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Abstract—Spectral clustering is a technique that uses the spectrum of a similarity graph to cluster data. Part of this procedure involves calculating the similarity between data points and creating a similarity graph from the resulting similarity matrix. This is ordinarily achieved by creating a k-nearest neighbour (kNN) graph. In this paper, we show the benefits of using a different similarity graph, namely the union of the kNN graph and the minimum spanning tree of the negated similarity matrix (kNN-MST). We show that this has some distinct advantages on both synthetic and real datasets. Specifically, the clustering accuracy of kNN-MST is less dependent on the choice of k than kNN is.

I. INTRODUCTION

Clustering is an important task in data exploration, with the aim being to group objects or observations in such a way that objects within the same group are broadly more similar to one another than they are to objects in other groups.

Spectral clustering is a group of methods that use the spectrum of a similarity matrix, or a matrix derived from it, to cluster the data. One of the most basic variants is described as follows [1]. Given a matrix \( A \in \mathbb{R}^{m \times n} \) containing \( m \) observations with \( n \) features, a similarity between each pair \( i, j \) of observations can be computed using a similarity measure. One such measure in common use is the Gaussian similarity measure [2]. Given that \( x_i \) is the \( i \)-th row of \( A \), then the similarity between observations \( i \) and \( j \) is defined as:

\[
s(x_i, x_j) = \exp \left( -\frac{||x_i - x_j||^2}{2\sigma^2} \right)
\]  

(1)

Where \( x_i, x_j \in \mathbb{R}^n \) and \( ||x_i - x_j|| \) is the Euclidean distance between the two vectors and \( \sigma \) controls the width of the neighbourhoods [1]. There are various suggestions for choosing \( \sigma \); a common choice is for \( \sigma \) to be the standard deviation of the observations \( ||x_i - x_j|| \), which is what we will use in this paper. [3]

In this way, the similarity between every pair \( i, j \) can be computed and a similarity matrix \( S \) can be constructed where \( S_{ij} = s(x_i, x_j) \).

At this point, a similarity graph \( G \) is constructed. The graph \( G = (V, E) \) consists of a finite set \( V = \{v_1, \ldots, v_n\} \) of vertices and a finite set \( E \) of edges, where each edge is an unordered pair connecting two vertices in \( V \). There are many ways in which to construct \( G \). The aim is to construct a graph such that similar observations are closely connected and dissimilar observations are not. The clustering problem then becomes that of finding a partition of the graph \( G \) such that edges between different clusters have low weight (low similarity).

One such similarity graph, and the one we focus on in this paper, is the k-nearest neighbour graph. For each \( i \in \{1, \ldots, m\} \), the edges to the \( k \) observations with highest similarities to \( i \) in \( S \) are added to the graph.

Due to useful spectral properties exhibited by the graph Laplacian [4], the spectrum is calculated for the Laplacian of the similarity graph, rather than on the similarity graph itself. Given that \( W \) be the adjacency matrix of \( G \), then the unnormalised Laplacian \( L_{un} \) of \( W \) is:

\[
L_{un} = D - W
\]  

(2)

Where \( D \) is the diagonal matrix containing the degree of each vertex on its diagonal. The degree \( d_i \) of vertex \( i \) is the sum of the entries in the \( i \)-th row of \( W \).

If we wish to divide the data into \( l \) clusters, we now compute the first \( l \) eigenvectors of \( L_{un} \), that is, the \( l \) eigenvectors corresponding to the \( l \) lowest eigenvalues. Let \( U \in \mathbb{R}^{m \times l} \) be the matrix containing the first \( l \) eigenvectors \((u_1, \ldots, u_l)\) as columns.

Let \( y_i \) be the \( i \)-th row of \( U \). The clusters are then found by using k-means clustering to cluster the points \((y_i)_{i=1, \ldots, m} \in \mathbb{R}^l \) into clusters \( C_1, \ldots, C_l \).

There are many varieties of spectral clustering, most notably, by changing the type of Laplacian used. Using the unnormalised Laplacian \( L_{un} \) as described above is known as unnormalised spectral clustering. Two popular variations that are based around normalised Laplacians are the normalised random walk spectral clustering [5] and normalised symmetric spectral clustering [2] that use \( L_{rw} \) and \( L_{sym} \) respectively:

\[
L_{rw} = D^{-1/2}L_{un}D^{-1/2}
\]  

(3)

\[
L_{sym} = D^{-1/2}L_{un}D^{-1/2}
\]  

(4)

We will use \( L_{sym} \) throughout this paper. The focus of this paper is the step of creating the similarity graph. We will show that by including the minimum spanning tree (MST)
of the negated similarity matrix into the k-nearest neighbour (kNN) similarity graph, the quality of the clusters detected by the spectral clustering procedure becomes less sensitive to the choice of $k$.

A. Sensitivity to $k$

When using kNN as the similarity graph in spectral clustering, the choice of $k$ can have a significant influence on the accuracy of the detected clusters when compared to the ground truth. A key reason for this is the connectivity of the similarity graph.

The following is true for the spectrum of all three Laplacians discussed, $L_{un}$, $L_{sym}$ and $L_{rw}$. If a similarity graph $G$ has $c$ disconnected components, then the multiplicity of the eigenvalue 0 is equal to the number of disconnected components $c$. The first $c$ eigenvectors work to separate those components. Therefore, if one were to look for $l$ clusters, but $c > l$, then this results in less accurate clustering as only the first $l$ eigenvectors are used in the spectral clustering algorithm.

Therefore, it is important to choose a good value for $k$ to ensure at the very least, that the kNN similarity graph is connected. In general, the optimal choice of $k$ does not have a closed form solution. But due to asymptotic connectivity results, it is common to choose $[\log n]$ for $k$ [6].

The key result in this paper is to make the search for the optimal $k$ less important. By adding the minimum spanning tree to the kNN similarity graph, connectivity is guaranteed. There will be no disconnected components. In this paper we will show empirically that doing this leads to more consistent results.

In section II we introduce the kNN-MST method and in section III we demonstrate our argument on the well known Iris dataset from Fisher. We then look at both synthetic and real datasets in section IV. The conclusion is outlined in section V.

II. Minimum Spanning Trees and k-Nearest Neighbours

In this paper we show that spectral clustering on a similarity graph consisting of the union of 1) the kNN similarity graph [7] and 2) the minimum spanning tree [8] of the negated complete similarity matrix, tends to perform better than kNN.

Given a similarity matrix $S$, we can produce the adjacency matrix $W^{(k)}$ corresponding to the $k$-nearest neighbour similarity graph with a simple procedure. Let $\left( x_i^{(p)}, p \in \{1 \ldots (n-1)\} \right)$ be the sequence of the $n-1$ neighbours of $i$ ordered from the closest to the furthest neighbour; from the most similar to the least similar. That is, $x_i^{(p)}$ is the index of the $p$-th most similar neighbour to $i$. We can then define the matrix $W^{(k)}$ as follows:

$$ W^{(k)}_{ij} = \begin{cases} S_{ij} & \text{if } j \in \{x_i^{(1)}, \ldots, x_i^{(k)}\} \\ 0 & \text{otherwise} \end{cases} $$

$W^{(k)}$ is the adjacency matrix corresponding to the kNN similarity graph for $S$.

A minimum spanning tree (MST) is a tree (connected undirected graph with no cycles) that spans a graph such that the sum of edge weights is less than or equal to that of any other tree of the graph. If we treat the negated complete similarity matrix $-S$ as an adjacency matrix for a graph, then that graph $G$ will be a complete graph. Let $G^{(MST)}$ be the MST for that graph, and let $W^{(MST)}$ be the adjacency matrix of $G^{(MST)}$. Note that the minimum spanning tree was taken for the negated similarity matrix so that the minimum spanning tree contains the edges representing the highest similarities.

The second MST can be found by removing from $-S$ the elements that are part of the MST of $-S$, and then finding the minimum spanning tree for that modified version of $-S$. We define kMST to be the similarity graph defined as the union of the first $k$ minimum spanning trees of $-S$.

We will look at kMST in our results to demonstrate that the use of the kNN-MST similarity graph leads to more accurate clusters both due to the MST portion and the kNN portion. That is, kNN-MST performs better than either kNN or kMST individually.

Let kNN-MST be the similarity graph created by taking the union of the first MST of the negated complete similarity matrix and the kNN similarity graph. That is, let the kNN-MST similarity graph be defined as $G^{(kNNMST)} = (V, E)$, where $V$ is the set of $n$ vertices and $E$ is defined as:

$$ E(G^{(kNNMST)}) = E(G^{(MST)}) \cup E(G^{(kNN)}) $$

Let $W^{(kNNMST)}$ be the adjacency matrix of $G^{(kNNMST)}$. In the next sections, we will show that the use of $G^{(kNNMST)}$ as a similarity graph in spectral clustering has advantages over using $G^{(kNN)}$.

III. An Example Using Fisher’s Iris Dataset

Fisher’s Iris Flower dataset consists of 150 observations with 4 features [9]. Each of the 150 observations is classified as being part of one of three species of Iris, namely Iris setosa, Iris Virginica and Iris versicolor.

We will use the adjusted Rand index to quantify the accuracy of the clusters determined by the spectral clustering algorithms [10]. A value of 1 indicates perfect agreement between the ground truth clusters and the clusters identified by the clustering algorithm. A value close to 0 indicates that any agreement is mostly due to chance.

Figure 1 shows the adjusted Rand index of the clusters identified by spectral clustering with the use of kNN, kMST and kNN-MST as the similarity graphs for the Iris dataset.

The key point is that while equally good clusters can be found using kNN spectral clustering, by combining it with the MST, the result is more stable, and less sensitive to the choice of $k$.

The primary reason that adding the MST to the kNN similarity graph improves the accuracy of spectral clustering is that it ensures that the resulting similarity graph is connected. Figure 2 shows the 2NN similarity graph and the 2NN-MST similarity graph for the Iris dataset. The fact that the 2NN
graph has seven disconnected components leads to the first seven eigenvectors of the Laplacian of that graph to separate the components. This does not work well if the goal is to find \( l \) clusters where \( l < 7 \). In figure 1 it is clear that with a low value of \( k \), the clusters found using spectral clustering on the kNN similarity graph are not accurate. But adding the MST to those same kNN graphs drastically improves the result.

Fig. 1: The adjusted Rand index of the three clusters found through spectral clustering when compared with the known ground truth for the Iris dataset. Comparing the use of the kNN, kMST and kNN-MST similarity graphs for various values of \( k \).

We will now show that the use of kNN-MST as the similarity graph yields similar benefits on a selection of other datasets.

IV. RESULTS

A. Datasets

We will use both synthetic and real datasets to demonstrate empirically the benefit of adding the MST to kNN similarity graphs in spectral clustering.

First, we will use the Fundamental Clustering Problem Suite (FCPS) \[11\], which consists of ten synthetic classification problems; each of these represents a different clustering challenge such as different cluster densities, clusters that are not linearly separable, data with outliers and so forth.

The size of the ten problems, and the challenges they exhibit are summarised in table I.

For these datasets, we performed spectral clustering with the kNN and the kNN-MST similarity graphs with \( k \in \{1, \ldots, 10\} \). In order to compare the Rand index for the various datasets, the adjusted Rand index results for each dataset are normalised through feature scaling. That is, given that \( x \in \mathbb{R}^{20} \) is the vector containing the 20 adjusted Rand index results (10 for kNN and 10 for kNN-MST) for one particular dataset, then we define the vector \( y \in \mathbb{R}^{20} \) to be:

\[
y = \frac{x - \min(x)}{\max(x) - \min(x)}
\]  

Figure 3 shows the mean normalised adjusted Rand index for all the datasets within the FCPS when spectral clustering was performed using the kNN and kNN-MST similarity graphs. kNN-MST outperforms kNN on its own, for every \( k \in \{1, \ldots, 10\} \), but significantly so for lower values of \( k \).
TABLE I: Summary of the problems within the Fundamental Clustering Problem Suite, where n is the number of data points, d is the number of dimensions and cl the number of clusters.

<table>
<thead>
<tr>
<th>Name</th>
<th>n</th>
<th>d</th>
<th>cl</th>
<th>Problem Summary</th>
</tr>
</thead>
<tbody>
<tr>
<td>Atom</td>
<td>800</td>
<td>3</td>
<td>2</td>
<td>Different variances. Linear not separable.</td>
</tr>
<tr>
<td>ChainLink</td>
<td>1000</td>
<td>3</td>
<td>2</td>
<td>Linear not separable.</td>
</tr>
<tr>
<td>EnzyTime</td>
<td>4096</td>
<td>2</td>
<td>2</td>
<td>Gaussian mixture overlap.</td>
</tr>
<tr>
<td>GolfBall</td>
<td>4002</td>
<td>3</td>
<td>1</td>
<td>One large cluster.</td>
</tr>
<tr>
<td>Hepta</td>
<td>212</td>
<td>3</td>
<td>7</td>
<td>Clearly defined clusters. Different variances.</td>
</tr>
<tr>
<td>Lsun</td>
<td>400</td>
<td>2</td>
<td>3</td>
<td>Different variances and inter-cluster distances.</td>
</tr>
<tr>
<td>Target</td>
<td>770</td>
<td>2</td>
<td>6</td>
<td>Outliers.</td>
</tr>
<tr>
<td>Tetra</td>
<td>400</td>
<td>3</td>
<td>4</td>
<td>Almost touching clusters.</td>
</tr>
<tr>
<td>TwoDiamonds</td>
<td>800</td>
<td>2</td>
<td>2</td>
<td>Cluster borders defined by density.</td>
</tr>
<tr>
<td>WingNut</td>
<td>1070</td>
<td>2</td>
<td>2</td>
<td>Density vs Distance</td>
</tr>
</tbody>
</table>

In this section we will look in detail at a synthetic ChainLink dataset (see figure 4). This dataset has two clusters; each is a ring in 3 dimensional space. The location of each data point on the circumference of the ring is determined from a uniform distribution. Each data point is then relocated from its position on the circumference. The magnitude and direction of this relocation for the three dimensions are determined by sampling from $N(0, \sigma^2)$. We will generate ChainLink datasets with various standard deviations to demonstrate how kNN and kNN-MST perform when the two clusters become less and less clearly defined; when each ring becomes noisier.

We have generated the ChainLink dataset using a normal distribution $N(0, \sigma^2)$ with $\sigma \in \{0, 0.01, \ldots, 0.19, 0.2\}$. For each $\sigma$ we generated 100 instances and averaged the adjusted Rand index results over those 100 instances.

Figures 4 and 5 show two examples of the ChainLink dataset, one with $\sigma = 0.1$ and the other with $\sigma = 0.2$. The heatmap in figure 6 shows the improvement of the adjusted Rand index when kNN-MST is used instead of kNN. This is shown for various values of $k$ and $\sigma$. 

![Fig. 4: A ChainLink dataset with standard deviation $\sigma = 0.1$.](image1)

![Fig. 5: A ChainLink dataset with standard deviation $\sigma = 0.2$.](image2)
It is clear that using kNN-MST instead of kNN alone leads to a significant improvement in the clusters found for lower values of $k$. As before, this is primarily due to the MST ensuring that the similarity graph is connected. For higher values of $k$, the resulting clusters from using kNN-MST are either better or equally as good as using kNN alone. This result holds both for ChainLink instances that are well separated and also for instances that are not well separated. However, it is especially noticeable in the well separated case.

![Heatmap showing the improvement in the clusters identified by spectral clustering when using kNN-MST instead of kNN, for ChainLink instances that are well separated (towards the top) and instances that are less well separated (towards the bottom). Dark red means there is no improvement. Lighter colours demonstrate that there is.](image)

Fig. 6: Heatmap showing the improvement in the clusters identified by spectral clustering when using kNN-MST instead of kNN, for ChainLink instances that are well separated (towards the top) and instances that are less well separated (towards the bottom). Dark red means there is no improvement. Lighter colours demonstrate that there is.

**D. Real Datasets**

Finally, in addition to the Iris dataset used in section III, we use four other real life datasets, namely the Leaf [12], Wine, Seeds, and Breast Cancer [13] datasets as found in the UCI machine learning repository [14].

Table II summarises the real life datasets we test the spectral clustering algorithms on.

<table>
<thead>
<tr>
<th>Name</th>
<th>n</th>
<th>d</th>
<th>cl</th>
</tr>
</thead>
<tbody>
<tr>
<td>Iris</td>
<td>150</td>
<td>4</td>
<td>3</td>
</tr>
<tr>
<td>Leaf</td>
<td>340</td>
<td>14</td>
<td>30</td>
</tr>
<tr>
<td>Wine</td>
<td>178</td>
<td>13</td>
<td>3</td>
</tr>
<tr>
<td>Seeds</td>
<td>210</td>
<td>7</td>
<td>3</td>
</tr>
<tr>
<td>Breast Cancer</td>
<td>683</td>
<td>9</td>
<td>2</td>
</tr>
</tbody>
</table>

Table II: Summary of the real life datasets, where $n$ is the number of data points, $d$ is the number of dimensions and $cl$ the number of clusters.

![Adjusted Rand Index comparing the ground truth for the datasets with the clusters found by spectral clustering. Comparing the use of kNN, kMST and kNN-MST as the similarity graphs in the spectral clustering algorithm.](image)

Fig. 7: Adjusted Rand Index comparing the ground truth for the datasets with the clusters found by spectral clustering. Comparing the use of kNN, kMST and kNN-MST as the similarity graphs in the spectral clustering algorithm.
V. CONCLUSION

The spectrum of the Laplacian of the kNN similarity graph is widely used for spectral clustering. One of the problems associated with this, is choosing a good value for \( k \). A common suggestion is to choose \( k = \lceil \log n \rceil \), but there is no guarantee that this will lead to a good clustering. Furthermore, the final quality of the clusters is very sensitive to the choice of \( k \), especially if the chosen value is low. This is primarily due to the similarity graph not being connected.

We have shown in this paper that the sensitivity to the choice of \( k \) can be reduced if the kNN-MST similarity graph is used in the spectral clustering algorithm, instead of kNN. The kNN-MST graph represents the union of the minimum spanning tree of the negated similarity matrix and the kNN similarity graph.

Our key result is that by adding the MST to the kNN similarity graph, the sensitivity to the choice of \( k \) is reduced. This is mostly due to ensuring that the final similarity graph is connected. Furthermore, our empirical tests show that adding the MST to the kNN similarity graph leads to good clustering accuracy even for a low choice of \( k \) that would ordinarily lead to especially inaccurate results. Therefore, we recommend using the kNN-MST similarity graph instead of kNN in spectral clustering.

The use of kNN-MST has been shown to detect clusters that are equally good, or better than, the clusters detected when kNN is used. This is the case in both synthetic and real datasets.

In future work, it is worth investigating kMST, as our work here shows that multiple minimum spanning trees also have the potential to work as good similarity graphs. Furthermore, while the primary reason that kNN-MST outperforms kNN as a similarity graph is due to the kNN-MST graph being connected, there are instances where kNN-MST continues to perform better even compared to connected kNN graphs. There is potential for investigating this.

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