Minor Sparsifiers and the Distributed Laplacian Paradigm

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Abstract

We study distributed algorithms built around edge contraction based vertex sparsifiers, and give sublinear round algorithms in the CONGEST model for exact mincost flow, negative weight shortest paths, maxflow, and bipartite matching on sparse graphs. For the maxflow problem, this is the first exact distributed algorithm that applies to directed graphs, while the previous work by [Ghaffari et al. SICOMP’18] considered the approximate setting and works only for undirected graphs. For the mincost flow and the negative weight shortest path problems, our results constitute the first exact distributed algorithms running in a sublinear number of rounds. These algorithms follow the celebrated Laplacian paradigm, which numerically solve combinatorial graph problems via series of linear systems in graph Laplacian matrices.

To enable Laplacian based algorithms in the distributed setting, we develop a Laplacian solver based upon the subspace sparsifiers of [Li, Schild FOCS’18]. We give a parallel variant of their algorithm that avoids the sampling of random spanning trees, and analyze it using matrix martingales. Combining this vertex reduction recursively with both tree and elimination based preconditioners leads to an algorithm for solving Laplacian systems on $n$ vertex graphs to high accuracy in $O(n^{o(1)}(\sqrt{n} + D))$ rounds. The round complexity of this distributed solver almost matches the lower bound of $\tilde{\Omega}(\sqrt{n} + D)$.

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1 Introduction

The steady growth of data makes it increasingly important to control and reduce the communication of algorithms. The CONGEST model [Pel00] is a widely studied model for low communication algorithms on large graphs and sparse matrices. In this model, each vertex/variable occupies a separate machine, and communicates in synchronous rounds by sending messages of length \( O(\log n) \) to its neighbors given by the edges of the underlying graph. This bandwidth restriction implies a polynomial lower bound in the round complexity for many fundamental graph problems [PR00, Elk06, SHK+12]. While early work on efficient algorithms in this model has focused on the minimum spanning tree problem [GHS83, GKP98, KP98], extensive work over the past few years has led to efficient algorithms for several more fundamental graph problems, such as approximate and exact single-source shortest paths [Nan14, HKN16, Elk20, GL18, FN18, CM20], approximate and exact all-pairs shortest paths [HW12, HNS17, ARKP18, EN18, LPP19, AR19, BN19, AR20], approximate and exact minimum cut [GK13, NS14, DHNS19, GNT20, DEMN20], approximate maximum flow [GKK+15], bipartite maximum matching [AKO18], triangle counting [IG17, CPZ19, CS19], and single-source reachability [GU15, JLS19].

A major development in sequential and parallel graph algorithms is the development of hybrid algorithms that combine numerical and combinatorial building blocks. This line of work was initiated by the seminal work of Spielman and Teng [ST14], which showed that a Laplacian linear system on a graph can be solved in nearly linear time. Here, the Laplacian of a weighted undirected graph \( G = (V, E) \) is defined as \( L(G) = D(G) - A(G) \), where \( D(G) \) is the diagonal weighted degree matrix, and \( A(G) \) is the weighted adjacency matrix. Equivalently, if \( \bar{w} \in \mathbb{R}_{\geq 0}^m \) are the edge weights,

\[
L(G)_{uv} = \begin{cases} 
\sum_{(u,z) \in E} \bar{w}_{uz} & \text{if } u = v, \\
-\bar{w}_{uv} & \text{otherwise}.
\end{cases}
\]

Since then, there has been extensive work towards giving more efficient and simpler Laplacian system solvers sequentially [KMP10, KMP11, KOSA13, CKM+14, KS16], as well as parallel versions [PS14, KLP+16]. These results have in turn been used to give the state-of-the-art runtimes for a variety of graph problems, including exact maximum flows, bipartite matchings, and mincost flows [Mad16, LS20b, LS20a, CMSV17, AMV20, BLN+20], approximate maximum flows [KLOS14, She13, Pen16], and approximate parallel shortest paths [Li20, ASZ20]. Ideas from the latter works have found application in the distributed setting, giving nearly optimal algorithms for approximate maxflows [GKK+15] and approximate shortest paths [BKKL17] in the CONGEST model.

Our main result is an algorithm for solving graph Laplacian linear systems in the CONGEST model in \( O(n^{o(1)}(\sqrt{n} + D)) \) rounds (Theorem 1), where \( D \) is the diameter of the underlying graph. This nearly matches a lower bound of \( \Omega(\sqrt{n} + D) \), which we show for completeness in Section A by reduction to [SHK+12].

**Theorem 1.** There is an algorithm in the CONGEST model that on a weighted graph \( G = (V, E, \bar{w}) \) with \( n \) vertices and diameter \( D \), vector \( \bar{b} \) on the vertices, and error \( \epsilon < 0.1 \), produces in \( O(n^{o(1)}(n^{1/2} + D) \log(1/\epsilon)) \) rounds a vector \( \bar{x} \) distributed over the vertices such that

\[
\left\| \bar{x} - L(G)^{\top} \bar{b} \right\|_{L(G)} \leq \epsilon \cdot \left\| \bar{b} \right\|_{L(G)}.
\]
Theorem 2. In the CONGEST model of computation, solving Laplacian systems to accuracy $\epsilon \leq \frac{1}{2}$ requires at least $\tilde{\Omega}(n^{1/2} + D)$ rounds of communication.

We give several applications towards designing hybrid algorithms for graph problems in the CONGEST model. Specifically, by combining our Laplacian solver with interior point methods [Mad16, CMSV17], we obtain the first algorithms for exact computation of maximum flows, bipartite matchings, and negative-weight shortest paths that run in a sublinear number of rounds in the CONGEST model on sparse graphs (Section 7).

At a high level, we build our CONGEST model algorithm by first building a parallel / PRAM algorithm for solving Laplacian systems that only works with minors of the original graph, and show that one round of communication necessary in our algorithm (such as matrix vector multiplication) between neighbors on a minor can be simulated in the original graph in $\tilde{O}(\sqrt{n} + D)$ rounds. Previous methods for computing low stretch spanning trees and approximate maximum flows [GKK+15] use a similar notion of considering a graph on clusters of nodes in the original graph, however we believe that we are the first to work explicitly with the notion of minors. We are optimistic that our approach based on minor vertex sparsifiers may provide a general framework for designing CONGEST model algorithms with near optimal complexities.

The main backbone of our algorithm for solving Laplacian systems that works with minors only is the parallel Laplacian solver of [KLP+16]. This algorithm eliminates onto Schur complements of the original graph while adding edges, leading to graphs that are not minors of the original graph. To resolve this, a major contribution of this paper is an efficient parallel algorithm to construct a spectral sparsifier for a Schur complement which is a minor of the original graph. While the existence of such a minor spectral sparsifier was known [LS18], the algorithm required sampling a random spanning tree, and hence could not be implemented in parallel. We instead show that a large batch of edges may be independently sampled via approximate leverage scores, providing an arguably simpler and more direct analysis than [LS18].

1.1 Applications to Flow Problems

We briefly discuss how our Laplacian solver can be applied to achieve results on maximum flow, bipartite matching, mincost flows, and negative weight shortest paths, and compare to previous complexities. We achieve our bounds by combining our Laplacian system solver in Theorem 1 with recent interior point methods of [Mad16, CMSV17].

For unit capacity graphs, the runtimes we achieve in Theorems 7, 8, and 9 for the maximum flow problem, mincost flow, and negative weight shortest path problems are

$$O(m^{3/7+o(1)}(n^{1/2}D^{1/4} + D)).$$

For sparse unweighted graphs with $m = O(n)$, and polynomially small diameter $D = n^{2/7-\Omega(1)}$, the algorithms in Theorems 7, 8, and 9 run in a sublinear number of rounds, i.e. $n^{1-\Omega(1)}$ rounds. To our knowledge, these are the first exact sublinear round algorithms for unit maximum flows, bipartite matchings, and negative weight shortest paths for any regime of diameter $D$. Our distributed maximum flow algorithm extends to directed graphs, while the previous work by Ghaffari et al. [GKK+15] considered the approximate setting and works only for undirected graphs. In fact, for the maximum flow problem, we believe that our results are the best known in all regimes – see Section 1.2 for further discussion.
At a high level, our runtime comes from two pieces. The results of [Mad16, CMSV17] show that in $\tilde{O}(m^{3/7})$ rounds of an interior point method, each round which involves solving a Laplacian system on the underlying graph with edge weights \( / \) resistances, we can reduce the amount of residual flow to $\tilde{O}(m^{3/7})$. The residual flow can routed combinatorially with $\tilde{O}(m^{3/7})$ rounds of an augmenting paths or shortest paths computation. Therefore, the total number of rounds required to implement the interior point method is $O(m^{3/7+o(1)}(n^{1/2} + D))$ using Theorem 1, and the shortest path path computations can be done in $\tilde{O}(m^{3/7}(n^{1/2}D^{1/4} + D))$ rounds using the results of [CM20]. Combining these gives the result.

1.2 Related Work

Distributed Graph Algorithms Previous works in distributed algorithms most related to our result are arguably the algorithms for simulating random walks and generating random spanning trees [DSNPT13, GB20]. On unweighted, undirected graphs with diameter \( D \), the algorithms by Das Sarma, Nanongkai, Pandurangan, and Tetali [DSNPT13] generate an \( \ell \)-step random walk in $\tilde{O}(\sqrt{D} + D)$ rounds, and a random spanning tree in $\tilde{O}(\sqrt{mD})$ rounds, respectively. There are well known connections between sampling a large number of random walks and Laplacian solving [DST17, DGT17]. However, it is not clear how to utilize these methods in the context of our algorithms, since many of the intermediate graph structures we deal with involve dealing with weighted random walks, which in turn leads to congestion issues when trying to simulate these walks in the distributed setting. We discuss how to overcome such obstacles in Section 3.

There also has been work in the distributed setting relating to spectral graph properties. This in particular includes distributed sparsification [KX16], PageRank [SMPU15], Laplacian solvers in well-mixing settings [GB20], and expander decomposition [CPZ19, CS19, CS20].

Continuous optimization methods have been used to give the state-of-the-art distributed algorithms for approximate max-flow [GKK+15] and approximate transshipment [BKKL17]. Note however that these approximation algorithms are tailored to undirected graphs and their running time depends polynomially on \( 1/\epsilon \) for a desired accuracy of \( \epsilon \). Our max-flow routine also works on directed graphs and only depends polylogarithmically on \( 1/\epsilon \), which allows for computing a high-accuracy solution and rounding it to an exact one. Furthermore, these prior approaches for \( \ell_\infty \) and \( \ell_1 \)-norm minimization, respectively, do not carry over to \( \ell_2 \)-norm minimization (as would be needed for solving Laplacian systems) because it is not known how to efficiently sample from a collection of trees when using an \( \ell_2 \) variant of tree-based graph approximations to build oblivious routing schemes.

In addition to these works, there are many papers related to the three problems we solve by applying our distributed Laplacian solver. There have been numerous results on exact and approximate shortest path computation in the past decade [HW12, Nan14, HKN16, EN19a, Elk20, HNS17, GL18, ARKP18, EN18, FN18, LPP19, EN19b, AR19, BN19, AR20, CM20]. For the single-source shortest paths (SSSP) problem all of these works assume non-negative or positive edge weights. It is well-known that the SSSP problem in presence of negative edge weights can be solved in $O(n)$ rounds by a variant of the Bellman-Ford algorithm. To the best of our knowledge no algorithm that improves upon this bound has been formulated (or implied) in the CONGEST model so far.

For distributed computations of maximum flows, Ghaffari, Karrenbauer, Kuhn, Lenzen, and Patt-Shamir [GKK+15] designed an algorithm that returns an \((1 + \epsilon)\)-approximation in $O((\sqrt{n} + D)n^{o(1)}\epsilon^{-3})$ rounds. In terms of exact algorithms, we are not aware of any paper claiming a sub-linear number of rounds in the CONGEST model (cf. [GKK+15] for a detailed discussion of maximum
flow for other distributed models). To the best of our knowledge, we need to compare ourselves with the following two approaches:

- The problem can trivially be solved in $O(m + D)$ rounds by collecting the whole graph topology in a single node and then solving the problem with internal computation.
- The Ford-Fulkerson algorithm [FF56] takes $|f^*|$ iterations (where $|f^*|$ is the value of a maximum flow) and the running time in each iteration is dominated by the time needed to perform an $s$-$t$ reachability computation (on a directed graph). The latter problem can be solved in $\tilde{O}(\sqrt{n}D^{1/4} + D)$ [GU15] or $\tilde{O}(\sqrt{n} + n^{1/3+o(1)}D^{2/3})$ rounds, respectively, which yields total running time of $\tilde{O}(|f^*|)(\sqrt{n}D^{1/4} + D)$ or $\tilde{O}(|f^*|)(\sqrt{n} + n^{1/3+o(1)}D^{2/3}))$ rounds, respectively. In unit-capacity ("unweighted") graphs, where $|f^*| \leq n$, this gives a total running time of $\tilde{O}(n^{3/2}D^{1/4} + nD)$ or $\tilde{O}(n^{3/2} + n^{4/3+o(1)}D^{2/3})$, respectively.

Due to a well-known reduction to maximum flow, the bipartite maximum matching problem is intimately connected to the maximum flow problem. In the CONGEST model, the fastest known algorithm for computing a bipartite maximum matching (of an unweighted graph) takes $O(n \log n)$ rounds [AKO18] – more precisely the algorithm takes $O(s^* \log s^*)$ rounds, where $s^*$ is the size of a maximum matching. Obtaining a subquadratic maximum matching algorithm for networks of arbitrary topology is a major open problem [AK20]. In addition, there are numerous works on computing approximate matchings, which are usually based on computing a maximal matching, using the framework of Hopcroft and Karp [HK73], or rounding a fractional matching (cf. [AK20] for an overview on approximate matching algorithms in the CONGEST model).

**Laplacian Solvers** Our algorithm combines both tree-based ultrasparsification algorithms [ST14, KMP10, KMP11, CKM+14] and elimination-based algorithms that utilize Schur complements [KLP+16, KS16, Kyn17]. Both types of algorithms were originally developed for the sequential model. The issue of round complexity was previously addressed in parallel Laplacian solving [BGK+14, PS14].

We believe that a variant of [BGK+14] tailored to the CONGEST model gives a round complexity of around $n^{3/4} + Dn^{1/4}$ as opposed to the bound in Theorem 1 to because the depth of the parallel algorithm of [BGK+14] is more than polylogarithmic. The polylogarithmic depth parallel algorithm from [PS14] is more difficult to convert to the CONGEST setting because it explicitly adds edges to the graph, which causes increased congestion.

The outer layer recursion of our algorithm is akin to the recursive construction of solvers and preconditioners present in Laplacian solving [Pen13, KLP+16], approximate max-flow [Pen16], and matrix sampling [CP15, CLM+15, CMM17].

Parallel Laplacian solvers and spectral algorithms have also motivated the study of (nearly) log space variants of these algorithms [MRSV17, MRSV19, AKM+19]. It’s an intriguing question to formally connect these low space algorithms with distributed algorithms, both of which stem from works on low iteration count algorithms.

**Vertex Sparsification** Critical to our result is the construction of minor based Schur complements by Li and Schild [LS18]. Minor based sparsification has been studied for distances [CGH16, KNZ14], and implicitly for cuts via hierarchical routing schemes [Räc02, RST14]. A more systematic treatment of uses of such sparsifiers, in dynamic graph algorithms, can be found in [Gor19]. Some of the cut preserving vertex sparsifiers [Moi09, LM10, CLLM10, EGK+10, KR13], as well as their recent variations in small cut settings [CDL+20] produce either minors or probability distribution over minors.
2 Preliminaries

We start by describing general notation we use throughout the paper.

**General notation.** Given a symmetric matrix $M$, we let \( \|M\|_2 = \max_{\|x\|=1} |x^\top M x| \) denote the maximum absolute value of any eigenvalue. For a vector $v$ and matrix $M$, we define \( \|v\|_M = \sqrt{v^\top M v} \). For positive real numbers $a, b$ we say that $a \approx_b b$ if \( \exp(-e) a \leq b \leq \exp(e) a \). We say that a matrix $M \in \mathbb{R}^{n \times n}$ is positive semidefinite if $x^\top M x \geq 0$ for all $x \in \mathbb{R}^n$. We say that matrices $A \preceq B$ if $B - A$ is positive semidefinite. For positive semidefinite matrices $A, B$ we say that $A \approx_e B$ if \( \exp(-e) A \preceq B \preceq \exp(e) A \).

**Schur complements and Cholesky factorization.** Our algorithms are based on Schur complements and sparsified Cholesky factorization. At a high level, the Schur complement of an $n \times n$ matrix provides a matrix which is equivalent under linear system solves on a subset of coordinates in $[n]$.

**Definition 2.1 (Schur complement).** For an $n \times n$ symmetric matrix $M$ and subset of terminals $T \subseteq [n]$, let $S = [n] \setminus T$. Permuted the rows / columns of $M$ to write

\[
\begin{pmatrix}
M_{S,S} & M_{S,T} \\
M_{T,S} & M_{T,T}
\end{pmatrix}
\]

Then the Schur complement of $M$ onto $T$ is denoted $\text{SC}(M, T) \overset{\text{def}}{=} M_{T,T} - M_{T,S} M_{S,S}^{-1} M_{S,T}$.

For a graph $G$ and subset $T \subseteq V(G)$, for simplicity we write $\text{SC}(G, T) \overset{\text{def}}{=} \text{SC}(L_G, T)$. It is well-known that $\text{SC}(G, T)$ is also a Laplacian.

**Lemma 2.2 (Cholesky factorization).** Given a matrix $M \in \mathbb{R}^{n \times n}$, a subset $T \subseteq [n]$, and $S = [n] \setminus T$, we have

\[
M^{-1} = \begin{pmatrix}
I & -M_{S,S}^{-1} M_{S,T} \\
0 & I
\end{pmatrix}
\begin{pmatrix}
M_{S,S}^{-1} & 0 \\
0 & \text{SC}(M, T)^{-1}
\end{pmatrix}
\begin{pmatrix}
I & 0 \\
-M_{T,S} M_{S,S}^{-1} & I
\end{pmatrix}.
\]

The Cholesky factorization directly implies that the Schur complement represents the inverse of the Laplacian on a subset of the coordinates.

**Lemma 2.3** (e.g. Fact 5.4 in [DKP+17]). Let $I$ be the identity matrix, and let $J$ be the all 1 matrix. For any graph $G$, and subset $T \subseteq V(G)$ we have that

\[
\text{SC}(G, T)^\dagger = (I - |T|^{-1} J)(L_G)^{\dagger}|_{T,T} (I - |T|^{-1} J).
\]

In addition, we have that

\[
\text{SC}(G, T)(L_G)^{\dagger}|_{T,T} \text{SC}(G, T) = \text{SC}(G, T).
\]

An equivalent view is that the quadratic form of the Schur complement gives the minimum energy extension of a vector on the terminals to the original vertex set, in the quadratic form of the original Laplacian $[\text{Gre96, MP13}]$.

**Lemma 2.4.** (Lemma B.2. of [MP13], matrix version in Appendix A.5.5 of [BBV04]) For a graph $G$ and a $T \subseteq V(G)$, the Schur complement of the Laplacian of $G$ onto $T$, $\text{SC}(G, T)$ satisfies for all vectors $\bar{x}[T]$

\[
\|\bar{x}[T]\|_{\text{SC}(G, T)} = \min_{\bar{x}[V \setminus T] \in \mathbb{R}^{V \setminus T}} \left\| \begin{bmatrix} \bar{x}[V \setminus T] \\ \bar{x}[T] \end{bmatrix} \right\|_{L(G)}.
\]
Matrix Analysis Tools Our algorithm for computing Schur complement sparsifiers which are minors requires computing and sampling via leverage scores.

Definition 2.5 (Effective resistance and leverage scores). For a graph $G$ with resistances $r_e$, define $\text{res}_G(e) \overset{\text{def}}{=} b_e^T L_G^{-1} b_e$ and $\text{lev}_G(e) \overset{\text{def}}{=} \text{res}_G(e) / r_e$.

Note that $0 \leq \text{lev}_G(e) \leq 1$ and $\sum_{e \in E(G)} \text{lev}_G(e) = n - 1$ for connected graphs $G$.

Let $G \setminus e$ and $G / e$ denote the graphs resulting respectively from deleting and contracting edge $e$.

Note that these correspond respectively to setting the resistance of edge $e$ to positive infinity or 0.

The Woodbury matrix formula allows us to understand changes in the quadratic form under resistance changes.

Lemma 2.6 (Woodbury matrix formula). For matrices $A$, $U$, $C$, $V$ of compatible sizes we have

$$(A + UCV)\dagger = A\dagger - A\dagger U(C^{-1} + VA\dagger U)^{-1}VA\dagger.$$

We use the following to understand the matrix martingales that arise in the analysis.

Lemma 2.7 (Freedman’s inequality for matrix martingales [Tro11]). Consider a matrix martingale $(Y^{(k)})_{k \geq 0}$ whose values are symmetric matrices with dimension $d$ and let $(X^{(k)})_{k \geq 1}$ be the difference sequence $X^{(k)} \overset{\text{def}}{=} Y^{(k)} - Y^{(k-1)}$. Assume that the difference sequence is uniformly bounded in that $\|X^{(k)}\|_2 \leq R$ almost surely for $k \geq 1$. Define the predictable quadratic variation random matrix $W^{(k)} \overset{\text{def}}{=} \sum_{j=1}^k \mathbb{E}[(X^{(j)})^2 | X^{(j-1)}]$.

Then for all $\epsilon \geq 0$ and $\sigma^2 > 0$ we have that

$$\Pr \left[ \exists k > 0 : \|Y^{(k)} - Y^{(0)}\|_2 \geq \epsilon \text{ and } \|W^{(k)}\|_2 \leq \sigma^2 \right] \leq 2d \cdot \exp \left( -\epsilon^2 / 3 \sigma^2 + R\epsilon / 3 \right).$$

The 2-norm of a symmetric matrix is bounded by its maximum row sum.

Lemma 2.8. For a symmetric matrix $M \in \mathbb{R}^{n \times n}$, we have that

$$\|M\|_2 \leq \max_{i \in [n]} \sum_{j \in [n]} |M_{ij}|.$$

Proof. For all vectors $\bar{x}$, note by the AM-GM inequality that

$$\bar{x}^T M \bar{x} = \sum_{1 \leq i, j \leq n} \bar{x}_i \bar{x}_j M_{ij} \leq \sum_{1 \leq i, j \leq n} \bar{x}_i^2 |M_{ij}| \leq \max_{i \in [n]} \sum_{j \in [n]} |M_{ij}| \sum_{i \in [n]} \bar{x}_i^2 \leq \max_{i \in [n]} \sum_{j \in [n]} |M_{ij}| \|x\|_2^2.$$

$\square$
**CONGEST model** In the CONGEST model [Pel00], we are given a communication network \( \mathcal{G} = (\mathcal{V}, \mathcal{E}) \) with \( \pi \) nodes modelling processors that have unique \( O(\log n) \)-bit IDs, \( m \) edges modelling bidirectional communication links between the processors, and diameter \( D \). Initially, each node knows its own ID and the IDs of its neighbors as well as the value of \( n \). Computation in this model is carried out in rounds synchronized by a global clock. In each round, every node sends to each of its neighbors an arbitrary \( O(\log n) \)-bit message, receives the messages of its neighbors, performs arbitrary internal computation, and stores arbitrary information for the next round. The main goal in this paper is to design algorithms for graph problems with a small number of rounds. For problems on directed graphs, the direction of each edge is known by both of its endpoints, but the corresponding communication link is still bidirectional. For problems on weighted graphs (involving, e.g., costs or capacities), the weight of each edge is known by both of its endpoints, but the corresponding communication link still allows for direct transmission of each message within a single round. In particular the diameter \( D \) always refers to the underlying undirected, unweighted communication network. Our running time bounds hold under the assumption that all weights are polynomial in \( n \), which is a standard assumption in the CONGEST model literature.

**3 Overview**

Here, we will give an overview of the concepts behind our algorithm. The core component of our algorithm is the distributed Laplacian solver given in Theorem 1. We then apply this algorithm to combinatorial graphs using ideas from the Laplacian paradigm, namely interior point methods for solving optimization problems using linear systems solves.

**Laplacian Solver: from PRAM to Distributed** Our algorithm is based on \( \text{poly}(\log n) \) round Laplacian system solvers in the PRAM model, namely the sparsified Cholesky algorithm from [KLP+16]. This algorithm repeatedly finds a constant fraction of the vertices on which the block minor is “almost independent” and hence easy to solve. The inverse of this block then gives the result of eliminating these vertices, which is the Schur complement on the rest of the vertices, which we view as the terminal vertices \( \mathcal{T} \).

More explicitly, this can be seen in the context of the Cholesky factorization in Lemma 2.2, where we let \( M = L(G) \) be the Laplacian. We find an “almost independent” set of vertices \( S \) so that computing \( M^{-1}[S,S] \) to high accuracy is simple using a preconditioned gradient descent method. Therefore, the remaining difficulty in computing \( M^{-1} \) is simply from computing and inverting the Schur complement: \( \text{SC}(M, \mathcal{T})^{-1} \). To do this, we first approximately compute the Schur complement \( \text{SC}(M, \mathcal{T}) \), which is again a Laplacian, and then recursively apply a Cholesky factorization to it again.

However, this resulting Schur complement may be dense, even if the original graph is sparse. For example, eliminating the center of a star results in a complete graph on the peripheral leaf vertices. To make this more efficient, sparsified elimination algorithms [KLP+16, CGP+18, DPPR20] seek to directly construct a sparse approximation of this Schur complement. This can be done in a variety of ways, but algorithmically one of the simplest interpretations is through the sampling of random walks. Indeed, matrix concentration bounds that the following procedure suffices for generating a good approximation of \( \text{SC}(G, \mathcal{T}) \) with high probability:

By picking \( \mathcal{T} \) so that \( V \setminus \mathcal{T} \) is almost independent, that is, each vertex not in \( \mathcal{T} \) has a constant fraction of its weight going to \( \mathcal{T} \), it can be ensured that the lengths of the walks don’t exceed \( O(\log n) \)
Algorithm 1: Approximate Schur Complement using Random Walks

1 Set $H \leftarrow \emptyset$
2 for each edge $e = uv$ in $G$ do
3     Repeat the following two steps $O(\epsilon^{-2} \log n)$ times:
4         Random walk both endpoints $u$ and $v$ until they are in $T$, to $t_u$ and $t_v$ respectively.
5         Add an edge to the approximate Schur complement $H$ between $t_u$ and $t_v$, with weight as
6 return $H$

with high probability. As a result, PRAM algorithms are able to construct low error Schur complements
by sampling about $O(\epsilon^{-2} \log n)$ walks of length $O(\log n)$ per edge. As the number of vertices in the
Schur complement decreases by a constant factor per step, this process yields a parallel solver with
another $O(\log n)$ factor overhead in parallel depth.

In the distributed setting, however, a major difficulty with using such approximate Schur comple-
ments is that they may highly congest some edges. Recall that in the CONGEST model, each edge can
only pass $O(\log n)$ bits per round. Random walks in weighted graphs on the other hand may severely
congest some edges: consider for example, a star with one very heavily weighted edge, and rest lightly
weighted. All the walks starting from the lightly weighted edges’ end points will likely utilize the heav-
ily weighted edge, leading to a congestion of $\Omega(n)$ in the worst case. This requires us to use a procedure
to estimate the congestion of an edge accumulated by such random walks, and to add their endpoints
back in as additional terminals.

However, a single elimination round only removes a constant fraction of the vertices, but perform-
ing $\Omega(\log n)$ elimination rounds would add a significant number of new edges to the graph. Hence, we
only perform $\Theta((\log \log n)^2)$ rounds of elimination between sparsification steps. A formal statement
of this elimination scheme is shown in Lemma 4.10.

Minor Sparsifiers and its Distributed Construction After that, the core component of our algo-
rrithm is that we must bring the Schur complement back to being a minor of the original graph, by
constructing a spectral sparsifier of the Schur complement which is a minor of the original graph (The-
orem 3). That is, the Schur complement results from contracting connected subsets of vertices in the
original graph and reweighting edges. Existence and efficient sequential constructions of these objects
were first shown by Li and Schild [LS18].

Minors are particularly useful for distributed algorithms because we can simulate one round of
communication between neighbors on a minor, such as multiplying by the incidence matrix, in $\tilde{O}(\sqrt{n}+D)$ rounds (Lemma 4.3). They interact particularly well with the parallel Laplacian solving algorithm
which is a short sequence of matrix-vector multiplies on submatrices.

Our construction algorithm is also a parallelized version of the Li-Schild algorithm [LS18]. The
algorithm works by contracting or deleting edges $e$ with probability given by its leverage score. The
main difference is that instead of sampling edges using a random spanning tree, we identify a large
subset of edges that can be sampled independent of each other, without affecting each edges’s sampling
probability too much. This is done via localization [SRS18], which provides an overall bound on the
total influence of edges’ effective resistances.

The algorithm then comprises of three main steps, and is analyzed via matrix martingales.
1. Calculating edges’ influence on the Schur complement (Lemma 5.15).

2. Computing the mutual influence of edges’s resistances, and picking a large set that has small mutual influence, which we term the steady set (Definition 5.1).

3. Among these steady edges, randomly contract / delete them with probability given by an approximation of their leverage scores.

Note that all steps actually require solving Laplacian systems in the original graph, which seems circular. We address this using the now well understood recursive approach of [Pen16], which we discuss below together with the overall algorithm.

**Overall Recursive Scheme**  Given a graph $G$, the goal of the algorithm is to return a chain of approximate Schur complements of $G$, each with 0.99 as many vertices as the previous. This chain has length $O(\log n)$, and after built, can be applied in $O(\log n)$ steps and $\tilde{O}(\sqrt{n} + D)$ rounds to solve a Laplacian system to high accuracy in the CONGEST model. The construction of the chain is as follows – pick $d = \Theta((\log \log n)^2)$ say, and run $d$ rounds of the sparsified Cholesky elimination scheme (Lemma 4.10) to reduce the graph size to $0.99^d |V(G)|$. Now, use the minor Schur complement algorithm (Theorem 3) to build a minor of $G$ which is a Schur complement sparsifier with respect to the remaining $0.99^d |V(G)|$ remaining vertices.

To compute the Schur complement sparsifier, we employ a separate recursion, because the Schur complement sparsifier construction requires Laplacian system solves to compute leverage scores (and other similar measures). To do this, we ultrasparsify the graph $G$, thus reducing the size by a factor of $k$ (Lemma 4.9), and build a Schur complement chain on the ultrasparsifier. Now, we can use this solver on the ultrasparsifier to precondition a solver on $G$ with $\tilde{O}(\sqrt{k})$ steps of preconditioned conjugate gradient to compute the desired leverage scores. We want to emphasize the final Schur complement chain we output for $G$ does not involve the ultrasparsifier, and hence can still be applied in parallel.

One final technical detail is that due to needing to solve submatrices of the Laplacian (Lemma 5.15) we require tracking graphs that embed with low congestion in the original graph, a slight generalization of minors (Definition 4.1). We ensure that the congestion stays as $n^{o(1)}$ throughout the algorithm, so it does not affect the final round complexity.

4 Full Algorithm and Analysis

The goal of this section is to formalize the notions and graph reduction algorithms described in Section 3, and provide a bound for the overall performance.

4.1 Distributed Communication on Minors of Overlay Networks

As described, we will work with graphs that are minors of the original graph, which doubles as the communication network. However, some of our linear systems reductions duplicate edges, leading to minors with slightly larger congestion. So we will need to incorporate such congestion parameters into our definition of minors. The following definition is a direct extension of the distributed $N$-node cluster graph from [GKK+15], with congestion incorporated, and the connection with graph minors stated more explicitly.

**Definition 4.1.** A graph $G$ is a minor of $H$ if we have the following mappings:
1. For each vertex of \( G \), \( u \in V(G) \):

   (a) A subset of vertices of \( H \), which we term a supervertex, \( S^{G \to H}(u) \subseteq V(H) \), with a root vertex \( V^{G \to H}_\text{map}(u) \in S^{G \to H}(u) \).

   (b) A connected subgraph of \( H \) on \( S^{G \to H}(u) \), which for simplicity we will keep as a tree, \( T^{G \to H}(u) \). Note that this requires \( S^{G \to H}(u) \) being connected in \( H \).

2. A mapping of the edges of \( G \) onto edges of \( H \), or self-loops on vertices of \( H \), such that for any \( u^G v^G = e^G \in E(G) \), the mapped edge \( E^{G \to H}_\text{map}(e^G) = e^H = u^H v^H \) satisfies \( u^H \in S^{G \to H}(u^G) \) and \( v^H \in S^{G \to H}(v^G) \).

and we say this minor has congestion \( \rho \), or \( G \) is a \( \rho \)-minor of \( H \) if:

1. Each vertex of \( H \) is contained in at most \( \rho \) supervertices \( V^{G \to H}_\text{map}(v^G) \) for some \( v^G \).

2. Each edge of \( H \) appears as the image of the edge map \( E^{G \to H}_\text{map}(\cdot) \), or in one of the trees connecting supervertices, \( T^{G \to H}(v^G) \) for some \( v^G \), at most \( \rho \) times.

Finally, we say a \( \rho \)-minor mapping is stored distributedly, or that \( G \) is \( \rho \)-minor distributed over \( H \) if it’s stored by having all the images of the maps recording their sources. That is, each \( v^H \in V(H) \) records

1. All \( v^G \) for which \( v^H \in V_{\text{map}}^{G \to H}(v^G) \),

2. For each edge \( e^H \) incident to \( v^H \) (including self loops that may not exist in original \( H \)):

   (a) All vertices for which \( e^H \) is in the corresponding tree
   
   \[ \{ v^G \mid e^H \in T^{G \to H}(v^G) \} \]

   (b) All edges \( e^G \) that map to it.

We will denote the original graph, which doubles as the overlay network, using \( \overline{G} \).

Note that the vertex mappings, or even the neighborhoods of \( G \), cannot be stored at one vertex in \( \overline{G} \). This is because both of these sets may have size up to \( \Omega(n) \), and passing that information to a single low degree vertex would incur too much communication.

We store vectors on \( G \) by putting the values at the root vertices of each of its corresponding supervertices. This notion of rooting can be made more explicit: we can compute directions for all edges in the spanning tree \( T^{G \to \overline{G}}(v^G) \) that point to the corresponding root vertex \( V^{G \to \overline{G}}_\text{map}(v^G) \).

**Lemma 4.2.** Given a graph \( G \) that’s \( \rho \)-minor distributed over a communication network \( \overline{G} = (\overline{V}, \overline{E}) \) with \( \overline{n} \) vertices, \( \overline{m} \) edges, and diameter \( D \), we can compute in \( O(t \rho \sqrt{\overline{n}} \log \overline{n} + D) \) rounds of communication on \( \overline{G} \), an orientation for each \( v^G \) and each edge \( e \in T^{G \to \overline{G}}(v^G) \) such that each vertex other than the root has exactly one edge pointing away from it, and following these edges leads us to the root.

We will make extensive usage of the following lemma, which we prove in Appendix B, about simulating communications on \( G \) using rounds of CONGEST communications in a graph that it \( \rho \)-minor distributes into.
Lemma 4.3. Let \( G = (V, E) \) be a graph with \( n \) vertices and \( m \) edges that \( \rho \)-minor distributes into a communication network \( \overline{G} = (\overline{V}, \overline{E}) \) with \( \overline{n} \) vertices, \( \overline{m} \) edges, and diameter \( D \). In the CONGEST model, the following operations can be performed with high probability using \( O(t \rho \sqrt{\overline{n}} \log \overline{n} + D) \) rounds of communication on \( \overline{G} \):

1. Each \( \overline{V}^{G \rightarrow \overline{G}}_{\text{map}}(v^G) \) sends \( O(t \log n) \) bits of information to all vertices in \( S^{G \rightarrow \overline{G}}(v^G) \).

2. Simultaneously aggregate the sum/minimum of \( O(t \log n) \) bits, from all vertices in \( S^{G \rightarrow \overline{G}}(v^G) \) to \( \overline{V}^{G \rightarrow \overline{G}}_{\text{map}}(v^G) \) for all \( v^G \in V(G) \).

One direct use of this communication result is that it allows us to efficiently compute matrix-vector products.

Corollary 4.4. Given a matrix \( A \) with nonzeroes supported on the edges of a graph \( G \) that’s \( \rho \)-minor distributed over a communication network \( \overline{G} = (\overline{V}, \overline{E}) \) with \( \overline{n} \) vertices, \( \overline{m} \) edges, and diameter \( D \), with values stored with endpoints of the corresponding edge, and a vector \( \overline{x} \in \mathbb{R}^{\overline{V}(G)} \) stored distributedly on the vertices \( \overline{V}^{G \rightarrow \overline{G}}_{\text{map}}(v^G) \), we can compute the vector \( A \overline{x} \), also stored at \( \overline{V}^{G \rightarrow \overline{G}}(v^G) \) using \( O(t \rho \sqrt{\overline{n}} \log \overline{n} + D) \) of communication in the CONGEST model, with high probability.

Proof. We first invoke Lemma 4.3 to pass \( \overline{x}^G \) to all of \( S^{G \rightarrow \overline{G}}(V^G) \). Then in \( O(\rho) \) round of distributed communication, we can pass these entries (multiplied by the weights of \( A \)) to the corresponding row index. That is, if \( E^{G \rightarrow \overline{G}}_{\text{map}}(u^G, v^G) = u^H v^H \), we pass

\[
A_{u^G, v^G} \overline{x}_{u^G}
\]

from \( v^H \) to \( u^H \). Running Lemma 4.3 again to sum together the passed values over each super vertex then brings the values to the root vertex. \( \square \)

Another implication is that that the Koutis-Xu distributed sparsification algorithm can also be simulated on \( \overline{G} \) in the CONGEST model, with a round overhead of \( \tilde{O}((\sqrt{\overline{n}} + D)) \) \cite{KX16}.

Corollary 4.5. There is an algorithm, \texttt{SpectralSparsifyKX}, that for a graph \( G \) that \( \rho \)-minor distributes into \( \overline{G} \), and some error \( 0 < \epsilon < 0.1 \), \texttt{SpectralSparsifyKX}(\( G, \overline{G}, \epsilon \)) with high probability returns in

\[
O\left(\left(\rho \sqrt{\overline{n}} \log \overline{n} + D\right) \log^8 \overline{n} / \epsilon^2\right)
\]

rounds, a graph \( \overline{G} \), distributed as a \( \rho \)-minor in \( \overline{G} \) such that:

1. \( \overline{G} \) is a (reweighted) subgraph of \( G \),

2. \( \mathbf{L}(G) \approx_{\epsilon} \mathbf{L}(\overline{G}) \).

Proof. The algorithm by \cite{KX16} is based on repeated spanner computations on subgraphs (which are obtained by removing edges from previous spanner computations and uniform sampling of edges). The spanner algorithm of \cite{BS07}, internally used in \cite{KX16}, iteratively grows clusters—organized as spanning trees rooted at center nodes—and adds edges to the spanner. In each iteration some of the existing clusters first are sampled at random, which is done by the respective center node who then forwards the information whether the cluster is sampled to all nodes in its cluster. Then each node decides whether it joins a cluster and if so which one and also decides which of its neighboring edges it adds to the spanner. These decisions are made by comparing the weights of its incident edges. Thus, all the operations performed by nodes in the algorithm of \cite{BS07}, and thus the algorithm of \cite{KX16} fit the description of operations supported by Lemma 4.3. \( \square \)
We will call this sparsification routine regularly, often as preprocessing. This is partly because subgraphs 1-minor distributes into itself trivially.

The minor property also compose naturally: a minor of a minor of \( G \) is also a minor of \( G \). This holds with \( \rho \)-minors too, up to multiplications of the congestion parameters. We prove the following general composition result in Appendix B.

**Lemma 4.6.** Given graphs \( G_1 \) and \( G_2 \) via a \( \rho_2 \)-minor distribution of \( G_2 \) into \( G \), and a \( \rho_1 \)-minor distribution of \( G_1 \) into \( G_2 \) stored on the root vertices of the supervertices of \( G_2 \), and images of \( G_2 \)'s edges in \( G \), we can, with high probability, compute using \( \tilde{O}(\rho_1 \rho_2 \cdot (\sqrt{n} + D)) \) rounds of communication in the CONGEST model a \( \rho_1 \cdot \rho_2 \)-minor distribution of \( G_1 \) into \( G \).

We will always work with congestions in the \( n^{o(1)} \) range: this essentially means we can perform distributed algorithms on \( G \), while paying an overhead of about \( \sqrt{n} + D \) in round complexity to simulate on the original graph.

The composition of minors from Lemma 4.6 implies, among others, that a subset of edges can be quickly contracted.

**Corollary 4.7.** Given \( G \) that’s \( \rho \)-minor distributed on \( G \), along with a subset of edges \( F \subseteq E(G) \) then we can obtain a \( \rho \)-minor distribution of \( G/F \) (\( G \) with \( F \) contracted), into \( G \) in \( \tilde{O}(\rho(\sqrt{n} + D)) \) rounds, under the CONGEST model of computation.

The proof of this requires running \( O(\log n) \) rounds of parallel contraction on the edges of \( F \). We defer it to Appendix B as well.

### 4.2 Laplacian Building Blocks

Some of our algorithm require working with submatrices of Laplacians, which may not be Laplacians anymore, but are still SDD. Fortunately, we can reduce solving an SDD matrix on a graph to solving a Laplacian on a 2-minor. The proof is straightforward and can be found in [Gre96] or [ST14].

**Lemma 4.8** (Greban [Gre96]). Given an \( n \)-by-\( n \) SDD matrix \( M \) that \( \rho \)-minor distributes into \( G \), we can construct a graph \( H \) on \( 2n \) vertices, along with a \( 2 \rho \)-minor distribution of \( H \) into \( G \) with vertices \( i \) and \( n + i \)’s roots mapping to the same root vertex, so that for any vector \( \bar{b} \in \mathbb{R}^n \) and any vectors \( \vec{x} \in \mathbb{R}^{2n} \) such that

\[
\| \vec{x} - L(H)^\dagger \begin{bmatrix} \bar{b} \\ \bar{b} \end{bmatrix} \|_{L(H)} \leq \varepsilon \| \begin{bmatrix} \bar{b} \\ \bar{b} \end{bmatrix} \|_{L(H)^\dagger},
\]

we have

\[
\| \vec{x}_{1:n} - \vec{x}_{n+1:2n} \|_M \leq \varepsilon \| \bar{b} \|_M^\dagger.
\]

In this section we outline the main pieces needed to prove Theorem 1. As described in Section 3, we require three main graph reduction procedures: ultrasparsification (Lemma 4.9), sparsified Cholesky (Lemma 4.10), and minor-based Schur complements (Theorem 3).

The ultrasparsification procedure, based on [ST14, KMP10], allows us to significantly reduce the size of the graph and maintain congestion, but incurs a large approximation error.
Lemma 4.9. There is a routine \textsc{UltraSparsify}(G, k) in the CONGEST model that given a graph G with \( n \) vertices and \( m \) edges, that \( \rho \)-minor distributes into the communication network \( \Gamma \), which has \( \pi \) vertices, \( \overline{m} \) edges, and diameter \( D \), along with a parameter \( k \), produces in \( O(n^{o(1)}(\rho\sqrt{\pi} + D)) \) rounds a graph \( H \) such that:

1. \( H \) is a subgraph of \( G \),
2. \( H \) has at most \( n - 1 + m2^{O(\sqrt{\log n \log \log n})}/k \) edges.
3. \( L(G) \preceq L(H) \preceq kL(G) \).

Furthermore, the algorithm also gives \( \hat{G}, Z_1, Z_2, C \) such that

1. \( \hat{G} \) 1-minor distributes into \( H \) such that \( \hat{G} = \text{SC}(H, C) \) with \( |C| = m2^{O(\sqrt{\log n \log \log n})}/k \).
2. There are operators \( Z_1 \) and \( Z_2 \) evaluable with \( O(\rho\sqrt{\pi} \log \pi + D) \) rounds of CONGEST communication on \( \hat{G} \) such that:

\[
L(H)^\dagger = Z_1^\top \begin{bmatrix} 0 & Z_2 \\ 0 & L(\hat{G})^\dagger \end{bmatrix} Z_1
\]

The elimination procedure, based on [KLP+16], incurs small approximation error, but significantly increases the congestion.

Lemma 4.10. There is a routine \textsc{Eliminate}(G, d, \epsilon) in the CONGEST model that given a graph \( G \) that \( \rho \)-minor distributes into a communication network \( \Gamma \), along with step count \( d \) and error \( \epsilon \), produces in \( O((\epsilon^{-6} \log^{14} n)^d(\rho\sqrt{\pi} \log \pi + D)) \) rounds a subset \( \mathcal{T} \) and access to operators \( Z_1 \) and \( Z_2 \) such that

1. \( |\mathcal{T}| \leq (\frac{4\epsilon}{1-\epsilon})^d |V(G)| \).
2. The cost of applying \( Z_1, Z_1^\top \) and \( Z_2 \) to vectors is \( O((\epsilon^{-6} \log^{14} \pi)^d(\rho\sqrt{\pi} \log \pi + D)) \) rounds of communication on \( \Gamma \).
3. \( L(G)^\dagger \) is \((1 \pm \epsilon)^d\)-approximated by a composed operator built from \( Z_1, Z_2, \) and the inverse of the of the Schur complement of \( L(G) \) onto \( C, \text{SC}(L(G), C) \):

\[
(1 - \epsilon)^d L(G)^\dagger \preceq Z_1^\top \begin{bmatrix} Z_2 \\ 0 \end{bmatrix} \text{SC}(L(G), C)^\dagger \preceq (1 + \epsilon)^d L(G)^\dagger
\]

Note that we cannot directly set \( d = \Omega(\log n) \) to finish with \( |\mathcal{T}| \) a constant: \( \log n \log n \) may be even larger than \( n \). So we need to bring the structure back to a minor of the original communication network. For this we use the construction of spectral vertex sparsifiers that are minors [LS18], modified to not use random spanning trees.

Theorem 3. There is a routine \textsc{ApproxSC}(G, \mathcal{T}, \epsilon) in the CONGEST model that takes a graph on \( n \) vertices, \( m \) edges, that \( \rho \)-minor distributes into the communication network \( \Gamma \), a subset of its vertices \( \mathcal{T} \), error \( \epsilon < 0.1 \), and access to a (distributed) Laplacian Solver \textsc{Solve} returns a graph \( H \), expressed as a distributed \( \rho \)-minor of \( \Gamma \) such that:
1. \( T \subseteq V(H) \),
2. \( H \) has \( O(|T|\epsilon^{-2} \log^2 n) \) edges (and hence at most that many vertices as well).
3. The Schur complements well approximate each other:
   \[
   \text{SC}(G, T) \approx \epsilon \text{SC}(H, T).
   \]

The cost of this computation consists of:
1. \( O(\epsilon^{-3} \log^{10} n) \) calls to \texttt{Solve} to accuracy \( 1/\text{poly}(n) \) on graphs that \( 2\rho \)-distribute into \( \overline{G} \).
2. An overhead of \( O(\rho(n^{1/2} + D)\epsilon^{-3} \log^{11} n) \) rounds.

### 4.3 Schur complement chains and a proof of Theorem 1

In this section, we formally show how to combine Lemma 4.10, Lemma 4.9, and Theorem 3 to efficiently construct a Schur complement chain and prove Theorem 1.

**Definition 4.11.** For a graph \( G \) of \( n \) vertices, \( \{(G_i, Z_{i,1}, Z_{i,2}, T_i)\}_{i=1}^t \) is a \((\gamma, \epsilon)\)-Schur-complement solver chain of \( G \) if the following conditions hold.

1. \( G_1 = G \).
2. \[
   (1 - \epsilon) L(G_i) \preceq Z_{i,1}^\top \begin{bmatrix} Z_{i,2} & 0 \\ 0 & \text{SC}(L(G_i), T_i)^\dagger \end{bmatrix} Z_{i,1} \preceq (1 + \epsilon) L(G_i)^\dagger
   \]
3. \( T_i \subset V(G_{i+1}) \subset V(G_i) \) and \( \text{SC}(G_i, T_i) \approx \epsilon \text{SC}(G_{i+1}, T_i) \)
4. \( |V(G_i)| \geq \gamma \cdot |V(G_{i+1})| \) if \( i < t \), and \( |V(G_t)| \leq \gamma \).

**Lemma 4.12.** Let \( \overline{G} \) be a communication network with \( \overline{n} \) vertices and \( \overline{m} \) edges. Let \( \{(G_i, Z_{i,1}, Z_{i,2}, T_i)\}_{i=1}^t \) be a \((\gamma, \epsilon)\)-Schur-complement solver chain of graph \( G \) for some \( \gamma \geq 2 \) and \( \epsilon \leq \frac{1}{C \log n} \) for large constant \( C \), satisfying the following conditions:

1. \( G_i \rho \)-minor distributes into \( \overline{G} \).
2. Linear operators \( Z_{i,1} \) and \( Z_{i,2} \) can be evaluated in \( O(\overline{n}^{\Theta(1)}(\overline{n}^{1/2} + D)) \) rounds.

Then for a given vector \( \overline{b} \), Algorithm \texttt{PSEUDOINVERSEMULTI} computes a vector \( \overline{x} \) in \( O(\rho \overline{n}^{\Theta(1)}(\overline{n}^{1/2} + D)) \) rounds such that
   \[
   \|\overline{x} - L(G)^\dagger \overline{b}\|_{L(G)} \leq 100\epsilon \cdot \log n \cdot \|\overline{b}\|_{L(G)^\dagger}.
   \]

The correctness proof rely heavily on the following conversion from operator guarantees to error guarantees.
Algorithm 2: Pseudoinverse multiplication

```plaintext
1  procedure PSEUDOINVERSEMULTI(\{(G_i, Z_{i,1}, Z_{i,2}, T_i)\}_{i=1}^t, \vec{b})
2      Create vector \vec{u} = Z_{1,1}\vec{b} with per-coordinate decomposition:
3          \vec{u}_{[V(G_1) \setminus T_i]} \leftarrow (Z_{1,1}\vec{b})_{[V(G_1) \setminus T_i]},
4          \vec{u}_{[T_i]} \leftarrow (Z_{1,1}\vec{b})_{[T_i]}.
5      if \( t = 1 \) then
6          solve \( L(\text{SC}(G_1, T_1))\vec{v} = \vec{u}_{[T_1]} \) exactly.
7      else
8          \vec{v} \leftarrow PSEUDOINVERSEMULTI(\{(G_i, Z_{i,1}, Z_{i,2}, T_i)\}_{i=2}^t, [\vec{u}_{[V(G_1) \setminus T_i]}, \vec{u}_{[T_i]}])
9      return \( Z_{1,1}^\dagger \begin{bmatrix} Z_{1,2}\vec{u}_{[V(G_1) \setminus T_i]} \\ \vec{v}_{[T_i]} - \langle \vec{v}_{[T_i]}, \vec{1}_{[T_i]} \rangle \vec{1}_{[T_i]} \end{bmatrix} \).
```

Lemma 4.13. (Lemma 1.6.7 of [Pen13]) If \( A \) and \( B \) are symmetric PSD matrices such that \( A \approx_\delta B \) for some \( 0 < \delta < 1 \), then for any vector \( \vec{b} \), we have

\[
\left\| A^{\dagger} \vec{b} - B^{\dagger} \vec{b} \right\|_A \leq 10\delta \left\| \vec{b} \right\|_A^{\dagger}.
\]

Proof of Lemma 4.12. We first show correctness. We prove by induction on \( t \) that for a \( t \)-level \((\gamma, \epsilon)\)-Schur-complement solver chain, the error is at most \( 100t\epsilon \left\| \vec{b} \right\|_{L(G_1)^\dagger} \): the result then follows from the bound \( t \leq \log n \).

To simplify notation, let \( M \) denote the middle matrix consisting of \( Z_{1,2} \) and the Schur complement of \( L(G_1) \) onto \( T_i \):

\[
M \overset{\text{def}}{=} \begin{bmatrix} Z_{1,2} & 0 \\ 0 & \text{SC}(L(G_1), T_i)^\dagger \end{bmatrix}.
\]

Then Condition 2 and Lemma 4.13 gives

\[
\left\| Z_{1,1}^\dagger M Z_{1,1} \vec{b} - L(G_1)^\dagger \vec{b} \right\|_{L(G_1)} \leq 10 \left\| \vec{b} \right\|_{L(G_1)^\dagger}.
\]

So by triangle inequality, and the definition of \( \vec{u} = Z_{1,1}\vec{b} \) on Line 2 of Algorithm 2 (PSEUDOINVERSEMULTI), it suffices to bound

\[
\left\| Z_{1,1}^\dagger M Z_{1,1} \vec{b} - Z_{1,1}^\top \begin{bmatrix} Z_{1,2}\vec{u}_{[V(G_1) \setminus T_i]} \\ \vec{v}_{[T_i]} - \langle \vec{v}_{[T_i]}, \vec{1}_{[T_i]} \rangle \vec{1}_{[T_i]} \end{bmatrix} \right\|_{L(G_1)}.
\]

Furthermore, inverting Condition 2 gives

\[
Z_{1,1} L(G_1)^\dagger Z_{1,1}^\top \approx_2 M^\dagger
\]
so up to a constant factor of 2, we can bound the solve cost error involving $\vec{u}$, in the $M$ norm, that is

$$\left\| M \vec{u} - \begin{bmatrix} \vec{Z}_{1,2} \vec{u}_{[V(G_1) \setminus T_1]} \\ \vec{v}_{[T_1]} - \langle \vec{v}_{[T_1]}, \vec{I}_{[T_1]} \rangle \vec{I}_{[T_1]} \end{bmatrix} \right\|_M$$

relative to $\| \vec{u} \|_M$.

Now incorporating the block structure of $M$ means we can consider errors in its two diagonal block separately. Specifically, the top left block is solved exactly, so we can disregard error in it, and bound

$$\left\| \text{SC} \left( L(G_1), T_1 \right)^\dagger \vec{u}_{[T_1]} - \vec{v}_{[T_1]} - \langle \vec{v}_{[T_1]}, \vec{I}_{[T_1]} \rangle \vec{I}_{[T_1]} \right\|_{\text{SC}(L(G_1), T_1)}$$

relative to

$$\| \vec{u}_{[T_1]} \|_{\text{SC}(L(G_1), T_1)}$$

The added all 1s vector is in the null space of $\text{SC}(L(G_1), T_1)$, so does not affect the norm. The Schur complement approximation given in Condition 3 also gives

$$\| \vec{u}_{[T_1]} \|_{\text{SC}(L(G_1), T_1)} \approx \epsilon \| \vec{u}_{[T_1]} \|_{\text{SC}(L(G_2), T_1)}$$

and

$$\left\| \text{SC} \left( L(G_1), T_1 \right)^\dagger \vec{u}_{[T_1]} - \text{SC} \left( L(G_2), T_1 \right)^\dagger \vec{u}_{[T_1]} \right\|_{\text{SC}(L(G_2), T_1)} \leq 10 \epsilon \| \vec{u}_{[T_1]} \|_{\text{SC}(L(G_2), T_1)}$$

Thus, by triangle inequality, it remains to bound the error between $\vec{v}_{[T_1]}$ and $\text{SC} \left( L(G_2), T_1 \right)^\dagger \vec{u}_{[T_1]}$.

Here $\vec{v}$ is obtained from the recursive call on Line 5 of Algorithm 2 (PSEUDOMATIC). So by the inductive hypothesis, we have

$$\left\| \vec{v} - L(G_2)^\dagger \begin{bmatrix} \vec{0} \\ \vec{u}_{[T_1]} \end{bmatrix} \right\|_{L(G_2)} \leq 100 (t - 1) \epsilon \left\| \begin{bmatrix} \vec{0} \\ \vec{u}_{[T_1]} \end{bmatrix} \right\|_{L(G_2)^\dagger}$$

We relate these conditions back to the error of Schur complements via the representation of the Schur complement as the minor of inverse as stated in Lemma 2.3. For the LHS we have:

$$\left( L(G_2)^\dagger \begin{bmatrix} \vec{0} \\ \vec{u}_{[T_1]} \end{bmatrix} \right)_{[T_1]} = \text{SC} \left( L(G_2), T_1 \right)^\dagger \vec{u}_{[T_1]}$$

which by the energy minimization extension definition of Schur complements from Lemma 2.4 gives:

$$\left\| \vec{v}_{[T_1]} - \text{SC} \left( L(G_2), T_1 \right)^\dagger \vec{u}_{[T_1]} \right\|_{\text{SC}(L(G_2), T_1)}$$

$$= \left\| \vec{v}_{[T_1]} - L(G_2)^\dagger \begin{bmatrix} \vec{0} \\ \vec{u}_{[T_1]} \end{bmatrix} \right\|_{\text{SC}(L(G_2), T_1)}$$

$$= \min_{\vec{y} \in \mathbb{R}^{V(G_2) \setminus T_1}} \left\| \begin{bmatrix} \vec{y} \\ \vec{v} - L(G_2)^\dagger \begin{bmatrix} \vec{0} \\ \vec{u}_{[T_1]} \end{bmatrix} \end{bmatrix} \right\|_{L(G_2)}$$

$$\leq \left\| \vec{v} - L(G_2)^\dagger \begin{bmatrix} \vec{0} \\ \vec{u}_{[T_1]} \end{bmatrix} \right\|_{L(G_2)} \leq 100 (t - 1) \epsilon \left\| \begin{bmatrix} \vec{0} \\ \vec{u}_{[T_1]} \end{bmatrix} \right\|_{L(G_2)^\dagger}.$$
Here the second to last inequality follows from setting $\tilde{y}$ to be the $V(G_2) \setminus T_1$ entries of the difference. For the RHS we have:

$$100(t-1)\epsilon \left\| \begin{bmatrix} \vec{0} \\ \bar{u}_{[T_1]} \end{bmatrix} \right\|_{L(G_2)} \leq 100(t-1)\epsilon \left\| \bar{u}_{[T_1]} \right\|_{SC(L(G_1), T_1)} \leq (100(t-1) + 10)\epsilon \left\| \bar{u}_{[T_1]} \right\|_{SC(L(G_1), T_1)}$$

where the last inequality follows from the assumption of $\epsilon < \frac{1}{100}$. So the inductive hypothesis holds for $t$ as well.

**Algorithm 3: Distributed Laplacian Solver**

```plaintext
1 procedure Solve(G)
2   $G' \leftarrow$ SpectralSparsifyKX(G)
3   $(G_1, Z_{1,1}, Z_{1,2}, T_1, G_2) \leftarrow$ UltraSparsify($G'$, k)
4   $\{(G_i, Z_{i,1}, Z_{i,2}, T_i)\}_{i=2}^l \leftarrow$ BuildChain($G_2, d, k$)
5   solve $L(G)\vec{x} = \vec{b}$ by preconditioned Chebyshev with $G_1$ as preconditioner s.t. $L(G_1)\vec{y} = \vec{c}$

6 procedure BuildChain(G, d, k)
7   if $|V(G)| \leq k$ then
8      return ;
9   $(Z_1, Z_2, C) \leftarrow$ Eliminate($G, d, \epsilon$).
10  $H \leftarrow$ ApproxSC($G, C, \epsilon$)
11  return $(G, Z_1, Z_1, C) \cup$ BuildChain($H, d, k$)
```

Proof. (of Theorem 1) The parameters are set as follow:

- $\epsilon = \left(\frac{1}{\log n}\right)^{10}$
- $d = (\log \log \pi)^2$
- $k = 2(\log \pi)^{2/3}$

The correctness of the algorithm is obtained by Lemma 4.3, Lemma 4.9, Lemma 4.10, Theorem 3 and Lemma 4.12.

Now we bound the number of rounds required. By Lemma 4.10, Theorem 3 we have that the Schur-complement chain obtained for graph $G$ satisfying the following conditions

$$|V(G)| = |V(G_1)| \geq |V(G_2)| \geq 2^O(\sqrt{\log n \log \log n})/k = |V(G_2)|/k^{1-o(1)}$$

and

$$|V(G_i)|/|V(G_{i+1})| \geq \epsilon^{-2} \log^2 n/0.99d = 2^{O((\log \log \pi)^2)}.$$
Hence, the Schur-complement chain obtained by the BUILDCHAIN algorithm is a \((2^{Θ((\log \log n)^2)}, \epsilon)\)-Schur-complement chain of length \(O(\log \pi / (\log \log n)^2)\). By Lemma 4.12, PSEUDOINVERSEMULTI for the Schur-complement chain takes \(O(\rho \pi^{(1)} (\pi^{1/2} + D))\) rounds.

Let \(f(n, \rho)\) denote the number of rounds required by Algorithm LAPLACIAN on a graph with \(n\) vertices that is \(\rho\)-minor distributes to \(\overline{G}\), and let \(g(n, \rho)\) denote the number of rounds of BUILDCHAIN with \(n\) vertices that is \(\rho\)-minor distributes to \(\overline{G}\).

Since preconditioned Chebyshev needs to call the Laplacian solver of the preconditioner \(O(\sqrt{k})\) times, by Lemma 4.3, Lemma 4.9, Lemma 4.10, Theorem 3, and Lemma 4.12 we have

\[
f(n, \rho) = O \left( (\log^{14} n \log^{60} \overline{n}) (\log \pi)^2 \left( \rho \pi^{1/2} \log \overline{n} + D \right) \right) + g \left( n/k^{1-o(1)}, \rho \right) + O \left( \sqrt{k} \rho \pi^{o(1)} (\pi^{1/2} + D) \right)
\]

\[
= O \left( \pi^{o(1)} (\rho \pi^{1/2} \log \overline{n} + D) \right) + g \left( n/k^{1-o(1)}, \rho \right).
\]

and

\[
g(n, \rho) = O \left( (\log^{14} n \log^{60} \overline{n}) (\log \pi)^2 \left( \rho \pi^{1/2} \log \overline{n} + D \right) \right) + f(n, 2\rho) \log^{10} n \cdot \epsilon^{-3} + g \left( n/k, 2\rho \right)
\]

\[
= O \left( \pi^{o(1)} (\rho \pi^{1/2} + D) \right) + \text{polylog} (\pi) f(n, 2\rho) + g \left( n/k, 2\rho \right).
\]

Since the depth of the recursion is \(O(\log \pi / (\log \log n)^2)\), the overall increase in congestion is at most

\[2^{O((\log \pi / (\log \log n)^2))} \rho \leq n^{o(1)}\]

so all the graphs constructed \(\pi^{o(1)} \rho\)-minor distribute into \(\overline{G}\).

Hence, the algorithm SOLVE takes \(\rho \pi^{o(1)} (\pi^{1/2} + D)\) rounds.

\[\square\]

5 Minor Schur Complement

In this section we give our algorithm for constructing minor based approximate Schur Complements. Due to the recursive invocation of this routine and solver constructions in Section 4, we can treat the calls to solvers for SDD or Laplacian matrices as black-boxes. The formal guarantees of our constructions are stated in Theorem 3, which we restate below.

**Theorem 3.** There is a routine APPROXSC\((G, \mathcal{T}, \epsilon)\) in the CONGEST model that takes a graph on \(n\) vertices, \(m\) edges, that \(\rho\)-minor distributes into the communication network \(\overline{G}\), a subset of its vertices \(\mathcal{T}\), error \(\epsilon < 0.1\), and access to a (distributed) Laplacian Solver SOLVE returns a graph \(H\), expressed as a distributed \(\rho\)-minor of \(\overline{G}\) such that:

1. \(\mathcal{T} \subseteq V(H)\),
2. \(H\) has \(O(|\mathcal{T}| \epsilon^{-2} \log^2 n)\) edges (and hence at most that many vertices as well).
3. The Schur complements well approximate each other:

\[
\text{SC} \ (G, \mathcal{T}) \approx \epsilon \ \text{SC} \ (H, \mathcal{T}).
\]
The cost of this computation consists of:

1. \( O(\epsilon^{-3} \log^{10} n) \) calls to \textit{solve} to accuracy \( 1/poly(n) \) on graphs that \( 2\rho \)-distribute into \( \overline{G} \).

2. An overhead of \( O(\rho(\overline{\pi}^{1/2} + D)\epsilon^{-3} \log^{11} \overline{\pi}) \) rounds.

Before we go into details, we first discuss high level connections and differences between our algorithm and that of Li-Schild [LS18].

\textbf{Comparison to [LS18]} Our starting point is the same as [LS18], which notes that randomly contracting an edge with probability equal to its leverage score (and deleting otherwise) is exactly a matrix martingale on the spectral form of the graph. This gives a natural algorithm – iteratively compute leverage scores of edges and sample them until the variance has accumulated, and then we stop. The analysis is then via a matrix martingale concentration inequality.

The main difference between these algorithm is how they obtain nearly-linear time running times. Because the resistances of all edges change as edges get sampled, a fast algorithm need to do better than recomputing the probabilities of all edges, after each edge gets sampled. The Li-Schild algorithm [LS18] addresses this by showing that a random spanning tree has the correct marginals, and uses the fast random spanning tree sampling algorithm [Sch18, ALGV20] to obtain such trees. While there are distributed algorithms for sampling spanning trees from unweighted graphs [DSNPT13], partial states of elimination algorithms, namely Schur complements, are naturally weighted. Furthermore, we are unable to directly extend fast random walk simulations to weighted graphs due to the higher congestions of weighted random walks.

Instead, we devise a parallel version of this algorithm based on sampling large subsets of edges \textit{independently}. We compute a large subset of \textit{steady edges} \( Z \) that are mostly uncorrelated, which we obtain from localization of electrical flows [SRS18]. We then identify such subsets, as well as compute all their effective resistances, using standard sketching methods, which are also highly parallel. By ensuring that the size of these sets is at least \( 1/poly(\log n) \) of the total number of edges, we are able to ensure the rapid convergence of this process.

The remainder of the section is organized as follows:

1. Formalize the notion of steady edges, and describe the overall edge sampling/contraction algorithm Section 5.1.

2. Analyze the approximation guarantees via matrix martingales in Section 5.2.

3. Present the sketching procedure for computing resistances and steady edges in Section 5.3.

In general, we track the cost of our algorithms via three quantities. The first is the number of Laplacian solves to principal minors of \( L(G) \) that we must call, and the second is how many additional rounds of communication between neighbors of \( G \) that are necessary, each of which can be simulated in \( O(\rho \sqrt{\overline{\pi}} \log \overline{\pi} + D) \) rounds in \( \overline{G} \) by Lemma 4.3. Finally, we must also ensure that the local computations on vertices \( v \in V(G) \) are actually simple minimum / sum aggregations, as the vertices \( v \in V(G) \) actually correspond to connected components in \( \overline{G} \). These can also be simulated in \( O(\rho \sqrt{\overline{\pi}} \log \overline{\pi} + D) \) rounds in \( \overline{G} \) by Lemma 4.3. We note that because the computations done to solve Laplacian systems and compute leverage scores, etc. all involve only matrix vector multiplication and sampling Bernoulli/Cauchy random variables, it can all be aggregated in a distributed manner.
Distributed storage conventions. In this section, we work with graphs $G$ which $\rho$-minor distribute into the original graph / communication network $\overline{G}$ and is stored distributedly (Definition 4.1). We work with vectors $\vec{x} \in \mathbb{R}^{V(G)}$. In this case, we assume that for a vertex $v \in V(G)$ (corresponding to a connected component in $\overline{G}$), the root $V_{\text{map}}^G(v) \in V(\overline{G})$ knows the values of $\vec{x}_v$. We also work with edge vectors $\vec{w} \in \mathbb{R}^{E(G)}$, such as edge resistances, or leverage scores. For an edge $(uv) = e \in \mathbb{R}^{E(G)}$, it corresponds to an edge in $G$ with endpoints $u^H, v^H$. We assume that these vertices know the value of $\vec{w}_e$. When an algorithm is said to compute vector or edge vectors, we use it to mean that these conditions are satisfied.

5.1 Sparsification Algorithm

We start by defining the key notion of steady edges, which are edges that intuitively don’t interact with each other much. Here, we emphasize that these steady edges are stochastic, not deterministic.

Definition 5.1. A stochastic subset of edges $Z \subseteq E$ is $(\alpha, \delta)$-steady if:

1. (Quadratic form) We have that $\mathbb{E}_{Z} \left[ \sum_{e \in Z} r_e^{-1} b_e^\top b_e \right] \leq \alpha L(H)$.
2. (Localization) For all $e \in Z$ we have $\sum_{e \neq f \in Z} \frac{|b_e^\top L(H)^\top b_f|}{\sqrt{r_e r_f}} \leq \delta$.
3. (Variance) For all $e \in Z$ we have that

$$r_e^{-1} b_e^\top L(H)^\top \begin{bmatrix} SC(H, T) & 0 \\ 0 & 0 \end{bmatrix} L(H)^\top b_e \leq \frac{32|T|}{m}.$$ 

Intuitively, the (Quadratic form) constraint guarantees that no edge is picked to be in the steady set with too high of a probability. The (Localization) constraint bounds the “correlation” between edges, by how much flow the electric flow of an edge $e$ puts on the remaining edges in $Z$. Finally, the (Variance) constraint says that the induced leverage score of edge $e$ on the Schur complement is bounded, and allows us to control the variance in our matrix martingale analysis.

The identification of such edges is implicit from the computation of edge-wise influence, and localization of flows, from [SRS18]. We show the following algorithm in Section 5.3.

Lemma 5.2. Given graph $G$ which $\rho$-minor distributes into $\overline{G}$, constant $\delta \in (0, 1)$, and terminals $T \subseteq V(G)$, and access to a distributed Laplacian solver SOLVE, there is an absolute constant $C_{\text{local}}$ such that calling FINDSTEADY ($G, T, \delta$) as shown in Algorithm 5 returns with probability at least $1 - m^{-10}$ a set of at least $\frac{\delta m}{2000C_{\text{local}} \log^2 n}$ edges in expectation which are $(\frac{\delta}{1000C_{\text{local}} \log^2 n}, \delta)$-steady. The cost of this computation consists of:

1. $O(\log^2 n)$ calls to SOLVE($G'$) to $1/\poly(n)$ error on graphs $G'$ that $2\rho$-minor distribute into $\overline{G}$.

An additional $O(\rho(\sqrt{n} + D) \log^2 n)$ rounds of communication on $\overline{G}$.

We describe our sparsification algorithm. The algorithm has a pre-processing step needed to ensure all edges have leverage scores bounded away from 0 or 1. This is done through two simple procedures – SPLIT and UNSPLIT, whose guarantees we state below.

Lemma 5.3 (SPLIT and UNSPLIT, [LS18] Proposition 3.4 and 3.5). Given graph $G$ which $\rho$-minor distributes into $\overline{G}$, and 1.1-approximate leverage scores $\tilde{\lev}_G(e)$ for the edges of $G$, calling SPLIT($G, \tilde{\lev}_G(e)$) returns in $O(\rho(\sqrt{n} + D))$ rounds a graph $H$ that:

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1. is electrically equivalent to $G$,
2. $2\rho$-minor distribute into $\overline{G}$,
3. has all edge leverage scores between $[3/16, 13/16]$.

Algorithm Unsplit$(G)$ takes $G$, and returns in $\tilde{O}(\rho(\sqrt{n} + D))$ rounds a graph resulting from collapsing paths and parallel edges and removing non-terminal leaves, along with a $\rho$-minor distribution into $\overline{G}$.

Proof. The implementation of Unsplit is straightforward: the resulting graph is a minor of $G$, and all changes happen on $O(1)$ sized neighborhoods, and only involve local endpoints of edges of $G$. So they can be implemented using $O(1)$ rounds of communications among neighbors of $G$, and in turn Lemma 4.6.

Algorithm Split proceeds as follows.
1. If $\ell_e < 1/4$, replace $e$ with a path of length 2, each edge with resistance $r_e/2$.
2. If $\ell_e \geq 3/4$ then replace $e$ with two parallel edges of resistance $2r_e$.

In the first case, both edges will have leverage score $\frac{1}{2} + \frac{\text{lev}_G(e)}{2} \in [3/16, 13/16]$, because we know that $\text{lev}_G(e) \leq 1/2$ by the approximation guarantee.

In the second case, both edges will have leverage score $\text{lev}_G(e)/2 \in [3/16, 13/16]$, as $\text{lev}_G(e) \geq 1/2$ by the approximation guarantee. Otherwise, we know $\text{lev}_G(e) \in [3/16, 13/16]$ by the approximation guarantee.

The bound on the cost and embeddability follows from each edge turning into at most a path of 2 vertices. The new vertex can be placed at either end points of $E_{\text{map}}(\overline{G}, e)$, and the congestion on both $e$ and endpoints of $e$ go up by a factor of 2. This 2-minor embedding of the new graph into $G$ then meets the definition of Lemma 4.6, which means we get its $2\rho$-minor distribution in $\overline{G}$ with an overhead of $\tilde{O}(\rho(\sqrt{n} + D))$.

This scheme, as well as our sampling process, depends on the leverage scores of the edges. We obtain such estimates using the now standard random projection scheme devised due to Spielman and Srivastava [SS11], whose guarantees we state below in terms of solves and overhead in CONGEST.

Lemma 5.4 (Leverage score estimate). Given graph $G$ with resistance $r_e$, which $\rho$-minor distributes into $\overline{G}$, and access to a distributed Laplacian solver Solve, there is an algorithm $\text{LevAp}(e, G, \delta)$ which for all edges $e \in E(G)$ approximates the quantity $\text{lev}_G(e) = \overline{r}_e^{-1}\overline{b}_e^\top L(G)^{-1}\overline{b}_e$ to within a factor of $1 + \delta$ with high probability. It requires

1. $O(\delta^{-2} \log n)$ calls to Solve($\cdot$) on graphs that $\rho$-minor distributes into $\overline{G}$, to accuracy $1/poly(n)$
2. An additional $O(\rho\delta^{-2} (\sqrt{n} + D) \log^2 n)$ rounds of communication in $\overline{G}$.

As this algorithm is sketching based, and also a direct distributed implementation of an existing sequential algorithm, we will present it together with Lemma 5.2 in Section 5.3.

We now describe our algorithm for computing a minor Schur complement. First, identify a set of steady edges and approximate compute all their leverage scores via a standard JL sketch (Lemma 5.4). Now, for each steady edge independently, contract it with probability given by the leverage score estimate, and delete otherwise. Now, repeat the process until the size of the remaining graph is small enough.
Algorithm 4: Finding sparsifier of Schur complement onto terminals, but with extra Steiner vertices

```
1 procedure APPROXSC(G, T, ε)
2    G(0) ← G and i ← 0.
3    δ ← \frac{ε}{C \log^2 m}. \quad \triangleright C \text{ is large constant}
4    while |E(G(i))| ≥ C|T| log^2 m do
5       H(i) ← SPLIT(G(i), LEVAPX(e, G, 0.01)). \quad \triangleright \text{Lemmas 5.3 and 5.4}
6       Z(i) ← FINDSTEADY(H(i), T, δ). \quad \triangleright \text{Lemma 5.2}
7       For all e ∈ Z(i), set p_e ← LEVAPX(H(i), e, δ). \quad \triangleright \text{Lemma 5.4}
8       For all e ∈ Z(i), contract e with probability p_e, and delete e with probability 1 − p_e,
9          and let the resulting graph be I(i). Perform the contractions and deletions via
10          Corollary 4.7.
11    G(i+1) ← UNSPLIT(I(i)) and i ← i + 1. \quad \triangleright \text{Lemma 5.3}
```

Return $G(i)$.

**Theorem 4.** Let $G$ be a graph with terminals $T$. Calling APPROXSC($G, T, \epsilon$) produces with probability at least $1 - m^{-5}$ a graph $H$ satisfying $|E(H)| \leq O(|T| \log^2 m/\epsilon^2)$ and SC($H, T$) $\approx \epsilon$ SC($G, T$).

Before proceeding, we first bound the number of phases of APPROXSC as in Algorithm 4.

**Lemma 5.5.** Let $G$ be a graph with terminals $T$. The while loop of APPROXSC($G, T, \epsilon$) as in Algorithm 4 executes at most $O(\alpha^{-1} \log m)$ times, for $\alpha = \frac{\delta}{1000C_{local} \log^2 m} = \frac{\epsilon}{1000C_{local} \log^4 m}$ with probability at least $1 - m^{-6}$.

**Proof.** To show the iteration count bound, it suffices to argue that

$$\mathbb{E}[|E(G^{(i+1)})|] \leq (1 - \Omega(\alpha))|E(G^{(i)})|,$$

because $\alpha = \frac{\epsilon}{1000C_{local} \log^4 m}$ by definition. Note that $\mathbb{E}[|Z^{(i)}|] \geq \alpha |E(G^{(i)})|/2$, so it suffices to argue that each original edge of $G^{(i)}$ is removed with at least constant probability, even considering the SPLIT operation. We break the analysis into cases.

**Case 1: $e$ split into two parallel edges $e_1, e_2$.** Recall that by the definition of SPLIT, both $e_1$ and $e_2$ have leverage score in $[3/16, 13/16]$. Therefore, if say $e_1 \in Z^{(i)}$, then it will be contracted with probability at least $3/16$. In that case both $e_1$ and $e_2$ disappear, as desired.

**Case 2: $e$ split into a path of length two consisting of $e_1, e_2$.** Recall that by the definition of SPLIT, both $e_1$ and $e_2$ have leverage score in $[3/16, 13/16]$. Therefore, if say $e_1 \in Z^{(i)}$, the it will be deleted with probability at least $3/16$. Then $e_2$ becomes a leaf, so it will be removed during the UNSPLIT operation, as desired.

We can combine everything now to show Theorem 3.
Proof of Theorem 3. The algorithm clearly returns a minor, as it only applies deletions and contractions. The correctness and number of edges bound follows from Theorem 4. Therefore, it suffices to bound the computation cost.

By Lemma 5.5, the number of iterations that the main while loop of APPROXSC (Line 4 of Algorithm 4) is \(O(\alpha^{-1}\log m) = O(\epsilon^{-1}\log^5 n)\). By Lemma 5.2, 5.4, 5.3, the cost of each iteration is dominated by line 7, which costs \(O(\delta^{-2}\log n) = O(\epsilon^{-2}\log^5 n)\) linear system solves and rounds of communication.

Therefore, the number of Laplacian system calls is at most

\[
O\left(\epsilon^{-1}\log^5 n \cdot \epsilon^{-2}\log^5 n\right) = O\left(\epsilon^{-3}\log^{10} n\right).
\]

The overhead cost can be bounded in the same way, multiplied by the \(O(\rho\sqrt{n}\log \bar{n} + D)\) overhead to simulate communication on \(G\) in \(\overline{G}\) by Lemma 4.3. \(\square\)

5.2 Matrix Martingale Analysis of Approximation

To show Theorem 4, we will define several stochastic sequences of matrices that capture the change of the quadratic form of the Schur complement. Let \(\tau\) denote the random variable denoting the final value of \(i\) in APPROXSC as in Algorithm 4. Define

\[
S_0 \overset{\text{def}}{=} [SC(G, T)]^{1/2} \cdot 0.
\]

For \(0 \leq i \leq \tau\) and \(0 \leq t < |Z^{(i)}|\) let \(e_{i,t}\) be the \(t\)-th edge in \(Z^{(i)}\), under an arbitrary ordering, and define

\[
\hat{Y}^{(i,0)} := S_0 L(H^{(i)}) \cdot S_0^T
\]

and

\[
\hat{Y}^{(i,t+1)} := \begin{cases} 
\hat{Y}^{(i,t)} + \sqrt{e_{i,t}}(1 - \overline{b}_{e_{i,t}})^{-1} S_0 L(H^{(i)}) \cdot \overline{b}_{e_{i,t}} L(H^{(i)}) \cdot S_0^T & \text{if } e_{i,t} \text{ is deleted,} \\
\hat{Y}^{(i,t)} - \sqrt{e_{i,t}} \overline{p}_{e_{i,t}} S_0 L(H^{(i)}) \cdot \overline{b}_{e_{i,t}} L(H^{(i)}) \cdot S_0^T & \text{if } e_{i,t} \text{ is contracted.}
\end{cases}
\]

Note that the sequence \(\hat{Y}^{(i,t)}\) is a martingale for fixed \(i\), but \(\hat{Y}^{(i+1,0)} \neq \hat{Y}^{(i,|Z^{(i)}|)}\). To capture this, for \(0 \leq i \leq \tau\) and \(0 \leq t < |Z^{(i)}|\) we define

\[
X^{(i,t)} = \begin{cases}
0 & \text{if } t = 0, \\
\hat{Y}^{(i,t)} - \hat{Y}^{(i,t-1)} & \text{if } t > 0.
\end{cases}
\] (1)

Let \(Y^{(i,t)}\) be the sequence with difference sequence \(X^{(i,t)}\), and \(Y^{(0,0)} = \hat{Y}^{(0,0)} = I\). Note that \(E[X^{(i,t)}|X^{(i,t-1)}] = 0\), hence \(Y^{(i,t)}\) is a martingale. To show Theorem 4, it suffices to prove that \(\|\hat{Y}^{(\tau,0)} - \hat{Y}^{(0,0)}\|_2 \leq \epsilon\). We prove this in two pieces, first by analyzing the martingale \(Y^{(i,t)}\), and then by analyzing the errors resulting from \(\hat{Y}^{(i+1,0)} - \hat{Y}^{(i,|Z^{(i)}|)}\). Throughout the analysis, we may assume that \(SC(H^{(i)}, T) \approx_{1,1} SC(G, T)\) for all \(i\), or else we can define the matrix sequences above to terminate whenever this condition is violated.
Lemma 5.6 (Operator bound for $X^{(i,t)}$). Assume that $SC(H^{(i)}, \mathcal{T}) \approx_{1.1} SC(G, \mathcal{T})$ for all $(i, t)$. Then for all $(i, t)$ we have that

$$\|X^{(i,t)}\|_2 \leq \frac{300 |\mathcal{T}|}{|E(H^{(i)})|}.$$ 

Proof. Note that $p_{e_{i,t}} \in [1/8, 7/8]$ by the guarantee that $\text{lev}_{H^{(i)}}(e_{i,t}) \in [3/16, 13/16]$ and the approximation guarantee of Lemma 5.4. Therefore, we have that

$$\|X^{(i,t)}\|_2 \leq \frac{1}{\overline{r}_{e_{i,t}} \min(\overline{p}_{e_{i,t}}, 1 - \overline{p}_{e_{i,t}}) S_0 L(H^{(i)}) \tilde{b}_{e_{i,t}, t} \tilde{b}_{e_{i,t}, t}^\top L(H^{(i)})^\top S_0^\top}) \leq 300 |\mathcal{T}| \frac{|\mathcal{T}|}{|E(H^{(i)})|}.$$ 

by the (Variance) condition of Definition 5.1.

We define the quadratic variation $W^{(i,t)}$ as in Lemma 2.7 for the martingale $Y^{(i,t)}$.

Lemma 5.7 (Variance bound for $Y^{(i,t)}$). Assume that $SC(H^{(i,t)}, \mathcal{T}) \approx_{1.1} SC(G, \mathcal{T})$ for all $(i, t)$. For sufficiently large $C$ in APPROXSC as in Algorithm 4, we have for all $(i, t)$ that

$$\Pr \left[ \|W^{(i,t)}\|_2 \geq \frac{\epsilon^2}{100 \log m} \right] \leq m^{-10}.$$ 

Proof. It suffices to argue that

$$\Pr \left[ \|W^{(\tau,0)}\|_2 \geq \frac{\epsilon^2}{100 \log m} \right] \leq m^{-10}.$$ 

Note that

$$W^{(i+1,0)} - W^{(i,0)} = \sum_{1 \leq t \leq |Z^{(i)}|} \mathbb{E} \left[ \left( X^{(i,t)} \right)^2 \mid X^{(i,t-1)} \right] \leq \|X^{(i,t)}\|_2 \sum_{1 \leq t \leq |Z^{(i)}|} \frac{1}{\overline{r}_{e_{i,t}} \min(\overline{p}_{e_{i,t}}, 1 - \overline{p}_{e_{i,t}}) S_0 L(H^{(i)}) \tilde{b}_{e_{i,t}, t} \tilde{b}_{e_{i,t}, t}^\top L(H^{(i)})^\top S_0^\top} \leq \frac{300 |\mathcal{T}|}{|E(H^{(i)})|} \sum_{1 \leq t \leq |Z^{(i)}|} \frac{1}{\overline{r}_{e_{i,t}} \min(\overline{p}_{e_{i,t}}, 1 - \overline{p}_{e_{i,t}}) S_0 L(H^{(i)}) \tilde{b}_{e_{i,t}, t} \tilde{b}_{e_{i,t}, t}^\top L(H^{(i)})^\top S_0^\top} \leq \frac{2400 |\mathcal{T}|}{|E(H^{(i)})|} S_0 L(H^{(i)})^\top \left( \sum_{f \in Z^{(i)}} \overline{r}_f \overline{b}_f \overline{b}_f^\top \right) L(H^{(i)})^\top S_0^\top.$$ 

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Therefore, the matrix process $U^{(i)} \overset{\text{def}}{=} W^{(i+1,0)} - W^{(i,0)}$ is an adapted process of positive semidefinite matrices. We know that
\[
\|U^{(i)}\|_2 \leq \frac{2400 |T|}{E(H^{(i)})} \left\| \begin{array}{c}
S_0L(H^{(i)}) \left( \sum_{f \in Z^{(i)}} \tilde{r}_f \tilde{b}_f \tilde{b}_f^\top \right) L(H^{(i)})^\top S_0^\top \end{array} \right\|_2
\]
\[
\leq \frac{2400 |T|}{E(H^{(i)})} \left\| S_0L(H^{(i)}) S_0^\top \right\|_2 \leq \frac{3000 |T|}{E(H^{(i)})}
\]
with probability 1, and
\[
\left\| E_{Z^{(i)}} \left[ U^{(i)} \mid U^{(i-1)} \right] \right\|_2
\]
\[
\leq \frac{2400 |T|}{E(H^{(i)})} \left\| S_0L(H^{(i)}) \mid E_{Z^{(i)}} \left[ \sum_{f \in Z^{(i)}} \tilde{r}_f \tilde{b}_f \tilde{b}_f^\top \right] L(H^{(i)})^\top S_0^\top \right\|_2 \leq \frac{3000 \alpha |T|}{E(H^{(i)})}
\]
by the (Quadratic form) condition of Definition 5.1.

For $1 \leq i \leq \tau$ define
\[
V^{(i)} \overset{\text{def}}{=} U^{(i)} - E \left[ U^{(i)} \mid U^{(i-1)} \right],
\]
so that $E[V^{(i)} \mid V^{(i-1)}] = 0$. Let $\hat{W}^{(i)}$ be the martingale with difference sequence given by $V^{(i)}$, and $\hat{W}^{(0)} = W^{(0)} = 0$. Note that by Equation (2)
\[
\|V^{(i)}\|_2 \leq 2 \|U^{(i)}\|_2 \leq \frac{6000 |T|}{E(H^{(i)})}
\]
and by Equation (3)
\[
\left\| E \left[ (V^{(i)})^2 \mid V^{(i-1)} \right] \right\|_2 \leq \left\| E \left[ (U^{(i)})^2 \mid U^{(i-1)} \right] \right\|_2 \leq \frac{3000 |T|}{E(H^{(i)})} \left\| E \left[ U^{(i)} \mid U^{(i-1)} \right] \right\|_2 \leq \frac{10^7 |T|^2 \alpha}{E(H^{(i)})^2}.
\]

By Lemma 5.5 we know that $\tau \leq O(\alpha^{-1} \log m)$, so let $C_{5.5}$ be the universal constant such that $\tau \leq C_{5.5} \alpha^{-1} \log m$ with high probability. Therefore, to analyze the process $\hat{W}^{(i)}$ using Lemma 2.7, we can set
\[
R = \frac{6000 |T|}{C |T| \log^2 m/e^2} \leq \frac{e^2}{5000 \log^2 m}
\]
and
\[
\sigma^2 = \frac{10^7 |T|^2 \alpha}{C^2 |T|^4 \log^4 m/e^4} \cdot C_{5.5} \alpha^{-1} \log m \leq \frac{e^4}{10^6 \log^4 m}
\]
for sufficiently large $C$. Therefore, Lemma 2.7 gives us that
\[
\Pr \left[ \left\| \hat{W}^\tau \right\|_2 \geq \frac{e^2}{200 \log^2 m} \right] \leq |T| \cdot \exp \left( \frac{e^4}{10^6 \log^4 m + \frac{e^4}{10^6 \log^4 m}} \right) \leq m^{-10}.
\]
To finish, note that
\[ \left\| W^{(r)} \right\|_2 \leq \left\| \tilde{W}^{(r)} \right\|_2 + \sum_{1 \leq i \leq \tau} \left\| \mathbb{E} \left[ U^{(i)} \right| U^{(i-1)} \right\|_2 \]
\[ \leq \frac{\epsilon^2}{200 \log m} + \frac{3000 \tau \alpha |T|}{|E(H^{(i)})|} \leq \frac{\epsilon^2}{200 \log m} + \frac{3000 C_{5,5} \epsilon^2}{C \log m} \leq \frac{\epsilon^2}{100 \log m} \]
for sufficiently large $C$.

Combining these gives us a bound on $Y^{(i,t)}$.

**Lemma 5.8 (Bound on $Y^{(i,t)}$).** With probability at least $1 - m^{-6}$ we have that \( \left\| Y^{(i,t)} - I \right\|_2 \leq \epsilon/2 \) for all $(i, t)$.

**Proof.** We can directly plug in Lemmas 5.6 and 5.7 into Lemma 2.7. In particular, we may set $R = \frac{300 |T|}{|E(H^{(i)})|} \leq \frac{\epsilon^2}{100 \log m}$ for sufficiently large $C$, and $\sigma^2 = \frac{\epsilon^2}{100 \log m}$. Therefore, as $Y^{(0,0)} = I$, we have that
\[
\Pr \left[ \exists (i, t) : \left\| Y^{(i,t)} - Y^{(0,0)} \right\|_2 \geq \epsilon/2 \text{ and } \left\| W^{(i,t)} \right\|_2 \leq \sigma^2 \right] \leq 2 |T| \cdot \exp \left( \frac{-\epsilon^2/12}{\sigma^2 + R \epsilon/6} \right) \leq 2 |T| \exp \left( \frac{-\epsilon^2/12}{\frac{\epsilon^2}{100 \log m} + \frac{\epsilon^2}{600 \log m}} \right) \leq m^{-7}.
\]
To finish, we can union bound this along with Lemma 5.7.

Define $\tilde{X}^{(i)} = \tilde{Y}^{(i+1,0)} - \tilde{Y}^{(i,|Z^{(i)}|)}$, so that
\[ \tilde{Y}^{(\tau,0)} = Y^{(\tau,0)} + \sum_{0 \leq i < \tau} \tilde{X}^{(i)}.
\]
We will apply the Woodbury matrix formula (Lemma 2.6) to calculate $\tilde{Y}^{(i+1,0)}$. We choose
\[ A = L \left( H^{(i)} \right), \quad U = B_{Z^{(i)}}^\top R_{Z^{(i)}}^{-1/2}, \quad V = U^\top,
\]
where $B_{Z^{(i)}}$ and $R_{Z^{(i)}}$ are the restriction of $B$ and $R$ to the rows corresponding to the set $Z^{(i)}$. Also, set $C^{(i)}$ as the following diagonal matrix:
\[ C^{(i)}_{ff} = \begin{cases} -1 & \text{if } f \in Z^{(i)} \text{ is contracted, and} \\ +\infty & \text{otherwise.} \end{cases}
\]
Note that then $L(I^{(i)}) = A + U C^{(i)} V$. Therefore, Lemma 2.6 gives us that
\[
\tilde{Y}^{(i+1,0)} - \tilde{Y}^{(i,0)} = -S_0 L \left( H^{(i)} \right)^\top B_{Z^{(i)}}^\top R_{Z^{(i)}}^{-1/2}\left( \left( C^{(i)} \right)^{-1} + R_{Z^{(i)}}^{-1/2} B_{Z^{(i)}} L \left( H^{(i)} \right)^\top B_{Z^{(i)}}^\top R_{Z^{(i)}}^{-1/2} \right)^{-1} R_{Z^{(i)}}^{-1/2} B_{Z^{(i)}} L \left( H^{(i)} \right)^\top S_0^\top.
\]
(4)
Define the diagonal matrix $P^{(i)}$ as follows.

$$P^{(i)}_{ff} = \begin{cases} \frac{1}{1 - \bar{\mu}_f} & \text{if } f \text{ was deleted, and} \\
\frac{1}{1 - \bar{\mu}_f} & \text{otherwise.} \end{cases}$$

**Lemma 5.9.** We have that

$$\left\| P^{(i)} + \left( \left( C^{(i)} \right)^{-1} + R_{Z^{(i)}}^{-1/2} B_{Z^{(i)}} L (H^{(i)})^\dagger B_{Z^{(i)}}^\top R_{Z^{(i)}}^{-1/2} \right) \right\|_2 \leq 40\delta.$$

**Proof.** Let diagonal matrix $D^{(i)}$ be the diagonal entries of the matrix

$$(C^{(i)})^{-1} + R_{Z^{(i)}}^{-1/2} B_{Z^{(i)}} L (H^{(i)})^\dagger B_{Z^{(i)}}^\top R_{Z^{(i)}}^{-1/2}.$$ That is

$$D^{(i)}_{ff} = \begin{cases} -1 + \text{lev}_{H^{(i)}} (f) & \text{if } f \text{ is deleted, and} \\
\text{lev}_{H^{(i)}} (f) & \text{otherwise.} \end{cases}$$

Define

$$Q^{(i)} = D^{(i)} - \left( \left( C^{(i)} \right)^{-1} + R_{Z^{(i)}}^{-1/2} B_{Z^{(i)}} L (H^{(i)})^\dagger B_{Z^{(i)}}^\top R_{Z^{(i)}}^{-1/2} \right).$$

Note that $Q^{(i)}$ is 0 on the diagonal, and for any $f \in Z^{(i)}$, the sum of off-diagonal entries in the row corresponding to $f$ is at most $\delta$:

$$\sum_{f \neq g \in Z^{(i)}} \left| Q^{(i)}_{fg} \right| \leq \sum_{f \neq g \in Z^{(i)}} \frac{\left| \hat{b}_f \right| L (H^{(i)})^\dagger \hat{b}_g}{\sqrt{\nu_f} \sqrt{\nu_g}} \leq \delta$$

by the (Localization) condition of Definition 5.1 used in Line 6 of APPROXSC (given in Algorithm 4). Therefore, Lemma 2.8 gives that $\| Q^{(i)} \|_2 \leq \delta$. Therefore, we have that

$$\left\| P^{(i)} + \left( \left( C^{(i)} \right)^{-1} + R_{Z^{(i)}}^{-1/2} B_{Z^{(i)}} L (H^{(i)})^\dagger B_{Z^{(i)}}^\top R_{Z^{(i)}}^{-1/2} \right) \right\|_2 \leq \left\| P^{(i)} \right\|_2 + \left\| D^{(i)} - Q^{(i)} \right\|_2 + \left\| \left( D^{(i)} - Q^{(i)} \right)^{-1} - \left( D^{(i)} \right)^{-1} \right\|_2.$$

The approximation guarantee in line 7 of APPROXSC as in Algorithm 4 and the fact that $p_e \in [1/8, 7/8]$ gives us

$$\left\| P^{(i)} + \left( D^{(i)} \right)^{-1} \right\|_2 \leq 10\delta.$$

For the second term, we use again that all leverage scores of $H^{(i)}$ are in $[3/16, 13/16]$ to get

$$\left\| \left( D^{(i)} - Q^{(i)} \right)^{-1} - \left( D^{(i)} \right)^{-1} \right\|_2 \leq \frac{16}{3} \delta \cdot \left( \frac{3}{16} - \delta \right)^{-1} \leq 30\delta.$$

Combining these bounds gives the desired result. □
We use this to give bounds on $\hat{X}^{(i)}$.

**Lemma 5.10.** Assume that $SC(H^{(i,t)}, T) \approx_{1,1} SC(G, T)$ for all $(i, t)$. Then for all $0 \leq i < \tau$ we have that

$$
\left\| \hat{X}^{(i)} \right\|_2 \leq 50\delta \quad \text{and} \quad \left\| E_{Z(i)} \left[ \left( \hat{X}^{(i)} \right)^2 \right] \right\|_2 \leq 2500\alpha\delta^2.
$$

**Proof.** Note that

$$
Y^{(i, |Z^{(i)}|)} = Y^{(i, 0)} + S_0 L \left( H^{(i)} \right)^\dagger B_{Z(i)}^\top R_{Z(i)}^{-1/2} P(i) R_{Z(i)}^{-1/2} B_{Z(i)} L \left( H^{(i)} \right)^\dagger S_0^\top.
$$

Combining this with Equation (4) and Lemma 5.9, gives that for the matrix:

$$
M^{(i)} \overset{\text{def}}{=} \left( (C^{(i)})^{-1} + R_{Z(i)}^{-1/2} B_{Z(i)} L \left( H^{(i)} \right)^\dagger B_{Z(i)}^\top R_{Z(i)}^{-1/2} \right)^{-1}.
$$

we have:

$$
\left\| \hat{X}^{(i)} \right\|_2 = \left\| S_0 L \left( H^{(i)} \right)^\dagger B_{Z(i)}^\top R_{Z(i)}^{-1/2} \left( P(i) + M^{(i)} \right) R_{Z(i)}^{-1/2} B_{Z(i)} L \left( H^{(i)} \right)^\dagger S_0^\top \right\|_2 \\
\leq 40\delta \left\| S_0 L \left( H^{(i)} \right)^\dagger B_{Z(i)}^\top R_{Z(i)}^{-1/2} B_{Z(i)} L \left( H^{(i)} \right)^\dagger S_0^\top \right\|_2 \leq 40\delta \left\| S_0 L \left( H^{(i)} \right)^\dagger S_0^\top \right\|_2 \leq 50\delta.
$$

Similarly, we have by (4), the above, Lemma 5.9, and the (Quadratic form) condition of Definition 5.1 that

$$
\left\| E_{Z(i)} \left[ \left( \hat{X}^{(i)} \right)^2 \right] \right\|_2 \leq \left\| \hat{X}^{(i)} \right\|_2 \cdot \left\| P(i) + M^{(i)} \right\|_2 \\
\quad \cdot \left\| E_{Z(i)} \left[ S_0 L \left( H^{(i)} \right)^\dagger B_{Z(i)}^\top R_{Z(i)}^{-1/2} B_{Z(i)} L \left( H^{(i)} \right)^\dagger S_0^\top \right] \right\|_2 \\
\leq 2000\delta^2 \left\| S_0 L \left( H^{(i)} \right)^\dagger E_{Z(i)} \left[ B_{Z(i)}^\top R_{Z(i)}^{-1} B_{Z(i)} \right] L \left( H^{(i)} \right)^\dagger S_0^\top \right\|_2 \\
= 2000\delta^2 \left\| S_0 L \left( H^{(i)} \right)^\dagger E_{Z(i)} \left[ \sum_{j \in Z(i)} \tilde{r}_f^{-1} \tilde{b}_j \tilde{b}_j^\top \right] L \left( H^{(i)} \right)^\dagger S_0^\top \right\|_2 \\
\leq 2000\alpha\delta^2 \left\| S_0 L \left( H^{(i)} \right)^\dagger S_0^\top \right\|_2 \leq 2500\alpha\delta^2.
$$

Finally, we bound $\hat{Y}^{(i, 0)} - Y^{(i, 0)}$.

**Lemma 5.11.** With probability $1 - m^{-6}$, we have that

$$
\left\| \hat{Y}^{(i, 0)} - Y^{(i, 0)} \right\|_2 \leq \epsilon/2
$$

for all $i$. 28
**Proof.** Note that $\tilde{Y}^{(i,0)} - Y^{(i,0)} = \sum_{0 \leq j < i} \tilde{X}^{(j)}$. Define the adapted sequence $U^{(i)} \equiv \tilde{X}^{(i)} - E_{Z^{(i)}}[\tilde{X}^{(i)}]$, so that $E_{Z^{(i)}}[U^{(i)}] = 0$. Let $V^{(i)}$ be the martingale with difference sequence $U^{(i)}$, and $V^{(0)} = 0$. By Lemma 5.10 we know that

$$\|U^{(i)}\|_2 \leq 2\|\tilde{X}^{(i)}\|_2 \leq 100\delta,$$

and

$$\left\|E_{Z^{(i)}}[(U^{(i)})^2]\right\|_2 \leq \left\|E_{Z^{(i)}}[(\tilde{X}^{(i)})^2]\right\|_2 \leq 2500\alpha\delta^2.$$

Now we apply Lemma 2.7 to the martingale $V^{(i)}$. We can set $R = 100\delta$, and because Lemma 5.5 tells us that $\tau \leq C_{5.5}\alpha^{-1} \log m$ with high probability for some universal constant $C_{5.5}$, we can set

$$\sigma^2 \leq 2500\alpha\delta^2 C_{5.5}\alpha^{-1} \log m \leq 2500C_{5.5}\delta^2 \log m.$$

$$\Pr \left[ \|V^{(i)}\|_2 \geq \epsilon/4 \right] \leq |\mathcal{T}| \cdot \exp \left( \frac{-\epsilon^2/48}{2500C_{5.5}\delta^2 \log m + 250\delta/3} \right) \leq m^{-10}$$

for sufficiently large choice of $C$ in APPROXSC as in Algorithm 4. To finish, note by Lemma 5.10

$$\left\|\tilde{Y}^{(i,0)} - Y^{(i,0)}\right\|_2 \leq \|V^{(i)}\|_2 + \sum_{0 \leq j < i} \left\|E_{Z^{(i)}}[\tilde{X}^{(i)}]\right\|_2 \leq \epsilon/4 + 50\alpha\delta \cdot C_{5.5}\alpha^{-1} \log m = \epsilon/4 + 50C_{5.5}\delta \log m \leq \epsilon/2$$

for sufficiently large constant $C$.

We can combine the above to show Theorem 4.

**Proof of Theorem 4.** The size bound follows vacuously. By Lemmas 5.8 and 5.11 we know that

$$\left\|\tilde{Y}^{(\tau,0)} - I\right\|_2 \leq \|I - \epsilon/2 + \epsilon/2 = \epsilon,$$

which implies that $\text{SC}(G^{(\tau)}, \mathcal{T}) \approx_{\epsilon} \text{SC}(G, \mathcal{T})$, as desired. \qed

### 5.3 Sketching and Construction of Steady Edges

Our algorithm requires approximating various quantities via sketches. The following are essentially proven in [LS18], and we more explicitly state the cost of solving the Laplacians and building the sketch matrices in these lemmas.

Our proof of Lemma 5.4 requires the Johnson-Lindenstrauss lemma.

**Lemma 5.12** (Johnson-Lindenstrauss lemma [JL84, Ach01]). Let $\vec{v} \in \mathbb{R}^d$ be a vector, and let $Q \in \mathbb{R}^{t \times d}$ be a random matrix with random $\pm 1$ entries for some $t \geq C\delta^{-2} \log(1/\delta_0)$. Then with probability $1 - \delta_0$ we have that

$$(1 - \delta) \|\vec{v}\|_2^2 \leq t^{-1} \|Q\vec{v}\|_2^2 \leq (1 + \delta) \|\vec{v}\|_2^2.$$
Proof of Lemma 5.4. Note that

$$\text{lev}_G(e) = \frac{1}{r_e} \left\| R^{-1/2} B L(G)^\dagger b_e \right\|_2^2. $$

By Lemma 5.12, for the matrix $Q \in \mathbb{R}^{t \times m}$ chosen with random $\pm 1$ entries for $t = C\delta^{-2} \log n$, we have that with high probability for all $e$

$$\text{lev}_G(e) \approx \delta \frac{t^{-1/2}}{r_e} \left\| Q R^{-1/2} B L(G)^\dagger b_e \right\|_2^2. $$

We first generate $Q$ on the end points of the edges’ images in $\overline{G}$. As $R^{-1/2}$ is a rescaling by the resistances of these edges, which are also stored together with their end points, we can have each edge $e \in E(G)$ store the corresponding column in $QR^{-1/2}$ in both of their end points. That is, both end points of $E_{\text{map}}(e) \rightarrow \overline{G}(e)$ stores $(QR^{-1/2})_{i,e}$.

Computing the matrix $QR^{-1/2}B \in \mathbb{R}^{t \times n}$ is then an aggregation of these entries, distributed over the super node, to its root vertex. Note that each edge can choose its direction arbitrarily: the direction factors back in when we apply the multiplication by $b_e$ at the end, which is also a local step. Lemma 4.3 allows us to perform this in $O(t \rho(\sqrt{n} + D))$ rounds of communication.

Solving a Laplacian system on each of the rows of this matrix then gives $QR^{-1/2}B \in \mathbb{R}^{t \times n}$. The $t$ solution value at each root vertex can then be broadcasted back to the edges again via Lemma 4.3. Taking sum of squares of differences of these values across each edge’s end points then gives the answer, after another $O(\delta^{-2} \log n)$ rounds of communication in $\overline{G}$. \hfill $\square$

Our identification of steady edges relies on a similar estimation procedure, but in the $\ell_1$ norm.

Lemma 5.13 (Analog to [LS18] Proposition 4.3). There is an algorithm \textsc{ColumnApx} that given a Laplacian solver \textsc{Solve}, a graph $G$ with resistances $r_e$ that $\rho$-minor distributes into $\overline{G}$, and a subset $W \subseteq E(G)$ specified via flags on their images in $\overline{G}$, \textsc{ColumnApx}(e, G, W) computes for all edge $e \in W$ returns the quantity

$$\sum_{e \in W \text{e} \neq f} \frac{\left| b_e^\dagger L(G)^\dagger b_f \right|}{\sqrt{r_e} \sqrt{r_f}}$$

to within a factor of 2 with high probability, stored on their images in $\overline{G}$’s end points. The cost of the algorithm is:

1. $O(\log^2 n)$ calls to \textsc{Solve} on graphs that $\rho$-minor distributes into $\overline{G}$, to accuracy $1/\text{poly}(n)$.
2. An additional $O(\rho(\sqrt{n} + D) \log^2 n)$ rounds of communication in $\overline{G}$.

This will use an $\ell_1$-sketch result.

Lemma 5.14 ([Ind06] Theorem 3 stated for $\ell_1$). Given an integer $d \geq 1$, $\delta \in (0, 1)$, $\epsilon \in (0, 1)$ there is a matrix $C \in \mathbb{R}^{t \times d}$ with $t = O(\epsilon^{-2} \log(1/\delta))$ and an algorithm \textsc{Recover}(u, d, \delta, \epsilon) satisfying the following properties.

- For any vector $v \in \mathbb{R}^d$ the algorithm \textsc{Recover}($Cu$, $d$, $\delta$, $\epsilon$) with probability $1 - \delta$ outputs an estimate $r$ with

$$ (1 - \epsilon) \|v\|_1 \leq r \leq (1 + \epsilon) \|v\|_1. $$

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• Every element of $C$ is independently sampled from a Cauchy distribution.

We use this sketch in the same way as Proposition 4.3 of [LS18] to prove Lemma 5.13.

Proof of Lemma 5.13. We will use the $\ell_1$-sketch in Lemma 5.14 in a way analogous to the $\ell_2$ resistance estimation procedure proven above in Lemma 5.4. However, to avoid estimating the diagonal entries, we will repeat $O(\log n)$ times the following: randomly make a partition $W = U \cup (W \setminus U)$, where each edge $e \in W$ is in $U$ with probability $1/2$, and note that for each edge $e \in U$ that

\[
\mathbb{E} \left[ \sum_{f \in (W \setminus U)} \frac{\| \tilde{b}_e \|^2}{\sqrt{\tilde{r}_e} \sqrt{r_f}} \right] = \frac{1}{2} \sum_{e \in W} \frac{\| \tilde{b}_e \|^2}{\sqrt{\tilde{r}_e} \sqrt{r_f}}.
\]

A similar equality follows for $e \in W \setminus U$. Therefore, by a Chernoff bound, taking the mean of $O(\log n)$ samples and scaling by a factor of 2 will result in a 0.1 approximation. This sampling, and aggregation can also be done locally per edge. So it suffices to provide a 0.1-approximation of the quantities

\[
\sum_{f \in (W \setminus U)} \frac{\| \tilde{b}_e \|^2}{\sqrt{\tilde{r}_e} \sqrt{r_f}}
\]

for a fixed partition $W = U \cup (W \setminus U)$ using Lemma 5.14.

Let $R_{W \setminus U}$ and $B_{W \setminus U}$ be the diagonal resistance and incidence matrix restricted to involving $W \setminus U$. Note that for $\tilde{v}_e = \tilde{r}_e^{-1/2} \cdot R_{W \setminus U}^{-1/2} B_{W \setminus U} L(G) \dagger \tilde{b}_e$, 

\[
\| \tilde{v}_e \|_1 = \sum_{f \in (W \setminus U)} \frac{\| \tilde{b}_e \|^2}{\sqrt{\tilde{r}_e} \sqrt{r_f}}.
\]

Now using Lemma 5.14, sample matrix $C \in \mathbb{R}^{t \times |U|}$ for $\epsilon = 0.1, \delta = n^{-6}$, and $t = O(\log n)$. For the matrix

\[
M \overset{\text{def}}{=} CR_{W \setminus U}^{-1/2} B_{W \setminus U} L(G) \dagger
\]

we have with probability $1 - \delta$ that

\[
\text{RECOVER} \left( M \left( \tilde{r}_e^{-1/2} \tilde{b}_e \right), |U|, \delta, \epsilon \right)
\]

outputs a 0.1-approximation of the quantity $\| \tilde{v}_e \|_1$, as desired.

Now we analyze the cost, which is analogous to the $\ell_2$ sketch described above in the proof of Lemma 5.4 above. To compute $M$, first compute $CR_{W \setminus U}^{-1/2} B_{W \setminus U}$ and store it at the root vertices of supernodes using $\tilde{O}(\rho(\sqrt{n} + D))$ rounds of communication in $\overline{G}$. Multiplying its rows by $L(G) \dagger$ requires calling the solver $t = O(\log n)$ times, once per row of $CR_{W \setminus U}^{-1/2} B_{W \setminus U}$. This gives $M$, which can be broadcasted back to the edges images in $\overline{G}$ using another $\tilde{O}(\rho(\sqrt{n} + D))$ rounds of communication in $\overline{G}$, after which RECOVER can be ran locally per edge.

Factoring in the additional $O(\log n)$ overhead in the number of times this procedure is applied gives the result. 

\[31\]
We also need a sketch for computing the effect of each edge on the Schur complement on $T$.

Lemma 5.15 (Difference sketch, [LS18] Lemma 1.4). There is a routine DIFFAPX that given a graph $G$ with resistances $r_e$ which $\rho$-minor distributes into $\overline{G}$, and a subset $T \subseteq V(G)$ specified via labels at the root vertices of super vertices, DIFFAPX($e, G$) returns approximations to

$$r_e^{-1} \vec{b}_e^T L(G)^+ \begin{bmatrix} \text{SC}(G, T) & 0 \\ 0 & 0 \end{bmatrix} L(G)^+ \vec{b}_e$$

that are within a factor of 2 with high probability, stored on the images into $\overline{G}$’s end points for all edges $e$. It requires

1. $O(\log n)$ calls to SOLVE to accuracy $1 / \text{poly}(n)$ on graphs that $2\rho$-minor distribute into $\overline{G}$.
2. An additional $\tilde{O}(\rho(\sqrt{n} + D))$ rounds of communication in $\overline{G}$.

Proof. By Lemma 2.3, we have that

$$r_e^{-1} \vec{b}_e^T L(G)^+ \begin{bmatrix} \text{SC}(G, T) & 0 \\ 0 & 0 \end{bmatrix} L(G)^+ \vec{b}_e = r_e^{-1} \left\| \begin{bmatrix} \vec{r} e^{-1} \vec{b}_e \\ 0 \end{bmatrix} \right\|_2^2.$$

Therefore, by Lemma 5.12, we can pick a random $\pm 1$ matrix $Q \in \mathbb{R}^{t\times m}$ for $t = O(\log n)$ with

$$r_e^{-1} \vec{b}_e^T L(G)^+ \begin{bmatrix} \text{SC}(G, T) & 0 \\ 0 & 0 \end{bmatrix} L(G)^+ \vec{b}_e \approx_{0.1} t^{-1} r_e^{-1} \left\| QR^{-1/2} BL(G)^+ \begin{bmatrix} \text{SC}(G, T) & 0 \\ 0 & 0 \end{bmatrix} L(G)^+ \vec{b}_e \right\|_2^2$$

for all $e$ with high probability. Therefore, it suffices to find the cost of computing the matrix

$$M = QR^{-1/2} BL(G)^+ \begin{bmatrix} \text{SC}(G, T) & 0 \\ 0 & 0 \end{bmatrix} L(G)^+.$$

First, we can compute $QR^{-1/2} B$ in $O(t) = O(\log n)$ rounds in $G$, and then multiply by $L(G)^+$ by building a solver for $L(G)^+$, and invoking it on each of the $t = O(\log n)$ rows of $QR^{-1/2} B$. Now, we can multiply by

$$\text{SC}(G, T) = L(G)|_{T,T} - L(G)|_{T,V \setminus T} \left( L(G)|_{V \setminus T,V \setminus T} \right)^{-1} L(G)|_{V \setminus T,T}$$

using $t$ solves involving the matrix $L(G)|_{V \setminus T,V \setminus T}$, plus three matrix-vector multiplications in subgraphs of $G$. The costs of the matrix multiplications is $\tilde{O}(t\rho(\sqrt{n} + D))$ rounds of communication in $\overline{G}$ by Corollary 4.4. For the solves, by Lemma 4.8 we can build a solver for $L(G)|_{V \setminus T,V \setminus T}$ by solving a Laplacian on a graph $G'$ that 2-minor distributes into $G$. Finally, we can multiply by $L(G)^+$ again by solving $t$ more systems, and propagate the resulting matrix back to the entire supervertex in $\overline{G}$ via another broadcast (using Lemma 4.3 again). Combining all these costs gives the result. □
Flow localization and steady edges.

**Theorem 5** (Flow localization [SRS18]). Given a graph $H$ with $m$ edges let

$$
\bar{s}_e = \sum_{f \in H} \left| \frac{\hat{b}_e^\top L(H)\hat{b}_f}{\sqrt{\tau_e} \sqrt{\tau_f}} \right|
$$

Then there is a universal constant $C_{\text{local}}$ such that $\sum_{e \in E} s_e \leq C_{\text{local}} m \log^2 m$.

We can use this result and a random sampling scheme to sample a set of edges which is steady analogously to a result in Li-Schild [LS18].

**Algorithm 5:** Given graph $H$ with terminals $T$, and parameter $\delta$ finds steady edges

1. procedure $\text{FINDSTEADY}(H, T, \delta)$
2. $\alpha \leftarrow \frac{1000 C_{\text{local}} \log m}{\delta}$.
3. For $e \in E$ let $\tilde{v}_e \leftarrow \text{DIFFAPX}(e, H, T)$. \hfill \triangleright \text{Lemma 5.15}
4. For $e \in E$ let $\tilde{s}_e \leftarrow \text{COLUMNAPX}(e, H, E(H))$. \hfill \triangleright \text{Lemma 5.14}
5. $Z_1 \leftarrow \{ e \in E : \tilde{v}_e \leq 16|T|/m \text{ and } \tilde{s}_e \leq 16 C_{\text{local}} \log^2 m \}$.
6. Construct $Z_2$ by including each edge $e \in Z_1$ in $Z_2$ with probability $\alpha$.
7. For $e \in Z_2$ let $\tilde{s}_e^{Z_2} \leftarrow \text{COLUMNAPX}(e, H, Z_2)$. \hfill \triangleright \text{Lemma 5.14}
8. $Z \leftarrow \{ e \in Z_2 : \tilde{s}_e^{Z_2} \leq \delta/2 \}$.
9. Return $Z$.

**Lemma 5.16.** For graph $H$ and terminals $T \subseteq V(H)$, we have that

$$
\sum_{e \in E(H)} r_e^{-1} \tilde{b}_e^\top L(H)^\dagger \begin{bmatrix} \text{SC}(H, T) & 0 \\ 0 & 0 \end{bmatrix} L(H)^\dagger \tilde{b}_e \leq |T|.
$$

**Proof.** Applying Lemma 2.3 gives:

$$
\sum_{e \in E(H)} r_e^{-1} \tilde{b}_e^\top L(H)^\dagger \begin{bmatrix} \text{SC}(H, T) & 0 \\ 0 & 0 \end{bmatrix} L(H)^\dagger \tilde{b}_e = \text{Tr} \left( L(H)^\dagger \begin{bmatrix} \text{SC}(H, T) & 0 \\ 0 & 0 \end{bmatrix} L(H)^\dagger \sum_{e \in E(H)} r_e^{-1} \tilde{b}_e^\top \tilde{b}_e \right)
$$

$$
= \text{Tr} \left( L(H)^\dagger \begin{bmatrix} \text{SC}(H, T) & 0 \\ 0 & 0 \end{bmatrix} \right) = |T| - 1 \leq |T|.
$$

We now show that $\text{FINDSTEADY}$ as in Algorithm 5 computes a set of steady edges with high probability.
Proof of Lemma 5.2. The $1 - m^{-10}$ success probability follow from the guarantees of Lemmas 5.15 and 5.14. The (Quadratic form) condition follows because \textsc{FindSteady} as in Algorithm 5 samples each edge $e \in Z_1$ with probability $\alpha$, hence

$$
\mathbb{E}_{Z_1} \left[ \sum_{e \in Z_1} r_e^{-1} b_e b_e^\top \right] = \alpha \sum_{e \in Z_1} r_e^{-1} b_e b_e^\top \preceq \alpha L(H).
$$

The (Localization) condition follows by the condition in line 8 and the approximation guarantee of Lemma 5.14. The (Variance) condition follows from the guarantees of Lemma 5.15 and line 3.

To show the (Size) condition, we first lower bound $|Z_1|$. Note that at most $m/8$ edges $e$ satisfy

$$
\sum_{f \in E(H)} \frac{|b_e^\top L(H)^\top b_f|}{\sqrt{T_e} \sqrt{T_f}} \geq 8\sqrt{2} m
$$

by Theorem 5. Therefore, we have by the approximation guarantees of Lemmas 5.15 and 5.14 that $|Z_1| \geq 3m/4$. Therefore, $\mathbb{E}[|Z_2|] \geq 3\alpha m/4$. By the approximation guarantee of Lemma 5.14, we know that for all $e \in Z_1$

$$
\mathbb{E}_{Z_2} \left[ \sum_{e \neq f \in Z_2} \frac{|b_e^\top L(H)^\top b_f|}{\sqrt{T_e} \sqrt{T_f}} \right] \leq 32\alpha C_{\text{local}} \log^2 m \leq \delta/30.
$$

By Markov’s inequality and the approximation guarantee of Lemma 5.14, we have that $\mathbb{E}[|Z|] \geq \mathbb{E}[|Z_2|] - \frac{\alpha m}{6} \geq \alpha m/2$ as desired.

To bound the total cost, we can simply sum the costs of Lemma 5.14 and 5.15, as the remaining steps of \textsc{FindSteady} as in Algorithm 5 can be trivially implemented.

\[\square\]

6 Vertex and Edge Reductions

Here we show our reductions via tree and elimination based preconditioners in Section 6.1 and Section 6.2 respectively. This will prove Lemmas 4.9 and 4.10.

6.1 Ultra-Sparsifier

We prove the high error reduction routine as stated in Lemma 4.9

Lemma 4.9. There is a routine $\textsc{UltraSparsify}(G, k)$ in the CONGEST model that given a graph $G$ with $n$ vertices and $m$ edges, that $\rho$-minor distributes into the communication network $\overline{G}$, which has $\overline{n}$ vertices, $\overline{m}$ edges, and diameter $D$, along with a parameter $k$, produces in $O(n^{o(1)}(\rho \sqrt{\overline{n}} + D))$ rounds a graph $H$ such that:

1. $H$ is a subgraph of $G,$

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2. \( H \) has at most \( n - 1 + m2^{O(\sqrt{\log n \log \log n})} / k \) edges.

3. \( \mathbf{L}(G) \preceq \mathbf{L}(H) \preceq k \mathbf{L}(G) \).

Furthermore, the algorithm also gives \( \hat{G}, Z_1, Z_2, C \) such that

1. \( \hat{G} \) 1-minor distributes into \( H \) such that \( \hat{G} = \text{SC}(H, C) \) with \( |C| = m2^{O(\sqrt{\log n \log \log n})} / k \).
2. There are operators \( Z_1 \) and \( Z_2 \) evaluable with \( O(\rho \sqrt{\pi} \log \pi + D) \) rounds of CONGEST communication on \( \hat{G} \) such that:

\[
\mathbf{L}(H)^\dagger = Z_1^\top \begin{bmatrix} Z_2 & 0 \\ 0 & \mathbf{L}(\hat{G})^\dagger \end{bmatrix} Z_1
\]

We follow the construction from [KMP10], which samples off-tree edges with any upper bound on their stretches. To find the tree, we utilize the distributed version of the Alon-Karp-Peleg-West (AKPW) low stretch spanning tree, due to Ghaffari, Karrenbauer, Kuhn, Lenzen, and Patt-Shamir [GKK+15]. They work with a definition of distributed \( N \)-node cluster graphs that was the basis of our definition of distributed \( \rho \)-minor. We start by restating this definition, and describe how we simulate it when \( G \) is itself embedded.

**Definition 6.1.** A distributed \( N \)-node cluster graph \( G = (\mathcal{V}, \mathcal{E}, \mathcal{L}, \mathcal{T}, \psi) \) is defined by a set of \( N \) clusters \( \mathcal{V} = \{S_1, \ldots, S_N\} \) partitioning the vertex set \( \mathcal{V} \), a set of weighted multiedges, a set of cluster leaders \( \mathcal{L} \), a set of cluster trees \( \mathcal{T} \), as well as a function \( \psi \) that maps the edges \( \mathcal{E} \) of the cluster graph to edges in \( \mathcal{E} \). Formally, the tuple \((\mathcal{V}, \mathcal{E}, \mathcal{L}, \mathcal{T}, \psi)\) has to satisfy the following conditions.

1. The clusters \( \mathcal{V} = \{S_1, \ldots, S_N\} \) form a partition of the set of vertices \( \mathcal{V} \).
2. For each cluster \( S_i \), \(|S_i \cap \mathcal{L}| = 1\). Hence, each cluster has exactly one cluster leader \( \ell_i \in \mathcal{L} \cap S_i \). The ID of the node \( \ell_i \) also serves as the ID of the cluster \( S_i \) and for the purpose of distributed computations, we assume that all nodes \( v \in S_i \) know the cluster ID and the size \( n_i := |S_i| \) of their cluster \( S_i \).
3. Each cluster tree \( T_i = (S_i, E_i) \) is a rooted spanning tree of the subgraph \( G[S_i] \) of \( G \) induced by \( S_i \). The root of \( T_i \) is the cluster leader \( \ell_i \in S_i \cap \mathcal{L} \).
4. The function \( \psi : \mathcal{E} \to \mathcal{E} \) maps each edge of \( \mathcal{E} \) to an (actual) edge of \( \mathcal{E} \) connecting the clusters.

As a consequence of Lemma 4.3, we get that shortest paths can be ran on distributed \( N \)-node cluster graphs of \( G \).

**Lemma 6.2.** Let \( G = (\mathcal{V}, \mathcal{E}) \) be a graph with \( n \) vertices and \( m \) edges that \( \rho \)-embed into the communication network \( \hat{G} = (\hat{\mathcal{V}}, \hat{\mathcal{E}}) \), and \( \mathcal{G} = (\mathcal{V}, \mathcal{E}, \mathcal{L}, \mathcal{T}, \psi) \) be a distributed cluster graph for \( G \).

Then we have the following algorithms:

1. For each cluster \( S_i \), the cluster leader \( \ell_i \) broadcasts \( O(\log \pi) \) bit message \( s_i \) to each vertex of \( S_i \) in \( O(\rho \pi^{1/2} \log \pi + D) \) rounds.
2. Assume every vertex \( v \in S_i \) for each \( S_i \in \mathcal{V} \), the corresponding vertex \( v' \in \hat{\mathcal{V}} \) holds a value \( f(v) \). Then computing \( \min_{v \in S_i} \{f(v)\} \) at node \( \ell_i \) for each \( S_i \in \mathcal{V} \) needs \( O(\rho \pi^{1/2} \log \pi + D) \) rounds if the tree \( T_i \) with root \( \ell_i \) is known.
Proof. The definition of distributed $N$-node cluster graphs implies that $G$ 1-minor distributes over $G$. Lemma 4.6 then gives that $G$ $\rho$-minor distributes over $G$, and this distributed mapping can be obtained using $O(\rho m^{1/2} \log \pi + D)$ rounds of computations. The broadcast, and the aggregation of minimums then follow from Lemma 4.3.

This in turn implies that the SplitGraph algorithm in [GKK+15] can be simulated on a graph that’s $\rho$-minor distributed into $G$ in $O(n o(1)(\rho m^{1/2} \log \pi + D))$ rounds. Putting it together gives our variant of the AKPW low stretch spanning tree algorithm, with the main difference being that it’s ran on a $\rho$-minor distributed over our overall communication network.

Lemma 4.6 then gives that $G$ $\rho$-minor distributes over $G$, and this distributed mapping can be obtained using $O(\rho m^{1/2} \log \pi + D)$ rounds of computations. The broadcast, and the aggregation of minimums then follow from Lemma 4.3.

These upper bounds are sufficient for sampling the edges by stretch. The following was shown in [KMP10], or Theorem 2.2.4 in [Pen13].

Lemma 6.3. Let $G = (V,E)$ be a graph with $n$ vertices and $m$ edges that $\rho$-embeds into the communication network $\overline{G} = (\overline{V}, \overline{E})$, and $G = (V,E,L,T,\psi)$ be a distributed cluster graph for $G$. It takes $O(n o(1)(\rho m^{1/2} \log \pi + D))$ rounds to construct a spanning tree $T$ of $G$, along with stretch upper bounds that sum to $m \cdot 2^{O(\sqrt{\log n \log \log n})}$.

We then need to contract the tree so that its size becomes similar to the number of off-tree edges.

Lemma 6.5. Let $H = (V,E)$ be a graph with $n$ vertices and $m$ edges that $\rho$-embeds into the communication network $\overline{G} = (\overline{V}, \overline{E})$. Let $T$ be a spanning tree of $H$ and $W = E - T$ be the set of off-tree edges of $H$ with respect to $T$. There is an algorithm to compute a graph $\hat{G}$ that’s 1-embeddable into $H$ satisfying the following conditions in $O(\rho m^{1/2} \log \pi + D)$ rounds:

1. $\hat{G}$ contains $O(|W|)$ vertices and edges.  
2. There are operators $Z_1$ and $Z_2$ that can be evaluated in $O(\rho m^{1/2} \log \pi + D)$ rounds with 

   $$L(H)^\dagger = Z_1^\dagger \begin{bmatrix} Z_2 & 0 \\ 0 & L(\overline{G})^\dagger \end{bmatrix} Z_1$$

Proof. We use the parallel elimination procedure from Section 6.3 of [BGK+14], specifically Lemma 26. At a high level, it eliminates degree 1 and 2 vertices by random sampling a subset of vertices which have degree 1 or 2, computing an independent set, and eliminating them. The algorithm requires $O(\log n)$ rounds in PRAM, and therefore can be implemented in $O(\log n (\rho \sqrt{\log \pi + D}))$ rounds in the CONGEST model by Lemma 4.3. The operators $Z_1, Z_2$ are computed as in Lemma 26 of [BGK+14].

We can combine these pieces to prove the main ultrasparsification claim.
Proof of Lemma 4.9. The algorithm to prove Lemma 4.9 is as follows.

1. Compute a low-stretch tree using Lemma 6.3.
2. Sample edges using Lemma 6.4 with $\alpha = m \cdot 2^{O(\sqrt{\log n \log \log n})}$.
3. Compute the operators $Z_1, Z_2$ using Lemma 6.5.

We verify the conditions of Lemma 4.9. The approximation guarantees and number of off-tree edges are given by Lemma 6.4. After eliminating degree 1 and degree 2 vertices, the resulting graph has size $O(\alpha n)$ by Lemma 6.5 Part 1, and $Z_1$ and $Z_2$ are computed by Lemma 6.5.

The round complexity in the CONGEST model follows by summing the round complexities in Lemmas 6.3, 6.4, 6.5.

6.2 Elimination / Sparsified Cholesky

The main goal of this section is to prove Lemma 4.10, which allows the elimination of large subsets of vertices under small error.

Lemma 4.10. There is a routine $\text{ELIMINATE}(G, d, \epsilon)$ in the CONGEST model that given a graph $G$ that $\rho$-minor distributes into a communication network $\overline{G}$, along with step count $d$ and error $\epsilon$, produces in $O((\epsilon^{-6} \log^{14} n)^d (\rho \sqrt{\pi} \log \pi + D))$ rounds a subset $T$ and access to operators $Z_1$ and $Z_2$ such that

1. $|T| \leq \left(\frac{49}{50}\right)^d |V(G)|$.
2. The cost of applying $Z_1, Z_1^T$ and $Z_2$ to vectors is $O((\epsilon^{-6} \log^{14} \pi)^d (\rho \sqrt{\pi} \log \pi + D))$ rounds of communication on $\overline{G}$.
3. $L(G)^\dagger$ is $(1 \pm \epsilon)^d$-approximated by a composed operator built from $Z_1, Z_2$, and the inverse of the of the Schur complement of $L(G)$ onto $C, \text{SC}(L(G), C)$:

$$(1 - \epsilon)^d L(G)^\dagger \preceq Z_1^\dagger \begin{bmatrix} Z_2 & 0 \\ 0 & \text{SC}(L(G), C)^\dagger \end{bmatrix} Z_1 \preceq (1 + \epsilon)^d L(G)^\dagger$$

To prove the above lemma, we present a distributed implementation of the Schur Complement Chain (SCC) construction due to Kyng, Lee, Peng, Sachdeva, and Spielman [KLP+16]. The key components to this construction are (i) an algorithm that finds a large near-independent set $F$, and approximates the inverse of the matrix restricted to entries in $F$ and (ii) a procedure for spectrally approximating the Schur complement with respect to $C = V \setminus F$. We next discuss how to implement these components in the CONGEST model.
Finding large $\alpha$-DD sets. When doing Gaussian elimination, the goal is to find a large subset of vertices $F$ such that we can approximate the inverse of $L_{[F,F]}$ by an operator $Z$ that can be constructed efficiently. Ideally, $F$ forms an independent set. Unfortunately, we are not able to find a large independent set but we can instead find a large, almost-independent set, as made precise in the following definition.

**Definition 6.6 ($\alpha$-DD).** A matrix $M$ is $\alpha$-diagonally dominant ($\alpha$-DD) if

$$
\forall i, \quad M_{i,i} \geq (1 + \alpha) \sum_{j \in [F], j \neq i} M_{i,j}.
$$

An index set $F$ is $\alpha$-DD if $M_{[F,F]}$ is $\alpha$-DD.

The algorithm due to [KLP+16] for finding $\alpha$-DD sets in a Laplacian proceeds as follows: (i) pick a random subsets of vertices and (ii) discard all those that do not satisfy the condition in Definition 6.6. The pseudocode for computing such sets is given in Algorithm 6. In the CONGEST model, the way the set is “stored” is that each vertex remembers whether it is in the set.

**Algorithm 6: Find an $\alpha$-DD subset $F$ of $L$.**

1. **procedure** $\text{DDS}(L, \alpha)$
2. Sample each index of $\{1, \ldots, n\}$ independently with probability $\frac{1}{4(1 + \alpha)}$ and let $F'$ be the resulting set of sampled indices.
3. Set

   $$
   F = \left\{ i \in F' : |L_{i,i}| \geq (1 + \alpha) \sum_{j \in F', j \neq i} |L_{i,j}| \right\}.
   $$

4. if $|F| < \frac{n}{8(1 + \alpha)}$ then
5. \hspace{1em} Goto Step 1.
6. return $F$.

We have the following lemma.

**Lemma 6.7.** Let $G = (V, E)$ be a graph that $\rho$-minor distributes into the communication network $\mathbb{G} = (\mathbb{V}, \mathbb{E})$. Let $L$ be the Laplacian matrix associated with $G$ and let $\alpha \geq 0$ be a parameter. Then $\text{DDS}(L, \alpha)$ computes an $\alpha$-DD subset $F$ of $L$ of size $n/(8(1 + \alpha))$ in $O(\rho \sqrt{n} \log n + D)$ rounds.

**Proof.** In [LPS15, Lemma 5.2] (and more generally in [KLP+16]), it is shown that Algorithm 6 computes an $\alpha$-DD subset $F$ of size $n/(8(1 + \alpha))$. To bound the round complexity of the algorithm, consider the following distributed implementation:

1. Include each index of $\{1, \ldots, n\}$ in $F'$ with probability $\frac{1}{4(1 + \alpha)}$.

2. Each node corresponding to $i \in F'$ sums up the values $|L_{i,j}|$ of the indices $j$ corresponding to its neighbors in $G$, and then decides whether $|L_{i,i}| \geq (1 + \alpha) \sum_{j \in F', j \neq i} |L_{i,j}|$ and if so declares itself as belonging to $F$.  


3. The size of $F$ is computed by an (arbitrarily decided) leader vertex, which aggregates the sum of the following values over all vertices $v$ in $G$: 1 if $v$ is in $F$ and 0 otherwise.

4. The leader checks whether $|F| < n/(8(1 + \alpha))$. If the latter holds, then the leaders informs all the vertices in $G$ to repeat the previous steps. Otherwise, the algorithm terminates.

In the CONGEST model, Step 1 requires no communication between the nodes: each root vertex of the supervertices does the sampling independently. In Step 2, each node computes an aggregate of values stored by its neighbors in $G$, which by Lemma 4.3 takes $O(\sqrt{n \log n} + D)$ rounds. It is well-known that Steps 3 and 4 can be carried out in $O(D)$ rounds by routing the messages via a BFS tree rooted at the leader. Together with the fact that Algorithm 6 terminates in at most 2 iterations in expectation (see [LPS15, Lemma 5.2]), it follows that the distributed implementation takes $O(\rho \sqrt{n} \log n + D)$ rounds in expectation.

**Jacobi Iteration on $\alpha$-DD matrices.** Using an $\alpha$-DD set $F$, we will construct an operator $Z$ that approximates $L_{[F,F]}^{-1}$, and can be applied efficiently to any vector. An important observation is that we can write $L_{[F,F]} = X_{[F,F]} + Y_{[F,F]}$, where $X_{[F,F]}$ is a diagonal matrix and $Y_{[F,F]}$ is a Laplacian matrix. We have the following lemma.

**Lemma 6.8.** Let $G = (V,E)$ be a graph that $\rho$-minor distributes into the communication network $\overline{G} = (\overline{V},\overline{E})$. Let $L$ be the Laplacian matrix associated with $G$ and let $F$ be a subset of $V$ such that $L_{[F,F]}$ is $\alpha$-DD for some $\alpha \geq 4$. Then $\text{Jacobi}(L_{[F,F]},\cdot,\epsilon)$ gives a linear operator $Z$ that over vectors given on the root vertices of the supervertices such that for any vector $\vec{b}$ given by storing $\vec{b}_{v,G}$ on $V_{map}^G(\cdot)$, returns in $O((\rho \sqrt{n} + D) \log(1/\epsilon))$ rounds $Z \vec{b}$ stored on the same vertices, for some matrix $Z$ such that

$$L_{[F,F]} \preceq Z^{-1} \preceq L_{[F,F]} + \epsilon \cdot SC(L,F).$$

Note that the matrix $Z$ is only used in the analysis, and is never explicitly constructed by the algorithm. We first give the pseudocode of this algorithm in the centralized setting, and then show its distributed implementation.

**Algorithm 7:** Solve $L_{[F,F]} \cdot \vec{x}_F = \vec{b}_F$ up to $\epsilon$ accuracy

```
1 procedure \text{Jacobi}(L_{[F,F]}, \vec{b}_F, \epsilon)
2     Set $L_{[F,F]} = X_{[F,F]} + Y_{[F,F]}$ such that $X_{[F,F]}$ is diagonal and $Y_{[F,F]}$ is a Laplacian.
3     Set $k$ to be an odd integer that is greater than $\log(3/\epsilon)$.
4     Set $\vec{x}_F^{(0)} = X_{[F,F]}^{-1} \vec{b}_F$.
5     for $i = 1, \ldots, k$ do
6         Set $\vec{x}_F^{(i)} = -X_{[F,F]}^{-1} Y_{[F,F]} \vec{x}_F^{(i-1)} + X_{[F,F]}^{-1} \vec{b}_F$.
7     return $\vec{x}_F^{(k)}$.
```

To measure the quality of the operator produced by $\text{Jacobi}$ procedure, we observe that $k$ iterations produce the operator

$$Z^{(k)} := \sum_{i=0}^{k} X_{[F,F]}^{-1} \left( -Y_{[F,F]} X_{[F,F]}^{-1} \right)^i$$

(5)
by induction. Concretely, suppose we have
\[ \bar{x}^{(k-1)}_F = \sum_{i=0}^{k-1} X^{-1}_{[F,F]} \left( -Y_{[F,F]} X^{-1}_{[F,F]} \right)^i \bar{b}_F, \]
then substituting this into the step in Line 6 gives
\[ \bar{x}^{(k)}_F = -X^{-1}_{[F,F]} Y_{[F,F]} \bar{x}^{(k-1)}_F + X^{-1}_{[F,F]} \bar{b}_F \]
\[ = X^{-1}_{[F,F]} \bar{b}_F + \left( -X^{-1}_{[F,F]} Y_{[F,F]} \right) \sum_{i=0}^{k-1} X^{-1}_{[F,F]} \left( -Y_{[F,F]} X^{-1}_{[F,F]} \right)^i \bar{b}_F \]
\[ = X^{-1}_{[F,F]} \bar{b}_F + \sum_{i=1}^{k} X^{-1}_{[F,F]} \left( -Y_{[F,F]} X^{-1}_{[F,F]} \right)^i \bar{b}_F = \sum_{i=0}^{k} X^{-1}_{[F,F]} \left( -Y_{[F,F]} X^{-1}_{[F,F]} \right)^i \bar{b}_F. \]

We next review two lemmas from [KLP+16] that help us prove the approximation accuracy of the Jacobi iteration on \( L_{[F,F]} \). The first shows that \( \alpha \)-DD matrices admit good diagonal preconditioners. The second gives a way to bound the error produced by \textsc{Jacobi}.

**Lemma 6.9** ([KLP+16], Lemma 3.6.). Let \( L_{[F,F]} \) be an \( \alpha \)-DD matrix which can be written in the form \( X_{[F,F]} + Y_{[F,F]} \) where \( X_{[F,F]} \) is diagonal and \( Y_{[F,F]} \) is a Laplacian. Then \( \frac{\alpha}{2} Y \preceq X. \)

**Lemma 6.10** ([KLP+16], Lemma E.1.). Let \( L_{[F,F]} \) be an \( \alpha \)-DD matrix with \( L_{[F,F]} = X_{[F,F]} + Y_{[F,F]} \) where \( 0 \preceq Y_{[F,F]} \preceq \beta X \) for some \( 0 < \beta < 1 \). Then, for any odd \( k \) and \( Z^{(k)} \) as defined in Eq. (5), we have
\[ X_{[F,F]} + Y_{[F,F]} \preceq (Z^{(k)})^{-1} \preceq X_{[F,F]} + (1 + \delta) Y_{[F,F]}, \]
where
\[ \delta = \beta^k \frac{1 + \beta}{1 - \beta^{k+1}}. \]

**Proof of Lemma 6.8.** Let \( Y_{[F,F]} \) be the matrix generated when calling \textsc{Jacobi} with \( L_{[F,F]} \). Since \( L_{[F,F]} \)
is an \( \alpha \)-DD matrix, by extending \( Y_{[F,F]} \) with zero entries, we have \( Y \preceq L \). This in turn implies that \( Y_{[F,F]} = \text{SC}(Y_{[F,F]}, F) \preceq \text{SC}(L, F). \)

Lemma 6.9 gives that \( \frac{\alpha}{2} Y \preceq X. \) As \( \alpha \geq 4 \), we can invoke Lemma 6.10 with \( \beta = 1/2 \), which in gives that \( (1 + \beta)/(1 - \beta^{k+1}) \leq 3. \) Therefore, our choice of \( k = \log(3/\epsilon) \) gives the desired error guarantee. To bound the round complexity of the algorithm, consider the following distributed implementation of \textsc{Jacobi}:

1. Store the values \( \bar{b}_F(u), \ X_{[F,F]}(u, u), \ Y_{[F,F]}(u, u) \) at \( V^G_{\text{Map}}(u) \). Store the off-diagonal entries of \( Y_{[F,F]} \), together with their weights, in the endpoints of mapped edges \( E^G_{\text{Map}}(e) \): this is possible because \( Y_{[F,F]} \) is a Laplacian.

2. Set \( k = \log(1/\epsilon) \) and \( \bar{x}^{(0)}_F(u) = X^{-1}_{[F,F]}(u, u) \cdot \bar{b}_F(u) \) for each \( u \in F \).

3. For \( i = 1, \ldots, k \) do...
(a) For each $u \in F$ set
\[
\tilde{x}_{F}^{(i)}(u) \leftarrow X_{[F,F]}^{-1}(u,u) \cdot \tilde{b}_{F}(u) - X_{[F,F]}^{-1}(u,u) \sum_{v \in F: Y_{[F,F]}(u,v) \neq 0} Y_{[F,F]}(v,u) \tilde{x}^{(i-1)}(v)
\]

using the matrix-vector multiplication primitive from Corollary 4.4, with results stored on all root vertices, $V_{map}^{G \to \overline{G}}(u)$.

4. Every vertex $u \in F$ returns $\tilde{x}_{F}^{(k)}(u)$.

The complexity of the algorithm is dominated by the number of rounds to implement Step 3, which is given by Corollary 4.4. As there are $k = \log(1/\epsilon)$ iterations, we have that Step 3 requires $O(\log(1/\epsilon))$ rounds in $G$.

By Lemma 4.3 and using the fact that weights of the network, and hence the solution vector’s magnitudes, are polynomially bounded, we can simulate the algorithm in the original communication network $\overline{G}$ in $O((\rho\sqrt{n} \log n + D) \log(3/\epsilon))$ rounds.

Approximating Schur complements using low congestion random walks. We next show that $\alpha$-DD sets are useful when approximating Schur complements. A key ingredient to our construction is the following combinatorial view of Schur complements.

It is well known that $SC(L, T)$ is a Laplacian matrix of a graph on vertices in $T = V \setminus F$. For our purposes, it will be useful interpret $SC(L, T)$ in terms of random walks. To this end, given a walk $W = u_0, \ldots, u_\ell$ of length $\ell$ in $G$ with a subset of vertices $T$, we say that $W$ is a terminal-free walk if $u_0, u_\ell \in T$ and $u_1, \ldots, u_{\ell-1} \notin T$.

Lemma 6.11. For any undirected, weighted graph $G$ and any subset of vertices $T$, the Schur Complement $SC(G, T)$ is given as a union over all multi-edges corresponding to terminal-free walks $u_0, \ldots, u_\ell$ with weight
\[
\frac{\prod_{0 \leq i < k} \tilde{w}_{u_i u_{i+1}}}{\prod_{0 \leq i < k} \sum_{u_1 \in E(G)} \tilde{w}_{u_1 u}}
\]

The theorem below allows us to efficiently sample from this distribution of walks while paying a small cost in the approximation quality.

Lemma 6.12 (Theorem 3.1 in [DGGP19]). Let $G = (V, E)$ be an undirected, weighted graph with a subset of vertices $T$. Let $\epsilon \in (0, 1)$ be an error parameter and $\mu = \Theta(\epsilon^{-2} \log n)$ be some parameter related to the concentration of sampling. Let $H$ be an initially empty graph, and for every edge $e = (u, v) \in G$, repeat $\mu$ times the following procedure, where a random step from a vertex is taken proportional to the edge weights of its adjacent edges.

1. Simulate a random walk starting from $u$ until it hits $T$ at vertex $t_1$.
2. Simulate a random walk starting from $v$ until it hits $T$ at vertex $t_2$.
3. Let $\ell$ be the total length of this combined walk (including edge $e$). Add the edge $(t_1, t_2)$ to $H$ with weight
\[
\frac{1}{\mu \sum_{i=0}^{\ell-1} (1/\tilde{w}_{u_i, u_{i+1}})}.
\]
The resulting graph $H$ satisfies $L(H) \approx \text{SC}(L(G), \mathcal{T})$ with high probability.

The main idea to make use of the above theorem is to compute an $\alpha$-DD set $F$ using Lemma 6.7 as this ensures that the random walks in the graph are short in expectation. However, since we are dealing with weighted graphs, there might be scenarios where the expected congestion of an edge is prohibitively large, which makes it difficult to recursively repeat the algorithm. To alleviate this, we add new vertices to the terminals, whenever they have too much congestion. Note that because $G$ is distributed over $\overline{G}$ as a minor, we can only accommodate small vertex congestion due to the need for each root node to inform the entire supervertex. As a result, our resulting congestion depends on the average degree, and we resolve this via calling sparsification (Corollary 4.5) at each step.

Let $W$ be the family of walks generated in Lemma 6.12. For $e \in E$, let $\text{cong}_W(e)$ denote the number of walks from $W$ that use the edge $e$. Algorithm 8 below computes $W$. Note that it can also be run implicitly to generate the congestion on every edge without exceeding the communication limit on any edges: we simply pass around the congestion on every edge.

**Algorithm 8: Generate random walks from each edge until they hit terminals**

```
1 procedure RANDOMWALK($G, \mathcal{T}, \mu$)
2    Set $W \leftarrow \emptyset$
3    for $e = (u, v) \in E(G)$ in parallel do
4        for $i = 1, \ldots, \mu$ in parallel do
5            Generate a random walk $W(u, i)$ from $u$ until it hits $\mathcal{T}$ at vertex $t_1$.
6            Generate a random walk $W(v, i)$ from $v$ until it hits $\mathcal{T}$ at vertex $t_2$.
7            Set $W(e, i) = W(u, i) \cup (u, v) \cup W(v, i)$ and $W = W \cup \{W(e, i)\}$.
8    return $W$.
```

**Implementation of random walks.** We implement the random walks in lines 5 and 6 of RANDOMWALK as in Algorithm 8 as follows. When a non-terminal vertex gets a random walk edge into it, first we do a sum aggregation so that the leader of the corresponding cluster / super-vertex knows how many out-edges to compute. Below, we will ensure that all non-terminal vertices have low congestion as intermediate vertices of random walks, so we will focus on sampling a single out-edge. To do this, the vertex aggregates the sum of weights of out-edges, which we will call $W_{\text{total}}$. Now, the leader samples a random real number $r \in [0, W_{\text{total}}]$. Finally, the leader vertex does a binary search on the label of the out-edge and aggregates sums to figure out which out-edge corresponds to the sample $r$. All these steps can be done by Lemma 4.3.

We now move on to giving the distributed random walk based algorithm that approximates Schur complements. Pseudocode of this routine is in Algorithm 9.

Here, we discuss subtleties in the distributed implementation of Algorithm RANDOMWALK and RANDOMWALKSCHUR (Algorithms 8 and 9) in the CONGEST model.

**Lemma 6.13.** Let $G = (V, E)$ be a graph that $\rho$-minor distributes into the communication network $\overline{G} = (\overline{V}, \overline{E})$. Let $F$ be an $\alpha$-DD set, $\mathcal{T} = V \setminus F$, $\epsilon \in (0, 1)$ be an error parameter and $\gamma \geq 1$ be a congestion parameter. Then the procedure RANDWALKSCHUR($G, \mathcal{T}, \epsilon, \gamma, \alpha$) outputs in $O(\alpha^2 \epsilon^{-2} \log n(\rho \sqrt{n}\log n + \ldots$
Algorithm 9: Distributed Approximate Schur complements using random walks

1. \textbf{procedure} \textsc{RandWalkSchur}(G, T, \epsilon, \gamma, \alpha)
2. \hspace{1em} Initialize $\hat{T} \leftarrow T$
3. \hspace{1em} Set $H \leftarrow \emptyset$ and $\mu \leftarrow O(\epsilon^{-2} \log n)$
4. \hspace{1em} Implicitly compute the expected congestion of $W = \text{RandomWalk}(G, T, \mu)$ by
   \hspace{2em} propagating the expected congestion on vertices and edges evenly to neighbors for
   \hspace{2em} $O(\alpha \log n)$ steps.
5. \hspace{1em} \textbf{for all} vertices $u$ with $\mathbb{E}[	ext{cong}_W(u)] > \gamma$, \textbf{in parallel} do
6. \hspace{2em} \hspace{1em} Add $u$ to $\hat{T}$, $\hat{T} \leftarrow \hat{T} \cup \{u\}$.
7. \hspace{1em} Set $W \leftarrow \text{RandomWalk}(G, \hat{T}, \mu)$ (with walks explicitly generated).
8. \hspace{1em} Initialize the minor distribution of $H$ into $\overline{G}$ by associating each terminal $t$ with all
   \hspace{2em} vertices involved in all random walks that ended at $t$, and building $T^{H \rightarrow \overline{G}}(t)$ to be a
   \hspace{2em} spanning tree of all edges used, plus $T^{G \rightarrow \overline{G}}(u)$ of all vertices on these walks.
9. \hspace{1em} \textbf{for} $W(e, i) \in W$ \textbf{do}
10. \hspace{2em} \hspace{1em} Let $(t_1, t_2)$ be the endpoints of the walk $W(e, i)$.
11. \hspace{2em} \hspace{1em} Let $\ell$ be the length of $W(e, i)$.
12. \hspace{2em} \hspace{1em} Set $H = H \cup (t_1, t_2, w(t_1, t_2))$ with $\bar{w}(t_1, t_2) := 1/(\mu \sum_{i=0}^{\ell-1} (1/\bar{w}_{u_i, u_{i+1}}))$.
13. \hspace{1em} \textbf{return} $H, \hat{T}$.

$D$) rounds a graph $H$ along with its $\alpha \gamma \log n$-minor distribution into $\overline{G}$ such that with high probability,

$$L(H) \approx_{\epsilon} \text{SC}(L(G), \hat{T})$$

for some (slightly larger) superset $\hat{T} \supseteq T$ of size at most $n - |F| + O(\alpha m \epsilon^{-2} \log n / \gamma)$.

\textbf{Proof.} The spectral guarantee $L(H) \approx_{\epsilon} \text{SC}(L(G), \hat{T})$ follows directly from Lemma 6.12. To bound the
size of $T$, first note that by definition $T = V \setminus F$ and thus $|T| = n - |F|$. Next, as $F$ is an $\alpha$-DD set, the
expected length of a random walk that starts at an endpoint of any edge in $G$ and hits a vertex in $T$ is
$O(\alpha)$. Our algorithm simulates $O(m \epsilon^{-2} \log n)$ random walks for each edge, thus the total congestion
generated by these walks is $O(\alpha m \epsilon^{-2} \log n)$.

The latter gives that there can be at most $O(\alpha m \epsilon^{-2} \log n / \gamma)$ vertices whose expected congestion
is larger than $\gamma$, and \textsc{RandWalkSchur} adds these vertices to the set $T$. It follows that $|\hat{T}| \leq n - |F| + O(\alpha m \epsilon^{-2} \log n / \gamma)$. For each vertex $u$, the congestion incurred by other edges are independent
random variables bounded by the length of the walks, which is $O(\log n)$. So by a Chernoff bound, the
congestion of all edges with expected congestion less than $\gamma > O(\log^2 n)$ is at most $O(\gamma)$ with
high probability. So after line 4 of Algorithm 9 adds all vertices with high expected congestion into
the terminals (to form $\hat{T}$), all subsequent vertices in $V \setminus \hat{T}$ have vertex congestion at most $O(\gamma)$ in the
second random walk in line 7 with high probability.

We next study the round complexity. To this end, recall that the expected length of each walk in
$W$ is $O(\alpha \log n)$ with high probability.

When we are only passing the congestion of a vertex to neighbors, that is, running the walks
implicitly, each round can be executed in one round of message passing as described in Lemma 4.3.
As we execute $\mu = O(\epsilon^{-2} \log n)$ rounds for each edge, we have that the round complexity of the congestion estimation part of RANDOMWALK is $O(\alpha \rho (\sqrt{n} + D) \epsilon^{-2} \log^2 n)$.

For the explicit generation, we aggregate, for all non-terminal vertices of $G$, the walks that reach them, to the root node of the corresponding super vertex in $\overline{G}$. Then we broadcast these walks outward, we simulate the choice of a random edge by total weights of edges in subtrees (which we compute via Lemma 4.2). As the node congestions are at most $2\gamma$, Lemma 4.3 lets us perform these propagations in $\overline{G}$ in $O(\alpha \gamma \epsilon^{-2} \log n (\rho \sqrt{n} \log n + D))$ rounds.

Finally, to create the minor distribution of $H$, the graph with the new random walk edges, into $\overline{G}$, we extend the terminals to include the supervertices of all intermediate (non-terminal) vertices. As the non-terminals have at most $O(\gamma)$ walks through them, the resulting mapping is still a $\rho \gamma$-minor distribution into $\overline{G}$.

We remark that in this scheme, the end points of the new edges in $H$ cannot actually know these edges in a centralized manner (e.g. aggregate them at root vertices). Instead, such walks are only passed to the vertices in $\overline{G}$ that correspond to the first and last edges of the corresponding walk. This is because we can only guarantee low node congestion of intermediate vertices. Note that that in turn necessitates us sparsifying the graph at every intermediate step as well.

**Vertex Sparsifier Chain.** Bringing together the above algorithmic components leads to an algorithm for computing a vertex sparsifier chain, whose pseudocode is given in Algorithm 10 below.

**Algorithm 10: Eliminate a large subset of vertices for $d$ rounds**

1. procedure $\text{ELIMINATE}(G,d,\epsilon)$
   2. Set $L^{(0)} \leftarrow L$ and $\overline{T}_0 = V$.
   3. Compute a spectral sparsifier $M^{(0)} \approx \epsilon L^{(0)}$ (Corollary 4.5).
   4. for $0 < i \leq d$ iteratively do
      5. Let $F_i$ be an $\alpha$-DD set of $M^{(i-1)}$ (Lemma 6.7).
      6. Construct an operator $(Z^{(i)})^{-1}$ that approximates $M_{[F,F]}^{(i-1)}$ (Lemma 6.8).
      7. Compute $\overline{M}^{(i+1)} \approx \epsilon SC(M^{i-1}, \overline{T}_i)$ (Lemma 6.13) with $\overline{T}_i = \overline{T}_{i-1} - F_i + U_i$, where $U_i$ is the set of extra vertices added to ensure low congestion.
      8. Compute an $\epsilon$-spectral sparsifier $M^{(i+1)}$ of $\overline{M}^{(i+1)}$. (Corollary 4.5)
   9. Let $Z_1, Z_2$ be stored implicitly as the product of matrices using the Cholesky factorization (Lemma 2.2).
   10. return $M^{(d)}, Z_1, Z_2$.

**Proof of Lemma 4.10.** We start by analyzing the round complexity of the algorithm. By Corollary 4.5, there is a distributed algorithm for computing a sparsifier with $O(n \log^5 n \epsilon^{-2})$ edges in $O(\epsilon^{-2} \log^7 n)$ rounds. We call RANDWALDSCHUR with $\gamma = 1000 \cdot c \cdot \alpha \epsilon^{-2} \log^6 n$, where $c$ is a large enough constant. By Lemma 6.7, we find a 4-DD set of size $n_i/(8(1 + 4)) = n/40$. These together imply that the number of vertices in $G^{(i)}$ after $i$ steps in our algorithm is

$$n_i \leq \left(1 - \frac{1}{40} + \frac{1}{1000}\right) n_{i-1} \leq \left(1 - \frac{1}{50}\right) n_{i-1}.$$
By induction, $n_d \leq \left(\frac{\rho}{\epsilon}\right)^d n$.

In an iteration of the algorithm, the dominating cost is (1) approximating the Schur complement and (2) computing the spectral sparsifier. By our choice of $\alpha$ and $\gamma$ and Lemma 6.13, the number of rounds required to implement the first one is $O(\epsilon^{-4} \log^7 n (\rho \sqrt{\tau} \log \tau + D))$. The second one introduces a $O(\epsilon^{-2} \log^4 n)$ overhead, which then gives a round complexity of $O(\epsilon^{-6} \log^4 n (\rho \sqrt{\tau} \log \tau + D))$ per one step in ELIMINATE. Thus after $d$ steps, the round complexity is:

$$O((\epsilon^{-6} \log^4 n)^d (\rho \sqrt{\tau} \log \tau + D)).$$

The error is $(1 \pm \epsilon)^d$ because we do $d$ rounds of elimination, and each round accumulates $(1 \pm \epsilon)$-multiplicative error by using Lemma 6.8 to bound the quality of the inverse of $Z^{(i)}$, Lemma 6.13 to bound the quality of the Schur complement, and Corollary 4.5 to spectrally sparsify the Schur complement, each of which accumulates error $\epsilon/4$. \hfill \Box

## 7 Implications in Graph Algorithms

In this section we use Theorem 1 to give improved algorithms for maximum flow, min-cost flow, and shortest paths with negative weights in the CONGEST model (Theorems 7, 8, and 9). Our goal is to show that our distributed Laplacian solver can be used to achieve improved complexities for these three problems, so we provide pseudocode in Algorithms 12, 13, 15 (full details of these algorithms are given in Appendix C), and analyze the runtimes in the distributed setting. Our runtimes come from using the Laplacian system solver in Theorem 1 to implement an interior point method until the graph has a low amount of residual flow remaining, which we then route with augmenting path [GU15] or shortest path with positive weights [CM20], both taking $\tilde{O}(n^{1/2} D^{1/4} + D)$ rounds per iteration. In the remainder of this section, we formalize this reasoning. There are several additional technical pieces, as the algorithms of [Mad16, CMSV17] require changing the graph by adding edges, etc. Throughout, we assume that our Laplacian system solvers are exact – it is justified in the papers [Mad16, CMSV17] that solving to accuracy $1/poly(m, U)$, where $U$ is the maximum weight / capacity suffices to implement the interior point methods.

In Section 7.2, we simulate Cohen’s flow rounding algorithm [Coh95] in the CONGEST model. Given an $s - t$ flow $\vec{f}$, it returns an integral $s - t$ flow $\vec{f}'$ in $O(\sqrt{m} \log m \cdot D \cdot \log(1/\Delta))$ rounds such that the flow value of $\vec{f}'$ is at least $\vec{f}$'s flow value. In Section 7.3, we implement maximum flow [Mad16] in the CONGEST model, which takes $\tilde{O}(\sqrt{m} U^{1/3} (\sqrt{n})^{1/2} + D) + \sqrt{n} D^{1/4} + D^{1/2})$ rounds. In Section 7.4, it takes $\tilde{O}(\sqrt{m} \sqrt{\tau} \sqrt{n}^{1/3} (n^{1/3} + D^{1/4}) + \sqrt{n} D^{1/2})$ rounds to execute the min-cost flow [CMSV17] that consists of Laplacian solver, flow rounding and single-source shortest path [CM20] in the CONGEST model. In addition, shortest paths with negative weights can be implemented in the same rounds since it utilizes min-cost flow to make edges non-negative and then compute the shortest paths [CM20].

### 7.1 Flow Preconditioned Minor

In this subsection, we define flow preconditioned minor. Compared with the definition of $\rho$ congestion, this definition does not bound the size of the pre-image of vertex mapping function $V_{\text{map}}^G \rightarrow H$. Instead, we only allow a bounded number of vertices of $G$ that maps to more than one vertex of $\tilde{H}$.

**Definition 7.1.** Let $G = (V, E)$ be a minor of $H = (V_H, E_H)$ (as Definition 4.1). We say this minor is $(\rho, \alpha)$-flow-preconditioned if:

...
1. Each edge of $H$ appears as the image of the edge map $E_{\text{map}}^{G\rightarrow H}(\cdot)$, or in one of the trees connecting supervertices, $T^{G\rightarrow H}(v^G)$ for some $v^G$, at most $\rho$ times.

2. There is a set $U \subset V$ with $|U| = \alpha$ such that the following two conditions hold:
   
   (a) $V_{\text{map}}^{G\rightarrow H}(v^G) = V_H$ and $T^{G\rightarrow H}(v^G)$ is a spanning tree of $H$ with depth at most the diameter of $H$ for each $v^G \in U$.
   
   (b) $|V_{\text{map}}^{G\rightarrow H}(v^G)| = 1$ for each $v^G \in V \setminus U$.

We say a $(\rho, \alpha)$-flow-preconditioned minor mapping is stored distributedly, or that $G$ is a $(\rho, \alpha)$-flow-preconditioned minor distributed over $H$ if it’s stored by having all the images of the maps recording their sources (the same as Definition 4.1).

**Lemma 7.2.** Let $G = (V, E)$ be a graph with $n$ vertices and $m$ edges that $(\rho, \alpha)$-flow-preconditioned minor distributes into a communication network $\overline{G} = (\overline{V}, \overline{E})$ with $\pi$ vertices, $\pi$ edges, and diameter $D$. In the CONGEST model, the following operations can be performed using $O(t \alpha D)$ rounds of communication on $\overline{G}$:

1. Each $V_{\text{map}}^{G\rightarrow \overline{G}}(v^G)$ sends $O(t \log n)$ bits of information to all vertices in $S^{G\rightarrow \overline{G}}(v^G)$.
2. Simultaneously aggregate the sum/minimum of $O(t \log n)$ bits, from all vertices in $S^{G\rightarrow \overline{G}}(v^G)$ to $V_{\text{map}}^{G\rightarrow \overline{G}}(v^G)$ for all $v^G \in V(G)$.

**Proof.** Both operations can be achieved by running a BFS or a reverse BFS on $T^{G\rightarrow \overline{G}}(v^G)$ for each $v^G \in V$. \qed

We say a vector $\vec{x} \in \mathbb{R}^V$ on $G$ is distributed on $\overline{G}$ if for each $v^G$, all the vertices of $S^{G\rightarrow \overline{G}}(v^G)$ records $\vec{x}_{v^G}$.

We say a vector $\vec{f} \in \mathbb{R}^E$ defined on edges of $G$ is distributed to $\overline{G}$ if for each $e \in E$, the two endpoints of $E_{\text{map}}^{G\rightarrow \overline{G}}(e)$ records $\vec{f}_e$. Sometimes, we treat a vector $\vec{f} \in \mathbb{R}^E$ defined on edges of $G$ as a matrix, denoted as $M_{\vec{f}}$, such that $M_{\vec{f}_{uv}} = \vec{f}_{uv}$ if $(u, v)$ is an edge in $E$, otherwise $M_{\vec{f}_{uv}} = 0$.

**Lemma 7.3.** Let $\overline{G} = (\overline{V}, \overline{E})$ be a communication network with $\pi$ vertices and $\pi$ edges, and $G = (V, E)$ be a graph that is $(\rho, \alpha)$-flow-preconditioned minor distributed to $\overline{G}$. Then, we have the following basic operations

1. Let $\vec{f}^{(1)}, \vec{f}^{(2)}, \ldots, \vec{f}^{(t)}$ be $t$ edge vectors of $G$ that are distributed on $\overline{G}$, and $\vec{f}$ be an edge vector of $G$ such that $\vec{f}_e$ is a function of $\vec{f}^{(1)}_e, \vec{f}^{(2)}_e, \ldots, \vec{f}^{(t)}_e$ for each edge $e \in E$, then $\vec{f}$ can be computed and distributed on $\overline{G}$ in $O(1)$ rounds.
2. Let $\vec{x}^{(1)}, \vec{x}^{(2)}, \ldots, \vec{x}^{(t)}$ be $t$ vertex vectors of $G$ that are distributed on $\overline{G}$, and $\vec{x}$ be a vertex vector of $G$ such that $\vec{x}_{v^G}$ is a function of $\vec{x}^{(1)}_{v^G}, \vec{x}^{(2)}_{v^G}, \ldots, \vec{x}^{(t)}_{v^G}$, then $\vec{x}$ can be computed and distributed on $\overline{G}$ in $O(1)$ rounds.
3. For a vertex vector $\vec{x} \in \mathbb{R}^V$ on $G$ distributed to $\overline{G}$ and a $p > 1$, $p$-norm of $\vec{x}$ can be computed and broadcast to every vertex of $\overline{G}$ in $O(D)$ rounds.
4. For an edge vector $\vec{f} \in \mathbb{R}^E$ of $G$ distributed on $G$ and an integer $k$, it takes $O(D \cdot \text{poly} (\log \beta))$ rounds to identify top $k$ coordinates of $\vec{f}$ with largest absolute value, where $\beta = \frac{\max_{e \in E} |\vec{f}_e|}{\min_{e,e' \in E: \vec{f}_e \neq \vec{f}_{e'}} |\vec{f}_e - \vec{f}_{e'}|}$.

5. For an edge vector $\vec{f} \in \mathbb{R}^E$ of $G$ and a vertex vector $\vec{x} \in \mathbb{R}^V$ on $G$ both distributed to $G$, $M \vec{f} \vec{x}$ can be computed and distributed to $G$ in $O(\rho + \alpha D)$ rounds if $G$ is a $(\rho, \alpha)$-flow-preconditioned minor distributed to $G$.

**Proof.** The first and the second operation can be computed locally by each vertex of $G$.

The third operation can be computed by a BFS on $G$ such that every $G$ by aggregating the sum of $\vec{x}_v$ for each $v \in V$.

The fourth operation can be computed by binary searching $k$-th largest absolute value of $\vec{f}_e$ among all the $e \in E$ and counting the number of coordinates with absolute value greater than or equal to the binary searched.

The fifth operation can be implemented by computing $\vec{f}_{uw} \vec{x}_u$ at one endpoint of $E_{\text{map}}^{G \rightarrow G}(uv)$ in $S_{\text{map}}^{G \rightarrow G}(u)$ and taking the sum of $\sum_{uv \in E} \vec{f}_{uv} \vec{x}_v$ for each $u$ by $T_{\text{map}}^{G \rightarrow G}(u)$. The number of rounds required is by Lemma 7.2.

**Lemma 7.4.** Let $G = (V, E)$ be a communication network $\pi$ vertices, $\overline{m}$ edges, and diameter $D$. Let $\mathcal{P}$ be a collection of paths/cycles of $G$ such that every edge of $G$ is used for at most $\rho$ times for some $\rho = O(\text{poly}(\overline{m}))$, and for every two consecutive edges $(u, v), (v, w)$ of some path in $\mathcal{P}$, $v$ knows that $(u, v), (v, w)$ are two consecutive edges of some path/cycle of $\mathcal{P}$. Then the following operations can be performed using $O(\rho \overline{m}^{1/2} \log \overline{m} + D)$ rounds of communication on $G$:

1. Every path/cycle of $\mathcal{P}$ is associated with a unique ID such that for each edge $e \in E$ in the path/cycle, the two endpoints of $E_{\text{map}}^{G \rightarrow G}(e)$ know the ID.

2. Let $\vec{f}$ be an edge vector of $G$. Compute the sum of $\vec{f}$ for each path/cycle of $\mathcal{P}$ and let the two endpoints of $E_{\text{map}}^{G \rightarrow G}(e)$ know the result for each edge $e$ in the path/cycle.

**Proof.** We view $\mathcal{P}$ as a graph such that every vertex and edge appears once by treating each appearance of the same vertex/edge as a new vertex/edge. The resulted graph, denoted as $G$, is a graph of $O(\rho \overline{m})$ vertices and edges that corresponds to a set of edge disjoint paths.

We sample each vertex of $G$ with probability $\log \overline{m} / \overline{m}^{1/2}$. At most $O(\rho \overline{m}^{1/2} \log \overline{m})$ vertices are sampled with high probability, and for each simple path of length $\overline{m}^{1/2}$ that corresponds to an induced subgraph of $G$, at least one vertex of the simple path is sampled. Each sampled vertices performs a BFS on $G$ until it reaches another sampled vertex or an endpoint of some path in $G$. We aggregate the result of the BFS (the visit of sampled vertices or endpoints of some path in $G$) to an arbitrary vertex of $G$ in $O(\rho \overline{m}^{1/2} \log \overline{m} + D)$ rounds, and the IDs for paths/cycles of length at least $\overline{m}^{1/2}$ can be assigned and broadcast to each of these paths/cycles in $O(\rho \overline{m}^{1/2} \log \overline{m} + D)$ rounds.

Then, all the paths/cycles in these paths/cycles that do not contain any sampled vertex initiate a BFS from each vertex in the paths/cycles such that if two BFS collide, only the one initiated by small vertex ID is kept. Since every path/cycle that does not contain any sampled vertex is of length at most $O(\overline{m}^{1/2})$, this step can be done in $O(\overline{m}^{1/2})$ rounds, and afterwards the IDs for these paths/cycles can be computed and broadcast to each vertex on these paths/cycles in $O(\overline{m}^{1/2})$ rounds.
Hence, the first operation can be done in $O(\rho m^{1/2} \log m + D)$ rounds. The second operation can also be done in $O(\rho m^{1/2} \log m + D)$ rounds using sampled vertices in a way similar as the first operation.

7.2 Flow Rounding

We simulate the flow rounding algorithm by Cohen [Coh95] in the CONGEST model as a subroutine for maximum flow and min-cost flow. The algorithm by Cohen [Coh95] is summarized as Algorithm 11.

**Lemma 7.5** (Proposition 5.3 of [Coh95]). Let $G = (V, E)$ be a graph with $n$ vertices and $m$ edges, $f : E \rightarrow \mathbb{R} \geq 0$ be a $s$-$t$ flow function, and $\Delta$ be a real value such that $1/\Delta$ is a power of 2 and $f(e)$ is an integral multiplication of $\Delta$ for every $e \in E$. Then

1. Algorithm `FLOWROUNDING` rounds $f$ on edge $e \in E$ to $\lfloor f(e) \rfloor$ or $\lceil f(e) \rceil$ such that the resulted flow has the total flow value not less than $f$.

2. If the total flow value of $f$ is integral and there is an integral cost function $c : E \rightarrow \mathbb{Z} \geq 0$, then Algorithm `FLOWROUNDING` rounds $f$ on edge $e \in E$ to $\lfloor f(e) \rfloor$ or $\lceil f(e) \rceil$ such that the resulted flow has the total flow value not less than $f$, and the total cost not more than $f$.

**Algorithm 11: Flow Rounding**

```plaintext
procedure FLOWROUNDING(G, s, t, f, c, \Delta)
if the total flow of f is not integral then
    Add an edge from t to s with flow value the same as total flow.
while \Delta < 1 do
    E' ← \{(u, v) \in E : f(u, v)/\Delta is odd\}
    Find an Eulerian partition of E' (ignoring the directions of the edges)
    for every cycle of the Eulerian partition of E' do
        if cycle contains the edge (t, s) then
            Traverse the cycle such that edge (t, s) is a forward edge.
        else if cost function c exists then
            Traverse the cycle such that the sum of costs on forward edges is no more than the sum of costs on backward edges.
        else
            Traverse the cycle arbitrarily.
    for every edge (u, v) E' do
        if (u, v) is a forward edge w.r.t the traversal of the path containing (u, v) then
            f(u, v) ← f(u, v) + \Delta
        else
            f(u, v) ← f(u, v) - \Delta
    \Delta ← 2\Delta
return f.
```
In this section, we present two distributed simulations of Cohen’s algorithm, one for maximum flow rounding, and another one for min-cost flow rounding. The underlying reason of two algorithms is that the strategy of choosing path directions (Line 8-11 of Algorithm 11) are different: the former one always choose direction that does not decrease the flow value, and later one always chooses the direction that does not increase total cost.

**Lemma 7.6.** Let $G = (\overrightarrow{V}, \overrightarrow{E})$ be a communication network with $\overrightarrow{m}$ vertices, $\overrightarrow{m}$ edges and diameter $D$, $G = (V, E)$ be a flow network containing two vertices $s$ and $t$ such that $G$ is a $((\rho, \alpha)$-flow-preconditioned minor distributed to $G$, $\Delta$ be a real value such that $1/\Delta$ is an integer that is a power of 2, and (for two vertices $s, t \in \overrightarrow{V}$) $\overrightarrow{f}$ : be a $s$-$t$ flow function on $G$ such that $f(e)$ is an integral multiplication of $\Delta$ for every $e \in \overrightarrow{E}$, and $\overrightarrow{f}$ is distributed to $\overrightarrow{G}$. Then

1. There is a distributed algorithm which runs in $O((\rho \sqrt{\overrightarrow{m}}(\log \overrightarrow{m})^2 + D) \cdot \log(1/\Delta))$ rounds to compute an integer $s$-$t$ flow function $\overrightarrow{f}' : E \rightarrow \mathbb{Z}^+$ such that the flow value of $\overrightarrow{f}'$ at least that of $\overrightarrow{f}$ and $\overrightarrow{f}'(e) \in \{\lfloor f(e) \rfloor, \lceil f(e) \rceil \}$ for every $e \in \overrightarrow{E}$.

2. If the total flow value of $\overrightarrow{f}$ is integral and there is an integral cost function $c$, then there is a distributed algorithm $O((\rho \sqrt{\overrightarrow{m}}(\log \overrightarrow{m})^2 + D) \cdot \log(1/\Delta))$ rounds to compute an integer $s$-$t$ flow function $\overrightarrow{f}' : E \rightarrow \mathbb{Z}^+$ such that the resulted flow has the total flow value not less than $\overrightarrow{f}$, and the total cost not more than $\overrightarrow{f}$.

Proof. We first extend $G$ such that $S_{\overrightarrow{G} \rightarrow \overrightarrow{G}}(s) = S_{\overrightarrow{G} \rightarrow \overrightarrow{G}}(t) = \overrightarrow{V}$. The resulted $G$ is a $(\rho + 2, \alpha + 2)$-flow-preconditioned minor distributed on $\overrightarrow{G}$. Then in $O(1)$ rounds, we can add edge $(t, s)$ to $G$ by letting $E_{\overrightarrow{G} \rightarrow \overrightarrow{G}}(t, s) = \overrightarrow{v}$ for an arbitrary vertex $\overrightarrow{v} \in \overrightarrow{V}$. The resulted $G$ is still a $(\rho + 2, \rho + 2)$-flow-preconditioned minor distributed on $\overrightarrow{G}$.

Throughout the algorithm, we view the flow function as a vector on $E$, denoted as $\overrightarrow{f}$, that is distributed to $\overrightarrow{G}$.

For a $\overrightarrow{f}$ and a fixed $\Delta$, $\overrightarrow{E}'$ which is an edge set of $G$ can be identified in $O(1)$ rounds. Now we show that an Eulerian partition of $\overrightarrow{E}'$ can be determined in $O((\alpha + 2)D)$ rounds. By the first and second condition of the lemma, for any vertex $v^G \in V$, the number of edges in $\overrightarrow{E}'$ incident to $v^G$ is always even. Hence, to construct an Eulerian partition of $\overrightarrow{E}'$, we only need to pair all the incident edges in $\overrightarrow{E}'$ for each vertex of $V$. This pairing process can be done in $O(1)$ rounds for all the vertices $v^G$ such that $|V_{\overrightarrow{G} \rightarrow \overrightarrow{G}}(v^G)| = 1$. For each vertex $v^G \in V$ such that $V_{\overrightarrow{G} \rightarrow \overrightarrow{G}}(v^G) = \overrightarrow{V}$, we simulate the following algorithm in $O(D)$ rounds such that the pairing process are simulated:

- Run a reverse BFS on $T_{\overrightarrow{G} \rightarrow \overrightarrow{G}}(v^G)$ such that for each $v^G \in \overrightarrow{V}$: Let $E'_{\overrightarrow{v} \overrightarrow{v}}$ be the union of the edges sent to $v^G$ from all the children of $v^G$ in $T_{\overrightarrow{G} \rightarrow \overrightarrow{G}}(v^G)$ and all the edges in $\overrightarrow{E}'$ incident to $v^G$ whose images by $E'_{\overrightarrow{G} \rightarrow \overrightarrow{G}}$ are edges incident to $v^G$. If $|E'_{\overrightarrow{v} \overrightarrow{v}}|$ is even, then pair all the edges in $E'_{\overrightarrow{v} \overrightarrow{v}}$ arbitrarily, otherwise, send one edge of $E'_{\overrightarrow{v} \overrightarrow{v}}$ to the parent of $v^G$ in $T_{\overrightarrow{G} \rightarrow \overrightarrow{G}}(v^G)$, and pair the remaining edges of $E'_{\overrightarrow{v} \overrightarrow{v}}$ arbitrarily.

In addition, since for every vertex $v^G$ of $T_{\overrightarrow{G} \rightarrow \overrightarrow{G}}(v^G)$, every child of $v^G$ with respect to $T_{\overrightarrow{G} \rightarrow \overrightarrow{G}}(v^G)$ sends the information of at most one edge to $v^G$, the Eulerian partition of $\overrightarrow{E}'$ corresponds to a union of cycles of $\overrightarrow{G}$ such that every edge of $\overrightarrow{G}$ is used at most $\rho + 2$ times.

Lemma 7.4 gives that for each cycle in the Eulerian partition, aggregating the required information takes $O(\overrightarrow{m}^{1/2} \log \overrightarrow{m} + D)$ rounds. And since all the cycles are edge disjoint for $G$, the traversal of each
cycle of the Eulerian partition can be determined in \( O(\rho \sqrt{m} \log m + D) \) rounds, and the direction of the traverse of each cycle can be broadcasted to each edge of the cycle in \( O(\rho \sqrt{m} \log m + D) \) rounds.

Hence, simulating one iteration of the while loop on Line 4 of Algorithm \textsc{FlowRounding} takes \( O(\rho \sqrt{m} \log m + D) \) rounds. So the overall number of rounds needed for Algorithm \textsc{FlowRounding} is \( O((\rho \sqrt{m} \log m + D) \cdot \log(1/\Delta)) \).

\section{Maximum Flow}

In this subsection, we present a distributed exact maximum flow algorithm for flow network with integral capacity in \( \tilde{O}(m^{3/7}U^{1/7}m^{1/1}D^{1/4} + D) \) rounds in the CONGEST model, where \( U \) is upper bound of the capacities among all the edges. Based on the distributed Laplacian solver, our algorithm simulate the sequential exact maximum flow algorithm by Madry [Mad16]. Madry’s sequential algorithm is briefly summarized in Algorithm 12, and the details are given in Section C.1.

**Theorem 6** ([GU15, CM20]). Let \( G \) be a (undirected or directed) graph with \( n \) vertices, \( m \) edges and undirected diameter \( D \), \( s \) be a vertex of \( G \), and \( \vec{w} \) be an edge vector such that for any edge \((u,v)\) of \( G \), vertices \( u \) and \( v \) know \( w_{u,v} \). Assume in each round, two vertices can send \( O(\log n) \) bit information to each other if there is an edge between them in \( G \) no matter the direction of the edge. Then there is a distributed SSSP algorithm that in \( \tilde{O}(n^{1/2}D^{1/4} + D) \) rounds computes the distances with respect to \( \vec{w} \) from \( s \) to all of its reachable vertices as well as an implicit shortest path tree rooted at \( s \) such that every vertex knows its parent in the shortest path tree.

In our distributed maximum flow and min-cost flow algorithm, the graph which we run single source shortest path (SSSP) algorithm on is different to the communication network, because the algorithms we want to simulate [Mad16, CMSV17] add additional vertices and edges to the graph. Hence, we show that this SSSP algorithm can be simulated efficiently if the graph is a flow preconditioned minor distributed to the communication network.

**Corollary 7.7.** Let \( \overline{G} = (\overline{V}, \overline{E}) \) be a communication network with \( \overline{m} \) vertices and \( \overline{m} \) edges, and \( G = (V, E) \) be a (undirected or directed) graph with \( n \) vertices, \( m \) edges and diameter \( D \) that is \((\rho, \alpha)\)-flow-preconditioned minor distributed to \( \overline{G} \). Then there is a distributed SSSP algorithm that, for any given source vertex \( s^G \in V \), performs \( \tilde{O}(\rho(\overline{m}^{1/2}D^{1/4} + D) \cdot \alpha^2) \) rounds.

**Proof.** We assume that excluding the \( \alpha \) vertices of \( G \) that are mapped to all the vertices of \( \overline{G} \), at most one vertex is mapped to any vertex of \( \overline{V} \). This is without loss of generality, because if multiple vertices are mapped to the same vertex of \( \overline{G} \), then any communication between these vertices are free. In the following, we will call vertices of \( G \) that are mapped to all the vertices of \( \overline{G} \) simulated vertices.

We first explain how the SSSP algorithm of [CM20] can be modified to work as an \( s-t \) shortest path algorithm in a setting where only the source (start) vertex \( s \) and the sink (target) vertex \( t \) (and their incident edges) are simulated vertices. In particular, we argue that the algorithm can be simulated in \( \tilde{O}(\rho(n^{1/2}D^{1/4} + D)) \) rounds (where the multiplicative \( \rho \) factor simply comes from the fact that every edge of \( \overline{G} \) is used at most \( \rho \) times for edges in \( G \)).

The algorithm of [CM20] consists of eight steps. In Steps 1 and 2, vertices sample themselves with certain probabilities. These steps require no communication and therefore can also be carried out in our setting in which source and sink are just simulated. We slightly modify Step 1 to ensure that the vertex \( t \) is never sampled. This does not affect the correctness of the algorithm if we are only interested
in computing the shortest path from \( s \) to \( t \) as shortest paths are simple and thus \( t \) will never be an inner vertex on this shortest path.

In Steps 4 and 7, certain auxiliary graphs are created implicitly in the sense that each vertex only knows its incident edges in the auxiliary graph and their respective edge weights. Therefore these steps also require no communication and therefore can also be carried out in our setting in which source and sink are just simulated.

To implement the rest of the algorithm we will rely on the following observation: whenever a step of the algorithm is performed solely by broadcasting or aggregating values via a global BFS tree of the network, then this step immediately can be carried out in our setting with the two simulated vertices as well. This is the case in Steps 5 and 6 of the algorithm.

In Step 8, a certain number of iterations of the Bellman-Ford algorithm is performed on a graph that in addition to the edges of the input graph contains edges from \( s \) to certain other vertices. Similar to [CM20], we carry out the first iteration of the Bellman-Ford algorithm – in which the neighbors of \( s \) set their tentative distance to the weight of the edge from \( s \) – by a global broadcast in \( O(D) \) rounds. In [CM20], the remaining iterations of Bellman-Ford are carried out in the standard way where vertices directly communicate with their neighbors. For our modification of [CM20] we do the same, but ignore the vertex \( t \) for these iterations. In the end, we explicitly need to ensure that the simulated vertex \( t \) also gets to know its distance from \( s \). We achieve this by additionally performing one iteration of the Bellman-Ford iteration in which only the incoming edges of \( t \) (and the corresponding neighbors of \( t \)) are considered. This can be carried out in \( O(D) \) rounds by broadcasting. This works because the incoming neighbors of \( t \) (which are part of the communication network and are not just simulated by it) already know their distance from \( s \) at this stage due to the previous iterations of Bellman-Ford.

This leaves only Step 3 of the algorithm. In Step 3, Lemma 2.4 of [FN18] is applied to compute approximate distances from each vertex of a set \( S \) (where \( S \) includes the vertex \( s \), but not the vertex \( t \)). Essentially this Lemma amounts to running a “weighted” version of the breadth-first-search algorithm for each vertex of \( S \) (which is repeated \( O(\log(nW)) \) times with a certain weight rounding applied to the edges in each iteration). The start times of these BFS algorithms are chosen with random delay to guarantee that the congestion at each vertex is low. For the BFS starting at vertex \( s \), the first iteration can be carried out by broadcasting the random delay of \( s \). The neighbors of \( s \) (knowing the weight of the edge from \( s \)) then know when the “weighted” BFS of \( s \) reaches them and can continue with it at the respective time. This gives an additional additive term of \( O(D) \) in the running time, which does not affect the asymptotic bounds stated in Lemma 2.4 of [FN18]. This concludes our discussion of the \( s \)-\( t \) shortest path algorithm.

Now observe that with the same approach we can obtain an SSSP algorithm in a setting where the source vertex \( s \) is the only vertex simulated by the network: we simply need to remove the special handling we had for vertex \( t \) in our approach above.

Note that these two algorithmic primitives are sufficient to compute SSSP in a setting where there are \( \alpha \) simulated vertices: Let \( S \) denote the set of vertices consisting of \( s^G \) and the simulated vertices. First, perform an SSSP computation from each vertex \( s \in S \) ignoring the other vertices of \( S \) (i.e., perform the SSSP computation in the graph \( G \setminus S \cup \{s\} \)). Then, perform an \( s \)-\( t \) shortest path computation for each pair of vertices \( s, t \in S \), ignoring the other vertices of \( S \) (i.e., perform the \( s \)-\( t \) shortest path computation in the graph \( G \setminus S \cup \{s, t\} \)). Now each vertex \( v \) can reconstruct its distance from \( s^G \) by the information it stored so far as the shortest path from \( s^G \) to \( v \) can be subdivided into subpaths between vertices of \( S \) containing no other vertices of \( S \). Overall, we perform \( O(\alpha) \) SSSP computations with at most one simulated vertex and \( O(\alpha^2) \) \( s \)-\( t \) shortest path computations with at most two simulated vertices.
vertices. Finally, note that as soon as each vertex $v$ knows its distance from $s^G$ an implicit shortest path tree (in which each vertex knows its parent in the tree) can be reconstructed by performing a single iteration of the Bellman-Ford algorithm. For the $\alpha$ simulated vertices, we perform this final step in $O(\alpha D)$ rounds by broadcasting via a global BFS tree.

**Theorem 7.** Let $G = (V, E)$ be a communication network with $\pi$ vertices, $\pi$ edges, and diameter $D$, $G_0$ be a graph and $c$ be an integral capacity function for each edge of $G_0$ with maximum capacity $U$ satisfying one of the following two conditions:

1. $G_0$ is the same as $G$, and for each edge $(u, v) \in E$, $u$ and $v$ know the capacity of edge $(u, v)$.

2. $G_0$ is a directed graph obtained by associating each edge of $G$ a direction such that for each edge $(u, v) \in E$, $u$ and $v$ know the direction of edge $(u, v)$ and its capacity.

Then there is a distributed algorithm to compute exact $s$-$t$ maximum flow for two vertices $s$ and $t$ of $G_0$ in $\tilde{O}(\pi^{3/7} U^{1/7} \pi^{\alpha(1)}(\pi^{1/2} D^{1/4} + D) + \pi^{1/2})$

rounds in the CONGEST model.

**Proof.** We simulate Algorithm 12. By [Mad16], the accuracy required throughout the algorithm is $1/\poly(\pi)$. Without loss of generality, we assume all the values throughout multiplied by $2^\gamma$ are integers for some $\gamma = O(\log \pi)$. Throughout the algorithm, we set $V_{\map}^{G_0 \to G}(s) = V_{\map}^{G_0 \to G}(t) = \overline{V}$, and for each vertex $v \in \overline{V} \setminus \{s, t\}$, $V_{\map}^{G_0 \to G}(v) = v$.

In the preconditioning edge step, we need to add $m$ parallel $(t, s)$ edges each with capacity $U$. This step can be simulated in $O(1)$ rounds by specifying an arbitrary vertex in $u^G \in \overline{V}$ such that $m$ parallel $(t, s)$ edges are mapped to selfloops of $u^G$.

In the initialization step, every edge $(u, v)$ with capacity $\alpha$ of $G_0$ is replace by three edges $(u, v)$, $(s, u)$ and $(v, t)$ with capacity $\alpha$. Let $G$ denote the graph after the preconditioning edge step and the initialization step. We always make sure that $E_{\map}^{G \to G}(s, u) = (u, u)$ and $E_{\map}^{G \to G}(v, t) = (v, v)$. Hence, $G$ is a $(3, 2)$-flow-preconditioned minor distributed to $G$.

In the progress step, if we replace an edge $(u, v)$ of $G$ by a path $(u, v_1, v_2, \ldots, v_\ell, v)$ in the Boosting subroutine, then we consider the following two cases:

1. If $E_{\map}^{G \to G}(u, v)$ is a selfloop of some vertex $x$ in $G$, then add all the new vertices $v_1, v_2, \ldots, v_\ell$ such that $V_{\map}^{G \to G}(v_i) = x$, and all the new edges are also selfloops on $x$.

2. If $E_{\map}^{G \to G}(u, v)$ corresponds to an edge of $G$, then add all the new vertices $v_1, v_2, \ldots, v_\ell$ such that $V_{\map}^{G \to G}(v_i) = V_{\map}^{G \to G}(u)$, and add edges such that $E_{\map}^{G \to G}(v_\ell, v) = E_{\map}^{G \to G}(u, v)$ and the remaining edges to be selfloops on $V_{\map}^{G \to G}(u)$.

Hence, we maintain the invariant that $G$ is $(3, 2)$-flow-preconditioned minor distributed to $G$.

Note that each execution of Augmentation, Fixing, and Boosting subroutines (see Appendix C.1 for details) takes a constant number of basic vector operations in Lemma 7.3 or solves a Laplacian system on graph $G$ (excepting replacing an edge by a path in the Boosting subroutine, which can be done in $O(1)$ rounds). By Lemma 7.3, each basic vector operation can be simulated in $O(D \log \pi)$ rounds.
Algorithm 12: MaxFlow($G_0, s, t, U, F$)

**Input:** directed graph $G_0 = (V, E_0, \vec{u})$ with each $e \in E_0$ having two non-negative integer capacities $u_e^-$ and $u_e^+$; $|V| = n$ and $|E_0| = m$; source $s$ and sink $t$; the largest integer capacity $U$; target flow value $F \geq 0$;

/* Preconditioning Edges */
1. Add $m$ undirected edges $(t, s)$ with forward and backward capacities $2U$ to $G_0$;

/* Initialization */
2. for each $e = (u, v) \in E_0$ do
3. replace $e$ by three undirected edges $(u, v), (s, v)$ and $(u, t)$ whose capacities are $u_e$;
4. Let the new graph be $G = (V, E)$;
5. Initialize $\vec{f} \leftarrow \vec{0}$ and $\vec{y} \leftarrow \vec{0}$;

/* Progress Step */
6. $\vec{f}, \vec{f}, \vec{y} \leftarrow \text{AUGMENTATION}(G, s, t, F)$;
7. Compute $\vec{\rho}$ by letting $\rho_e \leftarrow \frac{\vec{f} e}{\min\{u_e^+ - f_e, u_e^- + f_e\}}$;
8. $\vec{f}, \vec{y} \leftarrow \text{FIXING}(G, \vec{f}, \vec{y})$;
9. $\eta \leftarrow \frac{1}{14} - \frac{1}{4} \log m U - O(\log \log(mU))$, $\delta \leftarrow \frac{1}{m^{\frac{1}{2} - \eta}}$;
10. for $t = 1$ to $100 \cdot \frac{1}{\delta} \cdot \log U$ do
11. if $\|\vec{\rho}\|_3 \leq \frac{m^{\frac{1}{2} - \eta}}{3(1-\alpha)}$ then
12. $\delta \leftarrow \frac{1}{m^{\frac{1}{2} - \eta}}$;
13. $\vec{f}, \vec{f}, \vec{y} \leftarrow \text{AUGMENTATION}(G, s, t, F)$;
14. Compute $\vec{\rho}$ by letting $\rho_e \leftarrow \frac{\vec{f} e}{\min\{u_e^+ - f_e, u_e^- + f_e\}}$;
15. $\vec{f}, \vec{y} \leftarrow \text{FIXING}(G, \vec{f}, \vec{y})$;
16. else
17. let $S^*$ be the edge set that contains the $m^{4\eta}$ edges with the largest $|\rho_e|$;
18. $G \leftarrow \text{BOOSTING}(G, S^*, U, \vec{f}, \vec{y})$;
19. $\vec{f} \leftarrow \text{FLOWROUNDING}(G, \vec{f}, s, t)$;
20. while there is an augmenting path from $s$ to $t$ with respect to $\vec{f}$ for $G$ do
21. augment an augmenting path for $\vec{f}$;
To solve a Laplacian system, we first eliminate all the vertices that are added by the Boosting subroutine. Since all these vertices are of degree 2, this elimination can be done locally in each vertex of $\overline{G}$, and the resulted graph is $O(1)$-minor distributed to $G$. By Theorem 1, the Laplacian system can be solved in $\mathcal{O}(\log n \cdot (1/2 \log m + D))$ rounds. After the flow rounding, the difference between the flow value and maximum flow value is at most $O(\log m \cdot (1/2 \log m + D))$. By Corollary 7.7, each iteration of finding an augmenting path takes $\tilde{O}(D + \Omega(1/2 D^{1/4}))$ rounds. Hence, the additional augmenting step takes $O(\log m \cdot (1/2 \log m + D))$ rounds. 

### 7.4 Unit Capacity Minimum Cost Flow

In this subsection, we present a distributed minimum cost unit capacity flow algorithm with integral cost in $\tilde{O}(\log m \cdot (1/2 \log m + D))$ rounds in the CONGEST model. Based on the distributed Laplacian solver, our algorithm simulate the algorithm by Cohen et al. [CMSV17]. The sequential algorithm is briefly summarized in Algorithm 13, and the details of the algorithm are given in Section C.2.

**Theorem 8.** Let $\overline{G} = (\overline{V}, \overline{E})$ be a communication network with $m$ vertices and $\overline{m}$ edges, $G_0$ be a directed unweighted graph also defined on $\overline{V}$ such that each edge $(u, v)$ of $G_0$ is an edge of $\overline{E}$ if the direction of edge is ignored, and $u$ and $v$ in the communication network know the direction of edge $(u, v)$ and its cost. Then, given $G_0$ and a demand vector $\tilde{\sigma}$, there is a distributed algorithm to compute the minimum cost flow for graph $G_0$ with respect to $\tilde{\sigma}$ in $\tilde{O}(\log m \cdot (1/2 \log m + D))$ rounds in the CONGEST model.

**Proof.** We simulate Algorithm 13 and the accuracy required throughout the algorithm is $1/poly(m)$. Without loss of generality, we assume all the values throughout the algorithm multiplied by $2^\gamma$ are integers for some $\gamma = O(\log m)$.

To build graph $G_1$ as the INITIALIZATION subroutine required, we add vertex $v_{aux}$ to the graph such that $V_{map}^{G_1 \rightarrow \overline{G}}(v_{aux}) = \overline{V}$, and for all the vertices $v \in \overline{V}$, $V_{map}^{G_1 \rightarrow \overline{G}}(v) = v$. For each edge $(u, v)$ of $G_1$ with $u, v \in \overline{V}$, we let $E_{map}^{G_1 \rightarrow \overline{G}}(u, v) = (u, v)$. For edge $(v, v_{aux})$ or $(v_{aux}, v)$, we let $E_{map}^{G_1 \rightarrow \overline{G}}(v, v_{aux})$ to be a selfloop on $v$. The construction of $G_1$ can be done locally, and the resulted graph is a $(2, 1)$-flow-preconditioned minor distributed to $\overline{G}$.

To construct $G$, for each vertex $e_{uv} \in Q$, we map $e_{uv}$ to be one of $u$ and $v$ of $\overline{G}$ arbitrarily. For vertices of $P$, the mapping is the same as that of $G_1$. For each edge $(u, e_{uv})$ of $G$, if $V_{map}^{G \rightarrow \overline{G}}(u) = V_{map}^{G \rightarrow \overline{G}}(e_{uv})$, then we map edge $(u, e_{uv})$ to a selfloop on $V_{map}^{G \rightarrow \overline{G}}(u)$, otherwise, $(V_{map}^{G \rightarrow \overline{G}}(u), V_{map}^{G \rightarrow \overline{G}}(e_{uv}))$ is an edge of $\overline{G}$, and we set $E_{map}^{G \rightarrow \overline{G}}(u, e_{uv})$ to be the edge $(V_{map}^{G \rightarrow \overline{G}}(u), V_{map}^{G \rightarrow \overline{G}}(e_{uv}))$. Hence, the INITIALIZATION subroutine can be simulated in $O(1)$ rounds, and the resulting graph $G$ is a $(2, 1)$-flow-preconditioned minor distributed to $\overline{G}$.

To simulate Line 2, we add an additional vertex $v_0$ to $G$ such that $V_{map}^{G \rightarrow \overline{G}}(v_0) = \overline{V}$, and add edges $(v_0, v)$ for each $v \in P$ by setting $E_{map}^{G \rightarrow \overline{G}}(v_0, v)$ be a selfloop on $v$ if $v \in \overline{V}$, and $E_{map}^{G \rightarrow \overline{G}}(v_0, v)$ be a
Algorithm 13: MinCostFlow($G, \sigma, W$)

Input: directed graph $G_0 = (V_0, E_0, \tilde{c}_0)$ with each edge having unit capacity and cost $\tilde{c}_0$;
$|V_0| = n$ and $|E| = m$; integral demand vector $\sigma$; the absolute maximum cost $W$;
1. $G = (P \cup Q, E), \tilde{b}, \tilde{f}, \tilde{y}, \tilde{s}, \tilde{v}, \tilde{\mu}, c_p, c_T, \eta \leftarrow$ INITIALIZATION($G_0, \sigma$);
2. Add a new vertex $v_0$ and undirected edges $(v_0, v)$ for every $v \in P$ to $G$;
3. for $i = 1$ to $\varepsilon T \cdot m^{1/2 - 3\eta}$ do
4. for each $v \in P$ do
5. set resistance of edge $(v_0, v)$ for each $v \in P$ to be $r_{v_0 v} \leftarrow \frac{m^{1+2\eta}}{a(v)}$, where
6. for $j = 1$ to $m^{2\eta}$ do
7. while $\|\tilde{\rho}\|_{L_3} > c_p \cdot m^{1/2 - \eta}$ do
8. $\bar{\rho}, \bar{y}, \bar{s}, \bar{v} \leftarrow$ PERTURBATION($G, \bar{\rho}, \bar{f}, \bar{y}, \bar{s}, \bar{v}$);
9. $\bar{f}, \bar{s}, \bar{\rho}, \bar{\mu} \leftarrow$ PROGRESS($G, \sigma, \bar{f}, \bar{v}$);
10. Let $\tilde{b}^+$ be the demand vector corresponding to the current flow $\tilde{f}$ and construct the fractional flow vector $\tilde{f}^\leq$;
11. Add source $s$ and sink $t$ to $G$, and connect $s$ to each $v \in P$ with $f_s^\leq \leftarrow f^\leq(E(v))$, and connect each $v \in Q$ to $t$ with $f_t^\leq \leftarrow f^\leq(E(v))$ in $G$;
12. $\tilde{M} \leftarrow$ FLOWROUNDING($G, \tilde{f}^\leq, s, t$);
13. Remove $s, t$ from $G$ and related coordinates in $\tilde{f}^\leq$ and $\tilde{M}$;
14. for $i = 1$ to $\hat{O}(m^{3/7})$ do
15. Construct bidirectional graph $\tilde{G}_M$ using $G$, $M$ and $\tilde{c}$;
16. Set $F_M \leftarrow \{ v \in P \cup Q \mid M(v) < b_v \}$;
17. Compute a shortest path $\pi$ in $\tilde{G}_M$ from $P \cap M$ to $Q \cap M$;
18. for $u \in P \cup Q$ do
19. if $u$ can be reached from $P$ in $\tilde{G}_M$ then
20. if $u \in P$ then
21. $y_u \leftarrow y_u - D_{\tilde{G}_M}(P, u)$;
22. else
23. $y_u \leftarrow y_u + D_{\tilde{G}_M}(P, u)$;
24. Augment $\tilde{M}$ using the augmenting path $\pi$;
25. return $\tilde{M}$.
Algorithm 14: Initialization$(G, \tilde{\sigma})$

1. Create a new vertex $v_{aux}$ with $\sigma(v_{aux}) = 0$;
2. for each $v \in V_0$ do
   3. $t(v) \leftarrow \sigma(v) + \frac{1}{2} \deg_{\text{in}}^{G_0}(v) - \frac{1}{2} \deg_{\text{out}}^{G_0}(v)$;
   4. if $t(v) > 0$ then construct $2t(v)$ parallel edges $(v, v_{aux})$ with costs $\|\tilde{c}_0\|_1$;
   5. else if $t(v) < 0$ then $\ll$
   6. construct $|2t(v)|$ parallel edges $(v_{aux}, v)$ with costs $\|\tilde{c}_0\|_1$;
7. Let the new graph be $G_1 = (V_1, E_1, \tilde{c}_1)$;
8. Initialize the bipartite graph $G = (P \cup Q, E, \tilde{c})$ with $E \leftarrow \emptyset$, $P \leftarrow V_1$ and $Q \leftarrow \{e_{uv} \mid (u, v) \in E_1\}$ where $e_{uv}$ is a vertex corresponding to edge $(u, v) \in E_1$;
9. for each $(u, v) \in E_1$ do
   10. let $E \leftarrow E \cup \{(u, e_{uv}), (v, e_{uv})\}$ with $c(u, e_{uv}) = c_1(u, v)$ and $c(v, e_{uv}) = 0$, and set $b(u) \leftarrow \sigma(u) + \deg_{\text{in}}^{G_1}(u)$, $b(v) \leftarrow \sigma(v) + \deg_{\text{in}}^{G_1}(v)$ and $b(e_{uv}) \leftarrow 1$;
11. Initialize $\vec{f}, \vec{c}, \vec{\mu}, \tilde{\nu}, \tilde{\tau}, \eta$ based on $G, \tilde{c}, \tilde{b}$;
12. return $G, \tilde{b}, \vec{f}, \vec{c}, \vec{\mu}, \tilde{\nu}, \tilde{\tau}, c_T$ and $\eta$;

selfloop on an arbitrary vertex of $\overline{V}$ if $v = v_{aux}$. The resulting graph $G$ is a $(3, 2)$-flow-preconditioned minor distributed to $\overline{G}$.

For Line 3-9 of Algorithm 13, each execution of Perturbation and Progress subroutines (see Appendix C.2 for details) takes a constant number of basic vector operations in Lemma 7.3 or solves a Laplacian system on graph $G$. To solve a Laplacian system, since all the vertices of $Q$ have two incident edges, we eliminate vertices of $Q$ in $O(1)$ rounds, and the resulting graph is $O(1)$-minor distributed to $\overline{G}$. By Theorem 1, every Laplacian system can be solved in $\overline{O}(m^{2/7} + D)$ rounds. Based on the parameter setting, Line 3-9 can be simulated in $\overline{O}(m^{2/7} + (1/2 + D)\text{poly}(\log W))$ rounds.

Line 10 can be simulated locally. To simulate Line 11, we add $s$ and $t$ such that $V_{\text{map}}^G(s) = V_{\text{map}}^G(t) = \overline{V}$, and add edges by adding selfloops on vertices of $\overline{V}$. The resulting graph is a $(5, 4)$-flow-preconditioned minor distributed to $\overline{G}$. By Lemma 7.6, Line 12 can be simulated in $\overline{O}(m^{1/2} + D)$ rounds.

To simulate Line 14-24, all the operations except finding shortest path can be simulated in a way similar to that of Line 3-9. To find the shortest path from $P \cap F_M$ to $Q \cap F_M$, we add an additional vertex $v'$, and edges $(v', v)$ for each $v \in P \cap F_M$ with weight zero to $G$. The resulted graph is a $(4, 3)$-flow-preconditioned minor distributed to $\overline{G}$. By Corollary 7.7, the shortest path can be computed in $\overline{O}(D + m^{1/2}D^{1/4})$ rounds. The total number of rounds for Line 14-24 is $\overline{O}(m^{3/7}(D + m^{1/2}D^{1/4}))$ rounds.

Hence, the total number of rounds required is

$$\overline{O}(m^{3/7} + o(1))(m^{1/2} + D)\text{poly}(\log W)) + \overline{O}(m^{1/2} + D) + \overline{O}(m^{3/7}(D + m^{1/2}D^{1/4}))$$

$$= \overline{O}(m^{3/7} + o(1))(m^{1/2}D^{1/4} + D)\text{poly}(\log W)).$$

□
7.5 Negative shortest path

We now give a distributed algorithm for computing single source shortest path with negative weights. It is a direct use of the reduction from shortest paths with negative weights to min-cost flow by Cohen et al. [CMSV17]. Pseudocode of this algorithm is in Algorithm 15.

**Algorithm 15: ShortestPaths($G$, $s$, $W$)**

**Input:** directed graph $G = (V, E, w)$ with $|V| = n$ and $|E| = m$; source $s$; the absolute maximum weight $W$;

/* Reduction to a weighted perfect $\vec{1}$-matching problem */

1. Let the bipartite graph be $G_{12} = (V_1 \cup V_2, E_{12}, w_{12})$ with $V_1 = \{v_1 \mid v \in V\}$,
   $V_2 = \{v_2 \mid v \in V\}$, $E_{12} = \{u_1v_2 \mid uv \in E\} \cup \{v_1v_2 \mid v \in V\}$ and
   $w_{12}(u_1v_2) = \begin{cases} -w_{uv} & uv \in E, \\ 0 & u = v \end{cases};$

2. $(\vec{f}, \vec{y}) \leftarrow \text{MinCostFlow}(G_{12}, \vec{1}, W);$;

3. for each edge $(u, v) \in \overline{E}$ do
   4. $w_{uv} \leftarrow w_{uv} + y_u - y_v$;

5. Compute the shortest paths with source $s$ on $G = (V, E, w')$;

**Theorem 9.** Let $G = (V, E)$ be a communication network with $n$ vertices and $m$ edges, and $w : E \to \{-W, -W + 1, \ldots, -1, 0, 1, \ldots, W\}$ be an integral weight function. For a vertex $s \in V$, there is a distributed algorithm to compute the shortest path from $s$ to all the other vertices that has a shortest path from $s$ in

$$\tilde{O}\left(\frac{m^{3/7} + o(1)}{(n^{1/2}D^{1/4} + D)}\text{poly}(\log W)\right)$$

rounds in the CONGEST model.

**Proof.** The strategy of Algorithm 15 is that transferring the given $G$ to a bipartite graph $G_{12}$ and calling the Algorithm MinCostFlow on $G_{12}$ to obtain the dual solution $\vec{y}$, which is utilized to transform the original edge weights in $G$ to be non-negative, and then using the single source shortest path algorithm with non-negative weights on the new instance $G' = (V, E, w)$ to compute the shortest paths.

In constant round, we can construct $G_{12}$ based on $G$ that is a $(1, 0)$-flow-preconditioned minor distributed to $G$. By Theorem 8 and Corollary 7.7, Algorithm 15 can be simulated in rounds.

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A Lower Bound

Theorem 2. In the CONGEST model of computation, solving Laplacian systems to accuracy $\epsilon \leq \frac{1}{2}$ requires at least $\tilde{\Omega}(n^{1/2} + D)$ rounds of communication.

Proof. Note first that any low-accuracy solver with accuracy $\epsilon \leq \frac{1}{2}$ can be boosted to a solver with precision $\tilde{\epsilon}$ at the cost of increasing the running time by a factor of $O(\log \tilde{\epsilon}^{-1})$ (see Lemma 1.6.8. in [Pen13]). This is done by running $O(\log \tilde{\epsilon}^{-1})$ iterations of the iterative refinement method, which in each iteration performs one matrix-vector product, one vector subtraction, one vector addition, and one call to the low-accuracy solver. Excluding the call to the low-accuracy solver, all of these operations can be performed in a constant number of rounds. The running time of the iterative refinement method is therefore dominated by the $O(\log \tilde{\epsilon}^{-1})$ calls to the low-accuracy solver. We thus assume in the following that we are given a high-accuracy solver with accuracy $\epsilon = 1/poly(n)$, which as just argued requires only an overhead of $O(\log n)$ in the number of rounds compared to a low-accuracy solver.

To prove the lower bound we use the framework of Das Sarma et al. [SHK+12] who established an $\Omega(\sqrt{n}/(\log n) + D)$ lower bound for the following verification problem: Given a subgraph $H$ of the communication network $\overline{G}$ (which has $n$ nodes and diameter $D$) and two nodes $s$ and $t$, the network needs to decide whether $s$ and $t$ are connected (i.e., lie in the same connected component). In their lower bound construction, the distance between $s$ and $t$ in $\overline{G}$ is $O(\log n)$. We show that any algorithm for solving Laplacian Systems up to small enough error $\epsilon = 1/poly(n)$ can be used to give $s$ and $t$ the information whether they are connected in $H$ in additional $\text{dist}(s,t) = O(\log n)$ rounds. In particular, we exploit that such a solver can be used to compute an approximation to the effective $s$-$t$ resistance.

Define the weighted graph $H'$ (which we view as a resistor network) as having the same nodes and edges as $\overline{G}$ and resistances $r_e = 1$ for every edge $e \in E(H)$ and $r_e = n$ for every edge $e \notin E(H)$. Let $\chi_{s,t}$ be the $n$-dimensional vector 1 at the coordinate corresponding to $s$, -1 at the coordinate corresponding to $t$, and 0 otherwise. It is well known (see, e.g., [CKM+11, Vis12]) that $\text{res}_{H'}(s,t) =$
Thus, resistance in special vertices, picked randomly so that any vertex is within a distance of about $\sqrt{n}$, is upper-bounded by the length of the shortest path between them. Henceforth let $\phi = L(H')^\dagger \chi_{s,t}$, where $L(H')^\dagger$ is the right inverse of $L(H')$.

The combination of (7) and (8) together with the estimate $\text{res}_{H'}(s,t) \leq nR$ gives

$$\left\| \vec{\phi}' - \phi \right\|_\infty \leq \epsilon \cdot \text{poly}(n).$$

By setting $\epsilon$ to a small enough value inversely polynomial in $n$, (9) implies

$$\text{res}_{H'}(s,t) - 0.25 \leq \phi'(s) - \phi'(t) \leq \text{res}_{H'}(s,t) + 0.25.$$

In the rest of the proof, we argue that knowing the value $\phi'(s) - \phi'(t)$ suffices to decide whether $s$ and $t$ are connected in $H$. If $s$ and $t$ are connected in $H$, then – since effective resistances obey the triangle inequality – the effective $s$-$t$ resistance in $H'$ is upper-bounded by the length of the shortest path between $s$ and $t$ in $H$, i.e., $\text{res}_{H'}(s,t) \leq n - 1$. If $s$ and $t$ are not connected in $H$, then let $S$ be the connected component containing $s$ and let $e_1, \ldots, e_k$ (for some $k \leq m \leq n^2$) be the edges leaving $S$ in $H$. By the Nash-Williams inequality (see, e.g., [LP16], chapter 2.5), the effective $s$-$t$ resistance is at least

$$\text{res}_{H'}(s,t) \geq \frac{1}{\sum_{i=1}^k \frac{1}{v(e_i)}} = \frac{1}{\sum_{i=1}^k \frac{1}{(1+\epsilon)^2 n^3}} \geq (1 + \epsilon)^2 n.$$ 

Thus, $s$ and $t$ are connected in $H$ if and only if $\phi'(s) - \phi'(t) \leq n - 0.5.$

### B Building Blocks for Distributed Minors

Here we give the deferred proofs from Section 4.1. All of our algorithms are based on placing about $\sqrt{n}$ special vertices, picked randomly so that any vertex is within a distance of about $\sqrt{n}$ from these, and then aggregating information at these special vertices globally via a DFS tree in about $\sqrt{n} + D$ rounds.

For this, it is useful to define a tree decomposition scheme for the set of overlapping trees used to connect the supervertices.
Lemma B.1. There is an algorithm \textit{SpecialVertices} that takes an input a forest \( F \) specified with mappings of vertices and edges into a graph \( \overline{G} \) such that each vertex of \( \overline{G} \) appears in at most \( \rho \) trees of \( F \), and each edge of \( \overline{G} \) is used in at most \( \rho \) trees of \( F \), and returns after \( O(\rho \sqrt{n}) \) rounds of communication a collection of \( O(\rho \sqrt{n} \log n) \) special vertices of \( F \), labeled at their mapped vertices in \( \overline{G} \), such that with high probability, for each \( T \) in \( F \), we have:

1. either the diameter of \( T \) is at most \( \sqrt{n} \),
2. or for any vertex of \( T \),
   (a) it can reach at most 2 special vertices, without going through more special vertices.
   (b) its distance in \( T \) to closest special vertex is at most \( O(\sqrt{n}) \).

\textbf{Algorithm 16:} Partition all trees of a forest into low diameter pieces via special vertices

\begin{algorithm}
\begin{algorithmic}
  \Procedure{SpecialVertices}{$F, \overline{G}$}
    \State Sample \( K \) by including each vertex in each tree of \( F \) with probability \( \log n/n \).
    \For{\( O(\sqrt{n}) \) rounds}
      \State Each vertex propagate to all its neighbors whether taking that edge towards it leads to a vertex in \( K \).
      \State Add all vertices that can reach special vertices in three or more directions to \( K \).
    \EndFor
  \EndProcedure
\end{algorithmic}
\end{algorithm}

Proof. Consider \textit{SpecialVertices} in Algorithm 16.

The congestion bound gives that the total number of vertices among the trees is at most \( O(\rho n) \). This means picking \( O(\rho \sqrt{n} \log n) \) random vertices from

\[ \bigcup_{v^G \in V(G)} S_G^{\overline{G}}(v^G), \]

ensures that with high probability, the maximum distance to a special vertex along any tree path is \( O(\sqrt{n}) \) with high probability.

This means that in \( O(\sqrt{n}) \) rounds of propagation, we can find, for each edge in each tree, whether there is a special vertex in either direction. Formally, the local operation at each vertex is to check whether there are 2 or more directions from it that lead to special vertices: if there are, then all edges entering this vertex are part of paths that reach special vertices. Otherwise, all except that one direction that the path from a special vertex came from can be continued, so we push ‘possible’ along all except that direction.

By declaring all vertices with three or more edges leaving it that lead to special vertices as special themselves. As this only adds in the lowest common ancestors of the previous special vertices, it only increases the number of special vertices by a constant factor. This step is also completely local, so the total cost is dominated by the propagation steps.

This partition routing allows us to root all of the trees.
Proof. (of Lemma 4.2) We first run the partition scheme SpecialVertices on the forest that’s the unions of the spanning trees of the supernodes of \( G \). Lemma B.1 gives that each resulting piece consisting of edges reachable to each other without going through special vertices have diameter at most \( O(\sqrt{n}) \). So \( O(\sqrt{n}) \) rounds of propagation lets the root vertex inform all nodes in its piece. In this number of rounds, we also propagate the ID of these special vertices, as well as distances to them, to all vertices in each piece using another \( O(\sqrt{n}) \) rounds of communication.

Note this in particular allows the two special nodes on each piece to know their distance to each other. The amount of information aggregated at each special vertex may be large. Aggregating these information centrally along the BFS tree of \( \overline{G} \) then allows us to find the distance from the root vertex to all the special vertices in their piece in a centralized manner. Once this information is propagated back, each edge can just be oriented towards the direction of the special vertex closer to the root, giving the desired orientations.

This direction to the root is necessary for propagating information that cannot be duplicated, such as the sum of values. Using it, we can prove our main communication tool, Lemma 4.3.

Proof. (of Lemma 4.3) Once again we run the partition scheme SpecialVertices. Lemma B.1 ensures that all paths hit one such vertex after at most \( O(\sqrt{n}) \) steps. After that, we root the tree using Lemma 4.2.

For the push case, we repeatedly push information from vertices to their neighbors. Each such step costs \( O(\rho) \) due to the vertex congestion bound. By the bound above, after \( O(\sqrt{n}) \) such steps (which costs a total of \( O(\rho\sqrt{n}) \) rounds, this information either reached all nodes in the corresponding supernode, or some special vertex.

Getting all info on special nodes to a center node (along with the ID of \( v^G \) that they originated from) over a BFS tree takes \( O(\rho\sqrt{n}\log n + D) \) rounds, after which they can also be re-distributed to all special nodes. Then by the distance bound from Lemma B.1 another \( O(\rho\sqrt{n}) \) rounds of propagations to neighbors passes the information to everyone.

The aggregation of sum or minimum follows similarly. We repeat \( O(\rho\sqrt{n}) \) rounds of all non-special vertices propagating their sum, or min, up to their parents. In case of sum, once a value ‘floats’ to its parent, it’s set to 0 at the current vertex so we do not over count. Finally, all the information at the special vertices are aggregated via the global BFS tree, and passed to the corresponding root vertices.

Proof. (of Lemma 4.6) Let

\[
S_{G_1 \rightarrow G_2} : V(G_1) \rightarrow V(G_2)^*
\]

\[
V_{map}^{G_1 \rightarrow G_2} : V(G_1) \rightarrow V(G_2)
\]

\[
T_{G_1 \rightarrow G_2} : V(G_1) \rightarrow E(G_2)^*
\]

\[
E_{map}^{G_1 \rightarrow G_2} : E(G_1) \rightarrow E(G_2)
\]
be the maps from $G_1$ to $G_2$, and similarly, let the mapping from $G_2$ to $G$ be:

\[
\begin{align*}
S^{G_2 \rightarrow G} : V(G_2) &\rightarrow V(G) \ast \\
V_{\text{map}}^{G_2 \rightarrow G} : V(G_2) &\rightarrow V(G) \\
T^{G_2 \rightarrow G} : V(G_2) &\rightarrow E(G) \ast \\
E_{\text{map}}^{G_2 \rightarrow G} : E(G_2) &\rightarrow E(G)
\end{align*}
\]

We will construct the mapping from $G$ to $G$ from these. The edge and vertex maps are directly by composition:

\[
\begin{align*}
V_{\text{map}}^{G_1 \rightarrow G} (v^{G_1}) &= V_{\text{map}}^{G_2 \rightarrow G} (V_{\text{map}}^{G_1 \rightarrow G_2} (v^{G_1})) \\
E_{\text{map}}^{G_1 \rightarrow G} (e^{G_1}) &= E_{\text{map}}^{G_2 \rightarrow G} (E_{\text{map}}^{G_1 \rightarrow G_2} (e^{G_1}))
\end{align*}
\]

The edge mapping is a direct transfer of the pre-images, locally per edge. While the vertex label propagation is via one round of communication along new supervertex, via Lemma 4.3.

So we can focus on the construction of new supervertices and their spanning trees. For $v^{G_1} \in V(G_1)$, we let

\[
S^{G_1 \rightarrow G} (v^{G_1}) = \bigcup_{v^{G_2} \in S^{G_1 \rightarrow G_2} (v^{G_1})} S^{G_2 \rightarrow G} (v^{G_2})
\]

with corresponding spanning tree a subset of the edges

\[
\bigcup_{v^{G_2} \in S^{G_1 \rightarrow G_2} (v^{G_1})} T^{G_2 \rightarrow G} (v^{G_2}) .
\]

To compute this union we have each root vertex of each supervertex corresponding to some $v^{G_2}$ inform the entire supervertex of the new ID in $G_1$. As each vertex of $G_2$ corresponds to the image of at most $\rho_1 v^{G_1}$s, this mapping takes $O(\rho_1 \rho_2 (\sqrt{n \log n} + D))$ iterations.

Then the edges of $G$ with the new labels (of vertex ID from $G_1$) gives the corresponding supervertices. That is, $S^{G_1 \rightarrow G} (v^{G_1})$ is simply the set of vertices that received the label $v^{G_1}$ after we propagated from the root vertices of $G_2$ in $G$.

We then need to find spanning forests among these unions of trees. We do so with a variant of Brouvka’s algorithm, combined with parallel tree contraction. Specifically, we iterate the following $O(\log n)$ times:

1. Each remaining vertex (in each of the super vertices) pick a random priority.
2. Each vertex identify highest priority neighbor, computed on the minor with some edges already contracted using Lemma 4.3.
3. Contract either:
   a) all the leaf vertices (degree 1 nodes),
   b) a vertex disjoint subset of edges with both end points having degree 2.
In the second case of contracting edges with both endpoints having degree 2, we can find an independent set of such edges whose total size is at least a constant factor of all such edges by simply picking a random subset with probability $1/5$, and dropping the ones where another endpoint is picked.

This procedure reduces the number of vertices in each component a constant factor in expectation. This is because the first step ensures the edges found form a tree with edge count at least half the number of vertices. Then parallel tree contraction [MR89] ensures that either the number of leaves, or the number of edges with both endpoints degree 2 is at least a constant factor of the tree size. Contracting the larger set of these then gives the desired constant factor progress, so it terminates in $O(\log n)$ rounds.

Note that in subsequent rounds, the edges already identified to be part of $T^{G_1 \rightarrow G_\ell'(v^G)}$ form a spanning forest, and we’re working on the minor with this forest contracted. This means we need to invoke Lemma 4.3 repeatedly to do the neighborhood aggregations on this contracted graph.

Proof. (of Corollary 4.7) We want to simplify $F$ into a sequence of subgraphs that have simple 1-minor distributions into $G$. After that, we can invoke Lemma 4.6 to make progress.

We repeat the same tree contraction procedure used for finding spanning forests in Lemma 4.6 above. It gives that at each step, we’re computing $G/F$ for a set of vertex disjoint stars $F$.

Given such a $F$, we can then construct a 1-minor distribution of $G/F$ into $G$ by:

1. Having the center vertex generate the new vertex ID.
2. Propagate this ID to vertices in the corresponding supervertex in $G$.
3. Have edges that declared themselves part of $F$ pass this info from one end point to the other.
4. All leaf supervertices then pull this new vertex ID into their root vertex.

after which invoking Lemma 4.6 gives the minor. This halves the number of

C Max flow and minimum cost flow algorithm

In this section, we give the missing details of the max flow algorithm and minimum cost flow algorithm as Section 7.

C.1 Max flow algorithm

In this section, we give the missing subroutines of Algorithm 12.
Algorithm 17: **AUGMENTATION**\((G, s, t, F)\)

```plaintext
/* Augmentation Step */
1 For each \(e \in E\), let \(r_e \leftarrow \frac{1}{(u_e^s - f_e)^2} + \frac{1}{(u_e^t + f_e)^2}\) and \(w_e \leftarrow \frac{1}{r_e}\);
2 Solve Laplacian linear system \(L(G)\overrightarrow{\phi} = F \cdot \overrightarrow{\chi}_{s,t}\) where \(\overrightarrow{\chi}_{s,t}\) is the vector whose entry is \(-1\) (resp. 1) at vertex \(s\) (resp. \(t\)) and 0 otherwise;
3 For each \(e = (u, v) \in E\), let \(\overrightarrow{f}_e \leftarrow \overrightarrow{\phi}_v - \overrightarrow{\phi}_u\) and \(\overrightarrow{f}_e \leftarrow f_e + \delta \overrightarrow{f}_e\);
4 For each \(v \in V\), let \(\overrightarrow{y}_v \leftarrow y_v + \delta \overrightarrow{\phi}_v\);
5 return \(\overrightarrow{f}, \overrightarrow{f}, \overrightarrow{y}\);
```

Algorithm 18: **FIXING** \((G, \overrightarrow{f}, \overrightarrow{y})\)

```plaintext
/* Fixing Step */
1 For each \(e = (u, v) \in E\), let \(r_e \leftarrow \frac{1}{(u_e^s - f_e)^2} + \frac{1}{(u_e^t + f_e)^2}\) and \(w_e \leftarrow \frac{1}{r_e}\) and
   \[\theta_e \leftarrow w_e \left[ (\overrightarrow{y}_v - \overrightarrow{y}_u) - \left( \frac{1}{u_e^s - f_e} \right) \right] ;\]
2 \(\overrightarrow{f}' \leftarrow \overrightarrow{f} + \overrightarrow{\theta}\);
3 Let \(\overrightarrow{\delta}\) be \(\overrightarrow{\theta}\)'s residue vector;
4 For each \(e \in E\), let \(r_e \leftarrow \frac{1}{(u_e^s - f_e)^2} + \frac{1}{(u_e^t + f_e)^2}\) and \(w_e \leftarrow \frac{1}{r_e}\);
5 Solve Laplacian linear system \(L(G)\overrightarrow{\phi}' = -\overrightarrow{\delta}\);
6 For each \(e = (u, v) \in E\), let \(\theta'_e \leftarrow \frac{\overrightarrow{\phi}'_v - \overrightarrow{\phi}'_u}{r_e}\);
7 For each \(e \in E\), \(f_e \leftarrow f_e + \theta'_e\);
8 For each vertex \(v \in V\), \(y_v \leftarrow y_v + \phi'_v\);
9 return \(\overrightarrow{f}, \overrightarrow{y}\);
```
Algorithm 19: Boosting \((G, S^*, U, \vec{f}, \vec{y})\)

/* Boosting Step */
1 \textbf{for each} edge \(e = (u, v) \in S^*\) \textbf{do}
2 \hspace{1em} \(\beta(e) \leftarrow 2 + \left\lfloor \frac{2U}{\min\{u_e^{+} - f_e, u_e^{-} + f_e\}} \right\rfloor\);  
3 replace \(e\) with path \(u \xrightarrow{} v\) that consists of \(\beta(e)\) edges \(e_1, \cdots, e_{\beta(e)}\) oriented towards \(v\) and \(e_1, e_2 \leftarrow e;\)
4 \hspace{1em} \textbf{for} \(3 \leq i \leq \beta(e)\) \textbf{do}
5 \hspace{2em} \(u_{e_i}^+ \leftarrow +\infty\) and \(u_{e_i}^- \leftarrow \left(1 - \frac{1}{u_e^+ - f_e} - \frac{1}{u_e^- + f_e}\right)^{-1} (\beta(e) - 2) - f_e;\)
6 \hspace{1em} \textbf{for each} \(1 \leq i \leq \beta(e)\) \textbf{do}
7 \hspace{2em} \(y_{v_0} \leftarrow y_u;\)
8 \hspace{2em} \(y_{v_{\beta(e)}} \leftarrow y_v;\)
9 \hspace{2em} \(y_{v_1} \leftarrow y_v;\)
10 \hspace{2em} \(y_{v_2} \leftarrow y_v + \frac{1}{u_e^+ - f_e} - \frac{1}{u_e^- + f_e};\)
11 \hspace{1em} \textbf{for} \(3 \leq i \leq \beta(e)\) \textbf{do}
12 \hspace{2em} \(y_{v_i} - y_{v_{i-1}} = -\frac{1}{\beta(e) - 2} \left(\frac{1}{u_e^+ - f_e} - \frac{1}{u_e^- + f_e}\right);\)
13 \hspace{1em} \textbf{Update} \(G;\)

C.2 Unit Capacity Minimum Cost Flow Algorithm

We give the detailed unit capacity minimum cost flow algorithm proposed by Cohen et al. [CMSV17] in this subsection.

Algorithm 20: \textsc{MinCostFlow}(\(G, \vec{\sigma}, W\))

\textbf{Input:} directed graph \(G_0 = (V_0, E_0, c_0)\) with each edge having unit capacity and cost \(c_0;\)
\hspace{1em} |\(V_0| = n\) and |\(E_0| = m;\) integral demand vector \(\vec{\sigma};\) the absolute maximum cost \(W;\)
\hspace{1em} \(G = (P \cup Q, E), \vec{b}, \vec{f}, \vec{y}, \vec{s}, \vec{\nu}, \vec{\rho}, c_p, c_T, \eta \leftarrow \text{INITIALIZATION}(G_0, \vec{\sigma});\)
\hspace{1em} \text{Add a new vertex} \(v_0\) and undirected edges \((v_0, v)\) for every \(v \in P\) to \(G;\)
\hspace{1em} \textbf{for} \(i = 1\) to \(c_T \cdot m^{1/2 - 3\eta}\) \textbf{do}
\hspace{2em} \textbf{for each} \(v \in P\) \textbf{do}
\hspace{3em} \text{set resistance of edge} \((v_0, v)\) for each \(v \in P\) to be \(r_{v_0v} = \frac{m^{1/2 - 2\eta}}{a(v)},\) where \(a(v) \leftarrow \sum_{e \in Q, e = (v, u) \in E} \nu_e + \nu_{\mathbf{T}};\) \(\mathbf{T} = (\mathbf{T}, u)\) is \(e\)'s partner edge that is the unique edge sharing one common vertex from \(Q.\)
\hspace{2em} \textbf{for} \(j = 1\) to \(m^{2\eta}\) \textbf{do}
\hspace{3em} \textbf{while} \(\|\vec{\rho}\|_{\vec{\sigma},3} > c_p \cdot m^{1/2 - \eta}\) \textbf{do}
\hspace{4em} \(\vec{\rho}, \vec{y}, \vec{s}, \vec{\nu} \leftarrow \text{PERTURBATION}(G, \vec{\rho}, \vec{f}, \vec{y}, \vec{s}, \vec{\nu});\)
\hspace{4em} \(\vec{f}, \vec{s}, \vec{\rho}, \vec{\mu} \leftarrow \text{PROGRESS}(G, \vec{\sigma}, \vec{f}, \vec{\nu});\)
\hspace{2em} \text{REPAIRING}(G, \vec{f}, \vec{y});\)
Algorithm 21: **INITIALIZATION**($G, \bar{\sigma}$)

1. Create a new vertex $v_{aux}$ with $\sigma(v_{aux}) = 0$;
2. **for each** $v \in V_0$ **do**
   3. \( t(v) \leftarrow \sigma(v) + \frac{1}{2} \deg_{in} G_0(v) - \frac{1}{2} \deg_{out} G_0(v) \);
4. **if** $t(v) > 0$ **then** construct $2t(v)$ parallel edges $(v, v_{aux})$ with costs $\|\bar{c}_0\|_1$;
5. **else if** $t(v) < 0$ **then**
   6. construct $|2t(v)|$ parallel edges $(v_{aux}, v)$ with costs $\|\bar{c}_0\|_1$;
7. Let the new graph be $G_1 = (V_1, E_1, \bar{c}_1)$;
8. Initialize the bipartite graph $G = (P \cup Q, E, \bar{c})$ with $E \leftarrow \emptyset$, $P \leftarrow V_1$ and $Q \leftarrow \{e_{uv} \mid (u, v) \in E_1\}$ where $e_{uv}$ is a vertex corresponding to edge $(u, v) \in E_1$;
9. **for each** $(u, v) \in E_1$ **do**
   10. let $E \leftarrow E \cup \{(u, e_{uv}), (v, e_{uv})\}$ with $c(u, e_{uv}) = c_1(u, v)$ and $c(v, e_{uv}) = 0$, and set $b(u) \leftarrow \sigma(u) + \deg_{in} G_1(u)$, $b(v) \leftarrow \sigma(v) + \deg_{out} G_1(v)$ and $b(e_{uv}) \leftarrow 1$;
11. For each $v \in P$, set $y_v \leftarrow \|\bar{c}\|_\infty$, and for each $v \notin P$, set $y_v \leftarrow 0$;
12. For each $e = (u, v) \in E$, set $f_e \leftarrow \frac{1}{2}$, $s_e \leftarrow c_e + y_u - y_v$ and $\nu_e \leftarrow \frac{s_e}{\|\bar{c}\|_\infty}$;
13. Set $\hat{\mu} \leftarrow \|\bar{c}\|_\infty$, $c_\rho \leftarrow 400 \sqrt{3} \cdot \log^{1/3} W$, $c_T \leftarrow 3c_\rho \log W$ and $\eta \leftarrow \frac{1}{14}$;
14. **return** $G, \hat{b}, \hat{f}, \bar{y}, \bar{s}, \bar{v}, \hat{\mu}, c_\rho, c_T$ and $\eta$;

Algorithm 22: **PERTURBATION**($G, \hat{\mu}, \hat{f}, \bar{y}, \bar{s}, \bar{v}$)

```plaintext
/* Perturbation Step */
1. **for each** $v \in Q$ **do**
   2. let $e = (u, v)$ and $e = (u, v)$;
   3. $y_v \leftarrow y_v - s_e$;
   4. $\nu_e \leftarrow 2\nu_e$;
   5. $\nu_e \leftarrow \nu_e + \frac{\nu_e}{f_e}$;
6. Solve Laplacian linear systems in graph consisting of $G, v_0$ and the edge set $\{(v_0, v) \mid v \in P\}$;
```
Algorithm 23: Progress$(G, \bar{\sigma}, \tilde{f}, \tilde{v})$

/* Progress Step */
1 For each $e \in E$, let $r_e \leftarrow \frac{\nu_e}{|f_e|}$;
2 Solve Laplacian linear system $L(G)\tilde{\phi} = \bar{\sigma}$;
3 For each $e = (u, v) \in E$, let $\tilde{f}_e \leftarrow \frac{\phi_v - \phi_u}{r_e}$ and $\rho_e \leftarrow |\tilde{f}_e|$;
4 $\delta \leftarrow \min\left\{\frac{1}{8\|\rho\|_\infty}, \frac{1}{8}\right\}$;
5 Update $f'_e \leftarrow (1 - \delta)f_e + \delta \tilde{f}_e$ and $s'_e \leftarrow s_e - \frac{\delta}{1-\delta}(\phi_v - \phi_u)$;
6 For each $e \in E$, let $f'^*_e \leftarrow \frac{1}{s_e} f'_e$;
7 Obtain the flow vector $\tilde{\sigma}'$ corresponding to the residue $\tilde{f'} - \tilde{f'^*}$;
8 For each $e \in E$, let $r_e \leftarrow \frac{s'_e}{(1-\delta)f'_e}$;
9 Solve Laplacian linear system $L(G)\tilde{\phi} = \tilde{\sigma}'$;
10 For each $e = (u, v) \in E$, let $\tilde{f}_e \leftarrow \frac{\phi_v - \phi_u}{r_e}$;
11 Update $f_e \leftarrow f'^*_e + \tilde{f}_e$ and $s_e \leftarrow s'_e - \frac{s'_e \tilde{f}_e}{f'^*_e}$;
Algorithm 24: Repairing$(G, \vec{f}, \vec{y})$

/* Repairing Step */

1. Let $\vec{b}^+$ be the demand vector corresponding to the current flow $\vec{f}$;
2. For each $v \in P \cup Q$, set $b_v^{\leq} \leftarrow \min(b_v, b_v^+)$;
3. For each $v \in P \cup Q$, if $f(E(v)) > b_v^{\leq}$, set $\vec{f}$ on $E(v)$ such that $f(E(v)) = b_v^{\leq}$, and let the resulting vector be $\vec{f}^{\leq}$;
4. Add source $s$ and sink $t$ to $G$, and connect $s$ to each $v \in P$ with $f^{\leq}_{sv} \leftarrow f^{\leq}(E(v))$, and connect each $v \in Q$ to $t$ with $f^{\leq}_{vt} \leftarrow f^{\leq}(E(v))$ in $G$;
5. $\vec{M} \leftarrow \text{FlowRounding}(G, \vec{f}^{\leq}, s, t)$;
6. Remove $s, t$ and related coordinates on $\vec{f}^{\leq}$ and $\vec{M}$ from $G$;
7. for $i = 1$ to $\tilde{O}(m^{3/7})$ do
8. construct graph $\vec{G}_M = (P \cup Q, E_M, \vec{c}_M)$ using $G$, $\vec{M}$ and $\vec{c}$ such that for each $e = (u, v) \in E, \bar{c}_e = c_e - y_u - y_v$ and $E_M = \{(u, v) \in E \mid u \in P, v \in Q\} \cup \{(u, v) \mid u \in Q, v \in P, M_{uv} \neq 0\}$,
   $\bar{c}_M(u, v) = \begin{cases} 
   \bar{c}_{uv}, & u \in P, v \in Q \\
   -\bar{c}_{uv}, & u \in Q, v \in P
   \end{cases}$;
9. set $F_M \leftarrow \{v \in P \cup Q \mid M(v) < b_v\}$;
10. compute a shortest path $\pi$ in $\vec{G}_M$ from $P \cap F_M$ to $Q \cap F_M$;
   /* $D_{G_M}^\pi(P, u)$ is the distance from $P$ to $u$ in $\vec{G}_M$ */
   /* Edges that are reachable in $\vec{G}_M$ from $P \cap F_M$ have non-negative weights $\bar{c}_e$ */
11. for $u \in P \cup Q$ do
12. if $u$ can be reached from $P$ in $\vec{G}_M$ then
13. if $u \in P$ then
14. $y_u \leftarrow y_u - D_{G_M}^\pi(P, u)$;
15. else
16. $y_u \leftarrow y_u + D_{G_M}^\pi(P, u)$;
17. augment $\vec{M}$ using the augmenting path $\pi$;
18. return $\vec{M}$;