Efficient Eigen-updating for Spectral Graph Clustering
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Partitioning a graph into groups of vertices such that those within each group are more densely connected than vertices assigned to different groups, known as graph clustering, is often used to gain insight into the organisation of large scale networks and for visualisation purposes. Whereas a large number of dedicated techniques have been recently proposed for static graphs, the design of on-line graph clustering methods tailored for evolving networks is a challenging problem, and much less documented in the literature. Motivated by the broad variety of applications concerned, ranging from the study of biological networks to the analysis of networks of scientific references through the exploration of communications networks such as the World Wide Web, it is the main purpose of this paper to introduce a novel, computationally efficient, approach to graph clustering in the evolutionary context. Namely, the method promoted in this article can be viewed as an incremental eigenvalue solution for the spectral clustering method described by Ng, et al. (2001). The incremental eigenvalue solution is a general technique for finding the approximate eigenvectors of a symmetric matrix given a change. As well as outlining the approach in detail, we present a theoretical bound on the quality of the approximate eigenvectors using perturbation theory. We then derive a novel spectral clustering algorithm called Incremental Approximate Spectral Clustering (IASC). The IASC algorithm is simple to implement and its efficacy is demonstrated on both synthetic and real datasets modelling the evolution of a HIV epidemic, a citation network and the purchase history graph of an e-commerce website.

1 Introduction

Graph-mining has recently received increasing attention in the machine-learning literature, motivated by application domains such as the Internet, social net-
works, epidemics of transmissible infectious diseases, sensor and biological networks. A crucial task in exploratory analysis and data visualisation is graph clustering \[32, 10\], which aims to partition the vertices in a graph into groups or clusters, with dense internal connections and few connections between each other. There is a large body of work on graph clustering. A possible approach is to consider a certain measure that quantifies community structure and formulate the clustering issue as an optimisation problem (which is generally NP-hard), for which fairly good solutions can be obtained recursively or by using adequate metaheuristics, see \[24, 22, 9, 31\] for example. A variety of approaches to graph clustering exist, such as those based on modularity maximisation for instance, see \[24, 23\]. In this paper, focus is on the spectral clustering approach \[37\], which performs well empirically, is often simple to implement and benefits computationally from the availability of fast linear algebra libraries. The general principle of spectral clustering is to compute the smallest eigenvectors of some particular matrix \(L\) (refer to Section 2 for further details) and then cluster the vertices based on their representation in the corresponding eigen-space. The popularity of this approach arises from the fact that the obtained clustering of vertices is closely connected to the spectral relaxation of the minimisation of the normalised cut criterion, see \[34\].

In many applications such as communications networks (e.g. the Web and Internet), biological networks (of proteins, metabolic reactions, etc.), social networks or networks of scientific citations for instance, the graphs of interest slowly change with time. A naive approach to this incremental problem is to cluster each graph in the sequence separately, however this is computationally expensive for spectral clustering as the cost of solving the eigenvector problem is \(O(n^3)\) at each iteration, where \(n\) is the number of vertices. There has been some previous work on the incremental spectral clustering problem, for example \[36, 27, 26, 16\] however only \[27, 26\] update the eigen-system. In this paper we propose an efficient method for clustering a sequence of graphs which leverages the eigen-decomposition and the clustering on the previous graph to find the new clustering. Firstly, a fast approximation of a rank-\(k\) eigen-decomposition of \(L_{t+1}\) knowing that of \(L_t\) is derived from the Singular Value Decomposition (SVD) updating approach used for Latent Semantic Indexing in \[42\]. Here, the update is efficient to compute when the change (defined in the subsequent analysis) between \(L_t\) and \(L_{t+1}\) is small. Secondly the clustering of vertices is updated according to the new eigen-space. The efficiency of the complete approach, in terms of clustering accuracy, is demonstrated on synthetic and real data. We point out that the clustering approach was first outlined in \[5\]. Here we provide a theoretical analysis on the quality of the eigen-approximation, as well as a more extensive empirical study of the algorithm.

The paper is organised as follows. Standard spectral clustering and SVD updating approaches are recalled in Sections 2 and 3. Then Section 4 details the proposed eigen-decomposition update and Section 5 studies the accuracy of the resulting approximate eigenvectors using perturbation theory. In Section 6 we show how the eigen-decomposition updates can be applied to spectral clustering. Numerical results are gathered in Section 7, and the paper ends with Section 8 discussing results and planned future work.

**Notation:** A bold uppercase letter represents a matrix, e.g. \(X\), and a column vector is displayed using a bold lowercase letter, e.g. \(x\). The transpose of
a matrix or vector is written $X^T$. The concatenation of the columns of two matrices $A$ and $B$ is written $[A \ B]$. $A[I, I]$ represent the submatrix of $A$ formed using the row and columns indexed by $I$, and $A[I, :]$ and $A[(:, I)]$ are submatrices formed using the rows and columns $I$ respectively. The matrix $A_k$ is that formed using the largest $k$ eigenvectors and eigenvalues of $A$, however $I_p$ is the $p \times p$ identity matrix.

2 Graph Clustering

Consider an undirected graph $G = (V, E)$, composed of a set of vertices $V = \{v_1, \ldots, v_n\}$ and edges $E \subseteq V \times V$ such that for every edge $(v_i, v_j)$ there is also an edge $(v_j, v_i)$. One way of representing the edges of $G$ is using an adjacency matrix $A \in \{0, 1\}^{n \times n}$ which has $A_{ij} = 1$ if there is an edge from the $i$-th to the $j$-th vertex and $A_{ij} = 0$ otherwise. More generally, the weight matrix $W \in \mathbb{R}^{n \times n}$ allows one to assign nonzero numerical values on edges, and thus $W_{ij} \neq 0$ when there is an edge from the $i$-th to the $j$-th vertex.

In the perspective of spectral clustering, a useful way of representing $G$ is through its Laplacian matrix [3]. There are several definitions of the Laplacian matrix, however we are interested in the normalised Laplacian matrix, which is symmetric and positive semi-definite. We recall its definition below for clarity.

**Definition 2.1.** The unnormalised Laplacian matrix of a graph $G$ is defined as $L = D - W$ where $D$ is the degree matrix with supposedly nonzero diagonal entries $D_{ii} = \deg(v_i)$, denoting by $\deg(v_i) = \sum_j W_{ij}$ the degree of the $i$-th vertex, and zeros elsewhere, and $W$ is the weight matrix. The normalised Laplacian matrix of a graph $G$ is then defined as $\tilde{L} = D^{-\frac{1}{2}}LD^{-\frac{1}{2}}$.

The normalised Laplacian matrix is used in the spectral clustering approach of Ng et al. [25]. The algorithm computes the Laplacian and then finds the $k$ smallest eigenvectors which are used for clustering in conjunction with the $k$-means algorithm, see Algorithm 1. The Laplacian matrix is often sparse and one can use power or Krylov subspace methods such as the Lanczos method to find the eigenvectors. There are several variants of Algorithm 1, such as that of [34] which uses the so-called random walk Laplacian and clusters the smallest eigenvectors in a similar way. One of the motivations for these clustering methods is from the spectral relaxation of the minimisation of the normalised cut criterion.

**Algorithm 1** Spectral Clustering using the Normalised Laplacian [25]

**Require:** Graph $G$ with weight matrix $W \in \mathbb{R}^{n \times n}$, number of clusters $k$

1: Find $k$ smallest eigenvectors $V_k = [v_1, \ldots, v_k]$ of normalised Laplacian $\tilde{L}$
2: Normalise the rows of $V_k$, i.e. $V_k \leftarrow \text{diag}(V_kV_k^T)^{-\frac{1}{2}}V_k$
3: Cluster rows of $V_k$ with the $k$-means algorithm
4: **return** Cluster membership vector $c \in \{1, \ldots, k\}^n$

A naive approach to spectral clustering on a sequence of graphs has a large update cost due to the computation of the eigen-decomposition of $\tilde{L}_t$ at each
iteration $t$. A more efficient approach is to update relevant eigenvectors from one iteration to the next. In [36] the authors use the spectral clustering of [25] however they do not update the eigenvectors incrementally but instead the clustering directly. In [16] the authors cluster using the unnormalised Laplacian matrix. Eigenvectors are updated and then clusters are modelled using a set of representative points (a similar strategy is used in [40] for example to cluster points in $\mathbb{R}^d$). The algorithm is potentially costly since it uses the full eigen-decomposition in conjunction with the eigen-gap heuristic to estimate the number of clusters. Furthermore, the solution is updated with the addition of vertices only, and not edges. The iterative clustering approach of Ning et al. [27, 26] uses the spectral clustering method given in [34] and updates eigenvectors incrementally. The algorithm incrementally updates the solution of the generalised eigenproblem $Lv = \lambda Dv$ by finding the derivatives on the eigenvalues/vectors with perturbations in all of the quantities involved. An iterative refinement algorithm is given for the eigenvalues and eigenvectors given a change in the edges or vertices of a graph. One then clusters the resulting $k$ smallest eigenvectors using $k$-means clustering. In order to limit errors which can build up cumulatively the authors recompute the eigenvectors after every $R$-th graph in the sequence.

A disadvantage of the approach of [27, 26] lies in the fact that, to update an eigenvector, one must invert a small matrix for each weight change in the graph which makes updates costly. The size of this matrix is proportional to the number of neighbours of the vertices incident to the changed edge. To be more precise, the cost of updating an eigenvector in [27] is $O(\bar{N}^2n + \bar{N}^3)$ in which $\bar{N}$ is the average size of the “spatial neighbourhood” of an edge, i.e. the average number of rows/columns incident to the edge. It follows that to update $k$ eigenvectors following $r$ edge changes, the complexity is $O(rk(\bar{N}^2n + \bar{N}^3))$.

In contrast, the approach presented in our paper has a smaller update cost for a set of vertex or edge weight changes between $L_t$ and $L_{t+1}$, since changes are considered in a batch fashion, as will later become clear. A further problem with the Ning et al. approach is that eigenvectors are updated independently of one another and hence one loses the orthogonality $v_i^T D v_j = \delta(i, j)$, where $\delta$ is the Kronecker delta function (taking the value 1 if $i = j$ and 0 otherwise), and vectors can become correlated for example.

Another way of improving the efficiency of spectral clustering is to compute approximate eigen-decompositions at each iteration, for example by using the Nyström approach [39]. The Nyström method is used for spectral graph clustering in [11] in conjunction with image segmentation. To estimate the eigenvalues and eigenvectors of $A \in \mathbb{R}^{n \times n}$, one first finds a matrix $A[I, I]$ in which $I \in \{1, \ldots, n\}^m$ is a set of indices selected uniformly at random. If we assume that $A[I, I]$ is positive definite, we can take its square root $A[I, I]^{1/2}$. One then defines


where $\bar{I}$ is the complement of $I$, and diagonalises $S$ using the eigen-decomposition $S = U \Lambda U^T$. Let $V = A[\cdot, I] A[I, I]^{-1/2} U \Lambda^{-1/2}$, then it can be shown that the Nyström approximation of $A$, given by $\tilde{A}_I = A[\cdot, I] A[I, I]^{-1} A[I, \cdot]$, is equivalent to $V \Lambda V^T$. If $A$ is indefinite a more complicated two-step procedure is required, see [11] for details. The resulting approximation is applied to find
the first few eigenvectors of the normalised Laplacian matrix at a total cost of \( O(nm^2 + m^3) \). Several efficiency improvements based on this approach are proposed in [20] which finds the largest \( k \) approximate eigenvectors of \( D^{-\frac{1}{2}}WD^{-\frac{1}{2}} \) at a reduced time and space complexity.

The quality of the resulting approximation is determined by the extent the submatrix \( \mathbf{A}[I, I] \) is spanned by \( \mathbf{A}[I, :] \). One would naturally expect a good approximation for example when \( \mathbf{A}[I, :] \) spans the space of \( \mathbf{A} \). The choice of sampling of \( I \) also affects approximation quality, and empirical and theoretical work in [17] suggests random sampling without replacement over the non-uniform sampling in [7, 6]. Hence for our later empirical work, we will use uniform random sampling without replacement. In addition to [17], error bounds for the Nyström method are presented in [7, 21] in terms of the matrix approximation error using the Frobenius or spectral norm (given by \( \|\mathbf{A}\|_F = \sqrt{\text{tr}(\mathbf{A}^T\mathbf{A})} \) and \( \|\mathbf{A}\|_2 = \sqrt{\lambda_{\text{max}}(\mathbf{A}^T\mathbf{A})} \) respectively). One of the disadvantages of applying the Nyström based approaches of [11, 20] on graphs is that by sampling only a subset of columns of the normalised Laplacian one can exclude important edges which help define clusters. These approaches are naturally more effective when a set of points \( \{x_1, \ldots, x_n\} \in \mathbb{R}^d \) is mapped into a weighted graph using for example the Gaussian weighted distance \( W_{i,j} = \exp(-\|x_i - x_j\|^2)/2\sigma^2 \), for \( \sigma \in \mathbb{R}^+ \), which is very informative about the relative positions of the points.

Another class of algorithms which are potentially useful for spectral clustering is the randomised SVD [14]. Notice that the SVD of a symmetric positive semi-definite matrix is identical to its eigendecomposition and hence SVD algorithms can be applied to spectral clustering. We recount an algorithm from [14] which is used in conjunction with kernel Principal Components Analysis (KPCA, [33]) in [41]. Algorithm 2 provides the associated pseudo-code. The purpose of the first three steps is to find an orthogonal matrix \( \mathbf{Q} \) such that the projection of \( \mathbf{A} \) onto \( \mathbf{Q} \) is a good approximation of \( \mathbf{Y} \), whose columns are random samples from the range of \( \mathbf{A} \), under a rank-\( k \) projection. This projection is then used in conjunction with \( \mathbf{Q} \) so that one need only find the SVD of the smaller matrix \( \mathbf{B} \). The complexity of this approach is \( O((qr + r^2)(m + n)) \), which for a square matrix of size \( n \) collapses simply to \( O((qr + r^2)n) \).

### Algorithm 2 Randomised SVD [14]

<table>
<thead>
<tr>
<th>Require:</th>
<th>Matrix ( \mathbf{A} \in \mathbb{R}^{m \times n} ), number of projection vectors ( r ), exponent ( q )</th>
</tr>
</thead>
<tbody>
<tr>
<td>1:</td>
<td>Generate a random Gaussian matrix ( \Omega \in \mathbb{R}^{n \times r} )</td>
</tr>
<tr>
<td>2:</td>
<td>Create ( \mathbf{Y} = (\mathbf{A}\mathbf{A}^T)^q\mathbf{A}\Omega ) by alternative multiplication with ( \mathbf{A} ) and ( \mathbf{A}^T )</td>
</tr>
<tr>
<td>3:</td>
<td>Compute ( \mathbf{Y} = \mathbf{QR} ) using the QR-decomposition</td>
</tr>
<tr>
<td>4:</td>
<td>Form ( \mathbf{B} = \mathbf{Q}^\top\mathbf{A} ) and compute SVD ( \mathbf{B} = \mathbf{UV}^\top )</td>
</tr>
<tr>
<td>5:</td>
<td>Set ( \mathbf{U} = \mathbf{Q} \mathbf{U} )</td>
</tr>
<tr>
<td>6:</td>
<td><strong>return</strong> Approximate SVD ( \mathbf{A} \approx \mathbf{UV}^\top )</td>
</tr>
</tbody>
</table>

### 3 SVD Updating

An important aspect of the incremental clustering method described later lies in its ability to efficiently compute the eigenvectors of a matrix from the eigen-
vectors of a submatrix, and for clarity’s sake, we outline the SVD-updating algorithm of [42]. The SVD of \( A \in \mathbb{R}^{m \times n} \) is the decomposition

\[
A = P\Sigma Q^T,
\]

where \( P = [p_1, \ldots, p_m] \), \( Q = [q_1, \ldots, q_n] \) and \( \Sigma = \text{diag}(\sigma_1, \ldots, \sigma_r) \) are respectively the orthogonal matrices of left and right singular vectors and a diagonal matrix of singular values \( \sigma_1 \geq \sigma_2 \geq \ldots \geq \sigma_r \), with \( r = \min(m, n) \).

In [42], the authors use the SVD of \( A \) to approximate the SVD of \( [A B] \), with \( B \in \mathbb{R}^{m \times p} \), without recomputing the SVD of the new matrix. It is known that the best \( k \)-rank approximation of \( A \), using the Frobenius norm error, is given by its SVD, namely

\[
A_k = P_k\Sigma_kQ_k^T,
\]

where \( P_k, Q_k, \Sigma_k \) correspond to the \( k \) largest singular values. The general idea of the algorithm is to write

\[
[A_k B] = S\Lambda R^T,
\]

in which \( S \) and \( R \) are matrices with orthonormal columns. One then takes the rank-\( k \) SVD \( \Lambda = G_k\Gamma_kH_k^T \) and \( (SG_k)\Gamma_k(RH_k)^T \) is the rank-\( k \) approximation for \( [A B] \). The dimensionality of \( \Lambda \) is generally much smaller than that of \( [A_k B] \) and hence the corresponding SVD approximation is computationally inexpensive.

This approach is more accurate than those presented in [2, 28], however it comes at additional computational cost. Furthermore, the authors prove that when the matrix \( [A B]^T[A B] \) has the form \( X + \alpha I \) in which \( X \) is symmetric positive semi-definite with rank-\( k \) then the rank-\( k \) approximation of \( [A B] \) is identical to that of \( [A_k B] \).

4 Incremental Eigen-approximation

In this section we address three types of updating problem upon the largest \( k \) eigenvectors of a symmetric matrix. The updates are general operations, however they will be explained in the context of spectral clustering later in Section 6. Assuming that \( Y_1, Y_2 \in \mathbb{R}^{n \times p} \) and one does not have direct access to \( A \in \mathbb{R}^{m \times n} \) and \( B \in \mathbb{R}^{m \times p} \) but only to the matrices \( C = A^TA, A^TB \) and \( B^TB \), the updates are:

1. Addition of a low-rank symmetric matrix \( C \to C + U \) where \( U = Y_1Y_2^T + Y_2Y_1^T \)

2. Addition of rows and columns \( A^TA \to [A B]^T[A B] \)

3. Removal of rows and columns \( [A B]^T[A B] \to A^TA \)

We have written the updates above in terms of a symmetric matrix \( A^TA \) to improve notation. Note that any positive semi-definite symmetric matrix \( M \) can be decomposed into the form \( M = A^TA \), where \( A \) has real entries, for example by using an eigen-decomposition or incomplete Cholesky factorisation.
4.1 Addition of a Low-rank Symmetric Matrix

The first type of eigen-approximation we are interested in is the addition of a low-rank symmetric matrix. One computes the eigen-decomposition of \( \mathbf{C} \) and then approximates the rank-\( k \) decomposition of \( \mathbf{C}_k + \mathbf{U} \) where \( \mathbf{U} = \mathbf{Y}_1 \mathbf{Y}_2^T + \mathbf{Y}_2 \mathbf{Y}_1^T \) and \( \mathbf{C}_k \) is the approximation of \( \mathbf{C} \) using the \( k \) largest eigenvectors (also known as the best \( k \)-rank approximation of \( \mathbf{C} \)). A similar but not applicable update is considered for the SVD case in [42] in which the rank-\( k \) approximation of \( \mathbf{A}_k + \mathbf{Y}_1 \mathbf{Y}_2^T \) is found from \( \mathbf{A}_k \) in which \( \mathbf{Y}_1 \in \mathbb{R}^{m \times j} \) and \( \mathbf{Y}_2 \in \mathbb{R}^{n \times j} \) for some \( j \).

The general idea in our case is to find a matrix with orthonormal columns \( \tilde{\mathbf{Q}} \) such that \( \mathbf{C}_k + \mathbf{U} = \mathbf{Q} \mathbf{\Delta} \mathbf{Q}^T \) for a square matrix \( \mathbf{\Delta} \). To this purpose, we first project the columns of \( \mathbf{Y}_1 \) into the space orthogonal to the \( k \) largest eigenvectors \( \mathbf{Q}_k \) of \( \mathbf{C} \) (note the deviation from standard notation), a process known as deflation. Assuming that eigenvectors have unit norm, the matrix \( \mathbf{Y}_1 \) is thus deflated as follows:

\[
\mathbf{Y}_1 = (\mathbf{I} - \mathbf{Q}_k \mathbf{Q}_k^T) \mathbf{Y}_1,
\]

at a cost of \( \mathcal{O}(npk) \). Note that \( \mathbf{Y}_1 \mathbf{\Theta}_1 \) for some \( \mathbf{\Theta}_1 \), is orthogonal to \( \mathbf{Q}_k \) since \( \mathbf{Q}_k^T \mathbf{Y}_1 \mathbf{\Theta}_1 = (\mathbf{Q}_k^T \mathbf{Y}_1 - \mathbf{Q}_k^T \mathbf{Y}_1) \mathbf{\Theta}_1 = 0 \). If we take the SVD \( \mathbf{Y}_1 = \mathbf{P}_1 \mathbf{\Sigma}_1 \mathbf{\Omega}_1 \) then \( \mathbf{P}_1 \) is orthogonal to \( \mathbf{Q}_k \) since \( \mathbf{P}_1 = \mathbf{Y}_1 \mathbf{\Omega}_1 \mathbf{\Sigma}_1^{-T} \) assuming \( \mathbf{\Sigma}_1 \) has nonzero diagonal entries.

At the next stage we would like to orthogonalise the columns of \( \mathbf{Y}_2 \) with respect to both \( \mathbf{Q}_k \) and \( \mathbf{P}_1 \). Hence we deflate \( \mathbf{Y}_2 \) in the following way:

\[
\mathbf{Y}_2 = (\mathbf{I} - \mathbf{P}_1 \mathbf{P}_1^T - \mathbf{Q}_k \mathbf{Q}_k^T) \mathbf{Y}_2,
\]

at cost \( \mathcal{O}(npk) \), where we have used the fact that \( \mathbf{P}_1 \) is orthogonal to \( \mathbf{Q}_k \).

Proved in a similar way to the step used earlier, the matrix in the column space of \( \mathbf{Y}_2 \) is orthogonal to \( \mathbf{Q}_k \) for some \( \mathbf{\Theta}_2 \), is orthogonal to \( \mathbf{Q}_k \) and \( \mathbf{P}_1 \). Hence, we compute the SVD \( \mathbf{Y}_2 = \mathbf{P}_2 \mathbf{\Sigma}_2 \mathbf{\Omega}_2 \) and note that the matrices \( \mathbf{P}_1 \), \( \mathbf{P}_2 \) and \( \mathbf{Q}_k \) are mutually orthogonal and span the space spanned by \( \mathbf{C}_k + \mathbf{U} \). This allows one to write

\[
\mathbf{C}_k + \mathbf{U} = \mathbf{Q} \mathbf{\Delta} \mathbf{Q}^T
\]

as required with \( \mathbf{Q} = [\mathbf{Q}_k, \mathbf{P}_1, \mathbf{P}_2] \in \mathbb{R}^{n \times (k+2p)} \) and \( \mathbf{\Delta} = \mathbf{Q}^T (\mathbf{C}_k + \mathbf{U}) \mathbf{Q} \), or equivalently

\[
\mathbf{\Delta} = \begin{bmatrix}
\mathbf{\Omega}_k + \mathbf{Q}_k^T \mathbf{U} \mathbf{Q}_k & \mathbf{Q}_k^T \mathbf{U} \mathbf{P}_1 & \mathbf{Q}_k^T \mathbf{Y}_1 \mathbf{Q}_2 \mathbf{\Sigma}_2 \\ 
\mathbf{P}_1^T \mathbf{U} \mathbf{Q}_k & \mathbf{P}_1^T \mathbf{U} \mathbf{P}_1 & \mathbf{\Sigma}_1 \mathbf{Q}_1^T \mathbf{Q}_2 \mathbf{\Sigma}_2 \\ 
\mathbf{\Sigma}_2 \mathbf{Q}_2^T \mathbf{Y}_1 \mathbf{Q}_k & \mathbf{\Sigma}_2 \mathbf{Q}_2^T \mathbf{Q}_1 \mathbf{\Sigma}_1 & 0
\end{bmatrix},
\]

in which \( \mathbf{\Delta} \in \mathbb{R}^{(k+2p) \times (k+2p)} \). We take the rank-\( k \) eigen-decomposition \( \mathbf{\Delta}_k = \mathbf{H}_k \mathbf{\Pi}_k \mathbf{H}_k^T \) and then the final eigen-approximation is given by \( (\mathbf{Q} \mathbf{H}_k) \mathbf{\Pi}_k (\mathbf{Q} \mathbf{H}_k)^T \) in which it is easy to verify that the columns of \( \mathbf{Q} \mathbf{H}_k \) are orthonormal.

The deflation and SVD of \( \mathbf{Y}_1 \) and \( \mathbf{Y}_2 \) cost \( \mathcal{O}(npk) \) and \( \mathcal{O}(np^2) \) respectively and the eigen-decomposition of \( \mathbf{\Delta} \) is \( \mathcal{O}((k+2p)^3) \). In order to compute \( \mathbf{\Delta} \) one can reuse the computations of \( \mathbf{Q}_k^T \mathbf{Y}_1 \), \( \mathbf{Q}_k^T \mathbf{Y}_2 \), \( \mathbf{P}_1^T \mathbf{Y}_1 \) and \( \mathbf{P}_2^T \mathbf{Y}_2 \) which are used for deflations and also the matrices used for the SVD decompositions. Thus \( \mathbf{\Delta} \) is found in \( \mathcal{O}(p^3 + p^2k + pk^2) \), and the overall complexity of this algorithm is
\( O((k^2 + p^2)(p + k) + np(p + k)) \). Of note here is that \( n \) scales the complexity in a linear fashion, however costs are cubically related to \( k \) and \( p \).

### 4.2 Addition of Rows and Columns

In correspondence with the SVD-updating method given above we consider the case in which one has the eigen-decomposition of \( C \) and then wants to find the rank-\( k \) approximation of \( E = [A \ B]^T[A \ B] \). Such a process is useful not just in incremental clustering but also in incrementally solving kernel Principal Components Analysis (KPCA, [33]) for example. This update can be written in terms of that described above. This is seen by writing the former update \( A^T \ A \rightarrow [A \ B]^T[A \ B] \) in terms of the latter:

\[
\begin{bmatrix}
A^T A & 0 \\
0 & 0
\end{bmatrix} \rightarrow \begin{bmatrix}
A^T A & 0 \\
0 & 0
\end{bmatrix} + \begin{bmatrix}
0 & A^T B \\
B^T A & B^T B
\end{bmatrix}.
\]

The second term on the right-hand side can be written as \( Y_1 Y_1^T + Y_2 Y_2^T \) where \( Y_1 = [0 \ I_p]^T \) and \( Y_2 = [B^T A \ \frac{1}{2} B^T B]^T \). The eigenvectors of the first matrix on the right-hand side are found from those of \( A^T \ A \) by simply adding \( p \) zero rows to the existing eigenvectors, and the corresponding eigenvalues are identical. Additional eigenvectors are standard unit vectors spanning the \( p \) new rows with corresponding eigenvalues as zero. A useful insight is that the deflated matrix \( \tilde{Y}_1 = Y_1 \) and hence its SVD decomposition can be written directly as \( \tilde{Y}_1 = [0 \ I_p]^T I_p I_p \).

#### 4.2.1 Alternative Approach

Here we outline a simpler and more direct approach for the addition of rows and columns to a matrix. First let \( C = Q \Omega Q^T \) in which \( Q \) is a matrix of eigenvectors and \( \Omega \) is a diagonal matrix of eigenvalues. Note that \( \tilde{E} = [A_k \ B]^T[A_k \ B] \) can be written as \( Q \tilde{\Delta} Q^T \) for a square matrix \( \Delta \). In our case we have

\[
\tilde{Q} = \begin{bmatrix}
Q_k & 0 \\
0 & I_p
\end{bmatrix}
\quad \text{and} \quad
\Delta = \begin{bmatrix}
\Omega_k & Q_k^T A_k^T B \\
B^T A_k Q_k & B^T B
\end{bmatrix},
\]

noting that \( A_k Q_k Q_k^T = Q_k \Omega_k Q_k^T Q_k^T = A_k \) since \( Q_k Q_k^T = I \). Furthermore, note that \( Q_k^T A_k^T B = Q_k^T Q_k \Sigma_k \pi_k^T B = Q_k^T A_k^T B = Q_k^T A_k^T B \) and calculating \( Q_k^T A_k^T B \) is \( O(npk) \). It follows that \( \Delta \in \mathbb{R}^{(k+p) \times (k+p)} \) can be found using \( Q_k, A_k^T A, A_k^T B \) and \( B^T B \). In the final step, and analogous to the SVD case we take the rank-\( k \) eigen-decomposition \( \Delta = H_k \Pi_k H_k^T \) at a cost of \( O((k+p)^3) \) and then the rank-\( k \) eigen-approximation of \( \tilde{E} \) is given by \( (Q_k H_k) \Pi_k (Q_k H_k)^T \).

Notice that this eigen-update almost follows directly from the SVD update in [42] however one need not use the QR decomposition of the deflated \( B \).

Furthermore, the eigen-approximation above is identical to that at the start of Section 4.2 with \( Y_1 = [0 \ I_p]^T \) and \( Y_2 = [B^T A_k \ \frac{1}{2} B^T B]^T \). The difference between the method above and that of Section 4.2 is the former uses \( A_k \) as opposed to \( A \) in \( Y_2 \) which results in a greater error.

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4.3 Removing Rows and Columns

Observe that removing rows and columns is equivalent to zeroing the corresponding rows/columns:

\[
\hat{C} = \begin{bmatrix} A^T A & A^T B \\ B^T A & B^T B \end{bmatrix} \rightarrow \begin{bmatrix} A^T A & A^T B \\ B^T A & B^T B \end{bmatrix} - \begin{bmatrix} 0 & A^T B \\ B^T A & B^T B \end{bmatrix},
\]

and in this form one can see the connection to Section 4.2. Again, one can write the second term on the right-hand side as \( Y_1 Y_2^T + Y_2 Y_1^T \) where \( Y_1 = [0 \quad I_p]^T \) and \( Y_2 = -[\tilde{B}^T \tilde{A} \quad \frac{1}{2} \tilde{B}^T \tilde{B}]^T \) where \( \tilde{B}^T \tilde{A} \) and \( \tilde{B}^T \tilde{B} \) are found using the rank-\( k \) approximation of \( \hat{C} \). Since we are updating the rank-\( k \) approximation of \( \hat{C} \), the final eigen-approximation will have zero elements in the eigenvectors at rows corresponding to those row/columns that are deleted.

4.4 Discussion

A similar eigen-update to that of Section 4.2.1 is considered in [18] and used in conjunction with PCA [15] and KPCA in [43]. In PCA, one starts with a set of examples \( T = \{x_1, x_2, \ldots, x_n\} \) where \( x_i \in \mathbb{R}^d \). These examples are centred, and then one finds the \( k \) largest eigenvectors \( u_1, \ldots, u_k \) of the covariance matrix \( \mathbf{C} = \frac{1}{n} \sum_{i=1}^{n} \tilde{x}_i \tilde{x}_i^T \), where \( \tilde{x} \) is a centred example. KPCA functions similarly, except that one finds the largest eigenvectors of a kernel matrix \( \mathbf{K} \in \mathbb{R}^{n \times n} \) which is computed using the centred examples. In the incremental setting one approximates the eigenvectors of the covariance or kernel matrix on the addition of a new set of examples. In [43] the authors phrase the problem as a series of SVD updates, and we use a more direct approach in Section 4.2.1. Furthermore, whereas [43] examines the addition of rows and columns to a positive semi-definite matrix, our work is more general in that we additionally consider removal of rows and columns and addition of a low-rank symmetric matrix. One key novelty of this paper is the update of Section 4.1 which encapsulates all three of these updates and yet does not simply follow from the SVD updating work of [42].

To formalise this notion, the updates outlined above can be written in terms of the addition of a low-rank symmetric matrix \( \mathbf{U} \) to a positive semi-definite matrix \( \mathbf{C} \). Our approximation method computes \( \mathbf{C}_k + \mathbf{U} \) via its expression as \( \tilde{Q} \mathbf{H}_k \Pi_k \mathbf{H}_k^T \tilde{Q}^T \) where \( \tilde{Q} \) is a matrix with orthogonal columns spanning \( \mathbf{C}_k + \mathbf{U} \) and \( \mathbf{H}_k \) and \( \Pi_k \) represent the largest \( k \) eigenvectors and eigenvalues of a matrix \( \Delta = \tilde{Q}^T (\mathbf{C}_k + \mathbf{U}) \tilde{Q} \). The following lemma shows the consequence of this approach.

**Lemma 4.1.** Decompose a matrix \( \mathbf{Z} = \mathbf{F} \mathbf{A} \mathbf{F}^T \) where \( \mathbf{F} \) is any matrix with orthonormal columns \( \mathbf{F}^T \mathbf{F} = \mathbf{I} \). For some \( k \) find the best rank-\( k \) eigen-approximation \( \Lambda_k = \mathbf{H}_k \Pi_k \mathbf{H}_k^T \) in which \( \mathbf{H}_k \) and \( \Pi_k \) are the largest \( k \) eigenvectors and eigenvalues of \( \Lambda \), and let \( \hat{\mathbf{Z}} = \mathbf{F} \mathbf{H}_k \Pi_k \mathbf{H}_k^T \mathbf{F}^T \). Then the best rank-\( k \) eigen-decomposition of \( \mathbf{Z} \) is given by:

\[
\hat{\mathbf{Z}} = \mathbf{U}_k \mathbf{S}_k \mathbf{U}_k^T = \mathbf{Z}.
\]
Proof. Note that $\Lambda = F^T Z F$ due to the orthogonality of $F$. Let $u, s$ be the eigenvectors and eigenvalues of $Z$, and define $u = F v$ for some $v$ then $F^T Z F v = s v$. This implies that the eigenvalues of $\Lambda$ are the same as those of $Z$ and the eigenvectors are related by $U = F V$ which implies $Z = F H_k U_k H_k^T F^T = U_k S_k U_k^T = Z_k$ as required. The only condition on $F$ is that the eigenvectors of $Z$ are in the column space of $F$. Hence we have $V = H$ and $S = \Pi$ which implies $\hat{Z} = F H_k \Pi_k H_k^T F^T$. Hence the update $C_k + U$ of Section 4.1 is identical to the best rank-$k$ approximation of $C_k + U$.

5 Eigen-approximation Quality

This section aims to bound the quality of the proposed eigen-approximation approach. As mentioned in Section 4.4, the proposed approach replaces the expected best rank-$k$ approximation of a matrix $A + B$ by the best rank-$k$ approximation of $A_k + B$, where $A_k$ denotes the best rank-$k$ approximation of $A$. Hence, our objective is to control the difference between $(A + B)_k$ and $(A_k + B)_k$.

The result in Section 4.4 gives us a first insight into the approximation error of the updates described: the residual matrix is that corresponding to the eigenvectors and eigenvalues after $k$. One can see that for any matrix $C$

$$\|C - C_k\|_F^2 = \|C_{k+1}\|_F^2 = \sum_{i=k+1}^n \omega_i^2,$$

where $\omega_i$ is the $i$th eigenvalue of $C$ (unless otherwise stated eigenvalues are always given in descending order) and $C_{k+1}$ is the approximation of $C$ using eigenvectors/eigenvalues after $k$. This implies that $C$ is well approximated by the largest $k$ eigenvectors if the sum of the square of the remaining eigenvalues is small.

This certainly gives us insight into when our eigen-updating approach will be accurate, however the kind of matrices we will work with do not have this property in general. We now turn to matrix perturbation theory [35] in order to learn more about the approximated eigenvectors of the updated matrix. In a nutshell, it lies in controlling the angle between two invariant subspaces. Before giving the corresponding theorem, the following subsection introduces the necessary notions of invariant subspaces of a matrix and of the canonical angles between subspaces.

5.1 Invariant Subspaces and Canonical Angles

We begin by introducing the simple concept of an invariant subspace.

Definition 5.1. The subspace $\mathcal{X}$ is an invariant subspace of $A$ if $A \mathcal{X} \subset \mathcal{X}$.

It can also be shown that if the columns of $X$ form a basis for $\mathcal{X}$ of $A$ then there is a unique matrix $L$ such that $AX = XL$. The matrix $L$ is a representation of $A$ with respect to the basis $X$, and it has identical eigenvalues to $A$. A useful decomposition in perturbation theory is to reduce $A$ to a block diagonal form. Let $X_1$ be an orthogonal matrix which spans the invariant
subspace of $A$, $X_1$, and assume that we have a matrix $Y_2$ such that $[X_1Y_2]$ is unitary and $Y_2$ spans the space orthogonal to $X_1$, then this allows us to write

$$[X_1Y_2]^T A [X_1Y_2] = \begin{bmatrix} L_1 & H \\ 0 & L_2 \end{bmatrix},$$

where $L_1 = X_1^T A X_1$, $L_2 = Y_2^T A Y_2$ and $H = X_1^T A Y_2$. The above equation is known as the reduced form of $A$ with respect to $[X_1Y_2]$. The proof of why the bottom left block of this matrix is zero is straightforward, see [35] for details. This gives us the knowledge to define a simple invariant subspace.

**Definition 5.2.** Let $X$ be an invariant subspace of $A$ and consider the reduced form of Equation (1), then $X$ is a simple invariant subspace of $A$ if there are no common eigenvalues between $L_1$ and $L_2$.

Notice that a simple invariant subspace has a complementary space, defined as follows.

**Definition 5.3.** Let the simple invariant subspace $X_1$ have the reduced form of Equation (1) with respect to the orthogonal matrix $[X_1Y_2]$. Then there exist $X_2$ and $Y_1$ such that $[X_1X_2]^{-1} = [Y_1Y_2]^T$ and

$$A = X_1 L_1 Y_1^T + X_2 L_2 Y_2^T,$$

where $L_i = Y_i^T A X_i$, $i = 1, 2$. This form of $A$ is known as the spectral resolution of $A$ along $X_1$ and $X_2$.

This allows us to introduce a theorem essential to our main result. However, first we must define the notion of angle between two subspaces.

**Theorem 5.1.** Let $X_1, Y_1 \in \mathbb{R}^{n \times \ell}$ with $X_1^T X_1 = I$ and $Y_1^T Y_1 = I$. If $2 \ell \leq n$, there are unitary matrices $Q, U_{11}, V_{11}$ such that

$$Q X_1 U_{11} = \begin{pmatrix} I_{\ell} \\ 0 \\ 0 \end{pmatrix} \quad \text{and} \quad Q Y_1 V_{11} = \begin{pmatrix} \Gamma \\ \Sigma \\ 0 \end{pmatrix},$$

in which $\Gamma = \text{diag}(\gamma_1, \ldots, \gamma_\ell)$ with $0 \leq \gamma_1 \leq \ldots \leq \gamma_\ell$, $\Sigma = \text{diag}(\sigma_1, \ldots, \sigma_\ell)$ with $\sigma_1 \geq \cdots \geq \sigma_\ell \geq 0$, and $\gamma_i^2 + \sigma_i^2 = 1$, $i = 1, \ldots, \ell$. If $2 \ell > n$ then $Q, U_{11}, V_{11}$ can be chosen so that

$$Q X_1 U_{11} = \begin{pmatrix} I_{n-\ell} & 0 \\ 0 & I_{2\ell-n} \end{pmatrix} \quad \text{and} \quad Q Y_1 V_{11} = \begin{pmatrix} \Gamma & 0 \\ 0 & I_{2\ell-n} \end{pmatrix},$$

in which $\Gamma = \text{diag}(\gamma_1, \ldots, \gamma_{n-\ell})$ with $0 \leq \gamma_1 \leq \ldots \leq \gamma_{n-\ell}$, $\Sigma = \text{diag}(\sigma_1, \ldots, \sigma_{n-\ell})$ with $\sigma_1 \geq \cdots \geq \sigma_{n-\ell} \geq 0$, and $\gamma_i^2 + \sigma_i^2 = 1$, $i = 1, \ldots, n-\ell$.

Geometrically, let $X_1$ and $Y_1$ be subspaces of dimension $\ell$ and $Q$ be a unitary transformation. Then the matrices $Q X_1 U_{11}$ and $Q Y_1 V_{11}$, with $X_1 \in X_1$, $X_2 \in X_2$, $Q \in \mathbb{Q}$, form bases of $Q X_1$ and $Q Y_1$ and $\sigma_i$ and $\gamma_i$ can be regarded as sines and cosines of the angles between the bases. We now define a measure of similarity between subspaces.
Definition 5.4. Let $\mathcal{X}$ and $\mathcal{Y}$ be subspaces of the same dimension, then the canonical angles between the subspaces are the diagonal entries of the matrix $\sin^{-1} \Sigma = \text{diag}(\sin^{-1}(\sigma_1), \ldots, \sin^{-1}(\sigma_{n-k}))$ where $\Sigma$ is the matrix defined in Theorem 5.1.

It follows that $\Sigma$ is a measure of how two subspaces differ.

5.2 Angle Between Exact and Updated rank-$k$ Approximations

We are now ready to state the main theorem required for our result.

Theorem 5.2. Let $A$ be a Hermitian matrix with spectral resolution given by $[X_1 X_2]^T A [X_1 X_2] = \text{diag}(L_1, L_2)$ where $[X_1 X_2]$ is unitary. Let $Z \in \mathbb{R}^{n \times k}$ have orthonormal columns, $M$ be any Hermitian matrix of order $k$ and define the residual matrix as $R = AZ - ZM$. Let $\lambda(A)$ represent the set of eigenvalues of $A$ and suppose that $\lambda(M) \subset [\alpha, \beta]$ and for some $\delta > 0$, $\lambda(L_2) \subset \mathbb{R} \setminus [\alpha - \delta, \beta + \delta]$. Then, for any unitary invariant norm, we have:

$$\| \sin \Theta(\mathcal{R}(X_1), \mathcal{R}(Z)) \| \leq \frac{\| R \|_F}{\delta}$$

where $\mathcal{R}(\cdot)$ is the column space of a matrix.

Before we introduce the main result we present a result by Weyl [38] which characterises the perturbation in the eigenvalues of a matrix.

Theorem 5.3 (Weyl, [38]). Define $A \in \mathbb{R}^{n \times n}$ and let $\hat{A} = A + E$ be its perturbation. The eigenvalues of $A$ and $E$ are given by $\lambda_i$ and $\epsilon_i$, $i = 1, \ldots, n$, respectively. Then the eigenvalues of $\hat{A}$ are, for $i = 1, \ldots, n$, $\hat{\lambda}_i \in [\lambda_i + \epsilon_n, \lambda_i + \epsilon_1]$.

We can now present our main result which is closely related to the Davis-Kahan theorem [4].

Theorem 5.4. Consider a positive semi-definite matrix $A \in \mathbb{R}^{n \times n}$ with eigenvalues $\omega_1, \ldots, \omega_n$ and corresponding eigenvectors $Q = [q_1, \ldots, q_n]$. Let $B \in \mathbb{R}^{n \times n}$ be symmetric with eigenvalues $\epsilon_i$ and $(A + B)$ be positive semi-definite with eigen-decomposition $U \Sigma U^T$ where $\gamma_i$ are eigenvalues, $i = 1, \ldots, n$. Fix integer $k$, let $A_k + B$ have decomposition $V_k \Sigma_k V_k^T$ with eigenvalues $\pi_1, \ldots, \pi_k$ and assume $\gamma_k \neq \gamma_{k+1}$. Then the following bounds hold on the canonical angles between the subspaces defined by $U_k$ and $V_k$, assuming $\pi_k > \gamma_{k+1}$,

$$\| \sin \Theta(\mathcal{R}(U_k), \mathcal{R}(V_k)) \|_F \leq \frac{\sqrt{\text{tr}(V_k^T A_k^2 V_k)}}{\pi_k - \gamma_{k+1}}$$

(4)

$$\| \sin \Theta(\mathcal{R}(U_k), \mathcal{R}(V_k)) \|_2 \leq \frac{\lambda_{\text{max}}(V_k^T A_k^2 V_k)}{\pi_k - \gamma_{k+1}}$$

(5)

where $\gamma_{k+1} = \omega_{k+1} + \pi_{k+1}$. 

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Proof. We will start by considering the first bound. It is clear that $U_k$ is a simple invariant subspace for $(A + B)$. The spectral resolution of $A + B$ is given by $[U_k U_{k+1}]$ since this matrix is unitary and we have $A + B = U_k U_{k+1}^T (A + B) U_k U_{k+1}^T$. In the reduced form $L_1 = \Gamma_k$ and $L_2 = \Gamma_{k+1}$. Furthermore, we set $M = \Pi_k$ and $Z = V_k$. The residual matrix is given by $R = (A + B)V_k - V_k \Pi_k = A_k \perp V_k$ and

$$
\|R\|_F = \sqrt{\text{tr}(V_k^T A_k^2 \perp V_k)}.
$$

We know that the eigenvalues of $M$ fall within the range $[\pi_k, \pi_1]$ and those of $L_2$ are bounded using Theorem 5.3 in the range $\gamma_i \in [\pi_i + \omega_n, \pi_i + \omega_{k+1}]$ for $i = k + 1, \ldots, n$. Considering also the perturbation of eigenvalues of $A$ we can write $\gamma_i \leq \tilde{\gamma}_i = \min(\omega_{k+1} + \pi_{k+1}, \omega_{k+1} + \epsilon_1) = \omega_{k+1} + \pi_{k+1}$ given $\pi_{k+1} \leq \omega_{k+1} + \epsilon_1$. It follows that $\delta = \pi_k - \tilde{\gamma}_{k+1}$ and plugging into Theorem 5.2 gives the required result.

The second bound is proved similarly except in this case we have

$$
\|R\|_2 = \sqrt{\lambda_{\text{max}}(V_k^T A_k^2 \perp V_k)}.
$$

Thus we have a bound on the angle between the subspace of the first $k$ eigenvectors of $A_k + B$ and the corresponding eigenvectors of its perturbation $A + B$ without explicitly requiring the eigen-decomposition of $A + B$. Provided the eigenvalues of $V_k^T A_k^2 \perp V_k$ are small and the eigengap $\pi_k - \tilde{\gamma}_{k+1}$ is large one can be sure that the two subspaces have small canonical angles. One can see that under small perturbations $V_k$ is close to $Q_k$ and hence $V_k^T A_k^2 \perp V_k$ is small, resulting in tight bounds in the angles. In the case that the matrices involved correspond to normalised Laplacians this result corresponds well with similar results outlining a perturbation-based motivation of spectral clustering (see e.g. [37]) which state that if the value of $\gamma_k - \gamma_{k+1}$ is large then one might reasonably expect a good clustering. The bound becomes loose when this eigengap is small, however in this case the clusters are less distinct even when computing the exact eigenvectors.

6 Incremental Cluster Membership

We now return to the eigenproblem of Algorithm 1, $\tilde{L}v = \lambda v$, in which we are interested in the eigenvectors with the smallest eigenvalues. Define the shifted Laplacian as

$$
\tilde{L} = 2I - \tilde{L} = I + D^{-\frac{1}{2}} WD^{-\frac{1}{2}},
$$

which is positive semi-definite since $\tilde{L}$ is positive semi-definite with largest eigenvalue 2. Note that by negating a matrix one negates the eigenvalues, leaving the eigenvectors the same, and similarly an addition of $\sigma I$ increases the eigenvalues by $\sigma$ leaving the eigenvectors intact. Since we are interested in the smallest eigenvectors of $\tilde{L}$ they correspond exactly to the maximum eigenvectors of $\tilde{L}$ and we can use the eigen-update methods described above. Observe that the
shifting Laplacian is a normalised version of the signless Laplacian [13] defined as \( L^+ = D + W \), and this can be seen from \( \tilde{L} = D^{-\frac{1}{2}}(D + W)D^{-\frac{1}{2}} = D^{-\frac{1}{2}}L^+D^{-\frac{1}{2}} \).

Putting the ingredients together allows us to outline an efficient incremental method for performing graph clustering called Incremental Approximate Spectral Clustering (IASC), see Algorithm 3. At a high level the algorithm is quite simple: in the initialisation steps one computes the shifted Laplacian matrix for the first graph \( \tilde{L}_1 \) and then performs k-means clustering using the largest \( k \) eigenvectors of this matrix. For the \( t \)-th successive graph, \( t > 1 \), we use the eigen-update methods above to approximate the largest eigenvectors of \( \tilde{L}_t \) using the approximate eigenvectors computed at the previous iteration in step 10. For these updates, it is simple to recover the matrices \( Y_1 \) and \( Y_2 \) given a change in edge weights. Note that we use the first method of Section 4.2 to compute eigenvector upon the addition of row and columns. Furthermore, we store the first \( \ell \) eigenvectors of \( \tilde{L}_1 \), with \( \ell \geq k \), and recompute eigenvectors every \( T \) iterations in order to reduce cumulative errors introduced in the loop at the expense of increased computation. When \( \ell = n \) one recovers the exact eigenvectors at each iteration and Algorithms 3 and 1 become nearly equivalent. In the case that there is a significant eigengap between eigenvalues, one can fix both \( \ell \) and \( k \) according to Theorem 5.4.

**Algorithm 3** Incremental Approximate Spectral Clustering

**Require:** Graphs \( G_1, \ldots, G_T \) of sizes \( n_1, \ldots, n_T \), no. clusters \( k \), approximation rank \( \ell \geq k \), eigen-decomposition recomputation step \( R \)

1: Compute the shifted Laplacian for \( G_1, \tilde{L}_1 \)
2: Find \( \ell \) largest eigenvectors of \( \tilde{L}_1 \), \( V^{(1)}_k \), let \( V^{(1)}_k \) be the first \( k \) cols of \( V^{(1)}_k \)
3: Normalise the eigenvector rows, \( V^{(1)}_k \leftarrow \text{diag}(V^{(1)}_k(V^{(1)}_k)^T)^{-\frac{1}{2}}V^{(1)}_k \)
4: Use k-means on rows of \( V^{(1)}_k \) and store indicators \( c_1 \in \{1, \ldots, k\}^{n_1} \)
5: for \( t = 2 \rightarrow T \) do
6: Compute shifted Laplacian for \( G_t, \tilde{L}_t \)
7: if \( i \% R = 0 \) then
8: Recompute eigenvectors of \( \tilde{L}_t \)
9: else
10: Use rank-\( \ell \) eigen-approximation of Section 4
11: end if
12: Normalise rows of \( V^{(t)}_k \), \( V^{(t)}_k \leftarrow \text{diag}(V^{(t)}_k(V^{(t)}_k)^T)^{-\frac{1}{2}}V^{(t)}_k \)
13: Use k-means on \( V^{(t)}_k \) with initial centroids \( c_{t-1} \), store \( c_t \in \{1, \ldots, k\}^{n_t} \)
14: end for
15: return Cluster membership \( c_1 \in \{1, \ldots, k\}^{n_1}, \ldots, c_T \in \{1, \ldots, k\}^{n_T} \)

The complexity of Algorithm 3 is dictated by the sparsity of the graphs and the extent of the change between successive graphs. We ignore the steps before the for-loop since in general they do not impact the overall complexity. At iteration \( t \) and step 6 one can compute \( \tilde{L}_t \) from the weight and degree matrix at a cost of \( O(n_t + |E_t|) \). In the following step if there is a change between edges incident to vertices \( S = \{v_{t_1}, \ldots, v_{t_2}\} \) then the rows and columns corresponding to the union of the neighbours of \( S \), \( n(S) \), will change in the corresponding shifted Laplacian. In this case \( p = |n(S)| \) in \( Y_1, Y_2 \in \mathbb{R}^{n \times p} \) and if the neighbourhood of the vertices with changed edges \( n(S) \) is small, this
update can be efficiently computed as outlined in Section 4. The cost of $k$-means is $O(k^2ns)$ where $s$ is the number of iterations required for convergence.

7 Computational Results

In this section we study the clustering quality of five incremental strategies: the naive one which computes the exact eigenvectors at each iteration, denoted Exact, Ning et al.’s method (Ning) since it is a competing incremental strategy, IASC, Nyström, which uses the Nyström eigen-decomposition approximation of the shifted Laplacian, since it is often used in spectral clustering, and likewise the Randomised SVD method of Algorithm 2 denoted RSVD. First, however we observe the quality of the eigenvectors found using our eigen-updating approach.

7.1 Quality of Approximate Eigenvectors

We compare Nyström and RSVD approximations with our eigen-updating methods on a synthetic dataset generated in the following way: The initial graph contains 4 clusters of size 250 which are generated using an Erdős-Rényi [8] process with edge probability $p = 0.1$. The Erdős-Rényi process creates edges independently randomly with a fixed probability $p$ for all pairs of vertices. For each successive graph we then add 50 random edges to simulate “noise” in the clusters for a total of 100 graphs. We then compute the largest $k = 4$ eigenvectors of the shifted Laplacian using the full eigen-decomposition, the Nyström method, RSVD and the eigen-updating approach of Section 4. For the Nyström method we use $m = 900$ randomly selected columns, for RSVD we use $r = \{100, 900\}$ random projections, and for the eigen-updating approach we update based on $\ell = \{4, 300\}$ approximate eigenvectors and eigenvalues found for the previous graph. The quality of the approximations is measured by $\|\sin \Theta(R(U_k), R(V_k))\|_F$ where $U_k$ stands for the exact eigenvectors and $V_k$ corresponds to the eigenvectors returned by one of the three considered eigen-approximation strategies. Applying Corollary I.5.4 of [35], this norm is equal to $\|U_k^\perp V_k\|_F$. The experiment is repeated with results averaged over 20 iterations.

Observe that the quality of the approximation decreases with the noise amplitude, see Figure 1(a). For the three approximation strategies, the more eigenvectors/columns/random projection they consider, the smaller the canonical angles with the exact decomposition as one might expect. Nyström leads to a poor approximation whilst using 90% of the columns. RSVD has poor results with 100 random projections, but is close to the exact decomposition with 900 random projections. The eigen-update strategy always does a better job than keeping the initial decomposition, and in this case there does not appear to be a large gain when moving from 300 eigenvectors to just 4. In the case in which we use the complete set of eigenvectors we would expect IASC to coincide with exact eigen-decomposition.

7.1.1 Perturbation Bound

We now turn to demonstrating the effectiveness of the bound of Theorem 5.4 on a synthetic dataset containing 150 vertices. The initial graph contains 3
clusters of size 50 which are generated using an Erdős-Rényi process with edge probability \( p = 0.3 \), resulting in 1092 edges. For each successive graph we add 10 random edges and in total there is a sequence of 80 graphs. We then compute the bound of Theorem 5.4 to measure the difference between the canonical angles of the real and approximated eigenvectors of the shifted Laplacian. We compare the results to the bound using the real eigenvalues of the Laplacian, i.e. \( \delta = \pi_k - \gamma_{k+1} \). This process is repeated 50 times with different random seeds and the results are averaged.

![Graphs](image)

Figure 1: The left plot compares the canonical angles of the largest eigenvectors found using the approximation methods with the equivalent real eigenvectors. On the right we compare the perturbation bounds of Theorem 5.4. By “precise” we mean that we use \( \delta = \pi_k - \gamma_{k+1} \) in the theorem.

Figure 1(b) shows the resulting bounds for this sequence of graphs. The bound diverges slowly from the precise bound for many of the initial graphs, for example \( \| \sin \Theta(\mathcal{R}(U_k), \mathcal{R}(V_k)) \|_F \) is bounded by 0.616 ± 0.106 versus 0.352 ± 0.024 at the 50th graph and the equivalent results for the 2-norm are 0.434 ± 0.007 and 0.248 ± 0.02. Soon after this point, we see a large divergence in the precise and approximate bounds, although the approximate bounds become trivial after the 78th graph for the Frobenius norm and the 73rd for the 2-norm corresponding to the addition of 770 and 730 edges respectively to the initial graph. When we look at the precise bound, one can see that at the last graph \( \| \sin \Theta(\mathcal{R}(U_k), \mathcal{R}(V_k)) \|_2 \leq 0.39 \) despite nearly doubling the number of edges which points to the precision of the eigen-approximation in this case.

7.2 Clustering on Synthetic Data

To evaluate the clustering approaches, two separate synthetic datasets are considered, both of which are generated by an Erdős-Rényi process. The first dataset, called 3clust, is based on a graph of 3 clusters of 60 vertices each. This dataset allows us to compare the clustering quality as the clusters become more/less distinct and also the addition and removal of edges. In the corresponding graph, any possible edge between two points from the same cluster occurs with probability \( p_c = 0.3 \) and the probability of edges between vertices in different clusters is selected from \( p_g \in \{0.1, 0.2\} \). To generate a sequence of graphs we first allow only 20 vertices per cluster and then add 5 vertices to each cluster at a time until each one is of size 60. We then reverse the process so that
the 10th graph is the same as the 8th, and the 11th is the same as the 7th etc., which allows us to test the clustering methods upon the removal of vertices.

The second dataset, discrepancy, aims to examine the clustering methods on a more complex set of hierarchical clusters. Here, 180 vertices are split into 3 clusters of equal size, and inside each cluster there are 3 sub-clusters. The initial graph is empty, and at each iteration \( t \in \{1, \ldots, T\}, T = 23 \), an edge \( \{v_i, v_j\} \) is added with a probability \( p_\varepsilon + (p - p_\varepsilon)|t/T| \), where \( \ell \) is set to 1 if \( v_i \) and \( v_j \) are in the same subgroup, 0.5 if \( v_i \) and \( v_j \) are in the same group, and 0 otherwise. In our case we set \( p_\varepsilon = 0.0005 \) and \( p = 0.01 \).

For both datasets and the clustering approaches, \( k \)-means is run with \( k \) corresponding to the number of clusters (3 and 9 for 3clust and discrepancy respectively). For IASC we fix the number of eigenvectors \( \ell \in \{3, 6, 12, 24\} \) with the 3clust dataset and \( \ell \in \{9, 72\} \) with discrepancy in order to test the approximation quality as this parameter varies. With Nyström we sample \( m = 90 \) columns of the Laplacian matrix to find the approximate eigenvectors on 3clust and select \( m \in \{9, 72\} \) for discrepancy. RSVĐ uses \( r = 24 \) random projections on 3clust and \( r \in \{9, 72\} \) for discrepancy. On discrepancy, the approximation methods start with the 3rd graph in the sequence to allow the initial graph to contains enough edges. The experiments are repeated 50 times with different random graphs constructed using the methods described above, and the results are averaged. Clustering accuracy is measured through the Rand Index [29] between the finest true clustering \( C \) and the learned one \( \tilde{C} \). Rand Index corresponds to the proportion of true answers to the question “Are vertices \( v_i \) and \( v_j \) in the same cluster?”. More formally, it is given by

\[
\text{RandIndex}(C, \tilde{C}) = \frac{|\{v_i, v_j \in V : v_i \neq v_j, \delta(C(v_i), C(v_j)) \neq \delta(\tilde{C}(v_i), \tilde{C}(v_j))\}|}{|\{v_i, v_j \in V : v_i \neq v_j\}|},
\]

where \( C(v_i) \) stands for the cluster index of vertex \( v_i \) after clustering \( C \), \( \delta \) is the Kronecker delta function and \(|E|\) denotes the cardinality of any finite set \( E \).

The evaluation is completed by the computation of \( \|\sin\Theta(U_k, V_k)\|_F \), see Section 7.1 for more details. The canonical angles for Ning are not given as Ning uses the random-walk Laplacian which (i) differs from the normalised Laplacian used with other approaches and (ii) is not symmetric, leading to non-orthogonal eigenvectors.

The errors on 3clust are shown in Figure 2. The first point to note is that the errors decrease as the cluster size increases up until graph 8 and then increase again as vertices are removed. This is explained by the fact that as the cluster size increases there are more edges within each cluster relative to those between clusters, hence they becomes easier to identify. Notice also that the approximation methods perform worse (in terms of Rand Index) than exact except for Ning and IASC \( \ell = 3 \) when vertices are removed. This exception is due to a coincidence: for any iteration \( i \in 8, \ldots, 16 \), clustering results are better when using the eigen-decomposition at time 8 as they are based on more data. On the other hand, Ning and IASC, \( \ell = 3 \), do not accurately update the eigen-decomposition and hence the eigenvectors are close to those at graph 8. As a consequence, both approaches lead to better results than exact. Apart from this exception, IASC, \( \ell = 24 \), and RSVĐ have results close to exact, while Nyström leads to a bad Rand Index score for \( p = 0.2 \). Ning has non-smooth behaviour for any value of \( p \). Lastly, the results of Nyström when \( p = 0.1 \) demonstrate the
Figure 2: The mean Rand Index error of the learned clustering on 3clust. The numbers after IASC denote the number of eigenvectors computed.

limit of canonical angles to measure the clustering quality of an approximation approach: having large canonical angles values does not necessarily imply a Rand Index score far from exact.

It is worth noting that since $\ell$ is a fixed value for IASC, the approximation of the largest eigenvectors of the shifted Laplacian represents a smaller fraction of the total sum of eigenvalues as the cluster size increases. When using 24 eigenvectors for the approximation on the 2nd largest graph one requires approximately 14.5% of the dimensionality of the Laplacian, yet Rand Index is close to that of exact.

The results on discrepancy (Figure 3) show that Ning generally does not generate accurate clustering until the 12th graph when eigenvectors are recomputed, however it becomes competitive after this point. This is partly due to the fact that one solves a different eigen-system to the other methods which is not as effective at clustering for the initial graphs. As we increase the number of columns used for the Nyström approach or the random projections used by RSVD, the Rand Index values improve for the first few graphs in which the clusters are generally not well defined. In contrast, IASC results are relatively accurate with 72 eigenvectors. Surprisingly at the final graph we see that with just 9 eigenvectors IASC results in the most accurate clustering. At this point more eigenvectors
Figure 3: The mean Rand Index of the learned clusterings on discrepancy. For IASC and Nyström the numbers in the legend indicates the number of eigenvectors and columns sampled at each iteration.

seem to make the solution worse, with the exact eigen-decomposition being less accurate than IASC with 72 eigenvectors. One explanation is that an accurate eigen-decomposition fits \textquotedblleft noise\textquotedblright{} in the Laplacian which is excluded with IASC and a low value of $\ell$.

7.3 Real-world Graphs

We now apply the clustering methodology analysed previously on several real datasets.

7.3.1 Setup

Here we use the clustering methods on three real datasets. The first one represents individuals in Cuba who are detected as HIV positive between the period 1986 and 2004, see [1] for details of the related database. Edges in the graph indicate the occurrence of a sexual contact between two individuals as determined using contact tracing, whereby contacts of an infected person are identified and tested. The full sexual contact graph at the end of 2004 consists of 5389 people however it is strongly disconnected and we consider the growth of the largest component of size 2387. We find graphs of detected individuals at 1 month intervals, starting when the graph contains at least 500 vertices. At any particular time point, we consider the component containing first person detected in the largest component at the end of the recorded epidemic.

The next dataset is the high energy physics theory citation network from the Arxiv publications database [12] for the period February 1992 to March 2002. The full dataset contains 27,770 papers with 352,807 edges and a graph is generated as follows: if a paper cites another, an edge is made from the former to the latter. Not all of the papers present in the dataset have publication dates and hence we use only those with dates and label cited papers with the date of the oldest citing paper. Taking the largest component, the final resulting graph consists of 15,112 vertices and 193,826 edges. To track the evolution of the
connected component we start with the oldest paper, and consider the graphs of connected papers at 1 month intervals, starting with a graph of at least 500 vertices.

The final dataset, called Bemol, was first introduced in [30] and corresponds to the purchase history of an e-commerce website over a period of almost two years. The initial dataset is a bipartite graph between users and products composed of more than 700,000 users and 1,200,000 products. In the current experiment we focus on the first 10,000 users, and a graph is constructed between users with edge weights corresponding to the number of commonly purchased products between two users. Taking a maximum of 500 purchases per iteration in the graph sequence we focus on graphs 500 to 600.

To test the clustering methods we run each on the sequences of evolving graphs under a variety of parameters. As the selection of the number of clusters is a complex issue and outside the scope of this paper and we manually choose this value for each dataset. The experiment is run using \( k = 25 \) clusters for HIV, \( k = 50 \) for Citation and \( k = 100 \) for Bemol. For HIV and IASC, \( \ell \in \{25, 50, 100\} \) and \( R = 10 \) and for Ning we recompute exact eigenvectors after every 10 iterations. When using Citation and Bemol, eigenvector recomputations are performed every 20 iterations and \( \ell \in \{100, 200, 500\} \). With Nyström the number of columns is chosen from \( m \in \{1000, 1500\} \) for HIV and \( m \in \{2000, 5000\} \) for the other datasets. Finally, we apply the randomised SVD method with \( q = 2 \) and \( r \in \{1000, 1500\} \) for HIV and \( r \in \{2000, 5000\} \) for the remaining datasets. Note that in our implementation of \( k \)-means clustering, \( k \) represents an upper bound on the number of clusters found. To evaluate the learned clusters we use measures of modularity and \( k \)-way normalised cut. Let \( d_i = \sum_j W_{ij} \) and \( r = \sum_i d_i \), then the modularity is defined as

\[
Q = \frac{1}{2r} \sum_{i,j} \left( W_{ij} - \frac{d_i d_j}{2r} \right) \delta(c_i, c_j),
\]

where \( c \in \mathbb{R}^n \) is the cluster indicator vector and \( \delta \) is the Kronecker delta function. Intuitively modularity is the difference in the sum of edges weights within a cluster and the expected edge weights assuming the same weight distribution \( d \) for each vertex. The \( k \)-way normalised cut is

\[
N = \frac{1}{k} \sum_{\ell=1}^k \frac{\sum_{i,j} W_{ij} \delta(c_i, c_\ell)(1 - \delta(c_i, c_\ell))}{\sum_{i,j} W_{ij} \delta(c_i, c_\ell)}. \]

A cut between two clusters \( A \) and \( B \) is the sum of the weights between the clusters and the normalised cut is this sum divided by the sum of the weights of all edges incident to vertices in cluster \( A \). Hence the \( k \)-way normalised cut is the mean normalised cut between each cluster and its complementary vertices. To summarise, the greater the modularity, the better, and the lower the \( k \)-way normalised cut the better.

All experimental code is written in Python and we use an Intel Core i7-2600K at 3.40GHz with 16GB of RAM to conduct the simulations. The Laplacian matrices are stored in compressed sparse row representation and eigenvectors are found using Implicitly Restarted Lanczos Method in ARPACK [19] which computes only the required eigenvectors and not the full eigen-decomposition.
7.3.2 Results

Figure 4 shows the resulting modularities and $k$-way normalised cuts for all datasets however we begin by studying HIV. For both IASC and Ning, since eigenvectors are recomputed every 10 iterations, this can manifest itself as sudden changes in the modularities and $k$-way normalised cuts. These changes are more pronounced with Ning. We see a close correspondence of IASC and exact for both the modularity and normalised cut. As we observed with the toy datasets, a lower value of $\ell$ seems to improve results with the final graph having a cut of 0.09 with IASC $\ell = 100$ versus 0.10 for exact. Notice that IASC matches or improves results on exact, while keeping only 5% or fewer of the final number of eigenvectors. Ning fares badly in terms of the modularity of the resulting clustering with a value of 0.73 versus 0.82 for the exact approach at the final graph. However, with the cut measure Ning provides the best clustering, albeit with a more unstable curve than the other methods. Nyström does not provide a good approximation of the largest eigenvectors when the rank of the Laplacian exceeds the number of columns sampled. In contrast, RSVD is broadly competitive with IASC when using 1500 random projections. One of the reasons for the effectiveness of RSVD is Step 2 of Algorithm 2 which helps to ensure the column space of $Y$ is close to that generated using the largest eigenvectors of $A$, see [14] for further details.

A similar picture emerges with Citation and we see again that IASC is close to exact in terms of both measures. Note that it was too costly to compute clustering using Ning on this dataset and Bemol. We ran Ning on Citation for 337,920s before terminating the experiment: a computational time of at least 7.5 times more than the next most costly approach of RSVD, $r = 5000$, which took 44,880s. On Citation, when $\ell \in \{200, 500\}$ we obtain a close match to exact in general. The results are impressive when we consider the change in the graphs between eigenvector recomputations: the first graph is of size 555, and the 19th is 2855, an increase of 2300. With the 60th graph there are 10,063 vertices and 12,135 at the 79th. Looking at the Nyström curves, we again observe poor clustering performance even when $m = 5000$. Furthermore, RSVD $r = 5000$ can compete well with the exact method particularly when considering the cut measure although the modularity using RSVD suffers after approximately the 60th graph.

Finally consider the Bemol graphs in which it is difficult to find clear clusters, although they become more distinct over time. This is evident when looking at the exact curves for example: the modularity increases slightly from 0.27 to 0.32 whereas the $k$-way normalised cut falls from 0.74 to 0.65 from beginning to end. In contrast to the other datasets exact is improved upon by both IASC and Nyström (when $m = 5000$) respectively. One of the reasons that the Nyström and RSVD methods are effective on this data is because there are many edges and one can sample them out without affecting the clustering significantly. Note however that as we have seen in the other plots, Nyström is rather unstable compared to the other methods. Furthermore, the eigenvector updates every 20 iterations are noticeable in the cluster measures with IASC.

To conclude the analysis, Figure 5 shows the timings of the eigenvector computations of the clustering methods for the datasets. With HIV we can make a comparison with Ning and one can see that a cost of the method is a computation time which exceeds that of exact. IASC has a cumulative computation time
Figure 4: The performance of the clustering methods on the sequences of changing graphs.
Figure 5: The cumulative times taken by the eigenvector computations of the clustering methods.

of 62.0s, exact took 256.4s compared to 382.1s for Ning. With the Bemol and Citation datasets the Nyström approach costs the least in terms of computation when \( m = 2000 \) however exceeds the time taken for exact when \( m = 5000 \). In this case the time is dominated by the eigen-decomposition of a matrix in \( \mathbb{R}^{m \times m} \). Of note also is that RSVD with \( r = 5000 \) exceeds the time required for exact yet this was the number of random projections required for competitive performance. IASC improves over exact over the whole sequence of graphs as one does not recompute the eigenvectors at every iteration. Notice that the “staircase” effect in the IASC curves correspond the computation of the exact eigenvectors. Observe that on Bemol, IASC \( \ell = 200 \) takes 4,956 seconds in total for eigenvector computations versus 31,182 for exact, a speedup factor of 6.29 for a similar cluster quality. The equivalent improvement is 2.26 on Citation.

To emphasise the conditions in which IASC can be effective, we again cluster over the Citation data however graphs are recorded at 5 day intervals. The parameters are identical to those used above except we set \( R = 50 \). Figure 6 shows the resulting clustering qualities and eigenvector computation timings for IASC and RSVD. As one might expect we observe that RSVD is competitive to IASC, \( \ell = 100 \), when \( r = 5000 \), and only competitive with \( r = 2000 \) until approximately the 400th graph. The timings however show that IASC is faster
Figure 6: The $k$-way normalised cut and cumulative times taken by the eigenvector computations on Citation using 5 day intervals.

than RSVD for all values of $\ell$ over all graphs. Furthermore, when $\ell = 100$ IASC took 9640 seconds compared to 32,066 and 158,510 seconds with RSVD using $r$ values for 2000 and 5000 respectively.

8 Discussion

We have presented a novel incremental method for spectral graph clustering which updates the eigenvectors of the Laplacian in a computationally efficient way. Such an algorithm is useful for finding clusterings in time evolving graphs such as biological and social networks, and the Internet. A key part of the algorithm is a general way to approximate the first $k$ eigenvectors of a perturbed symmetric matrix given the eigenvectors of the original matrix. The resulting clustering algorithm, IASC, can be easily implemented using a standard linear algebra library.

We analysed IASC in both theoretical and empirical respects. Using perturbation theory, we showed when the canonical angles between the real and approximate subspaces generated by our update algorithm are close. Furthermore, IASC is examined empirically relative to the computation of exact eigenvectors for each graph, the method of Ning et al., randomised SVD, and the Nyström approach. On 2 toy and 3 real datasets we show that IASC can often match the cluster accuracy of the exact approach using a small fraction of the total number of eigenvectors and at a much reduced computational cost.

This work has opened up several perspectives for further study. The first is the analysis of the update of eigenvectors for a modularity matrix and other cluster quality criteria. As we have shown, the quality of the updates would depend on the spectrum of the matrices in question. Another interesting line of research is the issue of how to choose the number of clusters in the time evolving graphs.
References


