Large-Scale Distributed Graph Computing Systems: An Experimental Evaluation

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ABSTRACT
With the prevalence of graph data in real-world applications (e.g., social networks, mobile phone networks, web graphs, etc.) and their ever-increasing size, many distributed graph computing systems have been developed in recent years to process and analyze massive graphs. Most of these systems adopt Pregel’s vertex-centric computing model, while various techniques have been proposed to address the limitations in the Pregel framework. However, there is a lack of comprehensive comparative analysis to evaluate the performance of various systems and their techniques, making it difficult for users to choose the best system for their applications. We conduct extensive experiments to evaluate the performance of existing systems on graphs with different characteristics and on algorithms with different design logic. We also study the effectiveness of various techniques adopted in existing systems, and the scalability of the systems. The results of our study reveal the strengths and limitations of existing systems, and provide valuable insights for users, researchers and system developers.

1. INTRODUCTION
Many distributed graph computing systems have been proposed to conduct all kinds of data processing and data analytics in massive graphs, including Pregel [15], Giraph [2], GraphLab [13], PowerGraph [7], GraphX [24], Mizan [11], GPS [19], Giraph++ [23], Pregelix [4], Pregel+ [26], and Blogel [25]. These systems are all built on top of a shared-nothing architecture, which makes big data analytics flexible even on a cluster of low-cost commodity PCs.

The majority of the systems adopt a “think like a vertex” vertex-centric computing model [15], where each vertex in a graph receives messages from its incoming neighbors, executes the user-specified computation and updates its value, and then sends messages to its outgoing neighbors. The vertex-centric computing model makes the design and implementation of scalable distributed algorithms simple for ordinary users, while the system handles all the low-level details. There are also a few extensions to the vertex-centric model, e.g., the edge-centric model in PowerGraph [7] and the block-centric models in [23, 25], to address various limitations in the vertex-centric systems. However, these models also suffer from other problems such as longer graph partitioning time.

While many distributed graph computing systems have emerged recently, it is unclear to most users and researchers what the strengths and limitations of each system are and there is a lack of overview on how these systems compare with each other. Thus, it is difficult for users to decide which system is better for their applications, or for researchers and system developers to further improve the performance of existing systems or design new systems.

In this paper, we first give a survey on existing distributed graph computing systems in Section 3. Among these systems, we conduct comprehensive experimental evaluation on Giraph [2], GraphLab/PowerGraph [13, 7], GPS [19], Pregel+ [26], and GraphChi [12]. We do not conduct experiments on other existing systems for various reasons given in Section 3.7. In Section 4, we discuss a set of graph algorithms that are popularly used to evaluate the performance of various systems in existing works, and we classify them into five categories where each category represents a different logic of distributed algorithm design. Then, in Section 5, we conduct a comprehensive analysis on the performance of various systems focusing on the following key objectives:

• To evaluate the performance of various systems in processing large graphs with different characteristics, including skewed (e.g., power-law) degree distribution, small diameter (e.g., small-world), large diameter, (relatively) high average degree, and random graphs.

• To evaluate the performance of various systems with respect to different categories of algorithms presented in Section 4.

• To study the effects of individual techniques used in various systems on their performance, hence analyzing the strengths and limitations of each system. The techniques to be examined include message combiner, mirroring, dynamic repartitioning, request-respond API, asynchronous and synchronous computation, and support for graph mutation. We also study the effects of a few algorithmic optimizations [20] and compare with GraphChi [12] as a single machine baseline.

• To test the scalability of various systems by varying the number of machines and CPU cores, the number of vertices and edges in graphs with different degree distributions.

Related work. Guo et al. [8] proposed a benchmarking suite to compare the performance of various systems for distributed graph computation. Their benchmark defines a comprehensive evaluation process to select representative metrics, datasets and algorithm categories. While this is a pioneering work in benchmarking distributed graph computing systems, the authors also admit that their
method has limitations in terms of selection of metrics and algorithmic coverage. Satish et al. [21] evaluated the performance of a number of systems, both distributed and single-machine, and identified the potential performance gap between these systems and the hand-optimized baseline code (whose performance is close to hardware limits). The results were then used to provide insights on how the systems may be improved to better utilize hardware capacity (e.g., memory/network bandwidth). Very recently (after the submission of our paper), we noticed the work by Han et al. [9], which evaluated four systems for their performance and usability, and identified potential areas of improvement in each system. Although both [9] and our work evaluated Giraph, GraphLab and GPS, due to different focus some findings are different. For example, the performance results of GPS we obtained are very different as we found that GPS’s large fixed overhead can be easily eliminated by setting its polling time appropriately. We also evaluated a new system, Pregel+, which records better performance in many aspects, while [9] present results (e.g., memory usage, network I/O) that we do not report. Compared with [8] and [21], we focus on the performance evaluation of vertex-centric distributed graph computing systems, among which, only GraphLab and Giraph were evaluated in [8] and [21]. We also used more algorithms and among them, only three were used in [8], [9] and [21]. Besides large real graphs, we also used larger synthetic random and power-law graphs to analyze the scalability of the systems. We studied the effects of different system optimization techniques, as well as algorithmic optimizations [20], on the performance of the systems, which were not studied in [8], [9] and [21]. We analyzed in greater details the differences between GraphLab’s asynchronous and synchronous modes. We also compared with GraphChi [12] as a single machine baseline. Thus, though [8], [9] and [21] made significant contributions in the evaluation of graph-computing systems, we believe that our work has also made substantial new contributions.

2. PRELIMINARY

We first define some basic graph notations. Let $G = (V, E)$ be a graph, where $V$ and $E$ are the sets of vertices and edges of $G$. If $G$ is undirected, we denote the set of neighbors of a vertex $v \in V$ by $\Gamma(v)$. If $G$ is directed, we denote the set of in-neighbors (out-neighbors) of a vertex $v$ by $\Gamma_in(v)$ ($\Gamma_out(v)$). Each vertex $v \in V$ has a unique integer ID, denoted by $id(v)$. The diameter of $G$ is denoted by $\delta(G)$, or simply $\delta$ when $G$ is clear from the context.

The distributed graph computing systems evaluated in this paper are all based on a shared-nothing architecture, where data are stored in a distributed file system (DFS), e.g., Hadoop’s DFS. The input graph is stored as a distributed file in a DFS, where each line records a vertex and its adjacency list. A distributed graph computing system consists of a cluster of $k$ workers, where each worker $w_i$ keeps and processes a batch of vertices in its main memory. Here, “worker” is a general term for a computing unit, and a machine can have multiple workers in the form of threads/processes.

A job is processed by a graph computing system in three phases as follows. (1) **Loading**: each worker loads a portion of vertices from the DFS into its main-memory; then workers exchange vertices through the network (e.g., by hashing on vertex ID as in Pregel) so that each worker $w_i$ finally keeps all and only those vertices that are assigned (i.e., hashed) to $w_i$. (2) **Iterative computing**: in each iteration, each worker processes its own portion of vertices sequentially, while different workers run in parallel and exchange messages. (3) **Dumping**: each worker writes the output of all its processed vertices to the DFS. Most existing graph-parallel systems follow the above three-phase procedure.

## 3. A SURVEY ON EXISTING SYSTEMS

We briefly discuss existing distributed graph computing systems, and highlight their distinguished features.

### 3.1 Pregel

Pregel [15] is implemented in C/C++ and designed based on the bulk synchronous parallel (BSP) model. It distributes vertices to different machines in a cluster, where each vertex $v$ is associated with the set of $v$’s neighbors. A program in Pregel implements a user-defined `compute()` function and proceeds in iterations (called *supersteps*). In each superstep, the program calls `compute()` for each active vertex. The `compute()` function performs the user-specified task for a vertex $v$, such as processing $v$’s incoming messages (sent in the previous superstep), sending messages to other vertices (to be received in the next superstep), and making $v$ vote to halt. A halted vertex is reactivated if it receives a message in a subsequent superstep. The program terminates when all vertices vote to halt and there is no pending message for the next superstep.

#### Message Combiner

If $x$ messages are to be sent from a machine $M_i$ to a vertex $v$ in a machine $M_j$, and some commutative and associative operation is to be applied to the $x$ messages in $v$’compute() when they are received by $v$, then these $x$ messages can be first combined into a single message which is then sent from $M_i$ to $v$ in $M_j$. To achieve this goal, Pregel allows users to implement a `combine()` function to specify how to combine messages that are sent from machine $M_i$ to the same vertex $v$ in machine $M_j$.

### 3.2 Giraph

Giraph [2] is implemented in Java by Yahoo! as an open source of Google’s Pregel. Later, Facebook built its Graph Search services upon Giraph, and further improved the performance of Giraph by introducing the following optimizations:

**Multi-threading.** Facebook adds multithreading to graph loading, dumping, and computation. In CPU bound applications, a speedup near-linear with the number of processors can be observed by multi-threading.

**Memory optimization.** The initial release of Giraph by Yahoo! requires high memory consumption, since all data types are stored as separate Java objects. A large number of Java objects greatly degrades the performance of Java Virtual Machine (JVM). Since object deletion is handled by Java’s Garbage Collector (GC), if a machine maintains a large number of vertex/edge objects in main...
memory, GC needs to track a lot of objects and the overhead can severely degrade the system performance. To decrease the number of objects being maintained, Java-based systems maintain vertices in main memory in their binary representation. Thus, the later Giraph system organizes vertices and messages as main memory pages, where each page is simply a byte array object that holds the binary representation of many vertices.

3.3 GPS

GPS [19] is another open source Java implementation of Google’s Pregel, with additional features. GPS extends the Pregel API to include an additional function, master.compute(), which provides the ability to access to all of the global aggregated values, and store global values which are transparent to the vertices. The global aggregated values can be updated before they are broadcast to the workers. Furthermore, GPS also introduces the following two techniques to boost system performance:

Large adjacency-list partitioning (LALP). When LALP is applied, adjacency lists of high-degree vertices are not stored in a single worker, but they are rather partitioned across workers. For each partition of the adjacency list of a high-degree vertex, a mirror of the vertex is created in the worker that keeps the partition. When a high-degree vertex broadcasts a message to its neighbors, at most one message is sent to its mirror at each machine. Then, the message is forwarded to all its neighbors in the partition of the adjacency list of the high-degree vertex.

Dynamic repartitioning (DP). DP repartitions the graph dynamically according to the workload during the execution process, in order to balance the workload among all workers and reduce the number of messages sent over the network. However, DP also introduces extra network workload to reassign vertices among workers, and the overhead can exceed the benefits gained.

3.4 Pregel+

Pregel+ [26] is implemented in C/C++ and each worker is simply an MPI process. In addition to the basic techniques provided in existing Pregel-like systems, Pregel+ introduces two new techniques to further reduce the number of messages.

Mirroring. Mirroring is similar to LALP in GPS; but Pregel+ integrates both mirroring and message combiner, and the Pregel+ system selects vertices for mirroring based on a cost model that analyzes the tradeoff between mirroring and message combining. Thus, the integration of mirroring and message combiner in Pregel+ leads to significantly more effective message reduction than applying combiner alone. Pregel+ also supports an additional API that allows mirroring to be used in applications where message values depend on the edge fields (e.g., single-source shortest path computation), which is not supported by LALP in GPS.

Request-Respond API. This API allows a vertex $u$ to request another vertex $v$ for a value, $o(v)$, and the requested value will be available to $u$ in the next iteration. The technique can effectively reduce the number of messages passed, since all requests from a machine to the same target vertex $v$ are merged into one request (e.g., as Figure 1 shows, requests from all $u_i$ in machine $M_1$ are merged into one request sent to $v_j$ in machine $M_2$).

3.5 GraphLab and PowerGraph

GraphLab 2.2 (which includes PowerGraph) is implemented in C/C++. Unlike Pregel’s synchronous data-pushing model and message passing paradigm, GraphLab [13] adopts an Gather, Apply, Scatter (GAS) data-pulling model and shared memory abstraction. A program in GraphLab implements a user-defined GAS function for each vertex. To avoid the imbalanced workload caused by high-degree vertices in power-law graphs, a recent version of GraphLab, called PowerGraph [7], introduces a new graph partition scheme to handle the challenges of power-law graphs as follows.

In the Gather phase, each active vertex collects information from adjacent vertices and edges, and performs a generalized sum operation over them. This generalized sum operation must be commutative and associative, ranging from a numerical sum to the union of the collected information. In the Apply phase, each active vertex can update its value based on the result of the generalized sum and its old value. Finally, in the Scatter phase, each active vertex can activate the adjacent vertices. However, unlike Pregel’s message passing paradigm, GraphLab can only gather information from adjacent edges and scatter information to them, which limits the functionality of the GAS model. For example, the S-V algorithm to be described in Section 4.1 is hard to be implemented in GraphLab.

GraphLab maintains a global scheduler, and workers fetch vertices from the scheduler for processing, possibly adding the neighbors of these vertices into the scheduler. The GraphLab engine executes the user-defined GAS function on each active vertex until no vertex remains in the scheduler. The GraphLab scheduler determines the order to activate vertices, which enables GraphLab to provide with both synchronous and asynchronous scheduling.

Asynchronous execution. Unlike the behaviors in a synchronous model, changes made to each vertex and edge during the Apply phase are committed immediately and visible to subsequent computation. Asynchronous execution can accelerate the convergence of some algorithms. For example, the PageRank algorithm can converge much faster with asynchronous execution. However, asynchronous execution may incur extra cost due to locking/unlocking and intertwined computation/communication.

Synchronous execution. GraphLab also provides a synchronous scheduler, which executes the GAS phases in order as an iteration. The GAS function of each active vertex runs synchronously with a barrier at the end of each iteration. Changes made to the vertex value is committed at the end of each iteration. Vertices activated in each iteration are executed in the subsequent iteration.

Vertex-cut partitioning. PowerGraph partitions an input graph by cutting the vertex set, so that the edges of a high-degree vertex are handled by multiple workers. As a tradeoff, vertices are replicated across workers, and communication among workers are required to guarantee that the vertex value on each replica remains consistent.

3.6 GraphChi

GraphChi [12] is implemented in C/C++, which is a single-machine system that can process massive graphs from secondary storage. In addition to the vertex-centric model, GraphChi introduces two new techniques to process large graphs in a single PC.

Out-of-core computation. An innovative out-of-core data structure is used to reduce the amount of random access to secondary storage. The parallel sliding windows algorithm partitions the input graph into subgraphs, called shards. In each shard, edges are sorted by the source IDs and loaded into memory sequentially.

Selective scheduling. GraphChi supports selective scheduling in order to converge faster on some parts of the graph, particularly when the change on values is significant. Each vertex in the update() function (similar to apply() in GraphLab) can add its neighbors to the scheduler and conduct selective computation.

3.7 Other Systems

In this paper, we conduct experimental evaluation on Giraph [2], GraphLab/PowerGraph [13, 7], GPS [19], and Pregel+ [26], which
we have discussed in Sections 3.2-3.5 and we also give an overview of various features supported by these systems in Figure 2. There are also a number of other systems that we do not evaluate experimentally, which we explain in this subsection.

Mizan [11] is a C++ optimized Pregel system that supports dynamic load balancing and vertex migration, based on runtime monitoring of vertices to optimize the end-to-end computation. However, Mizan performs pre-partitioning separately which takes much longer compared with Giraph and GPS, and the overhead of pre-partitioning can exceed the benefits. For this reason, we could not run Mizan on some large graphs used in this paper and hence we do not include it in our experimental evaluation.

GraphX [24] is a graph parallel system, and it supports GraphLab and Pregel abstractions. Since GraphX is built upon the more general data parallel Spark system [28], the end-to-end performance of pipelined jobs can be superior when they are all implemented in Spark. Consider the task of first extracting the link graph from Wikipedia, and then computing PageRank on the link graph. Compared with implementing and running both jobs in Spark (with the second job done by GraphX), the traditional method of running the first (second) job by Hadoop (by a vertex-centric system) is more costly since it requires that the output of the first job be dumped to HDFS and reloaded by the second job. However, if only graph computation time is considered, GraphX is generally slower than GraphLab as reported in [24].

4. ALGORITHMS

We use seven graph algorithms, which are classified into five categories based on the behaviors of the compute function in Pregel-like systems and the GAS function in GraphLab and PowerGraph.

Always-active. An algorithm is always-active if every vertex in every superstep sends messages to all its neighbors. Thus, the distribution of messages sent/received by all active vertices is the same across supersteps, i.e., the communication workload of every machine remains the same. Typical examples include PageRank [15] in synchronous computation and Diameter Estimation [10].

GraphLab’s async algorithms. This category is specifically for algorithms that are designed to run on GraphLab’s (also PowerGraph’s) asynchronous computation model. Such algorithms add vertices to the scheduler, and then workers fetch vertices from the scheduler and pull data from neighboring edges and vertices for processing. The asynchronous execution can accelerate the convergence of algorithms such as the asynchronous PageRank and Graph Coloring [20] algorithms for GraphLab and PowerGraph.

Graph traversal. Graph traversal is a category of graph algorithms for which there is a set of starting vertices, and other vertices are involved in the computation based on whether they receive messages from their in-neighbors. Attribute values of vertices are updated and messages are propagated along the edges as the algorithm traverses the graph. Algorithms such as HashMin [18] and Single-Source Shortest Paths [15] are in this category.

Multi-phase. For this category of algorithms, the entire computation can be divided into a number of phases, and each phase consists of some supersteps. For example, in Bipartite Maximal Matching [1], there are four supersteps to simulate a three-way handshake in each phase. The SV [22] algorithm also falls into this category, since it simulates tree hooking and star hooking in each phase.

Graph mutation. Algorithms in this category need to change the topological structure of the input graph through edges and/or vertices addition and/or deletion. For example, the greedy graph coloring algorithm that removes a maximal independent set from the current graph iteratively falls into this category. Systems that do not support edge deletion, such as GraphLab and PowerGraph, cannot straightforwardly support these algorithms.

4.1 Algorithm Description

We now describe the seven graph algorithms.

4.1.1 PageRank

Given a directed web graph $G = (V, E)$, where each vertex (page) $v$ links to a list of pages $\Gamma_{\text{out}}(v)$, the problem is to compute the PageRank value, $pr(v)$, of each $v \in V$.

The typical PageRank algorithm [15] for Pregel-like systems works as follows. Each vertex $v$ keeps two fields: $pr(v)$ and $\Gamma_{\text{out}}(v)$. In superstep 0, each vertex $v$ initializes $pr(v) = 1$ and sends each out-neighbor of $v$ a message with a value of $pr(v)/\lvert \Gamma_{\text{out}}(v) \rvert$. In superstep $i (i > 0)$, each vertex $v$ sums up the received PageRank values, denoted by $\text{sum}$, and computes $pr(v) = 0.15 + 0.85 \times \text{sum}$. It then distributes $pr(v)/\lvert \Gamma_{\text{out}}(v) \rvert$ to each of its out-neighbors. This process terminates after a fixed number of supersteps or the PageRank distribution converges.

The asynchronous version of PageRank for GraphLab and PowerGraph, named as Async-PageRank, works as follows. Each vertex $v$ keeps three fields: $pr(v)$, $\Gamma_{\text{in}}(v)$ and $\Gamma_{\text{out}}(v)$, where $pr(v)$ is initialized as 1. We define the generalized sum in the GAS function as a numerical sum. Then for each active vertex $v$ fetched from the scheduler, in the Gather phase, the values $pr(u)/\lvert \Gamma_{\text{out}}(u) \rvert$ for all neighbors $u \in \Gamma_{\text{in}}(v)$ are gathered and summed up as $\text{sum}$. In the Apply phase, we update $pr(v) = 0.15 + 0.85 \times \text{sum}$. In the Scatter phase, if the change in value of $pr(v)$ is greater than $\epsilon$ (typically, $\epsilon$ is set to 0.01), we add each vertex $u$ in $\Gamma_{\text{out}}(v)$ to the scheduler. This process terminates after a fixed number of supersteps.

4.1.2 Diameter Estimation

Given a graph $G = (V, E)$, we denote the distance between $u$ and $v$ in $G$ by $d(u, v)$. We define the neighborhood function $N(h)$ for $h = 0, 1, \ldots, \infty$ as the number of pairs of vertices that can reach each other in $h$ hops or less.

Each vertex $v$ keeps two fields: $N[h; v]$ and $\Gamma_{\text{out}}(v)$. where $N[h; v]$ indicates the set of vertices $v$ can reach in $h$ hops. In superstep 0, each vertex $v$ sets $N[0; v]$ to $\{v\}$ and broadcasts $N[0; v]$ to each $u \in \Gamma(v)$. In superstep $i (i > 0)$, each vertex $v$ receives messages from its neighbors and set the value of $N[i; v]$ as the union of $N[i − 1; v]$ and $N[i − 1; u]$ for all $u \in \Gamma(v)$. A global aggregator is used to compute the total pairs of vertices, denoted by $N(i)$, that can be reached from each other after superstep $i$. The algorithm terminates if the following stop condition is true: in superstep $i$, $N(i)$ is less than or equal to $(1 + \epsilon) \times N(i − 1)$.

To handle the large volume of each vertex’s neighborhood information, i.e., $N[h; v]$, the algorithm applies the idea of Flajolet-Martin [5], which was also used in the ANF algorithm [17].

4.1.3 Single-Source Shortest Paths (SSSP)

Let $G=(V, E)$ be a weighted graph, where each edge $(u, v) \in E$ has length $\ell(u, v)$. The length of a path $P$ is equal to the sum of the length of all the edges on $P$. Given a source $s \in V$, the SSSP algorithm computes a shortest path from $s$ to every other vertex $v \in V$, denoted by $SP(s, v)$, as follows. Each vertex $v$ keeps two fields: $(\text{prev}(v), \text{dist}(v))$ and $\Gamma_{\text{out}}(v)$, where $\text{prev}(v)$ is the vertex preceding $v$ on $SP(s, v)$ and $\text{dist}(v)$ is the length of $SP(s, v)$. Each out-neighbor $u$ in $\Gamma_{\text{out}}(v)$ is also associated with $\ell(v, u)$.

Initially, only $s$ is active with $\text{dist}(s) = 0$, and $\text{dist}(v) = \infty$ for any other vertex $v$. In superstep 0, $s$ sends a message $(s, \text{dist}(s) + \ell(s, u))$ to each $u \in \Gamma_{\text{out}}(s)$, and votes to halt. In superstep $i$
(i > 0), if a vertex \( v \) receives messages \([u, d(w)]\) from any of
\( v \)'s in-neighbor \( w \), then \( v \) finds the in-neighbor \( w^* \) such that \( d(w^*) \)
is the smallest among all \( d(w) \) received. If \( d(w^*) < dist(v) \), \( v \)
updates \([\text{prev}(v), \text{dist}(v)] = (w^*, d(w^*))\), and sends a message
\([v, \text{dist}(v) + \ell(v, u)]\) to each out-neighbor \( u \in \Gamma_{\text{out}}(v) \). Finally, \( v \)
votes to halt.

4.1.4 HashMin

Assume each CC \( C \) in an undirected graph \( G \) has a unique ID, and for each vertex \( v \) in \( C \), let \( cc(v) \) be the ID of \( C \). Given \( G \),
HashMin [18] computes \( cc(v) \) for each \( v \) in \( G \), and hence all CCs for
\( G \), as all vertices with the same \( cc(v) \) form a CC.

Each vertex \( v \) keeps two fields: \( min(v) \) and \( \Gamma(v) \), where \( min(v) \)
is initialized as the ID of the vertex itself. HashMin broadcasts the
smallest vertex ID seen so far by each vertex \( v \) as follows. In super-
step 0, each vertex \( v \) sets \( min(v) \) to be the smallest ID among \( id(v) \) and
\( id(u) \) of all \( u \in \Gamma(v) \), broadcasts \( min(v) \) to all its neighbors,
and votes to halt. In superstep \( i \) \((i > 0)\), each vertex \( v \) receives
messages from its neighbors; let \( min^* \) be the smallest ID received,
if \( min^* < min(v) \), \( v \) sets \( min(v) = min^* \) and broadcasts \( min^* \)
to its neighbors. All vertices vote to halt at the end of a superstep.
When the process converges, \( min(v) = cc(v) \) for all \( v \).

4.1.5 Shiloach-Vishkin’s Algorithm (SV)

The HashMin algorithm requires \( O(d) \) steps for computing CCs,
which is too slow for graphs with a large diameter, such as
spatial networks where \( d \approx O(\sqrt{n}) \). The SV algorithm [22]
can be translated into a Pregel algorithm which requires \( O(\log n) \)
supersteps [27], which can be much more efficient than HashMin for
computing CCs in general graphs.

The SV algorithm groups vertices into a forest of trees, so that
all vertices in each tree belong to the same CC. The tree here is
relaxed to allow the root to have a self-loop. Each vertex \( v \) keeps
two fields: \( D[u] \) and \( \Gamma_{\text{out}}(u) \), where \( D[u] \) points to the parent of \( u \)
in the tree and is initialized as \( u \) (i.e., forming a self loop at \( u \)).

The SV algorithm proceeds in phases, and in each phase, the
pointers are updated in the following three steps: (1) for each edge
\( (u, v) \), if \( u \)'s parent \( w \) is the root, set \( w \) as a child of \( D[v] \),
which merges the tree rooted at \( w \) into \( v \)'s tree; (2) for each edge
\( (u, v) \), if \( u \) is in a star, set \( u \)'s parent as a child of \( D[v] \); (3) for each vertex \( v \),
set \( D[v] = D[D[v]] \). We perform Steps (1) and (2) only if
\( D[u] < D[v] \), so that if \( u \)'s tree is merged into \( v \)'s tree due to edge
\( (u, v) \), then edge \( (u, v) \) will not cause \( v \)'s tree to be merged into
\( u \)'s tree again. The algorithm ends when every vertex is in a star.

4.1.6 Bipartite Maximal Matching (BMM)

Given a bipartite graph \( G = (V, E) \), this algorithm computes a
BMM, i.e., a matching to which no additional edge can be added
without sharing an end vertex. The algorithm [15] proceeds in
phases, and in each phase, a three-way handshake is simulated.

Each vertex \( v \) keeps three fields: \( S[v], M[v] \) and \( \Gamma_{\text{out}}(v) \), where
\( S[v] \) indicates which set the vertex is in (L or R) and \( M[v] \) is the
name of its matched vertex (initialized as \(-1\) to indicate that \( v \)
not yet matched). The algorithm computes a three-way handshake in four super-
steps: (1) each vertex \( v \), where \( S[v] = L \) and \( M[v] = -1 \), sends a
message to each of its neighbors \( u \in \Gamma_{\text{out}}(v) \) to request a match;
(2) each vertex \( v \), where \( S[v] = R \) and \( M[v] = -1 \), randomly chooses
one of the messages \( w \) it receives, sends a message to \( w \)
granting its request for match, and sends messages to other
requesters \( w' \neq w \) denying its request; (3) each vertex \( v \), where
\( S[v] = L \) and \( M[v] = -1 \), chooses one of the grantors \( w \) it receives
sends an acceptance message to \( w \); (4) each vertex \( v \), where \( S[v] = R \) and \( M[v] = -1 \), receives at most one acceptance
message, and then changes \( M[v] \) to the acceptance message’s
value. All vertices vote to halt at the end of a superstep.

4.1.7 Graph Coloring (GC)

Given an undirected graph \( G = (V, E) \), GC computes a color
for every vertex \( v \in V \), denoted by \( color(v) \), such that if \( (u, v) \in E \),
\( color(u) \neq color(v) \).

The GC algorithm for Pregel-like systems normally adopts the
greedy GC algorithm from [6]. The algorithm iteratively finds a
maximal independent set (MIS) from the set of active vertices, as-
signs the vertices in the MIS a new color, and then removes them
from the graph, until no vertices are left in the graph. Each iterative
phase is processed as follows, where all vertices in the same MIS
are assigned the same color \( c \): (1) each vertex \( v \in V \) is selected as a
tentative vertex in the MIS with a probability \( 1/(2 \* |\Gamma(v)|) \); if
a vertex has no neighbor (i.e., an isolated vertex or becoming iso-
lated after graph mutation), it is a trivial MIS; each tentative vertex
\( v \) then broadcasts \( id(v) \) to all its neighbors; (2) each tentative vertex
\( v \) receives messages from its tentative neighbors; let \( min^* \) be
the smallest ID received, if \( min^* > id(v) \), then \( v \) is included in the
MIS and \( color(v) = c \), and \( id(v) \) is broadcast to its neighbors;
(3) if a vertex \( v \) receives messages from its neighbors (that
have been included in the MIS in superstep (2)), then for each such
neighbor \( u \), delete \( v \) from \( \Gamma(u) \).

4.2 Algorithmic Optimizations

Apart from algorithm categorization, we also describe three al-
gorithmic optimizations [20] here that can improve the performance
of distributed graph computing systems on certain algorithms.

Finishing Computations Serially (FCS). Some algorithms may
run for a large number of supersteps, even though the later super-
steps are merely executing on a small fraction of the graph, called
the active-subgraph. FCS monitors the size of the active-subgraph
and sends it to the master for serial computation as soon as the size
is below a threshold (5M edges by default [20]), so as to terminate
the computation earlier without running a prolonged number of
supersteps. The results computed in the master are then sent back
to the workers. FCS can be applied to algorithms in which an inac-
tive vertex will not be activated again in later process. Among
the algorithms we discussed, BMM and GC have this property.

Edge Cleaning On Demand (ECOD). Edge cleaning in an al-
gorithm removes edges from the graph. ECOD delays the operation
of edge cleaning and regards the edges as stale edges until they
are involved in later computation where they are demanded to be
removed.

Single Pivot (SP). SP is a heuristic for speeding up the compu-
tation of connected components (CC). SP first samples a vertex \( v \)
called the pivot, and runs the cheaper BFS algorithm instead of the
CC algorithm from \( v \). Then, a standard CC algorithm is run on
the graph excluding the CC that contains \( v \). Theoretically, \( v \) has a
higher probability to be in the giant CC of the graph, and the graph
excluding the giant CC may be much smaller.

5. EXPERIMENTAL EVALUATION

We now evaluate the performance of Giraph, GPS, Pregel+,
GraphLab (we use GraphLab 2.2 which includes all the features
of PowerGraph), and use GraphChi as a single machine baseline.
We release all the source codes of the algorithms used in our evalu-
ation in www.cse.cuhk.edu.hk/pregelplus/exp, while the source codes of the different systems can be found in their own
websites.
Datasets. We used six large real-world datasets, which are from four different domains as shown in Figure 3: (1) web graph: WebUK, Friendster and Twitter have average degree relatively higher than other large real-world graphs; USA and WebUK have a large diameter, while Friendster, Twitter and BTC have a small diameter.

We also used synthetic datasets for scalability tests, where we generate power law graphs using Recursive Matrix (R-MAT) model [3] and random graphs using Przzer algorithm [16].

Experimental settings. We ran our experiments on a cluster of 15 machines, each with 48 GB DDR3-1,333 RAM, two 2.0GHz Intel(R) Xeon(R) E5-2620 CPU, a SATA disk(6Gb/s, 10k rpm, 64MB cache) and a Broadcom Gigabit Ethernet BCM5720 network adapter, running 64-bit CentOS 5.5 with Linux kernel 2.6.32. Giraph 1.0.0 and GPS (rev. 112) are built on JDK 1.7.0 Update 45, the hadoop DFS is built on Apache Hadoop 1.2.1. Pregel+, GraphChi (2014.4.30) and GraphLab 2.2 are compiled using GCC 4.4.7 with -O2 option enabled, and MPICH 3.0.4 is used.

Unless otherwise stated, we use all 15 machines for all distributed systems for all experiments; and we use 8 cores in each machine for GraphChi, Giraph and GraphLab which run with multithreading, and 8 processes (also using 8 cores) in each machine for GPS and Pregel+. There is no limit set on the amount of memory each system can use, i.e., all the systems have access to all the available memory (48GB) in each machine. GPS’s polling time is set to 10 ms in order to reduce the fixed overhead in each superstep.
5.1 System Performance Overview

We first evaluate (1) the performance of the various systems w.r.t. different algorithm categories (discussed in Section 4), (2) the performance of the various systems on graphs with different characteristics, and (3) the performance of a distributed system compared with that of a baseline single-machine system. The results to be presented in Subsections 5.1.1–5.1.3 give readers an overview on the performance of the various systems.

In the experiments in Sections 5.1.2–5.1.3, we enable the techniques of the various systems that give the best performance, while in Sections 5.2–5.5 we analyze the effects of each individual technique. We run the systems on every algorithm-graph combination that makes sense (except for HashMin, SV, and Coloring which are applicable on undirected graphs only, and Bipartite Maximal Matching which runs on bipartite graphs only). We also do not report all system-algorithm combinations, since it is not clear how pointer jumping in SV and edge deletion in graph coloring can be implemented in GraphLab.

With limited space, we only report 16 figures (in Figures 4(a)–4(f)) that are sufficient to reflect the overall performance over all the figures (reported in Figures 3–9 in our technical report [14]).

5.1.1 Performance on Different Algorithms

We first analyze the performance of the various systems w.r.t. different algorithm categories.

Performance on always-active algorithms. Figures 4(a)–4(f) report the performance of the systems on two representative always-active algorithms, i.e., synchronous PageRank and Diameter Estimation. It is difficult to draw a clear conclusion on which system has the best performance. Overall, GPS and Pregel+ have better performance in most cases. Between Giraph and GraphLab, GraphLab has the best performance in some cases while Giraph ran out of memory for diameter estimation on WebUK. Moreover, taking into account all the results in Figures 3 and 4 in [14], GraphLab is faster than Giraph in more cases.

Performance on graph-traversal algorithms. Figures 4(g)–4(k) report the performance of the systems on two representative graph-traversal algorithms, i.e., HashMin and SSSP. The results show that Pregel+ clearly outperforms all the other systems (Figures 5 and 6 in [14] show the same trend). GPS has better performance than Giraph and GraphLab in most cases.

Performance on multi-phase algorithms. Figures 4(l)–4(n) report the performance of the systems on two representative multi-phase algorithms, i.e., BMM and SV. Pregel+ always has the best performance for this category of algorithms, while GPS is faster than Giraph and GraphLab.

Performance on graph mutation. To test the performance on graph mutation, we use the graph coloring algorithm. We do not report GraphLab since it does not support edge deletion. Figures 4(o) and 4(p) (also Figure 9 in [14]) show that Pregel+ is much faster than both GPS and Giraph for graph coloring.

GraphLab supports both synchronous and asynchronous execution and we evaluate the performance as follows. First, for a graph with a large diameter, e.g., USA road network, Figures 5(a)–5(b) show that asynchronous execution is significantly faster than synchronous execution. This is because changes made to each vertex and edge during the apply phase in asynchronous mode are committed immediately and visible to subsequent computation; while in synchronous mode, the change commits are delayed till the end of each superstep, leading to slower convergence. However, for processing graphs with a small diameter as shown in Figures 5(c)–5(d), the overhead of locking/unlocking is not paid off by the faster convergence of asynchronous execution.

However, for some algorithms, asynchronous execution can lead to faster convergence even for small-diameter graphs. For example, for asynchronous PageRank, most vertices can converge after only a small number of updates. On the contrary, in synchronous execution, all vertices need to update their PageRank values and distribute their new values to neighbors. In each superstep, there are $O(n)$ updates made and $O(m)$ messages transmitted. In asynchronous execution, the global scheduler only maintains the vertices that need to be updated. If there is a significant change in some vertex’s PageRank value, then it activates its neighbors and puts them into the global scheduler. Figures 6(a)–6(b) show that PageRank takes only 554.4 seconds on the small-diameter Twitter graph using 508,251,513 updates, and takes 1,037.9 seconds on the large-diameter WebUK graph using 847,312,369 updates. However, the synchronous PageRank uses 4,679,591,698 and 11,893,340,560 updates, respectively, and are also much slower.
Overall performance. Figure 7 gives a ranking on the overall performance of the systems for different algorithm categories. Pregel+ and GPS have superior performance for always-active algorithms, because those algorithms generate a lot of messages, which is effectively addressed by Pregel+ and GPS’s message reduction techniques (see Section 5.3). Graph-traversal algorithms also generate a large number of messages from active vertices (especially high-degree ones), usually in the preceding supersteps. Pregel+ has better performance than GPS because of its integration of mirroring and combiner, which is more effective than LALP alone in GPS. Although vertex replica in GraphLab is similar to mirroring, such replica is constructed for every vertex instead of just high-degree vertices and so the extra overhead is not paid off. For SV, Pregel+’s request-response technique is effective (see Section 5.4). But in general, no system has any specific technique to improve its performance on multi-phase algorithms, though the message reduction techniques still help improve the performance. For graph coloring, GPS and Giraph require users to subclass a separate Edge Class during the graph loading phase and edge deletion requests must be made in the `compute()` function; while in Pregel+, the edge information of a graph is stored with vertices, which simplifies the API and enables faster edge addition/deletion.

C++ vs. Java. Among the systems, Giraph and GPS are implemented in Java, and GraphLab and Pregel+ are implemented in C++. While it is difficult to tell from our results which language, C++ or Java, leads to better performance, Figure 4(d) shows that the Java-based systems ran out of memory on the large WebUK graph for diameter estimation (GPS also ran out of memory on Friendster as shown in Figure 4(b) in [14]). This is mainly because a Java object takes more space than a C++ object; moreover, Java uses Garbage Collector to automatically handle object deletion, which cannot keep the pool of objects small in an optimal manner and hence often leads to larger memory usage. As for running time, none of the distributed systems has an implementation in both Java and C++, and hence we cannot make a comparison that gives a clear conclusion. But Java-based systems incur extra (de)serialization cost for processing objects in binary representation in memory. Moreover, the single-machine system GraphChi was implemented in both Java and C++, and [12] remarks that the Java implementation ran 2-3 times slower than the C++ implementation. Thus, we believe Pregel+’s C++ implementation also contributes to its superior overall performance.

5.1.2 Performance on Different Graphs

Next we analyze the performance of the various systems on graphs with different characteristics.

Performance on graphs with skewed degree distribution. Figures 4(a), 4(b), 4(c), 4(d), 4(e), 4(g), 4(h), 4(j), 4(m) and 4(p) report the performance of the systems on graphs with skewed degree distribution (i.e., WebUK, L1, BTC, and Twitter). The results show that Pregel+ (thanks to its mirroring and request-respond techniques) has the best performance in most cases, while GPS (with the help of LALP) also has good performance in most of the cases. GraphLab is faster than Giraph in about half of the cases but is slower in the rest. Overall, no system is always better than the others in processing graphs with skewed degree distribution, but Pregel+ and GPS are the better choices as they exhibit good performance in most of the cases tested.

Performance on graphs with a large diameter. Figures 4(a), 4(d), 4(f), 4(h), 4(k) and 4(m) report the performance of the systems on graphs with a large diameter (e.g., WebUK and USA Road). Again, each system has better performance in some cases, but overall Pregel+ has the best performance in more cases. GPS and GraphLab beat each other in roughly equal number of cases, while Giraph has the worst performance in most cases.

Performance on graphs with a small diameter. Figures 4(b), 4(c), 4(e), 4(g), 4(i), 4(j), 4(l), 4(o) and 4(p) report the performance of the systems on graphs with a small diameter, e.g., Friendster, Twitter and BTC. Pregel+ has the best performance in most cases, while GPS also has good performance in most cases. Giraph has poorer performance than GPS, but is better than GraphLab overall.

Performance on graphs with high average degree. Figures 4(a), 4(b), 4(d), 4(h), 4(i), 4(j), 4(l) and 4(o) report the performance of the systems on graphs with a relatively high average degree (e.g., WebUK, Friendster and Twitter). Pregel+ has the best performance in most cases. GPS is generally faster than GraphLab, while both of them are faster than Giraph in most cases.

Overall performance. Figure 8 gives a ranking on the overall performance of the systems for different types of graphs. Pregel+ has the best performance in most cases mainly because of its integration of mirroring and combiner, which effectively addresses load balancing in skewed-degree graphs and reduces messages in graphs with high average degree. Similarly, GPS’s LALP is also effective in addressing load balancing and in message reduction. GraphLab’s vertex-cut partitioning effectively addresses load balancing, but it has an overhead of locking/unlocking (which is required even in synchronous mode, e.g., to prevent more than one vertices from scattering values to the same vertex), and hence its overall performance is not much better than Giraph. For handling large-diameter graphs which usually require a large number of supersteps, our results reveal that systems like Giraph, which have a large constant overhead (hundreds of ms) per superstep, can be very slow. On the contrary, Pregel+ has a very small constant overhead per superstep, while GPS and GraphLab have a slightly larger constant overhead per superstep than that of Pregel+. Finally, small diameter usually leads to faster convergence and hence for systems like GraphLab, which has a larger start-up overhead in vertex-cut partitioning, can be slower than other systems.

5.1.3 Comparison with GraphChi

We also compare with a single-machine baseline, GraphChi [12]. Due to space limitation, we only report the performance of Pregel+ and GraphChi in Figures 9(a)–9(e). Performance of other systems can be compared by referring to the performance of Pregel+
and other systems in Figures 4(a)–4(p) (and Figures 3–9 in [14]), GraphChi needs to pre-sort the graph, the cost of which is reported in Figure 16 in [14].

We do not run SV and GC since it is not clear to us how pointer jumping in SV and edge deletion in GC can be implemented in GraphChi. We also note that GraphChi took much longer to run SSSP on WebUK and we killed the job after its running time is three orders of magnitude longer than Pregel+’s.

The results show that Pregel+ is about 10 times faster than GraphChi when processing the large graphs, WebUK, Friend, Twitter and BTC. But for the two smaller graphs, LJ and USA, which can fit in the memory of a single machine, GraphChi uses fully main-memory mode and its running time is closer to that of Pregel+ (and even faster in two cases on USA). Thus, GraphChi is a reasonable choice for moderate-sized graphs. But when the graph is large and sufficient computing resources are available, a distributed system can achieve much better performance than a single-machine system. To be more specific, Pregel+ requires 4, 3, 2 and 2 machines to process WebUK, Friend, Twitter and BTC in memory, respectively; and given such number of machines, Pregel+ is already much faster than GraphChi (see details in Section 5.1.3 in [14]). We note that Java-based systems such as Giraph and GPS may require more machines as they use more memory.

5.2 Effects of Message Combiner

We now study the effects of message combiner. GPS does not perform sender-side message combining, as the authors claim that very small performance difference can be observed whether combiner is used or not [19]. To verify whether this claim is valid, we first analyze how many messages can be combined by applying a message combiner as follows (the proof is detailed in [14]).

**THEOREM 1.** Given a graph \( G = (V, E) \) with \( n = |V| \) vertices and \( m = |E| \) edges, we assume that the vertex set is evenly partitioned among \( M \) machines (i.e., each machine holds \( n/M \) vertices). We further assume that the neighbors of a vertex in \( G \) are randomly chosen among \( V \), and the average degree \( \deg_{avg} = m/n \) is a constant. Then, at least \( \left( 1 - \exp\left( -\deg_{avg}/M \right) \right) \) fraction of messages can be combined using a combiner in expectation.

According to Theorem 1, if a large number of machines are available and the average degree is small, then indeed applying combiner may not improve the performance much as claimed in [19]. For example, if \( M = 1000 \) and \( \deg_{avg} = 10 \), then only 1% of the messages can be combined. However, in many applications and for many datasets (e.g., for all the algorithms we discuss in this paper and graphs with more than 5 billions of edges we used here), one may not require or use thousands of machines. When \( M \) is smaller, combiners can effectively reduce the number of messages to be sent over the network and hence improve the performance of the systems, which we verify as follows.

We assess the effect of combiner by testing the two systems, Giraph and Pregel+, that support combiner. We use two versions for each system, Giraph vs Giraph− and Pregel+ vs Pregel+−, where the superscript ‘−’ indicates that combiner is not applied. As shown in Figures 10(b) and 10(d), there is an obvious reduction on the total number of messages sent over the network when combiner is applied. As the number of machines increases, less messages are combined but the number is still considerably smaller than that without combiner.

Figures 10(a) and 10(c) further show that the running time of both systems with combiner is shorter than that without combiner. In conclusion, applying combiner can always reduce the total number of messages and shorten the running time. Although the improvement is not so obvious in some cases, there also exist cases where the improvement is quite significant, e.g., running PageRank in Pregel+− on WebUK. This conclusion has also been verified on many other algorithms on the datasets we used, and we have not found a case where applying combiner leads to worse performance than without combiner.

5.3 Effects of LALP and Mirroring

We now study the effects of LALP in GPS and mirroring in Pregel+. We report the performance of GPS and Pregel+, with and without LALP/mirroring, for running Diameter Estimation on LJ in Figure 11 and for running HashMin on BTC in Figure 12. The results show that applying LALP/mirroring reduces the running time in both cases. For running Diameter Estimation on LJ, using LALP in GPS is 1.3 times faster and using mirroring in Pregel+ is 2.2 times faster than without using the techniques. For running HashMin on BTC, the reduction in running time is not as significant, because the number of supersteps that involve a large number of redundant messages is only 4, and so message reduction is signifi-
the entire process, and therefore, the performances degrades due to
14(d). In all cases, DP does not obtain a good graph partition in
HashMin on BTC, and BMM on LiveJournal in Figures 14(a)–
implied in all algorithms and on all graph types. We report the per-
distribute vertices across workers. The DP technique can be ap-

5.5 Effects of Dynamic Repartitioning

request-respond technique.

significantly smaller than in Diameter Estimation where many supersteps
involves a large number of redundant messages. Figures 22 and 23
in [14] further show the skewed distribution in the number of mes-
sents sent by the workers of GPS/Pregel+ is evened by applying
LALP/mirroring. In addition, there is also a significant reduction in
the number of messages sent by each worker.

5.4 Effects of Request-Respond API

We next study the effects of the request-respond technique in
Pregel+, which can address the imbalanced workload created by
algorithm logic such as in the SV algorithm. We test SV on the
USA road network and the BTC graph. Figure 13(a) shows that the
running time of Pregel+ is almost reduced by half after applying the
request-respond technique, which can be explained by the signifi-
cant reduction in the number of messages as shown in Figure 13(b).
Figure 24 in [14] further shows that the imbalanced communication
workload caused by the logic of SV is effectively eliminated by the
request-respond technique.

5.5 Effects of Dynamic Repartitioning

GPS also adopts a dynamic repartitioning (DP) technique to re-

distribute vertices across workers. The DP technique can be ap-
plied in all algorithms and on all graph types. We report the per-
formances of PageRank on Twitter, Diameter Estimation on BTC,
HashMin on BTC, and BMM on LiveJournal in Figures 14(a)–
14(d). In all cases, DP does not obtain a good graph partition in
the entire process, and therefore, the performances degrades due to

5.6 Effects of Algorithmic Optimizations

We next study the effects of the algorithmic optimizations de-
scribed in Section 4.2. Figures 15(a)–15(c) report the effects of ap-
plying FCS and ECOD in Giraph, GPS and Pregel+ (more results
can be found in Section 5.6 in [14]). Note that FCS and ECOD
cannot be applied in GraphLab since it does not support edge dele-
tion and Pregel-like aggregator. The results show that FCS con-
siderably improves the performance of Graph Coloring (GC) in all
the systems, since GC takes a large number of supersteps to con-
verge on Friendster and BTC, and hence FCS can significantly re-
duce the number of supersteps. Figure 15(c) shows that FCS also
improves the performance of the systems on BMM. On the other
hand, the results show that ECOD degrades the performance. This
is mainly because GC removes every stale edge in later computa-
tion and hence ECOD does not reduce the total workload. More-
ever, ECOD incurs additional overhead. Thus, ECOD is only ef-
fective in algorithms in which many stale edges will not be touched
again after they are decided to be removed [20].

Figures 15(d)–15(e) (more results can be found in Section 5.6
in [14]) report the effects of applying SP in different systems for
computing connected components (CC), where we use HashMin to
compute CCs in the remaining graph after BFS from the pivot. The
performance of the systems on Friendster is significantly improved,
but degrades on BTC. The reason is that, all vertices in Friendster
constitutes a single giant CC, and so no matter which vertex is cho-

en as the pivot, the computation will terminate after running BFS
from the pivot. In other words, less costly BFS is ran on the whole
graph instead of HashMin. However, the largest component of BTC
consists of only around 3% of all the vertices. Thus, SP can only
label a small fraction of the graph and the overhead exceeds the
gain obtained by SP.

5.7 Scalability of Various Systems

We evaluate the scalability of the systems on both real-world
graphs and synthetic graphs.

5.7.1 Effects of Number of Machines/CPU-Cores

We first report the performance of the systems by varying the
number of machines or CPU cores.

Effects of machine number. We vary the number of machines
from 6 to 15 in this experiment, and fix the number of CPU cores
in each machine to 8. GPS and Pregel+ run 8 processes on each
machine, while Giraph and GraphLab can take advantages of all
the computing resource from the 8 cores by multithreading.

We first consider PageRank on WebUK, Figure 16(a) shows that
only Giraph scales linearly with the number of machines, though it
is significantly slower than the other systems. Pregel+ scales almost
linearly (note that Figures 16(a)–16(d) are in logarithmic scale) and
it is the fastest system in all settings. For GraphLab, we can only
obtain the results when there are at least 12 machines in the clus-
ter, as the total aggregate memory of the cluster is not sufficient for
running GraphLab on this large web graph when there are less than
12 machines. The situation is similar for GPS, but for the smaller
BTC graph, we obtain their results for all cases as reported in Figure
16(b). For processing the BTC graph, Giraph, GraphLab, and
Pregel+ all scale linearly with the number of machines, but GPS’s
running time does not change much as the number of machines increases.

**Effects of CPU core number.** We vary the number of CPU cores in each machine from 4 to 16 in this experiment, and fix the number of machines in the cluster to 15. The number of processes in GPS and Pregel+ is the same as the number of CPU cores.

For running PageRank on WebUK, Figure 16(c) shows that only GraphLab scales sub-linearly with the number of CPU cores. This is because GraphLab can take advantage of multithreading. Giraph also uses multithreading but the effect is not obvious. The running time of GPS and Pregel+ decreases considerably (about 1.5 times) when the number of processes in each machine increases from 4 to 8, but further increasing the number of processes does not improve the performance since the overhead of network communication also
increases with the number of processes.

For processing BTC, Figure 16(d) shows that multithreading in Giraph and GraphLab becomes even less effective since the dataset is much smaller. Pregel+ scales only linearly when the number of processes in each machine doubles from 4 to 8, while the performance of GPS even degrades when the number of processes in each machine increases.

5.7.2 Effects of Graph Size

We now report the performance of the systems by varying the number of vertices and edges using synthetically generated random graphs [16] and power-law graphs [3]. We set the number of machines in the cluster to be 15 and the number of CPU cores or processes in each machine to be 8.

Effects of vertex number. We vary the number of vertices in the synthetic graphs from 50M to 400M, while we fix the average vertex degree of each graph to 20. As Figures 17(a)–17(d) show, the running time of all the systems increases approximately linearly with the number of vertices. However, GPS ran out of memory when running PageRank on the two largest graphs, while GraphLab ran out of memory when running both PageRank and HashMin on the two largest graphs. Giraph and Pregel+ can run on all graphs, but Giraph has poorer scalability than Pregel+ when the graph size becomes larger.

Effects of edge number. We fix the number of vertices in each graph to 100M, and vary the average vertex degree in the synthetic graphs from 10 to 80 (i.e., the number of edges changes from 1 billion to 8 billion). For this set of experiments, Figures 18(a)–18(d) show that the running time of Giraph, GPS, and Pregel+ increases sub-linearly with the number of edges, which indicates that the systems have good scalability (except for GPS which ran out of memory for running PageRank on the two largest graphs). GraphLab has the best scalability among all systems for running PageRank, but it exhibits the worse scalability for running HashMin.

6. CONCLUSIONS

We evaluated the performance of Giraph [2], GraphLab [13, 7], GPS [19], and Pregel+ [26], w.r.t. various graph characteristics, algorithm categories, optimization techniques, and system scalability. Our results show that, while there is no single system that has superior performance in all cases, Pregel+ and GPS have better overall performance than Giraph and GraphLab. Pregel+ has better performance mainly thanks to its combination of mirroring, message combining, and request-respond techniques, while GPS also benefits significantly from its LALP technique. GraphLab uses vertex-cut partitioning for load balancing, but it incurs extra overhead in locking/unlocking, which may degrade its performance. Giraph generally has the poorer performance because it does not employ any specific technique for handling skewed workload and mainly relies on combiner for message reduction. Finally, we believe that the efficiency of Pregel+ also comes from its C++ implementation, which at least uses less memory, and according to [12] can be 2-3 times faster, than a Java implementation.

Acknowledgments. We thank the reviewers for giving us many constructive comments, with which we have significantly improved our paper. We thank our teammates, Xiaotong Sun and Jishi Zhou, for their help with some of the experiments. This research is supported by SHIAE Grant No. 8115048.

7. REFERENCES


