Streaming Graph Computations with a Helpful Advisor

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Abstract

Motivated by the trend to outsource work to commercial cloud computing services, we consider a variation of the streaming paradigm where a streaming algorithm can be assisted by a powerful helper that can provide annotations to the data stream. We extend previous work on such *annotation models* by considering a number of graph streaming problems. Without annotations, streaming algorithms for graph problems generally require significant memory; we show that for many standard problems, including all graph problems that can be expressed with totally unimodular integer programming formulations, only a constant number of hash values are needed for single-pass algorithms given linear-sized annotations. We also obtain a protocol achieving *optimal* tradeoffs between annotation length and memory usage for matrix-vector multiplication; this result contributes to a trend of recent research on numerical linear algebra in streaming models.

1 Introduction

The recent explosion in the number and scale of real-world structured data sets including the web, social networks, and other relational data has created a pressing need to efficiently process and analyze massive graphs. This has sparked the study of graph algorithms that meet the constraints of the standard streaming model: restricted memory and the ability to make only one pass (or few passes) over adversarially ordered data. However, many results for graph streams have been negative, as many foundational problems require either substantial working memory or a prohibitive number of passes over the data [1]. Apparently most graph algorithms fundamentally require flexibility in the way they query edges, and therefore the combination of adversarial order and limited memory makes many problems intractable.

To circumvent these negative results, variants and relaxations of the standard graph streaming model have been proposed, including the Semi-Streaming [2], W-Stream [3], Sort-Stream [4], Random-Order [1], and Best-Order [5] models. In Semi-Streaming, memory requirements are relaxed, allowing space proportional to the number of vertices in the stream but not the number

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of edges. The W-Stream model allows the algorithm to write temporary streams to aid in computation. And, as their names suggest, the Sort-Stream, Random-Order, and Best-Order models relax the assumption of adversarially ordered input. The Best-Order model, for example, allows the input stream to be re-ordered arbitrarily to minimize the space required for the computation.

In this paper, our starting point is a relaxation of the standard model, closest to that put forth by Chakrabarti *et al.* [6], called the *annotation model*. Motivated by recent work on outsourcing of database processing, as well as commercial cloud computing services such as Amazon EC2, the annotation model allows access to a powerful advisor, or *helper* who observes the stream concurrently with the algorithm. Importantly, in many of our motivating applications, the helper is not a trusted entity: the commercial stream processing service may have executed a buggy algorithm, experienced a hardware fault or communication error, or may even be deliberately deceptive [5, 6]. As a result, we require our protocols to be *sound*: our *verifier* must detect any lies or deviations from the prescribed protocol with high probability.

The most general form of the annotation model allows the helper to provide additional annotations in the data stream *at any point* to assist the verifier, and one of the cost measures is the total length of the annotation. In this paper, however, we focus on the case where the helper's annotation arrives as a single message after both the helper and verifier have seen the stream. The helper's message is also processed as a stream, since it may be large; it often (but not always) includes a re-ordering of the stream into a convenient form, as well as additional information to guide the verifier. This is therefore stronger than the Best-Order model, which only allows the input to be reordered and no more; but it is weaker than the more general *online* model, because in our model the annotation appears only after the input stream has finished.

We argue that this model is of interest for several reasons. First, it requires minimal coordination between helper and verifier, since it is not necessary to ensure that annotation and stream data are synchronized. Second, it captures the case when the verifier uploads data to the cloud as it is collected, and later poses questions over the data to the helper. Under this paradigm, the annotation must come *after* the stream is observed. Third, we know of no non-trivial problems which separate the general online and our "at-the-end" versions of the model, and most prior results are effectively in this model.

Besides being practically motivated by outsourced computations, annotation models are closely related to Merlin-Arthur proofs with space-bounded verifiers, and studying what can (and cannot) be accomplished in these models is of independent interest.

Relationship to Other Work. Annotation models were first explicitly studied by Chakrabarti *et al.* in [6], and focused primarily on protocols for canonical problems in numerical streams, such as Selection, Frequency Moments, and Frequent Items. The authors also provided protocols for some graph problems: counting triangles, connectedness, and bipartite perfect matching. The Best-Order Stream Model was put forth by Das Sarma et al. in [5]. They present protocols requiring logarithmic or polylogarithmic space (in bits) for several problems, including perfect matching and connectivity. Historical antecedents for this work are due to Lipton [7], who used fingerprinting methods to verify polynomial-time computations in logarithmic space. Recent work verifies shortest-path computations using cryptographic primitives, using polynomial space for the verifier [8].

Our Contributions. We identify two qualitatively different approaches to producing protocols for problems on graphs with n nodes and m edges. In the first, the helper directly proves matching upper and lower bounds on a quantity. Usually, proving one of the two bounds is trivial: the helper

provides a feasible solution to the problem. But proving *optimality* of the feasible solution can be more difficult, requiring the use of structural properties of the problem. In the second, we simulate the execution of a non-streaming algorithm, using the helper to maintain the algorithm's internal data structures to control the amount of memory used by the verifier. The helper must provide the contents of the data structures so as to limit the amount of annotation required.

Using the first approach (Section 3), we show that only constant space and annotation linear in the input size m is needed to determine whether a directed graph is a DAG and to compute the size of a maximum matching. We describe this as an (m, 1) protocol, where the first entry refers to the annotation size (which we also call the hcost) and the second to the memory required for the verifier (which we also call the vcost). Our maximum matching result significantly extends the bipartite perfect matching protocol of [6], and is tight for dense graphs, in the sense that there is a lower bound on the product of hcost and vcost of hcost \cdot vcost = $\Omega(n^2)$ bits for this problem. Second, we define a streaming version of the linear programming problem, and provide an (m, 1) protocol. By exploiting duality, we hence obtain (m, 1) protocols for many graph problems with totally unimodular integer programming formulations, including shortest *s*-*t* path, max-flow, min-cut, and minimum-weight bipartite perfect matching. We also show all are tight by proving lower bounds of hcost \cdot vcost = $\Omega(n^2)$ bits for all four problems. A more involved protocol obtains optimal tradeoffs between annotation cost and working memory for dense LPs and matrix-vector multiplication; this complements recent results on approximate linear algebra in streaming models (see e.g. [9, 10]).

For the second approach (Section 4), we make use of the idea of "memory checking" due to Blum et al. [11], which allows a small-space verifier to outsource data storage to an untrusted server. We present a general simulation theorem based on this checker, and obtain as corollaries tight protocols for a variety of canonical graph problems. In particular, we give an (m, 1) protocol for verifying a minimum spanning tree, an $(m+n \log n, 1)$ protocol for single-source shortest paths, and an $(n^3, 1)$ protocol for all-pairs shortest paths. We provide a lower bound of hcost \cdot vcost = $\Omega(n^2)$ bits for the latter two problems, and an identical lower bound for MST when the edge weights can be given incrementally. While powerful, this technique has its limitations: there does not seem to be any generic way to obtain the same kind of tradeoffs observed above. Further, there are some instances where direct application of memory checking does not achieve the best bounds for a problem. We demonstrate this by presenting an $(n^2 \log n, 1)$ protocol to find the diameter of a graph; this protocol leverages the ability to use randomized methods to check computations more efficiently than via generating or checking a deterministic witness. In this case, we rely on techniques to verify matrix-multiplication in quadratic time, and show that this is tight via a nearly matching lower bound for diameter of hcost \cdot vcost = $\Omega(n^2)$.

In contrast to problems on numerical streams, where it is often trivial to obtain (m, 1) protocols by replaying the stream in sorted order, it transpires that achieving linear-sized annotations with logarithmic space is more challenging for many graph problems. Simply providing the solution (e.g. a graph matching or spanning tree) is insufficient, since we have the additional burden of demonstrating that this solution is *optimal*. A consequence is that we are able to provide solutions to several problems for which no solution is known in the best-order model (even though one can reorder the stream in the best-order model so that the "solution" edges arrive first).

2 Model and Definitions

Consider a data stream $\mathcal{A} = \langle a_1, a_2, \ldots, a_m \rangle$ with each a_i in some universe \mathcal{U} . Consider a probabilistic verifier \mathcal{V} who observes \mathcal{A} and a deterministic helper \mathcal{H} who also observes \mathcal{A} and can send a message h to \mathcal{V} after \mathcal{A} has been observed by both parties. This message, also referred to as an annotation, should itself be interpreted as a data stream that is parsed by \mathcal{V} , which may permit \mathcal{V} to use space sublinear in the size of the annotation itself. That is, \mathcal{H} provides an annotation $h(\mathcal{A}) = (h_1(\mathcal{A}), h_2(\mathcal{A}), \ldots, h_{\ell}(\mathcal{A})).$

We study randomized streaming protocols for computing functions $f(\mathcal{A}) \to \mathbb{Z}$. Specifically, assume \mathcal{V} has access to a private random string \mathcal{R} and at most w(m) machine words of working memory, and that \mathcal{V} has one-way access to the input $\mathcal{A} \cdot h$, where \cdot represents concatenation. Denote the output of protocol \mathcal{P} on input \mathcal{A} , given helper h and random string \mathcal{R} , by $out(\mathcal{P}, \mathcal{A}, \mathcal{R}, h)$. We allow \mathcal{V} to output \perp if \mathcal{V} is not convinced that the annotation is valid. We say that h is *valid* for \mathcal{A} with respect to \mathcal{P} if $Pr_{\mathcal{R}}(out(\mathcal{P}, \mathcal{A}, \mathcal{R}, h) = f(\mathcal{A})) = 1$, and we say that h is δ -invalid for \mathcal{A} with respect to \mathcal{P} if $Pr_{\mathcal{R}}(out(\mathcal{P}, \mathcal{A}, \mathcal{R}, h) \neq \perp) \leq \delta$. We say that h is a valid helper if h is valid for all \mathcal{A} . We say that \mathcal{P} is a valid protocol for f if

- 1. There exists at least one valid helper h with respect to \mathcal{P} and
- 2. For all helpers h' and all streams \mathcal{A} , either h' is valid for \mathcal{A} or h' is $\frac{1}{3}$ -invalid for \mathcal{A} .

Conceptually, \mathcal{P} is a valid protocol for f if for each stream \mathcal{A} there is *at least* one way to convince \mathcal{V} of the true value of $f(\mathcal{A})$, and \mathcal{V} rejects all other annotations as invalid (this differs slightly from [6] to allow for multiple h's that can convince \mathcal{V}). The constant $\frac{1}{3}$ can be any constant less than $\frac{1}{2}$.

Let h be a valid helper chosen to minimize the length of $h(\mathcal{A})$ for all \mathcal{A} . We define the help cost hcost(\mathcal{P}) to be the maximum length of h over all \mathcal{A} of length m, and the verification cost vcost(P) = w(m), the amount of working memory used by the protocol P. All costs are expressed in machine words of size $\Theta(\log m)$ bits, i.e. we assume any quantity polynomial in the input size can be stored in a constant number of words; in contrast, lower bounds are expressed in bits. We say that \mathcal{P} is an (h, v) protocol for f if \mathcal{P} is valid and hcost(\mathcal{A}) = O(h + 1), vcost(\mathcal{A}) = O(v + 1). While both hcost and vcost are natural costs for such protocols, we often aim to achieve a vcost of O(1) and then minimize hcost. In other cases, we show that hcost can be decreased by increasing vcost, and study the tradeoff between these two quantities.

In some cases, f is not a function of \mathcal{A} alone; instead it depends on \mathcal{A} and h. In such cases, \mathcal{V} should simply *accept* if convinced that the annotation has the correct properties, and output \bot otherwise. We use the same terminology as before, and say that \mathcal{P} is a valid protocol if there is a valid helper and any \mathfrak{h}' that is not valid for \mathcal{A} is $\frac{1}{3}$ -invalid for \mathcal{A} .

In this paper we primarily consider graph streams, which are streams whose elements are edges of a graph G. More formally, consider a stream $\mathcal{A} = \langle e_1, e_2, \ldots, e_m \rangle$ with each $e_i \in [n] \times [n]$. Such a stream defines a (multi)graph G = (V, E) where $V = \{v_1, \ldots, v_n\}$ and E is the (multi)set of edges that naturally corresponds to \mathcal{A} . We use the notation $\{i : m(i)\}$ for the multiset in which *i* appears with multiplicity m(i). Finally, we will sometimes consider graph streams with directed edges, and sometimes with weighted edges; in the latter case each edge $e_i \in [n] \times [n] \times \mathbb{Z}_+$.

2.1 Fingerprints

Our protocols make careful use of *fingerprints*, permutation-invariant hashes that can be efficiently computed in a streaming fashion. They determine in small space (with high probability) whether

two streams have identical frequency distributions. They are the workhorse of algorithms proposed in earlier work on streaming models with an untrusted helper [5, 6, 7, 12]. We sometimes also need the fingerprint function to be linear.

Definition 2.1 (Fingerprints). A fingerprint of a multiset $M = \{i : m(i)\}$ where each $i \in [q]$ for some known upper bound q is defined as a computation over the finite field with p elements, \mathbb{F}_p , as $\mathfrak{f}_{p,\alpha}(M) = \sum_{i=1}^q m(i)\alpha^i$, where α is chosen uniformly at random from \mathbb{F}_p . We typically leave p, α implicit, and just write $\mathfrak{f}(M)$.

Some properties of \mathfrak{f} are immediate: it is linear in M, and can easily be computed incrementally as elements of [q] are observed in a stream one by one. The main property of \mathfrak{f} is that $\Pr[\mathfrak{f}(M) = \mathfrak{f}(M')|M \neq M'] \leq q/p$ over the random choice of α (due to standard properties of polynomials over a field). Therefore, if p is sufficiently large, say, polynomial in q and in an (assumed) upper bound on the multiplicities m(i), then this event happens with only polynomially small probability. For cases when the domain of the multisets is not [q], we either establish a bijection to [q] for an appropriate value of q, or use a hash function to map the domain onto a large enough [q] such that there are no collisions with high probability (whp). In all cases, p is chosen to be O(1) words.

A common subroutine of many of our protocols forces \mathcal{H} to provide a "label" l(u) for each node upfront, and then replay the edges in E, with each edge (u, v) annotated with l(u) and l(v) so that each instance of each node v appears with the same label l(v).

Definition 2.2. We say a list of edges E' is label-augmented if (a) E' is preceded by a sorted list of all the nodes $v \in V$, each with a value l(v) and deg(v), where l(v) is the label of v and deg(v) is claimed to be the degree of v; and (b) each edge e = (u, v) in E' is annotated with a pair of symbols l(e, u) and l(e, v). We say a list of label-augmented edges E' is valid if for all edges e = (u, v), l(e, u) = l(u) and l(e, v) = l(v); and E' = E, where E is the set of edges observed in the stream A.

Lemma 2.3 (Consistent Labels). There is a valid (m, 1) protocol that accepts any valid list of label-augmented edges.

Proof. \mathcal{V} uses the annotation from Definition 2.2 (a) to make a fingerprint of the multiset $S_1 := \{(u, l(u)) : \deg(u)\}$. \mathcal{V} also maintains a fingerprint f_1 of all (u, l(e, u)) pairs seen while observing the edges of L. If $f_1 = \mathfrak{f}(S_1)$ then (whp) each node u must be presented with label l(e, u) = l(u) every time it is reported in an edge e (and moreover u must be reported in exactly deg(u) edges), else the multiset of observed (node, label) pairs would not match S_1 . Finally, \mathcal{V} ensures that E' = E by checking that $\mathfrak{f}(E) = \mathfrak{f}(E')$.

3 Directly Proving Matching Upper and Lower Bounds

3.1 Warmup: Topological Ordering and DAGs

A (directed) graph G is a DAG if and only if G has a topological ordering, which is an ordering of V as $v_1, \ldots v_n$ such that for every edge (v_i, v_j) we have i < j [13, Section 3.6]. Hence, if G is a DAG, \mathcal{H} can prove it by providing a topological ordering. If G is not a DAG, \mathcal{H} can provide a directed cycle as witness.

Theorem 3.1. There is a valid (m, 1) protocol to determine if a graph is a DAG.

Proof. If G is not a DAG, \mathcal{H} provides a directed cycle C as $(v_1, v_2), (v_2, v_3) \dots (v_k, v_1)$. To ensure $C \subseteq E, \mathcal{H}$ then provides $E \setminus C$, allowing \mathcal{V} to check that $\mathfrak{f}(C \cup (E \setminus C)) = \mathfrak{f}(E)$.

If G is a DAG, let $v_1, \ldots v_n$ be a topological ordering of G. We require \mathcal{H} to replay the edges of G, with edge (v_i, v_j) annotated with the ranks of v_i and v_j i.e. i and j. We ensure \mathcal{H} provides consistent ranks via the Consistent Labels protocol of Lemma 2.3, with the ranks as "labels". If any edge (v_i, v_j) is presented with j > i, \mathcal{V} rejects immediately.

3.2 Maximum Matching

We give an (m, 1) protocol for maximum matching which leverages the combinatorial structure of the problem. Previously, matching was only studied in the bipartite case, where an (m, 1) protocol and a lower bound of hcost \cdot vcost = $\Omega(n^2)$ bits for dense graphs were shown [6, Theorem 11]. The same lower bound applies to the more general problem of maximum matching, so our protocol is tight up to logarithmic factors.

The protocol shows matching upper and lower bounds on the size of the maximum matching. Any feasible matching presents a lower bound. For the upper bound we appeal to the Tutte-Berge formula [14, Chapter 24]: the size of a maximum matching of a graph G = (V, E) is equal to $\frac{1}{2}\min_{V_S \subseteq V}(|V_S| - \operatorname{occ}(G - V_S) + |V|)$, where $G - V_S$ is the subgraph of G obtained by deleting the vertices of V_S and all edges incident to them, and $\operatorname{occ}(G - V_S)$ is the number of components in the graph $G - V_S$ that have an odd number of vertices. So for any set of nodes V_S , $\frac{1}{2}(|V_S| - \operatorname{occ}(G - V_S) + |V|)$ is an upper bound on the size of the maximum matching, and there exists some V_S for which this quantity equals the size of a maximum matching M. Conceptually, providing both V_S and M, \mathcal{H} proves that the maximum matching size is M. Additionally, \mathcal{H} has to provide a proof of the value of $\operatorname{occ}(G - V_S)$ to \mathcal{V} .

Theorem 3.2. There is a valid (m, 1) protocol for maximum matching. Moreover, any protocol for max-matching requires hcost \cdot vcost = $\Omega(n^2)$ bits.

Proof. To prove a lower bound of k on the size of the maximum matching, \mathcal{H} provides a matching $M = (V_M, E_M)$ of size $|E_M| = k$, and then proves that M is indeed a matching. It suffices to prove that $|V_M| = 2|E_M|$ and $M \subseteq E$. First, \mathcal{H} lists E_M , and \mathcal{V} fingerprints the nodes present as $f(V_M)$. \mathcal{H} then presents V'_M which is claimed to be V_M in sorted order, allowing \mathcal{V} to easily check no node appears more than once and that $\mathfrak{f}(V_M) = \mathfrak{f}(V'_M)$. Next, \mathcal{H} provides $E \setminus M$, allowing \mathcal{V} to check that $\mathfrak{f}(M \cup (E \setminus M)) = \mathfrak{f}(E)$. Hence M is a matching.

To prove an upper bound of k on the size of the maximum matching, \mathcal{H} sends a (sorted) set $V_S \subseteq V$, where $\frac{1}{2}(|V_S| - \operatorname{occ}(G - V_S) + |V|) = k$. Both $|V_S|$ and |V| are computed directly; for $\operatorname{occ}(G - V_S)$, \mathcal{H} sends a sequence of (sub)graphs $C_i = (V_i, E_i) \subseteq V \times E$ claimed to be a partition of $G - V_S$ into connected components. \mathcal{V} can easily compute c, the number of C_i 's with an odd number of nodes. To ensure that the C_i 's are indeed the connected components of $G - V_S$, it suffices to show that (a) each C_i is connected in $G - V_S$; (b) $V \setminus V_S$ is the disjoint union of the V_i 's; and (c) there is no edge $(v, w) \in E$ s.t. $v \in V_i, w \in V_j, i \neq j$.

To prove Property (a), \mathcal{H} presents the (sub)graph C_i as $V_i \subset V$ (in sorted order) where each v is paired with its degree deg(v); followed by $E_i \subset E$ (in arbitrary order). Fingerprints are used to ensure that the multiset of nodes present in E_i matches the claimed degrees of nodes in V_i . If these fingerprints agree, then (whp) $E_i \subseteq V_i \times V_i$. Then \mathcal{H} uses the connectivity protocol from [6, Theorem 5.6] on the (sub)graph $C_i = (V_i, E_i)$ to prove that C_i is connected. Each of these checks on C_i has hcost $O(|E_i|)$. Note that \mathcal{V} requires only a constant number of fingerprints for these

checks, and can use the same working memory for each different C_i to check that $E_i \subseteq V_i \times V_i$ and that C_i is connected. The total vcost over all C_i is a constant number of fingerprints; the total hcost is O(m).

Property (b) is checked by testing $f((\cup_i V_i) \cup V_S) = f(V)$, where the unions in the LHS count multiplicities; if the fingerprints match then whp $V \setminus V_S$ is the *disjoint* union of the V_i 's. For (c), it suffices to ensure that each each edge in $E \setminus (\bigcup_i E_i)$ is incident to at least one node in V_S , as we have already checked that no edges in $\bigcup_i E_i$ cross between V_i and V_j for $i \neq j$. To this end, we use the "Consistent Labels" protocol of Lemma 2.3, with l(u) = 1 indicating $u \in V_S$ and l(u) = 0 indicating $u \notin V_S$, to force \mathcal{H} to replay all of E with each edge (u, v) annotated with l(u) and l(v). This allows \mathcal{V} to identify the set E_S of edges incident to at least one node in V_S . \mathcal{V} checks that $f((\cup_i E_i) \cup E_S) = f(E)$, which ensures (whp) that Property (c) holds and that over the entire partition of G no edges are added or omitted. Finally, provided all the prior fingerprint tests pass, the protocol accepts if c, the number of C_i 's with an odd number of nodes, satisfies $\frac{1}{2}(|S| - c + |V|) = k$.

3.3 Linear Programming and TUM Integer Programs

We present protocols to solve linear programming problems in our model leveraging the theory of LP duality. This leads to non-trivial schemes for a variety of graph problems.

Definition 3.3. Given a data stream \mathcal{A} containing entries of vectors $\mathbf{b} \in \mathbb{R}^{b}$, $\mathbf{c} \in \mathbb{R}^{c}$, and non-zero entries of a $b \times c$ matrix A in some arbitrary order, possibly interleaved. Each item in the stream indicates the index of the object it pertains to. The LP streaming problem on \mathcal{A} is to determine the value of the linear program min{ $\mathbf{c}^{T}\mathbf{x} \mid A\mathbf{x} \leq \mathbf{b}$ }.

We present our protocol as if each entry of each object appears at most once (if an entry does not appear, it is assumed to be zero). When this is not the case, the final value for that entry is interpreted as the *sum* of all corresponding values in the stream.

Theorem 3.4. There is a valid (|A|, 1) protocol for the LP streaming problem, where |A| is the number of non-zero entries in the constraint matrix A of A.

Proof. The protocol shows an upper bound by providing a primal-feasible solution \mathbf{x} , and a lower bound by providing a dual-feasible solution \mathbf{y} . When the value of both solutions match, \mathcal{V} is convinced that the optimal value has been found.

From the stream, \mathcal{V} fingerprints the sets $S_A = \{(i, j, A_{i,j})\}$, $S_B = \{(i, \mathbf{b}_i)\}$ and $S_C = \{(i, \mathbf{c}_j)\}$. Then \mathcal{H} provides all pairs of values $\mathbf{c}_j, \mathbf{x}_j, 1 \leq j \leq c$, with each \mathbf{x}_j additionally annotated with $|A_{\cdot j}|$, the number of non-zero entries in column j of A. This allows \mathcal{V} to fingerprint the multiset $S_X = \{(j, \mathbf{x}_j) : |A_{\cdot j}|\}$ and calculate the solution cost $\sum_{j=1}^{b} \mathbf{c}_j \mathbf{x}_j$.

To prove feasibility, for each row *i* of *A*, A_{i} , \mathcal{H} sends \mathbf{b}_i , then (the non-zero entries of) A_i . so that A_{ij} is annotated with \mathbf{x}_j . This allows the *i*th constraint to be checked easily in constant space. \mathcal{V} fingerprints the values given by \mathcal{H} for A, \mathbf{b} , and \mathbf{c} , and compares them to those for the stream. A single fingerprint of the multiset of values presented for \mathbf{x} over all rows is compared to $\mathfrak{f}(S_X)$. The protocol accepts \mathbf{x} as feasible if all constraints are met and all fingerprint tests pass.

Correctness follows by observing that the agreement with $\mathfrak{f}(A)$ guarantees (whp) that each entry of A is presented correctly and no value is omitted. Since \mathcal{H} presents each entry of **b** and **c** once, in index order, the fingerprints $\mathfrak{f}(S_B)$ and $\mathfrak{f}(S_C)$ ensure that these values are presented correctly. The claimed $|A_{j}|$ values must be correct: if not, then the fingerprints of either S_X or S_A will not match the multisets provided by \mathcal{H} . $\mathfrak{f}(S_X)$ also ensures that each time \mathbf{x}_j is presented, the same value is given (similar to Lemma 2.3).

To prove that \mathbf{x} is primal-optimal, it suffices to show a feasible solution \mathbf{y} to the dual A^T so that $c^T \mathbf{x} = b^T \mathbf{y}$. Essentially we repeat the above protocol on the dual, and check that the claimed values are again consistent with the fingerprints of S_A, S_B, S_C .

For any graph problem that can be formulated as a linear program in which each entry of A, **b**, and **c** can be derived as a linear function of the nodes and edges, we may view each edge in a graph stream \mathcal{A} as providing an update to values of one or more entries of A, **b**, and **c**. Therefore, we immediately obtain a protocol for problems of this form via Theorem 3.4. More generally, we obtain protocols for problems formulated as *totally unimodular integer programs* (TUM IPs), since optimality of a feasible solution is shown by a matching feasible solution of the dual of its LP relaxation [15].

Corollary 3.5. There is a valid (|A|, 1) protocol for any graph problem that can be formulated as a linear program or TUM IP in which each entry of A, b, and c is a linear function of the nodes and edges of graph.

This follows immediately from Theorem 3.4 and the subsequent discussion: note that the linearity of the fingerprinting builds fingerprints of S_A , S_B and S_C , so \mathcal{H} presents only their (aggregated) values, not information from the unaggregated graph stream.

Corollary 3.6. Shortest s - t path, max-flow, min-cut, and minimum weight bipartite perfect matching (MWBPM) all have valid (m, 1) protocols. For all four problems, a lower bound of hcost \cdot vcost = $\Omega(n^2)$ bits holds for dense graphs.

Proof. The upper bound follows from the previous corollary because all the problems listed possess formulations as TUM IPs and moreover the constraint matrix in each case has O(m + n) non-zero entries. For example, for max-flow, **x** gives the flow on each edge, and the weight of each edge in the stream contributes (linearly) to constraints on the capacity of that edge, and the flow through incident nodes.

The lower bound for MWBPM, max-flow, and min-cut holds from [6, Theorem 11] which argues hcost \cdot vcost = $\Omega(n^2)$ bits for bipartite perfect matching, and straightforward reductions of bipartite perfect matching to all three problems, see e.g. [13, Theorem 7.37]. The lower bound for shortest s - t path follows from a straightforward reduction from INDEX, for which a lower bound linear in hcost \cdot vcost was proven in [6, Theorem 3.1]. Given an instance (x, k) of INDEX where $x \in \{0, 1\}^{n^2}, k \in [n^2]$, we construct graph G, with $V_G = [n+2]$, and $E_G = E_A \cup E_B$. Alice creates $E_A = \{(i, j) : x_{f(i,j)=1}\}$ from x alone, where f is a 1-1 correspondence $[n] \times [n] \to [n^2]$. Bob creates $E_B = \{(n+1, i), (j, n+2)\}$ using f(i, j) = k. The shortest path between nodes n + 1 and n + 2is 3 if $x_k = 1$ and is 4 or more otherwise. This also implies that any approximation within $\sqrt{4/3}$ requires hcost \cdot vcost = $\Omega(n^2)$ (better inapproximability constants may be possible).

Conceptually, the above protocols for solving the LP streaming problem are straightforward: \mathcal{H} provides a primal solution, potentially repeating it once for each row of A to prove feasibility, and repeats the protocol for the dual. There are efficient protocols for the problems listed in the corollary since the constraint matrices of their IP formulations are sparse. For dense constraint matrices, however, the bottleneck is proving feasibility. We observe that computing $A\mathbf{x}$ reduces to

computing b inner-product computations of vectors of dimension c. There are $(c^{\alpha}, c^{1-\alpha})$ protocols to verify such inner-products [6]. But we can further improve on this since one of the vectors is held constant in each of the tests. This reduces the space needed by \mathcal{V} to run these checks in parallel; moreover, we prove a lower bound of hcost \cdot vcost = $\Omega(\min(c, b)^2)$ bits, and so obtain an *optimal* tradeoff for square matrices, up to logarithmic factors.

Theorem 3.7. Given a $b \times c$ matrix A and a c dimensional vector \mathbf{x} , the product $A\mathbf{x}$ can be verified with a valid $(bc^{\alpha}, c^{1-\alpha})$ protocol. Moreover, any such protocol requires $hcost \cdot vcost = \Omega(min(c, b)^2)$ bits for dense matrices.

Proof. We begin with the upper bound. The protocol for verifying inner-products which follows from [6] treats a c dimensional vector as an $h \times v$ array F, where $hv \geq c$. This then defines degree c polynomials f over a suitably large field, so that for each polynomial $f(x, y) = F_{x,y}$. For an innerproduct between two vectors, we wish to compute $\sum_{x \in [h], y \in [v]} F_{x,y} G_{x,y} = \sum_{x \in [h], y \in [v]} f(x, y)g(x, y)$ for the corresponding arrays F, G and polynomials f, g. These polynomials can then be evaluated at locations outside $[h] \times [v]$, so in the protocol \mathcal{V} picks a random position r, and evaluates f(r, y)and g(r, y) for $1 \leq y \leq v$. \mathcal{H} then presents a degree h polynomial s(x) which is claimed to be $\sum_{y=1}^{v} f(x, y)g(x, y)$. \mathcal{V} checks that $s(r) = \sum_{y=1}^{v} f(r, y)g(r, y)$, and if so accepts $\sum_{x=1}^{h} s(x)$ as the correct answer.

In [6] it is shown how \mathcal{V} can compute f(r, y) efficiently as F is defined incrementally in the stream: each addition of w to a particular index is mapped to $(x, y) \in [h] \times [v]$, which causes $f(r, y) \leftarrow f(r, y) + wp(r, x, y)$, where p(r, x, y) depends only on x, y, and r. Equivalently, the final value of f(r, y) over updates in the stream where the *j*th update is $t_j = (w_j, x_j, y_j)$ is $f(r, y) = \sum_{t_j: y_j = y} w_j p(r, x_j, y)$.

To run this protocol over multiple vectors in parallel naively would require keeping the f(r, y) values implied by each different vector separately, which would be costly. Our observation is that rather than keep these values explicitly, it is sufficient to keep only a fingerprint of these values, and use the linearity of fingerprint functions to finally test whether the polynomials provided by \mathcal{H} for each vector together agree with the stored values.

In our setting, the $b \times c$ matrix A implies b polynomials of degree c. We evaluate each polynomial at (r, y) for $1 \leq y \leq v$ for the same value of r: since each test is fooled by \mathcal{H} with small probability, the chance that none of them is fooled can be kept high by choosing the field to evaluate the polynomials over to have size polynomial in b + c. Thus, conceptually, the parallel invocation of binstances of this protocol require us to store $f_i(r, y)$ for $1 \leq y \leq v$ and $1 \leq i \leq b$ (for the b rows of A), as well as $f_x(r, y)$ for $1 \leq y \leq v$ (where f_x is the polynomial derived from \mathbf{x}). Rather than store this set of bv values explicitly, \mathcal{V} instead stores only v fingerprints, one for each value of y, where each fingerprint captures the set of b values of $f_i(r, y)$.

From the definition of our fingerprints, this means over stream updates $t_j = (w_j, i_j, x_j, y_j)$ of weight w_j to row i_j and column indexed by x_j and y_j we compute a fingerprint

$$\mathfrak{f}(A,y) = \sum_{i=1}^{b} f_i(r,y)\alpha^i = \sum_{i=1}^{b} \sum_{t_j: y_j = y, i_j = i} w_j p(r_j, x_j, y)\alpha^i$$

for each $y, 1 \le y \le v$. Observe that for each y this can be computed incrementally in the stream by storing only r and the current value of $\mathfrak{f}(A, y)$.

To verify the correctness, \mathcal{V} receives the *b* polynomials s_i , and builds a fingerprint of the multiset of $S = \{i : s_i(r)\}$ incrementally. \mathcal{V} then tests whether

$$\sum_{y=1}^{v} \mathfrak{f}(A, y) f_x(r, y) = \mathfrak{f}(S)$$

To see the correctness of this, we expand the lhs, as

$$\sum_{y=1}^{v} \mathfrak{f}(A,y) f_x(r,y) = \sum_{y=1}^{v} \left(\sum_{i=1}^{b} f_i(r,y) \alpha^i \right) f_x(r,y)$$
$$= \sum_{y=1}^{v} \left(\sum_{i=1}^{b} f_x(r,y) f_i(r,y) \alpha^i \right)$$
$$= \sum_{i=1}^{b} \left(\sum_{y=1}^{v} f_x(r,y) f_i(r,y) \right) \alpha^i$$

Likewise, if all s_i 's are as claimed, then

$$\mathfrak{f}(S) = \sum_{i=1}^{b} s_i(r) \alpha^i = \sum_{i=1}^{b} (\sum_{y=1}^{v} f_x(r,y) f_i(r,y)) \alpha^i$$

Thus, if the s_i 's are as claimed, then these two fingerprints should match. Moreover, by the Schwartz-Zippel lemma, and the fact that α and r are picked randomly by \mathcal{V} and not known to \mathcal{H} , the fingerprints will not match with high probability if the s_i 's are *not* as claimed, when the polynomials are evaluated over a field of size polynomial in (b + c).

To analyze the vcost, we observe that \mathcal{V} can compute all fingerprints in O(v) space. As \mathcal{H} provides each polynomial $s_i(x)$ in turn, \mathcal{V} can incrementally compute $\mathfrak{f}(S)$ and check that this matches $\sum_{y=1}^{v} \mathfrak{f}(A, y) f_x(r, y)$. At the same time, \mathcal{V} also computes $\sum_{i=1}^{b} \sum_{x=1}^{h} s_i(x)$, as the value of $A\mathbf{x}$. Note that if each s_i is sent one after another, \mathcal{V} can forget each previous s_i after the required fingerprints and evaluations have been made; and if h is larger than v, does not even need to keep s_i in memory, but can instead evaluate it term by term in parallel for each value of x. Thus the total space needed by \mathcal{V} is dominated by the v fingerprints and check values.

Setting $h = c^{\alpha}$ and $v = c^{1-\alpha}$, the total size of the information sent by \mathcal{H} is dominated by the *b* polynomials of degree $h = c^{\alpha}$.

To prove the lower bound, we give a simple reduction of INDEX to matrix-vector multiplication. Suppose we have an instance (x, k) of INDEX where $x \in \{0, 1\}^{n^2}$, $k \in [n^2]$. Alice constructs an $n \times n$ matrix A from x alone, in which $A_{i,j} = 1$ if $x_{f(i,j)=1}$, where f is a 1-1 correspondence $[n] \times [n] \to [n^2]$, and $A_{i,j} = 0$ otherwise. Assume f(i, j) = k. Bob then constructs a vector $\mathbf{x} \in \mathbb{R}^n$ such that $\mathbf{x}_i = 1$ and all other entries of \mathbf{x} are 0. Then the j'th entry of $A\mathbf{x}$ is 1 if and only if $x_{f(i,j)=1}$, and therefore the value of $x_{f(i,j)}$ can be extracted from the vector $A\mathbf{x}$. Therefore, if we had an (h, v) protocol for verifying matrix-vector multiplication given an $n \times n$ matrix A (even for a stream in which all entries of A come before all entries of \mathbf{x}), we would obtain a (\sqrt{h}, \sqrt{v}) protocol for INDEX. The lower bound for matrix-vector multiplication thus holds by a lower bound for INDEX given in [6, Theorem 3.1].

Corollary 3.8. For $c \ge b$ there is a valid $(c^{1+\alpha}, c^{1-\alpha})$ protocol for the LP streaming problem.

Proof. This follows by using the protocol of Theorem 3.7 to verify $A\mathbf{x} \leq \mathbf{b}$ and $A^T\mathbf{y} \geq \mathbf{c}$ within the protocol of Theorem 3.4. The cost is $(bc^{\alpha} + cb^{\alpha}, c^{1-\alpha} + b^{1-\alpha})$, so if $c \geq b$, this is dominated by $(c^{1+\alpha}, c^{1-\alpha})$ (symmetrically, if b > c, the cost is $(b^{1+\alpha}, b^{1-\alpha})$).

Our protocol for linear programming relied on only two properties: strong duality, and the ability to compute the value of a solution \mathbf{x} and check feasibility via matrix-vector multiplication. Such properties also hold for more general convex optimization problems, such as quadratic programming and a large class of second-order cone programs. Thus, similar results apply for these mathematical programs, motivated by applications in which weak peripheral devices or sensors perform error correction on signals. We defer full details from this presentation.

Theorem 3.7 also implies the existence of protocols for graph problems where *both* hcost and vcost are sublinear in the size of the input (for dense graphs). These include:

- An $(n^{1+\alpha}, n^{1-\alpha})$ protocol for verifying that λ is an eigenvalue of the adjacency matrix A or the Laplacian L of G: \mathcal{H} provides the corresponding eigenvector x, and \mathcal{V} can use the protocol of Theorem 3.7 to verify that $Ax = \lambda x$ or $Lx = \lambda x$.
- An $(n^{1+\alpha}, n^{1-\alpha})$ protocol for the problem of determining the effective resistance between designated nodes s and t in G where the edge weights are resistances. The problem reduces to solving an $n \times n$ system of linear equations [16].

4 Simulating Non-Streaming Algorithms

Next, we give protocols by appealing to known non-streaming algorithms for graph problems. At a high level, we can imagine the helper running an algorithm on the graph, and presenting a "transcript" of operations carried out by the algorithm as the proof to \mathcal{V} that the final result is correct. Equivalently, we can imagine that \mathcal{V} runs the algorithm, but since the data structures are large, they are stored by \mathcal{H} , who provides the contents of memory needed for each step. There may be many choices of the algorithm to simulate and the implementation details of the algorithm: our aim is to choose ones that result in smaller annotations.

To make this concrete, consider the case of requiring the graph to be presented in a particular order, such as depth first order. Starting from a given node, the exploration retrieves nodes in order, based on the pattern of edges. Assuming an adjacency list representation, a natural implementation of the search in the traditional model of computation maintains a stack of edges (representing the current path being explored). Edges incident on the current node being explored are pushed, and pops occur whenever all nodes connected to the current node have already been visited. \mathcal{H} can allow \mathcal{V} to recreate this exploration by providing at each step the next node to push, or the new head of the stack when a pop occurs, and so on. To ensure the correctness of the protocol, additional checking information can be provided, such as pointers to the location in the stack when a node is visited that has already been encountered.

We provide protocols for breadth first search and depth first search in Appendix A, based on this idea of "augmenting a transcript" of a traditional algorithm. However, while the resulting protocols are lightweight, it rapidly becomes tedious to provide appropriate protocols for other computations based on this idea. Instead, we introduce a more general approach which argues that any (deterministic) algorithm to solve a given problem can be converted into a protocol in our model. The running time of the algorithm in the RAM model becomes the size of the proof in our setting.

Our main technical tool is the off-line memory checker of Blum et al. [11], which we use to efficiently verify a sequence of accesses to a large memory. Consider a *memory transcript* of a sequence of read and write operations to this memory (initialized to all zeros). Such a transcript is *valid* if each read of address *i* returns the last value written to that address. The protocol of Blum et al. requires each read to be accompanied by the timestamp of the last write to that address; and to treat each operation (read or write) as a read of the old value followed by the write of a new value. Then to ensure validity of the transcript, it suffices to check that a fingerprint of all write operations (augmented with timestamps) matches a fingerprint of all read operations (using the provided timestamps), along with some simple local checks on timestamps. Consequently, any valid (timestamp-augmented) transcript is accepted by \mathcal{V} , while any invalid transcript is rejected by \mathcal{V} with high probability.

We use this memory checker to obtain the following general simulation result.

Theorem 4.1. Suppose P is a graph problem possessing a non-randomized algorithm \mathcal{M} in the random-access memory model that, when given G = (V, E) in adjacency list or adjacency matrix form, outputs P(G) in time t(m, n), where m = |E| and n = |V|. Then there is an (m + t(m, n), 1) protocol for P.

Proof sketch. \mathcal{H} first repeats (the non-zero locations of) a valid adjacency list or matrix representation G, as writes to the memory (which is checked by \mathcal{V}); \mathcal{V} uses fingerprints to ensure the edges included in the representation precisely correspond to those that appeared in the stream, and can use local checks to ensure the representation is otherwise valid. This requires O(m) annotation and effectively initializes memory for the subsequent simulation. Thereafter, \mathcal{H} provides a valid augmented transcript T' of the read and write operations performed by algorithm \mathcal{M} ; \mathcal{V} rejects if T' is invalid, or if any read or write operation executed in T' does not agree with the prescribed action of \mathcal{M} . As only one read or write operation is performed by \mathcal{M} in each timestep, the length of T' is O(t(m, n)), resulting in an (m + t(m, n), 1) protocol for P.

Although Theorem 4.1 only allows the simulation of deterministic algorithms, \mathcal{H} can nondeterministically "guess" an optimal solution S and prove optimality by invoking Theorem 4.1 on a (deterministic) algorithm that merely checks whether S is optimal. Unsurprisingly, it is often the case that the best-known algorithms for verifying optimality are more efficient than those finding a solution from scratch (see e.g. the MST protocol below); therein lies much of the power of the simulation theorem.

Theorem 4.2. There is a valid (m, 1) protocol to find a minimum cost spanning tree; a valid $(m + n \log n, 1)$ protocol to verify single-source shortest paths; and a valid $(n^3, 1)$ protocol to verify all-pairs shortest paths.

Proof. We first prove the bound for MST. Given a spanning tree T, there exists a linear-time algorithm \mathcal{M} for verifying that T is minimum (see e.g. [17]). Let \mathcal{M}' be the linear-time algorithm that, given G and a subset of edges T in adjacency matrix form, first checks that T is a spanning tree by ensuring |T| = n - 1 and T is connected (by using e.g. breadth-first search), and then executes \mathcal{M} to ensure T is minimum. We obtain an (m, 1) protocol for MST by having \mathcal{H} provide a minimum spanning tree T and using Theorem 4.1 to simulate algorithm \mathcal{M}' .

The upper bound for single-source shortest path follows from Theorem 4.1 and the fact that there exist implementations of Djikstra's algorithm that run in time $m + n \log n$. The upper bound for all-pairs shortest paths also follows from Theorem 4.1 and the fact that the Floyd-Warshall algorithm runs in time $O(n^3)$.

We now provide near-matching lower bounds for all three problems.

Theorem 4.3. Any protocol for verifying single-source or all pairs shortest paths requires $hcost \cdot vcost = \Omega(n^2)$ bits. Additionally, if edge weights may be specified incrementally, then an identical lower bound holds for MST.

Proof. The lower bounds for single-source and all-pairs shortest paths are inherited from shortest s - t path (Corollary 3.6).

To prove the lower bound for MST, we present a straightforward reduction from an instance of INDEX, (x, k), where $x \in \{0, 1\}^{n^2}$, $k \in [n^2]$. Alice will construct a graph G, with $V_G = [n]$, and $E_G = E_A$. Bob will then construct two graphs, G_1 and G_2 , with $E_{G_1} = E_A \cup E_{B_1}$ and $E_{G_2} = E_A \cup E_{B_2}$. If edge (i, j) is in $E_A \cap E_{B_1}$, then we interpret this to mean that the weight of edge (i, j) in E_{G_1} is the sum of its weights in E_A and E_{B_1} . Below, we will write (i, j, w) to denote an edge between nodes i and j with weight w.

Alice creates $E_A = \{(i, j, 1) : x_{f(i,j)=1}\}$ from x alone, where f is a bijection $[n] \times [n] \to [n^2]$. Bob creates $E_{B_1} = \{(u, v, 3) : f(u, v) \neq k\}$, and $E_{B_2} = E_{B_1} \cup \{(i, j, 1)\}$, where (i, j) is the edge satisfying f(i, j) = k. Edge (i, j), if it exists, is the lowest-weight edge in E_{G_1} , and hence (i, j) is in any min-cost spanning tree of G_1 if and only if $x_k = 1$. In contrast, (i, j) is always in the min-cost spanning tree of G_2 . Therefore, if $x_k = 1$, then the minimum spanning tree of G_2 will be of higher cost than that of G_1 , because the weight of (i, j) is 1 in E_{G_1} and 2 in E_{G_2} . And if $x_k = 0$, then the minimum spanning tree of G_2 will be of lower cost than that of G_1 , because the weight of edge (i, j) will be ∞ in G_1 and 1 in G_2 . Thus, by comparing the cost of the MSTs of G_1 and G_2 , Bob can extract the value of x_k . The lower bound now follows from the hardness of INDEX [6, Theorem 3.1].

Diameter. The diameter of G can be verified via the all-pairs shortest path protocol above, but the next protocol improves over the memory checking approach.

Theorem 4.4. There is a valid $(n^2 \log n, 1)$ protocol for computing graph diameter. Further, any protocol for diameter requires hcost \cdot vcost = $\Omega(n^2)$ bits.

Proof. [6, Theorem 5.2] gives an $(n^2 \log l, 1)$ protocol for verifying that $A^l = B$ for a matrix A presented in a data stream and for any positive integer l. Note that if A is the adjacency matrix of G; then $(I + A)_{ij}^l \neq 0$ if and only if there is a path of length at most l from i to j. Therefore, the diameter of G is equal to the unique l > 0 such that $(I + A)_{ij}^l \neq 0$ for all (i, j), while $(I + A)_{ij}^{l-1} = 0$ for some (i, j). Our protocol requires \mathcal{H} to send l to \mathcal{V} , and then run the protocol of [6, Theorem 5.2] twice to verify that l is as claimed. Since the diameter is at most n-1, this gives an $(n^2 \log n, 1)$ protocol.

We prove the lower bound via a reduction from an instance of INDEX, (x, k), where $x \in \{0, 1\}^{n^2/4}$, $k \in [n^2/4]$. Alice creates a bipartite graph G = (V, E) from x alone: she includes edge (i, j) in E if and only if $x_{f(i,j)} = 1$, where f is a bijection from edges to indices. Bob then adds to G two nodes L and R, with edges from L to each node in the left partite set, edges from R to each node in the

right partite set, and an edge between L and R. This ensures that the graph is connected, with diameter at most 3. Finally, Bob appends a path of length 2 to node i, and a path of length 2 to node j, where f(i, j) = k. If $x_k = 0$, then the diameter is now 7, while if $x_k = 1$, the diameter is 5. The lower bound follows from the hardness of INDEX [6, Theorem 3.1] (this also shows that any protocol to approximate diameter better than $\sqrt{1.4}$ requires hcost \cdot vcost $= \Omega(n^2)$ bits; no effort has been made to optimize the inapproximability constant).

5 Conclusion and Future Directions

In this paper, we showed that a host of graph problems possess streaming protocols requiring only constant space and linear-sized annotations. For many applications of the annotation model, the priority is to minimize vcost, and these protocols achieve this goal. However, these results are qualitatively different from those involving numerical streams in the earlier work [6]: for the canonical problems of heavy hitters, frequency moments, and selection, it is trivial to achieve an (m, 1) protocol by having \mathcal{H} replay the stream in sorted ("best") order. The contribution of [6] is in presenting protocols obtaining optimal tradeoffs between hcost and vcost in which both quantities are sublinear in the size of the input. There are good reasons to seek these tradeoffs. For example, consider a verifier with access to a few MBs or GBs of working memory. If an (m, 1) protocol requires only a few KBs of space, it would be desirable to use more of the available memory to significantly reduce the running time of the verification protocol.

In contrast to [6], it is non-trivial to obtain (m, 1) protocols for the graph problems we consider, and we obtain tradeoffs involving sublinear values of hcost and vcost for some problems with an algebraic flavor (e.g. matrix-vector multiplication, computing effective resistances, and eigenvalues of the Laplacian). We thus leave as an open question whether it is possible to obtain such tradeoffs for a wider class of graph problems, and in particular if the use of memory checking can be adapted to provide tradeoffs.

A final open problem is to ensure that the work of \mathcal{H} is scalable. In motivating settings such as Cloud computing environments, the data is very large, and \mathcal{H} may represent a *distributed* cluster of machines. It is a challenge to show that these protocols can be executed in a model such as the MapReduce framework.

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A Direct Protocols for Breadth First and Depth First Search

In this appendix, we consider the problem of "verifying" a breadth first or depth first search. Our goal is to force \mathcal{H} to provide to \mathcal{V} the edges of G in the order they would be visited by a non-streaming BFS or DFS algorithm, despite limiting \mathcal{V} to logarithmic space. We can accomplish this with linear annotation simply by invoking Theorem 4.1 on a BFS or DFS algorithm; however we now give "direct" protocols for this problem. Our protocols improve over Theorem 4.1 by constant factors, because we avoid having to replay the entire contents of memory at the beginning and end of the protocol. As an immediate corollary of our BFS protocol, we obtain an (m, 1) protocol for bipartiteness.

To make precise the notion of "verifying a BFS", we define the concept of a BFS transcript, using the concept of a label-augmented list of edges defined in Definition 2.2.

Definition A.1. Given a connected undirected graph G, a BFS transcript T rooted at s is a labelaugmented list of edges E', with each label l(e, u) claimed to be the distance from s to u. Let $d_e := \min(l(e, u), l(e, v))$. A BFS transcript is valid if: (a) Edges are presented in increasing order of d_e ; (b) E' is a valid list of label-augmented edges; and (c) For all u, l(u) is the (hop) distance from s to u.

For clarity, we make explicit the labels of each edge; a more concise protocol would group edges in order of l(e, u) + l(e, v). However, this does not alter the asymptotic cost.

It is easy to see that any valid BFS transcript corresponds to the order edges are visited in an actual BFS of G. We therefore say a protocol verifies a BFS if it accepts any valid BFS transcript (possibly with additional annotation interleaved) and rejects any invalid BFS transcript with high probability. With this in mind, we now define augmented BFS transcripts.

Any valid BFS transcript corresponds to a possible order of edges being visited in an actual BFS of G. So we seek a protocol that accepts any valid BFS transcript (possibly with additional annotation interleaved), and rejects an invalid transcript (whp). This leads us to define an augmented BFS transcript.

Let T be a BFS transcript. We partition the edges by their distance, so that $E_l = \{e : d_e = l\}$. Likewise, we define $V_l = \{u | \exists e = (u, v) \text{ s.t. } l(e, u) = l\}$. In a valid BFS transcript, the sets V_l partition V since l(e, u) gives the same distance from s to u every time is is listed; an invalid transcript may not have this property.

Definition A.2. An augmented BFS transcript T' is a BFS transcript T where additionally before each E_l , V_{l+1} is presented, along with degrees $\deg_l(v)$, defined as $\deg_l(v) = |\{u|(u,v) \in E, u \in V_l, v \in V_{l+1}\}|$.

The augmented BFS transcript is valid if the underlying BFS transcript is valid and all claimed degrees are truthful.

Theorem A.3. There is a valid (m, 1) protocol to accept a valid augmented BFS transcript.

Proof. To ensure Definition A.1 (a), \mathcal{V} rejects if the edges are not presented in increasing order of d_e , which is trivial to check since l(e, u) and l(e, v) are presented for every edge. \mathcal{V} ensures Definition A.1 (b) using the "Consistent Labels" protocol of Lemma 2.3.

To verify the augmented transcript, fingerprints ensure multiset $\{v \in V_{l+1} : \deg_l(v)\}$ matches the multiset of edges e = (u, v) with l(e, u) = l, l(e, v) = l + 1. \mathcal{V} (re)uses the same working memory to store these fingerprints for each level l in turn. To ensure Definition A.1 (c), \mathcal{V} uses the augmented annotation at each edge-level to reject if a node at level l+1 is not incident to any nodes at level l, or if an edge is ever presented with |l(e, u) - l(e, v)| > 1. An inductive argument ensures that l(e, u) is the distance from s to u for all u and all e incident to u (the base case is just l(s) = 0). For the vcost, note that any valid augmented BFS transcript has length O(n + m) = O(m).

From this theorem, we obtain an (m, 1) protocol for bipartiteness: G is bipartite if and only if there is no edge (u, v) with l(e, u) = l(e, v), which is easily checked given a valid augmented BFS transcript. Further, any online protocol for bipartiteness requires $hcost \cdot vcost = \Omega(n)$ bits, even when m = O(n); this lower bound follows by reducing bipartiteness to INDEX [18], which has the same lower bound [6, Theorem 3.1].

Our approach for Depth First Search is similar, but more involved. Analogously to BFS, we now define DFS transcripts.

Definition A.4. Given a connected undirected graph G, a DFS transcript consists of m + 2n rows, where each row is of the form edge(u, v), push(u), or pop(u). Each symbol push(u) (resp. pop(u)) represents a simulated push (pop) of node u on a (simulated) stack. Reading the transcript in order, the most recently pushed vertex that has not yet been popped is termed the "top" node. A DFS transcript is valid for G = (V, E) if

(a) Every $v \in V$ is pushed exactly once, immediately after it first appears in an edge;

(b) Every $e \in E$ is presented exactly once, and is incident to the top node;

(c) Every $v \in V$ is popped exactly once, when it is the top and all edges incident to it have already appeared in the transcript.

The order edges are visited in an actual DFS of G always corresponds to the ordering of edges in a valid DFS transcript, and each simulated push and pop operation corresponds to actual push and pop operations in the same DFS. Therefore, just as in the BFS case, we seek a protocol that accepts any valid DFS transcript (possibly with additional annotation on each event) and rejects invalid DFS transcripts whp. We write t(event, u) = t to denote that the t'th row of the transcript is event(u). Thus, the top node at time t is $top(t) = \arg \max_u(t(\text{push}, u)|t(\text{push}, u) \le t < t(\text{pop}, u))$, and we say the stack height at time t is $height(t) = |\{u|t(\text{push}, u) \le t < t(\text{pop}, u)\}|$.

Definition A.5. An augmented DFS transcript T' is a DFS transcript where in addition:

- (a) before the edges, triples (u, level(u), ntop(u)) are listed for each node in order;
- (b) every edge e = (u, v) is annotated with (t(push, u), t(pop, u), t(push, v), t(pop, v)).
- (c) each pop(u) with t(pop, u) = t is annotated with (v, t(push, v), t(pop, v)) where v = top(t+1)

Definition A.6. We say an augmented DFS transcript is valid if

(a) the underlying DFS transcript is valid;

(b) for all u, level(u) = height(t(push, u)) and ntop(u) is $|\{t|top(t) = u\}|$.

- (c) all t(push, u) and t(pop, u) annotations are consistent;
- (d) the triple after each pop event correctly identifies the new top.

Theorem A.7. There is a valid (m, 1) protocol to accept a valid augmented DFS transcript.

Proof. To ensure Definition A.6 (b), \mathcal{V} initially computes a fingerprint f_1 of the multiset $\{(u, level(u)) : ntop(u)\}$. Throughout the protocol, \mathcal{V} also tracks height(t), and maintains a fingerprint f_2 of all (top(t), height(t)) pairs observed while reading the augmented DFS transcript. If property (b) is satisfied, f_1 and f_2 will match, otherwise they will differ with high probability. This is essentially the "Consistent Labels" protocol applied to this setting.

For Definition A.6 (c), we use the "Consistent Labels" protocol of Lemma 2.3 to ensure that the claimed push times are consistent for all nodes. This protocol presents a list of nodes in sorted order accompanied by their t(push, u) values. This list is also used to check to see that exactly one push event is claimed for each node: a fingerprint of the list can then be compared to one generated directly from the transcript. The case for pop is identical.

For Definition A.6 (d), \mathcal{V} can reject if the triple of Definition A.5 (c) has $t(\mathsf{push}, v) > t$ or $t(\mathsf{pop}, v) < t$. Observe that the stack must have the same height every time v is claimed to be the top, else the fingerprints f_1 and f_2 will differ (whp). Thus we can assume that v is always reported as top at the same height, which is $h = height(t(\mathsf{push}, v))$. We argue that at any timestep t there can be only one node v at height h such that $t(\mathsf{push}, v) \le t < t(\mathsf{pop}, v)$. If not, then let u and v be a "witness" pair of nodes both claimed to be at height h such that $t(\mathsf{push}, v) < t(\mathsf{push}, u) < t(\mathsf{pop}, v)$: if there are many such nodes, then pick v with the smallest value of $t(\mathsf{push}, v)$, and u likewise with the smallest $t(\mathsf{push}, u)$ such that the condition is met. For this to happen, there must have been a **pop** event to bring the height below h; either this is to v itself, contradicting the assumptions, or to some other node, w. Assuming that $t(\mathsf{push}, w) < t(\mathsf{pop}, w)$ (else our other checks identify this error), then v and w form a witness pair with $t(\mathsf{push}, w) < t(\mathsf{pop}, w) < t(\mathsf{push}, u)$, contradicting the calimet the earliest witness pair.

Next, we ensure that the underlying DFS transcript is valid. Definition A.4 (a) is easy to check, given Definition A.6 (c), by having \mathcal{V} reject if the t'th row of the transcript is $\mathsf{edge}(u, v)$ and is annotated with $t(\mathsf{push}, v) > t + 1$. Definition A.4 (b) is also easy to check given Definition A.6 (d): \mathcal{V} tracks top(t) implicit from push operations or from annotations on pop events. This leaves Definition A.4 (c) i.e. to ensure that nodes are popped at the correct times. To accomplish this, it suffices to have \mathcal{V} reject if the t'th row of the transcript is $\mathsf{edge}(u, v)$ and is annotated with $t(\mathsf{pop}, u) < t$ or if the t'th row is $\mathsf{pop}(u)$ and $top(t) \neq u$. These rules ensure $\mathsf{pop}(v)$ cannot occur "too early", i.e. before all edges incident to v have been presented: when such an edge is presented in row t, \mathcal{V} will reject due to $t(\mathsf{pop}, v) < t$. Likewise, v cannot be popped "too late", since if top(t) = v and there are no more edges incident on v, the next edge will not be incident on top(t) (required by Definition A.4 (b)) unless v is popped first.