

PDTL: Parallel and Distributed Triangle Listing for Massive Graphs

Ilias Giechaskiel*[†], George Panagopoulos*, and Eiko Yoneki*

*University of Cambridge, Cambridge, UK

[†]University of Oxford, Oxford, UK

Email: Ilias.Giechaskiel@{cs.ox.ac.uk, cl.cam.ac.uk}, gmp37@cam.ac.uk, Eiko.Yoneki@cl.cam.ac.uk

Abstract—This paper presents the first distributed triangle listing algorithm with provable CPU, I/O, Memory, and Network bounds. Finding all triangles (3-cliques) in a graph has numerous applications for density and connectivity metrics, but the majority of existing algorithms for massive graphs are sequential, while distributed versions of algorithms do not guarantee their CPU, I/O, Memory, or Network requirements. Our Parallel and Distributed Triangle Listing (PDTL) framework focuses on efficient external-memory access in distributed environments instead of fitting subgraphs into memory. It works by performing efficient orientation and load-balancing steps, and replicating graphs across machines by using an extended version of Hu et al.’s Massive Graph Triangulation algorithm. PDTL suits a variety of computational environments, from single-core machines to high-end clusters, and computes the exact triangle count on graphs of over 6B edges and 1B vertices (e.g. Yahoo graphs), outperforming and using fewer resources than the state-of-the-art systems PowerGraph, OPT, and PATRIC by $2\times$ to $4\times$. Our approach thus highlights the importance of I/O in a distributed environment.

Keywords—Triangle Listing, Triangle Counting, Big Data, Massive Graphs, I/O-Efficient Algorithm, Distributed Algorithm, Parallel Algorithm

I. INTRODUCTION

Graphs are important abstractions to model a range real-world situations, but they are becoming increasingly massive and will soon reach billions of vertices and trillions of edges, making in-memory algorithms insufficient for computing graph properties. One such property that has gained the attention of the graph processing community is the *number of triangles* in the graph, which can be seen as a special case of counting cycles of given length, or finding complete subgraphs. Finding all triangles in a graph is crucial for metrics such as the *clustering coefficient* [25] and the similar *transitivity ratio* [19], which can be used to find high-density nodes, and to detect fake accounts in social networks [26], as well as web spam and content quality [5]. Triangle enumeration is also necessary for *dense neighborhood discovery* [24], *triangular connectivity* [4], and finding the *k-trusses* of graphs [23]. However, research has been limited on either external memory considerations, or the creation of parallel frameworks (Section II). In this paper we show that it is possible to combine both approaches with substantial improvements in performance. Our Parallel and Distributed Triangle Listing (PDTL) framework (Section IV) extends Hu et al.’s Massive Graph Triangulation (MGT) algorithm

[14] in order to work in the distributed environment by duplicating the graph across each machine, and has provable bounds on CPU, I/O, Memory, and Network utilization. By further parallelizing the orientation step, and by intelligently distributing the load across each processor, our algorithm computes the exact triangle counts on graphs with billions of edges or vertices $2\times$ to $4\times$ faster than the state-of-the-art frameworks, using considerably fewer resources, and exhibiting scalability across multiple processors and machines (Section V). In summary, our contributions are as follows:

- We create a general framework for triangle listing and counting for both distributed and single-machine systems. Our algorithm is the first triangle listing algorithm that provides efficient and well-understood bounds on CPU, I/O, Memory, and Network utilization, across multiple environments (Theorem IV.3).
- We uncover hidden assumptions in the proofs and implementation of the closed-source MGT algorithm. We modify the algorithm to correspond to its implementation, and prove that our modifications do not alter its theoretical efficiency (Section IV-A).
- We introduce further optimizations in the orientation and distribution steps of our algorithm to reduce bottlenecks without adding complexity.
- We conduct extensive experiments that show our algorithm is highly scalable across multiple cores and machines, with low memory requirements, even for graphs with hundreds of millions of vertices, and multiple billion edges (Section V). Over the standard Twitter dataset [16], our algorithm is 4 times faster than PATRIC [3], 3 times faster than OPT [15], and 2 times faster than PowerGraph [11], the state-of-the-art frameworks in distributed and multicore triangle counting.

II. RELATED WORK

Dementiev [9] and Menegola [18] first introduced **external-memory** algorithms for triangle counting, but their algorithms had high I/O overheads. The first algorithms with reasonable performance for triangle listing were introduced by Chu and Cheng [8], and relied on *graph partitioning* to achieve an I/O complexity of $\mathcal{O}\left(\frac{|E|^2}{MB} + \frac{T}{B}\right)$, under certain assumptions on the structure of the graph. However, MGT

by Hu et al. [14] exhibits the same performance without any additional assumptions, and was superior in practice. Finally, Pagh and Silvestri recently proposed a new algorithm for triangle counting (but not listing) which has an I/O complexity of $\mathcal{O}\left(\frac{|E|^{1.5}}{\sqrt{MB}}\right)$, and improves the given bound [20].

The first **dedicated parallel** triangle counting framework, PATRIC [3], uses graph partitioning and message passing. It is not I/O-efficient, but proposes multiple load balancing mechanisms, which are calculated in parallel and do not pose a bottleneck. Even so, PATRIC requires that each partition fits in memory, and targets datacenters, with hundreds of processors and high dedicated RAM per processor. OPT [15] is a disk-based, single machine system that exploits I/O and multi-core CPU parallelism, and performs favorably compared to distributed triangle-counting frameworks.

In terms of **general-purpose frameworks**, there are multiple MapReduce algorithms for counting triangles, the best of which is CTTTP [21]. MapReduce algorithms produce too much intermediate networking data, and are considerably slow: CTTTP takes $2\times$ longer on the Twitter dataset [16] using 40 nodes compared a single-core MGT. PowerGraph [11] is a general-purpose vertex-oriented framework that is the fastest for triangle counting among existing alternatives, while PSgL [22] proposes novel methods for generic sub-graph listing, but is $6\times$ slower than PowerGraph.

Overall, we see that there is a divide between using external memory and parallelizing the algorithm, but as we show in Section V, by combining the two approaches, PDTL is $4\times$ faster than PATRIC, $3\times$ faster than OPT, and $2\times$ faster than PowerGraph, while providing theoretical guarantees, and not running out of memory for larger graphs.

III. PRELIMINARIES

A. Definitions

All graphs $G = (V, E)$ on $n = |V|$ vertices and $m = |E|$ edges are assumed to be undirected and simple. For every $u \in V$, $N_G(u) = \{v : (u, v) \in E\}$ denotes the set of its *neighbors* and $d_G(u) = |N_G(u)|$ its *degree*. Note that we may omit the qualifier G when doing so is clear. Finally, for simplicity, we also identify V with $[n] = \{0, \dots, n-1\}$.

Definition III.1 (Triangle). *Given an undirected graph $G = (V, E)$, a triangle is a set of three vertices $\{u, v, w\} \subseteq V$, such that all of (u, v) , (v, w) and (w, u) are edges in E .*

Finding the set K of all such triangles is *triangle listing*, and reporting on their number $T = |K|$ is *triangle counting*.

Definition III.2 (Degree-Based Order, Orientation). *Given an undirected graph $G = (V, E)$, the degree-based order \prec on V is defined as follows: $u \prec v$ if and only if $d(u) < d(v)$ or $d(u) = d(v)$ and $u < v$. We define the directed graph $G^* = (V, E^*)$, called G 's orientation, by $(u, v) \in E^*$ if and only if $(u, v) \in E$ and $u \prec v$.*

Because \prec is a strict total order, the orientation uniquely associates $\{u, v, w\}$ where $u \prec v \prec w$ with (u, v, w) :

Definition III.3 (Cone Vertex, Pivot Edge [14]). *Given a triangle (u, v, w) with $u \prec v \prec w$ in G^* , we call u its cone vertex, and (v, w) its pivot edge.*

The arboricity $\alpha(G)$ of a graph G is the minimum number of edge-disjoint forests needed to cover its edges and satisfies:

Theorem III.4 (Arboricity bounds [7]). *The arboricity of a graph $G = (V, E)$ satisfies:*

- 1) $\alpha \leq \lceil \sqrt{|E|} \rceil$
- 2) $\alpha = \mathcal{O}(1)$ if G is planar
- 3) $\sum_{(u,v) \in E} \min\{d(u), d(v)\} \leq \mathcal{O}(\alpha |E|)$

Note that the $T \leq \frac{1}{3} \sum_{(u,v) \in E} \min\{d(u), d(v)\}$, where T is the number of triangles, as any edge can appear in at most $\min\{d(u), d(v)\}$ triangles, so $T = \mathcal{O}(\alpha |E|)$. As a result, it is beneficial to have a runtime dependent on $\alpha(G)$, because it is at most $\lceil \sqrt{|E|} \rceil$, but can be $\mathcal{O}(1)$ for planar graphs.

Finally, we remind the reader of Aggarwal and Vitter's I/O complexity analysis methodology [2], which depends on the block size B : in accessing N elements in order, the disk performs $scan(N) = \Theta(N/B)$ I/Os, whereas random access can require $\Omega(N)$ I/Os in the worst case. Sorting takes $sort(N) = \Theta\left(\frac{N}{B} \log_{M/B} \frac{N}{B}\right)$ I/Os by external mergesort, where M is the memory size.

IV. PDTL

We assume a computational environment of N nodes, each of which has P processors, with M bytes of memory for each of the processors, so that for appropriate parameters, we can model a high-end data center, with multiple processors per machine, or even just a single computer with low available memory. In Section IV-A we explain the baseline single-core MGT algorithm and our modifications, while in Section IV-B we explain our parallel and distributed PDTL system and prove its theoretical properties.

A. Massive Graph Triangulation

Algorithm 1 presents the MGT algorithm [14] under the *small-degree assumption*: that every vertex $v \in V$ has $d_{G^*}(v) \leq cM/2$ for some implementation-specific constant $c < 1$.¹ The idea behind MGT is that given an oriented graph G^* , one can find all triangles by loading consecutive edges into memory and iterating over all vertices u and their out-edges to find all triangles with cone vertex u and pivot edge loaded into memory. By using hash structures on the loaded edges and $N(u)$ this can be done in a CPU- and I/O- efficient way. However, as we illustrate in Section IV-A1, the high-level algorithm does not correspond to its implementation,

¹We note that our code does not make such assumption, and refer the reader to [14] for a way to remove it.

so we modify MGT, and show in Section IV-A2 that our modifications do not alter the algorithm's efficiency.

Algorithm 1 MGT

Input: An oriented $G^* = (V, E^*)$
Output: All triangles in G
while there are edges in E^* to be read **do**
 Read the next cM edges into memory
 Create hash structures on the edges
 for $u \in V$ **do**
 Read $N(u)$ from disk
 Construct hash structures on $N(u)$
 Report triangles with cone u and pivot in memory
 Release the structures on $N(u)$

1) *Modifications:* Although only a binary for MGT is available at [12], during our experimentation we hypothesized that the implementation of MGT does not use explicit sets, but arrays. Indeed, if the adjacency list for any given vertex is not sorted, the given implementation misses triangles, though the manual [13] does not make mention of such requirements. Clearly, if any types of sets were constructed, this need for a sorted adjacency list would not be present. This belief was further verified by our own implementation, where using sets and maps of any kind made our implementation more than $10\times$ slower.

Consequently, our implementation deviates from Algorithm 1 by utilizing sorted arrays instead of sets. Specifically, it is assumed (for compatibility with the MGT implementation [12]) that the file format for the graph is such that if $v < w$ then $N_G(v)$ comes before $N_G(w)$ and if additionally $v, w \in N_G(u)$ then v comes before w in $N_G(u)$.

Following the notation in [14], let E_{mem} denote the set of edges in memory, V_{mem} its endpoints, V_{mem}^+ those $v \in V_{mem}$ that have *outgoing* edges in E_{mem} , and $N^+(u) = N(u) \cap V_{mem}^+$. In other words, $V_{mem}^+ = \{u \in V \mid \exists v \in V : (u, v) \in E_{mem}\}$, and for symmetry let $V_{mem}^- = \{u \in V \mid \exists v \in V : (v, u) \in E_{mem}\}$, with $V_{mem} = V_{mem}^+ \cup V_{mem}^-$. Additionally let $v_{low} = \min_{v \in V_{mem}^+} v$ and $v_{high} = \max_{v \in V_{mem}^+} v$.

Because the graph is sorted, we know that if $v < v_{low}$ or $v > v_{high}$, then $v \notin V_{mem}^+$. As a result, we can split E_{mem} into two arrays: *edg* which stores the sequence of out-neighbors and *ind* that stores the degree of v and offset into *edg* at location $v - v_{low}$. In other words, the out-edges of v (provided it is in memory) are stored at $E_v = \text{edg}[\text{ind}[v - v_{low}]]$.

Moreover, because $|N^+(u)| \leq |N(u)| \leq d_{max}^*$, and because we know d_{max}^* from the orientation step, $N(u)$ and $N^+(u)$ can also be represented by static arrays of size d_{max}^* , called *nm* and *nmp* respectively.

As a result, the modified MGT (Algorithm 2) works as follows on the sorted and oriented graph: it loads the next

$\Theta\left(\frac{|E|}{M}\right)$ edges into *edg* and *ind* as indicated above. Then, it iterates over the entire graph vertex by vertex, and for each vertex u , it does the following:

- 1) Stores $N(u)$ into the array *nm*
- 2) Evaluates $N^+(u)$ into the array *nmp* by iterating over $v \in N(u)$ and checking *ind* for out-neighbors
- 3) For each $v \in N^+(u)$, it reports triangle (u, v, w) with cone u and pivot w for each $w \in N(u) \cap E_v$

Algorithm 2 Modified MGT

Input: A sorted, oriented $G^* = (V, E^*)$
Output: All triangles in G
while there are edges in E^* to be read **do**
 Read the next $c'M$ out-neighbors into *edg* and store in *ind* the degrees and offsets
 for $u \in V$ **do**
 Read $N(u)$ from disk to array *nm*
 Write $N^+(u)$ to *nmp* using *nm* and *ind*
 for $v \in \text{nmp}$ **do**
 for $w \in \text{nm} \cap \text{edg}[\text{ind}[v - v_{low}]]$ **do**
 Report (u, v, w)
 Clear *nm* and *nmp*

2) *Analysis:* First of all, because $\Theta(M)$ edges are loaded at each step there are $h = \Theta(|E|/M)$ iterations, and each iteration performs $\frac{|E|}{B}$ I/Os to read over the graph. Additionally, the cost of outputting T triangles is $\frac{T}{B}$, for a total I/O complexity of $\mathcal{O}\left(\frac{|E|^2}{MB} + \frac{T}{B}\right)$.

For the CPU complexity, we note that checking whether $v \in V_{mem}^+$ amounts to checking whether $\text{ind}[v - v_{low}]$ has a positive degree, which is a $\mathcal{O}(1)$ operation, so construction of E_{mem} , and V_{mem}^+ (together with clearing it) takes $\Theta(|E_{mem}|) = \Theta(M)$ time. Construction of $N(u)$ and $N^+(u)$ thus also takes $\Theta(|N(u)|) = \Theta(d_{G^*}(u))$ time. Since each edge is examined once in a single iteration, each iteration incurs time $\mathcal{O}(|E|)$ for construction of these structures, for a total of $\Theta(|E|^2/M)$ time.

Set intersection of two ordered sets of size m, n takes time $\mathcal{O}(m + n)$ using a naive set intersection, thus the total complexity for the triangle operations is

$$\sum_{i=1}^h \sum_{u \in V} \sum_{v \in N_i^+(u)} (d_{G^*}(u) + d_{G^*}(v))$$

where $N_i^+(u)$ denotes $N^+(u)$ in the i -th iteration. First, note that any v is in at most 2 (consecutive) $N_i^+(u)$ for any given u . This is due to the small degree assumption, because if the adjacency is split the first time, the second time it will entirely fit in memory. Thus, we can reorder as follows:

$$\sum_{i=1}^h \sum_{u \in V} \sum_{v \in N_i^+(u)} = \sum_{u \in V} \sum_{i=1}^h \sum_{v \in N_i^+(u)} \leq 2 \sum_{u \in V} \sum_{v \in N^+(u)}$$

Examining each term separately:

$$\sum_{u \in V} \sum_{v \in N^+(u)} d_{G^*}(u) = \sum_{u \in V} d_{G^*}^2(u)$$

Additionally,

$$\begin{aligned} & \sum_{u \in V} \sum_{v \in N^+(u)} d_{G^*}(v) \\ &= \sum_{v \in V} d_{G^*}(v) (d_G(v) - d_{G^*}(v)) \\ &= \sum_{v \in V} d_G(v) \cdot d_{G^*}(v) - \sum_{v \in V} d_{G^*}^2(v) \end{aligned}$$

because $d_G(v) - d_{G^*}(v)$ represents the number of incoming vertices to v . The sums of $d_{G^*}^2(v)$ cancel out, so we need to calculate $\sum_{v \in V} d_G(v) \cdot d_{G^*}(v)$. This is where the arboricity becomes useful (Theorem IV.1 is adapted from — but is not identical to — the one given in [14]):

Theorem IV.1 (Ordering). $\sum_{v \in V} d_G(v) \cdot d_{G^*}(v) = \mathcal{O}(\alpha|E|)$

Proof:

$$\begin{aligned} \sum_{v \in V} d_G(v) \cdot d_{G^*}(v) &= \sum_{v \in V} \sum_{u \in N^+(v)} d_G(v) \\ &= \sum_{(v,u) \in E^*} d_G(v) \\ \text{(by orientation)} &\leq \sum_{(v,u) \in E} \min\{d(v), d(u)\} \\ \text{(by Theorem III.4)} &= \mathcal{O}(\alpha|E|) \end{aligned}$$

Note that the sorting of the original file takes $\mathcal{O}(\text{sort}(|E|))$ I/Os and $\mathcal{O}(|E| \ln |E|)$ CPU time [2], while the orientation itself takes $\mathcal{O}(\text{scan}(|E|))$ I/Os and $\mathcal{O}(|E|)$ CPU time, provided that the entire degree array can fit in memory.² If not, in the worst case (e.g. for the complete graph K_n), a vertex has a neighbor in every block. As a result, for each node, there must be $\mathcal{O}(|V|/B)$ I/Os, for a total of $\mathcal{O}(|V|^2/B)$ I/Os, just for the degree file. Because $|E| = \mathcal{O}(|V|^2)$, the total complexity is $\mathcal{O}(\text{scan}(|V|^2))$ I/Os and $\mathcal{O}(|E|)$ CPU time. This does not make a difference in dense graphs (except for the asymptotic constant), but it is still a point of omission for the analysis presented in [14]. Consequently, the overall complexity is identical to that of the baseline MGT, and is summarized in Theorem IV.2.

Theorem IV.2 (MGT Complexity). *In summary, our implementation of MGT has an I/O complexity of*

$$\mathcal{O}\left(\frac{|E|^2}{MB} + \frac{T}{B}\right)$$

and CPU complexity of

$$\mathcal{O}\left(\frac{|E|^2}{M} + \alpha|E|\right)$$

If the graph is not already sorted, an additional $\mathcal{O}(\text{sort}(|E|))$ I/Os and $\mathcal{O}(|E| \log |E|)$ computations are needed, and if $|V| < M$, $\mathcal{O}(\text{scan}(|V|^2))$ I/Os are necessary to orient it.

²The degrees and adjacency lists for all vertices are stored in separate files of sizes $|V|$ and $|E|$ respectively (Section V-B).

B. Distributed Framework

For our distributed protocol, every machine is sent a copy of the entire graph, and every available processor is allocated a (contiguous) set of edges S , and is responsible for finding all triangles in the graph which contain pivot edges in S , by using MGT. This is significantly different from the existing parallel triangle-counting systems, where different machines are responsible for different subsets of the vertices.

1) *Description:* In our framework, a *master* machine delegates responsibility to the N *client* machines (including itself), and combines their results. Because the orientation step need only occur once, it is the responsibility of the master to apply the degree-based order to the graph in question, before sending it over the network. The master then sends the oriented graph to each client, together with the indices that each processor is responsible for. Each core processes the adjacency list between the specified indices. The client combines the triangle counts (and possibly the triangle lists if necessary), and sends these back to the master, which atomically sums the results.

Our PDL framework is oblivious to how the orientation step is performed, and what specific (contiguous) subset of edges is assigned to each processor. In a naive implementation, orientation is performed sequentially, and edges are split equally to all processors. However, our master parallelizes the orientation, and includes a load-balancing step to equalize the time taken for triangle counting in each of the processors. More concretely, for **multicore orientation**, the master reads the entire degree array into memory (provided $|V| < PM$), and each core performs the orientation on a contiguous set of edges, which are then concatenated. **Load balancing** similarly calculates the number of in-edges for each vertex after orientation (equal to $d_G(v) - d_{G^*}(v)$), and splits the edges equally amongst the processors so that they are still contiguous, and the sum of these in-degrees are approximately the same among all processors. This provides an estimate for the average size of $N^+(u)$, and thus the number of required intersections.

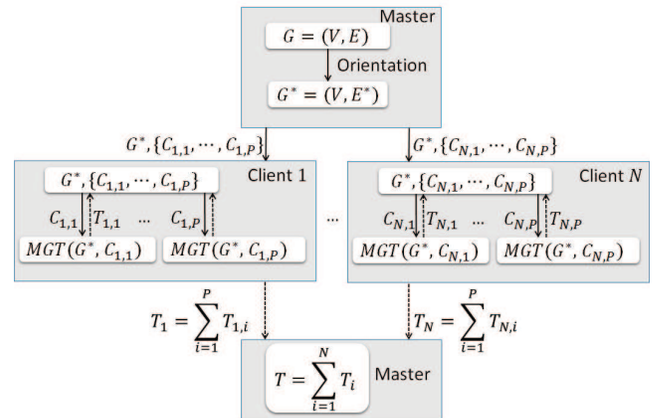


Figure 1: PDL protocol overview

Our protocol is illustrated in Figure 1. For clarity, the master process is duplicated, and is shown to run on a separate machine from the clients. Boxes represent different processes and clients, while ovals within boxes represent threads. Lines between boxes represent network traffic, with solid lines representing requests, and dotted lines answers. Finally, $C_{i,j}$ represents the “configuration” for processor j on machine i : the memory allocated for that thread, together with the section of the graph for which the processor is responsible. Note that the master starts the triangle counting operations before the network transfer has finished, so sending the graph does not pose a big bottleneck in practice.

2) *Analysis*: A problem with distributed algorithms using graph partitioning is that they assume each partition can fit in memory. Though for smaller graphs this may be the case, in dense graphs, such as the complete K_n , this is no longer true. Such algorithms require $\Theta(n^2)$ memory on *each* of the processors, and each of the NP processors must receive the entire graph. However, our algorithm requires memory proportional to the maximum degree, and the graph is only duplicated once per the N nodes. As a result, PDTL has lower network traffic and is preferable for dense graphs, and is also able to accommodate more computational environments. More concretely, PDTL incurs $\Theta(N \cdot (P + |E|) + T)$ **network** traffic in total, where T is the total number of triangles in the graph (or 0 for triangle counting), due to the communication cost per processor, and the duplication across the N nodes.

Since the **master** is responsible for orienting the graph according to the degree-based order, it incurs $\mathcal{O}(\text{scan}(|E|))$ I/Os and $\mathcal{O}(|E|)$ CPU time, assuming there is enough memory to hold V , as explained in Section IV-A2. This is true even for multicore orientation, as the graph is read once over all cores, but with an additional $\mathcal{O}(P)$ term: one for each of the cores. For load balancing, the vertex degrees are read once, with $PM = \mathcal{O}(|E|)$ edges sampled, and the results are stored for NP processors using $\mathcal{O}(\text{scan}(|V|))$ I/Os and $\mathcal{O}(|V| + \max(|E|, PM)) = \mathcal{O}(|E|)$ CPU time. The master is also responsible for adding the triangle counts received (in parallel) and also concatenating the triangle listing (sequentially), for a CPU complexity of $\mathcal{O}(N + T)$ and an I/O complexity of $\mathcal{O}((T + N)/B)$, as there might be an additional block for each of the N machines.

Since each processor is responsible for a unique (contiguous) section of the graph, there are no repeated **computations**. The chunk that each processor is responsible for has size $S = \frac{|E|}{NP}$, and each processor must make $R = \lceil \frac{S}{M} \rceil$ iterations over the graph.³ During these iterations, the graph is read once for creation of the vertex structures, and contributes $\mathcal{O}(|E|)$ processing time. Though it would be impossible to calculate exactly the amount of computations performed in each iteration for counting triangles as it de-

³For the load-balanced approach, this is only true in summation.

pends closely on the graph structure, we know by the proof of Theorem IV.2 that over all processors, these computations sum to $\mathcal{O}(\alpha \cdot |E|)$.⁴ Consequently, total computations across all processors are $\mathcal{O}\left(NP \cdot \lceil \frac{|E|}{NPM} \rceil \cdot |E| + \alpha \cdot |E|\right) = \mathcal{O}\left(NP \cdot |E| + \frac{|E|^2}{M} + \alpha \cdot |E|\right)$ as $\lceil \frac{|E|}{NPM} \rceil \leq \frac{|E|}{NPM} + 1$.

The **I/O** complexity is also easy to find. As above, each processor makes $R = \lceil \frac{|E|}{NPM} \rceil$ iterations over the graph, and outputs a variable number of triangles t , making its I/O complexity equal to $\mathcal{O}(R \cdot \text{scan}(|E|) + \text{scan}(t))$. As a result, the total I/O over all processors is $\mathcal{O}\left(NP \frac{|E|}{B} + \frac{|E|^2}{MB} + \frac{T}{B}\right)$. One of the important distinctions between PDTL and frameworks which load entire subgraphs in memory is that in PDTL, $|E|$ can still be larger than the total amount of available memory NPM . Moreover, we see that when $NPM > |E|$, we can reduce M to $\frac{|E|}{NP}$ without affecting any individual processor, whereas the total amount of memory needed in frameworks like PATRIC and PowerGraph can exceed $|E|$, due to overlapping subgraphs. Finally, it is important to note that the limiting factor after the graphs have been sent to all machines is the processor responsible for the highest number of triangles, so increasing the total number of processors is usually preferable, even with the same amount of total memory, as we also identify in Section V. Our findings are summarized in Theorem IV.3:

Theorem IV.3 (PDTL Complexity). *Letting T represent the number of triangles in the case of triangle listing and 0 in the case of triangle counting, and assuming $\frac{|E|}{NP} > d_{max}^*$, PDTL incurs across all cores a total of:*

- $\Theta(NP + N|E| + T)$ Network traffic
- $\mathcal{O}\left(NP|E| + \frac{|E|^2}{M} + \alpha|E|\right)$ CPU computations
- $\mathcal{O}\left(NP \frac{|E|}{B} + \frac{|E|^2}{MB} + \frac{T}{B}\right)$ I/Os

V. EVALUATION

Due to the wide range of environments in which PDTL can run, our extensive experiments cover single-core, limited memory machines to multi-machine, multi-core, large memory clusters. We discuss our setup and methodology in Section V-A, and introduce our datasets in Section V-B. We discuss the pre-processing and orientation operations in Section V-C, and compare them to those of competing algorithms. In Section V-D we discuss the core properties of our PDTL algorithm, in both the local and distributed environments, including the effects of load balancing. In Section V-E, we compare our algorithm against MGT, OPT, and PowerGraph extensively, and show that PDTL demonstrates superior performance.

⁴If S is smaller than the maximum degree, the complexity changes to $\mathcal{O}(NP \cdot \alpha \cdot |E|)$ as a single vertex can be split across NP machines.

A. Setup and Methodology

To illustrate the breadth of environments in which PDDL supports triangle counting, we conducted experiments in multiple different clusters and machine configurations:

- **Amazon EC2:** We used 4 Amazon EC2 `c3.8xlarge` instances, each of which contained 32 vCPU units, 60GB of memory, and were connected using a 10 Gigabit Ethernet network. For the PowerGraph measurements, we rented 4 Amazon EC2 `r3.8xlarge` that are similar to `c3.8xlarge` instances, but have 244GB of memory in order to satisfy PowerGraph’s memory requirements.
- **Local Cluster:** More distributed experiments were conducted in a local 4-node Linux cluster machine running 8 Virtual Xen nodes, each with 4 cores of an Intel Xeon E5607, 40GB of memory and a Samsung 840 SSD.
- **Local Multicore:** Additional multicore experiments were conducted in a local machine running Linux with 2 AMD Opteron 6344 CPUs for a total of 24 cores, 256GB of memory, and a Samsung 840 SSD.
- **Local Multicore Windows:** Since we only had access to an OPT [15] Windows binary, we used a Windows box, with performance similar to the above, having 2 Intel Xeon E5-2420 CPUs with support for 24 concurrent threads, 128GB memory, and a Samsung 840 SSD.

Our code was compiled with G++, using the `-O3` optimization option, and explicitly cleared disk caches before each experiment was run. Though our code fully supports triangle listing, our experiments only measured counting time, to allow comparison with alternatives. To account for random variation and general fluctuations, we repeated each experiment 3 times, and present the averages here. Full results can be found in our technical report [10].

B. Datasets

Table I lists the real and synthetic graphs used for our experiments. Our synthetic RMat graphs are scale-free graphs produced by the RMat generator [6], such that $\text{RMat-}n$ contains 2^n vertices and 2^{n+4} edges. Our triangle counts for real graphs have been verified to be correct by comparing to SNAP [17] and the results in OPT [15].

Our PDDL framework assumes that graphs are in binary, bi-directional format, with degrees of vertices and their out-edges in separate files. Moreover, we assume that edges are sorted by source and destination, partly for compatibility with the original MGT binary [12]. Since all efficient graph storage techniques operate on binary data, and all counting algorithms require efficient access to neighbors of a vertex, we exclude any time to convert a graph to this format from our discussion.⁵ However, because the degree-based ordering is non-standard, we consider the orientation cost separately,

⁵Note that OPT [15] requires that the input be sorted by vertex degree which is not included in the measurements, so this is a fair starting point.

and include it in our measurements. Similarly, we include copying costs from the master to the clients, to illustrate that our algorithm runs faster, even with graph duplication. Though other architectures such as NFS or HDFS were considered, we store a graph copy locally, since each graph is read at least once per processor. As seen in Table II, the average copying time is up to $10\times$ less than total time.

C. Preprocessing

Table III presents the time orientation took in our Local Multicore machine with 24 cores, compared to PowerGraph’s setup time and OPT’s database creation, whose pre-processing steps are much slower. Figure 2 shows a $5.2\times$ speed-up of multicore orientation over the single-core solution, and shows that our SSD is capped at 500MB/s.

Graphs	d_{max}^*	PDDL	PowerGraph	OPT
LiveJ1	687	1.4	-	106.8
Orkut	535	3.6	25.7	43.6
Twitter	4,102	32.8	233.2	437.6
Yahoo	1,540	235.6	-	-
RMAT-26	2,964	29.3	213.0	910.3

Table III: Preprocessing time (s): PDDL (Orientation), PowerGraph (Setup), OPT (Database Creation)

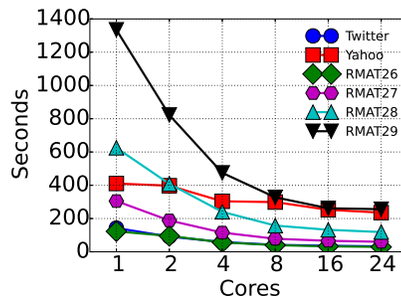


Figure 2: PDDL in Local Multicore: Orientation

D. PDDL Properties

In this section we examine the properties exhibited by PDDL without comparing it to other systems.

1) *Local:* We tested weak scaling of PDDL in the Local Multicore machine, by increasing the number of cores, but keeping the total amount of memory constant at 128GB, as shown in Figure 3. Specifically, using 2 cores halves calculation times, and this effect persists at a decreasing rate. Synthetic graphs exhibit better speedups due to their scale-free nature, as does the Twitter graph. However, due to its structure, the Yahoo graph only exhibits a $5\times$ speedup at 24 cores, compared to a $13\times$ speedup for the other graphs.

2) *Distributed:* We also ran distributed experiments in Amazon EC2, with 1GB of memory/core, as shown in Figure 4. We observe the following:

- The Twitter graph shows good scalability, while the Yahoo graph, being sparser and having a low average

Graph	Nodes	Edges	Triangles	Size	AvDeg	STD	MaxDeg	Source
soc-LiveJournal1	4.8M	68.0M	285,730,264	365MB	17.8	52	20,334	[17]
com-Orkut	3.1M	117.2M	627,584,181	917MB	76.0	155	33,313	[17]
Twitter	61.6M	1.5B	34,824,916,864	9.4GB	57.7	402	2,997,487	[16]
Yahoo	1.4B	6.6B	85,782,928,684	59GB	17.9	279	7,637,656	[1]
RMAT-26	67.1M	1.1B	51,559,452,522	8.4GB	61.2	632	430,269	[6]
RMAT-27	134.2M	2.1B	114,007,006,286	17GB	63.6	601	676,199	[6]
RMAT-28	268.4M	4.3B	251,913,686,661	34GB	66.0	660	1,062,289	[6]
RMAT-29	536.9M	8.6B	556,443,109,053	68GB	69.0	782	1,665,635	[6]

Table I: Graphs used for the experiments

Graph	1 node	2 nodes		3 nodes		4 nodes	
	Total time	Total time	Avg copy time	Total time	Avg copy time	Total time	Avg copy time
Twitter	164.2	127.4	13.5	116.0	16.2	109.0	19.1
Yahoo	397.9	364.9	106.0	338.6	112.4	433.8	186.4
RMAT-26	370.4	209.7	14.7	166.8	16.7	151.3	19.0
RMAT-27	841.1	479.5	27.9	373.1	27.9	306.6	33.3
RMAT-28	1876.8	1077.4	52.1	814.4	57.2	672.1	68.3
RMAT-29	4644.5	2531.0	106.6	1860.3	138.4	1565.0	154.6

Table II: PDTL in EC2: Total time and average copy time per remote node (s)

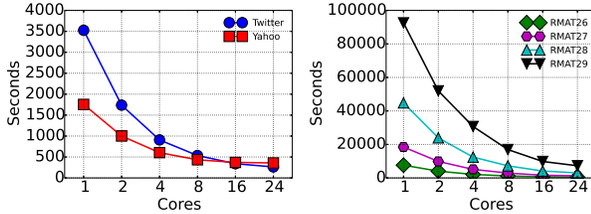


Figure 3: PDTL in Local Multicore: Total Time

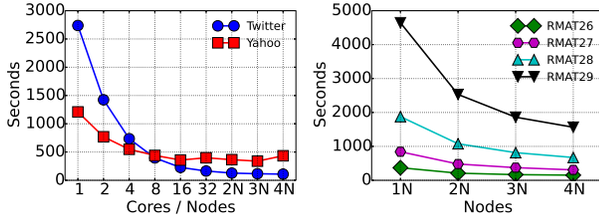


Figure 4: PDTL in EC2: Total Time

degree, does not benefit from more than 16 cores. Consequently, the structure of each graph heavily affects processing time, as also indicated by our analysis.

- RMAT graphs are much denser and more computationally intensive. This results in good scalability even up to 128 cores (4 nodes), as copy overhead is negligible.
- Comparing total computation time (Table II) and orientation time (Table III) further illustrates the unusual behavior of the Yahoo graph. Specifically, even though orientation only represents a small proportion of overall runtime for Twitter and RMAT, it comes close to 50% for the Yahoo graph on 3 nodes.
- Table II also details the copy times (averaged over the number of non-master nodes) for each of our graphs. As expected, this number scales with increasing graph size (recall Yahoo is larger than RMAT-28, but smaller than RMAT-29), and with increasing number of nodes (due to more limited network bandwidth). The Yahoo

graph also presents an anomaly in the copying of the graph which results in higher than expected increase in copy time for 4 nodes due to improper I/O balancing of the master node (Section V-D4): since the Yahoo graph results in heavy I/O due to its structure, it causes an initial I/O bottleneck when the master is performing its computations while also copying the graph.

3) *Memory*: To identify the effect of limited memory, we ran experiments in our Local Cluster varying the number of nodes, and the total amount of memory available per node (fixing $P = 4$ cores/node). As can be seen in Figure 5, the effect of limiting memory is negligible, and as a matter of fact more memory can lead to slightly higher costs due to array initialization overhead, as indicated in Section IV-B2.

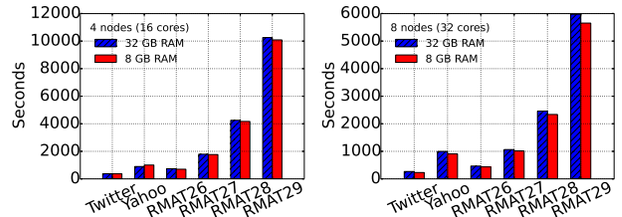


Figure 5: PDTL in Local Cluster: Memory vs. Calc Time

4) *I/O and CPU*: Despite the fact that PDTL is an external-memory algorithm, in our Amazon EC2 experiments we discovered that it is not I/O-bound. Specifically, measuring the total I/O for different numbers of cores and nodes on the Yahoo and Twitter graphs, we found that it represents a small percentage of the computation time (Figure 6). As explained in Section IV-B2, the absolute time spent on I/O operations increases as the number of cores increases, and is tied to the concrete graph structure, as indicated by the difference between Twitter and Yahoo. Figures 7 and 8 show the per node I/O and CPU breakdown for Twitter and Yahoo respectively. For the Twitter graph, our load-balancing mechanism works fairly well, and there is

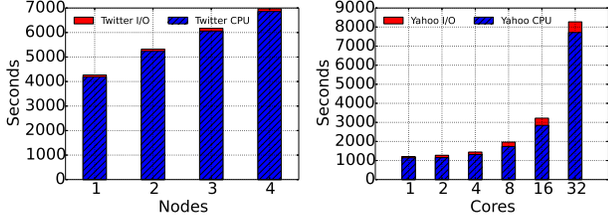


Figure 6: PDDL in EC2: Total CPU and I/O breakdown for various number of cores and nodes

no correlation between the CPU and the I/O operation times. However, the Yahoo graph is heavily skewed, and higher I/Os appear at the nodes with highest computation times, further illustrating the point that the concrete graph structure heavily influences the overall runtime of our algorithm.

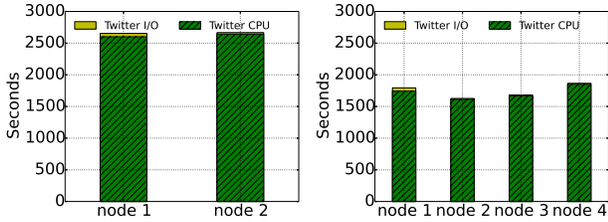


Figure 7: PDDL in EC2: Twitter CPU and I/O breakdown

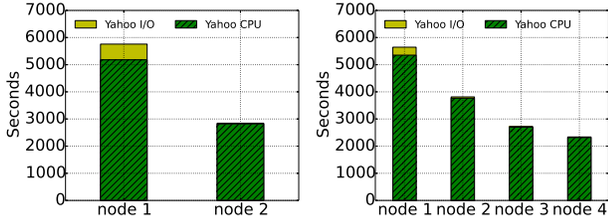


Figure 8: PDDL in EC2: Yahoo CPU and I/O breakdown

5) *Load Balancing*: We compared in our Local Multicore machine the naive approach of allocating the same number of edges for each core to our load balancing solution. Figure 9 contains our findings for 16 and 24 cores, and clearly illustrates up to a $3\times$ improvement on the calculation time, even for the Yahoo graph. Table IV details the total I/O and CPU computations for all processors within each node in Amazon EC2. As can be seen, our load-balancing mechanism leaves room for improvement, since the discrepancies between nodes increase as more nodes are added: even though there is only a 1% difference for 2 Twitter nodes, the difference increases to 13% for 4 nodes, while for Yahoo, the number increases from 87% to 130%.

E. PDDL Comparisons

In this section we examine our PDDL algorithm in the context of competing frameworks.

1) *MGT*: To compare PDDL against single-core MGT, we conducted experiments in Amazon EC2 nodes, looking at just the calculation times. Figure 10 shows that using just 2 processors halves the processing time for all real

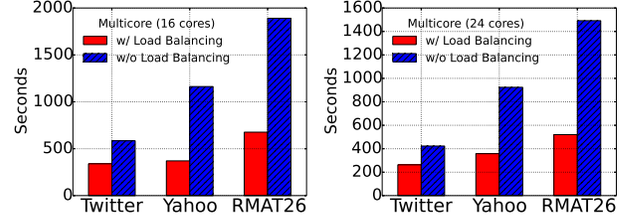


Figure 9: PDDL in Local Multicore: Load Balancing

graphs, and using 32 cores provides a $16\times$ speedup for the Twitter graph. Figure 11 similarly shows that the speedup for calculations of distributed, multicore PDDL over MGT reaches up to $55\times$ with 4 nodes. This effect is especially pronounced for scale-free RMAT graphs, whereas speedups reach $30\times$ for Twitter, but only $4\times$ for Yahoo. It should be noted that the comparison here is against our implementation of MGT, because the provided MGT binary [12] misreported triangle counts for some of the larger graphs.⁶ As a result, we cannot directly compare our implementation to the baseline one, but for completeness we note that for small graphs the performance was similar to the given binary.

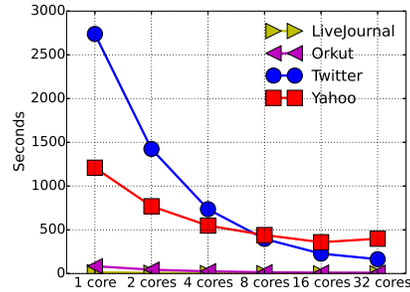


Figure 10: PDDL in EC2: Single Node Performance

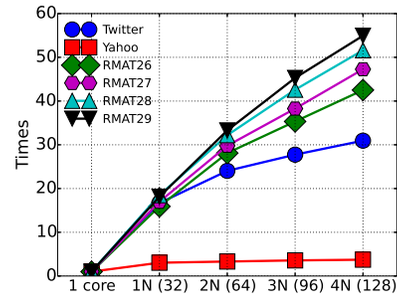


Figure 11: PDDL in EC2: Speedup over MGT

2) *OPT*: We compared our multi-core algorithm to OPT [15] in our two Local Multicore machines. We measured setup time (orientation for PDDL and database creation for OPT) and calculation time separately, and report our results when using 24 cores in Table V. With the exception of the LiveJournal dataset, our calculation time is always (and up

⁶MGT reported 627,506,739 triangles for Orkut and 559,420,538 triangles for Twitter, compared to 627,584,181 and 34.8B respectively.

Graph	2 nodes		3 nodes			4 nodes			
CPU: Twitter	2599.9	2643.0	2016.7	1920.4	2132.1	1741.0	1613.1	1663.5	1848.8
Yahoo	5177.7	2817.8	5296.2	3545.4	2412.4	5352.0	3765.8	2698.7	2317.0
RMAT-26	5173.9	4115.3	3863.1	3181.3	2758.7	3237.1	2674.3	2414.3	2099.3
I/O: Twitter	53.5	23.7	54.4	22.4	23.0	53.5	13.3	17.8	17.3
Yahoo	580.8	28.3	422.4	32.6	24.7	293.1	48.1	24.9	22.4
RMAT-26	155.2	29.2	143.5	22.1	20.9	85.2	19.6	18.4	15.3

Table IV: PDTL in EC2: Per node total CPU and I/O breakdown (s)

to $2\times$) faster than OPT’s calculations, and our setup time is up to $75\times$ faster. When looking at the total time, PDTL is up to $3.5\times$ faster for large graphs (and $7.8\times$ faster for LiveJournal). As can be seen in Figure 12, these effects remain for any number of cores, though they are even more pronounced for fewer ones. We should note that the OPT binary we received occasionally gave inconsistent triangle counts and could not run with $M = 128GB$, hence the memory discrepancy in our testing.

Graph	PDTL		OPT	
	Orientation	Calc	Database	Calc
LiveJ1	1.4	12.4	106.8	3.3
Orkut	3.6	11.4	43.6	11.7
Twitter	32.8	262.9	235.2	437.6
Yahoo	235.6	357.9	-	-
RMAT-26	29.3	520.4	910.3	1011.2

Table V: Local Multicore: PDTL and OPT Performance (s)

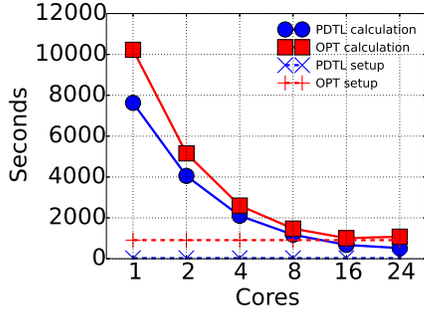


Figure 12: Local Multicore: PDTL (128GB) and OPT (100GB) on RMAT-26

3) *PowerGraph*: We also compared our distributed framework to PowerGraph [11] in Local Cluster and Amazon EC2. For a fair comparison, we consider two measures: the total runtime of both programs (including orientation in PDTL’s case), and the pure calculation time (including load-balancing costs for PDTL). For PowerGraph, the calculation time is the reported time of the triangle counting algorithm. For PDTL, the overall calculation time corresponds to the maximum individual calculation time between the different nodes. This is because the nodes start calculating as soon as they receive the files and, thus, the calculation time of the “straggler” node determines entirely the overall calculation time. The value *total*–*calc* thus represents the setup time for PowerGraph, while it represents a combination of network costs and workload imbalance for PDTL.

Figure 13 shows that although calculation times are similar (with PDTL presenting an advantage as the graphs become bigger), with setup times, PDTL is more than $2\times$ faster. Table VI illustrates this point more clearly, and also highlights that for larger graphs, PowerGraph runs out of memory. This is especially noteworthy, given that PowerGraph experiments were run on nodes with 244GB of memory **each**, for a total of 976GB, while our PDTL experiments were run using only 1GB/core (with much lower requirements) for a total of 128GB of memory. This validates our analysis in Section IV-B2, illustrating that partitioning-based approaches do not work for large graphs, and that external-memory algorithms like PDTL are needed.

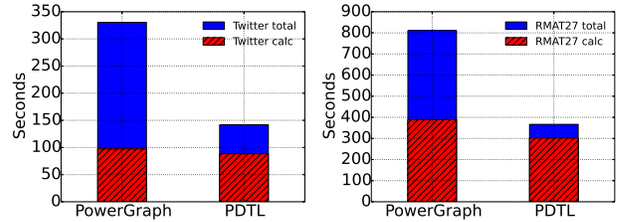


Figure 13: EC2 (4N): PDTL and PG breakdowns

Graph	PDTL		PowerGraph	
	Calc	Total	Calc	Total
Orkut	6.9	11.7	4.9	30.6
Twitter	88.5	141.8	97.3	330.5
Yahoo	323.9	669.4	OOM	OOM
RMAT-26	138.6	180.6	176.7	389.7
RMAT-27	302.1	366.7	389.5	810.5
RMAT-28	672.1	790.9	OOM	OOM
RMAT-29	1533.5s	1821.2	OOM	OOM

Table VI: PDTL and PowerGraph in EC2 (s). OOM represents an out-of-memory exception

4) *Other Frameworks*: Although we could not obtain a copy of the PATRIC binary, the original paper [3] indicates that PATRIC counts the triangles in the Twitter graph in 564s using 200 cores, and 4GB of memory/core. In another recent experiment [15], PATRIC was run in a cluster of 31 nodes with 12 threads per node (372 threads total) and 2GB of memory/core for a time of 608s. In either case, PDTL is $4\times$ faster using only 96 cores and 1GB of memory/core, and is still faster even when the number of cores is reduced to 8, again highlighting our fast performance under low memory requirements. Finally, it is worth briefly mentioning that MapReduce-based algorithms such as CTPP [21] are not competitive, spending 5520s calculating Twitter triangles using 40 nodes with 4GB of memory/node.

VI. CONCLUSIONS AND FUTURE WORK

In this paper, we presented our Parallel and Distributed Triangle Listing (PDTL) framework, the first distributed triangle listing and counting algorithm that focuses on external-memory I/O efficiency, but also provides theoretical CPU and Network guarantees. Our framework works well in a variety of computational environments, and is based on the recent MGT algorithm [14]. Key to our engineering approach is the combination of both a distributed setting and external memory. This gives us a high amount of parallelism whilst freeing us from the usual distributed constraint of fitting entire subgraphs in memory. Our resulting implementation is scalable and performs especially well in low-memory scenarios. As graphs become larger, the requirement of fitting even parts of a graph in memory will no longer be viable, as we also verified experimentally: our algorithm was able to accommodate for massive graphs containing over 8 billion edges with little memory, while competing partitioning-based frameworks ran out of memory even using almost 1TB of RAM.

More generally, our extensive experiments demonstrate that PDTL is highly scalable across multiple cores and machines, with low memory requirements, even for graphs with hundreds of millions of vertices, and billions of edges. Over the Twitter data set [16], our algorithm is faster than all of the state-of-the-art algorithms in distributed and parallel triangle counting: PDTL is $4\times$ faster than PATRIC [3], $3\times$ faster than OPT [15], and $2\times$ faster than PowerGraph [11].

Future work could focus investigation on different types of disks and file systems (for instance distributed file systems, or lazy evaluation), as a means of removing any copying bottlenecks that may exist. Such research could be informed by PowerGraph's general-framework, which fares better compared to triangle-specific systems. Even though its high memory requirements influence the results, it would be interesting to more formally investigate this. Additionally, more detailed investigations could try different techniques of load balancing, and provide a better understanding of the optimal number of machines and cores for any given graphs. As we identified in our experiments, scale-free graphs scale extremely well, even up to 8 machines, while the real-world Yahoo graph [1] exhibits a slowdown at even 4 nodes.

Overall, our framework provides a starting point towards many directions, including dynamic or approximate triangle counting, but more importantly for investigating other graph algorithms and processing systems which can benefit from our disk-based approach for large datasets.

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