## Randomized Approximation Schemes for Cuts and Flows in Capacitated Graphs

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#### Abstract

We improve on random sampling techniques for approximately solving problems that involve cuts and flows in graphs. We give a near-linear-time construction that transforms any graph on n vertices into an  $O(n \log n)$ -edge graph on the same vertices whose cuts have approximately the same value as the original graph's. In this new graph, for example, we can run the  $\tilde{O}(m^{3/2})$ -time maximum flow algorithm of Goldberg and Rao to find an *s*-*t* minimum cut in  $\tilde{O}(n^{3/2})$  time. This corresponds to a  $(1 + \epsilon)$ -times minimum *s*-*t* cut in the original graph. In a similar way, we can approximate a sparsest cut to within  $O(\log n)$  in  $\tilde{O}(n^2)$  time using a previous  $\tilde{O}(mn)$ -time algorithm. A related approach leads to a randomized divide and conquer algorithm producing an approximately maximum flow in  $\tilde{O}(m\sqrt{n})$  time.

### 1 Introduction

Previous work [Kar94, Kar99, Kar00] has shown that random sampling is an effective tool for problems involving cuts in graphs. A *cut* is a partition of a graph's vertices into two groups; its *value* is the number, or in weighted graphs the total weight, of edges with one endpoint in each side of the cut. Many problems depend only on cut values. The maximum flow that can be routed from s to t is the minimum value of any cut separating s and t [FF56]. A minimum bisection is the smallest cut that splits the graph into two equal-sized pieces. The *connectivity* or *minimum cut* of the graph, which we denote throught by c, is equal to the minimum value of any cut.

Random sampling "preserves" the values of cuts in a graph. If we pick each edge of a graph G with probability p, we get a new graph in which every cut has expected value exactly p times it value in G. A theorem by Karger [Kar99] shows that if the graph has unit-weight edges and minimum cut c, then sampling with probability roughly  $1/\epsilon^2 c$  gives cuts that are all, with high probability, within  $1 \pm \epsilon$  of their expected values. In particular, the minimum cut of the sampled graph corresponds to a  $(1 + \epsilon)$ -times minimum cut of the original graph. Similarly, an s-t minimum cut of the sampled graph is a  $(1 + \epsilon)$ -times minimum s-t cut of the original graph. Since the sampled graph has fewer edges (by a factor of 1/c for any fixed  $\epsilon$ ),

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minimum cuts can be found in it faster than in the original graph. Working through the details shows that an approximately minimum cut can be found roughly  $c^2$  times faster than an exact solution.

A variant of this approach finds approximate solutions to flow problems via randomized divide and conquer. If we randomly partition the edges of a graph into roughly  $\epsilon^2 c$  subsets, each looks like the sample discussed in the previous paragraph, so has approximately accurate cuts. In other words, random division is a good approximation to evenly dividing up the capacities of all the cuts. By max-flow min-cut duality [FF56], this means that the *s*-*t* max-flow of *G* is also approximately evenly divided up. We can find a maximum flow in each of the subgraphs and add them together to get a flow in *G* that is at least  $(1 - \epsilon)$  times optimal. Again, detailed analysis shows that finding this approximate flow can be done *c* times faster than finding the exact maximum flow.

Unfortunately, the requirement that  $p = \Omega(1/c)$  limits the effectiveness of this scheme. For cut approximation, it means that in a graph with m edges, we can only reduce the number of edges to m/c. Similarly for flow approximation, it means we can only divide the edges into c groups. Thus, when c is small, we gain little. Results can be even worse in weighted graphs, where the ratio of total edge weight to minimum cut value is unbounded.

### 1.1 Results

In this paper, we show how nonuniform sampling can be used to remove graph sampling's dependence on the minimum cut c. Our main results are twofold: one for cut problems, and one for flow problems. For cuts, we show that by sampling edges nonuniformly, paying greater attention to edges crossing small cuts, we can produce accurate samples with far less than m/c edges—rather, the resulting compressed graph has only  $\tilde{O}(n/\epsilon^2)$  edges, regardless of the number of edges in the original graph.<sup>1</sup> In consequence, we show that a  $(1 + \epsilon)$ -times minimum s-t cut can be found in  $\tilde{O}(n^{3/2}/\epsilon^3)$  time in general capacity graphs (as compared to the  $\tilde{O}(m^{3/2})$  exact bound) and  $\tilde{O}(nv/\epsilon^2)$  time in unit-capacity graphs with flow value v (as compared with the O(mv) exact bound). Similarly, a nonuniform divide-and-conquer approach can be used to find a  $(1 - \epsilon)$ times maximum flow in  $\tilde{O}(m\sqrt{n}/\epsilon)$  time. Our approach works for undirected graphs with arbitrary weights (capacities).

Even ignoring the algorithmic aspects, the fact that any graph can be approximated by a sparse graph is of independent combinatorial interest.

In addition to proving that such sampling works, we give fast algorithms for determining the importance of different edges and the correct sampling probabilities for them. This involves an extension of the *sparse certificate* technique of Nagamochi and Ibaraki [NI92b].

Using these results, we demonstrate the following:

**Theorem 1.1.** Given a graph G and an error parameter  $\epsilon$ , there is a graph G' such that

- G' has  $O(n \log n / \epsilon^2)$  edges and
- the value of every cut in G' is  $(1 \pm \epsilon)$  times the value of the corresponding cut in G.

G' can be constructed in  $O(m \log^2 n)$  time if G is unweighted and in  $O(m \log^3 n)$  time if G is weighted.

It follows that given any algorithm to (even approximately) solve a cut problem, if we are willing to accept an approximate answer, we can substitute  $n \log n$  for any factor of m in the running time. Our applications of this result are the following:

**Corollary 1.2.** In an undirected graph, a  $(1 + \epsilon)$  times minimum s-t cut can be found in  $\tilde{O}(n^2/\epsilon^2)$  or  $\tilde{O}(n^{3/2}/\epsilon^3)$  time.

**Corollary 1.3.** In an undirected graph,  $a(1+\epsilon)$  times minimum s-t cut of value v can be found in  $O(nv/\epsilon^2)$  time.

<sup>&</sup>lt;sup>1</sup>The notation  $\tilde{O}(f)$  denotes O(f polylog I) where I is the input problem size.

**Corollary 1.4.** An  $O(\log n)$ -approximation to the sparsest cut in an undirected graph can be found in  $\tilde{O}(n^2/\epsilon^2)$  time.

These corollaries follow by applying our sampling scheme to (respectively) the maximum flow algorithms of Goldberg and Tarjan [GT88] and Goldberg and Rao [GR97], the classical augmenting-paths algorithm for maximum flow [FF56, AMO93], and the Klein-Stein-Tardos algorithm for approximating the sparsest cut [KST90].

A related approach helps solve flow problems: we divide edges crossing small cuts into several parallel pieces, so that no one edge forms a substantial fraction of any cut it crosses. We can then apply a randomized divide and conquer scheme. If we compute a maximum flow in each of the subgraphs created by the random division using the Goldberg-Rao algorithm, and then add the flows into a flow in G, we deduce the following corollary:

**Corollary 1.5.** A  $(1-\epsilon)$  times maximum flow can be found in  $\tilde{O}(m\sqrt{n}/\epsilon)$  time.

The work presented here combines work presented earlier by Karger and Benczur [BK96] and by Karger [Kar98]. The presentation is simplified and slight improvements are given.

### 1.2 Method

The previous work on sampling for cuts is basically an application of the Chernoff bound. Our goal in cut sampling is to estimate the total weight (or number, in the case of unit-weight graphs) of edges crossing each cut of the graph. We motivate our approach by considering a simpler problem—that of estimating a single cut. Consider a set of m weights  $w_e$ , and suppose that we wish to estimate the sum  $S = \sum w_e$ . A natural approach is random sampling: we choose a random subset of the weights, add them, and scale the result appropriately. A somewhat easier to analyze approach is to choose each weight independently with some probability p, compute their sum S', and estimate S = S/p. Since we choose only pm weights in expectation, this sampling approach saves time. But we must analyze its accuracy. The Chernoff bound is a natural tool.

**Lemma 1.6 (Chernoff [Che52]).** Given any set of random variables  $X_i$  with values distributed in the range [0,1], let  $\mu = E[\sum X_i]$  and let  $\epsilon < 1$ . Then

$$\Pr[\sum X_i \notin (1 \pm \epsilon)\mu] \le 2e^{-\epsilon^2 \mu/3}.$$

The lemma's requirement that  $X_i \leq 1$  is in force to prevent any one random variable from "dominating" the outcome of the sampling experiment. For example, if one variable takes on value S with probability 1/S and 0 otherwise, while all other variables are uniformly 0, then the (relatively rare, but still occasional) outcome of taking on value S will dramatically skew the sum away from is expectation of 1.

We can model our sampling experiment so as to apply the Chernoff bound. For now, let us assume that each  $w_e \leq 1$ . Let  $X_e$  be a random variable defined by setting  $X_e = w_e$  with probability p and  $X_e = 0$ otherwise. Note that  $\sum X_e$  is the value of our sampling experiment of adding the weights we have chosen to examine. Also,  $E[\sum X_e] = \sum pw_e = pS$ . The variables  $X_e$  satisfy the conditions of the Chernoff bound, letting us deduce that the probability that  $\sum X_e$  deviates by more than  $\epsilon$  from its expectation is  $e^{-\epsilon^2 pS/3}$ . Note that this deviation is exponentially unlikely as a function of the expected sample value pS.

We now note some slack in this sampling scheme. If some  $w_e \ll 1$ , then its random sample variable  $X_e$ , which takes on values 0 or  $w_e$ , is far away from violating the requirement that each  $X_e \in [0, 1]$ . We can afford to apply a more aggressive sampling strategy without violating the Chernoff bound assumptions. Namely, we we can set  $X_e = 1$  with probability  $pw_e$  and 0 otherwise. We have chosen this probability because it keeps the expected value of each  $X_e$ , and thus  $E[\sum X_e]$ , unchanged while making each variable "tight" against the  $X_e \leq 1$  limit of the Chernoff bound. Since this fits the preconditions of the lemma, we preserve the  $(1 \pm \epsilon)$ concentration around the mean shown by the Chernoff bound. However, under this scheme, the expected number of sampled values drops from pm to  $\sum pw_e$  (which is less since we assume each  $w_e < 1$ ). This is a noteworthy quantity: it is equal to the expected value  $\mu = E[\sum X_e]$ . Since the probability of error in the Chernoff bound is itself a function only of  $\mu$ , it follows that under this scheme the expected number of samples  $\mu$  needed to guarantee a certain error probability  $\delta$  is a function only of the desired bound (namely,  $\mu = 3(\ln 1/\delta)/\epsilon^2$ ), and not of the number of variables m or their values  $w_e$ . Note further that since the  $w_e$ do not affect the analysis, if our  $w_e$  violate the assumption that  $w_e \leq 1$ , we can scale them all by dividing by max  $w_e$  and apply the same result. So the restriction  $w_e \leq 1$  was actually irrelevant.

The key feature of this scheme is that an item's greater weight is translated into an increased probability of being sampled: this lets it contribute more to the expectation of the sample without contributing too much to its variance.

One might object that in order to apply the above scheme, we need to know the weights  $w_e$  in order to decide on the correct sampling probabilities. This would appear to imply a knowledge of the very quantity we wish to compute. It is at this point that we invoke the specifics of our approach to avoid the difficulty.

We modify a uniform sampling scheme developed previously [Kar99]. That scheme sampled all graph edges with the same probability and showed the following.

**Lemma 1.7 ([Kar99]).** Let G be a graph in which the edges have mutually independent random weights, each distributed in the interval [0,1]. If the expected weight of every cut in G exceeds  $\rho_{\epsilon} = 3(d+2)(\ln n)/\epsilon^2$ for some  $\epsilon$  and d, then with probability  $1 - 1/n^d$  every cut in G' has value within  $(1 \pm \epsilon)$  of its expectation.

The intuition behind this theorem is the same as for the Chernoff bound. In the sampled graph, the expected value of each cut is  $\Omega((\log n)/\epsilon^2)$ , while each edge contributes value at most 1 to the sampled cuts it is in. Thus, the contribution of any one edge to the possibile deviation of a cut from its mean is negligible.<sup>2</sup>

As in our above discussion, we now observe that an edge that only crosses large-valued cuts can have its sampled weight scaled up (and its probability of being sampled correspondingly scaled down) without making that edge dominate any of the samples it is in. Consider a k-connected induced subgraph of G with k > c. Lemma 1.7 says that we can sample the edges of this subgraph with probability  $\tilde{O}(1/k)$  (and scale their weights up by  $\tilde{O}(k)$  to preserve expectations) without introducing significant error in the cut values. More generally, we can sample edges in any subgraph with probability inversely proportional to the connectivity of that subgraph. We will generalize this observation to argue that we can simultaneously sample each edge with probability inversely proportional to the maximum connectivity of any subgraph containing that edge.

To take advantage of this fact, we will show that almost all the edges are in components with large connectivities and can therefore be sampled with low probability—the more edges, the less likely they are to be sampled. We can therefore construct an  $O(n \log n)$ -edge graph that, regardless of the minimum cut value, accurately approximates all cut values.

### **1.3** Definitions

We use the term "unweighted graph" to refer to a graph in which all edges have weight 1. In the bulk of this paper, G denotes an unweighted undirected graph with n vertices and m edges; parallel edges are allowed. We also consider weighted graphs. By scaling weights, we can assume the minimum edge weight is at least one. For the purpose of correctness analysis when running times are not relevant, it is often convenient to treat an edge of weight w as a set of w parallel edges with the same endpoints.

A cut C is a partition of the vertices into two subsets. The value VAL(C, G) of the cut in unweighted (resp. weighted) graph G is the total number (resp. weight) of edges with endpoints in different subsets.

We simplify our presentation with a vector notation. The term  $x_E$  denotes a vector assigning some value  $x_e$  to each  $e \in E$ . All operations on vectors in this paper are coordinatewise. The interpretation of  $x_E + y_E$  is standard, as is the product  $\gamma x_E$  for any constant  $\gamma$ . However, we let  $x_E \times y_E$  denote the product  $z_E$  with  $z_e = x_e y_e$ . Similarly, let  $1/x_E$  denote the vector  $z_E$  such that  $z_e = 1/x_e$  (pointwise inverse). More generally, let  $y_E/x_E$  be the vector  $z_E$  with  $z_e = y_e/x_e$ .

 $<sup>^{2}</sup>$ This theorem is nontrivial, as the exponential number of cuts means that events which are very unlikely on one cut still seem potentially probable over all cuts. But it can be shown that most cuts are so large in expectation that their deviation is exponentially unlikely.

A weighted graph G can be thought of as the vector (indexed by edge set E) of its edge weights. (An unweighted graph has value 1 in all coordinates.) Applying our vector notation, when  $r_E$  is a vector over the edge set, we let  $r_E \times F$  denote a graph with edge weight vector  $r_E F$ . Similarly, if G and H are graphs, then G + H denotes the graph whose edge weight vector is the sum of those graphs'.

We also introduce a sampling notation. As is traditional, we let G(p) denote a graph in which each edge of G is incorporated with probability p. Generalizing, we let  $G(p_E)$  denote a random subgraph of G generated by included each edge e of G (with its original weight) independently with probability  $p_e$ . We define the *expected value graph*  $E[G(p_E)] = p_E \times G$ , since the expected value of any edge in  $G(p_E)$  is equal to the value of that edge in  $p_E \times G$ . This means that expected cut values are also captured by the expected value graph.

We say that an event occurs with high probability if its probability is  $1 - O(n^{-d})$  for some constant d. The constant can generally be modified arbitrarily by changing certain other constants hidden in the asymptotic notation.

### 1.4 Outline

In Section 2 we define the *strong connectivity* measure that is used to determine the relative impact of different edges on cut samples, and show that samples based on this strong connectivity measure have good concentration near their mean. Our application to s-t min-cuts is immediate. In Section 3 we introduce graph smoothing, a variation on compression that can be used for flow approximation. Finally, in Section 4, we show how the strong connectivities needed for our sampling experiments can actually be estimated quickly.

### 2 Approximating Cuts via Compression

As was stated above, we aim to sample edges with varying probabilities. To preserve cut values, we compensate for these varying sampling probabilities using *compression*. To define the appropriate sampling probability for each edge, we introduce the notion of *strong connectivity*. For the bulk of this section, we will focus on unweighted graphs, though we will occasionally make reference to edge weights for future use.

### 2.1 Compression

Sampling edges with different probabilities means that cut values no longer scale linearly. To make the expected cut value meaningful, we counterbalance the varying sampling probabilities by introducing edge weights on the sampled edges.

**Definition 2.1.** Given an unweighted graph G and compression probabilities  $p_e$  for each edge e, we build a compressed graph  $G[p_E]$  by including edge e in  $G[p_E]$  with probability  $p_e$ , and giving it weight  $1/p_e$  if it is included.

In our notation above, the compressed graph  $G[p_E] = 1/p_E \times G(p_E)$ . Since the expected weight of any edge in the graph is 1, every cut's expected value is equal to its original value, regardless of the  $p_e$ . That is,  $E[1/p_E \times G(p_E)] = G$ . However, the expected number of edges in the graph is  $\sum p_e$ . We would therefore like to make all the  $p_e$  as small as possible. We are constrained from doing so, however, by our need to have all the cut values tightly concentrated around their expectations. An edge compressed with probability  $p_e$  has variance  $(1 - p_e)/p_e$ , and the large variances produced by small  $p_e$  work against our wish for tight concentration. The key question, then, is how small we can make our  $p_e$  values (and thus our expected number of sampled edges) while preserving tight concentration of cut values.

### 2.2 Strong Connectivity

In this section, we formalize the notion of subgraphs with large connectivities. As was discussed above, if we identify a subgraph with connectivity  $k \gg c$ , then we might hope, based on Lemma 1.7, to sample

edges in this subgraph with probability roughly 1/k, producing a graph much sparser than if we sample with probability 1/c.

**Definition 2.2.** A graph G is *k*-connected if the value of each cut in G is at least k.

**Definition 2.3.** A k-strong component of G is a maximal k-connected vertex-induced subgraph of G.

It follows that the k-strong components partition the vertices of a graph and each (k+1)-strong component is contained in a single k-strong component—that is, that the partition into (k+1)-strong components refines the partition into k-strong components.

**Definition 2.4.** The strong connectivity or strength of an edge e, denoted  $k_e$ , is the maximum value of k such that a k-strong component contains (both endpoints of) e. We say e is k-strong if its strong connectivity is k or more, and k-weak otherwise.

Note that the definition of strong connectivity of an edge differs from the standard definition of connectivity:

**Definition 2.5.** The *(standard) connectivity* of an edge e is the minimum value of a cut separating its endpoints.

Consider the graph with unit-weight edges  $(s, v_i)$  and  $(v_i, t)$  for i = 1, ..., n. Vertices s and t have (standard) connectivity n but only have strong connectivity 1. An edge's strong connectivity is always less than its connectivity since an edge in a k-strong component cannot be separated by any cut of value less than k.

### 2.3 The Compression Theorem

We now use the above definitions to describe our results. We will use a fixed *compression factor*  $\rho_{\epsilon}$  chosen to satisfy a given error bound  $\epsilon$ :

$$\rho_{\epsilon} = 3(d+4)(\ln n)/\epsilon^2$$

**Theorem 2.6 (Compression).** Let G be an unweighted graph with edge strengths  $k_e$ . Given  $\epsilon$  and a corresponding  $\rho_{\epsilon}$ , for each edge e, let  $p_e = \min\{1, \rho/k_e\}$ . Then with probability  $1 - n^{-d}$ ,

- 1. The graph  $G[p_E]$  has  $O(n\rho)$  edges, and
- 2. every cut in  $G[p_E]$  has value between  $(1 \epsilon)$  and  $(1 + \epsilon)$  times its value in G.

In particular, to achieve any constant error in cut values with high probability, one can choose  $\rho$  to yield  $O(n \log n)$  edges in the compressed graph.

We now embark on a proof of the Compression Theorem.

#### 2.3.1 Bounding the number of edges

To prove the first claim of the Compression Theorem we use the following lemma:

**Lemma 2.7.** In a weighted graph with edge weights  $u_e$  and strengths  $k_e$ ,

$$\sum u_e/k_e \le n-1.$$

*Proof.* Define the cost of edge e to be  $u_e/k_e$ . We show that the total cost of edges is at most n-1. Let C be any connected component of G and suppose it has connectivity k. Then there is a cut of value k in

C. On the other hand, every edge of C is in a k-strong subgraph of G (namely C) and thus has strength at least k. Therefore,

$$\sum_{e \text{ crossing } C} u_e/k_e \leq \sum_{e \in V} u_e/k$$
$$= k/k$$
$$= 1$$

Thus, by removing the cut edges, of total cost at most 1, we can break C in two, increasing the number of connected components of G by 1.

If we find and remove such a cost-1 cut n-1 times, we will have a graph with n components. This implies that all vertices are isolated, meaning no edges remain. So by removing n-1 cuts of cost at most 1 each, we have removed all edges of G. Thus the total cost of edges in G is at most n-1.

This lemma implies the first claim of the Compression Theorem. In our graph compression experiment, all edge weights are one, and we sample each e with probability  $\rho/k_e$ . It follows that the expected number of edges is  $\rho \sum 1/k_e \leq \rho(n-1)$  by the previous lemma. The high probability claim follows by a standard Chernoff bound [Che52, MR95].

#### 2.3.2 Proving cuts are accurate

We now turn to the proof that cuts are accurate in the compressed graph. Once again, we apply a useful property of edge strengths.

**Lemma 2.8.** If graph G has edge strengths  $k_e$  then the graph  $1/k_E \times G$  has minimum cut exactly 1.

*Proof.* Consider any minimum cut in G, of value c. Each edge in the cut has strength c, giving it weight 1/c in  $1/k_E \times G$ . Thus, the cut has value 1 in  $1/k_E \times G$ . It follows that the minimum cut in  $1/k_E \times G$  is at most 1.

Now consider any cut, of value k in G. Each edge crossing the cut has strength at most k, meaning it gets weight at least 1/k in  $1/k_E \times G$ . Since k edges cross this cut, it follows that the cut has weight at least  $k(1/k) \ge 1$ . This shows that the minimum cut in  $1/k_E \times G$  is at least 1.

Combining these two arguments yields the claimed result.

Recall that for graph compression, we initially assign weight  $k_e$  to edge e, producing a weighted graph  $k_E \times G$ . We then produce a random graph by choosing edge e of  $k_E \times G$  with probability  $\rho/k_e$ , generating the graph  $k_E \times G(\rho/k_E)$  (we assume for the moment that all  $k_e \ge \rho$  so the sampling probability is at most 1). Our goal is to show that the resulting graph has cuts near their expected values.

Our basic approach is to express  $k_E \times G$  as a weighted sum of graphs, each of which, when sampled, is easily proven to have cut values near their expectations. It will follow that the sampled  $k_E \times G(\rho/k_E)$  also has cut values near its expectations.

We now define the decomposition of G. There are at most m distinct edge-strength values in G, one per edge (in fact it can be shown there are only n-1 distinct values, but this will not matter). Number these values  $k_1, \ldots, k_r$  in increasing order, where  $r \leq m$ . Now define the graph  $F_i$  to be the edges of strength at least  $k_i$ —in other words,  $F_i$  is the set of edges in the  $k_i$ -strong components of G. Write  $k_0 = 0$ . We now observe that

$$k_E \times G = \sum_i (k_i - k_{i-1}) \times F_i.$$

To see this, consider some edge of strength exactly  $k_i$ . This edge appears in graphs  $F_1, F_2, \ldots, F_i$ . The total weight assigned to that edge in the right hand of the sum above is therefore

$$(k_1 - k_0) + (k_2 - k_1) + \dots + (k_i - k_{i-1}) = k_i - k_0 = k_i$$

as is required to produce the graph  $k_E \times G$  which has weight  $k_e$  on edge e.

We can now examine the effect of compressing G by examining its effect on the graphs  $F_i$ . Our compression experiment flips an appropriately biased coin for each edge of  $k_E \times G$  and keeps it if the coin shows heads. We can think of these coin flips as also being applied to the graphs  $F_i$ . We apply the same coin flip to all the  $F_i$ : edge e of strength  $k_i$ , present in  $F_1, \ldots, F_i$ , is kept in all of the respective samples  $F_i(\rho/k_E)$  if the coin shows heads, it is discarded from all if the coin shows tails. Thus, the samples from the graphs  $F_i$  are not independent. However, if we consider a *particular*  $F_i$ , then the sampling outcomes of edges are mutually independent in that particular  $F_i$ .

Let us first consider graph  $F_1$  (which is simply the graph G since all  $k_e \ge 1$ ). As was discussed in Section 1.3, the expected value  $E[G(\rho/k_E)] = \rho/k_E \times G$  has cut values equal to the expectations of the corresponding cuts of the sampled graph  $G(\rho/k_E)$ . We saw above that the graph  $1/k_E \times G$  has minimum cut 1. It follows that the expected value graph  $\rho/k_E \times G$  has minimum cut  $\rho$ . This suffices to let us apply the basic sampling result (Lemma 1.7) and deduce that every cut in  $F_1$  has value within  $(1 \pm \epsilon)$  of it expectation with high probability. Scaling the graph preserves this: the graph  $(k_1 - k_0) \times F_1(1/k_E)$  has cut values within  $(1 \pm \epsilon)$  of their expectations with high probability.

Now consider any other  $F_i$ . The subgraph  $F_i$  consists of all the edges inside the  $k_i$ -strong components of G. Consider one particular such component C, and an edge  $e \in C$ . Since C is  $k_i$ -connected, we know that  $k_e \geq k_i$ . By definition, edge e is contained in some  $k_e$ -connected subgraph of G. As was argued above in Section 2.2, the  $k_e$ -connected subgraph that contains e must be wholly contained in C. Thus, the strength of edge e with respect to the graph C is also  $k_e$ .<sup>3</sup> Our argument of the previous paragraph for graph G therefore applies to the graph C, implying that the sampled version of C in  $F_i(\rho/k_E)$  has cuts within  $(1 \pm \epsilon)$  of their expected values with high probability. Since this is true for each component C, it is also true for the graph  $F_i$  (since each cut of  $F_i$  is a cut of components of  $F_i$ ).

This completes our argument. We have shown that each  $F_i(\rho/k_E)$  has all cuts within  $(1 \pm \epsilon)$  of their expected values with probability  $1-1/n^{d+2}$  (the quantity d+2 follows from our choice of  $\rho$  and the application of Lemma 1.7). Even though the  $F_i(1/k_E)$  are not independent, it follows from the union bound that all (possibly  $n^2$ ) distinct  $F_i$  samples are near their expectation with probability  $1-1/n^d$ . If this happens, then the sample  $k_E \times G(1/k_E) = \sum (k_i - k_{i-1}) \times F_i(1/k_E)$  has all cuts within  $1 \pm \epsilon$  of their expected values (this follows because all multipliers  $k_i - k_{i-1}$  are positive). Of course, the expected graph  $E[k_E \times G(1/k_E)] = G$ .

Our analysis has assumed all edges are sampled with probability  $\rho/k_e$ , which is false for edges with  $k_e < \rho$  (their sampling probability is set to 1 in the Compression Theorem). To complete the analysis, consider the  $\rho$ -strong components of G. Edges outside these components are not sampled. Edges inside the components are sampled with probabilities at most 1. We apply the argument above to each  $\rho$ -strong component separately, and deduce that it holds for the entire compressed graph.

### 2.4 Weighted Graphs

For simplicity, our compression analysis was done in terms of unweighted graphs. However, we can apply the same analysis to a weighted graph. If the weights are integers, we can think of a weight u edge as a set of u parallel unit-weight edges and apply the analysis above. Given the strengths  $k_e$ , we would take each of the u edges with probability  $1/k_e$  and give it weight  $k_e$  if taken. Of course, if u is large it would take too much time to perform a separate coin flip for each of the u edges. However, we can see that the number of edges actually taken has a binomial distribution with parameters  $u_e$  and  $\rho/k_e$ ; we can sample directly from that binomial distribution. Note that the number of edges produced is  $O(n \log n)$  regardless of the  $u_e$ .

<sup>&</sup>lt;sup>3</sup>This proof step is the sole motivation for the introduction of strong connectivity. The nesting of strong components lets us draw conclusions about the graphs  $F_i$  that *cannot* be drawn about standard connectivity. The set of edges with standard connectivity exceeding k does *not* form a k-connected graph, which prevents our proof from going through when we use standard connectivity.

Nonetheless, it is conceivable that standard connectivity is a sufficient metric for our sampling algorithm. We have found no counterexample to this possibility.

To handle noninteger edge weights, imagine that we multiply all the edge weights by some large integer z. This uniformly scales all the cut values by z. It also scales all edge strengths by z. If we now round each edge down to the nearest integer, we introduce an additive error of at most m to each cut value (and strength); in the limit of large z, this is a negligible relative error. To compress the resulting graph, the approach of the previous paragraph now says that for a particular edge e with original weight  $u_e$ , we must choose from a binomial distribution with parameters  $\lfloor zu_e \rfloor$  (for the number of edges, which has been multiplied by z and rounded) and  $\rho/zk_e$  (since all edges strengths have also been multiplied by z). In the limit of large z, it is well known [Fel68] that this binomial distribution converges to a Poisson Distribution with parameter  $\lambda = \rho u_e/k_e$ . That is, we produce s sample edges with probability  $e^{-\lambda}\lambda^s/s!$ . Under the compression formula, their weights would each be  $zk_e/\rho$ . Recall, however that we initially scaled the graph up by z; thus, we need to scale back down by z to recover G; this produces edge weights of  $k_e/\rho$ .

From an algorithmic performance perspective, we really only care whether the number of sampled edges is 0 or nonzero since, after sampling, all the sampled edges can be aggregated into a single edge by adding their weights. Under the Poisson distribution, the probability that the number of sampled edges exceeds 0 is  $1 - e^{-\rho u_e/k_e} \approx \rho u_e/k_e$ . It is tempting to apply this simplified compression rule to the graph (take edge ewith probability  $\rho u_e/k_e$ , giving it weight  $k_e/\rho$  if taken). A generalized Compression theorem in the appendix shows that this approach will indeed work.

### 2.5 Using Approximate Strengths

Our analysis above assumed edge strengths were known. While edge strengths can be computed exactly, the time needed to do so would make them useless for cut and flow approximation algorithms. Examining the proofs above, however, shows that we do not need to work with exact edge strengths.

**Definition 2.9.** Given a graph G with n vertices, edge weights  $u_e$ , and edge strengths  $k_e$ , a set of edge value  $\tilde{k}_e$  are *tight strength bounds* if

1.  $k_e \leq k_e$  and

2. 
$$\sum u_e/k_e = O(n)$$

**Theorem 2.10.** The Compression Theorem remains true even if tight strength bounds are used in place of exact strength values.

*Proof.* The proof of cut accuracy relied on the fact that each sampled edge had small weight compared to its cuts. The fact that  $\tilde{k}_e \leq k_e$  means that the weights of included edges are smaller than they would be if true strengths were used, which can only help.

The bound on the number of edges in the compressed graph followed directly from the fact that  $\sum u_e/k_e \leq n$ ; for tight strength bounds this summation remains asymptotically correct.

Tight strength bounds are much easier to compute than exact strengths.

**Theorem 2.11.** Given any m-edge, n-vertex graph, tight strength bounds can be computed in  $O(m \log^2 n)$  time for unweighted graphs and  $O(m \log^3 n)$  time for weighted graphs.

*Proof.* See Section 4.

### 2.6 Applications

We have shown that graphs can be compressed based on edge strengths while preserving cut values. This suggests that cut problems can be approximately solved by working with the compressed graph as a surrogate for the original graph. We now prove the application corollaries from the introduction.

#### 

#### 2.6.1 Minimum s-t cuts.

As discussed above, we can compute tight strengths bounds in  $\tilde{O}(m)$  time and generate the resulting compressed graph  $G[p_E]$  as described in the Compression Theorem. The graph will have  $O(\rho n) = O(n(\log n)/\epsilon^2)$  edges.

Let us fix a pair of vertices s and t. Let  $\hat{v}$  be the value of a minimum cut separating s from t in the compressed graph  $G[p_E]$ . We show that the minimum s-t cut value v in G is within  $(1 \pm 3\epsilon)\hat{v}$ . By the Compression Theorem, with high probability the s-t minimum cut C in G has value at most  $(1 + \epsilon)v$  in  $G[p_E]$ . Thus  $\hat{v} \leq (1 + \epsilon)v$ . Furthermore, with high probability every cut of G with value exceeding  $(1 + 3\epsilon)v$  in G will have value at least  $(1 - \epsilon)(1 + 3\epsilon) \geq (1 + \epsilon)v$  in  $G[p_E]$  and therefore will not be the minimum cut of  $G[p_E]$ .

We can find an approximate value  $\hat{v}$  of the minimum s-t cut (and an s-t cut with this value) by computing a maximum flow in the  $O(n \log n / \epsilon^2)$ -edge graph  $G[p_E]$ . The maximum flow algorithm of Goldberg and Tarjan [GT88] has a running time of  $O(nm \log(n^2/m))$  which leads to a running time of  $O(n^2 \log^2 n / \epsilon^2)$ after compression. Similarly, the Goldberg-Rao algorithm [GR97], which runs in  $\tilde{O}(m^{3/2})$  time, leads to a running time of  $\tilde{O}(n^{3/2}/\epsilon^3)$  after compression.

In an integer-weighted graph with small flow value, we may wish to apply the classical augmenting path algorithm [FF56, AMO93] that finds a flow of value v in v augmentations. As described, the graph-compression process can produce noninteger edge weights  $\rho/k_e$ , precluding the use of augmenting paths in the smoothed graph. However, if we decrease each compression weight to the next lower integer (and increase the sampling probability by an infinitesimal amount to compensate) then compression will produce an integer-weighted graph in which the augmenting paths algorithm can be applied to find an s-t cut of value at most  $(1 + \epsilon)v$  in time  $O(nv \log n / \epsilon^2)$ .

#### 2.6.2 Sparsest cuts

A sparsest cut of a graph G minimizes the ratio between the cut value and the product of number of vertices on the two sides. It is  $\mathcal{NP}$ -hard to find the value of a sparsest cut. To find an  $\alpha$ -approximate value of a sparsest cut, we use the approach of the previous subsection: we compute a  $\beta$ -approximate sparsest cut in the compressed graph  $G[p_E]$ . This cut is then an  $\alpha = (1 + \epsilon)\beta$ -approximate sparsest cut of G.

An algorithm of Klein, Stein and Tardos [KST90] finds an  $O(\log n)$ -approximation to a sparsest cut in  $O(m^2 \log m)$  time. By running their algorithm on  $G[p_E]$ , we will find an  $O(\log n)$ -approximate sparsest cut in  $O(n^2 \log^3 n / \epsilon^4)$  time. Our small cut-sampling error is lost asymptotically in the larger error of the approximation algorithm.

Our approach been applied in a similar way to improve the running time of a spectral partitioning algorithm [KVV00].

### 3 Approximating Flows by Graph Smoothing

Until now we have focused on cut problems. Our compression scheme produces a graph with nearly the same cut values as the original, so that cut problems can be approximated in the compressed graph. But consider a maximum flow problem. It would seem natural to try to approximate this maximum flow by finding a maximum flow in the compressed graph. By providing an approximately minimum s-t cut, this approach does indeed give an approximation to the *value* of the maximum flow. But since edges in the compressed graph have *larger* capacity than the original graph edges, a feasible flow in the compressed graph will probably not be feasible for the original graph.

Previous work [Kar99] tackled the flow approximation problem with a divide-and-conquer approach. The edges of G are randomly divided into a number of groups, producing several random subgraphs of G. Lemma 1.7 is applied to deduce that each subgraph has cut values near their expectations. By computing a flow in each subgraph and adding the flows, we find a flow of value  $(1 - \epsilon)$  times the maximum flow in G.

This approach suffers the same limitation as the uniform sampling approach for cuts: the probability of each edge occurring in each subgraph must be  $\Omega(1/c)$  to preserve cut values. This translates into a limit that we divide into O(c) groups, which limits the power of the scheme on a graph with small minimum cuts. Graph compression's nonuniform sampling approach does not seem to provide an immediate answer: clearly we cannot simultaneously divide each edge with strength  $k_e$  among  $k_e$  distinct subgraphs. Instead we need a consistent rule that divides all edges among a fixed number of subgraphs. Each subgraph must therefore look like a *uniform* sample from the original graph.

In this section we introduce *graph smoothing*—a technique that lets us apply uniform sampling, and through it analyze randomized divide and conquer algorithms, for graphs with small minimum cuts, yielding fast approximation algorithms for flows in such graphs. The approach applies equally well to weighted graphs.

Our approach again starts with Lemma 1.7. The sampling proof used a Chernoff bound, which relied on individual edges having only a small impact on the outcome of the experiment. In particular, since the graph had minimum cut c, and every edge was being chosen with probability p, every cut had expected value at least pc. Thus, the presence or absence of a single (weight 1) edge could affect that value of a cut by at most a 1/pc-fraction of its expected value.

If we want to be able to sample more sparsely, we run into a problem of certain edges contributing a substantial fraction of the expected value of the cuts they cross, so that the Chernoff bound breaks down. A fix is to divide such edges into a number of smaller-weight edges so that they no longer dominate their cuts. Dividing *all* the graph edges is quite pointless: splitting all edges in half has the effect of doubling the minimum cut (allowing us to sample at half the original rate while preserving approximate cut values), but since we double the number of edges, we end up with the same number of sampled edges as before.

The approach of k-strong components lets us circumvent this problem. We use k-strong components to show that only a small fraction of the graph's edges are large compared to their cuts. By dividing *only* those edges, smoothing the highest-variability features of the sample, we allow for a sparser sample that still preserves cut values. Since only a few edges are being divided, the random subgraphs end up with fewer edges than before, making algorithms based on the samples more efficient.

### 3.1 Smooth Graphs

For the study of graph compression, we focused on unweighted graphs. For smoothing we focus on weighted graphs. In keeping with standard terminology for flows, we will refer to weights as *capacities*. It is easy to extend the notation G(p) to denote taking each capacitated edge with probability p, but somewhat harder to prove that sampling does the right thing. As discussed above, the problem is that a single capacitated edge might account for much of the capacity crossing a cut. The presence or absence of this edge has a major impact on the value of this cut in the sampled graph. However, the idea of edge strength described above gives us a useful bound on how much impact a given edge can have.

**Definition 3.1.** A graph G with edge capacities  $u_e$  and edge strengths  $k_e$  is c-smooth if for every edge,  $k_e \ge cu_e$ .

Note that a graph with integer edge weights and minimum cut c has smoothness at most c but possibly much less. We now argue that smoothness is the criterion we need to apply uniform sampling to weighted graphs.

**Theorem 3.2.** Let G be a c-smooth graph. Let  $p = \rho_{\epsilon}/c$  where  $\rho_{\epsilon} = O((\log n)/\epsilon^2)$  as in the Compression Theorem. Then with high probability, every cut in G(p) has value in the range  $(1 \pm \epsilon)$  times its expectation (which is p times its original value).

*Proof.* We use a variation on the proof of the Compression Theorem. Given the graph G, with edge capacities  $u_e$ , let  $k_i$  be a list of the at most m strengths of edges in G in increasing order, and let  $F_i$  denote the graph whose edge set is the  $k_i$ -strong edges of G, but with edge e assigned weight  $cu_e/k_e$ . It follows, just as

was argued above, that  $G = \sum (k_i - k_{i-1})F_i$ . So if we prove that each  $F_i$  can be accurately sampled with probability  $p = \rho/c$ , then the same will apply to G.

So consider graph  $F_i$ . Since we have assigned weights  $cu_e/k_e$ , the minimum cut in  $F_i$  is c, as was argued in Lemma 2.8. At the same time, edge e, if present in this graph, has weight  $cu_e/k_e \leq 1$  by the smoothness property. It follows that we can apply Lemma 1.7 to each component of the graph  $F_i$  and deduce that all cuts are within  $(1 \pm \epsilon)$  of their expectation, as desired. The remainder of the proof goes as for the Compression Theorem.

### 3.2 Making Graphs Smooth.

We have shown that a smooth graphs can be sampled uniformly, which will lead to good flow algorithms. We now give algorithms for transforming any graph into a smooth one.

**Lemma 3.3.** Given an m edge capacitated graph, a smoothness parameter c and the strengths  $k_e$  of all edges, we can transform the graph into an m + cn-edge c-smooth graph in  $\tilde{O}(m)$  time.

*Proof.* Divide edge e into  $\lceil cu_e/k_e \rceil$  parallel edges, each of capacity  $u_e / \lceil cu_e/k_e \rceil \le k_e/r$  but with total capacity  $u_e$ . These edges remain  $k_e$  strong, but now satisfy the smoothness criterion.

It remains to prove that this division creates at most nr new edges. The number of edges in our smoothed graph is

$$\sum_{e} \lceil cu_e/k_e \rceil \leq m + \sum cu_e/k_e$$
$$= m + c \sum_{e} u_e/k_e$$
$$\leq m + cn$$

where the last line follows from Lemma 2.7.

**Corollary 3.4.** Given edge strengths, in O(m) time we can transform any m-edge capacitated graph into an O(m)-edge capacitated (m/n)-smooth graph.

Choosing the smoothness parameter m/n is in some sense optimal. Any smaller smoothness parameter leads to worse sampling performance without decreasing the asymptotic number of edges (which is always at least m). A larger smoothness parameter provides better sampling behavior, but linearly increases the number of edges such that the gains from sparser sampling are lost.

### 3.3 Approximate Max-Flows

To approximate flows, we use the graph smoothing technique. As was argued in Theorem 2.10, graph smoothing works unchanged even if we use tight strength bounds, rather than exact strengths, in the computation.

After computing tight strength bounds in O(m) time (as will be discussed in Section 4), we can apply Lemma 3.5. This shows that in any c-smooth graph, sampling with probability p produces a graph in which with high probability all cuts are within  $(1 \pm \epsilon)$  of their expected values. This fact is the only one used in the uncapacitated graph flow algorithms of [Kar99]. Therefore, those results immediately generalize to the smooth graphs defined here—we simply replace "minimum cut" with "smoothness" in all of those results. The generalization is as follows:

**Lemma 3.5.** Let T(m, n, v, c) be the time to find a maximum flow in a graph with m edges, n vertices, flow v and smoothness c. Then for any  $\epsilon$ , the time to find a flow of value  $(1 - \epsilon)v$  on an m-edge, n-vertex, smoothness-c graph is

$$\tilde{O}(\frac{1}{p}T(pm,n,pv,pc))$$

where  $p = \Theta((\log n)/\epsilon^2 c)$ .

*Proof.* Divide the graph edges into 1/p random groups. Each defines a graph with pm edges. Since the minimum *s*-*t* cut of *G* is *v*, the minimum expected *s*-*t* cut in each group is pv. By the Smoothing Theorem, each sample has minimum *s*-*t* cut, and thus maximum *s*-*t* flow, at least  $(1 - \epsilon)pv$ . Find flows in each piece, and combine the results. This total flow will be  $(1/p)(1 - \epsilon)pv = (1 - \epsilon)v$ .

**Corollary 3.6.** In any undirected graph, given edge strengths, a  $(1 - \epsilon)$ -times maximum flow can found in  $\tilde{O}(m\sqrt{n}/\epsilon)$  time.

*Proof.* Begin by converting the graph to an O(m)-edge (m/n)-smooth graph, as discussed in Lemma 3.3. The Goldberg-Rao flow algorithm [GR97] gives  $T(m,n) = \tilde{O}(m^{3/2})$  for the previous lemma. (Since we are already giving up a factor of  $\epsilon$ , we can assume without loss of generality that all edge capacities are polynomial, thus eliminating the capacity scaling term in their algorithm.) Plugging this in gives a time bound of  $\tilde{O}(m\sqrt{n}/\epsilon)$ .

Unlike for minimum cuts, it is not possible to use the standard augmenting paths algorithm to find a flow in  $\tilde{O}(nv/\epsilon^2)$  time. The graph smoothing process would subdivide unit-cost edges, producing variable cost edges to which unit-capacity augmenting flows cannot be applied.

In previous work [Kar98], Karger used the above techniques to compute exact flows more quickly than before; however, this work has been superseded by better algorithms (also based on edge strength) [KL02b].

### 4 Finding strong connectivities

To efficiently compress and smooth graphs we would like to efficiently find the strong connectivities of edges. Unfortunately, it is not clear that this can be done (*n* maximum flow computations are one slow solution). But as discussed in Theorem 2.10, we do not require the exact values  $k_e$ . We now show that it is possible to find tight strength bounds  $\tilde{k}_e$  that satisfy the two key requirements of that Theorem: that  $\tilde{k}_e \leq k_e$  and  $\sum 1/\tilde{k}_e = O(n)$ . These suffice for the cut and flow algorithms described above.

Our basic plan begins with the following lemma.

**Lemma 4.1.** The total weight of a graph's k-weak edges is at most k(n-1). In particular, any unweighted graph with more than k(n-1) edges has a nontrivial k-strong component (which may be the entire graph).

*Proof.* Let S be the set of k weak edges, and suppose that the total weight of edges in S exceeds k(n-1). Then

$$\sum u_e/k_e \ge \sum_{e \in S} u_e/k_e$$
$$> \sum_{e \in S} u_e/k$$
$$> k(n-1)/k$$
$$= n-1$$

which contradicts Lemma 2.7.

We apply this lemma first to unweighted graphs. Lemma 4.1 says that any unweighted graph with k(n-1) or more edges has a k-strong component. It follows that at most k(n-1) edges are k-weak (that is, have strong connectivity less than k). For otherwise the subgraph consisting of the k-weak edges would have a k-strong component, a contradiction. For each value  $k = 1, 2, 4, 8, \ldots, m$ , we will find a set of k(n-1) edges containing all the k-weak edges (note that every edge is m-weak). We set  $k_e = k/2$  for all edges that are in

the k-weak set but not the k/2-weak set, thus establishing lower bounds for which the Compression Theorem works. The expected number of edges sampled under this basic scheme would be

$$\sum_{i=0}^{\log m} 2^i (n-1)(\rho/2^i) = O(\rho n \log m).$$

We will eventually describe a more sophisticated scheme that eliminates the factor of  $\log m$ . It will also let us handle weighted graphs efficiently.

### 4.1 Sparse Certificates

A basic tool we use is *sparse certificates* defined by Nagamochi and Ibaraki [NI92b].

**Definition 4.2.** A sparse k-connectivity certificate, or simply a k-certificate, for an n-vertex graph G is a subgraph H of G such that

- 1. *H* has k(n-1) edges, and
- 2. H contains all edges crossing cuts of value k or less.

The certificate edges are related to k-weak edges, but are not quite equivalent. Any edge crossing a cut of value less than k is k-weak, but certain k-weak edges will not cross any cut of value less than k. We will show, however, that by finding k-certificate edges one can identify k-weak edges.

Nagamochi and Ibaraki gave an algorithm [NI92b] that constructs a sparse k-connectivity certificate in O(m) time on unweighted graphs, independent of k.

### 4.2 Finding *k*-weak edges

Although a sparse k-certificate contains all edges with *standard* connectivity less than k, it need not contain all edges with *strong* connectivity less than k, since some such edges might not cross any cut of value less than k. We must therefore perform some extra work. In Figure 1 we give an algorithm WeakEdges for identifying edges with  $k_e < k$ . It uses the Nagamochi-Ibaraki Certificate algorithm as a subroutine.

```
procedure WeakEdges(G, k)
do \log_2 n times
E' \leftarrow Certificate(G, 2k)
output E'
G \leftarrow G - E'
end do
```

Figure 1: Procedure WeakEdges for identifying  $k_e < k$ 

**Theorem 4.3.** WeakEdges outputs a set containing all the k-weak edges of G.

*Proof.* First suppose that G has no nontrivial k-strong components, i.e. that  $k_e < k$  for all edges. Then by Lemma 4.1, there are at most k(n-1) edges in G; hence at least half of the vertices have at most 2k incident edges (which define a cut of value at most 2k with a single vertex on one side). In an iteration of the loop in WeakEdges, these vertices become isolated after removing the sparse certificate edges. We have thus shown that in a single loop iteration half of the non-isolated vertices of G become isolated. The remaining graph still has no k-strong edges, so we can repeat the argument. Hence in  $\log_2 n$  rounds we isolate all vertices of G, which can only be done by removing all the edges. Thus all the edges of G are output by WeakEdges.

In the general case, let us obtain a new graph H by contracting each k-strong component of G to a vertex. Any sparse 2k-certificate of G contains the edges of a sparse 2k-certificate of H as well. Thus by the previous paragraph, all edges of H are output by WeakEdges. But these are all the k-weak edges of G.

### 4.3 Sparse partitions

Algorithm WeakEdges can clearly be implemented via  $O(\log n)$  calls to the Nagamochi-Ibaraki Certificate algorithm. It follows that it runs in  $O(m \log n)$  time on unweighted graphs and outputs a set of at most  $k(n-1) \log n$  edges.<sup>4</sup> In this section, we eliminate a  $\log n$  factor in this approach by finding edge sets that are "sparser" than the Nagamochi–Ibaraki certificate.

The first observation we use is that a given k-certificate E' may contain edges that are inside a connected component of G - E'. The edges in G - E' do not cross any cut of value at most k (by definition of a sparse certificate), so the same holds for any edge of E' whose endpoints are connected by a path in G - E'. We can therefore remove any such edge from E' and put it back in G without affecting the correctness of the proof of Theorem 4.3.

We can find the specified reduced edge set by contracting all edges not in E', yielding a new graph G'. This effectively contracts all (and only) edges connected by a path in G - E'. But now observe that any edge crossing a cut of value at most k in G also crosses such a cut in G' since we contract no edge that crosses such a small cut. Thus we can find all edges crossing a small cut via a certificate in G'. Since G' has fewer vertices, the certificate has fewer edges. We can iterate this procedure until all edges in the certificate cross some cut of value at most k or until G' becomes a single vertex. In the latter case, the original graph is k-connected, while in the former, if the current contracted graph has n' vertices, it has at most k(n'-1)edges. This motivates the following definition:

**Definition 4.4.** A sparse k-partition, or k-partition, of G is a set E' of edges of G such that

- 1. E' contains all edges crossing cuts of value k or less in G, and
- 2. If G E' has r connected components, then E' contains at most 2k(r-1) edges.

In fact, the construction just described yields a graph with at most k(r-1) edges, but we have relaxed the definition to 2k(r-1) edges to allow for an efficient construction.

Procedure Partition in Figure 2 outputs a sparse partition. It uses the Nagamochi–Ibaraki Certificate algorithm and obtains a new graph G' by contracting those edges not in the certificate. It repeats this process until the graph is sufficiently sparse.

#### procedure Partition(G, k)

input: An *n*-vertex *m*-edge graph G

 $\begin{array}{l} \textbf{if } m \leq 2k(n-1) \textbf{ then} \\ \textbf{output } \textbf{the edges of } G \\ \textbf{else} \\ E' \leftarrow \texttt{Certificate}(G,k) \\ G' \leftarrow \texttt{contract all edges of } G - E' \\ \texttt{Partition}(G',k) \end{array}$ 

### Figure 2: Partition finds low-connectivity edges

**Lemma 4.5.** Partition outputs a sparse k-partition partition in O(m) time on unweighted graphs.

<sup>&</sup>lt;sup>4</sup>It also follows that a  $k \log n$  sparse-certificate will contain all k-weak edges, so they can be found with a single Certificate invocation. This gives a better running time. Indeed, since the Nagamich Ibaraki algorithm "labels" each edge with the value k for which it vanishes, we can use those labels (divide by  $\log n$ ) as strength lower-bounds, producing a complete result in  $O(m + n \log n)$  time. However, this approach produces an extra  $\log n$  factor in the edge bound (or worse in weighted graphs) that we have been unable to remove.

*Proof.* Correctness is clear since no edge crossing a cut of value less than k is ever contracted and at termination  $m \leq 2k(n-1)$ ; we need only bound the running time. If initially m < k(n-1) then the algorithm immediately terminates. So we can assume  $m \geq k(n-1)$ .

Suppose that in some iteration m > 2k(n-1). We find a sparse connectivity certificate with  $m' \le k(n-1)$  edges and then contract the graph to n' vertices. If n'-1 > (n-1)/2 then in the following iteration we will have  $m' \le k(n-1) < 2k(n'-1)$  and the algorithm will terminate. It follows that the number of vertices (minus one) halves in every recursive call except the last.

A single iteration involves the O(m)-time sparse-certificate algorithm [NI92b]. At each recursive call, the edges remaining are all k-certificate edges from the previous iteration. The number of such certificate edges is at most k times the number of vertices—thus the (upper bound on the) number of edges halves in each recursive call. It follows that after the first call we have T(n) = O(kn) + T(n/2) = O(kn). This is O(m) since  $m \ge k(n-1)$  by assumption.

**Lemma 4.6.** If Partition is used instead of Certificate in a call to WeakEdges(G, k) (meaning we invoke Partition(G, 2k) instead of Certificate(G, 2k)), then algorithm WeakEdges runs in  $O(m \log n)$  time on unweighted graphs and returns a partition of G into r components for some r. There are at most 4k(r-1) cross-partition edges and they include all the k-weak edges of G.

Note that the partition output by WeakEdges is itself almost a sparse k-partition; it simply has twice as many edges as the definition allows. On the other hand, it contains all k-weak edges; not just the ones crossing small cuts.

*Proof.* The running time follows from the previous lemma. To prove the edge bound, consider a particular connected component H remaining in a particular iteration of WeakEdges. A call to Partition(H, 2k) returns a set of 4k(s-1) edges that breaks that component into s subcomponents (the multiplier 4 arises from the fact that we look for a 2k-partition). That is, it uses at most 4k(s-1) edges to increase the number of connected components by s-1. We can therefore charge 4k edges to each of the new components that gets created. Accumulating these charges over all the calls to Partition shows that if WeakEdges outputs 4k(r-1) edges then those edges must split the entire graph into at least r components.

### 4.4 Assigning Estimates

We now give an algorithm Estimation in Figure 3 for estimating strong connectivities. We use subroutine WeakEdges to find a small edge set containing all edges e with  $k_e < k$  but replace the Nagamochi-Ibaraki Certificate implementation with our algorithm Partition to reduce the number of output edges.

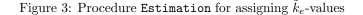
We assign values  $k_e$  as follows. In the first step, we run WeakEdges on G with k = 2; we set  $k_e = 1$  for the edges in the output edge set  $E_0$ . Then we delete  $E_0$  from G; this breaks G into connected components  $G_1, \ldots, G_\ell$ . Note that each edge in  $G_i$  has  $k_e \ge 2$  in G, though possibly not in  $G_i$ . Then we recursively repeat this procedure in each  $G_i$ , by setting k = 4 in WeakEdges and labeling all output edges with  $\tilde{k}_e = 2$ , then with  $k = 8, 16, \ldots, m$ . At the  $i^{th}$  step, all as-yet unlabeled edges have  $k_e \ge 2^i$ ; we separate all those with  $k_e < 2^{i+1}$  and give them (valid lower bound) label  $\tilde{k}_e = 2^i$ . Thus we find all  $\tilde{k}_e$ -values in at most  $\log m$ iterations since m is the maximum strength of an edge in an unweighted graph.

**Lemma 4.7.** If H is any subgraph of G, then Estimation(H, k) assigns lower bounds  $\tilde{k}_e \leq k_e$  for all edges  $e \in H$  with  $k_e \geq k$  in G.

**Corollary 4.8.** After a call to Estimation(G, 1), all the labels  $\tilde{k}_e$  satisfy  $\tilde{k}_e \leq k_e$ .

*Proof.* We prove the lemma by induction on the size of H. The base case of a graph with no edges is clear. To prove the inductive step we need only consider edges e with  $k_e \ge k$ . We consider two possibilities. If e is in the set E' returned by WeakEdges(H, 2k) then it receives label k, which is a valid lower bound for any edge with  $k_e \ge k$ . So the inductive step is proved for  $e \in E'$ . On the other hand, if  $e \notin E'$ , then e is in some H' upon which the algorithm is invoked recursively. By the correctness of WeakEdges we know  $k_e \ge 2k$  (in

procedure Estimation(H, k)input: subgraph H of G  $E' \leftarrow$  WeakEdges(H, 2k)for each  $e \in E'$   $k_e \leftarrow k$ for each nontrivial connected component  $H' \subset H - E'$ Estimation(H', 2k)



H, and thus in G) in this case. Thus, the inductive hypothesis applies to show that e receives a valid lower bound upon invocation of WeakEdges(H', 2k).

**Lemma 4.9.** Assume that in procedure WeakEdges, procedure Certificate is replaced by Partition. Then the values  $\tilde{k}_e$  output by Estimation(G, 1) are such that  $\sum 1/\tilde{k}_e = O(n)$ .

Proof. The proof is similar to the proof that  $\sum u_e/k_e \leq n$ . Define the cost of edge e to be  $1/\tilde{k}_e$ . We prove that the total cost assigned to edges is O(n). Consider a call to Estimation(H,k) on some remaining connected component of G. It invokes WeakEdges(H,k), which returns a set of 4k(r-1) edges whose removal partitions H into r connected components. (Note that possibly r = 0 if H is k-connected.) The algorithm assigns values  $\tilde{k}_e = k$  to the removed edges. It follows that the total cost assigned to these edges is 4(r-1). In other words, at a cost of 4(r-1), the algorithm has increased the number of connected components by r-1. Ultimately, when all vertices have been isolated by edge removals, there are n components; thus, the total cost of the component creations is at most 4(n-1).

In summary, our estimates  $\tilde{k}_e$  satisfy the necessary conditions for our Compression and Smoothing applications:  $\tilde{k}_e \leq k_e$  and  $\sum 1/\tilde{k}_e = O(n)$ .

**Lemma 4.10.** Estimation runs in  $O(m \log^2 n)$  time on an unweighted graph.

*Proof.* Each level of recursion of Estimation calls subroutine WeakEdges on graphs of total size m. An unweighted graph has maximum strong connectivity m and therefore has  $O(\log m)$  levels of recursion.

### 4.5 Weighted graphs

Until now, we have focused on the estimation of edge strengths for unweighted graphs. When graphs are weighted, things are more difficult.

Nagamochi and Ibaraki give an  $O(m+n\log n)$ -time weighted-graph implementation of their Certificate algorithm [NI92a]. (In weighted graphs, the k-sparse sparse certificate has an upper bound of k(n-1) on the total weight of edges incorporated.) We can use the Nagamochi-Ibaraki weighted-graph algorithm to implement Partition(G, k) in  $O(m \log n)$  time for any value of k. Unlike the unweighted case, the repeated calls to Certificate need not decrease the number of edges substantially (though their total weight will decrease). However, the claimed halving in vertices still happens. Thus algorithm Partition satisfies a recurrence  $T(m,n) = O(m+n\log n) + T(m,n/2) = O(m\log n)$ . Since Partition runs in  $O(m\log n)$  time, we deduce that WeakEdges runs in  $O(m\log^2 n)$  time.

A bigger problem arises in the iterations of Estimation. In a weighted graph with maximum edge weight W, the  $k_e$  values may be as large as  $n^2W$ , meaning that  $\Omega(\log nW)$  levels of recursion will apparently be required in Estimation. This can be a problem if W is superpolynomial. To deal with this problem, we show how to localize our computation of strong connectivities to a small "window" of relevant connectivity values.

We begin by computing a rough underestimate for the edge strengths. Construct a maximum spanning tree (MST) for G using the weights  $u_e$ . Let  $d_e$  be the minimum weight of an edge on the MST-path between the endpoints of e. The quantities  $d_e$  can be determined in O(m) time using an MST sensitivity analysis algorithm [DRT92] (practical algorithms run in  $O(m \log n)$  time and will not dominate the running time). Since the MST path between the endpoints of e forms a (nonmaximal)  $d_e$ -connected subgraph containing e, we know that  $k_e \ge d_e$ . However, if we remove all edges of weight  $d_e$  or greater, then we disconnect the endpoints of e (this follows from maximum spanning tree properties [Tar83]). There are at most  $\binom{n}{2}$  such edges, so the weight removed is at most  $n^2 d_e$ . Therefore,  $k_e \le n^2 d_e$ . This gives us an initial factor-of- $n^2$  estimate  $d_e \le k_e \le n^2 d_e$ .

Our plan is to compute the  $\tilde{k}_e$  in a series of phases, each focusing on a set of edges with narrow range of  $d_e$  values. In particular, we will contract all edges with  $d_e$  above some upper bound, and delete all edges with  $d_e$  below some lower bound. Then we will use Estimation to assign  $\tilde{k}_e$  labels to the edges that remain.

# **Lemma 4.11.** If we contract a set of edges, all of which have weights at least W, then the strengths of edges with original strength less than W are unchanged.

Proof. Consider an edge e with strength  $k_e$ , and suppose that its strength is  $k'_e$  in the contracted graph. It follows that there is some maximal  $k'_e$ -connected component H' containing e in the contracted graph. Consider the preimage H of this component in G—that is, the set of vertices that get contracted into H'. This component is at best  $k_e$ -connected in G by the definition of  $k_e$ . It follows that there is some cut of value  $k_e$  in this component. The edges of this cut have value at most  $k_e$ , so contracting edges of value exceeding  $k_e$  cannot destroy this cut. Thus, the connectivity of H' is at most  $k_e$ . It follows that  $k'_e \leq k_e$ . Since contracting edges cannot decrease connectivities, we deduce  $k'_e = k_e$ .

We label our edges in a series of phases. In a phase, let D be the maximum  $d_e$  on any unlabelled edge. Since  $k_e \leq n^2 d_e$ , the maximum strength of any unlabelled edge is at most  $n^2 D$ . Our goal in one phase is to (validly) label all edges with  $d_e \geq D/n$ . We begin by contracting all edges of weight exceeding  $n^2 D$ . By the previous lemma, the contractions do not affect strengths of edges with  $k_e \leq n^2 D$  (which includes all unlabelled edges). In the resulting graph, let us delete all edges with  $d_e < D/n$  (since  $d_e \leq k_e$ , no edge we want to label is deleted). The deletions may decrease certain strengths but not increase them. It follows that every unlabelled edge (all of which have  $k_e \leq Dn^2$ ) has strength in the modified graph *no greater* than in G.

On each connected component H induced by the remaining edges, execute Estimation(H, D/n). By Lemma 4.7, this assigns valid lower-bound labels to all edges e with strength at least D/n (in the modified graph). In particular, the labels are valid for all e with  $d_e \ge D/n$  (since any edge with  $d_e \ge D/n$  is connected by a path of edges of value at least D/n, none of which get deleted in the phase). These labels are valid lower bounds for strengths in the modified graph; however, as discussed in the previous paragraph, all unlabelled edges have the strengths in the subgraph no greater than their strength in G. Thus, the computed labels can be used as valid labels for all the unlabelled edges with  $d_e \ge D/n$ .

The approach just described has computed labels for each unlabelled edge with  $d_e \ge D/n$ . We have therefore reduced the maximum  $d_e$  on any unlabelled edge by a factor of n. We iterate this process, continuously decreasing the maximum unlabelled d(e), until all edges are labelled.

Summarizing our discussion above gives the algorithm WindowEstimation listed in Figure 4.

### **Lemma 4.12.** Procedure WindowEstimation can be implemented to run in $O(m \log^2 n)$ time.

*Proof.* The contractions in WindowEstimation can be implemented using a standard union-find data structure [CLR90]. Each time an edge is contracted, a union is called on its endpoints. Each time an edge is added from L, find operations can identify its endpoints. Therefore, the additions and contractions of edges do not affect the running time. Instead, the running time is determined by the repeated calls to Estimation.

Consider a particular iteration of the loop with some D value. We initially contract all edges with  $d_e > n^2 D$ , so that the maximum strength in the resulting graph is at most  $n^4 D$ . We invoke Estimation

```
procedure WindowEstimation(G)

Sort the edges in decreasing order of d_e into a list L

initialize G' as an empty graph on the vertices of G

repeat

let D \leftarrow \text{maximum } d_e among unlabelled edges in L

contract every e \in G' with d(e) > n^2D

move every edge e \in L with d_e \ge D/n to G'

call Estimation(G', D/n) to get labels \tilde{k}_e

for the new edges added from L in this phase

until no edges remain
```

Figure 4: WindowEstimation for weighted graphs

with a starting strength argument of D/n, which means that it terminates in  $O(\log n)$  iterations (the number of argument doublings from D/n to  $n^4D$ ). As to the size of the problem, recall that we contracted all edges with with  $d_e \ge n^2D$  and deleted all edges with  $d_e < D/n$ . It follows that our running time is proportional to  $m' \log^3 n$  where m' is the number of edges with  $D/n \le d_e \le D$ .

Now we can bound the running time over all phases. An edge d(e) is present (neither contracted nor deleted) if and only if  $D/n \leq D < n^2 D$ . Since the threshold D decreases by a factor of n each time, this means that edge e contributes to the size of the evaluated subgraph in at most 3 iterations. In other words, the sum of m' values over all iterations of our algorithm is 3m. It follows that the overall running time of these iterations is  $O(\sum m' \log^3 n) = O(m \log^3 n)$ .

**Lemma 4.13.** Procedure WindowEstimation assigns labels such that  $\sum u_e/\tilde{k}_e = O(n)$ 

*Proof.* Recall the definition of cost of edge e as  $u_e/\tilde{k}_e$ . Our algorithm incorporates some of the labels computed by Estimation in each phase, contributing their cost (in that phase) to the final total cost. We show that the total cost of *all* labels computed over all the phases is O(n).

We invoke the concept of *rank*. The rank of a graph is equal to the number of edges in a spanning tree of the graph. Inspection of Partition shows that the total weight of edges returned by Partition(G, k) is at most 4 times the rank of G. Similarly, inspection of Estimation show that on a rank-r graph, its results satisfy  $\sum u_e/\tilde{k}_e = O(r)$ .

In a phase, we contract all edges of weight exceeding  $Dn^2$  and delete all edges with weight less than D. By the properties of maximum spanning trees, the resulting graph is precisely spanned by the set of MST edges with weights in this range. That is, the rank of this graph is equal to the number  $r_D$  of such MST edges. It follows that the total cost  $\sum u_e/\tilde{k}_e$  of Estimation labels in this phase is  $O(r_D)$ . Now note that each MST edge contributes to  $r_D$  only when its weight is between D and  $Dn^2$ , which happens in at most 3 phases since D decreases by n each phase. Thus, each edge contributes to  $3 r_D$  values, so  $\sum r_D \leq 3(n-1)$ . This bounds the total cost by  $O(\sum r_D) = O(n)$ , as desired.

### 5 Conclusion

We have given new, stronger applications of random sampling to problems involving cuts in graphs. The natural open question is whether these approximation algorithms can be made exact. An initial step towards the answer was given in [Kar99], but it only gives a useful speedup for graphs with large minimum cuts. More recently, sampling has led to an exact linear-time algorithm for minimum cuts [Kar00]; however, the techniques used there appear to be specialized to that particular problem. Karger and Levine [KL02a] have

recently given very fast algorithms for flows in unweighted graphs; the important remaining question is to develop fast exact algorithms for weighted graphs.

A more limited open question has to do with the use of strong connectivity. We introduced strong connectivity in order to make our theorems work. Many of the intuitions about our theorems apply even to the standard connectivity notion in which the connectivity of edge (u, v) is defined to be the minimum u-v cut in G. We have no counterexample to the conjecture that using these weak connectivities would suffice in our algorithms. Such a change would likely simplify our algorithms and presentation (though the time bounds are unlikely to change).

### A The General Weighted Sampling Theorem

For possible future use, we give a general theorem on when a weighted random graph has all cut values tightly concentrated near their expectation. The compression theorem and smooth graph sampling theorems are special cases of this theorem.

**Theorem A.1.** Let G be a random graph in which the weight  $U_e$  of edge e has a probability distribution with expectation  $u_e$  and maximum value  $m_e$ . Let  $k_e$  be the strength of edge e in the graph where each edge e gets weight  $E[U_e]$ . If for every edge,  $k_e \geq 2m_e(\ln n)/\epsilon^2$ , then with high probability, every cut in G has value within  $(1 \pm \epsilon)$  times its expectation.

*Proof.* Order the distinct edge strengths  $k_1, \ldots, k_r$  in H in increasing order. Let  $F_i$  be the graph consisting of all of  $k_i$ -strong edges in H, with edge e given weight  $U_e/k_e$  (so  $F_i$  is a random graph). Observe that  $G = \sum (k_i - k_{i-1})F_i$ . So if every  $F_i$  is near its expectation, it follows that G is near its expectation.

So consider (a component of) graph  $F_i$ . The expected value of a cut in  $F_i$  has the form  $\sum u_e/k_e \ge 1$  by Lemma 2.7. In other words, the minimum cut in  $E[F_i]$  is at least 1. On the other hand, the maximum value attained by any edge in  $F_i$  is  $U_e/k_e \le m_e/k_e$ . By Lemma 1.7, it follows that  $F_i$  has all cuts within  $(1 \pm \epsilon)$  of its expectation with high probability.

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