COMS 4232 Advanced Algorithms: Spectral Sparsification

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May 10th 2023

1 Introduction

Spectral sparsification of a graph is the procedure of removing edges from it such that the graph spectral properties are preserved. Although there are other notions of graph similarity, spectral metrics generalize beyond combinatorial ones thanks to the connections underlying Cheegar's inequality. In addition, spectral methods come packaged with optimized algorithms for common spectral tasks that can be applied to sparsification. To provide a mathematical definition of spectral sparsification, we consider the general case of the undirected and weighted graph G = (V, E, w) with n vertices and m edges such that V = [n] is the set of nodes, $E \subset \{(u, v) \mid u, v \in V\}$ is the edge set, and weight function $w: V \times V \to \mathbb{R}^+$. The Laplacian matrix of G can be thought of as the adjacency matrix subtracted from the degree matrix and is defined to be an $n \times n$ matrix L where

$$L_G(u, v) = \begin{cases} -w(u, v) & \text{if } u \sim v \\ deg(u) & \text{if } u = v \\ 0 & \text{otherwise} \end{cases}$$

where $deg(u) = \sum_{u \sim v} w(u, v)$. With this, it can be shown that the Laplacian Matrix may be defined by its quadratic form

$$Q_G(x) = x^T L_G x = \sum_{u \in v} w(u, v)(x_u - x_v)^2 \ge 0$$
 for any $x \in \mathbb{R}^n$

The quadratic form of a graph is connected to its spectral properties and its cut boundaries. Considering $x \in \mathbb{R}^n$, such that, ||x|| = 1, the maximum and minimum quadratic forms of a matrix are the largest and smallest eigenvalues of the matrix respectively. When components of x represent 1-0 membership of a set $S \subset V$ ($x_i = \mathbb{1}[i \in S]$), then, $Q_G(x) = 2|\partial S|$ where ∂S represents the set of edges that connect a vertex in S with a vertex in S. Note the conductance of a cut S of S is,

$$\Phi_G(S) \triangleq \frac{|\partial S|}{\min\{\operatorname{Vol}(S), \operatorname{Vol}(V - S)\}}$$

where $\operatorname{Vol}(S) = \sum_{i \in S} \operatorname{deg}(i)$ Since the conductance of a cut can be understood as the ratio of the number of edges that cross the boundary of a cut of V to the number of edges contained in a cut of V, we can also define the conductance of a graph G by

$$\Phi_G \triangleq \min_{\emptyset \neq S \subset V} \Phi_G(S).$$

Even without a proof, it can be seen that there is a connection between the number of edges connecting a min cut and the eigenvalues of the Laplacian. Consider the normalized Laplacian

$$\hat{L}_G \triangleq D^{-1/2} L_G D^{1/2}$$

where D is the diagonal matrix representing the degree of each vertex. It turns out that the first non-zero eigenvalue of the normalized Laplacian represents the conductance of a graph such if the magnitude of this eigenvalue is zero, it implies the graph contains two disconnected components and if the magnitude of this eigenvalue is one, it implies the conductance of any cut is high. This is formalized in Cheegar's inequality which states that

$$2/\Phi_G \le \mu_2 \le 2\Phi_G$$

where μ_2 is the second smallest eigenvalue of \hat{L}_G . With this, we can introduce an intuitive definition of spectral similarity by defining a notion of similarity that preserves the eigenvalues of the normalized Laplacian of a graph G and its sparsifier \tilde{G} . With an understanding of the relationship between the quadratic form and the spectral properties of a matrix, it is natural to define that \tilde{G} is a σ -spectral approximation of G if for all $x \in \mathbb{R}^n$

$$\frac{1}{\sigma} x^T L_{\widetilde{G}} x \le x^T L_{G} x \le \sigma x^T L_{\widetilde{G}} x$$

We may condense this definition by saying the sub-graph \widetilde{G} is a σ -spectral approximation of G if

$$\frac{1}{\sigma}L_{\widetilde{G}} \preceq L_G \preceq \sigma L_{\widetilde{G}},$$

where the notation, $A \leq B$, implies that for $A, B \in \mathbb{R}^{n \times n}$, for all $x \in \mathbb{R}^n$, $x^T A x \leq x^T B x$. To further understand this definition, consider a graph that has two connected components that are very dense, but there are only a few edges connecting the two components (shown in fig.1). Intuitively, removing

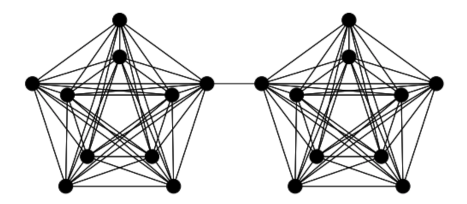


Figure 1: Two highly connected graphs connected by a single edge taken from [ST11].

an edge from the boundary of the two equal-volume connected components will have a greater impact on similarity than removing an edge from one of the connected components. Through the relationship between conductance and spectrum defined above, we can see that removing certain boundary edges such as the edge connecting two dense components will also impact the spectral properties of the sparsifier. This example not only helps to explain the definition of spectral sparsification, but also suggests a method of sparsifying graphs by removing edges that approximately maintain the conductance of the original graph.

2 Preliminaries

Throughout this paper we will assume the graph construction described in the in Introduction. For any matrix, let $\lambda_{\max}(A)$ and $\lambda_{\min}(A)$ be the maximum and minimum eigenvalues of A. The condition number of matrix A is defined by $\lambda_{\max}(A)/\lambda_{\min}(A)$. For any two matrices A and B, we write $A \leq B$ to represent that B - A is positive semidefinite and $A \prec B$ to represent that B - A is positive definite. For any two matrices A and B of equal dimensions, let $A \cdot B = tr(A^T B)$. For any function f, we write $\tilde{O}(f) \triangleq O(f \cdot log^{O(1)}f)$. For matrices A and B, we write $A \approx_{\epsilon} B$ if $(1 - \epsilon) \cdot A \leq B \leq (1 + \epsilon)A$.

3 Nearly Linear Time Sparsifiers

The first spectral sparsification procedure, taken from Spielman and Teng [ST11], introduces a simple extension of the ideas presented in the introduction. First, partition a graph into subgraphs of high conductance. Second, randomly sample edges to remove from these graphs from the high conductance subgraphs but preserve the boundaries. Lastly, if there are too many edges remaining in the boundary, consider a new graph where each subgraph is a node connected to other subgraph nodes with boundary edges from the original graph. Then recursively run the algorithm on this new graph to remove edges from the boundary. As this rather simple idea leads to a complex implementation that approximates the NP-hard exact algorithm, we will focus on the case of sparsifying an unweighted graph and we will omit many details from the proof of correctness. According to the authors, their main result is showing the existence of good spectral sparsifiers with a nearly linear number of edges and that their discovery is feasible in better than polynomial time.

3.1 Notation and Tools

For graphs G and H, we will use the notation G+H to refer to the graph with Laplacian L_G+L_H . For graphs G and H, it is immediate that $G+H \preceq \widetilde{G}+\widetilde{H}$. Note that when a subgraph is constructed, its node numeric identities will be preserved allowing the sum of subgraphs to yield results about the original graph. We define the boundary of a vertex set S with respect to another vertex set S to be the set of edges from S into S denoted S denoted S or S. We define the conductance of a set S in the subgraph induced by $S \subseteq V$ to be,

$$\Phi_B^G(S) \triangleq \frac{|\partial_B(S)|}{\min\{\operatorname{Vol}(S), \operatorname{Vol}(B-S)\}}$$

and we define

$$\Phi_B^G \triangleq \min_{S \subset B} \Phi_B^G(S).$$

We also define that $\Phi_B^G(\emptyset) = 1$ and for $|B| = 1, \Phi_B^G = 1$. We will assume that if a graph has high conductance, then it may be sparsified by a random sampling procedure such that $E[\widetilde{A}] = A$ where \widetilde{A}

and A are the adjacency matrices of the sparsified graph and the original graph respectively. While we include the specific sampling algorithm here we do not analyze its specifics as there are a variety that can be chosen. This algorithm in particular is tuned such that there is zero probability of removing all edges from a node. The internal parameter γ controls the number of edges expected in the samples graph. Note that if $\gamma = n \triangleq |V|$, then the sampled graph will be exactly the same as the original. The algorithm below is taken directly from [ST11]. Theorem 1 ensures that this procedure applied to graph of high

Algorithm 2 Algorithm to randomly sample from graph of high conductance

```
1: \widetilde{G} = \text{Sample}(G, \epsilon, p, \lambda)
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2: Set
$$k = \max\{\log_2(3/p), \log_2(n)\}$$

3: Set $\gamma = \left(\frac{12k}{\epsilon\lambda}\right)^2$

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4: for every edge (i, j) in G do

Set the probability $p_{i,j} = \min\{1, \frac{\gamma}{\min\{d_i, d_i\}}\}$

Place an edge of weight $1/p_{i,j}$ between vertices (i,j) into \widetilde{G} 6:

7: end for

conductance ensures a good approximation.

Theorem 1. Let $\epsilon, p \in (0, 1/2)$ and let G = (V, E) be an unweighted graph whose smallest non-zero normalized Laplacian eigenvalue is at least λ . Let $S \subset G$ and let F be the edges in G(S). Suppose $(S,\widetilde{F}) = \text{Sample}((S,F),\epsilon,p,\lambda)$ and let $\widetilde{G} = (V,\widetilde{F} \cup (E-F))$. Then with probability at least 1-p,

- 1. \widetilde{G} is a $(1+\epsilon)$ -approximation of G, and
- 2. The number of edges in \widetilde{F} is at most

$$\frac{288 \max\{\log_2(3/p), \log_2(n)\}^2}{(\epsilon \lambda)^2} |S|$$

Lets define a decomposition of G into vertex sets $(A_1, ..., A_k)$ for some k as a ϕ -decomposition if $\Phi_{A_i}^G \geq \phi$ for all i. We define the boundary of a decomposition as the set of edges between different vertex sets in the partition denoted,

$$\partial(A_1,...,A_k) \triangleq E \cap \bigcup_{i \neq j} (A_i \times A_j).$$

Considering the conductance of a subgraph, the boundary of a decomposition gives us a decomposition where each subgraph has conductance at least ϕ . We also define a decomposition as a λ -spectral decomposition if the smallest non zero normalized Laplacian eigenvalue of $G(A_i)$ is at least λ , for all i. Using Cheegar's inequality we have that every ϕ -decomposition is a $(\phi^2/2)$ -spectral decomposition.

3.2 Existence of Sparsifiers

In this section, we will show the existence of sparsifiers for the procedure for generating sparse graphs with our desired properties. Theorem 3 is central to the existence of sparsifiers and in its proof there will be ideas on how to construct sparsifiers. The proof of Theorem 3 follows quickly from lemma 2 which makes a statement connecting the size of a sparse cut to its conductance.

Lemma 2. Let G = (V, E) be a graph and let $\phi \leq 1$. Let $B \subseteq V$ and let $S \subset B$ be a set maximizing Vol(S) among those satisfying:

- 1. $Vol(S) \leq Vol(B)/2$, and
- 2. $\Phi_B^G(S) \leq \phi$

If $Vol(S) = \alpha Vol(B)$ for $\alpha \leq 1/3$, then

$$\Phi_{B-S}^G \ge \phi \left(\frac{1 - 3\alpha}{1 - \alpha} \right)$$

Proof. Consider B a vertex set of some graph G. Let S maximize $\operatorname{Vol}(S)$ and satisfy condition 1 and 2 of Lemma 2. S is the largest volume vertex set in B of bounded conductance. Let $\beta = \frac{1-3\alpha}{1-\alpha}$. Suppose contrarily that $\Phi_{B-S}^G < \phi\beta$. Suppose R is the set that minimizes Φ_{B-S}^G . Then $\Phi_{B-S}^G(R) < \phi$ and $\operatorname{Vol}(R) \leq \operatorname{Vol}(B-S)/2$. Let $T = R \cup S$. Note that

$$|E(T, B - T)| \le |E(S, B - S)| + |E(R, B - S - R)| \tag{1}$$

$$<\phi Vol(S) + \phi\beta Vol(R)$$
 (2)

Consider 2 cases

1. Suppose $\operatorname{Vol}(T) \leq \operatorname{Vol}(B)/2$. This implies that $\operatorname{Vol}(S) \leq \operatorname{Vol}(B)/2$ because of T has strictly more connections than S. Thus

$$|E(T, B - T)| < \phi(\operatorname{Vol}(S) + \beta \operatorname{Vol}(R))$$

$$< \phi(\operatorname{Vol}(S) + \operatorname{Vol}(R)) = \phi \operatorname{Vol}(T)$$

$$\Rightarrow \frac{|E(T, B - T)|}{\operatorname{Vol}(T)} = \Phi_B^G(T) < \phi$$

However, this is a contradiction because we stated that S was the largest set that satisfied these properties and T is strictly larger than S.

2. Suppose that Vol(T) > Vol(B)/2. Note that

$$Vol(T) = Vol(S) + Vol(R)$$

$$\leq Vol(S) + (1/2)(Vol(B) - Vol(S))$$

$$= (1/2)Vol(B) + (1/2)Vol(S)$$

$$= \frac{1+\alpha}{2}Vol(B)$$

because $\operatorname{Vol}(S) = \alpha \operatorname{Vol}(B)$. From the definition of conductance, $\operatorname{Vol}(S) \leq (1/2)\operatorname{Vol}(B)$ and that $\operatorname{Vol}(R) \leq (1/2)(\operatorname{Vol}(B-S))$ This implies that

$$Vol(B - T) = Vol(B) - Vol(T)$$

 $\geq Vol(B)(1 - \alpha)/2$

Now consider the conductance of T,

$$\begin{split} |E(T,B-T)| &< \phi \mathrm{Vol}(S) + \phi \beta \mathrm{Vol}(R) \text{ by line } 2 \\ &\leq \phi \mathrm{Vol}(S) + \phi \beta (\mathrm{Vol}(B) - \mathrm{Vol}(S))/2 \\ &= \phi \alpha \mathrm{Vol}(B) + \phi \beta (\mathrm{Vol}(B) - \alpha \mathrm{Vol}(B))/2 \\ &= \phi \mathrm{Vol}(B)(\alpha + \beta (1-\alpha)/2) \end{split}$$

Thus, we have an upper bound on the conductance of T within the subgraph induced by B

$$\Phi_B^G(T) = \frac{|E(T, B - T)|}{\min\{\text{Vol}(T), \text{Vol}(B - T)\}}$$

$$= \frac{|E(T, B - T)|}{\text{Vol}(B - T)} \text{ by case 2 assumption}$$

$$\leq \frac{\phi \text{Vol}(B)(\alpha + \beta(1 - \alpha)/2)}{\text{Vol}(B)(1 - \alpha)/2}$$

$$= \phi$$

T contradicts the constructive assumption of the conditions of S.

Therefore, Lemma 2 holds.

Theorem 3. Let G = (V, E) be a graph and let m = |E|. Then, G has a $(6 \log_{4/3} 2m)^{-1}$ -decomposition with $|\partial(A_1, ..., A_k)| \le |E|/2$

Proof. To show Theorem 3, we will show that Algorithm 3 yields the decomposition that is claimed to exist. First we see that this procedure must terminate because the recursive call is always passed a strictly smaller set than the calling function was passed and $\Phi_B^G = 1$ when |B| = 1 and $\phi < 1$. Let $(A_1, ..., A_k)$ be the output of idealDecomp(V). Note that set A_i can be returned when $\Phi_{A_i}^G \ge \phi$ or when $\Phi_{A_i}^G \ge \phi^{1-3\alpha}$. So in either case $\Phi_{A_i}^G \ge \phi/3$ for each i. Now consider the depth of the recursion. In the worst case the $\operatorname{Vol}(S) \le \operatorname{Vol}(S)/4$ at every iteration because that maximizes the size of the set passed to idealDecomp. Therefore, at each iteration the volume of the set passed recursively will be at least 3/4 times the volume of set passed to the caller. Therefore, because the volume of a set cannot be less than 1 we have that $1 \le \operatorname{Vol}(V)(3/4)^d$ where d is the worst case depth of the recursion. Therefore,

$$d \le \log_{4/3} \operatorname{Vol}(V)$$
.

Now supplement this finding with the fact that every time a set is split into S and B-S we add edges to $\partial(A_1,...,A_k)$. However, since S satisfies Lemma 2, we know that $\Phi_B^G(S) \leq \phi$. Therefore, every time we split a set we are adding at most $|E|\phi$ edges to $\partial(A_1,...,A_k)$ because |E| is always larger than $\min\{\operatorname{Vol}(S),\operatorname{Vol}(S)\}$. Therefore,

$$|\partial(A_1,...,A_k)| \le |E|\phi \log_{4/3} \operatorname{Vol}(V) \le |E|/2$$

Since $\operatorname{Vol}(V) = 2m$, we have that $\phi/3 \le \left(2\log_{4/3} 2m\right)^{-1}$ implies

$$\phi \le \left(6\log_{4/3} 2m\right)^{-1}$$

Algorithm 3 Algorithm to partition graph with desired properties

```
1: Set \phi = \left(2\log_{4/3} \operatorname{Vol}(V)\right)^{-1}
```

- 2: Note, the algorithm is called initially with B = V
- 3: idealDecomp (B, ϕ)
- 4: If $\Phi_B^G \ge \phi$ then return B. Otherwise proceed.
- 5: Let S be the subset of B maximizing Vol(B) satisfying the conditions of lemma 2

- 6: If $Vol(S) \leq Vol(B)/4$, return the decomposition $(B S, idealDecomp(S, \phi))$
- 7: else, return the decomposition (idealDecomp $(B S, \phi)$, idealDecomp (S, ϕ))

Since edges are bounded by n^2 for n = |V|, then Theorem 3 shows the decomposition with minimum conductance of $(6 \log_{4/3} 2m)^{-1}$. Since the conductance is $\Omega(1/\log(n))$, then by Cheegar's inequality, the second lowest non-zero eigenvalue is $\Omega(1/\log^2(n))$. Therefore, we can apply Theorem 1 to each set of the partition to ensure that the number of edges in these partitions are $\widetilde{O}(n)$. All that remains is to ensure that the edges in the boundary of the decomposition are also $\widetilde{O}(n)$. This can be accomplished by collapsing all the nodes in each partition into a single node of a new graph and recursively partitioning this new graph with Theorem 3. Everytime we use Theorem 3, the number of edges in the resulting boundary is divided by 2. Therefore, we will re-curse to a depth logarithmic in |E|. The crux of this proof is the procedure of partitioning a graph into connected components of "high" conductance. Due to Lemma 2, we have a set of conditions that will guarantee that the complement of the set that satisfies them has high enough conductance. Those necessary conditions are numbered 1 and 2 in Lemma 3 as well as the size limit on the set produced. Our idealized decomposition algorithm extracts a set of high conductance if all the conditions are met and re-runs the algorithm on the rest. If the size condition is not met, it simply re-runs the algorithm on both the set and its complement.

3.3 Approximation

Unfortunately, the previous section skips over one important detail. It is not specified exactly how we can find a set that satisfies the conditions of Lemma 2. If this were clear we would have a perfectly functional algorithm for spectral sparsification. Line 5 of Algorithm 3 is where this breaks down. Finding such a set is an NP-hard problem, and therefore must be approximated. We will not go into the details, but there exists an algorithm that can generate these sets approximately in linear time:

$$D = \operatorname{ApproxCut}(G, \phi, p),$$

where G is the graph to find the cut within, $\phi \in (0,1)$ is the conductance, and $p \in (0,1)$ is the probability of failiure where success is defined as the output set $D \subset V$ as meeting conditions.

- 1. $Vol(D) \ge (1/29)Vol(V)$ or
- 2. there exists a set $W \supset V D$ for which $\Phi_W^G \ge f_2(\phi)$ where $f_2(\phi) \triangleq \frac{c_2\phi^2}{\log^4 m}$ for some constant c_2

Note that condition 2 is essentially the same thing as saying that instead of the complement of D having high conductance, there exists a subset of the complement of D that has high conductance. Also D is guaranteed to satisfy,

```
1. Vol(D) \le (23/25)Vol(V) and
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2. for
$$D \neq \emptyset$$
, $\Phi_G(D) \leq \phi$

The expected runtime of ApproxCut is $O(\phi^{-4}m\log^9 m\log(1/p))$. Now using this approximation of line 5 of Algorithm 3, we introduce Algorithm 4 which sparsifies an unweighted graph using approximate sparsest cuts and our random sampling algorithm from the first section. In simple english, this algo-

Algorithm 4 Approximate algorithm to sparsify an unweighted graph

```
1: G = \text{UnwtedSparsify}(G, \epsilon, p)
               If Vol(V) \leq c_3 \epsilon^{-2} n \log^3 0(n/p), return G
               Set \phi = (2\log_{29/28} \text{Vol}(V))^{-1}, \hat{p} = p/6n\log_2 n, and \hat{\epsilon} = \frac{\epsilon(\ln 2)^2}{(1+2\log_{29/28} n)(2\log n)}
 3:
               Set (\widetilde{G}_1, ..., \widetilde{G}_k) = PartitionAndSample(G, \phi, \hat{\epsilon}, \hat{p}).
 4:
               Let V_1, ..., V_k be the vertex sets of \widetilde{G}_1, ..., \widetilde{G}_k respectively, and let G_0 be the graph with vertex
      set V and edge set \partial(V_1, ..., V_k).
               Set \widetilde{G} = \sum_{i=0}^{k} \widetilde{G}_i
     (\widetilde{G}_1, ..., \widetilde{G}_k) = \text{PartitionAndSample}(G = (V, E), \phi, \hat{\epsilon}, \hat{p})
               Set \lambda = f_2(\phi)^2/2
 8:
 9:
               Set D = \operatorname{ApproxCut}(G, \phi, \hat{p})
               If D = \emptyset return G_1 = \text{Sample}(G, \hat{\epsilon}, \hat{p}, \lambda)
10:
               Else, if Vol(V) \leq (1/29)Vol(V)
11:
                        Set \widetilde{G}_1 = \text{Sample}(G(V - D), \hat{\epsilon}, \hat{p}, \lambda)
12:
                        Return (\widetilde{G}_1, \operatorname{PartitionAndSample}(G(D), \phi, \hat{\epsilon}, \hat{p}))
13:
               Else
14:
                        Set \widetilde{H}_1, ..., \widetilde{H}_k = \text{PartitionAndSample}(G(V - D), \phi, \hat{\epsilon}, \hat{p})
Set \widetilde{I}_1, ..., \widetilde{I}_j = \text{PartitionAndSample}(G(D), \phi, \hat{\epsilon}, \hat{p})
15:
16:
                        Return (\widetilde{H}_1,...,\widetilde{H}_k,\widetilde{I}_1,...,\widetilde{I}_i)
17:
```

rithm uses a subroutine, PartitionAndSample that uses ApproxCut to find high conductance vertex sets. Whenever it does find one, it instantly calls Sample on it. When ApproxCut does not directly yield a high conductance cut, (when the set it returns is large) PartitionAndSample is recursively called on the cut and its complement. The caller of PartitionAndSample, UnwtedSparsify, sparsifies a given partition and then acts recursively on the remaining edges in boundary set of the partition. The specifics of this algorithm are included more for completeness than understanding and we do not analyze it fully. In addition, the constants in Algorithm 4 make it entirely impractical for reasonable sized graphs. However, it does produce a $(1 + \epsilon)$ -approximation of G with at most $c_4 \epsilon^{-2} n \log^{31}(n/p)$ edges, for some constant c_4 with probability p. It achieves an expected runtime of $O(m \log(1/p) \log^{15} n)$

4 Graph Sparsification by Effective Resistances

4.1 Introduction

In 2008, Spielman and Srivastava [SS08] introduce a nearly-linear time algorithm which produces sparsifiers with $O(nlogn/\epsilon^2)$ edges by using random sampling over the edges of G. For an edge $e \in E$, they introduce a notion of resistance which is equal to $1/w_e$, and a notion of effective resistance R_e which is the potential difference induced across it when a unit current is injected at one end and extracted at the other. It is known to be proportional to the commute time of an edge, which is the expected time it takes for a natural random walk starting at one end to travel to the other end and back.

4.2 Main algorithm

We denote q to be the sample size.

Algorithm 6 Effective Resistances Sparsifier Algorithm

```
1: Initially, set \tilde{G} = (V, \tilde{E} = \emptyset, \tilde{w})

2: for i in range(q) do

3: Pick a random edge e from G with probability p_e proportional to w_eR_e

4: if e is not already in \tilde{E} then

5: Add e to \tilde{E} with weight \tilde{w}_e = w_e/qp_e

6: else

7: Set \tilde{w}_e = \tilde{w}_e + w_e/qp_e

8: end if

9: end for
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If $q = 4C^2 n log n/\epsilon^2$, where C is a constant from a concentration theorem by Rudelson & Vershynin [RV06], then \tilde{G} is a sparsifier with probability at least 1/2. This is proven using Markov inequality on

$$\tilde{L} = B^T W^{1/2} S W^{1/2} B$$

where $S, W \in \mathbb{R}^{m \times m}$ are such that $\forall e \in E, S(e, e) = \frac{\tilde{w}_e}{w_e}, W(e, e) = w_e$, and B is the signed edge-vertex incidence matrix of G defined as

$$B(e, v) = \begin{cases} 1 & \text{if } v \text{ is } e \text{'s head} \\ -1 & \text{if } v \text{ is } e \text{'s tail} \\ 0 & \text{otherwise} \end{cases}$$

after orienting the edges arbitrarily if G is undirected. Once the effective resistances have been computed, the main algorithm runs in time $O(nlogn/\epsilon^2)$.

4.3 Computing the effective resistances

The key part of this algorithm is approximating the effective resistances $\{R_e\}_{e\in E}$ needed for sampling in nearly-linear time. Since we are comparing the effective resistances in the main algorithm, it is enough to approximate them all up to the same common factor. The authors use the *Moore-Penrose Pseudoinverse* L_G^+ of L_G defined as

$$L_G^+ = \sum_{i=1}^{n-1} \frac{1}{\lambda_i} u_i u_i^T$$

where $\lambda_1, ..., \lambda_{n-1}$ are the non-zero eigenvalues of L_G and $u_1, ..., u_{n-1}$ a corresponding set of orthonormal eigenvectors, to build the set

$$\{W^{1/2}BL_G^+\chi_v\}_{v\in V}$$

where χ_u is the elementary vector with coordinate 1 in position u. They prove that for $u, v \in V$, the effective resistance R_e of the edge e between u and v is the distance between $W^{1/2}BL_G^+\chi_u$ and $W^{1/2}BL_G^+\chi_v$. Using the Johnson-Linderstrauss Lemma [JL84], these vectors are projected onto $\mathbb{R}^{O(\log n)}$ to accelerate the distance computation. The resulting set is

$$\{QW^{1/2}BL_G^+\chi_v\}_{v\in V}$$

where $Q \in \mathbb{R}^{k \times n}$ is a random $\pm 1/\sqrt{k}$ matrix, and k is given by the Achlioptas Lemma [Ach01] in order to maintain ϵ -approximate distances.

Algorithm 8 Computing the Effective Resistances

- 1: Compute such a random $Q \in \mathbb{R}^{k \times n}$ for $k = 24 log n/\epsilon^2$
- 2: Compute $Y = QW^{1/2}B$
- 3: for i in range(k) do
- 4: Compute z_i where $z_iL = y_i$ where y_i is the *i*-th row of Y by calling the Spielman-Teng solver [ST08]
- 5: end for
- 6: Return the matrix Z whose i-th row is z_i

The matrix Z is equal to $QW^{1/2}BL_G^+$, and its columns can be subtracted 2 by 2 to find distances between the vectors of the set and the effective resistances. Computing Y takes time $2m \times 24 log n/\epsilon^2 + m = \tilde{O}(m/\epsilon^2)$ since B has 2m entries and $W^{1/2}$ is diagonal. Each call to the Spielman-Teng solver takes $\tilde{O}(m)$ time, leading to the construction of Z in time $\tilde{O}(m/\epsilon^2)$. Thus, for each pair $u, v \in V$, we can compute the effective resistance R_e of the edge between u and v in time O(log n).

5 Twice-Ramanujan Sparsifiers

5.1 Introduction

Ramanujan graphs are connected d-regular graphs H, for which

$$\forall \lambda \in Sp(L_H), \lambda \neq 0 \implies \lambda \in [d - 2\sqrt{d-1}, d + 2\sqrt{d-1}]$$

Building a Ramanujan graph on n vertices and multiplying the weight of every edge by $n/(d-2\sqrt{d-1})$ give us a graph which κ -approximates the complete graph for

$$\kappa = \frac{d + 2\sqrt{d - 1}}{d - 2\sqrt{d - 1}}$$

5.2 The Main Result

In their paper [BSS14] from 2009, Batson, Spielman and Srivastava show that every graph can be approximated at least this well for d > 1 by a graph with only twice as many edges as a Ramanujan graph. In other words, they show this very theorem:

Theorem 4. For every d > 1, every undirected weighted graph G = (V, E, w) on n vertices contains a

weighted subgraph $H = (V, F, \tilde{w})$ with $\lceil d(n-1) \rceil$ edges (i.e., average degree at most 2d) that satisfies:

$$x^T L_G x \le x^T L_H x \le \left(\frac{d+1+2\sqrt{d}}{d+1-2\sqrt{d}}\right) \cdot x^T L_G x \quad \forall x \in \mathbb{R}^V$$

The main result which leads to Theorem 4 is the following:

Theorem 5. Suppose d > 1, and $v_1, v_2, ..., v_m$ are vectors in \mathbb{R}^n with

$$\sum_{i \le m} v_i v_i^T = i \boldsymbol{d}_{R^n}$$

Then there exists scalars $s_i \geq 0$ with $|\{i: s_i \neq 0\}| \leq dn$ so that

$$I_n \preceq \sum_{i \leq m} s_i v_i v_i^T \preceq \left(\frac{d+1+2\sqrt{d}}{d+1-2\sqrt{d}} \right) \cdot id_{R^n}$$

We define

$$V_{n \times m} = (L_G^+)^{1/2} B^T W^{1/2}$$

where B and L_G^+ are respectively the signed edge-vertex incidence and the Moore-Penrose Pseudoinverse matrices defined in the previous section and $W \in \mathbb{R}^{m \times m}$ is the diagonal matrix where $\forall e \in E, W(e, e) = w_e$. Because $L_G L_G^+ = L_G^+ L_G = \mathbf{id}_{im(L_G)}, V_{n \times m}$ yields columns $\{v_i\}_{i \in [m]}$ such that

$$\begin{split} \sum_{i \leq m} v_i v_i^T &= V V^T \\ &= (L_G^+)^{1/2} B^T W B (L_G^+)^{1/2} \\ &= (L_G^+)^{1/2} L_G (L_G^+)^{1/2} \\ &= \mathbf{id}_{im(L_G)} \end{split}$$

Using Theorem 5, we obtain the diagonal matrix $S_{m\times m}$ such that $\forall i\in[m],\ S(i,i)=s_i$ and set

$$L_{\tilde{G}} = B^T W^{1/2} S W^{1/2} B$$

which is the Laplacian of the subgraph \tilde{G} of G with weights $\{\tilde{w}_i = w_i s_i\}_{i \in E}$; \tilde{G} has at most d(n-1) edges since at most that many of the s_i are nonzero. Using the Courant-Fischer Theorem, we can prove

$$1 \le \frac{x^T L_{\tilde{G}} x}{x^T L_{G} x} \le \left(\frac{d+1+2\sqrt{d}}{d+1-2\sqrt{d}}\right) \quad \forall x \perp \mathbf{1}$$

as desired. The correct set of weights $\{s_i \in R^+ \mid i \in [m]\}$ is picked using an algorithm described below, which gives a constructive proof to Theorem 5 and thus concludes the proof to Theorem 4.

5.3 The Algorithm

The first observation to be made is that for $A \in \mathbb{R}^{n \times n}$ and $v \in \mathbb{R}^n$, the eigenvalues of $A + vv^T$ interlace those of A. The eigenvalues of A can be seen as charged particles on a slope, resting on chargeless barriers because of gravity. Adding vv^T to A corresponds to adding a charge $\langle v, u_j \rangle^2$ on the barrier corresponding to the j-th eigenvalue, thus repelling the initial eigenvalue forward on the slope. The idea is that by adding sufficiently enough vv^T for v picked at random, all of the eigenvalues move up at the same pace and the condition number $\frac{\lambda_{max}(A)}{\lambda_{min}(A)}$ is bounded.

The algorithm of Theorem 4 takes its inspiration from this model and builds $A^{(q)}$ from $A^{(q-1)}$ iteratively by picking $i \in [m]$, $s_i > 0$ and adding $s_i v_i v_i^T$ while keeping all of the eigenvalues between two barriers. To that end, the authors introduce two functions that describe how far the eigenvalues of A are from the lower and upper barriers $l, u \in \mathbb{R}$:

$$\Phi^{u}(A) = Tr(uI - A)^{-1}$$
 (Upper potential)
$$\Phi_{l}(A) = Tr(A - lI)^{-1}$$
 (Lower potential)

It is very important that the interval between $\lambda_{min}(A)$ and $\lambda_{max}(A)$ does not increase as we iterate, and for this reason the Both Barriers Lemma is introduced. It proves that for a given A and values $\delta_L > 0$, $\delta_U > 0$, there always exists an $i \in [m]$ and $s_i > 0$ such that we can shift the lower and upper barriers respectively by δ_L and δ_U forward and maintain the upper and lower potentials, i.e.

$$\Phi^{u+\delta_U}(A + s_i v_i v_i^T) \le \Phi^u(A)$$

$$\Phi_{l+\delta_L}(A + s_i v_i v_i^T) \le \Phi_l(A)$$

The index i and value s_i are picked using special matrices referred to as $U_{A^{(q)}}$ and $L_{A^{(q)}}$; these matrices result from a computation involving A, the potentials and the barriers.

Algorithm 10 Twice-Ramanujan Sparsifiers Algorithm

- 1: Compute the columns $\{v_i\}_{i\in[m]}$ of $V_{n\times m}=(L_G^+)^{1/2}B^TW^{1/2}$
- 2: Initially, set $A^{(0)} = 0$, the barriers $u = u_0$ and $l = l_0$, and the potentials

$$\Phi_0^u(A^{(0)}) = \epsilon_U$$
 and $\Phi_{l_0}(A^{(0)}) = \epsilon_L$

- 3: **for** q in range(dn) **do**
- 4: Set $u = u + \delta_U$ and $l = l + \delta_L$
- 5: Set $A^{(q+1)} = A^{(q)} + s_i v_i v_i^T$, where v_i and s_i are picked so that the potentials don't increase.
- 6: end for

After Q = dn steps, we have

$$\frac{\lambda_{max}(A^{(Q)})}{\lambda_{min}(A^{(Q)})} \le \frac{u_0 + dn\delta_u}{l_0 + dn\delta_L}$$

By choosing the right values $u_0, l_0, \delta_U, \delta_L, \epsilon_U$, and ϵ_L , we get

$$\frac{\lambda_{max}(A^{(Q)})}{\lambda_{min}(A^{(Q)})} \le \frac{d+1+2\sqrt{d}}{d+1-2\sqrt{d}}$$

as desired by Theorem 5, which leads to the sparsifier G.

Computing the vectors v_i is done in time $O(n^2m)$. For each iteration of the algorithm, computing the matrices $U_{A(q)}$ and $L_{A(q)}$ needed to pick the next v_i to add can be performed in time $O(n^3)$. Finally, we need to compute $U_{A^{(q)}}(v_i)$ and $L_{A^{(q)}}(v_i)$ for each edge v_i , which can be done in time $O(n^2m)$. As we run for dn iterations, the total time for the algorithm is $O(dn^3m)$.

6 Fast Construction of Linear-size Spectral Sparsifers

Constructing a Randomize Algorithm for Fast Spectral Sparsification

It is known that constructing a spectral sparsifier for graphs is a special case of sparsifying the sum of rank-1 PSD matrices. With this, [LS18] goal was to create a near-linear time algorithm such that for any vectors $v_1, ... v_m$ with $\sum_{i=1}^m v_i v_i^T = I$, it finds scalars $\{s_i\}_{i=1}^m$ satisfying

$$|\{s_i: s_i \neq 0\}| = O\left(\frac{qn}{\epsilon^2}\right)$$

such that

$$(1 - \epsilon) \cdot I \leq \sum_{i=1}^{m} s_i v_i v_i^T \leq (1 + \epsilon) \cdot I$$

For inspiration, [LS18] builds upon [BSS14] by improving on a randomized version of their algorithm, which is shown below. As described previously, [BSS14] uses two barrier values u_j and l_j and defines two

Algorithm 11 Randomized [BSS14] Algorithm for spectral sparsification

- 1: j = 0
- 2: $l_0 = -(2n)/\epsilon$, $u_0 = (2n)/\epsilon$, $A_0 = 0$
- 3: while $u_j l_j < 8n/\epsilon$ do
- Let $t = tr(u_i I A_i)^{-1} + tr(A_i = l_i I)^{-1}$
- Sample a vector v_i with probability 5:

$$p_i = \left(v_i^T (u_j I - A_j)^{-1} + v_i^T (A_j - l_j I)^{-1} v_i\right) / t$$

- $A_{j+1} = A_j + \frac{\epsilon}{t} \cdot \frac{1}{p_i} \cdot v_i v_i^T$ $u_{j+1} = u_j + \frac{\epsilon}{t \cdot (1-\epsilon)}, l_{j+1} = l_j + \frac{\epsilon}{t \cdot (1+\epsilon)}$
- 8:
- 9: end while
- 10: return A_j

potential functions $\Phi^u(A) \triangleq tr(uI - A)^{-1}$ and $\Phi_l(A) \triangleq tr(A - lI)^{-1}$ to determine if some eigenvalue of A is close to the barriers u and l. During an iteration j, their Algorithm always find a vector $\{v_i\}_{i=1}^m$ and updates u_j, l_j such that the invariant $l_j I \prec A_j \prec u_j I$ holds in each iteration. After $T = \Theta(n/\epsilon^2)$

iterations, they would find a matrix $A_T \approx_{O(\epsilon)} I$. However, areas of improvement can be made to the randomized [BSS14] algorithm.

- 1. We need a fast algorithm to approximate the probabilities $\{p_i\}_{i=1}^m$ used to choose vectors
- 2. We need to use the computed $\{p_i\}_{i=1}^m$ to choose multiple vectors in each phase to reduce the total number of phases

With this, [LS18] improves on both points for their algorithm by adopting the potential function

$$\Phi_{u,l}(A) \triangleq tr(uI - A)^{-q} + tr(A - lI)^{-q}$$

to maintain that the eigenvalues of A do not get close to the barriers u and l in order to approximate the probabilities $\{p_i\}_{i=1}^m$ efficiently (since constant q is large) and by showing that the potential function still works as before as long as the sampling probability satisfies

$$p_i \ge C \cdot \frac{v_i^T (uI - A)^{-1} v_i + v_i^T (A - lI)^{-1} v_i}{\sum_{j=1}^m \left(v_j^T (uI - A)^{-1} v_j + v_j^T (A - lI)^{-1} v_j \right)}$$

for some constant C>0. With this, the algorithm only needs to recompute $\{p_i\}_{i=1}^m$ after every $\Omega(n^{1-1/q})$ iterations. As the algorithm chooses $\Theta(n/\epsilon^2)$ vectors in total, the algorithm only recomputes the probabilities $\Theta(\frac{n^{1/q}}{\epsilon^2})$ times. In essence, [LS18] is constructed such that it uses the same probability $\{p_i\}_{i=1}^m$ to choose multiple vectors before the algorithm increases the barrier values u_j, l_j for each iteration. The algorithm is shown below:

Algorithm 12 [LS18]'s algorithm for fast-construction of spectral sparsifiers

```
Require: \epsilon \le 1/120, q \ge 10
```

- 1: j = 0
- 2: $l_0 = -(2n)^{1/q}, u_0 = (2n)^{1/q}, A_0 = 0$
- 3: while $u_j l_j < 4 \cdot (2n)^{1/q}$ do
- 4: $W_i = 0$
- 5: Compute $R_i(A_i, u_i, l_i)$ for all vectors v_i
- 6: Sample N_j vectors independently with replacement, where every v_i is chosen with probability proportional to $R_i(A_j, u_j, l_j)$. For every sampled v, add $\epsilon/q \cdot (R_i(A_j, u_j, l_j))^{-1} \cdot vv^T$ to W_j
- 7: $A_{j+1} = A_j + W_j$
- 8: $u_{j+1} = u_j + \Delta_{l,j}, l_{j+1} = l_j + \Delta_{l,j}$
- 9: j = j + 1
- 10: end while

To analyze the algorithm further, given the initial values j, l_0 , u_0 , A_0 , ϵ and q set in the formal framework of the algorithm, the algorithm proceeds in iterations where in each iteration j, it computes the relative effective resistance $R_i(A_j, u_j, l_J)$ of vectors $\{v_i\}_{i=1}^m$ where

$$R_i(A_j, u_j, l_j) \triangleq v_i^T (u_j I - A_j)^{-1} v_i + v_i^T (A_j - l_j I)^{-1} v_i$$

After this, the algorithm samples N_j vectors independently with replacement, where vector v_i is chosen with probability proportional to $R_i(A, u, l)$. The number of samples N_j in iteration j is defined by

$$N_j \triangleq \frac{1}{n^{2/q}} \left(\sum_{i=1}^m R_i(A_j, u_j, l_j) \min\{\lambda_{\min}(u_j I - A_J), \lambda_{\min}(A_j - l_j I)\} \right)$$

After each iteration j the algorithm updates u_j with $\Delta_{u,j}$, l_j with $\Delta_{l,j}$, and A_j with W where W is defined to be

$$W = \sum_{\text{sampled vector } v_i} \frac{\epsilon}{q} \cdot \frac{1}{R_i(A_j, u_j, l_j)} \cdot v_i v_i^T$$

Using the expected value of W defined as $E[W] = \frac{\epsilon}{q} \cdot \frac{N_j}{R_i(A_j, u_j, l_j)} \cdot I$, the algorithm increases the barrier values u_j and l_j by $\Delta u, j$ and $\Delta l, j$ respectively where

$$\Delta_{u,j} \triangleq (1+3\epsilon) \cdot \frac{\epsilon}{q} \cdot \frac{N_j}{\sum_{i=1}^m R_i(A_j, u_j, l_j)} \quad \text{and} \quad \Delta_{l,j} \triangleq (1-3\epsilon) \cdot \frac{\epsilon}{q} \cdot \frac{N_j}{\sum_{i=1}^m R_i(A_j, u_j, l_j)}$$

The choice of N_j , $\Delta_{u,j}$ and $\Delta_{l,j}$ ensures that $0 \leq W \leq \frac{1}{2} \cdot (u_j I - A_j)$ holds with high probability and that the invariant (shown below) holds for any phase j

$$l_j I \prec A_j \prec u_j I$$

In essence, by the last iteration T, $l_T I \prec A_T \prec u_T I$ will hold, resulting in the a matrix A_T that is linear-sized where $A_T \approx_{O(\epsilon)} I$.

6.2 Run-time of LS18's Fast Spectral Sparsifier algorithm

We want to show that the algorithm's run-time for producing a $(1+O(\epsilon))$ -spectral sparsifier is near-linear; specificially we want to show the algorithm's runtime being $\tilde{O}\left(\frac{qm}{\epsilon^2}\cdot n^{\omega-1+3/q}\right)$. For this construction, we use $v_1,...,v_N$ to denote the N sampled vectors and we denote the reweighted vectors for any $1 \le i \le N$ as $w_i \triangleq \sqrt{\frac{\epsilon}{q \cdot R_i(A,u,l)}} \cdot v_i$. With this, we let matrix W be $W \triangleq \sum_{i=1}^N w_i w_i^T$.

Lemma 6. Assume that the number of sampled vectors satisfies $N < \frac{2}{n^{2/q}} \Big(\sum_{i=1}^m R_i(A, u, l) \cdot \lambda(uI - A) \Big)$. It holds that

$$\mathbb{E}[W] = \mathbb{E}\left[\sum_{1}^{N} w_{i} w_{i}^{T}\right] = \frac{\epsilon}{q} \cdot \frac{N}{\sum_{i=1}^{m} R_{i}(A, u, l)} \cdot I$$

and

$$Pr[0 \leq W \leq \frac{1}{2} \cdot (uI - A)] \geq 1 - \frac{\epsilon^2}{100an}$$

Lemma 7. The output matrix A_k has a condition number at most $1 + O(\epsilon)$

Lemma 8. The following statements hold true:

- 1. The algorithm finishes in $\frac{10qn^{3/q}}{\epsilon^2}$ phases with probability at least 4/5
- 2. The algorithm chooses at most $\frac{10qn}{\epsilon^2}$ vectors with probability at least 4/5

Although we will not prove Lemma 7, its importance allows us to show that the Algorithm does produce a linear-sized $(1 + O(\epsilon))$ -spectral sparsifier by showing that the condition number of A_k is small. This is done by noticing that the barrier gap $u_j - l_j$ increases after every iteration j for point 1 and that the barrier gap $\Delta_{u,j} - \Delta l, j$ increases on average for every vector chosen in iteration j for point 2. Now, we want to construct an assumption for approximating the required quantities quickly for constructing graph sparsifiers. For this, let us assume the following: let L and \tilde{L} be the Laplacian matrices of graph G and its subgraph after re-weighting, let $A = L^{-1/2}\tilde{L}L^{-1/2}$, and assume that

$$(l+|l|\eta) \cdot I \prec A \prec (1-\eta)u \cdot I$$

holds for some $0 < \eta < 1$. Thus, the next two lemmas hold under this assumption:

Lemma 9. Let $A = \sum_{i=1}^{m} v_i v_i^T$ and suppose that A satisfies the assumption. Then, we can compute $\{r_i\}_{i=1}^m$ and $\{t_i\}_{i=1}^m$ in $\tilde{O}(\frac{m}{\epsilon^2 \eta})$ such that

$$(1 - \epsilon)r_i \le v_i^T (ul - A)^{-1} v_i \le (1 + \epsilon)r_i$$
 and $(1 - \epsilon)t_i \le v_i^T (A - lI)^{-1} v_i \le (1 + \epsilon)t_i$

Lemma 10. We can compute values α, β in $\tilde{O}(\frac{m}{ne^3})$ time such that

$$(1 - \epsilon)\alpha \le \lambda_{\min}(uI - A) \le (1 + \epsilon)\alpha$$
 and $(1 - \epsilon)\beta \le \lambda_{\min}(A - lI) \le (1 + \epsilon)\beta$

Using the assumption and the lemmas that follow from it, we can prove the following theorem:

Theorem 11. Let G = (V, E, w) be an undirected and weighted graph with n vertices and m edges, and let $q \ge 10, 0 < \epsilon \le 120$ be constants. Then there is an algorithm that outputs a $(1 + \epsilon)$ -spectral sparsifier of G with $O(qn/\epsilon^2)$ edges. The algorithm runs in $\tilde{O}(\frac{q \cdot m \cdot n^{5/q}}{\epsilon^{4+4/q}})$ time.

Proof. We want to construct our proof by firstly proving that the Algorithm does output a linear-size spectral sparsifer. WLOG, consider the graph G=(V,E,2) defined earlier. Suppose for every edge $e=(u,v)\in E$ where $u,v\in V$, we define the vector $b_e\in\mathbb{R}^n$ to be $b_e(x)=1$ if $x=u,b_e(x)=-1$ if x=v, and 0 otherwise. The Laplacian matrix of graph G would be $L=\sum_{e\in E}b_eb_e^T$. If we let $v_e=L^{-1/2}b_e$, we can show that constructing a spectral sparsifier of G is equivalent to sparsifying the matrix $\sum_{e\in E}v_ev_e^T$. Lemma 7 and 8 allow us to say that A_k is a linear-sized spectral sparsifier, partly as it allows us to prove that A_k is a $(1+O(\epsilon)$ approximation of I. To analyze the runtime of the algorithm, we know from Lemma 6 that with probability at least 9/10, all matrices picked in $k=\frac{10qn^{3/q}}{\epsilon^2}$ phases satisfy $W_j \leq \frac{1}{2}(u_jI-A_j)$. If this even occurs, then it holds for any phase j that

$$\mathbb{E}\Big[\Phi_{u_j,l_j}(A_j)|\forall j:W_j \leq \frac{1}{2}\big(u_jI - A_J\big)\Big] \leq 2$$

Thus, by Markov's inequality, for all iterations j, $\Phi_{u_j,l_j}(A_j) = O(qn^{3/q}/\epsilon^2)$ holds with high probability. Notice that for eigenvalues $\lambda_1, ..., \lambda_n$ of matrix A, it holds that

$$(u - \lambda_j)^{-q} \le \sum_{i=1}^n (u - \lambda_i)^{-q} < \Phi_{u,l}(A)$$

for any $1 \le j \le n$. This implies that $\lambda_j < u - (\Phi_{u,l}(A))^{-1/q}$ and $\lambda_j > l - (\Phi_{u,l}(A))^{-1/q}$. Thus, we have

$$\left(l_j + O\left(\left(\frac{\epsilon^2}{qn^{3/4}}\right)^{1/q}\right)\right)I \preceq A_j \preceq \left(u_j - O\left(\left(\frac{\epsilon^2}{qn^{3/4}}\right)^{1/q}\right)I$$

Since u_j and l_j are of the order $O(n^{1/q})$, we can set $\eta = O((\epsilon/n)^{2/q})$ to get

$$(l_j + |l_j|\eta)I \leq A_j \leq (1 - \eta)u_jI$$

We can apply Lemma 9 and 10 to compute all required quantities in each phase for a constant approximation in time.

$$\tilde{O}\left(\frac{m}{\epsilon^2} \cdot \eta\right) = \tilde{O}\left(\frac{m \cdot n^{2/q}}{\epsilon^{2+2/q}}\right)$$

Since Lemma 8 says the algorithm finishes in $\frac{10qn^{3/q}}{\epsilon^2}$ phases with probability 4/5, the total run-time of the algorithm is

$$\tilde{O}\left(\frac{m \cdot n^{2/q}}{\epsilon^{2+2/q}}\right) \cdot \frac{10qn^{3/q}}{\epsilon^2} = \tilde{O}\left(\frac{q \cdot m \cdot n^{5/q}}{\epsilon^{4+4/q}}\right)$$

which completes our proof

Now we want to show the goal of [LS18]: to find a near-linear time algorithm for outputting scalars $\{s_i\}_{i=1}^m$ with $|\{s_i:s_i\neq 0\}|=O(qn/\epsilon^2)$ such that

$$(1 - \epsilon) \cdot I \leq \sum_{i=1}^{m} s_i v_i v_i^T \leq (1 + \epsilon) \cdot I$$

Using Lemma 6, Theorem 11, and fast matrix multiplication, all required quantities can be computed for each iterature in $\tilde{O}(m \cdot n^{\omega-1})$ time (where ω is a matrix-multiplication constant). Thus, the total run-time is

$$\tilde{O}\left(\frac{qm}{\epsilon^2} \cdot n^{\omega - 1 + 3/q}\right) \tag{3}$$

which proves our run-time claim for the algorithm and that the algorithm runs in near-linear time.

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