral Clustering diverge (Guattery and Miller 1998). Related to this problem is the somewhat arbitrary choice to employ $k$-means to binarize the relaxed result. In principle, also other discretization methods could and have been applied (Von Luxburg 2007), yet there is no theoretical justification for any of these.

### 2.4 The Robustness Issue

Beyond theoretical concerns, an often reported issue with Spectral Clustering results is the noise sensitivity. This might be attributed to its relation with the minimum cut paradigm, where few edges suffice to let two clusters appear as one.

The state-of-the-art in Spectral Clustering research includes two approaches aiming at increasing robustness towards noise. The one is to incorporate ideas from density-based clustering, such as requiring that every node in a cluster has a minimal node degree (Bojchevski, Matkovic, and Günnemann 2017). Minimizing the cut subject to the minimal degree constraint resembles finding a connected component of core points, as performed by DBSCAN. The drawback of this approach is that it introduces a new parameter to Spectral Clustering, influencing the quality of the result. The other is to learn the graph representation simultaneously with the clustering (Bojchevski, Matkovic, and Günnemann 2017; Kang et al. 2018; Nie et al. 2017). This is generally achieved by alternating updates of the clustering and the graph representation, requiring the computation of a truncated eigen-decomposition in every iteration step. This results in higher computational costs. A final alternative is the approach based on Dominant Sets, as outlined in (Hou, Gao, and Li 2016). This approach is a sequential method, iteratively deriving the single best cluster and reducing the dataset accordingly, unlike SPECTACL, which simultaneously optimizes all clusters.

Of these newer methods, we compare our method against the Robust Spectral Clustering algorithm (Bojchevski, Matkovic, and Günnemann 2017), since it incorporates both the notion of density and the learning of the graph representation.

### 3 Spectral Averagely-Dense Clustering

We propose a cluster definition based on the average density (i.e., node degree) in the subgraph induced by the cluster. Let $W$ be the adjacency matrix. Since we are interested in deviations of the nodes’ degree, we employ the $\epsilon$-neighborhood graph to determine $W$. We strive to solve the following problem, maximizing the average cluster density:

$$\max_{Y \in \mathbb{R}^{m \times r}} \text{tr}(Y^T W Y (Y^T Y)^{-1}) = \sum_s \delta(Y_s, W).$$  \hspace{1cm} (6)

The objective function returns the sum of average node degrees $\delta(Y_s, W)$ in the subgraph induced by cluster $s$, i.e.:

$$\delta(Y_s, W) = \frac{Y_s^T W Y_s}{\|Y_s\|^2} = \frac{1}{\|Y_s\|^2} \sum_{j: Y_{js} = 1} W_{jj},$$  \hspace{1cm} (7)
Note that our Objective (6) is equivalent to minimum cut if the matrix \(W\) is normalized. In this case, the Laplacian is given as \(L = I - \tilde{W}\), and subtracting the identity matrix from Equation (6) does not change the objective.

We derive a new relationship for the solution of Objective (6) and the spectrum of the adjacency matrix. Thereby, we establish a connection with the application of k-means to the spectral embedding. As a result, our method encompasses the same steps as Spectral Clustering, and hence it can be efficiently computed even for large scale data.

### 3.1 Dense Clusters and Projected Eigenvectors

The function \(\delta\) from Equation (7) is also known as the Rayleigh quotient. The values of the Rayleigh quotient depend on spectral properties of the applied matrix. As such, the density \(\delta(y, W) \in [\lambda_m, \lambda_1]\) is bounded by the smallest and largest eigenvalues of the matrix \(W\) (Collatz 1978). The extremal densities are attained at the corresponding eigenvectors. A simple calculation shows that the eigenvectors \(V_1, \ldots, V_d\) to the \(d\)-largest eigenvalues span a space whose points have a minimum density \(\delta(y, W) \geq \lambda_d\). Thus, the projection of \(W\) onto the subspace spanned by the first eigenvectors reduces the dimensionality of the space in which we have to search for optimal clusters.

This insight suggests a naive approach, where we approximate \(W\) by its truncated eigendecomposition. We restate in this case the optimization problem to find averagely dense clusters from Equation (6) as

\[
\max_{Y \in \mathbb{R}^{m \times r}} \sum_k \frac{Y_s^\top V(d)\Lambda(d)V(d)^\top Y_s}{|Y_s|}. \tag{8}
\]

A comparison with the \(k\)-means objective from Equation (2) shows that the objective above is a trace maximization problem which is equivalent to \(k\)-means clustering on the data matrix \(U = V(r)(\Lambda(r))^{1/2}\).

Unfortunately, applying \(k\)-means to the matrix \(U\) does not yield acceptable clusterings. The objective of \(k\)-means is nonconvex and has multiple local solutions. The number of local solutions increases with every eigenvector which we include in the eigendecomposition from \(W\), due to the orthogonality of eigenvectors. That is, with every included eigenvector, we increase the dimension of the subspace, in which we search for suitable binary cluster indicator vectors. As a result, \(k\)-means returns clusterings whose objective function value approximates the global minimum, but which reflect seemingly haphazard groupings of the data points.

The eigenvectors are not only orthogonal, but also real-valued, having positive as well as negative values. The first eigenvectors have a high value of the density function \(\delta\), but the mixture of signs in their entries makes an interpretation as cluster indicators difficult. If the eigenvectors would be nonnegative, then they would have an interpretation as fuzzy cluster indicators, where the magnitude of the entry \(V_{jk}\) indicates the degree to which point \(j\) is assigned to the fuzzy cluster \(k\). Applying \(k\)-means to a set of highly dense fuzzy cluster indicators would furthermore reduce the space in which we search for possible cluster centers to the positive quadrant. One possibility is to approximate the matrix \(W\) by a symmetric nonnegative matrix factorization, but this replaces the polynomially solvable eigendecomposition with an NP-hard problem (Vavasis 2009). We conduct another approach, showing that fuzzy cluster indicators are derivable from the eigenvalues by a simple projection.

**Observation 1.** Let \(W\) be a symmetric real-valued matrix, and let \(v\) be an eigenvector to the eigenvalue \(\lambda\). Let \(v = v^+ - v^-\), with \(v^+, v^- \in \mathbb{R}_+^m\) be the decomposition of \(v\) into its positive and negative parts. The nonnegative vector \(u = v^+ + v^-\) has a density

\[
\delta(u) \geq |\lambda|.
\]

**Proof.** Application of the triangle inequality shows that the entries of an eigenvector having a high absolute value, play an important role for achieving a high Rayleigh quotient. Let \(v\) be an eigenvector to the eigenvalue \(\lambda\) of \(W\), then:

\[
|\lambda||v_j| = |W_{j\cdot}v| = \sum_l W_{jl}v_l \leq \sum_l W_{jl}|v_j|. \tag{9}
\]

Since \(u_j = |v_j|\), from the inequality above follows that \(Wu \geq \lambda u\). Multiplying the vector \(u^+\) from the left and dividing by \(\|u\|^2\) yields the stated result. \(\square\)

We refer to the eigenvectors, whose entries are replaced with their absolute values, i.e., \(u_j = |v_j|\) for \(1 \leq j \leq m\), as **projected eigenvectors**. The projected eigenvectors have locally similar values, resulting in the gradually changing shapes, illustrated on the two circles dataset in Figure 2. We see that the projected eigenvectors have an interpretation as fuzzy cluster indicators: a strongly indicated dense part of one of the two clusters fades out along the circle trajectory. Furthermore, we see that the projected eigenvectors are not orthogonal, since some regions are repeatedly indicated over multiple eigenvectors. These multiple views on possible dense and fuzzy clusters might be beneficial in order to robustly identify the clustering structure.

Applying \(k\)-means to the first \(d\) projected eigenvectors yields a set of \(r\) binary basis vectors, which indicate clusters.

![Figure 2: Visualization of every fifth eigenvector for the two circles dataset. The size of each point is proportional to the absolute value of the corresponding entry in the eigenvector.](image-url)
in the subspace of points with an average density larger than \( \lambda_d \). We summarize the resulting method SpectACL (Spectral Averagely-dense Clustering) with the following steps:

1. compute the adjacency matrix \( W \);
2. compute the truncated eigendecomposition \( W \approx V^{(d)} \Lambda^{(d)} V^{(d)\top} \);
3. compute the projected embedding \( U_{jk} = |V^{(d)}_{jk}|^{1/2} \);
4. compute a \( k \)-means clustering, finding \( r \) clusters on the embedded data \( U \).

These steps perform a minimization of an upper bound on the average cluster density function with respect to \( W \) if \( d = m \). In the general case \( d \ll m \), we optimize an approximation of the objective function, replacing \( W \) with the nonnegative product \( U U^\top \). Certainly, a similar argument could be made for the pipeline of Spectral Clustering, considering the approximate objective (8) for \( d = r \). However, the difference is that the result of Spectral Clustering deteriorates with an increasing accuracy of the eigendecomposition approximation \( (d \rightarrow m) \) due to orthogonality and negative entries in the eigenvectors as outlined above. We show in our experimental evaluation that the result of our method does in average not deteriorate with an increasing dimension of the embedding.

The matrix \( W \) from the first step is here calculated by the \( \epsilon \)-neighborhood graph. However, our method is in principle applicable to any provided (weighted) adjacency matrix. We recall that the average density objective is equivalent to the ratio cut objective if we normalize the matrix \( W \). We refer to this version as normalized SpectACL. In the normalized case, we compute the adjacency matrix according to the \( k \)-nearest neighbor graph, which is suggested for applications of Spectral Clustering.

Hence, our method only depends on the parameters \( r \) and \( \epsilon \), respectively \( k \). How to determine the number of clusters, which is a general problem for clustering tasks, is beyond scope of this paper. We do provide a heuristic to determine \( \epsilon \) and evaluate the sensitivity to the parameters \( \epsilon \) and \( k \).

### 4 Experimental Evaluation

We conduct experiments on a series of synthetic datasets, exploring the ability to detect the ground truth clustering in the presence of noise. On real-world data, we compare the Normalized Mutual Information (NMI) of obtained models with respect to provided classes. A small qualitative evaluation is given by means of a handwriting dataset, illustrating the clusters obtained by SpectACL.

#### 4.1 Synthetic Datasets

We generate benchmark datasets, using the renowned scikit library. For each shape—moons, circles, and blobs—and noise specification we generate \( m = 1500 \) data points. The noise is Gaussian, as provided by the scikit noise parameter; cf. http://scikit-learn.org. Our experiments pit five competitors against each other: two new ones presented in this paper, and three baselines. We compare our method SpectACL, using the \( \epsilon \)-neighborhood graph represented by \( W \) (denoted SpectACL) and the symmetrically normalized adjacency matrix \( \text{diag}(W_{knn}^{-1/2}W_{knn})^{-1/2} \) where \( W_{knn} \) is the adjacency matrix to the \( k \)-nearest neighbor graph (denoted SpectACL (Normalized), or (N) as shorthand). The parameter \( \epsilon \) is determined as the smallest radius such that 90\% of the points have at least ten neighbors and the number of nearest neighbors is set to \( k = 10 \). We compare against the scikit implementations of DBSCAN (denoted DBSCAN) and symmetrically normalized Spectral Clustering (denoted SC). For DBSCAN, we use the same \( \epsilon \) as for SpectACL and set the parameter \( \text{minPts} = 10 \), which delivered the best performance on average. We also compare against the provided Python implementation of Robust Spectral Clustering (Bojchevski, Matkovic, and Günnemann 2017) (denoted RSC), where default values apply. Unless mentioned otherwise, our setting for the embedding dimensionality is \( d = 50 \).

**Evaluation** We assess the quality of computed clusterings by means of the ground truth. Given a clustering \( Y \in \{0, 1\}^{m \times r} \) and the ground truth model \( Y^* \in \{0, 1\}^{m \times r} \), we compute the \( F \)-measure. The bijection of matching clusters
Comparing these results with Robust Spectral Clustering (RSC), learning the graph structure together with the Spectral Clustering does not drastically affect the result. Given the two moons or the two circles dataset, the $F$-measure of RSC is close to that of SC. The example clusterings in the top row of Figure 4 show that RSC is able to correct the vanilla Spectral Clustering, where a chunk from the wrong moon is added to the blue cluster. Yet, in the case of the two circles with different densities, both methods fail to recognize the circles (cf. Figure 4, middle row). Surprisingly, on the easiest-to-cluster dataset of the three blobs, RSC attains notably lower $F$-measures than Spectral Clustering.

The plots for DBSCAN present the difficulty to determine a suitable setting of parameters $\epsilon$ and $\text{minpts}$. The method to determine these parameters is suitable if the noise is low. However, beyond a threshold, DBSCAN decides to pack all points into one cluster. We also adapted the heuristic for the parameter selection, determining $\epsilon$ as the minimum radius such that $70 - 90\%$ of all points have at least $\text{minpts} = 10$ neighbors. This does not fundamentally solve the problem: it merely relocates it to another noise threshold.

Parameter Sensitivity We present effects of the parameter settings for SPECTACL, namely the embedding dimension $d$, the neighborhood radius $\epsilon$ and the number of nearest neighbors $k$. We summarize the depiction of these experiments in Figure 5. The plots on the top row show that the results of SPECTACL are robust to the parameters determining the adjacency matrix. The fluctuations of the $F$-measure when $\epsilon > 0.6$ are understandable, since the coordinates of points are limited to $[0, 3]$. That is, $\epsilon = 0.6$ roughly equates the diameter of the smaller blob on the top left in Figure 4 and is quite large. The plots on the bottom of Figure 5 indicate that SPECTACL in the unnormalized and normalized version generally only require the dimension of the embedding to be large enough. In this case, setting $d > 25$ is favorably. The only exception is the two circles dataset when using normalized SPECTACL, where the $F$-measure starts to fluctuate if $d > 75$. A possible explanation for this behavior is that eigenvalues in the normalized case lie between zero and one. When we inspect the first 50 projected eigenvectors of the normalized adjacency matrix, then we see that the last 19 eigenvectors do not reflect any sensible cluster structure, having eigenvalues in $[0.4, 0.54]$. In unnormalized SPECTACL, the unhelpful eigenvectors have a very low density and thus also a very low eigenvalue, being barely taken into account during the optimization.

### 4.2 Real-World Datasets

We conduct experiments on selected real-world datasets, whose characteristics are summarized in Table 1. The Pulsar dataset contains samples of Pulsar candidates, where the positive-class of real Pulsar examples poses a minority against noise effects. The Sloan dataset comprises measurements of the Sloan Digital Sky Survey, where every observation belongs either to a star, a galaxy or a quasar. The MNIST...
The neighborhood radius $\epsilon$ (top left) and the number of nearest neighbors $k$ (top right) determine the adjacency matrix for SPECTACL and its normalized version. The parameter $d$ (bottom left and right) specifies the number of projected eigenvectors. We plot the $F$-measure (the higher the better) against the variation of the parameter.

<table>
<thead>
<tr>
<th>Data</th>
<th>$m$</th>
<th>$n$</th>
<th>$r$</th>
<th>$\mu \left( \frac{X(j)}{m} \right)$</th>
<th>Algorithm</th>
<th>Pulsar</th>
<th>Sloan</th>
<th>MNIST</th>
<th>SNAP</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pulsar</td>
<td>17898</td>
<td>9</td>
<td>2</td>
<td>0.11 ± 0.08</td>
<td>SPECTACL</td>
<td>0.151</td>
<td>0.224</td>
<td>0.339</td>
<td>0.232</td>
</tr>
<tr>
<td>Sloan</td>
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<td>16</td>
<td>3</td>
<td>0.09 ± 0.07</td>
<td>SPECTACL(N)</td>
<td>0.005</td>
<td>0.071</td>
<td>0.756</td>
<td>0.104</td>
</tr>
<tr>
<td>MNIST</td>
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<td>784</td>
<td>10</td>
<td>0.05 ± 0.06</td>
<td>DBSCAN</td>
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<td>0.320</td>
<td>0.005</td>
<td>–</td>
</tr>
<tr>
<td>SNAP</td>
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<td>–</td>
<td>42</td>
<td>0.05 ± 0.06</td>
<td>SC</td>
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<td>0.098</td>
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<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>RSC</td>
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<td>0.027</td>
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<td>–</td>
</tr>
</tbody>
</table>

Table 1: Real-world dataset characteristics; number of samples $m$, features $n$, clusters (classes) $r$ and relative average number of neighbors in the $\epsilon$-neighborhood graph $\mu$.

The SNAP dataset (Lecun et al. 1998) is a well-known collection of handwritten ciphers. The SNAP dataset refers to the Email EU core network data (Leskovec and Krevl 2014), which consists of an adjacency matrix (hence this dataset has no features in the traditional sense). For this dataset, we only compare the two versions of SPECTACL and Spectral Clustering, since Robust Spectral Clustering and DBSCAN do not support a direct specification of the adjacency matrix. Table 2 summarizes the normalized mutual information between the found clustering and the clustering which is defined by the classification. We observe that the unnormalized version of SPECTACL obtains the highest NMI on the pulsar dataset, the second-highest on the Sloan dataset, a significantly lower NMI than RSC, SC and the normalized version on the MNIST sample and a similar NMI on the SNAP dataset. The Sloan and MNIST datasets pose interesting cases. The notoriously underperforming DBSCAN obtains the highest NMI at the Sloan dataset while obtaining the lowest $F$-measure of 0.09, where SPECTACL obtains the highest $F$-measure of 0.52. The datasets Pulsar and Sloan are clustered in accordance with the classes when using the density-based approach, yet the MNIST clustering corresponds more with the clustering when using the minimum-cut associated approaches.

To understand why SPECTACL fails to return clusters in accordance with the MNIST classes, we display some exemplary points for three SPECTACL clusters in Figure 6. On the one hand, some identified clusters indeed contain only one cipher, e.g., there are clusters for zeros, ones and sixes exclusively. On the other hand, the clusters depicted in the figure contain a mixture of digits, where the digits are written in a cursive style or rather straight. Hence, instead of identifying the numbers, SPECTACL identifies clusters of handwriting style, which is interesting information discovered from the dataset, albeit not the information the MNIST task asks for. To be fair, it is far from impossible that a similar rationale can be given for the clusters SPECTACL’s competitors find on the Pulsar or Sloan dataset; we lack the astrophysical knowledge required to similarly assess those results.

5 Conclusions

We introduce SPECTACL (Spectral Averagely-dense Clustering); a new clustering method that combines benefits from Spectral Clustering and the density-based DBSCAN algorithm, while avoiding some of their drawbacks.

By computing the spectrum of the weighted adjacency matrix, SPECTACL automatically determines the appropriate density for each cluster. This eliminates the specification of the $\text{minpts}$ parameter which is required in DBSCAN, and as we have seen in Figure 1, this specification is a serious hurdle for a DBSCAN user to overcome. On two concentric circles with the same number of observations (and hence with different densities), a DBSCAN run with $\text{minpts} = 25$ lumps all observations into one big cluster. When increasing $\text{minpts}$ by one, DBSCAN jumps to four clusters, none of

Figure 5: Variation of latent parameters used for SPECTACL.

Figure 6: SPECTACL clusters individual handwriting styles instead of cipher shapes.
which are appropriate. SPECTACL, conversely, can natively handle these nonconvex clusters with varying densities.

In both Spectral Clustering and SPECTACL, the final step is to postprocess intermediate results with $k$-means. Whether this choice is appropriate for Spectral Clustering remains open to speculation. However, from the objective function of SPECTACL, we derive an upper bound through the eigenvector decomposition, whose optimization we show to be equal to $k$-means optimization. Hence, for SPECTACL, we demonstrate that this choice for $k$-means postprocessing is mathematically fundamentally sound.

In comparative experiments, competing with DBSCAN, Spectral Clustering, and Robust Spectral Clustering, we find on synthetic data that the unnormalized version of SPECTACL is the most robust to noise (cf. Figure 3), and finds the appropriate cluster structure in three scenarios while the competitors all fail to do so at least once (cf. Figure 4). On real-life data, SPECTACL outperforms the competition on the inherently noisy Pulsar dataset and the Sloan dataset, it performs similarly to Spectral Clustering on the SNAP dataset, and it is defeated by both Spectral Clustering and Robust Spectral Clustering on the MNIST dataset (cf. Table 2). The latter observation is explained when looking at the resulting clusters: as Figure 6 illustrates, SPECTACL focuses on clusters representing handwriting style rather than clusters representing ciphers, which is not unreasonable behavior for an unsupervised method such as clustering; this uncovered information is merely not reflected in the NMI scores displayed in the table.

Figure 5 provides evidence that SPECTACL is robust with respect to its parameter settings. Hence, in addition to its solid foundation in theory and good empirical performance on data, SPECTACL provides a clustering method that is easy to use in practice. Hence, SPECTACL is an ideal candidate for outreach beyond data mining experts.

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