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<td>Author(s)</td>
<td>Guo, Tao; Cao, Xin; Cong, Gao; Lu, Jiaheng; Lin, Xuemin</td>
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Distributed Algorithms on Exact Personalized PageRank

Tao Guo  
Nanyang Technological University  
Singapore  
tguo001@e.ntu.edu.sg

Xin Cao  
The University of New South Wales  
Australia  
xin.cao@unsw.edu.au

Gao Cong  
Nanyang Technological University  
Singapore  
gaocong@ntu.edu.sg

Jiaheng Lu  
University of Helsinki  
Finland  
jiahenglu@gmail.com

Xuemin Lin  
The University of New South Wales  
Australia  
lxue@cse.unsw.edu.au

ABSTRACT
As one of the most well known graph computation problems, Personalized PageRank is an effective approach for computing the similarity score between two nodes, and it has been widely used in various applications, such as link prediction and recommendation. Due to the high computational cost and space cost of computing the exact Personalized PageRank Vector (PPV), most existing studies compute PPV approximately. In this paper, we propose novel and efficient distributed algorithms that compute PPV exactly based on graph partitioning on a general coordinator-based share-nothing distributed computing platform. Our algorithms take three aspects into account: the load balance, the communication cost, and the computation cost of each machine. The proposed algorithms only require one time of communication between each machine and the coordinator at query time. The communication cost is bounded, and the work load on each machine is balanced. Comprehensive experiments conducted on five real datasets demonstrate the efficiency and the scalability of our proposed methods.

CCS Concepts
• Mathematics of computing → Graph algorithms; • Theory of computation → Parallel computing models;

Keywords
Personalized PageRank; random walks

1. INTRODUCTION
Measuring the similarities between nodes is a fundamental graph computation problem. Many random surfer model based measures have been proposed to capture the node-to-node proximities \[2,9,22,23,28\]. The Personalized PageRank (PPR) \[23\] is one of the most widely used measures and has gained extensive attention because of its effectiveness and theoretical properties. It has been utilized in various fields of applications, such as web search \[8,11\], community detection \[3,20\], link prediction \[4\], anomaly detection \[39\], and recommendation \[25,45\].

Different from the PageRank model \[35\], PPR allows users to specify a set of preference nodes \(\mathcal{P}\). The result of PPR w.r.t. \(\mathcal{P}\) is called the Personalized PageRank Vector (PPV), which is denoted by \(r_{\mathcal{P}}\). Note that different preference vectors will yield different PPVs, and thus PPV needs to be computed in an online manner, which is different from PageRank. The PPR model can be simulated by numerous “random surfers.” Initially, these surfers are distributed to the nodes in \(\mathcal{P}\) equally. Next, in each step, a random surfer either jumps to a random outgoing neighbor with a probability of \(1 - \alpha\), or teleports back to a node \(u\) in \(\mathcal{P}\) according to the specified preference of \(u\) with a probability of \(\alpha\) (\(\alpha\) is called the teleport probability). The procedure is performed repeatedly until it converges to a steady state, i.e., the number of surfers on each node does not change after each iteration. The final distribution of the random surfers on the nodes represents the PPV of \(\mathcal{P}\).

The PPR model can also be interpreted as a linear system. We denote \(\mathbf{u}_{\mathcal{P}}\) as the preference vector w.r.t. \(\mathcal{P}\) and \(\mathbf{A}\) as the normalized adjacency matrix of the graph. PPV \(r_{\mathcal{P}}\) can be computed as:

\[
r_{\mathcal{P}} = (1 - \alpha)A^T\mathbf{r}_{\mathcal{P}} + \alpha\mathbf{u}_{\mathcal{P}}. \tag{1}
\]

Hence, \(r_{\mathcal{P}}\) can be computed using the power iteration method, i.e., \(r_{\mathcal{P}}^{k+1} = (1 - \alpha)A^T r_{\mathcal{P}}^{k} + \alpha\mathbf{u}_{\mathcal{P}}\), where \(r_{\mathcal{P}}^{0}\) represents the vector computed in the \(k^{th}\) iteration, and \(r_{\mathcal{P}}^{0}\) is set to a uniform distribution vector. Finally, \(r_{\mathcal{P}}\) converges to the PPV of \(\mathcal{P}\).

A. Challenges. Although computing PPVs has been extensively studied since the problem is first proposed in the work \[23\], it is still a very challenging task to compute exact PPVs efficiently for different online applications. One straightforward method is to adopt the power iteration approach by following Equation \(1\) which is however computationally expensive, and it is not practical for the online applications of PPR.

In the work \[23\], the linearity property of PPV is studied for computing PPVs exactly. If we pre-compute the PPV for each node, at query time, given a node set \(\mathcal{P}\), according to the property, the PPV of \(\mathcal{P}\) can be constructed using the pre-computed PPVs of the nodes in \(\mathcal{P}\). Unfortunately, this method is impractical because of both the expensive pre-computation time and the offline storage requirements (where \(O(|\mathcal{V}|^2)\) space is needed). In order to reduce the complexity and space cost, the work \[23\] selects some important nodes as the “hub nodes”, and the user preference nodes.
can only be from the hub set, thus losing the generality. Due to the hardness of computing the exact PPVs, most of existing studies (e.g., [5, 14, 40, 46]) focus on computing approximate PPVs. These methods sacrifice the accuracy to accelerate PPV computation.

Facing the challenges of computing the exact PPV for any arbitrary user preference node set, a natural question raised is how to perform the computation in parallel on multiple machines to gain efficiency. However, it is known that graph algorithms often exhibit poor locality and incur expensive network communication cost [22]. Therefore, it is a challenging task to design an efficient and scalable distributed algorithm for computing PPVs that has low communication cost and is able to achieve load balance.

The power iteration method can be implemented on the general distributed graph processing platforms to compute PPVs in a distributed way. However, this would not be suitable because of the unavoidable high communication cost. In the power iteration method, computing the vector in the current iteration requires the vector of the previous iteration. Thus, no matter how we distribute the computation, one machine always has to ask for some data from the other machines in each iteration before convergence, thus incurring expensive network communication cost. For example, the graph processing engine Pregel [34] is based on the general bulk synchronous parallel (BSP) model [41]. In each iteration of the BSP execution, Pregel applies a user-defined function on each vertex in parallel. The communications between vertices are performed with message passing interfaces. The block-centric system Blogel [43] distributes subgraphs to machines as blocks, and messages between blocks are transmitted over the network. Both Pregel and Blogel need many rounds of communications to compute PPVs, which incurs expensive communication cost, thus suffering from low efficiency.

B. Our Proposal. In this work, we design novel distributed algorithms utilizing the graph partitioning for computing exact PPVs, which can be implemented on a general coordinator-based share-nothing distributed computing platform. We take into account three aspects in designing our algorithms: the load balance, the communication cost, and the computation cost on each machine. A salient feature of the proposed algorithms is that each machine only needs to communicate with the coordinator machine once at query time, and we prove that the communication cost of our method is bounded. The main idea of our algorithms is outlined as below.

Observation 1: We can separate the graph into disjoint subgraphs of similar sizes to distribute the PPV computation.

Based on this observation, we propose the graph partition based algorithm, denoted by GPA. We prove that the benefit from the balanced disjoint graph partitioning is two-fold: first, this guarantees a total space cost of $O((|V| - |H|)^2 + 2|V| |H| + |H|^2)$, where $V$ represents the nodes of the graph, $m$ is the number of subgraphs, and $H$ is the set of hub nodes separating the subgraphs. Note that $|H|$ is much smaller than $|V|$ (see more analysis in Appendix E), and thus this space cost is significantly smaller than applying the method [23] directly. Second, this enables us to compute PPVs in parallel on separate machines without incurring communication cost between machines. At query time, based on the pre-computed values, each machine constructs part of the PPV, and only communicates with the coordinator once (i.e., sending part of the PPV to the coordinator). If we use $n$ machines, the communication cost of GPA is $O(n|V|)$.

Observation 2: We notice that if we treat each subgraph as an individual graph, we can use GPA to compute the “local” PPV of each node w.r.t. the subgraph itself, and these local PPVs can be used to construct the final “global” PPV. This motivates us to further partition each subgraph and we get a tree-like graph hierarchy. Then, we can recursively apply GPA along the hierarchy of subgraphs. We prove that the pre-computation space cost can be further reduced and bounded by utilizing the graph hierarchy.

Observation 3: Since the sizes of subgraphs in different levels of the hierarchy are different, simply distributing the subgraphs to machines cannot achieve load balance. We design a method to distribute the PPV computation evenly based on hub nodes partitioning to solve this problem.

Based on the aforementioned observations, we propose the hierarchical graph partition based algorithm, denoted by HGPA, which greatly reduces the space cost, and we prove that the communication cost of HGPA is $O(n|V|)$ as well.

C. Contributions. To the best of our knowledge, this is the first work that is able to compute exact PPVs efficiently in a distributed manner with reasonable space cost. In this paper, we propose novel distributed algorithms based on graph partitioning, and the salient features of new algorithms can be summarized as follows:

- **Efficiency**: Our proposed algorithm HGPA is 10 ~ 100 times faster than the power iteration approach running on general distributed graph processing platforms Pregel [44] and Blogel [43], and can meet the efficiency need of online applications. The experimental study also shows that HGPA is faster than the power iteration and one state-of-the-art approximate PPV computation algorithm [46] even under the centralized setting.

- **Accuracy**: Our proposed algorithms, namely GPA and HGPA, are able to obtain the same result as that of the work [23], and thus have guaranteed exactness (Theorems 1 and 3).

- **Load Balance**: Our distributed algorithms GPA and HGPA are load balanced and HGPA scales very well with both the size of datasets and the number of machines. As shown in the experiments, the runtime can be reduced nearly by half if we double the number of machines.

- **Low Communication Cost**: Our algorithms only require one time of communication between each machine and the coordinator, and no communication between any two machines is needed for pre-computation and query processing. As shown in the experiments, our method only costs about 1.5 MB network communication to compute a PPV on a graph containing 3M nodes using 10 machines.

The rest of the paper is organized as follows. Section 2 provides the preliminary of this work. Sections 3 and 4 present the distributed algorithms GPA and HGPA, respectively. Section 5 explains our distributed pre-computation. We perform comprehensive experiments to evaluate the efficiency and scalability of our methods on 5 real datasets, and the results are reported in Section 6. Section 7 summarizes the related work on computing PPVs. Finally, Section 8 offers conclusions and future research directions.

2. PRELIMINARY

2.1 PPV Decomposition

As proved by Jeh and Widom [23], the PPV score $r_u(v)$ is equal to the corresponding Inverse P-distance from a query node $u$ to $v$, which is measured by all the weighted tours from $u$ to $v$:

$$r_u(v) = \sum_{t : u \leadsto v} p(t),$$

(2)

where each tour $t : u \leadsto v$ represents a random surfer that consists of a sequence of edges starting from $u$ and ending at $v$. Note that it is allowed to teleport back to $u$, and thus there may exist cycles in
Table 1: Definition of main symbols.

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<th>Symbol</th>
<th>Definition</th>
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<tr>
<td>$P$</td>
<td>User preference nodes set</td>
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<tr>
<td>$r_u$</td>
<td>PPV of a single preference node $u$</td>
</tr>
<tr>
<td>$\alpha$</td>
<td>Teleport(Restart) probability, $0 &lt; \alpha &lt; 1$</td>
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<tr>
<td>$H$</td>
<td>Hub node set</td>
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$t : u \sim v$ A random tour from node $u$ to $v$

$P(t)$ Weight of tour $t$

$L(t)$ Length of tour $t$

$r^H(u)$ PPV of tours from $u$ passing at least one node in $H$

$s^H_{\alpha}(v, u)$ (Adjusted) Partial vector w.r.t. node $u$ and hub nodes $H$

$x_u$ Basic vector with zero filled except $x_u(u) = 1$

$G^{h}_{m}$ A hierarchical partitioned subgraph in the $m^{th}$ level

$H(G^{h}_{m})$ Hub nodes of subgraph $G^{h}_{m}$

the tour. The weight of a tour $P(t)$ is the probability that this tour walks from $u$ to $v$ along $t$:

$$P(t) = \alpha(1 - \alpha)^L(t) \prod_{i=1}^{L(t)} \frac{1}{|Out(w_i)|},$$

where nodes $w_1, w_2, \cdots, w_L$ (i.e., $v$) comprise the path of length $L(t)$, and $|Out(w_i)|$ is the outdegree of node $w_i$. Although the concept of inverse P-distance intuitively explains the distribution of random walk, it is impossible to sum up all the tours to obtain the PPV values. The reason is that there may exist loops in the tours, and thus the number of tours could be infinite.

In order to compute $r_u$, it is divided into two types of vectors w.r.t. a set of hub nodes $H$, i.e., partial vectors and hub skeleton vectors.

**Definition 1: Partial vector:** Given a node $u$, its partial vector $p^H_u$ is defined as a vector of random walk results computed using tours passing through no hub nodes. That means, given a node $v$,

$$p^H_u(v) = \sum_{u \in H \land w_u \sim v} P(t).$$

**Definition 2: Hubs skeleton vector:** Given a node $u$, its hubs skeleton vector $s^H_u$ is defined as a vector of $|H|$ dimensions composed of all hub nodes’ PPV values w.r.t. $u$. Given a hub node $h$, $s^H_u(h) = r_u(h)$.

Figure 1: Random Surfer Example

If a tour passes no hub node, it contributes to the partial vector. If a tour stops at a hub node, it contributes to the skeleton vector. For ease of understanding, we explain the meaning of the two types of vectors with the graph shown in Figure 1 where two nodes $u_2$ and $u_3$ are selected as hub nodes. We next illustrate the partial and skeleton vectors with node $u_3$ as an example.

The partial vector $p^H_{u_3}$ consists of two tours $t_1$ and $t_2$, where $t_1 = u_3 \sim u_4 \sim u_2$ and $t_2 = u_3 \sim u_4 \sim u_5 \sim u_3$. Note that all other tours are blocked by either $u_2$ or $u_3$ (hub nodes), and thus any gray node in this example is not reachable from $u_1$. We have $p^H_{u_3}(u_4) = P(t_1)$ and $p^H_{u_3}(u_2) = P(t_2)$.

For the skeleton vector $s^H_{u_3}$, we only consider the tours stopping at a hub node. Different from the partial vector, the tour can contain one or more hub nodes. From $u_1$ to hub node $u_2$, there exist two tours $t_3 = u_1 \sim u_3$ and $t_4 = u_1 \sim u_3 \sim u_5 \sim u_2$. From $u_1$ to $u_3$ there exist three tours $t_5 = u_1 \sim u_3 \sim u_2$, $t_6 = u_1 \sim u_4 \sim u_3$, and $t_7 = u_1 \sim u_4 \sim u_5 \sim u_2 \sim u_3$. Therefore, $s^H_{u_3}(u_2) = P(t_3) + P(t_4)$ and $s^H_{u_3}(u_3) = P(t_5) + P(t_6) + P(t_7)$.

Note that because of cycles in the tours, it is not feasible to compute all possible tours, which may be infinite. This simple example is only used to illustrate the intuitive meaning of the two types of vectors.

2.2 PPV Construction

The PPV of $u$ (i.e., $r_u$) can be constructed by the partial vectors and hubs skeleton vectors. We use $r^H_u$ to denote the vector of random walk results computed using tours passing through at least one hub node. That is, give a node $v$, $r^H_u(v) = \sum_{u \rightarrow H \rightarrow v} P(t)$. We can combine $r^H_u$ and the partial vector of $u$ to obtain the full PPV of $u$: $r^H_u + p^H_u = r_u$. As proved by Jeh and Widom [23], $r^H_u$ can be computed according to the following equation:

$$r^H_u = \frac{1}{\alpha} \sum_{h \in H} (s^H_u(h) - \alpha f_u(h)) \cdot (p^H_h - \alpha x_h),$$

where $f_u(h) = 1$ if $u = h$, and 0 otherwise. $f_u(h)$ is used to deal with the special case when $u$ is a hub node. $x_u$ is a vector that has value 1 at $h$ and 0 everywhere else. It is used to deal with the special case when $h$ is the target node.

In order to make the equation more clear, we define the adjusted partial vector for each hub node $h$ by:

$$D^H_h = p^H_h - \alpha x_h,$$

and the adjusted hubs skeleton vector $S^H_h$ for each node $u$ by:

$$S^H_u(h) = s^H_u(h) - \alpha f_u(h).$$

After the partial vectors and skeleton vectors have been pre-computed, the exact PPV of a given query node $u$ can be constructed by:

$$r_u = \frac{1}{\alpha} \sum_{h \in H} S^H_u(h) \cdot D^H_h + p^H_u$$

In order to reduce the complexity, the work [23] only considers the construction of PPV for hub nodes. According to Equation 4, we need to pre-compute the partial vectors of all hub nodes, which requires space cost $O((|V| - |H|) \cdot |H|)$ in the worst case, and the skeleton vectors of all hub nodes, which requires $O(|H|^2)$ space.

2.3 A Brute-force Extension

It is too restricted to consider the hub set from only preference nodes as done in work [23]. In fact, it can be extended to compute the PPV for any given query node, as indicated by Equation E. However, it incurs huge space cost as explained below: first, pre-computing the partial vectors for non-hub nodes requires worst case space cost $O((|V| - |H|)^2)$, which happens when every node can reach every other node without passing any hub node; second, pre-computing the partial vectors for hub nodes requires $O((|V| - |H|)|H|)$ space in the worst case; third, pre-computing the skeleton vectors for all nodes requires space cost $O(|V| \cdot |H|)$. Thus, the total pre-computation space cost is $O(|V|^2)$, which is equivalent to pre-computing the PPVs of all nodes. In practice, the vectors may be sparse and the space cost is usually not that large, but it is still not applicable for large graphs. We denote this algorithm by PPV-JW.
3. ALGORITHM GPA

Although it is usually difficult to efficiently parallelize graph algorithms, we propose the graph partition based algorithm, denoted by GPA, to distribute the PPV computation. We present the algorithm details in Section 3.1. We adopt a general coordinator-based share-nothing distributed computing platform. For the convenience of presentation, we call each machine handling some subgraphs as “machine” and the machine aggregating the final result as “coordinator.” In addition, in Section 3.2, we prove that GPA reduces the space cost significantly compared with the method PPV-JW presented in Section 2.3.

3.1 Distributed PPV Construction

Graph partition is commonly used in parallel computing, but it is always challenging to decouple the computation dependencies. We aim to distribute the PPV computation to multiple machines and each machine can independently compute part of the result. We use a balanced graph partition algorithm (METIS [24]) to divide the graph into m disjoint subgraphs. Figure 2 shows an example, where the graph is partitioned into two disjoint subgraphs \( G_1 \) and \( G_2 \). The bridging nodes between subgraphs form the hub nodes. In the example, \( u_1 \) and \( u_2 \) are selected as the hub nodes.

After the graph is partitioned into \( m \) subgraphs, we distribute the subgraphs to multiple machines evenly. The pre-computed partial vector and skeleton vector of each node is stored in the machine where the node resides.

![Figure 2: Example of graph partition and hub nodes](image)

Recall that the PPV construction of the method [23] is based on Equation 4. Now the pre-computed vectors are stored on \( n \) machines \( M_1, ..., M_n \), and we can distribute the computation according to the following equation:

\[
r_u = \frac{1}{n} \sum_{i=1}^{n} \sum_{h \in H(M_i)} S^H_u(h) \cdot P^H_u + P^H_u,
\]

where \( H(M_i) \) denotes the set of hub nodes assigned to \( M_i \).

Assume that the partial and skeleton vectors have already been pre-computed and stored. We introduce the details of this step in Section 5. Equation 5 indicates that we can do the distributed PPV construction at query time as follows: after a query node \( u \) is given, the coordinator first detects the machine \( M_i \) that stores the partial vector of \( u \). Then, \( M_i \) computes the following vector:

\[
v_u = \frac{1}{n} \sum_{h \in H(M_i)} S^H_u(h) \cdot P^H_u + P^H_u.
\]

Simultaneously, each of the other machines \( M_j \) (\( 1 \leq j \leq n, j \neq i \)) computes a vector \( v_j = \frac{1}{n} \sum_{h \in H(M_j)} S^H_u(h) \cdot P^H_u \). The coordinator receives the vectors computed from all machines, and computes the final PPV as \( r_u = \sum_{i=1}^{n} v_i \). Therefore, at query time, each machine communicates with the coordinator exactly once.

**Theorem 1:** GPA can obtain the same results as computed by the work proposed by Jeh and Widom [23].

**Proof:** \( \sum_{i=1}^{n} \sum_{h \in H(M_i)} = \sum_{h \in H} \) and thus Equation 5 can compute the same vector as Equation 4. \( \square \)

**Communication Cost.** Each machine needs to compute a vector of size \( |V| \), and then sends it to the coordinator. Thus, if \( n \) machines are employed in GPA, the total communication cost of GPA is \( O(n|V|) \).

**Time Complexity.** According to Equation 5 to compute a node \( u \)'s PPV, we need to fetch the partial vector of each hub node \( h \) reachable from node \( u \) (i.e., \( P^H_u \)), the skeleton vector of \( u \) (i.e., \( S^H_u \)), and the partial vector of \( u \) (i.e., \( P^H_u \)). The partial vector of each hub node \( h \) gets its weight \( S^H_u(h) \) from the skeleton vector of \( u \), and they are then aggregated together with \( P^H_u(h) \). Therefore, we need to read at most \( O(|H|) \) vectors of dimension \( |V| \) and \( O(|H|) \) vectors.

Assume that we use \( n \) machines and the hub nodes are distributed evenly to these machines. In the worst case, on each machine we only need to sum up \( O(|H|/n) \) vectors, and on the coordinator we need to sum up \( n \) vectors. Thus, the time complexity of GPA is \( O(\frac{1}{n}|H||V| + n|V|) \).

3.2 Space Cost of GPA

We proceed to show that the space cost of GPA is reduced significantly compared with PPV-JW, the extension of the method [23] as presented in Section 2.3. We use an example as shown in Figure 3 to briefly explain the reason. In the work [23], nodes with high PageRank values are chosen as hub nodes, since most random walks have high probability to visit these nodes. As a result, nodes \( u_1 \) and \( u_3 \) are selected to be the hub nodes. The support of vector \( p^H_u \) (the partial vector of \( u_4 \)) can be as large as the size of the graph, since there exists a tour from \( u_4 \) to each other node in the graph. If we select the nodes \( u_1 \) and \( u_3 \) as the hub nodes, they are able to partition the graph into two disjoint subgraphs. In this way, every tour from node \( u_4 \) to a node in the other part has to pass either \( u_1 \) or \( u_3 \), and thus the tours between different subgraphs are blocked by the hub nodes. The support of \( p^H_u \) is reduced to the size of the subgraph containing \( u_4 \).

As illustrated in the example, after the graph partition, we select the bridging nodes between subgraphs as the hub nodes. The random walks that represent partial vectors are restricted within each individual subgraph by the hub nodes. Thus, the support of the partial vector of non-hub node \( p^H_u \) is reduced to \( |V| - |H| \) to the size (the number of nodes) of the subgraph containing \( u \). In PPV-JW, the space cost of partial vectors of non-hub nodes, i.e., \( O(|V| - |H|)^2 \), is the major cost. In GPA, if we assume that the graph is partitioned into \( m \) subgraphs of equal size, the size of each subgraph is \( O(|V| - |H|)/m) \), and the space cost is \( O(|V| - |H|)^2/m) \). In the worst case, the space cost of storing the partial vectors of the hub nodes is \( O(|V| - |H|)|H|/m) \), and storing the skeleton vectors of non-hub nodes costs \( O(|V||H|) \). In conclusion, the total space cost of GPA is \( O(|V| - |H|)^2/m + 2|V||H| - |H|^2/m) \), based on the balanced graph partition.

Note that for most graphs, the number of hub nodes that can divide the graphs into different components is always much smaller than the total number of nodes, i.e., \( |H| \ll |V| \). Therefore, the space cost of GPA is much smaller than \( O(|V|^2) \), the cost of PPV-JW to compute PPV for an arbitrary node in a centralized setting.

4. ALGORITHM HGPA

In GPA, the computation of partial vectors is restricted to each individual subgraph, and the total space cost of all machines is consequently reduced compared with that required by the centralized extension of the method [23]. To further reduce the space cost and achieve better load balance and efficiency, we propose a new approach based on a hierarchy of subgraphs. This algorithm is inspired by the observation that, the computation of a node \( u \)'s par-
tial vector is equivalent to the computation of the “local” PPV of \( u \) w.r.t. the subgraph that contains \( u \).

We first introduce how we can compute the partial vectors using the way of computing “local” PPVs in Section 4.1. Based on this property, we partition the graph into a hierarchy of subgraphs as presented in Section 4.2. Then, we introduce how to get PPV utilizing the graph hierarchy in Section 4.3 and we design the distributed PPV computation method to achieve load balance in Section 4.4. We prove that the space cost benefits from the hierarchical graph partitioning in Section 4.5.

### 4.1 Partial Vector vs Local PPV

In GPA, we partition the graph into disjoint subgraphs. Recall that as presented in Section 2, PPV can be computed by the random surfers following all possible paths, and the partial vector of a node is the result computed using random surfers passing no hub node. Therefore, the computation of the partial vector of a node is only related to the subgraph containing the node. This motivates us to think whether it is possible to use the local PPV of a node w.r.t. a subgraph to compute the partial vector of this node. Consider the toy graph in Figure 3 which is separated by hub node \( u_2 \). Figure 4 shows the isolated subgraph \( SG \). Suppose the query node is \( u_5 \), we would like to know whether the partial vector \( p_{u_5}^H \) in \( G \) is equal to the local PPV of \( u_5 \) in \( SG \), i.e., \( r_{u_5}[SG] \).

Recall that the computation of \( P(t) \) requires the out-degrees of nodes in the tour. The out-degree of \( u_5 \) is 2 in \( G \) but it is 1 in subgraph \( SG \), and obviously the probability of a random surfer walking from \( u_5 \) to \( u_4 \) is different in the two graphs. Therefore, \( p_{u_5}^H \neq r_{u_5}[SG] \). In order to solve this problem, we introduce the following definition.

**Definition 3: Virtual subgraph:** After partitioning a graph into smaller subgraphs, for each subgraph \( SG \), we create a virtual node \( VN \), and for each edge that connects a hub node and a node \( u \) in \( SG \), we create an edge between \( u \) and \( VN \). We call the graph composed of subgraph \( SG \) and its virtual node as well as the edges connecting them the **virtual subgraph**, which is denoted by \( SG \).

Figure 5 shows the corresponding virtual subgraph of \( SG \) as shown in Figure 3. Using the concept of the virtual subgraph, we have the following theorem.

**Theorem 2:** Given a graph \( G \), a set of hub nodes \( H \), and a node \( u \) in a subgraph \( SG \), the partial vector of \( u \), i.e., \( p_{u}^H \), is equivalent to \( u \)’s PPV vector w.r.t. the virtual subgraph of \( SG \), i.e., \( r_{u}[SG] \).

**Proof:** Given any node \( v \) in \( G \), the value of \( p_{u}^H (v) \) can be computed using the tours from \( u \) to \( v \) without passing through any hub node in \( H \). If \( v \notin SG \), then \( p_{u}^H (v) = 0 \). Otherwise, \( p_{u}^H (v) = \sum_{t:u \leadsto v} P(t) \) where \( t:u \leadsto v \) represents a tour from \( u \) to \( v \) within \( SG \).

According to Equation 2, \( r_{u}[SG](v) \) can also be computed by \( \sum_{t:u \leadsto v} P(t) \). Note that all tours are within \( SG \), the virtual node in \( SG \) has no outgoing edge. Hence, for each tour \( t \), the value of \( P(t) \) is the same for computing both \( p_{u}^H (v) \) and \( r_{u}[SG](v) \), and this means that \( p_{u}^H (v) \) is equal to \( r_{u}[SG](v) \).

Using virtual subgraph \( SG \), we guarantee that the partial vector equals to the local PPV in the virtual subgraph, i.e., \( p_{u}^H = r_{u}[SG] \). For the simplification of presentation, we use “subgraph” to indicate “virtual subgraph” in the rest of the paper.

### 4.2 Hierarchical Graph Partitioning

Based on Theorem 2, we can obtain the partial vector for a node by computing the local PPV in the subgraph containing the node. To compute the local PPV for a subgraph, we can recursively apply GPA within the subgraph, i.e., we further partition the subgraph into lower level subgraphs, and for each lower level subgraph we apply Theorem 2 and the procedure can be repeated until we hit a specified level. To realize the idea, we recursively partition the whole graph from top to down into a hierarchy. For ease of presentation, we partition the graph into a hierarchy of two-way partitions. As shown in Figure 6 the root of the hierarchy is \( G \) itself. Generally, in the \( m \)th level \((0 \leq m \leq l)\) level, \( G \) is partitioned into \( 2^m \) disjoint subgraphs, and we denote a subgraph in the \( m \)th level by \( G_m \), where \( 0 \leq i < 2^m \). In this hierarchy, given a subgraph \( G_m \), its parent subgraph is \( G_{m+1} \), and its two child subgraphs are \( G_{2i+1} \) and \( G_{2i+3} \). We denote the hub nodes separating \( G_{2i+1} \) and \( G_{2i+3} \) by \( H(G_m) \).

![Figure 6: Hierarchy of the original graph](image_url)
4.3 PPV Construction on Hierarchy

We proceed to explain the procedure of PPV computation based on the hierarchy. Consider the hub nodes set $H_0$ in the first level of the hierarchy, which separates $G$ into two subgraphs $G^1_1$ and $G^1_2$. Given a query node $u \in G^1_1$, according to Equation 4 to obtain $r_u$ in GPA, we need the partial vector of node $u$, i.e., $p^{h_0}_u$, the partial vectors of hub nodes, i.e., $p^{h_0}_h (h \in H_0)$, and the skeleton vectors $s^{h_0}_u$.

Here, $p^{h_0}_u$ and $s^{h_0}_u$ can be computed similarly as in GPA. According to Theorem 2, the partial vector of $u$ ($p^{h_0}_u$) is identical to $u$’s PPV vector w.r.t. the virtual subgraph of $G^1_1$. Thus, we can compute the local PPV of $u$ w.r.t. $G^1_1$ as $p^{h_0}_u$. That is, we construct $p^{h_0}_u$ by using the local skeleton vector of lower level hub nodes in $G^1_1$ and the local partial vector of $u$ in $G^1_1$, which can be further computed based on lower level subgraphs, and thus the procedure can be repeated until we reach the leaf-level subgraphs.

Assume that all partial and skeleton vectors have been processed and stored (see more details in Section 5). Formally, the construction at query time can be described by the following equation:

$$r_u = \sum_{m=0}^{l-1} \frac{1}{\alpha} \sum_{h \in H(G^m_u)} S^H_{G^m_u}[G^{(u)}_m](h) \cdot P^H_{G^m_u}[G^{(u)}_m] + r_u[G^{(u)}_0],$$

where $G^{(u)}_m$ denote the subgraph in the $m^{th}$ level that contains $u$, $S^H_{G^m_u}[G^{(u)}_m]$ is the adjusted skeleton vector of $u$ w.r.t. the subgraph $G^{(u)}_m$, and $P^H_{G^m_u}[G^{(u)}_m]$ is the adjusted partial vector of a hub node w.r.t. $G^{(u)}_m$. Note that the subgraphs we visit are $G^{(u)}_1, G^{(u)}_{l-1}, G^{(u)}_{l-2}, \cdots, G^{(u)}_0$, and thus the number of graphs visited during this procedure is exactly $l$.

**Theorem 3:** The vector computed by Equation 5 is exactly the same as that computed by GPA.

**Proof:** $u$’s local partial vector at level $m$ can be computed as:

$$r_u[G^{(u)}_m] = \frac{1}{\alpha} \sum_{h \in H(G^m_u)} S^H_{G^m_u}[G^{(u)}_m](h) \cdot P^H_{G^m_u}[G^{(u)}_m] + r_u[G^{(u)}_{m-1}].$$

Therefore, Equation 6 can be written as:

$$r_u = \sum_{m=0}^{l-2} \frac{1}{\alpha} \sum_{h \in H(G^m_u)} S^H_{G^m_u}[G^{(u)}_m](h) \cdot P^H_{G^m_u}[G^{(u)}_m]$$

$$+ \frac{1}{\alpha} \sum_{h \in H(G^m_u)} S^H_{G^m_u}[G^{(u)}_m](h) \cdot P^H_{G^m_u}[G^{(u)}_m] + r_u[G^{(u)}_1]$$

$$= \sum_{m=0}^{l-2} \frac{1}{\alpha} \sum_{h \in H(G^m_u)} S^H_{G^m_u}[G^{(u)}_m](h) \cdot P^H_{G^m_u}[G^{(u)}_m] + r_u[G^{(u)}_1]$$

$$= \cdots$$

$$= \frac{1}{\alpha} \sum_{h \in H(G^m_u)} S^H_{G^m_u}[G^{(u)}_m](h) \cdot P^H_{G^m_u}[G^{(u)}_m] + r_u[G^{(u)}_1]$$

Note that $G^{(u)}_0$ is the whole graph, and $G^{(u)}_0$ can be viewed as a subgraph in GPA, and thus the result of Equation 6 is exactly the same as Equation 5.

4.4 Distributed PPV Computation

Based on the graph hierarchy, we can compute the PPV of a graph from the pre-computation results w.r.t. its hub nodes and the PPV of its child graphs. However, it is still challenging to design a distributed algorithm, especially when the graph is large and the pre-computation vectors are too large to be saved in memory.

To address these challenges, we propose HGPA, the hub-distributed hierarchical graph partition based algorithm, which is load balanced and scalable because the computation in each level can be evenly distributed. We use one coordinator machine to collect the results from other machines for HGPA. We note that the computation of $r_u$ relies on all hub nodes in all levels as shown in Equation 4. This inspires us to divide the hub node set of each subgraph in each level into disjoint components to distribute the computation. Specifically, given a subgraph $S_G$, we divide $H(S_G)$ into $s$ disjoint subsets equally $H^1(S_G), H^2(S_G), \cdots, H^s(S_G)$, where $H(S_G) = \cup_{i=1}^{s} H^i(S_G)$. We do this on each subgraph in each level. As a result, the computation of $r_u$ can be interpreted as Equation 7 based on the balanced hub nodes partition.

$$r_u = \sum_{m=0}^{l-2} \frac{1}{\alpha} \sum_{i=1}^{s} \sum_{h \in H^i(G^m_u)} S^H_{G^m_u}[G^{(u)}_m](h) \cdot P^H_{G^m_u}[G^{(u)}_m] + r_u[G^{(u)}_1]$$

It is obvious that the first component of Equation 7 can compute the same results as the first component in Equation 5.

Based on Equation 7 for each subgraph in each level, we divide its hub node set evenly into $s$ disjoint subsets and store them in $s$ machines. We also distribute the leaf level subgraphs evenly to $s$ machines. Each machine only maintains the partial and skeleton vectors of nodes stored on it. Given a query node $u$, the $i^{th}$ machine computes a vector using the pre-computation results stored on it, i.e., $\sum_{h \in H^i(G^m_u)} S^H_{G^m_u}[G^{(u)}_m](h) \cdot P^H_{G^m_u}[G^{(u)}_m]$, and then sends the vector to the coordinator. The partial vectors of all non-hub nodes w.r.t. the leaf level subgraphs (e.g., $r_u[G^{(u)}_1]$) are also distributed to $s$ machines evenly. The coordinator sums up all the vectors received from the machines to construct the PPV. Figure 8 illustrates the idea of HGPA.

It is obvious that algorithm HGPA is load balanced. The computation on each machine is presented by Algorithm 4.

**Theorem 4:** If the hub node set of each subgraph is divided into $n$ disjoint subsets, the communication cost can be bounded by $O(n|V|)$. 
Algorithm 1: HGPA processor($q$, $l$, $\alpha$)

1. $p \leftarrow 0$
2. foreach subgraph $G_s$ do
   3. foreach hub node $h$ in $H^l(G_s)$ do
      4. foreach non-zero entry $p_{ij}^h(G_s)(k)$ do
         5. $ppv(k) \leftarrow pppv(k) + S_{ji}^h(G_s)(h) \cdot P^l_i(G_s)(k)/\alpha$
   6. if $q$ is not a hub node and $P^l_i(G_s^h)(k)$ on machine $i$ then
      7. foreach non-zero entry $P^l_i(G_s^h)(k)$ do
         8. $pppv(k) \leftarrow pppv(k) + p_{ij}^h(G_s^h)(k)$
   9. send $pppv$ to coordinator;

PROOF. In HGPA, each machine computes a vector of size at most $|V|$, and sends it to the coordinator. Hence, the total communication cost is $O(n|V|)$.

4.5 Space Cost of HGPA

We assume that the graph is partitioned in balance in each level. Let $l$ denotes the level of the graph hierarchy, and there are $2^l$ leaf level subgraphs. We have the following theorem that shows the space complexity of HGPA.

Theorem 5: The space cost of HGPA is $O((|V| - |H|)/2^l + \sum_{i=0}^{l} |H|(\sum_{i=0}^{l} |H|)/2^i + (|V| - |H|) \sum_{i=0}^{l} |H|_{max}^i)$, where $H_i$ is the set of hub nodes in the $i^{th}$ level, and $|H|_{max}^i$ is the maximum number of hub nodes in a subgraph in the $i^{th}$ level.

PROOF. In the leaf level, we compute and store the PPV of non-hub nodes w.r.t. each subgraph, and the size of the local PPV is $(|V| - |H|)/2^l$. Therefore, the total space cost of storing PPVs for leaf level subgraphs is $O((|V| - |H|)/2^l)$.

In the $i^{th}$ level, there are $2^i$ subgraphs. For each subgraph $G^l_i$, we need to compute the partial vectors of the hub nodes in it. Hence, the space cost of storing the partial vectors of hub nodes for the level is $O(|H|((|V| - |H|)/2^i))$. For all levels, the total space cost of this part is $O(\sum_{i=0}^{l} |H|(\sum_{i=0}^{l} |H|)/2^i)$.

We also need to compute the skeleton vectors of the non-hub nodes in $G^l_i$, and the space cost for $G^l_i$ is $O((|V| - |H|)/2^l |H|_{max}^l)$, where $H_i(G^l_i)$ is the set of hub nodes in $G^l_i$. In the $i^{th}$ level, the total space cost is $O((|V| - |H|)/2^i \sum_{j=0}^{l} |H|_{max}^j) < O((|V| - |H|)/2^i \sum_{j=0}^{l} |H|_{max}^j) = O((|V| - |H|)|H|_{max}^i)$. Thus, for all levels, the total cost of this part is $O((|V| - |H|) \sum_{i=0}^{l} |H|_{max}^i)$.

To compare the space cost of GPA with HGPA in an intuitive way, we consider that in GPA the graph is partitioned as the leaf level of the graph hierarchy in HGPA, i.e., the leaf level subgraphs in HGPA are the subgraphs in GPA. We can conclude that HGPA has smaller space cost than GPA. The two algorithms have the same space cost of storing the partial vectors of non-hub nodes. However, HGPA has smaller space cost of storing the partial vectors of hub nodes and the skeleton vectors of non-hub nodes than that in GPA, because $O(|V| |H|) > O(\sum_{i=0}^{l} |H|(|V| - |H|)/2^i)$ and $O((|V| - |H|)|H|) > O(|V| |H|) \sum_{i=0}^{l} |H|_{max}^i)$.

5. DISTRIBUTED PRE-COMPUTATION

We proceed to present how to pre-compute the partial and the skeleton vectors in a distributed manner for our algorithms.

5.1 Distributed Partial Vectors Computation

We adopt the selective expansion algorithm [23] as introduced in Appendix F.1 to compute the partial vectors for all nodes. As shown in Equation 10, the iteration requires the information of the graph structure. We keep a copy of the graph structure on each machine. As a result, no communication is required in the pre-computation of partial vectors. The computation can be done for each node separately, and each machine only needs to handle the nodes assigned to it.

In GPA, each machine only computes the partial vector $p_{ij}^h$ if node $u$ is assigned to it. When computing the partial vectors for a non-hub node, only the structure of the subgraph containing the node is required, because the computation can be restricted by the subgraph.

In HGPA, the partial vectors of nodes are computed by doing iterations w.r.t. the subgraphs, rather than w.r.t. the whole graph as in GPA, thus reducing the space and time complexity required during the iteration for a node. Given a node $u$, if $u$ is a non-hub node, $M$ computes its partial vector w.r.t. the leaf-level subgraph containing $u$. Otherwise, if $u$ is a hub node at level $m$, $M$ computes its partial vector w.r.t. the subgraph at level $m$ containing $u$.

5.2 Distributed Skeleton Vectors Computation

We improve the dynamic skeleton algorithm [23] as introduced in Appendix F.2 in order to distribute the skeleton vectors computation. As shown in Equation 10 for a given node $u$, two vectors $D_h(u)$ and $E_h(u)$ are used to do the iteration, and finally $D_h(u)$ would converges to $s^H_h(u)$, the skeleton vector of $u$. However, this algorithm incurs huge space cost, and the intermediate vectors could likely be larger than available main memory, and it is suggested to be implemented as a disk-based version [23]. The problem of the original method is that, in each step, the update of $D_h(u)$ relies on $D_h(v)$, if there exists an edge $(u, v)$. Thus, the skeleton vectors of $u$ cannot be computed without computing the skeleton vectors of other nodes. In addition, the computation of a skeleton vector of a node needs to consider all hub nodes, and we have to maintain the value of $s^H_h(u)$ for each node $u$ in memory. As a result, the original algorithm cannot work in parallel and consumes huge memory.

To distribute the computation and reduce memory cost, we consider computing the skeleton vector score w.r.t. a single hub node each time, i.e., $D_h(u)$ for each node $u$. Our algorithm can be explained by Equation 10. Initially, $F_0 = 0$. It is easy to show that after convergence, $F_k(u)$ is equal to $D_h(u)$, which is $s^H_h(u)$.

$$F_{k+1}(u) = (1 - \alpha) \sum_{v \in Out(u)} F_k(v) / |Out(u)| + \alpha s^H_h(u) \quad (8)$$

Theorem 6: $F_k(u)$ is equal to $s^H_h(u)$ when the iteration converges.
PROOF. We prove the correctness of Equation 8 by demonstrating that $F_k(u)$ is equal to $D_k[u](h)$ of Equation 10 in every steps. In the first round, $F_1(u) = \alpha x_u(u)$ and $D_1[u](h) = \alpha x_u(u)$, which are identical. In step $k + 1$, $F_{k+1}(u) = (1 - \alpha) \sum_{v \in Out(u)} \frac{F_k(v)}{|Out(u)|} + \alpha x_u(u)$, meanwhile we have $D_{k+1}[u](h) = (1 - \alpha) \sum_{v \in Out(u)} D_k[v](h) + \alpha x_u(h)$, and we can conclude that they are identical in each step.

The space complexity of the improved skeleton computation method is only $O(|V|)$. To obtain the full vector skeleton for a node $u$, we need to run this algorithm for each $h \in H$ ($|H|$ times in total).

In GPA, given a hub node $h$, we compute $s^H(h)$ w.r.t. the whole graph for each node $u$ according to Equation 8. In HGPA, given a hub node $h$ at level $m$, we first identify the subgraph at level $m$ where $h$ resides ($G^m_h$), and we compute $s^H(h)$ for each node $u$ in $G^m_h$. Note that based on Equation 8, there is no dependency among machines, and each machine can do the computation independently for the nodes assigned to it, and thus there is no network communication required.

6. EXPERIMENTS

6.1 Experimental Setup

Datasets. We conduct experiments on five public real-life network datasets:

Email [1]. This dataset is generated using email data from a large European research institution. This graph comprises 265,214 nodes and 420,045 edges.

Web [2]. This dataset is generated using web pages from the Google programming contest in 2002. This graph comprises 875,713 nodes and 5,105,039 edges.

Youtube [3]. This graph is generated from the video and user information from the video sharing website Youtube, and it comprises 1,134,890 nodes and 2,987,624 edges.

PLD [4]. This dataset is extracted using hyperlink pages from the Web Crawler released by the Common Crawl Foundation in 2014. As the whole dataset is very large, we extract a sample graph comprising 3,000,000 nodes and 18,185,350 edges. We also report the results on the full graph in Appendix C, which contains 101M nodes and 1.94B edges and is denoted by PLD_full.

Meetup. This dataset is a social graph crawled from Meetup. We collect various sizes of events to build the graphs of different sizes for studying scalability of our algorithm. The detail of this dataset is shown in Section 6.2.7.

Algorithms. We evaluate the performance of the distributed algorithm HGPA (Section 4.4). We also compare with the algorithm GPA (Section 3) and the distributed PPV computation using power iteration based on Pregel+ [44] and Blogel [43]. Note that there is no better baseline for distributed PPV computation. We evaluate the performance of these algorithms from the following aspects: the space cost, the efficiency of computing the PPV for a query node, and the communication cost during the online PPV computing. We also compare HGPA with power iteration and a state-of-the-art approximate PPV computing method [46] in a centralized setting.

Accuracy Metric. To show the accuracy of our proposed algorithms, we compare with the power iteration method using average $L_1$-norm and $L_{\infty}$-norm metrics, which are also used to evaluate PageRank algorithm performance [7]. Given the PPV vectors $r_u$ and $r_v$ computed by different algorithms, the average $L_1$-norm is defined as $L_1(r_u, r_v) = \sum_{v \in V} |r_u(v) - r_v(v)| / |V|$ and the $L_{\infty}$-norm is defined as $L_{\infty}(r_u, r_v) = \max_{v \in V} |r_u(v) - r_v(v)|$.

Query Generation. We randomly choose 1000 nodes as query nodes for each graph, and report the average performance over all queries. In all experiments, we only focus on single node queries. Parameters. By default, we set the number of machines as 6. We use the two-way hierarchical partition method [24] for HGPA. We set $\epsilon = 10^{-3}$ by default, following the previous work [23]. We set teleport probability $\alpha = 0.15$ for all experiments, as it is widely used in previous work.

Setup. All algorithms are implemented in C++ compiled with GCC 4.8.2 and run on Linux. The experiments are conducted in a cluster consisting of 10 machines, each machine with a 2.70GHz CPU and 64GB of main memory. The machines are interconnected by a 100MB TP-LINK switch. All the pre-computations are performed using 4 threads on each machine. For each experiment, we run our algorithm 100 times and report the average.

6.2 Experimental Results

6.2.1 Number of Hub nodes in HGPA

For HGPA, we perform the hierarchical partitioning until no edges exist within each subgraph. This is because further partitioning cannot gain more improvement. We show the effect of partitioning levels in Section 6.2.2. We partition Email into 5 levels (yielding $2^5 = 32$ leaf-level subgraphs), Web into 12 levels (4,096 leaf-level subgraphs), and Youtube and PLD are partitioned into 15 levels (32,768 leaf-level subgraphs).

The pre-computation space and time cost of HGPA depends on the ratio of hub nodes (hub nodes number / |V|) in each level. We list the number of hub nodes in each level obtained by multi-way hierarchical partitioning of the four datasets in Tables 2-5, where the original graph is at level 0 and the leaf subgraphs are in the maximum level. It can be observed that the number of hub nodes is much smaller than the total number of nodes in all datasets.

Table 2: Ratio of Hub Nodes in Each Level on Email

<table>
<thead>
<tr>
<th>Level</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hub Percentage (%)</td>
<td>0.43</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
</tr>
</tbody>
</table>

Table 3: Ratio of Hub Nodes in Each Level on Web

<table>
<thead>
<tr>
<th>Level</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
<th>5</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hub Percentage (%)</td>
<td>0.77</td>
<td>0.24</td>
<td>0.13</td>
<td>&lt;0.1</td>
<td>&lt;0.1</td>
<td>0.42</td>
</tr>
</tbody>
</table>

Table 4: Ratio of Hub Nodes in Each Level on Youtube

<table>
<thead>
<tr>
<th>Level</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hub Percentage (%)</td>
<td>0.79</td>
<td>0.79</td>
<td>1.43</td>
<td>0.78</td>
<td>0.67</td>
</tr>
</tbody>
</table>

Table 5: Ratio of Hub Nodes in Each Level on PLD

<table>
<thead>
<tr>
<th>Level</th>
<th>0</th>
<th>1</th>
<th>2</th>
<th>3</th>
<th>4</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hub Percentage (%)</td>
<td>1.4</td>
<td>0.79</td>
<td>0.67</td>
<td>0.39</td>
<td>0.36</td>
</tr>
</tbody>
</table>

\(^1\text{http://snap.stanford.edu/data/email-EuAll.html}\)
\(^2\text{http://snap.stanford.edu/data/web-Google.html}\)
\(^3\text{http://snap.stanford.edu/data/com-Youtube.html}\)
\(^4\text{http://webdatacommons.org/hyperlinkgraph}\)
\(^5\text{http://www.meetup.com}\)
6.2.2 Comparison of GPA and HGPA

This experiment is to compare the performances of GPA and HGPA using default parameters. We report the maximum runtime across all machines as the overall query processing time. For the space cost, we report the maximum space used among all machines. The pre-computation time is evaluated as the maximum time across all machines. The communication cost is reported as the size of all the data received by the coordinator during the query processing.

![Figure 9: Algorithms Comparison on Web](image)

Figure 9 shows the results of the comparison between GPA and HGPA on Web. We observe from the results that GPA runs a bit slower than HGPA, because HGPA is more load balanced. The maximum space cost and offline pre-computation time of HGPA are better than that of GPA, and this is consistent with the theoretical analysis in Section 4.5. The theoretical communication cost of HGPA and GPA is the same, while HGPA takes less network cost than GPA as shown in Figure 9. Similar results are observed on the other datasets, and are thus not reported.

Since HGPA outperforms GPA in terms of all the aspects concerned, we omit the results of GPA in the subsequent experiments.

6.2.3 Effects of Number of Machines

This experiment is to study the effect of machine number on HGPA.

![Figure 10: HGPA Runtime](image)

![Figure 11: HGPA Space Cost](image)

Figure 11(a)–11(c) show the space cost of HGPA. Note that each machine only stores the pre-computed vectors of nodes assigned to it. We report the maximum space cost over all machines. As expected, the maximum space cost is reduced when the number of machines increases. There is no redundant information shared between different machines.

![Figure 12: HGPA Pre-Computation Time](image)

The pre-computation time is shown in Figures 12(a)–12(c). Each machine only needs to do the pre-computation for the nodes stored on it. HGPA is load balanced, and thus the space cost of pre-computation is nearly linear to the number of machines used.

![Figure 13: HGPA Communication Cost](image)

Figures 13(a)–13(c) show the communication cost of HGPA. We notice even for the largest dataset HGPA only has less than 2MB network cost using on 10 machines. It can be observed that the communication cost increases as more machines are employed, which is consistent with our analysis in Theorem 4.

6.2.4 Effect of Partitioning Levels

This set of experiments is to study the effect of the level of the hierarchy of subgraphs on the performance of HGPA. Figures 14(a)–14(c) show the runtime for computing PPVs for query nodes on Email, Web, and YouTube, respectively. Figures 15 and 16 show the space and time cost of pre-computation on the three datasets.

The space and time of pre-computation drop significantly as we increase the number of levels of the graph hierarchy. As the number of levels increases, the number of subgraphs in the hierarchy increases exponentially, and thus the size of the subgraphs in leaf-level decreases greatly. However, at a certain level there exists few or no edges within each subgraph, further partitioning is unnecessary because it cannot reduce space cost any more. According to Equation 7 using more levels needs more computation to construct the PPV, and thus causes slightly longer query processing time, as observed in Figure 14. The communication cost is almost not affected by the number of partitioning levels, and thus is not reported.
6.2.5 Effect of Multi-way Partitioning

This experiment is to study the effect of the partitioning strategy. We report results on Web, and we observe quantitatively similar results on the other datasets. We partition Web into 2, 4, 8, 16, 64 subgraphs in each level and we study the time and space cost for both pre-computation and query processing.

![Figure 16: HGPA Pre-Computation Time](image)

Figure 16: HGPA Pre-Computation Time

6.2.6 Effect of Tolerance $\epsilon$

This experiment is to study the effect of tolerance $\epsilon$ on the performance of HGPA. For the offline pre-computation, the tolerance decides when the iteration terminates. Again, we report results on Web, and we observe similar results on the other datasets. Figures 17(a) and 17(c) show the query processing time, the space and time cost of the pre-computation. We can see that with more partitions in each level, the cost of pre-computation increases greatly, while the query processing time does not reduce much. Therefore, we choose the 2-way partitioning strategy as the default partitioning strategy. Note that the partition strategy almost does not affect the communication cost, and thus we do not report it.

![Figure 17: Web (HGPA)](image)

Figure 17: Web (HGPA)

6.2.7 Scalability

![Figure 18: Web (HGPA)](image)

Figure 18: Web (HGPA)

![Figure 19: $L_\infty$ norm Accuracy](image)

Figure 19: $L_\infty$ norm Accuracy

![Figure 20: Scalability (Meetup)](image)

Figure 20: Scalability (Meetup)

This experiment is to study the scalability of HGPA with the size of graphs. The difficulty is that there does not exist a group of graphs of different sizes but with similar properties. To this end, we build graphs of different sizes by taking different number of meetup events. The sizes of these graphs are listed in Table 6. In this experiment, we fix the number of employed machines to be 10.

![Table 6: Graph sizes for scalability study (Meetup)](image)

Table 6: Graph sizes for scalability study (Meetup)

<table>
<thead>
<tr>
<th>Graph ID</th>
<th># Nodes</th>
<th># Edges</th>
</tr>
</thead>
<tbody>
<tr>
<td>M1</td>
<td>997,304</td>
<td>82,966,338</td>
</tr>
<tr>
<td>M2</td>
<td>1,197,009</td>
<td>107,393,088</td>
</tr>
<tr>
<td>M3</td>
<td>1,396,054</td>
<td>129,774,158</td>
</tr>
<tr>
<td>M4</td>
<td>1,596,455</td>
<td>163,320,390</td>
</tr>
<tr>
<td>M5</td>
<td>1,796,226</td>
<td>194,083,414</td>
</tr>
</tbody>
</table>

![Figure 21: Scalability (Meetup)](image)

Figure 21: Scalability (Meetup)
the space cost and time of pre-computation of HGPA for graphs of different sizes. We can see that both the space cost and time increase almost linearly as we increase the size of graphs.

### 6.2.8 Exact vs Approximate

This experiment extends the study of Section 6.2.10 to show the advantage of exact PPVs compared to approximate PPVs. We use two accuracy metrics, i.e., Precision and Kendall’s τ (by following the work [11, 46]), to compare the top-100 nodes obtained from each algorithm with the result of the power iteration method. In a nutshell, Precision is based on the value of top-k PPVs results, and Kendall is based on the percentage of node pairs with correct ordering [11]. Figure 21 shows the results on Email and Web. We observe that both metrics HGPA performs much better than FastPPV [46], and even the approximate algorithm HGPA_ad achieves nearly full score. This indicates that about 30% of the top-100 nodes returned by FastPPV are wrong, and about 10% node pairs are ordered incorrectly.

<table>
<thead>
<tr>
<th>Number of Machines</th>
<th>Communication Cost (KB)</th>
<th>HGPA</th>
<th>Pregel+</th>
<th>Blogel</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>10^2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>10^3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>10^4</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>8</td>
<td>10^5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>10^6</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 22: Runtime

<table>
<thead>
<tr>
<th>Number of Machines</th>
<th>Communication Cost (KB)</th>
<th>HGPA</th>
<th>Pregel+</th>
<th>Blogel</th>
</tr>
</thead>
<tbody>
<tr>
<td>2</td>
<td>10^2</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>10^3</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>10^4</td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>8</td>
<td>10^5</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>10</td>
<td>10^6</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

Figure 23: Communication Cost

### 6.2.9 Comparison with General Distributed Graph Processing Systems

**Baseline.** To the best of our knowledge, there exists no work for distributed exact PPV computation. Many distributed graph computation platforms [43, 44] take the PageRank (PR) problem as a basic graph computing application. In these platforms, the PR problem is usually solved by the power iteration method. Since the personalized PageRank is derived from the PR problem [35], we can compute PPV by implementing the power iteration method on these platforms, which are used as baselines.

We compare HGPA with the power iteration method implemented on Pregel+ [44] and Blogel [43], which are well-known open-source distributed graph computation platforms (we denote the two algorithms by Pregel+ and Blogel, respectively). It is shown [44] that Pregel+ outperforms other Pregel systems such as Giraph [19] and GPS [37]. Blogel [43] breaks the bottlenecks of vertex-centric models such as Pregel. We compare HGPA with Pregel+ and Blogel in terms of the runtime and communication cost under the same tolerances, and the result is shown in Figures 22(a) and 23(b).

It can be seen that our algorithm is faster than Pregel+ and Blogel by orders of magnitude on Web and Youtube, and HGPA outperforms Pregel+ by at least two orders of magnitude in terms of communication cost. We observe that the runtime and communication cost of Pregel+ and Blogel increase when the number of machines increases. The reason is as follows: Pregel+ is designed based on the general bulk synchronous parallel (BSP) model. It sends messages from vertex to vertex in each iteration of the BSP step, and when the vertices are on different machines it needs a lot of communications between machines. As the number of machines increases, the number of messages increases which costs more communication time. We observe the same phenomenon on Blogel. However it always outperforms Pregel+ in terms of the runtime and communication cost. This is because Blogel is based on the block-centric model, and it sends messages from block to block. Our proposed algorithm HGPA significantly outperforms the algorithms implemented on Pregel+ and Blogel. We observe that the runtime of HGPA decreases significantly as the number of machines increases. This is achieved by avoiding the huge communication costs, and thus HGPA is more suitable for online PPV applications.

HGPA can also be implemented on a single machine. We compare the runtime of HGPA in a centralized setting with the power iteration method, under the same error tolerance, and the result is shown in Figure 24. It can be seen that our algorithm is at least 3.5 times faster than the power iteration method. On Email and Web the speedup is much more significant. It demonstrates that HGPA can achieve comparable performance in terms of runtime as the algorithm proposed in the work [33] on a single machine.

We also compare our work with the state-of-the-art approximate PPV computation method FastPPV [46]. In FastPPV, the PPV scores less than 10^{−5} are discarded. It is shown that removing the small values only sacrifices little accuracy [11, 12]. To compare with the approximate method, we also discard the offline scores that are less than 10^{−4}, and this adapted method is denoted by HGPA_ad. Since in FastPPV the number of hub nodes is a parameter that affects the trade-off between accuracy and runtime, we compare with FastPPV using different numbers of hub nodes.

Figure 25 shows the runtime on Email and Web datasets. We use Fast_h to denote the method of FastPPV using h hub nodes. We notice that our exact method HGPA is faster than FastPPV on small dataset and slower than FastPPV on the large one. We notice that the adapted approximate method HGPA_ad runs faster than FastPPV by orders of magnitude on both datasets. Note that in this set of experiments our algorithm is implemented on a single machine.
machine. On the distributed computing platforms, our algorithm will be much faster.

We next evaluate the accuracy of the four algorithms, and where use the result computed by power iteration as exact. We still use average $L_1$ and $L_{\infty}$ as we used (in Section 6.2.6). Figure 26 show the accuracy comparison on Email and Web datasets. On all measures, we observe that HGPA is much better than FastPPV since the computing model of HGPA is exact. We notice that HGPA$_{ad}$ also consistently outperforms FastPPV in terms of accuracy.

In summary, our proposed distributed algorithms GPA and HGPA have low communication cost and are load-balanced with acceptable pre-computation and space cost. HGPA consistently outperforms GPA, and it outperforms the power iteration method implemented on general graph processing platforms such as Pregel+ and Blogel by orders of magnitude. In the centralized setting, HGPA can achieve comparable query time with the exact method [33] and the approximate method [46]. Its adapted version HGPA$_{ad}$ is able to outperform the approximate method [46] in terms of both efficiency and accuracy.

7. RELATED WORK

Exact PPV computation. The personalized PageRank was first proposed by Jeh and Widom [23]. It usually needs to be computed in an online manner, and thus has a high requirement for efficiency. A straightforward way of computing the Personalized PageRank Vector for a given set of nodes is the power iteration method, which is prohibitively expensive in time and is not suitable even for offline scenarios.

Due to the hardness of computing the exact PPV, Jeh and Widom propose to limit the preference nodes in a subset of a specified hub node set [23]. Therefore, this approach cannot be used to compute the exact PPV for arbitrary preference node set.

Machaira et al. [33] propose an iteration-based method that computes exact PPVs by exploiting graph structures. They decompose a graph into a core and a small tree-width graph. The two components are processed differently, and the PPV is constructed using the processed results. Unfortunately, this approach is not able to compute exact PPVs in a distributed manner. According to the experimental study, this method is about five times faster than the power iteration method. As shown in the experimental study, our distributed algorithm HGPA can achieve similar runtime compared with [33] under centralized settings.

Approximate methods for PPV. Most of existing studies compute PPVs approximately to trade for efficiency. Some proposals (e.g., [6][14]) utilize the Monte Carlo simulation methods, while some other studies (e.g., [40]) utilize the matrix factorization. Zhu et al. [46] propose an approximate method based on the concept of the inverse P-distance [23]. They first partition the tours into different tour sets according to their importance. Then, they design an algorithm to aggregate the contribution of tours from the most important ones to less important ones. Our experimental results show that our exact algorithm HGPA has similar query time as the approximate algorithm [46] while obtaining much better accuracy, and an adapted version of HGPA outperforms the approximate algorithm [46] in terms of both efficiency and accuracy.

Top-k and node-to-node search for PPV. Some approaches [16]-[18][42] aim to find a small part of the PPV. That is, for a query node, they only identify its top-k relevant nodes and omit the other nodes. Lofgren et al. [29][30] study how to estimate the node-to-node PPV score. Given a query node $u$, a target node $v$ and threshold $\delta$, they estimate whether $r_u(v) > \delta$ is true. All these methods cannot be used for computing the whole PPV w.r.t. a given query node set. However, finding top-k or estimating node-to-node PPV value is insufficient for many applications (e.g., [4][6][11]) which require the PPV scores of all nodes.

Distributed PPV computation. There also exist studies on distributed computation of approximate PPVs. In particular, Bahmani et al. [6] proposes a distributed algorithm based on MapReduce utilizing Monte Carlo simulation, which has no guaranteed error bound. The general graph processing engines, such as Pregel [34], Pregel+ [44] and Blogel [53], can be used for various distributed graph processing. As shown in the work [34], the power iteration method can be implemented on Pregel to compute PageRank, and thus PPVs. However, using these engines always induces multiple rounds of communications between machines; therefore, the communication cost is large and the query processing is slow, which make them impractical for applications of PPVs that have a high requirement on efficiency. In contrast, our proposed algorithms only require the communication between the machines and the coordinator once at query time.

8. CONCLUSION

In this paper, we propose novel and efficient distributed algorithms to compute the exact PPV for all nodes. The proposed algorithms can be implemented on a general coordinator-based share-nothing distributed computing platform. The processors only need to communicate with the coordinator once at query time in our algorithms. We first develop the algorithm GPA that works based on subgraphs. To further improve the performance, we propose HGPA based on a hierarchy of subgraphs, which has smaller space cost and better load balance and efficiency than GPA. The experimental study shows that HGPA has excellent performance in terms of efficiency, space cost, communication cost, and scalability. HGPA outperforms the power iteration method implemented on two distributed graph processing systems by orders of magnitude. Moreover, an approximate version of HGPA outperforms FastPPV [46] in terms of both efficiency and accuracy.

9. ACKNOWLEDGMENT

This work was carried out at the Rapid-Rich Object Search(ROSE) Lab at the Nanyang Technological University, Singapore. The ROSE Lab is supported by the National Research Foundation, Singapore, under its Interactive Digital Media(IDM) Strategic Research Programme. This work is also supported in part by a Tier-2 grant (MOE-2016-T2-1-137) awarded by Ministry of Education Singapore, and a grant awarded by Microsoft. Xuemin Lin is supported by NSFC61232006, ARC DP150102728, DP140103578 and DP170101628.
10. REFERENCES


APPENDIX

A. SCALABILITY STUDY ON GENERAL GRAPH PROCESSING SYSTEMS

This experiment is to study the scalability of the power iteration method on Pregel+ and Blogel. We use the same datasets used in Section 6.2.7 for studying the scalability of HGPA. Figure 27(a) shows the runtime of computing PPV on the two platforms compared to our HGPA method. We observe that the runtime of Pregel+ and Blogel increases linearly with the size of graphs, and HGPA is orders of magnitude faster than them. Figure 27(b) reports the communication cost during the query processing. Similarly, we can observe that the communication cost of Pregel+ and Blogel grows linearly with the graph size. It is because the communication of Pregel+ and Blogel is based on edges, and thus it is linear to the number of edges.

![Figure 27: Scalability (Meetup)](image)

B. VARYING QUERY LOAD

This experiment is to study how the query load affects the performance of PPV computing. We vary the number of queries from 1k to 1m. Figures 28 and 29 show the average performance on Web and Youtube. We observe that on both datasets the average runtime and communication cost is stable when varying the query load. We can see that choosing 1,000 random nodes is sufficient for the study.

![Figure 28: Queryload on Web](image)

![Figure 29: Queryload on Youtube](image)

C. EXPERIMENTS ON LARGE GRAPH

This experiment is to study the performance of HGPA on the large graph data set PLD_full, which contains 101 millions of nodes and 1.94 billions of edges. To conduct this experiment, we deploy our algorithm on Amazon EC2 using up to 24 instances, each of which has 64 processors (1500 processors in total). We set $\epsilon$ as $10^{-2}$ to save the cost of resources, and other settings are same with our other experiments.

The result is shown in Figure 30 as the number of processors is varied from 500 to 1500. We observe that: although the dataset PLD_full is over 100 times larger than the sampled PLD dataset, our algorithm HGPA is still able to perform well in terms of query time, pre-processing cost, and communication cost. We also observe that although the network communication cost is high, the runtime is still in the magnitude of seconds. This indicates that even for large graphs the network communication does not affect the runtime very much, because our algorithm only requires one time of communication between the coordinator and each machine.

![Figure 30: HGPA Performance on PLD_full](image)

D. POWER ITERATION IMPLEMENTATION

In this section, we introduce how the power iteration algorithm is implemented. We present the centralized version, and it can be easily modified to work in parallel on Pregel+ and Blogel. We utilize the adjacency list to implement the power iteration algorithm, and this method is also used in the work [46]. The pseudo code is shown in Algorithm 2.

To improve the efficiency, we use a queue valuedNodes to track the nodes with non-zero value, and only the nodes in valuedNodes and their neighbors are visited. In each iteration, the nodes in the queue either teleport to the query node(line 12) or randomly surf to their out-neighbors(lines 17–21). Note that in lines 14–16, we deal


with the dangling nodes (nodes without out-neighbors) by adding an arc to query node $q$. At last, we check whether the stop condition is fulfilled (lines 25–28).

### E. HUB NODE SELECTION

The graph partitioning problem aims to remove minimum number of edges/nodes from a given graph such that the graph becomes disconnected. It is shown that finding such edge separators or vertex separators is NP-hard [10]. The best approximate algorithm of finding the vertex separators has an approximation ratio of $O(\log n)$ [13] with high computational complexity. For planar graphs, there exists $O(\sqrt{n})$ theoretical bound [27] for the number of vertex separators. For general real-world networks, the number of hub nodes is usually small, e.g., social graphs are often organized into communities with large internal density of edges and sparse edges between communities [15, 26]. The assumption that the number of hub nodes is small is also exploited by the matrix-based methods for PPV computation [16, 38], in which the matrix is reordered by partitioning the graph into subgraphs (each of them is a small matrix) and leaving the hub nodes in a matrix.

Due to the hardness of this problem, most graph partitioning algorithms aim to minimize the number of hub edges approximately.

It is shown [24] that the METIS method performs well on minimizing the number of hub edges and balancing the size of subgraphs. We first adopt this partition algorithm to minimize the number of hub edges, and we then try to find the minimum number of hub nodes from these hub edges. The problem of finding the minimum number of hub nodes from hub edges is equivalent to the well-known Vertex Cover problem, which is also NP-hard. Specifically, given the hub edges, we choose some endpoints of the hub edges to be hub nodes, such that each edge is covered by at least one hub node. In this way, if we remove all the hub nodes, every hub edge will also be removed and the graph is partitioned into $k$ parts. We use the approximate vertex cover algorithm proposed in the work [36] to approximately find the minimum set of nodes that can cover these hub edges.

It is worth mentioning that though we use approximate algorithm to find hub nodes, our algorithm for PPV computation is still exact only if these nodes can separate the graph. A better way of selecting the hub nodes is an area of future work.

### F. PARTIAL & SKELETON VECTORS COMPUTATION

Both the partial and hubs skeleton vectors are computed in iterative ways. Note that these vectors could be rounded to any given error, known as tolerance, which is also used in the power iteration algorithm. The precision can be guaranteed, and such algorithms are regarded as exact in the literatures [23, 36]. In this paper, exact PPV means that we achieve the same results as the algorithms proposed by Jeh and Widom [23]. Details about our distributed pre-computation of partial vectors and skeleton vectors are referred to Section 5.

#### F.1 Partial Vector Computation

We introduce the selective expansion algorithm [23] to compute $p^H_u$ w.r.t. node $u$. Two intermediate vectors $D_k[u]$ and $E_k[u]$ are maintained, where $k$ represents the $k^{th}$ iteration. Initially, $D_0[u] = 0$ and $E_0[u] = x_u$, where $x_u$ is a basic vector with zero filled except $x_u(u) = 1$. Then, Equation 7 illustrates the iteration method, where $Out(v)$ denotes the set of out-going neighbors of $v$.

$$D_{k+1}[u] = D_k[u] + \sum_{v \in V-H} \alpha E_k[u](v)x_v$$

$$E_{k+1}[u] = E_k[u] - \sum_{v \in V-H} E_k[u](v)x_v$$

$$+ \sum_{v \in V-H} \frac{1 - \alpha}{|Out(v)|} \sum_{i=1}^{\frac{|Out(v)|}{2}} E_k[u](v)x_{Out_i(v)}$$

During the iterations, $D_k[u]$ would converge to $p^H_u$, and $E_k[u]$ would approach 0, which represents the difference between $D_k[u]$ and $p^H_u$. $D_0[u]$ is a lower-approximation of $p^H_u$. The computation is terminated when the value of $E_k[u]$ approaches 0, which means we can obtain $D_k[u]$ to any arbitrary precision. More precisely, we set an error tolerance bound $\epsilon$ and terminate the iteration when $E_k[u](v) \leq \epsilon, \forall u, v \in V$.

#### F.2 Skeleton Vector Computation

We introduce the basic dynamic programming algorithm [23] that computes $s^H_u$ w.r.t. node $u$. We also maintain two intermediate vectors $D_s[u]$ and $E_s[u]$, where initially $D_s[u] = 0$ and
$E_0[u] = x_u$. Then, the iteration is done as follows:

\[
D_{k+1}[u] = 1 - \alpha \frac{|Out(u)|}{|Out(u)|} \sum_{i=1}^{Out(u)} D_k[Out_i(u)] + \alpha x_u
\]

\[
E_{k+1}[u] = 1 - \alpha \frac{|Out(u)|}{|Out(u)|} \sum_{i=1}^{Out(u)} E_k[Out_i(u)]
\]

During the iterations, $D_k[u]$ would converge to $s^H_u$, and $E_k[u]$ would approach 0, which represents the difference between $D_k[u]$ and $s^H_u$. We terminate the computation when the value of $E_k[u]$ approaches 0. That is, we set an error tolerance $\epsilon$, and it is treated as converged when each value in $E_k$ is less than $\epsilon$. 