# Efficient Personalized PageRank Computation: A Spanning Forests Sampling Based Approach

#### **ABSTRACT**

Computing the personalized PageRank vector is a fundamental problem in graph analysis. In this paper, we propose several novel algorithms to efficiently compute the personalized PageRank vector with a decay factor  $\alpha$  based on an interesting connection between the personalized PageRank values and the weights of random spanning forests of the graph. Such a connection is derived based on a newly-developed matrix forest theorem on graphs. Based on this, we present an efficient spanning forest sampling algorithm via simulating loop-erased  $\alpha$ -random walks to estimate the personalized PageRank vector. Compared to all existing methods, a striking feature of our approach is that its performance is insensitive w.r.t. (with respect to) the parameter  $\alpha$ . As a consequence, our algorithm is often much faster than the state-of-the-art algorithms when  $\alpha$  is small, which is the demanding case for many graph analysis tasks. We show that our technique can significantly improve the efficiency of the state-of-the-art algorithms for answering two well-studied personalized PageRank queries, including single source query and single target query. Extensive experiments on five large real-world graphs demonstrate the efficiency of the proposed method.

#### **ACM Reference Format:**

#### 1 INTRODUCTION

Given a graph G = (V, E), two nodes  $s, t \in V$ , and a decay factor  $\alpha$ , the Personalized PageRank (PPR)  $\pi(s,t)$  is defined as the probability that a random surfer starts from s stops at t when applying an  $\alpha$ -random walk, where a random surfer randomly stops at the current node with probability  $\alpha$  or travels to a neighbor of the current node with probability  $1 - \alpha$ . Clearly, by this definition, the PPR value  $\pi(u,v)$  naturally measures the importance of node t w.r.t. (with respect to) s. That is, after randomly surfing, if s stops at t with a high probability, then t is important to s. Based on such a nice property, PPR has been widely used in web search related applications [28].

However, a recent trend is to employ PPR for many graph analysis tasks, such as graph clustering [4, 40], graph embedding [51], and graph neural networks [13]. In particular, PPR values can express the graph structure information by considering all paths between two

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nodes, which can be viewed as a type of information propagation procedure across the graph [42]. This new trend brings significant efficiency and effectiveness improvement to graph analysis tasks, but also introduces new computational challenges.

The new applications of PPR in graph analysis often require a small decay factor  $\alpha$ . For example, for local graph clustering application, the optimal parameter setting for the decay factor is  $\alpha=0.01$  as reported in [40]. The same optimal parameter setting of  $\alpha$  can also be found in graph neural network application [13]. The reason could be that with a small  $\alpha$ , the  $\alpha$ -random walk can explore a large portion of the graph, thus can obtain more information compared to the case with a large  $\alpha$ . But unfortunately, all existing algorithms for computing the PPR values with a small  $\alpha$  (e.g.,  $\alpha=0.01$ ) are not very efficient on large graphs, which motivates us to develop more efficient algorithms to handle the small  $\alpha$  case.

Specifically, previous PPR computation methods can be divided into two categories, including deterministic methods [3, 4, 10, 28] and Monte Carlo algorithms [7, 32]. In our case, of particular interest is the Monte Carlo algorithm. To estimate the  $\pi(s,t)$  value, a classic Monte Carlo algorithm first simulates  $\alpha$ -random walks from s and then counts the fraction of walks that terminates at t as an estimation. However, the major drawback of this method is that it only cares about the end-node of each random walk, and other nodes in the random walk are totally ignored. Moreover, the efficiency of such a method is heavily dependent on the decay factor  $\alpha$ . For a small  $\alpha$ , such a classic Monte Carlo method is inefficient, because it often takes a long time to simulate an  $\alpha$ -random walk in this case.

To tackle the issues, we propose a novel solution based on an interesting connection between PPR and random spanning forests of the graph. We first establish a novel PageRank matrix forest theorem, which gives a new combinatorial explanation of PPR value  $\pi(s,t)$  as the probability that s is rooted in t in a rooted random spanning forest. Such a combinatorial explanation motivates us to design efficient PPR computation algorithms by sampling spanning forests. To this end, we propose a new loop-erased  $\alpha$ -random walk technique to generate random spanning forests via extending the classic Wilson algorithm [48]. Compared to the previous methods, our spanning forest sampling based technique has several appealing features: (1) each step in the  $\alpha$ -random walk except loops provides valuable information for estimating PPR values, and (2) the expected running time of our technique will not grow rapidly as  $\alpha$  decreases, thus it is insensitive w.r.t.  $\alpha$ .

We apply the newly-developed technique to improve previous algorithms for answering two PPR queries: single source PPR query and single target PPR query. For the single source PPR query, the state-of-the-art algorithm is a two-stage algorithm [46, 49] that combines deterministic forward push and Monte Carlo. Although several optimizations are made to accelerate the deterministic forward push

<sup>&</sup>lt;sup>1</sup>A rooted spanning forest partitions the graph into several connected components (each tree is a connected component). If a node s is located in a tree that has a node t as its root, we call that node s is rooted in t.

stage [31, 49], there is little work focusing on optimizing the Monte Carlo stage. In the Monte Carlo stage, all these methods just simply simulates  $\alpha$ -random walk, which is inefficient when  $\alpha$  is small. We show that our technique can improve the Monte Carlo stage by replacing the  $\alpha$ -random walk sampling with the proposed random spanning forest sampling. Note that our random spanning forest sampling technique is orthogonal to those optimizations made on deterministic forward push [31, 49], thus they can also be applied to optimize our solutions. For the single target PPR query, the stateof-the-art algorithm is the backward push algorithm [3]. Similarly, the backward push algorithm is also slow when  $\alpha$  is small especially for high-degree nodes. To overcome this issue, we also propose a two-stage algorithm combining backward push and sampling spanning forests. We show that our algorithm can achieve a relative error guarantee in theory. Finally, we conduct extensive experiments using 7 large real-world graphs to evaluate our algorithms. The results show that (1) for the single source PPR query, our best algorithm can achieve one order of magnitude speedup over the state-of-the-art algorithm [49] on large graphs when  $\alpha = 0.01$ , and (2) for the single target PPR query, our best algorithm is around 3× faster than the state-of-the-art algorithm. To summarize, the main contributions of this paper are as follows.

New theoretical results. We develop three novel matrix forest theorems, based on which the PPR value  $\pi(s,t)$  between two nodes s and t can be explained as the probability that s is rooted in t in a random spanning forest. We show that the proposed matrix forest theorems can be applied to develop an efficient algorithm to estimate the PPR values by sampling spanning forests in a graph. We believe that such a novel combinatorial explanation for the PPR values could be of independent interest.

New algorithms for PPR queries. We first propose a new algorithm to sample spanning forests based on a loop-erased  $\alpha$ -random walk technique. We show that such a loop-erased  $\alpha$ -random walk technique is insensitive w.r.t.  $\alpha$ . Based on this technique, we develop a new two-stage algorithm which combing forward push (backward push) and spanning forest sampling to efficiently answer the single source (target) PPR query.

Extensive experiments. We conduct extensive experiments using 7 real-world graphs to evaluate the efficiency of the proposed algorithms. The results show that our algorithms significantly outperform the state-of-the-art algorithms on all datasets. For reproducibility purpose, the source code of this paper is released at an anonymous link https://anonymous.4open.science/r/RSFPPR-5767.

#### 2 PRELIMINARIES

Let G = (V, E, W) be a weighted graph, where V(n = |V|) is a set of vertices, E(m = |E|) is a set of edges, and  $w_{uv} \in W$  denotes the weight of an edge e = (u, v). Denote by A, the adjacency matrix of G with  $A_{uv} = w_{uv}$  if  $(u, v) \in E$ ,  $A_{uv} = 0$  otherwise. Let L = D - A be the Laplacian matrix of G, where D is a diagonal matrix with each entry  $D_{ii} = \sum_j A_{ij}$ . For a node  $u \in V$ , the weighted degree of u, denoted by  $d_u$ , is equal to  $D_{uu}$ . If the graph is unweighted, the weighted degree is exactly equal to the number of neighbors. For easy understanding of our results, we assume that the graph G is undirected in the following sections. It is important to note that the

main technique and all theoretical results presented in Section 3 and Section 4 still work for directed graphs.

Given a source node s, a target node t, and a decay factor  $\alpha$ , the personalized PageRank (PPR) of t w.r.t. s, denote by  $\pi(s,t)$ , is the probability that an  $\alpha$ -random walk starting from s terminates at t. Here an  $\alpha$ -random walk is a random walk where in each step the random walk stops at the current node with probability  $\alpha$  and travels to a random neighbor with probability  $1-\alpha$ . With this definition, we mainly focus on two types of personalized PPR computation problem in this paper.

Given a source node s, the single source PPR query is to compute  $\pi(s,v)$  for each node  $v \in V$ . The answer of this query is a row vector  $p_s \in R^{1 \times n}$ . Similarly, given a target node t, the single target PPR query is to compute  $\pi(v,t)$  for each  $v \in V$ , and the answer of this query is a column vector  $p_t \in R^{n \times 1}$ . Below, we focus mainly on describing the concepts of single source PPR query, and similar concepts can also be applied for the single target PPR query.

Let  $P = D^{-1}A$  be the probability transition matrix where each row is normalized by the weighted degree. The PPR vector  $p_s$ , which is PPR value of all nodes w.r.t. the source node s, satisfying the following linear equation

$$p_s = \alpha e_s + (1 - \alpha)p_s \cdot P,\tag{1}$$

where  $e_s \in \mathbb{R}^{1 \times n}$  is the unit vector with 1 on the *s*-th element and 0 on others. Then, we have

$$\pi(s, v) = p_s[v] = \alpha [I - (1 - \alpha)P]_{sv}^{-1}.$$
 (2)

Note that Eq. (1) can be easily reformulated as an equivalent linear system:

$$\tilde{p_s}(L + \beta D) = \beta e_s,\tag{3}$$

where  $\beta = \alpha/(1-\alpha)$ ,  $\tilde{p_s} = p_s D^{-1}$  and L is the Laplacian matrix. Based on Eq. (3), we can easily derive that

$$\pi(s, v) = p_S[v] = [(L + \beta D)^{-1} \beta D]_{Sv}. \tag{4}$$

Based on Eq. (4), we define the  $\beta$ -Laplacian matrix as follows.

Definition 2.1. (β-Laplacian) Given a graph G = (V, E) and its Laplacian matrix L = D - A. Let  $\alpha$  be the decay factor of the  $\alpha$ -random walk. The  $\beta$ -Laplacian of G with parameter  $\alpha$  is defined as  $L_{\beta} = (\beta D)^{-1}(L + \beta D)$ , where  $\beta = \frac{\alpha}{1-\alpha}$ .

Clearly, by Eq. (4) and Definition 2.1, we have  $\pi(s,t)=(L_{\beta}^{-1})_{st}$ . Thus, answering the PPR queries is equivalent to computing the inverse of the  $\beta$ -Laplacian matrix. Clearly, the answer of the single source query is a row of  $L_{\beta}^{-1}$ , while the answer of the single target query is a column of  $L_{\beta}^{-1}$ . Computing both the single source and single target queries are often very costly for large graphs, thus many approximation algorithms with a relative error guarantee have been proposed in the literature [31, 46, 49]. Below, we formally define such two approximate query processing problems which will be served as two applications of the proposed technique.

Definition 2.2. (Approximate single source PPR query) Given a relative error threshold  $\epsilon > 0$ , a PPR threshold  $\mu$  and a source node s, an approximate single source PPR query problem aims to compute an estimation  $\tilde{\pi}(s,v)$  for each node  $v \in V$  with  $\pi(s,v) \geq \mu$  such that  $|\tilde{\pi}(s,v) - \pi(s,v)| \leq \epsilon \pi(s,v)$  with a low failure probability  $p_f$ .

Definition 2.3. (Approximate single target PPR query) Given an relative error threshold  $\epsilon > 0$ , a PPR threshold  $\mu$  and a target node t, an approximate single target PPR query problem aims to compute an estimation  $\tilde{\pi}(v,t)$  for each node  $v \in V$  with  $\pi(v,t) \geq \mu$  such that  $|\tilde{\pi}(v,t) - \pi(v,t)| \leq \epsilon \pi(v,t)$  with a low failure probability  $p_f$ .

The two parameters,  $\mu$  and  $p_f$ , are used to control the estimation quality. The parameter  $\mu$  is a threshold such that we can achieve a relative error guarantee if the exact PPR value exceeds  $\mu$ ;  $p_f$  is the failure probability that the algorithm will produce a wrong result. Following previous studies [31, 46, 49], we set  $\mu = \frac{1}{n}$  and  $p_f = \frac{1}{n}$  to guarantee a relatively precise result. With these settings,  $\mu$  and  $p_f$  are both small enough to achieve a high estimating precision.

#### 3 PAGERANK MATRIX FOREST THEOREM

The classic matrix tree (or matrix forest) theorem connects the number of spanning trees (or forests) to the determinant of the Laplacian matrix of a graph which is perhaps the most well-known result in spectral graph theory [16]. In this section, we establish a novel matrix forest theorem, referred to as the PageRank matrix forest theorem, based on the  $\beta$ -Laplacian matrix defined in Definition 2.1.

#### 3.1 New matrix forest theorems

For a forest F, the weight of F is defined as the product of all weights of edges in F, that is  $w(F) = \prod_{e \in F} w_e$ . For unweighted graphs, we simply have  $w_e = 1$ , and thus w(F) is also equal to 1 for all F. A spanning forest of G is a forest including all nodes in G. Note that a forest may have several connected tree components. A rooted spanning forest is a spanning forest where we specify one node as a root in each connected component. For convenience, if a node s belongs to a tree  $\mathcal{T}$  (each tree is a connected component in the forest) which has a node t as its root, we say that s is rooted in t in the spanning forest. We denote  $\rho(F)$  as the set of roots of t. The following result shows a connection between the determinant of t and the weights of the rooted spanning forests.

THEOREM 3.1. (PageRank matrix forest theorem) Given a graph G = (V, E, W), for  $\beta \in (0, \infty)$ , the determinant of  $L_{\beta}$  is related to the rooted spanning forests as follows:

$$det(L_{\beta}) = \frac{1}{\beta^n \prod_{u \in V} d_u} \sum_{F \in \mathcal{F}} w(F) \prod_{u \in \rho(F)} \beta d_u,$$

where  $\mathcal{F}$  denotes the set of all rooted spanning forests in G.

**PROOF.** For a rooted spanning forest F, we define its weight as the product of all edge weights divided by the product of  $\beta d_u$  for all node u that is not a root. With this definition, the above theorem states that  $det(L_\beta)$  equals the sum of weights of all rooted spanning forests. Clearly, to prove the theorem, a direct way is to expand the determinant to see whether it is equal to the sum of weights of all rooted spanning forests. Below, we show that with a proper expansion, there is a one-to-one mapping between each term in the determinant and each rooted spanning forest; and each term in the determinant exactly equals the weight of the corresponding rooted spanning forest.

To achieve this, we expand the determinant by Leibniz formula and the resulting terms are related to the subgraphs of *G*. We can prove that terms correspond to a subgraph containing loops have the

same number of opposite signs, thus will be eliminated in the final result. All remaining terms correspond to the subgraphs that have no loop, which are exactly the rooted spanning forests. To help readers easily follow the proof argument, we give an intuitive example which is illustrated in Fig. 1. In Fig. 1(a), there is a simple graph G with 3 nodes. We can easily derive that G has 8 rooted spanning forests in total as shown in Fig. 1(b). In Fig. 1(b), the green nodes denote the roots of the spanning forest and the weight of each rooted spanning forest is also given under each subgraph. If there is no edge in a rooted spanning forest, we set the weight of the corresponding rooted spanning forest as 1. As a result, we need to show:  $det(L_{\beta}) =$  $1 + \frac{w_{12}}{\beta d_2} + \frac{w_{31}}{\beta d_3} + \frac{w_{21}w_{31}}{(\beta d_2)(\beta d_3)} + \frac{w_{12}w_{31}}{(\beta d_1)(\beta d_3)} + \frac{w_{12}}{\beta d_1} + \frac{w_{31}}{\beta d_1} + \frac{w_{13}w_{21}}{(\beta d_1)(\beta d_2)}.$ By definition,  $L_{\beta} = (\beta D)^{-1}(L + \beta D)$ , and we have  $det(L_{\beta}) = (\beta D)^{-1}(L + \beta D)$ .  $det(\beta D)^{-1}det(L+\beta D) = det(L+\beta D)/(\beta^n \prod_{u\in V} d_u)$ . Next, we consider the matrix  $M = L + \beta D$ . Let  $m_{ij}$  denote the i, j-th element of M. Note that this matrix is almost the same as the Laplacian matrix L = D - A, with each non-diagonal item  $m_{ij} = -w_{ij}$ ; the only difference is that each diagonal term is  $d_u + \beta d_u$  instead of  $d_u$  for all  $u \in V$ . For example, the  $L + \beta D$  matrix of the graph G in Fig. 1 is:

$$L + \beta D = \begin{pmatrix} m_{11} & m_{12} & m_{13} \\ m_{21} & m_{22} & 0 \\ m_{31} & 0 & m_{33} \end{pmatrix}$$

$$= \begin{pmatrix} \beta d_1 + w_{12} + w_{13} & -w_{12} & -w_{13} \\ -w_{21} & \beta d_2 + w_{21} & 0 \\ -w_{31} & 0 & \beta d_3 + w_{31}. \end{pmatrix} (5)$$

Next, we expand the determinant  $det(L + \beta D)$ , and the complete expansion procedure of the running example in Fig. 1 are given as follows:

$$det(L + \beta D) = (-1)^{\sigma_1} m_{11} m_{22} m_{33} + (-1)^{\sigma_3} m_{12} m_{21} m_{33} + (-1)^{\sigma_4} m_{13} m_{22} m_{31}$$

$$(6)$$

$$= (-1)^{\sigma_1} (\beta d_1 + w_{12} + w_{13}) (\beta d_2 + w_{21}) (\beta d_3 + w_{31})$$

$$+ (-1)^{\sigma_3} (-w_{21}) (-w_{12}) (\beta d_3 + w_{31})$$

$$+ (-1)^{\sigma_4} (-w_{13}) (-w_{31}) (\beta d_2 + w_{21})$$

$$= (-1)^{\sigma_1} (\beta d_1) (\beta d_2) (\beta d_3) + (-1)^{\sigma_1} (\beta d_1) w_{21} (\beta d_3)$$

$$+ (-1)^{\sigma_1} (\beta d_1) (\beta d_2) w_{31} + (-1)^{\sigma_1} (\beta d_1) w_{21} w_{31}$$

$$+ (-1)^{\sigma_1} w_{12} (\beta d_2) (\beta d_3) + (-1)^{\sigma_1} w_{12} w_{21} (\beta d_3)$$

$$+ (-1)^{\sigma_1} w_{12} (\beta d_2) w_{31} + (-1)^{\sigma_1} w_{12} w_{21} w_{31}$$

$$+ (-1)^{\sigma_1} w_{13} (\beta d_2) (\beta d_3) + (-1)^{\sigma_1} w_{13} w_{21} (\beta d_3)$$

$$+ (-1)^{\sigma_1} w_{13} (\beta d_2) (\beta d_3) + (-1)^{\sigma_1} w_{13} w_{21} (\beta d_3)$$

$$+ (-1)^{\sigma_3} (-w_{12}) (-w_{21}) (\beta d_3) + (-1)^{\sigma_3} (-w_{12}) (-w_{21}) w_{31}$$

$$+ (-1)^{\sigma_4} (-w_{13}) (\beta d_2) (-w_{31}) + (-1)^{\sigma_4} (-w_{13}) w_{21} (-w_{31})$$

$$= (\beta d_1) (\beta d_2) (\beta d_3) + (\beta d_1) w_{21} (\beta d_3)$$

$$+ (\beta d_1) (\beta d_2) (\beta d_3) + w_{12} (\beta d_2) w_{31} + w_{13} (\beta d_2) (\beta d_3) + w_{13} w_{21} (\beta d_3)$$

$$+ (\beta d_1) (\beta d_2) (\beta d_3) + w_{12} (\beta d_2) w_{31} + w_{13} (\beta d_2) (\beta d_3) + w_{13} w_{21} (\beta d_3)$$

Recall that the Leibniz formula expresses the determinant of a matrix in terms of permutations of the matrix elements:

$$det(M) = \sum_{\sigma} sgn(\sigma) \prod_{k=1}^{n} m_{k,\sigma(k)}.$$
 (10)

For convenience, we refer to the right hand side of Eq. (10) as A1, which is, Eq. (6) in the running example. A permutation is an arrangement of elements in an arbitrary ordering. In graph theory, it is well-known that a permutation  $\sigma$  can be decomposed into loops; and it has a unique correspondence to a linear factor [41], which consists of each arc  $(k, \sigma(k))$ . Fig. 2(a) and (b) are two examples of permutations and the corresponding linear factors. Basically, there are two types of loops: self-loops and non-trivial loops. The sign of a permutation  $\sigma$ , denoted by  $sqn(\sigma)$ , is given by evenness of n minus the number of loops l. Let  $l_s$  be the number of self-loops,  $l_n$  be the number of non-self loops. Since each diagonal element in M has positive sign and each non-zero non-diagonal element has negative sign, the sign of each term in Eq. (10) is determined by  $(-1)^{(n-l_s)+(n-l)} = (-1)^{l-l_s} = (-1)^{l_n}$ , which is the number of nontrivial loops. The permutations and the signs of the running example is illustrated in Fig. 1(c). As there is no edge between  $v_2$ and  $v_3$ , there are only 3 out of 6 terms in the expanded result, see

The second step is to substitute each diagonal element, which corresponds to a self-loop in a linear factor, in Eq. (10) with non-diagonal elements using the following formula

$$m_{uu} = \beta d_u + d_u = \beta d_u + \sum_{v \neq u} |m_{uv}|.$$

After substituting and removing parentheses, we will obtain a summation of several terms, which we refer to as A2 (also Eq. (8) in the running example). Each term in A2 corresponds to a subgraph of G with no self-loop. Notice that each term in the result in Eq. (8) is written in the same order as each rooted spanning forest in Fig. 1(d) w.r.t. the correspondence relation. To see this, suppose that there are s self-loops on vertices  $L = \{k_1, \cdots, k_s\}$  in a linear factor, then the product of diagonal elements is

$$\prod_{l=1}^{s} (\beta d_{k_l} + \sum_{v \neq k_l} |m_{k_l v}|).$$

After removing parentheses, each node  $k_l \in L$  will provide a term which is either a term  $m_{k_lv}$  (implying there is an out-going arc  $(k_l,v)$ ) or a term  $\beta d_{k_l}$  (implying there is no out-going arc from  $k_l$ ). If the term  $\beta d_{k_l}$  is provided, we will mark  $k_l$  as a root node. For example, Fig. 2(c) shows a possible term that we may obtain, where  $v_3$  provides the  $\beta d_3$  term and is marked as a root, other three nodes have out-going arcs  $(v_1,v_2),(v_2,v_3),(v_4,v_1)$ . Note that the left acyclic subgraph in Fig. 2(c) must be produced by self loops. Thus, the products of self-loops will produce either nontrivial loops or acyclic subgraphs in which a node is marked as a root. This establishes the correspondence between a subgraph that has no self-loop and a term in A2.

The third step is to reduce similar terms in A2. Note that in A2, similar terms with positive sign and terms with negative sign will be added to zero. After removing such zero terms from A2, we can obtain a *simplified term* called A3 (also Eq. (9) in the running example). We will see that any subgraph with nontrivial loops will

have the same number of terms with positive sign as that of terms with negative signs in A2, which will be absent in A3. To see this, any nontrivial loops in A2 comes from two cases: (1) the original linear factor, and (2) produced by self loops. Suppose that there are p nontrivial loops in one term in A2. Then, if k of the p loops come from a linear factor, then there will be  $C_p^k$  such similar terms with sign  $(-1)^k$  (recall that the sign of each term in A1 is determined by the evenness of the number of nontrivial loops in a linear factor). Thus, the coefficient of all such terms is equal to

$$\sum_{k=0}^{\lfloor p/2\rfloor} C_p^{2k} - \sum_{k=0}^{\lfloor (p-1)/2\rfloor} C_p^{2k+1} = \sum_{q=0}^p (-1)^q C_v^q = (1-1)^p = 0.$$

The above equality follows by applying a binomial formula. As a result, each final term in A3 has no loops which corresponds to a spanning forest. Each connected component has one node marked as a root. Thus, each spanning forest F corresponds to a term in A3 with weight  $w(F) \prod_{u \in \rho(F)} \beta d_u$ , which completes the proof. We can also see that for each subgraph that has a loop (plot in red box in Fig. 1(d)), the number of corresponding terms that has positive signs is equal to the number of terms that has negative signs. After reducing the same terms, we obtain the final 8 terms, which corresponds to 8 rooted spanning forests. Divided by  $det(\beta D)$ , each term is exactly the weight of that rooted spanning forest.

Based on Theorem 3.1, we can further develop two matrix forest theorems based on the minors of the matrix  $L_{\beta}$  as follows.

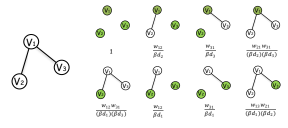
THEOREM 3.2. Given a graph G = (V, E, W), for  $\beta \in (0, \infty)$ , the determinant of the principle minor  $L_{\beta}^{(v)}$ , obtained by deleting the v-th row and column is related to the rooted spanning forests as follows:

$$det(L_{\beta}^{(v)}) = \frac{1}{\beta^n \prod\limits_{u \in V} d_u} \sum\limits_{F \in \mathcal{F}_v} w(F) \prod\limits_{u \in \rho(F)} \beta d_u,$$

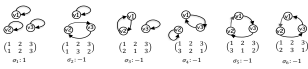
where  $\mathcal{F}_v$  denotes the set of all rooted spanning forests in G having v as a root.

PROOF. We can use a similar argument as used in the proof of Theorem 3.1 to prove this theorem. After deleting v-th row and column,  $det(L_{\beta}^{(v)}) = det((\beta D^{(v)})^{-1})det((L+\beta D)^{(v)}) = \frac{\beta d_v}{\beta^n \prod_{v \in V} d_u} det((L+\beta D)^{(v)})$ 

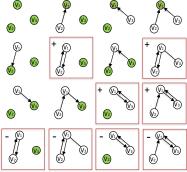
 $(\beta D)^{(v)}$ ). Next, we focus on the matrix  $det((L+\beta D)^{(v)})$ . Note that this matrix can also be expanded like Eq. (10). The only difference is that there is no  $m_v$ , term in the product. As a result, each term corresponds to a linear factor that must have a self-loop on v [38]. Then, the sign of each term still relates to the number of nontrivial loops following the proof of Theorem 3.1. After that, we can also substitute diagonal terms, remove parentheses and reduce similar terms to prove the theorem. Note that there is an additional  $\beta d_v$  term in the numerator of  $det(\beta D^{(v)})^{-1}$ . We can multiply this term into the weights. This is identical to the former proof where we choose the  $\beta d_v$  in the v-th parentheses instead of any arc out-going from v. Since all spanning forests with an out-going arc from v will be absent here, all final terms will correspond to one spanning forest in which v is marked as a root.



(a) G (b) All rooted spanning forests of G and their weights

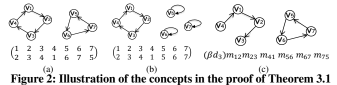


(c) All 3-permutations and their signs



(d) All terms after removing the parentheses

Figure 1: An intuitive example for proving Theorem 3.1. There is a one-to-one mapping between the determinant expansion terms and the rooted spanning forests. (a) An example graph G and (b) all 8 rooted spanning forests of G (green nodes are roots).  $det(L_B)$ equals the sum of the weights of all rooted spanning forests, which are given under each subgraph. Figures (c)-(d) illustrate the detailed proving procedure. Expand the determinant, there are 16 terms. All terms that has a loop (in the red box in (d)) will be absent in the final result. The remaining terms correspond to the rooted spanning forests.



THEOREM 3.3. Given a graph G = (V, E, W), for  $\beta \in (0, \infty)$ , given two distinct vertices u,v, the determinant of the minor  $L_{\rho}^{(u,v)}$ , obtained by deleting the u-th row and v-th column is related to the rooted spanning forests as follows:

$$det(L_{\beta}^{(u,v)}) = \frac{1}{\beta^n \prod_{i \in V} d_i} \sum_{F \in \mathcal{F}_{v,u}} w(F) \prod_{u \in \rho(F)} \beta d_u,$$

where  $\mathcal{F}_{v,u}$  denotes the set of all rooted spanning forests in G in which u and v are in the same connected component and u is a root.

PROOF. After deleting 
$$u$$
-th row and  $v$ -th column,  $det(L_{\beta}^{(u,v)}) = det((\beta D^{(u)})^{-1})det((L+\beta D)^{(u,v)}) = \frac{\beta d_u}{\beta^n \prod_{i \in \mathcal{U}} d_i} det((L+\beta D)^{(u,v)}).$ 

Next, we focus on the matrix  $det((L + \beta D)^{(u,v)})$ . This matrix can also be expanded like Eq. (10). The only difference is that there is no  $m_{u,\cdot}$  or  $m_{\cdot,v}$  term in the product. As a result, each term corresponds to a linear factor that must have an arc (u, v) [38]. The sign of each term still relates to the number of nontrivial loops following the previous proofs. Note that the arc (u, v) must belong to a nontrivial loop in a linear factor. Then, we can also substitute diagonal terms, remove parentheses and reduce similar terms to prove the theorem. Note that there is an additional  $\beta d_u$  term in the numerator of  $det((\beta D^{(u)})^{-1})$ . We can multiply this term into the weights. There is no term with an arc out-going from u, thus u is marked as a root, and v must be included in that connected component. All final terms will correspond to a spanning forest in which v is rooted in u. This completes the proof.

Note that although we focus mainly on undirected graphs, all results presented in the above theorems can be easily extended to directed graphs by using the concept of diverging forests as used in the traditional matrix forest theorem for directed graphs [1, 36]. Moreover, the extended results can also be proved by applying the same arguments based on the Leibniz formula as we used in the above theorems.

# 3.2 PPR computation by spanning forests

Recall that  $\pi(s,s)=(L_{\beta}^{-1})_{ss}$  and  $\pi(s,t)=(L_{\beta}^{-1})_{st}$ . By Cramer's rule, we can obtain  $\pi(s,s) = det(L_{\beta}^{(s)})/det(L_{\beta})$  for the diagonal term and  $\pi(s,t) = det(L_{\beta}^{(t,s)})/det(L_{\beta})$  for the non-diagonal term. Then, by the matrix forest theorem developed in Section 3, we can compute the PPR values by the weights of spanning forests. Formally, we have the following results.

THEOREM 3.4. Given a graph G = (V, E), a source node s and a decay factor  $\alpha$ , the PPR value satisfies

$$\pi(s,s) = \frac{\sum_{F \in \mathcal{F}_s} w(F) \prod_{u \in \rho(F)} \beta d_u}{\sum_{F \in \mathcal{F}} w(F) \prod_{u \in \rho(F)} \beta d_u},$$

where  $\beta = \alpha/(1-\alpha)$ ,  $\mathcal{F}$  is the set of all rooted spanning forests in G,  $\mathcal{F}_s$  denotes the set of all rooted spanning forests in G having s as a root.

**PROOF.** According to Theorem 3.1, Theorem 3.2 and  $\pi(s, s) =$  $det(L_{\beta}^{(s)})/det(L_{\beta})$ , the result follows by a simple substitution.  $\Box$ 

THEOREM 3.5. Given a graph G = (V, E), a source node s and a decay factor  $\alpha$ , the PPR value satisfies

$$\pi(s,v) = \frac{\sum\limits_{F \in \mathcal{F}_{v,s}} w(F) \prod\limits_{u \in \rho(F)} \beta d_u}{\sum\limits_{F \in \mathcal{F}} w(F) \prod\limits_{u \in \rho(F)} \beta d_u},$$

for every node  $v \in V$ , where  $\beta = \alpha/(1-\alpha)$ ,  $\mathcal{F}$  is the set of all rooted spanning forests in G,  $\mathcal{F}_{v,s}$  denotes the set of all rooted spanning forests in G in which s and v are in the same connected component and v is a root.

**PROOF.** The theorem can be easily proved by the results of Theorem 3.1, Theorem 3.3 and  $\pi(s,t) = det(L_{\beta}^{(t,s)})/det(L_{\beta})$ .

Based on the above two theorems,  $\pi(s, t)$  equals the proportion of weights of spanning forests in which s is rooted in t to weights of all spanning forests. Clearly, the weights of all spanning forests form a *weight distribution*. Let  $\Pr(s \text{ rooted in } t)$  be the probability that a node s rooted in a node t in a spanning forest randomly sampled from such a weight distribution. Then, we have the following results.

THEOREM 3.6.  $\pi(s, t) = \Pr(s \text{ rooted in } t)$ .

**PROOF.** According to Theorem 3.4 and Theorem 3.5,  $\pi(s,t)$  equals the sum of weights of rooted spanning forests that s is rooted in t, divided by the sum of weights of all rooted spanning forests. Since the rooted spanning forests are sampled from the distribution  $\Pr(F) \propto w(F) \prod_{u \in \rho(F)} \beta d_u$ , which is proportional exactly to the weight of rooted spanning forest, the probability that s is rooted in t equals  $\pi(s,t)$ .

Note that a spanning forest may contain several connected components (each tree is a connected component), which forms a partition of the nodes in the graph. Once a spanning forest F is generated, its corresponding partition  $\phi$  is determined. Note that for a fixed partition  $\phi$  of G, there may be many spanning forests in G that can produce the partition  $\phi$ . Interestedly, we find that if a partition  $\phi$  is given, the conditional probability that a node s is rooted in a node t (in a random spanning forest) conditioned on  $\phi$ , denoted by P(s rooted in  $t|\phi$ ), can be explicitly determined as follows.

THEOREM 3.7. Given a graph G = (V, E), a spanning forest F and its partition  $\phi = (V_1, \dots, V_k)$ . Suppose, without loss of generality, that s, t are two distinct vertices and t belongs to  $V_t$ . Let  $d_v$  be the weighted degree of node v. Then, the conditional probability that s is rooted in t conditioned on the partition  $\phi$  equals  $\frac{d_t}{\sum_{v \in V_t} d_v}$  if  $s \in V_t$ , equals 0 otherwise.

PROOF. Let w(F) denote the weight of a forest F. Let  $\tilde{w}(F) = w(F) \frac{1}{\prod_{u \in V} \beta d_u} \prod_{u \in \rho(F)} \beta d_u$  denote the weight of a rooted forest F. Let  $\mathcal{F}_{s,t}$  denote the set of rooted spanning forests that s is rooted in t. Recall that a rooted spanning forest can partition node into several connected components, we use  $\mathcal{F}^{\phi}$  to denote the set of rooted spanning forests that have the partition  $\phi$ . Let  $\mathcal{F}_{s,t}^{\phi}$  denote the set of rooted spanning forests that s is rooted in t, and has a node partition  $\phi$  at the same time. Then the conditional probability can be written as:

$$\Pr(F \in \mathcal{F}_{s,t}^{\phi} \mid \phi) = \frac{\Pr(F \in \mathcal{F}_{s,t}^{\phi})}{\Pr(F \in \mathcal{F}^{\phi})}$$
$$= \frac{\sum_{F \in \mathcal{F}_{s,t}^{\phi}} \tilde{w}(F)}{\sum_{F \in \mathcal{F}^{\phi}} \tilde{w}(F)}$$
$$= \frac{d_t}{\sum_{u \in V_t} d_u}$$

The last equality follows because  $\tilde{w}(F) = w(F) \frac{1}{\prod_{u \in V} \beta d_u} \prod_{u \in \rho(F)} \beta d_u$ .

On undirected graphs, for each  $F \in \mathcal{F}_{s,t}^{\phi}$ , in the connected component

 $V_t$  any node besides t can also be assigned as a root. So, there exists  $|V_t|$  rooted spanning forests F' that has weight  $\tilde{w}(F')$  differs from  $\tilde{w}(F)$  by only a  $d_t$  term.

Armed with Theorem 3.7, we can further obtain a different method to compute the PPR values. Let  $X_{st}$  be an indicator random variable that equals 1 if s and t are contained in the same component in a random spanning forest, equals 0 otherwise. Then, we have the following results.

THEOREM 3.8. 
$$\pi(s,t) = E\left[\frac{d_t}{\sum_{u \in V_t} d_u} X_{st}\right].$$

PROOF. Let  $\phi$  be an arbitrary partition of V. Then, we have the following results:

$$\pi(s,t) = \Pr(s \text{ rooted in } t)$$

$$= \sum_{\phi} \Pr(s \text{ rooted in } t | \phi) \Pr(\phi)$$

$$= \sum_{\phi} \frac{d_t X_{st}}{\sum_{u \in V_t} d_u} \Pr(\phi)$$

$$= \sum_{\phi} \frac{d_t X_{st}}{\sum_{u \in V_t} d_u} \sum_{Fs.t.\phi(F) = \phi} \Pr(F)$$

$$= \sum_{F} \frac{d_t X_{st}}{\sum_{u \in V_t} d_u} \Pr(F)$$

$$= E[\frac{d_t X_{st}}{\sum_{u \in V_t} d_u}].$$

## 4 SAMPLING SPANNING FORESTS

Note that by Theorem 3.6, we can estimate the PPR values via sampling spanning forests. Specifically, if we can sample spanning forests according to its weights, that is,  $P(F) \propto w(F) \prod_{u \in \rho(F)} \beta d_u$ , then an unbiased estimator of  $\pi(s,t)$  can be easily derived. For example, suppose that we have drawn N random spanning forests. If n of which has a component such that s is rooted in t, then we can estimate  $\pi(s,t)$  as  $\frac{n}{N}$ .

The remaining question is how can we sample spanning forests from such a weight distribution P(F)? To solve this problem, we propose a loop-erased  $\alpha$ -random walk approach to sample random spanning forests based on the weight distribution P(F). Our technique is a nontrivial extension of the classic Wilson algorithm for sampling spanning trees on graphs [48].

#### 4.1 The loop-erased $\alpha$ -random walk

The loop-erased  $\alpha$ -random walk does the same thing as the traditional  $\alpha$ -random walk, but erasing all loops in the random walk trajectory. Below, we first discuss the concept of the traditional loop-erased random walk as introduced in [48].

Given a graph G and a random walk trajectory  $\gamma = (v_1, \cdots, v_l)$  on G, we define the loop-erased trajectory as  $LE(\gamma) = (v_{i_1}, \cdots, v_{i_j})$  by deleting all loops in  $\gamma$ . Formally,  $i_j$  is defined by the following inductive procedure:  $i_1 = 1$  and  $i_{j+1} = max\{i|v_i = v_{i_j}\} + 1$ . Suppose that  $i_j$  is the max index by the above definition. Then,  $LE(\gamma)$  contains

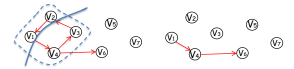


Figure 3: A random walk trajectory  $\gamma = (v_1, v_4, v_3, v_2, v_1, v_4, v_6)$  and its loop-erased trajectory  $LE(\gamma) = (v_1, v_4, v_6)$ 

 $i_j$  vertices and  $i_j-1$  directed edges. Loop-erased random walk is also self-avoiding; it will terminate when it hits the former trajectories. Initially, we set a node as a root and stop the first random walk when we hit the root. For example, in Fig. 3, suppose that  $v_6$  is the root, and there is a random walk  $\gamma = (v_1, v_4, v_3, v_2, v_1, v_4, v_6)$  stopping when it hits  $v_6$ . Then, its loop-erased trajectory is  $LE(\gamma) = (v_1, v_4, v_6)$ , by erasing the loop  $(v_1, v_4, v_3, v_2, v_1)$ . The process of erasing loops can be efficiently implemented by recording the next node in the random walk procedure. When a random walk stops, we retrace the trajectory, by starting from the first node, walking to the recorded next node until hitting the former trajectory. Note that the next node may be re-written many times, but after retracing it stores a unique next node in the final loop-erased trajectory.

The following results can be easily derived from [35].

THEOREM 4.1. Let  $\gamma = (v_{i_1}, \dots, v_{i_j})$  be the final random walk trajectory after erasing loops. Denote by the former trajectory set  $\Delta_0$  and let  $\Delta_k = \Delta_0 \cup \{v_1, \dots, v_k\}$ . We define  $w(\gamma) = \prod_{i=1}^k w_{i_{k-1}, i_k}$ . Then, the probability that  $\gamma$  is produced is

$$\Pr(\Gamma = \gamma) = w(\gamma) \frac{det(L^{\Delta_k})}{det(L^{\Delta_0})},$$

where L = D - A is the Laplacian matrix.

PROOF. By the definition of the loop-erased random walk, at each step  $(v_{i_k}, v_{i_{k+1}})$  of  $LE(\gamma)$ , a node from  $v_{i_k}$  may travel arbitrary number of loops before it finally walks one step from  $v_{i_k}$  to  $v_{i_{k+1}}$ . Let P be the transition matrix of the random walk. The probability that  $v_{i_k}$  returns to itself is  $\sum_{k=0}^{\infty} P_{v_{i_k}, v_{i_k}}^k = (I-P)_{v_{i_k}, v_{i_k}}^{-1}$ . During the random walk process, the random surfer is prevented from hitting the former trajectories  $\Delta_{k-1} = \{v_{i_1}, \cdots, v_{i_{k-1}}\}$ . For  $v_k$ , if it hits  $\Delta_{k-1}$ , the loop-erased random walk will stop, and thus the probability becomes  $(I-P)_{v_{i_k}, v_{i_k}}^{\Delta_{k-1}}$ , where  $A^{\Delta}$  denotes the submatrix by deleting rows and columns index by  $\Delta$  in A. The theorem follows from an application of Cramer's rule. As the probability that a node u travels to itself before hitting  $\Delta$  is given by  $G(u, u, \Delta) = (I-P)_{u,u}^{\Delta}$ , according to the Carmer's rule, it can also be written as  $\frac{\det(I-P)^{\Delta \cup \{u\}}}{\det(I-P)^{\Delta}}$ . It follows that the probability is:

$$\Pr(\Gamma = \gamma) = \prod_{k=0}^{k=j} G(v_{i_k}, v_{i_k}, \Delta) p_{v_{i_{k-1}}, v_{i_k}}.$$

The theorem follows by  $p_{v_{i_{k-1}},v_{i_k}} = \frac{w_{i_{k-1},i_k}}{d_{v_{i_k-1}}}$  and substituting each probability (note that most determinants will be eliminated by multiplying numerators and denominators).

The loop-erased  $\alpha$ -random walk is a nontrivial extension of the traditional loop-erased random walk. At each step, the loop-erased  $\alpha$ -random walk has a probability  $\alpha$  to stop. Suppose that it stops at

**Algorithm 1:** Sampling spanning forests by loop-erased  $\alpha$ -random walk

```
Input: Graph G = (V, E), a decay factor \alpha
   Output: Root[u] for all u \in V
1 InForest[u] \leftarrow false, Next[u] \leftarrow -1, Root[u] = -1 \text{ for } u \in V;
2 Fix an arbitrary ordering (v_1, \dots, v_n) of V;
3 for i = 1 : n \text{ do}
        u = v_i;
        while !InForest[u] do
6
             if rand() < \alpha then
              [nForest[u] \leftarrow true, Root[u] \leftarrow u;
7
8
                  Next[u] \leftarrow RandomNeighbor(u);
                  u \leftarrow Next[u];
10
        r \leftarrow Root[u], u \leftarrow v_i;
11
        while !InForest[u] do
12
             InForest[u] \leftarrow true, Root[u] \leftarrow r;
13
             u \leftarrow Next[u];
15 return Root[u] for all u \in V;
```

a node u, then u is marked as a root. Note that for the loop-erased  $\alpha$ -random walk, each loop-erased trajectory contains a root node when the  $\alpha$ -random walk stops. We can derive the probability that a loop-erased trajectory  $\gamma = (v_{i_1}, \cdots, v_{i_j})$  is generated when the loop-erased  $\alpha$ -random walk stops at  $v_{i_j}$ .

Theorem 4.2. Let  $\gamma = (v_{i_1}, \cdots, v_{i_j})$  be a loop-erased trajectory generated by a loop-erased  $\alpha$ -random walk which stops at  $v_{i_j}$ . Denote by the former trajectory set  $\Delta_0$ , and let  $\Delta_k = \Delta_0 \cup \{v_1, \cdots, v_k\}$ . We define  $w(\gamma) = \prod_{i=1}^k w_{i_{k-1}, i_k}$ . Then the probability that  $\gamma$  is produced is

$$\Pr(\Gamma = \gamma) = \beta d_{v_j} \frac{\det((L + \beta D)^{\Delta_k})}{\det((L + \beta D)^{\Delta_0})} w(\gamma).$$

PROOF. In this case, the probability that a node u travels to itself before hitting  $\Delta$  is given by  $(I-(1-\alpha)P)^{\Delta}_{u,u}$ , if the random walk does not stop with probability  $\alpha$ . According to the Carmer's rule, this can also be written as  $\frac{\det(I-(1-\alpha)P)^{\Delta\cup\{u\}}}{\det(I-(1-\alpha)P)^{\Delta}}$ . It follows that the probability  $\Pr(\Gamma=\gamma)$  is:

$$\alpha \frac{\det(I - (1 - \alpha)P)^{\Delta_{j-1}}}{\det(I - (1 - \alpha)P)^{\Delta_j}} \prod_{k=0}^{k=j-1} \frac{\det(I - (1 - \alpha)P)^{\Delta_{k-1}}}{\det(I - (1 - \alpha)P)^{\Delta_k}} \frac{(1 - \alpha)w_{v_{k-1}, v_k}}{d_{v_{k-1}}}$$

The theorem follows by eliminating numerators and denominators and by substituting  $\beta = \frac{\alpha}{1-\alpha}$  (note that  $L + \beta D = \frac{1}{1-\alpha}D - A = \frac{1}{1-\alpha}D^{-1}(I - (1-\alpha)P)$ ).

#### 4.2 Algorithm for sampling spanning forests

Here we present our algorithm for sampling spanning forests. The intuition is that by iteratively performing loop-erased  $\alpha$ -random walks until all nodes in G are traveled, the trajectory exactly constructs a rooted spanning forest. We will see that the probability of each spanning forest F generated by our algorithm is exactly proportional to its weights, i.e.,  $\Pr(F) \propto w(F) \prod_{u \in \alpha(F)} \beta d_u$ .

The implementation details of the loop-erased  $\alpha$ -random walk based sampling algorithm is outlined in Algorithm 1, which is an extension of the classic Wilson algorithm [48]. Specifically, Algorithm 1 starts by initializing an empty set F. We use a bool vector InForest to record whether node u has been added into F or not, a vector Next to record the next node in random walk step, and a vector *Root* to record the root of each node in the sampled spanning forest. The three vectors are initialized as false, -1 and -1 respectively (Line 1). Then, the loop-erased  $\alpha$ -random walks are performed iteratively from a node u following a pre-fixed node ordering, and the resulting loop-erased trajectory is added into F until all nodes are covered (Line 3-14). Specifically, in each step, the random walk will stop in two cases, either (1) terminates at the current node with probability  $\alpha$  (Line 6-7), or (2) terminates when hitting the former trajectories maintained by F (Line 5). If the loop-erased  $\alpha$ -random walk stops with the first case, the vertex u is assigned as a root and added into F (Line 7). After the random walk stops, we retrace the walk by the Next array, and add the loop-erased trajectory into F (Line 13-14). The algorithm terminates when all nodes are processed (Line 3), and F is returned as a rooted spanning forest sampled from the weight distribution (Line 15). Note that since we only use the root information of the sampled spanning forest, it suffices to return the *Root* vector to represent a rooted spanning forest.

THEOREM 4.3. Let  $\rho(F)$  be the root set of a rooted spanning forest F. Each F of G is sampled by Algorithm 1 with probability proportional to  $w(F) \prod_{\alpha(F)} \beta d_u$ , that is

$$\Pr(\gamma = F) = \prod_{u \in \rho(F)} \beta d_u \cdot \frac{w(F)}{\det(L + \beta D)} \propto w(F) \prod_{u \in \rho(F)} \beta d_u.$$

PROOF. Note that in Algorithm 1, after a loop-erased  $\alpha$ -random walk stops, we will start a new one outside from it, until all nodes are processed. Clearly, all nodes in V will be divided into l subsets  $\{V_1, \cdots, V_l\}$  by l loop-erased trajectories (each  $V_i$  is a node set of a loop-erased trajectory). Let  $\Delta_k = \bigcup_{i=1}^k V_i$ . Then, by Theorem 4.2, the probability  $\Pr(\gamma = F)$  is a product probability over all l loop-erased trajectories, in which each term is with  $\Delta_k$  on the numerator and  $\Delta_{k-1}$  on the denominator. As a result, the theorem is easily established by eliminating numerators and denominators.  $\square$ 

Complexity anlaysis. The time complexity of Algorithm 1 can be derived by analyzing the number of operations on the *Next* array (Line 10). For the loop-erased random walk, when a loop is generated, the *Next* value of a node will be revised. The total number of random walk steps is mainly determined by the total number of revision of the *Next* array in Line 10, because the cost spent in the retrace process (Line 11-14) is dominated by the cost taken by the random walk process (Line 4-10). Note that the expected number of revision of the *Next* array in Line 10 equals the sum of the expected number of visits of each node in the entire  $\alpha$ -loop erased walk process, which is denoted by  $\tau$ . As a result, the time complexity of Algorithm 1 is  $O(\tau)$ . Below, we analyze the the expected number of visits for each node in the  $\alpha$ -loop erased walk process.

Suppose that  $v_1$  is the first node in the pre-fixed node ordering. Then, the expected number of visits to  $v_1$  is given by  $\frac{1}{\alpha}\pi(v_1, v_1)$ ,

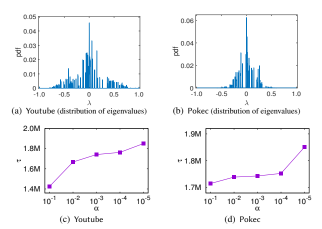


Figure 4: The distribution of eigenvalues of the matrix  $P(P = D^{-1}A)$  and the results of  $\tau$  with varying  $\alpha$ 

which can be derived by the power expansion  $\sum_{k=0}^{\infty} (1-\alpha)^k P^k_{v_1,v_1}$   $(P=D^{-1}A)$  is the probability transition matrix). In each step k, the expected probability mass that the  $\alpha$ -random walk passes through  $v_1$  is  $(1-\alpha)^k P^k_{v_1,v_1}$ , if the random walk does not stop. After that, the  $\alpha$ -loop erased walk will no longer pass through  $v_1$ . Furthermore, the most important property of the loop-erased walk is that the node ordering is irrelevant to the final result [48]. That is, any node can be the first node and the final trajectory maintains the same distribution [48]. As a result, during the whole  $\alpha$ -loop erased walk process, the expected number of visits to a node u is  $\frac{1}{\alpha}\pi(u,u)$ . By summing over all nodes, we can obtain  $\tau = \frac{1}{\alpha}\sum_{u \in V}\pi(u,u)$ . Note that since  $\pi(u,u) = \sum_{k=0}^{\infty}\alpha(1-\alpha)^k P^k_{uu}$ , we have  $\tau = \sum_{u \in V}(\sum_{k=0}^{\infty}(1-\alpha)^k P^k_{uu})$ . The following lemma show that  $\tau$  is closely related to the spectrum of the probability transition matrix  $P(P=D^{-1}A)$ .

LEMMA 4.4. Let  $1 = \lambda_1 > \lambda_2 \ge \cdots \ge \lambda_n > -1$  be the eigenvalues of the probability transition matrix P. Then, we have

$$\tau = \sum_{i=1}^{n} \frac{1}{1 - (1 - \alpha)\lambda_i}.$$
 (11)

PROOF. Recall that the trace of a matrix is the sum of its diagonal elements and also equals the sum of its eigenvalues. Thus, we have:  $\tau = \sum_{u \in V} (\sum_{k=0}^{\infty} (1-\alpha)^k P_{uu}^k) = \sum_{k=0}^{\infty} (1-\alpha)^k Trace(P^k) = \sum_{k=0}^{\infty} (1-\alpha)^k \sum_{i=1}^n \lambda_i^k = \sum_{i=1}^n \sum_{k=0}^{\infty} (1-\alpha)^k \lambda_i^k = \sum_{i=1}^n \frac{1}{1-(1-\alpha)\lambda_i}$ 

Armed with Lemma 4.4, we can explain why  $\tau$  is insensitive to  $\alpha$  as follows. To estimate all nodes' personalized PageRank values, traditional  $\alpha$ -random walk based methods need to simulate  $\alpha$ -random walks from all nodes, which takes  $O(\frac{n}{\alpha})$  per sample. For our looperased  $\alpha$ -random walk, the total time complexity is  $O(\tau)$ , which is related to the spectrum of P. There are a large number of previous studies on the spectrum of transition probability matrix on real world graphs [18, 21, 22]. Note that  $\tau$  is the sum of n terms, each term  $\frac{1}{1-(1-\alpha)\lambda_i}$  falls in the range  $(\frac{1}{2-\alpha},\frac{1}{\alpha}]$ , because  $|\lambda_i|<1$ . Therefore, we have  $\tau<\frac{n}{\alpha}$ . Moreover, we can see that when  $\lambda_i$  is close to 1,  $\frac{1}{1-(1-\alpha)\lambda_i}$  is close to  $\frac{1}{\alpha}$ . When  $\lambda_i$  is close to 0,  $\frac{1}{1-(1-\alpha)\lambda_i}$  is close to 1, which is independent on  $\alpha$ . As reported in [18], most of the eigenvalues of P in real-world graphs are concentrated around 0.

#### **Algorithm 2:** The Forward Push Algorithm

That is to say, most terms in Eq. (11) are independent on  $\alpha$ , thus  $\tau$  is insensitive to  $\alpha$  in real-world graphs. We also conduct experiments to compute the spectrum of P on 7 real world graphs (see Table 1) using the method proposed in [18]. The results on Youtube and Pokec are shown in Fig. 4(a-b). Similar results can also be observed on the other datasets. As can be seen, the distribution of eigenvalues are indeed concentrated around 0, implying that the time overhead of Algorithm 1 is insensitive w.r.t.  $\alpha$ . We also study how the  $\tau$  changes as  $\alpha$  decreases. As shown in Fig. 4(c-d),  $\tau$  increases smoothly w.r.t.  $\alpha$  (note that in the horizontal axis,  $\alpha$  decreases exponentially), which further confirms that our algorithm is insensitive to  $\alpha$ .

## 5 SINGLE SOURCE PPR QUERY

In this section, we apply the proposed techniques to improve the performance of the existing algorithms for answering the single source PPR query. Below, we first briefly review the forward push algorithm and the state-of-the-art two-stage algorithms which combine deterministic forward push and  $\alpha$ -random walk sampling. Then, we present our solutions by replacing the traditional  $\alpha$ -random walk sampling with the proposed loop-erased  $\alpha$ -random walk sampling.

#### **5.1** Existing solutions

The forward push algorithm. The forward push algorithm is an efficient local method to compute single source PPR vector which was first proposed in [4]. As shown in Algorithm 2, the forward push algorithm maintains two vectors, including a reserve vector  $q_s(v)$  and a residual vector r(v) for all  $v \in V$ . Specifically, a push procedure (Line 4-8) is applied for each v with r(v) larger than  $d_v \cdot r_{max}$  until no such v exists. During the entire procedure, the following invariant is maintained for all  $v \in V$  [4]:

$$\pi(s, v) = q_s(v) + \sum_{u \in V} r(u)\pi(u, v).$$
 (12)

The algorithm runs in  $O(\frac{1}{\alpha r_{max}})$  time. When  $r_{max}$  tends to 0,  $q_s(v)$  converges to  $\pi(s,v)$ . However, a major limitation of the forward push algorithm is that there is no additive or relative error guarantee on  $q_s(v)$  for a fixed  $r_{max}$ .

The  $\alpha$ -random walk sampling algorithm. The single source PPR query can be efficiently estimated by simulating  $\alpha$ -random walks. The algorithm generates a number of random walks from s, then counts the fraction of random walks that terminates at v as an estimation of  $\pi(s,v)$ . The major drawback of this algorithm is that

to obtain a precise estimation, the number of samples can be very large. According to [7], to guarantee a relative error  $\epsilon$ , it needs to generate  $O(\frac{nlogn}{\epsilon^2})$   $\alpha$ -random walks. As the expected length of each  $\alpha$ -random walk is  $\frac{1}{\alpha}$ , the algorithm takes  $O(\frac{nlogn}{\alpha\epsilon^2})$  time.

Combining forward push and  $\alpha$ -random walk sampling. To overcome the limitations of the forward push and the  $\alpha$ -random walk sampling algorithms, Wang et al. [46] proposed a two-stage algorithm, called FORA, which combines a deterministic forward push stage and a Monte Carlo stage by sampling  $\alpha$ -random walks. Let  $W = \frac{nlogn}{\epsilon^2}$ . To achieve a relative error  $\epsilon$ , FORA first performs forward push with threshold  $r_{max}$ , and then runs r(v)W random walks from each node v. The total  $\alpha$ -random walks needed can be bounded by  $nlogn \cdot r_{max}W$ . Then,  $r_{max}$  is set to minimize the complexity. As a result, FORA reduces the time complexity of the  $\alpha$ -random walk sampling algorithm from  $O(\frac{nlogn}{\alpha\epsilon^2})$  to  $O(\frac{nlogn}{\alpha\epsilon})$ . Recently, ResAcc [31] and SPEEDPPR [49] improves FORA by accelerating the forward push algorithm. In 9 improves FORA by accelerating the forward push algorithm. SPEEDPPR admits a time complexity  $O(\frac{1}{\alpha}nlognlog\frac{1}{\epsilon} + \frac{nlogn}{\alpha})$  which is the state-of-the-art algorithm. However, for all the two-stage algorithms, no existing optimization technique has been done for the Monte Carlo stage.

#### 5.2 Our solutions

In this subsection, we present our solutions based on the idea of replacing traditional  $\alpha$ -random walks with loop-erased  $\alpha$ -random walks in the state-of-the-art algorithms. We find that implementing such an idea is nontrivial, and there are two technical challenges needed to be tackled. Below, we first describe two challenges and the high-level ideas of our solutions to tackle these challenges.

Challenges and high-level ideas of our solutions. First, recall that in FORA, the number of  $\alpha$ -random walks needed to sample from node u is r(u)W, which is different for each node. This is because the threshold used in the forward push algorithm for each node is different. The high-degree node may admit a very large residue, thus requires a large number of  $\alpha$ -random walks. The total number of  $\alpha$ -random walk in FORA can be bounded by  $nr_{max}W$ . However, in the context of sampling spanning forests using loop-erased  $\alpha$ -random walks, the number of samples are the same for all nodes. Suppose that  $d_{max}$  is the largest degree over all nodes. Then, by applying the Chernoff bound, it requires  $d_{max}r_{max}W$  loop-erased  $\alpha$ -random walks, which makes the algorithm inefficient.

To circumvent this issue, we propose a new forward push algorithm called balanced forward push, which adapts the threshold for each node u from  $d_u r_{max}$  to  $r_{max}$ . The detailed implementation is shown in Algorithm 3. Although the balanced forward push algorithm only changes the threshold (compared to the traditional forward push algorithm), it is nontrivial to analyze its time complexity. Moreover, it is also very challenging to analyze the number of samples needed in our two-stage PPR computation algorithm when using such a balanced forward push as the push stage. We will tackle this by introducing an improved estimator together with carefully applying the Chernoff bound (see Theorem 5.3). Our result shows that it is sufficient to sample  $r_{max}W$  random spanning forests without losing theoretical guarantee. Note that sampling a random spanning forest is roughly equivalent to draw n  $\alpha$ -random walk samples. Since

**Algorithm 3:** The Balanced Forward Push Algorithm

sampling a random spanning forest by loop-erased  $\alpha$ -random walk is often much faster than sampling n  $\alpha$ -random walks, we can achieve significantly speedup over FORA, as confirmed in our experiments.

Second, to estimate  $\pi(s, t)$ , a basic estimator only needs the information of the spanning forests in which s is rooted in t based on Theorem 3.6. Let  $X_i$  be a random variable that represents whether a node i is rooted in a target node t in a random spanning forest. It is easy to verify that random variables  $X_1, \dots, X_n$  are dependent, which violates the condition of applying Chernoff inequality to bound the sample size. To tackle this challenge, we propose an improved estimator based on Theorem 3.8. The key idea of the improved estimator is based on the so-called conditional Monte Carlo estimation technique [37], because our spanning forests sampling method can obtain the root probability conditioned on a fixed partition of the graph. By using the conditional probabilities, we can reduce the variance of the estimator based on the result of the total variance formula Var[X] = Var[E[X|Y]] + E[Var[X|Y]], and Var[X] > Var[E[X|Y]] since a variance is always non-negative. More intuitively, compared to the basic estimator, the improved estimator based on Theorem 3.8 can use much more additional information of a sampled spanning forest (i.e., the information of two nodes in the same connected component), instead of only using the root information as used in the basic estimator, thus can reduce the variance. Note that such a variance reduction trick can reduce the number of samples needed for a desired accuracy guarantee. Moreover, we will show that we are able to bound the sample size based on such a technique.

The proposed algorithm. Based on the above high-level ideas, we present our algorithms FORAL and FORALV in Algorithm 4, which corresponds to the algorithm with the basic estimator and the improved estimator respectively. The algorithm first invokes the balanced forward push to obtain the residual r(s, u) and reserve  $q_s[u]$  for all  $u \in V$  (Lines 1-2). After that, the algorithm sets the parameters W and  $\omega$  to guarantee the approximate accuracy (Lines 3-4). Then,  $\omega$  random spanning forests are sampled independently by simulating loop-erased  $\alpha$ -random walks to estimate the PPR values (Lines 5-14). Let  $F_i$  be the *i*-th sampled random spanning forest (Line 7). Then, the estimator is updated in two different ways, with or without applying the variance reduction technique. In particular, in FORAL (the algorithm with the basic estimator),  $a_v$  is computed as the sum over residuals on the subset of the connected component which v belongs to (Line 14). However, in FORALV (the algorithm with the improved estimator),  $a_v$  is computed by weighted averaging

#### **Algorithm 4:** FORAL (FORALV)

```
Input: Graph G = (V, E), source node s, decay factor \alpha, push
             threshold r_{max}, relative error threshold \epsilon, PPR value
             threshold \mu
   Output: Estimate PPR \hat{\pi}(s, v) for all v \in V
1 Invoke balanced forward push with parameters G, s, \alpha and r_{max};
2 Let r(s, v_i), \hat{\pi}(s, t) be the returned residue and reserve for all
                \scriptstyle (2\epsilon/3+2)\cdot log(2/p_f)
4 Let \omega = \lceil r_{max} \cdot W \rceil;
5 for i = 1 : \omega do
        Simulate loop-erased \alpha-random walks on G;
        Let F_i be the returned random spanning forest;
        for each node t \in V do
              Let V_v be subset of the partition \Phi(F_i) which v belongs to;
              if apply variance reduction then
10
                   a_v = \frac{d_v \sum_{u \in V_v} r(u)}{\sum_{u \in V_v} d_u};
11
12
                a_v = \sum_{u \in V_v} r(u);
              \hat{\pi}(s,v) = \hat{\pi}(s,v) + a_v/\omega;
15 return \hat{\pi}(s, v) for all v \in V;
```

the residual in that subset according to the conditional probability (Line 11). Finally, the estimation  $\hat{\pi}(s, v)$  is returned for each  $v \in V$  as the query result (Line 15).

Note that in Line 1 of Algorithm 4, we can also use the improved forward push algorithm proposed in [49]. We refer to Algorithm 1 with the improved forward push algorithm as SPEEDL (with a basic estimator) and SPEEDLV (with an improved estimator) respectively. Below, we analyze the correctness and sample complexity of the proposed algorithms.

**Analysis of the algorithm.** First, we formally define the proposed estimators. Let r(u) be the residual of u returned by forward push algorithm,  $V_v$  be the vertex set that is rooted in the same node as v. Then, to estimate  $\sum_{u \in V} r(u)\pi(u,v)$  in Eq. (12), we can define two estimators  $\bar{r}(v)$  and  $\tilde{r}(v)$  for all  $u \in V$ , where  $\bar{r}(v) \triangleq \sum_{u \in V_v} r(u)$  is a basic estimator and  $\tilde{r}(v) \triangleq \frac{d_v \sum_{u \in V_v} r(u)}{\sum_{u \in V_v} d_u}$  is an improved estimator.

Let  $X_s$  be an indicator random variable that equals 1 if s is rooted in t in a spanning forest, equals 0 otherwise. Then, by Theorem 3.6, we have  $E[X_s] = \pi(s,t)$ . In other words,  $\bar{r}(v)$  is an unbiased estimator of  $\sum_{v \in V} r(v)\pi(v,t)$ , thus the correctness of FORAL and SPEEDL can be guaranteed.

Let  $X_{st}$  be the co-occurrence random variable that equals 1 when s and t are in the same connected component of a spanning forest, equals 0 otherwise. By Theorem 3.8, we have  $\pi(s,t) = E[\frac{d_t X_{st}}{\sum_{u \in V} d_v}]$ . Let  $Y = \sum_{v \in V} r(v) \frac{d_t X_{vt}}{\sum_{v \in V_s} d_t}$ . Then, we can derive that  $E[Y] = \sum_{v \in V} r(v) \pi(v,t)$  by the linearity of expectation. As a consequence,  $\tilde{r}(v)$  is an unbiased estimator of  $\sum_{v \in V} r(v) \pi(v,t)$ , which guarantees the correctness of FORALV and SPEEDLV.

The following lemma shows that the improved estimator has a smaller variance compared to the basic estimator.

```
Lemma 5.1. Var[\tilde{r}(v)] \leq Var[\bar{r}(v)] for all v \in V.
```

PROOF. For each node  $u \in V_v$ , we have  $\frac{d_v}{\sum_{u \in V_v} d_u} \leq 1$ . As a result,  $\frac{d_v \sum_{u \in V_v} r(u)}{\sum_{u \in V_v} d_u} \leq \sum_{u \in V_v} r(u)$ . We have  $E[\tilde{r}^2(v)] \leq E[\tilde{r}^2(v)]$ , the variance inequality follows.

Note that the relative error of Algorithm 4 with the basic estimator is hard to bound due to the dependency of random variables  $X_s$ . However, the practical performance of our FORAL and SPEEDL algorithms are comparable to the state-of-the-art algorithms as confirmed in our experiments. Interestingly, unlike the basic estimator, we find that Algorithm 4 with the improved estimator can obtain a relative error guarantee. For our analysis, we need the following Chernoff bound [17].

THEOREM 5.2. (Chernoff bound) Let  $X_i (1 \le i \le n)$  be independent random variables satisfying  $X_i \le E[X_i] + M$  for  $1 \le i \le n$ . Let  $X = \frac{1}{n_r} \sum_{i=1} X_i$ . Assume that E[X] and Var[X] be the expectation and variance of X. Then we have

$$\Pr(|X - E[X]| \ge \lambda) \le 2exp(-\frac{\lambda^2 n_r}{2Var[X] + 2M\lambda/3}).$$

By the Chernoff bound, we can derive the following theorem.

THEOREM 5.3. For any node t with  $\pi(s,t) > \mu$ , Algorithm 4 returns an approximate PPR value  $\hat{\pi}(s,t)$  satisfying  $|\pi(s,t) - \hat{\pi}(s,t)| \le \epsilon d_t \pi(s,t)$  with probability at least  $1 - p_f$ .

**PROOF.** Let  $Y = \sum_{u \in V} r(s, u) \frac{d_t X_{ut}}{\sum_{u \in V_s} d_t}$ . An important property of *Y* is that:

$$Y = \sum_{u \in V} r(s, u) \frac{d_t X_{ut}}{\sum_{u \in V_t} d_u} = \frac{d_t \sum_{u \in V_t} r(s, u)}{\sum_{u \in V_t} d_u} \le d_t r_{max}.$$

Let  $Z=\frac{Y}{d_t r_{max}}$ , we have  $Z\leq 1$ ,  $Z^2\leq 1$ . As a result, we have  $Var[Z]=E[Z^2]-E[Z]^2\leq E[Z^2]\leq E[Z]\leq \frac{\pi(s,t)}{d_t r_{max}}$ . Also, we have  $Var[Y]\leq d_t r_{max}\pi(s,t)$ . By substituting  $M=d_t r_{max}$  in Theorem 5.2, we have

$$\begin{split} \Pr(|Y-\mu| \geq \lambda) \leq 2exp(-\frac{\lambda^2 n_r}{d_t r_{max}(2\pi(s,t)+2\lambda/3)}). \\ \text{Let } \lambda = \epsilon d_t \pi(s,t). \text{ Then, by } \hat{\pi}(s,t) - \pi(s,t) = Y - E[Y], \text{ we have} \\ \Pr(|\pi(s,t)-\hat{\pi}(s,t)| \geq \epsilon d_t \pi(s,t)) \leq 2exp(-\frac{\epsilon^2 \cdot n_r \cdot \pi(s,t)}{r_{max}(2+2\epsilon/3)}) \leq p_f. \\ \text{The last inequality follows by substituting } n_r > \frac{r_{max}(2\epsilon/3+2)log(2/p_f)}{\epsilon^2 \cdot \mu} \end{split}$$

The last inequality follows by substituting  $n_r > \frac{\tau_{max}(ze_f) + \tau_{2} \log(z_f p_f)}{\epsilon^2 \cdot \mu}$  and  $\pi(s,t) > \mu$ .

Similar to the time complexity analysis of the forward push algorithm [4], we can easily derive that the time complexity of our balanced forward push is  $O(\frac{\bar{d}}{\alpha r_{max}})$ , where  $\bar{d}$  is the average degree. It can be further simplified as  $O(\frac{\log n}{\alpha r_{max}})$  on scale-free graphs when  $\bar{d} = \frac{\sum_{u \in V} d_u}{n} = \frac{2m}{n} = O(\log n)$ .

LEMMA 5.4. Let  $\bar{d}$  be the average degree. The time complexity of the balanced forward push can be bounded by  $O(\frac{\bar{d}}{\alpha r_{max}})$ .

PROOF. Let  $c_{push}$  be cost of each push operation. In each push operation of Algorithm 3, we pick a node u with residual r(s,u) larger than  $r_{max}$ , transfer a mass of  $\alpha r(s,u)$  to its reserve  $q_s[u]$ , and push the remaining probability mass  $(1-\alpha)r(s,u)/d_u$  to its

neighbors. According to Eq. (12), the reserve  $q_s[u]$  is always smaller than  $\pi(s,u)$ , thus the number of pushes is bounded by  $\frac{q_s[u]}{\alpha r_{max}} \leq \frac{\pi(s,u)}{\alpha r_{max}}$ . Note that each push operation processes  $d_u$  nodes and thus costs  $O(d_u)$ , which is denoted by  $c_{push}$ . Consequently, the cost of pushes on u is bounded by  $\frac{\pi(s,u)c_{push}}{\alpha r_{max}}$ , and the total cost is bounded by  $O(\sum_{v \in V} \frac{\pi(s,u)c_{push}}{\alpha r_{max}}) = O(\frac{c_{push}}{\alpha r_{max}})$ . Following the results in [33], the cost of push  $c_{push}$  can be amortized as the average degree  $\bar{d}$ , which completes the proof.

Based on Lemma 5.4, we can analyze the time complexity for all the proposed methods. In particular, the time costs of Algorithm 4 consist of two parts, the forward push stage and the Monte Carlo stage. For a fixed  $r_{max}$ , the time complexity of the deterministic forward push is bounded by  $O(\frac{\log n}{\alpha r_{max}})$ . The Monte Carlo stage samples  $r_{max}W$  random spanning forests. The cost of sampling a spanning forest is  $\tau$ ; and the estimation process takes O(n) in total, which is typically lower than  $\tau$ . Therefore, the Monte Carlo stage takes  $O(r_{max}W\tau)$  time. As a result, the total time complexity of Algorithm 4 is  $O(\frac{\log n}{\alpha r_{max}} + r_{max}W\tau)$ . This can be minimized by

setting  $r_{max} = \frac{\epsilon}{\sqrt{\alpha n \tau}}$ , which results in an  $O(\frac{1}{\epsilon} \sqrt{\frac{n \log n \tau}{\alpha}})$  complexity. Similarly, for SPEEDL and SPEEDLV, the time costs include two parts: the time spent for forward push and the time taken for sampling spanning forests. By a similar analysis shown in [49], we can easily derive that the total time complexity of SPEEDL and SPEEDLV is  $O(\frac{1}{\alpha} n \log n \log \frac{1}{\epsilon} + \log n \tau)$ . As can be seen, the time complexity of our algorithms has a weak dependency on the parameter  $\alpha$ , compared to the complexity of FORA [46] which is  $O(\frac{n \log n}{\alpha \epsilon})$ , and the complexity of SPEEDPPR [49] which is  $O(\frac{1}{\alpha} n \log n \log \frac{1}{\epsilon} + \frac{n \log n}{\alpha})$ . Therefore, our algorithms can be much faster than the previous algorithms when  $\alpha$  is small, which are also confirmed in our experiments.

## 5.3 Indexing spanning forests

Note that an optimization of FORA and SPEEDPPR is to pre-compute  $\alpha$ -random walks, and then maintain the end-node for each  $\alpha$ -random walk as an index. Such index-based methods are called FORA+ [46] and SPEEDPPR+ [49], respectively. To answer the single source PPR query, both FORA+ and SPEEDPPR+ can use the index to estimate PPR without simulating  $\alpha$ -random walks online. For space overhead, FORA+ requires  $d_v/\epsilon$   $\alpha$ -random walks for each node v. Thus, the total number of  $\alpha$ -random walks is  $\sum_{u \in V} d_u/\epsilon = 2m/\epsilon$ . The index size of FORA+ can be further bounded by  $O(\frac{n \log n}{\epsilon})$  with a relative error  $\epsilon$ , given that  $m = O(n \log n)$  on the scale free graphs. For SPEEDPPR+, it only requires  $d_v$  random walks for each v, thus its space overhead is  $O(n \log n)$  [49].

Similar to FORA+ and SPEEDPPR+, we can also devise indexbased variants of our online algorithms FORALV+ and SPEEDLV+. Specifically, we can first generate  $O(\log n)$  random spanning forests. Note that similar to SPEEDPPR+, we can derive that  $O(\log n)$  random spanning forests is sufficient to obtain a good estimation accuracy. Then, for each spanning forest, we maintain the root information for each node as the index. To implement the improved estimator in Algorithm 4 (Lines 10-11), we further maintains the total degree information in each connected component of a spanning

#### **Algorithm 5:** The Backward Push Algorithm

forest. The total space overhead of our index is  $O(n \log n)$ . Note that to estimate PPR, sampling a spanning forest by loop-erased  $\alpha$ -random walk is roughly equal to sampling n  $\alpha$ -random walks from each node. Because we can get n "samples of (i rooted in j)" for a spanning forest, while for an  $\alpha$ -random walk we only get one sample, i.e., the end node of the random walk. Thus, the number of samples needed by our algorithm is around 1/n times FORA+ and SPEEDPPR+. Since sampling a spanning forest ( $\tau$ ) is much faster than sampling n  $\alpha$ -random walks ( $\frac{n}{\alpha}$ ) (especially for a small  $\alpha$ ), the index construction time of our algorithm is much lower than FORA+ and SPEEDPPR+, as confirmed in our experiments. As a result, compared to FORA+ and SPEEDPPR+, the key advantage of our index-based methods is that they can significantly save index-building time especially when  $\alpha$  is small (e.g.  $\alpha = 0.01$ ), which is also confirmed in our experiments.

## **6 SINGLE TARGET PPR QUERY**

In this section, we focus on the single target PPR query. Compared to the single source PPR query, there are several differences which require more effort to design the single target PPR computation algorithm.

#### **6.1** Existing solutions

**Backward push.** The backward push algorithm is an analogy of the forward push algorithm [3]. As shown in Algorithm 5, it also maintains two vectors reserve  $q_t(v)$  and residual r(v). Then, a slightly different push procedure is applied for each node with  $r(v) > r_{max}$ , until there is no node meeting  $r(v) > r_{max}$  (Lines 4-8). The following invariant is maintained during the backward push process:

$$\pi(v,t) = q_t[v] + \sum_{u \in V} \pi(v,u) r(u). \tag{13}$$

Suppose that a push operation consumes time  $c_{push}$ , which is roughly  $O(\log n)$  in an average case [3], the algorithm runs in  $O(\frac{\pi(t)c_{push}}{\alpha r_{max}})$  time for a target node t. Unlike the forward push algorithm, the backward push algorithm can guarantee an additive error [3].

**Randomized backward push.** The randomized backward push algorithm introduces a sampling procedure into Algorithm 5 [43]. Specifically, in each push operation, a node sends residuals only to a small fraction of its randomly sampled neighbors, thus it can improve the efficiency. The time complexity of this algorithm is  $(\frac{n\pi(t)}{\alpha\epsilon})$  for a relative error  $\epsilon$ . The limitation of this algorithm is that

it needs to take additional cost for sampling in each push operation. Moreover, its time complexity still depends on  $\alpha$ .

## 6.2 The proposed algorithm

Here we develop two two-stage algorithms to answer the single target PPR query based on backward push and the proposed random forests sampling technique. Compared to the algorithms for processing single source PPR query, there are two differences in designing an algorithm for answering the single target PPR query. First, the time complexity of the backward push algorithm depends on  $\pi(t)$ of the target node t and it varies heavily over all nodes. For nodes with small  $\pi(t)$  (the low-degree nodes often have a small  $\pi(t)$ ), the backward push procedure terminates very fast. Consequently, there is no need to apply any sampling technique to speed up the algorithm for those nodes. For nodes with large  $\pi(t)$  (the high-degree nodes often have a large  $\pi(t)$ ), the backward push procedure often takes a long time especially when  $\alpha$  is small. Therefore, in this case, we can devise two-stage algorithms based on backward push and sampling random spanning forests. Second, unlike the forward push algorithm used in the single source PPR query problem, an additive error  $r_{max}$  can be guaranteed by the backward push algorithm. To achieve a relative error, we can set  $r_{max}$  as  $\frac{\epsilon}{n}$ , resulting in that the time complexity of the backward push is  $O(\frac{n\pi(t)c_{push}}{\alpha\epsilon})$ .

Implementation details. The pseudo code of our algorithms is shown in Algorithm 6. Algorithm 6 includes two stages including deterministic backward push and sampling random spanning forests. First, Algorithm 6 performs backward push to compute the residual and reserve for each node (Lines 1-2). Then, Algorithm 6 simulates the loop-erased  $\alpha$ -random walk technique to sample random spanning forests (Lines 5-14). Similar to Algorithm 4, Algorithm 6 can also use the basic estimating technique (Lines 10-11) and the improved estimating technique (Lines 12-13) to achieve unbiased estimations of PPR values. For convenience, Algorithm 6 with the basic estimator and the improved estimator are referred to as BACKL and BACKLV respectively.

Analysis of the algorithm. For a node v, let  $X_u$  be an indicator random variable that equals 1 if v is rooted in u, equals 0 otherwise. Let  $X_{st}$  be the co-occurrence random variable that equals 1 when s and t are in the same connected component of a spanning forest, equals 0 otherwise.  $Y_1 = \sum_{u \in V} r(u)X_u$  for BACKL and  $Y_2 = \sum_{u \in V} r(u)X_{uv} \frac{d_u}{\sum_{k \in V_v} d_k}$  for BACKLV. Similar to our previous analysis for the single source PPR query, we can easily derive that  $E[Y_1] = E[Y_2] = \sum_{u \in V} \pi(v,u)r(u)$ . Therefore, the variable  $a_v$  used in BACKL and BACKLV is an unbiased estimator of  $\sum_{u \in V} \pi(v,u)r(u)$ . Below, we apply the Chernoff bound to analyze the relative error guarantee of our algorithms.

THEOREM 6.1. For any node v with  $\pi(v,t) > \mu$ , both BACKL and BACKLV return an approximate PPR value  $\hat{\pi}(v,t)$  satisfying  $|\pi(v,t) - \hat{\pi}(v,t)| \le \epsilon \pi(v,t)$  with probability at least  $1 - p_f$ .

PROOF. Let  $Y_1$  and  $Y_2$  as defined before. After performing backward push, we have  $Y_1 \leq r_{max}$  and  $Y_2 = \frac{\sum_{u \in V_t} r(v) d_u}{\sum_{u \in V_t} d_u} \leq r_{max}$ .

#### **Algorithm 6:** BACKL (BACKLV)

15 **return**  $\hat{\pi}(v, t)$  for all  $v \in V$ ;

```
Input: Graph G = (V, E), target node t, decay factor \alpha, threshold
            r_{max}, relative error threshold \epsilon, PPR value threshold \mu
   Output: The estimated PPR \hat{\pi}(v, t) for all v \in V
1 Invoke backward push with input parameter G, t, \alpha and r_{max};
2 Let r(v), \hat{\pi}(v, t) be the returned residue and reserve for all v \in V;
3 Let W = \frac{(2\epsilon/3+2) \cdot log(2/p_f)}{2}.
4 Let \omega = \lceil r_{max} \cdot W \rceil;
5 for i = 1 : \omega do
        Simulate a loop-erased walk on G;
6
        Let F_i be the returned random spanning forests;
7
        for each node v \in V do
8
              Let V_t be subset of the partition \Phi(F_i) which t belongs to
               and the root is u;
10
             if apply variance reduction then
                  a_v = \frac{\sum_{u \in V_t} r(u) d_u}{\sum_{u \in V_t} d_u};
11
              else
12
13
                  a_v = r(u);
             \hat{\pi}(v,t) = \hat{\pi}(v,t) + a_v/\omega;
14
```

The following proof is the same for both  $Y_1$  and  $Y_2$ . We take  $Y_1$  as an example. Let  $Z=Y_1/r_{max}$ . Clearly,  $Z\in[0,1]$ . Then, we have  $Var[Z]=E[Z^2]-E[Z]^2\leq E[Z^2]\leq E[Z]\leq \frac{\pi(s,t)}{r_{max}}$ , and thus  $Var[Y_1]\leq r_{max}\pi(s,t)$ . We can apply Chernoff bound in Theorem 5.2 to analyze the concentration behaviour of  $Y_1$ . By substituting  $M=r_{max}$ , we have

$$\Pr(|Y_1 - \mu| \ge \lambda) \le 2exp(-\frac{\lambda^2 n_r}{r_{max}(2\pi(s,t) + 2\lambda/3)}).$$

Let  $\lambda = \epsilon \cdot \pi(s,t)$ ,  $n_r = \frac{r_{max}(2\epsilon/3+2) \cdot log(2/p_f)}{\epsilon^2 \cdot \mu}$ . By  $\hat{\pi}(s,t) - \pi(s,t) = r_{max}(Y_1 - E[Y_1])$ , for each node v with  $\pi(v,t) > \mu$ , the algorithm produces  $\hat{\pi}(v,t)$  which satisfies  $\hat{\pi}(v,t) - \pi(v,t) \le \epsilon \pi(v,t)$  with probability at least  $1 - p_f$ .

Note that although both BACKL and BACKLV can guarantee the same relative error as shown in Theorem 6.1, BACKLV has a smaller variance based on the improved estimating technique. Therefore, we focus mainly on the BACKLV algorithm in the remaining of this paper. The time complexity of BACKLV for a target node t consists of two parts. In the backward push stage, BACKLV takes  $O(\frac{c_{push}\pi(t)}{\alpha r_{max}})$  time, while in the Monte Carlo stage, BACKLV needs to sampling  $r_{max}W$  spanning forests which consumes  $O(r_{max}W\tau)$  time in total. Thus, the time complexity of BACKLV is  $O(\frac{c_{push}\pi(t)}{\alpha r_{max}} + \frac{c_{push}\pi(t)}{c_{push}\pi(t)})$ 

 $r_{max}W\tau$ ), which can be minimized to  $O(\frac{1}{\epsilon}\sqrt{\frac{c_{push}nlogn\tau}{\alpha}})$  by setting an appropriate  $r_{max}$ .

#### 7 EXPERIMENTS

#### 7.1 Experimental setup

**Datasets and query sets.** We use 5 real-life datasets including Youtube, Pokec, LiveJournal, Orkut and Twitter, which are widely

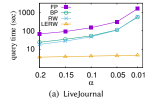
**Table 1: Datasets** 

Туре	Dataset	n	m	ā
	Youtube	1,134,890	2,987,624	5.27
unweighted	Pokec	1,632,803	22,301,964	27.32
graphs	LiveJournal	4,846,609	42,851,237	17.68
	Orkut	3,072,441	117,185,083	76.28
	Twitter	41,652,230	1,202,513,046	57.74
weighted	DBLP	1,824,701	8,344,615	32.32
graphs	Stack Overflow	2,584,164	28,142,395	37.02

used in previous studies [31, 43, 46, 49]. We also include 2 reallife general weighted graphs DBLP and StackOverflow. Specifically, DBLP is a collaboration network where each node represents an author, each edge represents collaboration relationship and the edge weight is the number of co-authored papers. StackOverflow is a user interaction network from the StackExchange site. Each node represents a user, each edge denotes an interaction relationship and the edge weight is the number of user interactions. The detailed statistics of these datasets are summarized in Table 1 . All these datasets can be obtained from [30]. For the single source query problem, as used in [46], we perform queries using 50 source nodes generated uniformly at random for all competitors and take the average query time as the final result. For the single target query, the query time is highly dependent on the chosen target node. For the low-degree nodes, it terminates fast by only applying backward push, while for the high-degree nodes, it spends a long time for the backward push such that sampling technique is necessary in this case. We perform queries on 50 target nodes generated uniformly at random from the top 10% highest degree nodes and again take the average query time as the final result.

**Different algorithms.** For single source PPR query, we compare our algorithms with the state-of-the-art algorithms FORA [46] and SPEEDPPR [49]. We do not include other previous algorithms in the experiments because all of them are outperformed by SPEEDPPR [49]. For FORA and SPEEDPPR, we use their original implementation in [46] and [49] respectively. For our solutions, we implement 4 different algorithms which are FORAL, FORALV, SPEEDL, and SPEEDLV. FORAL and FORALV denote Algorithm 4 with the basic estimator and the improved estimator respectively. Both of them use the balanced forward push in Line 1 of Algorithm 4. SPEEDL (SPEEDLV) is an improved algorithm for FORAL (FORALV) which is equipped with an improved forward push algorithm proposed in [49].

For single target PPR query, we compare our two-stage algorithm with the state-of-the-art algorithms BACK and RBACK. BACK is the backward push algorithm which can guarantee an additive error  $\epsilon$ . To achieve a relative error, we only need to set the threshold as  $\epsilon/n$  for BACK. RBACK [43] is the randomized backward push which can prune nodes with small residues in each push operation. However, RBACK needs to take additional time to preform random sampling. We implement BACK and RBACK by ourselves, as no available implementation of these algorithms are provided. For our algorithm, we implement BACKLV which is Algorithm 6 with an improved estimator. Note that since BACKL is clearly worse than BACKLV, we did not implement BACKL in our experiments.



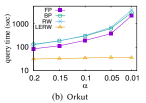


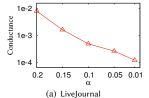
Figure 5: Runtime of four basic algorithms (vary  $\alpha$ )

**Parameters.** Since we focus mainly on small  $\alpha$ , we set the parameter  $\alpha=0.01$  in our experiments. Moreover, many existing PPR-based graph mining algorithms often work very well when  $\alpha=0.01$  [13, 40, 50]. We will study the performance of our algorithms with varying  $\alpha$  and also with very small  $\alpha$  (e.g.,  $\alpha=10^{-5}$ ). In addition, since the parameter  $\alpha$  is typically set to 0.2 in most existing algorithms [43, 46, 49], we also consider this parameter setting for a fair comparison with those baseline algorithms. For the approximate single source/target query, there is a parameter  $\epsilon$  which controls the relative error. We set  $\epsilon$  as 0.5 by default; and vary  $\epsilon$  from 0.1 to 0.5.

## 7.2 Effect of the parameter $\alpha$

We start by studying the performance of four basic algorithms with varying  $\alpha$ , including the forward push (FP), backward push (BP),  $\alpha$ -random walks (RW) and loop-erased  $\alpha$ -random walks (LERW), because FORA, SPEEDPPR, and our algorithms are based on these four basic algorithms. For both forward push and backward push, we set  $r_{\rm max} = 10^{-12}$ . We run  $\alpha$ -random walks from all nodes and one looperased  $\alpha$ -random walk for 10 times and report the total time as the results. We show the results on two large datasets: LiveJournal and Orkut. The results on the other datasets are consistent. As shown in Fig. 5, the runtime of the forward push algorithm, the backward push algorithm, and the  $\alpha$ -random walk algorithm grow rapidly when  $\alpha$ decreases, while the running time of our loop-erased  $\alpha$ -random walk is insensitive w.r.t.  $\alpha$ . These results confirm our previous analysis that (1) both the performance of the local push methods and the Monte Carlo methods by simulating  $\alpha$ -random walk have a strong dependency on  $\alpha$ , and (2) our loop-erased  $\alpha$ -random walk based algorithm is robust w.r.t.  $\alpha$ . We can also observe that when  $\alpha = 0.01$ , the running time of the  $\alpha$ -random walk algorithm is three orders of magnitude slower than that of the loop-erased  $\alpha$ -random walk algorithm. These results suggest that the Monte Carlo methods by simulating loop-erased  $\alpha$ -random walk is more efficient than by simulating  $\alpha$ -random walk when  $\alpha$  is small (e.g.,  $\alpha = 0.01$ ). Note that personalized PageRank with a small  $\alpha$  may be useful in many graph analysis tasks. Below, we demonstrate the effectiveness of the personalized PageRank with a small  $\alpha$  in a local graph clustering application.

For local graph clustering, we first compute personalized PageR-ank values from a source node s, and then sort the nodes based on their degree-normalized personalized PageRank values. After that, a sweep cut procedure is applied to produce clusters [4]. We make use of the well-known conductance metric [4] to measure the quality of the local clustering (a good clustering often has a small conductance value). We randomly choose 50 nodes as the source nodes and take the average conductance over 50 nodes as the final result. Fig. 6 shows the conductance with varying  $\alpha$  from 0.2 to 0.01 on LiveJournal and Orkut. Similar results can also be observed in the



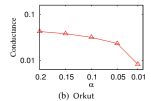


Figure 6: Conductance with varying  $\alpha$ 

other datasets. As can be seen, the conductance decreases sharply as  $\alpha$  decreases. These results indicate that a smaller  $\alpha$  will produce a better clustering, which are also confirmed in [40]. Therefore, it is important to study the problem of computing personalized PageRank with a small  $\alpha$ .

## 7.3 Single source query

In this experiment, we compare the performance of different algorithms for answering the single source query. The results on five datasets are reported in Fig. 7. For a better understanding of these results, we first focus on comparing the performance of FORA, FORAL, and FORALV. We observe that compared to FORA, both FORAL and FORALV obtain around 10× speedups when  $\alpha=0.2$  and around 100× speedups when  $\alpha=0.01$  on all five datasets. FORALV spends slightly more time than FORAL because it includes an additional computational cost of the sum over partitions. For large datasets, for example on Twitter, FORA runs out of 24 hours while both FORAL and FORALV take only thousands of seconds. In general, the runtime of all algorithms increase with decreasing  $\epsilon$ , because all algorithms take more time to achieve a small error.

Second, we compare the runtime of SPEEDPPR, SPEEDL and SPEEDLV. Note that SPEEDPPR applies an optimized version of forward push which is often more efficient, but it cannot apply the theoretical bound to balance the time spent in the Monte Carlo phase and the forward push phase (as FORA does). Alternatively, SPEEDPPR balance the time spent in the two phases based on an estimation of the  $\alpha$ -random walk time. SPEEDL and SPEEDLV also adopt a similar idea to balance the time. From Fig. 7, we can see that SPEEDL is the fastest algorithm among all the methods for both  $\alpha = 0.2$  and  $\alpha = 0.01$ . On the largest dataset Twitter, both SPEEDL and SPEEDLV are an order of magnitude faster than SPEEDPPR for  $\alpha = 0.01$ . These results confirm our theoretical analysis that by replacing traditional  $\alpha$ -random walks with loop-erased  $\alpha$ -random walks can achieve a considerable speedup (especially for the cases of small  $\alpha$  values).

To further evaluate the effectiveness of our loop-erased  $\alpha$ -random walk based algorithms, we compare the  $L_1$  errors of different algorithms. Here the  $L_1$  error is deified as  $\sum_{v \in V} |\tilde{\pi}(s,v) - \pi(s,v)|$  for all  $v \in V$ , where  $\tilde{\pi}(s,v)$  is the personalized PageRank value estimated by different algorithms and  $\pi(s,v)$  is the exact personalized PageRank value. Note that since both FORA and SPEEDPPR achieve the same relative error bounds, we focus mainly on comparing FORA, FORAL, and FORALV to see whether the loop-erased  $\alpha$ -random walk based algorithms can also achieve a good estimating accuracy. Similar results can also be obtained by comparing SPEEDPPR, SPEEDL, SPEEDLV. Fig. 8 shows the results on LiveJournal and Orkut. The results on the other datasets are consistent. As can be seen, FORALV achieves the smallest  $L_1$  errors for both  $\alpha=0.2$  and  $\alpha=0.01$ , followed by FORA and FORAL. Note that FORAL is worse than FORA,

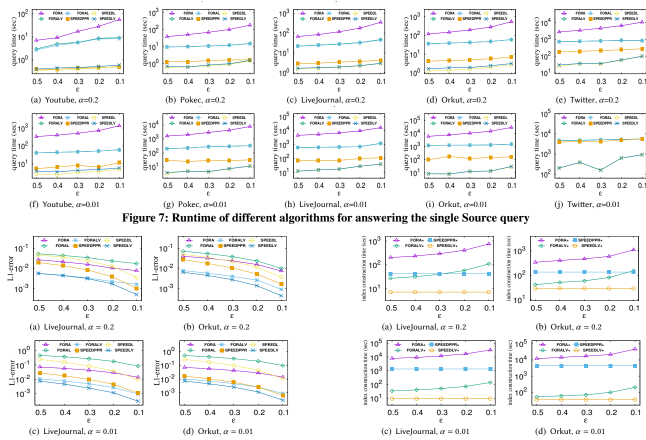


Figure 8: Comparison of the  $L_1$  error of different algorithms

because the the random variables used in FORAL is dependent. However, after using our variance reduction technique, the accuracy can be significantly improved. SPEEDPR, SPEEDL, and SPEEDLV follow a similar trend; their L1-errors are slightly smaller than FORA, FORAL, and FORALV respectively, because they conduct more deterministic computing. These results suggest that our loop-erased  $\alpha$ -random walk based algorithms can achieve a very good estimating accuracy.

## 7.4 Index-based method for single source query

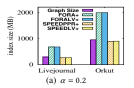
In this experiment, we evaluate the performance of different index-based algorithms. The index-based variants for FORA and SPEEDPPR are denoted by FORA+ [46] and SPEEDPPR+ [49] respectively. We implement two index-based variants for our algorithms FORALV and SPEEDLV, denoted by FORALV+ and SPEEDLV+ respectively, because FORALV (SPEEDLV) is shown to be better than FORAL (SPEEDLV).

We first compare the index construction time and index size of different algorithms. Note that both FORA+ and SPEEDPPR+ determine the index size based on theoretical results [46, 49]. FORA+ maintains  $O(n \log n/\epsilon)$   $\alpha$ -random walks [46], while SPEEDPPR+ stores around  $O(n \log n)$   $\alpha$ -random walks [49]. Since the performance of sampling n  $\alpha$ -random walks (from n nodes) is similar to that of sampling a spanning forest, FORALV and SPEEDLV maintain  $O(\log n/\epsilon)$  and  $O(\log n)$  spanning forests respectively. Fig. 9

Figure 9: Comparison of index construction time

shows the index construction time on LiveJournal and Orkut. The results on other datasets are consistent. As can be seen, SPEEDLV+ achieves the lowest index construction time under all parameter settings, followed by FORALV+, SPEEDPPR+, and FORA+. When  $\alpha = 0.01$ , SPEEDLV+ is around an order of magnitude faster than SPEEDPPR+. These results demonstrate that our index-based algorithms are much more efficient than the state-of-the-art algorithms to construct the index, which also confirm our analysis in Section 5.3. Fig. 10 reports the index size of different algorithms on LiveJournal and Orkut. From Fig. 10, we can see that our index-based algorithms can achieve similar index size as the state-of-the-art algorithms. This is because for a spanning forest sample, we need to store the root for each node, while for a random walk sample, we only need to store the end node. Thus, although the number of samples needed by FORALV+ (SPEEDLV+) is around 1/n times FORA+ (SPEEDPPR+), the total space costs of them are nearly the same. Moreover, the space usages of all the index-based algorithms are comparable w.r.t. the graph size. These results further confirm that our index-based algorithms are space-efficient.

Second, we evaluate the query processing time of different indexbased algorithms. The results are shown in Fig. 11. As can be seen, FORALV+ and SPEEDLV+ can achieve similar performance as FORA+ and SPEEDPPR+, respectively. Moreover, we also add the online algorithms FORALV and SPEEDLV for comparison. We can observe that all the index-based algorithms are faster than their



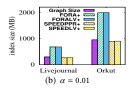


Figure 10: Comparison of index size

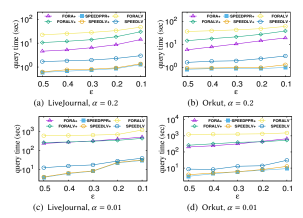


Figure 11: Runtime of different index-based algorithms

online versions. Note that FORALV+ and SPEEDLV+ are slightly slower than FORA+ and SPEEDPPR+ respectively, because our algorithms takes additional cost to sum the results over the partitions of random spanning forests.

In summary, our index-based algorithms can achieve similar query processing time and similar index size over the state-of-the-art index-based algorithms. However, our index can be constructed within much lower time than those of the state-of-the-art algorithms.

#### 7.5 Single target query

In this experiment, we compare the performance of BACK, RBACK, and BACKLV for answering the single target query. Fig. 12 shows the runtime of these three algorithms on five datasets. As shown in Fig. 12, when  $\alpha = 0.2$ , BACKLV degrades to BACK. This is because to achieve high precision, performing a backward push is often faster than sampling a random sampling forest when  $\alpha$  = 0.2. Therefore, we conclude that in this case Monte Carlo is not necessary for getting a precise result. When  $\alpha = 0.01$ , we can see that BACKLV is significantly faster than BACK and RBACK. In general, BACKLV can achieve  $1 \times \sim 3 \times$  speedups over BACK on all datsets under most parameter settings. We also observe that RBACK is worse than BACK. The reason could be that (1) RBACK needs to use additional computational cost for sampling, and (2) to achieve a high precision, RBACK needs to set a small sampling threshold so that its performance is similar to that of the power method, which is often worse than the backward push algorithm. These results indicate that for the small  $\alpha$  case, our loop-erased  $\alpha$ -random walk based technique can also be useful for processing the single target query.

## 7.6 Results on real-life weighted graphs

In all previous experiments, we only consider unweighted graphs (a special case of weighted graph with all edge weights equaling 1) for

a fair comparison with the state-of-the-art algorithms. In this experiment, we study the performance of different algorithms on general weighted graphs. To this end, we re-implement all the baseline methods as the available implementations in [43, 46, 49] cannot support general weighted graphs. The results of single source experiments on DBLP and StackOverflow are shown in Fig.13 and Fig.14. Similar results can also be observed on the other datasets. In general, the results on weighted graphs are consistent with our previous results on unweighted graphs. FORAL and FORALV (SPEEDL and SPEEDLV) have significantly less query time than FORA (SPEEDPPR). Our best algorithm SPEEDLV is at least one order of magnitude faster than the state-of-the-art algorithm (SPEEDPPR). The comparison of empirical error is also similar to that on unweighted graphs. Our SPEEDLV is clearly the winner with all parameter settings, and it is much more accurate than SPEEDPPR. Note that although there is a  $d_t$  term in the error bound of FORALV and SPEEDLV, the practical error performance is significantly better than that of FORA and SPEEDPPR as shown in Fig.14. The results of single target experiments are depicted in Fig.15. When  $\alpha = 0.01$ , BACKLV achieves a 2× speed-up on both datasets, which is consistent with the previous results on unweighted graphs.

## 7.7 Results with various query node distributions

Here we study the effect of query node distributions. To this end, we consider three different node distributions to study how the degree of query node affects the query time. Specifically, we independently sample nodes uniformly from the whole node set, the top 10% highdegree node set and the top 10% low-degree node set respectively. In this experiment, we choose the best methods, SPEEDLV and BACKLV, as two representative methods for single source query and single target query respectively; the results for other methods are consistent. To see the distribution of query time, we conduct experiments on 1000 nodes and use box-plot to show the results. The results are depicted in Fig.16. As can be seen, the query time of single source algorithms always have a very small variance. However, the query time of the single target algorithms are highly dependent on the node degree; and the query time of low-degree nodes is significantly lower than that of the high-degree nodes. For example, on Pokec when  $\alpha = 0.01$ , it takes around 400s for high-degree nodes, while all low-degree nodes take less than 1s to perform a query. These results indicate that for single target query, we only need to perform sampling to improve the query efficiency for high-degree query nodes, while for low-degree nodes, there is no need to do sampling.

#### 7.8 Results with very small $\alpha$

As discussed before, our algorithms are faster than existing methods especially when  $\alpha$  is small. Previously, we only consider the case when  $\alpha$  is relatively small ( $\alpha=0.01$ ). In this experiment, we study the case when  $\alpha$  is very small. Note that if  $\alpha$  tends to zero, the single source PPR vector tends to a degree-weighted uniform distribution  $\pi(s,u)=\frac{d_u}{2m}$  and the single target PPR vector tends to a constant distribution  $\pi(u,t)=\frac{d_t}{2m}$  for all  $u\in V$ . Therefore, the degree-weighted uniform distribution vector is a very simple baseline for computing single source PPR vector when  $\alpha$  is very small. Note that although the PPR vector is very close to a degree-weighted uniform

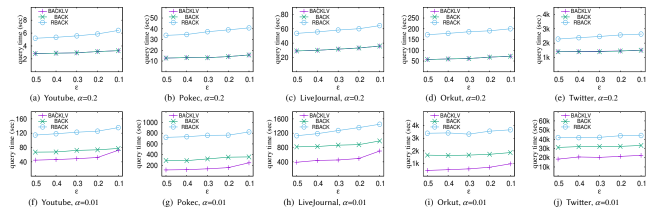


Figure 12: Runtime of different algorithms for answering the single target query

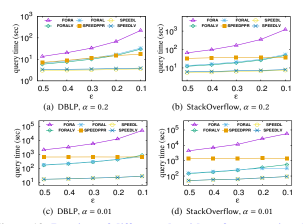


Figure 13: Runtime of different algorithms for answering the single source query on general weighted graphs

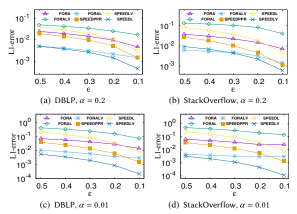


Figure 14: Comparison of  $L_1$  error for answering the single source query on general weighted graphs

distribution when  $\alpha$  is very small, it can still provide more useful information than such a degree-weighted uniform distribution for node ranking and clustering due to the subtly difference between them [50]. For example, when we consider the degree normalized vector  $\frac{\pi(s,u)}{d_u}$ , the simple baseline will be degraded as a constant

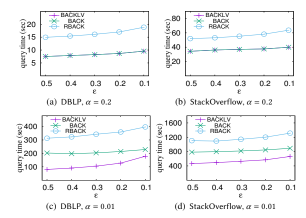


Figure 15: Runtime of different algorithms for answering the single target query on general weighted graphs

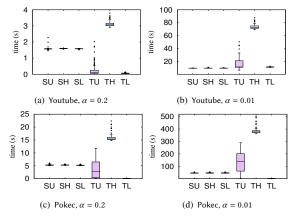


Figure 16: Query time distribution of single source query (SPEEDLV) and single target query (BACKLV). SU denotes single source query, choosing nodes uniformly. SH (SL) denotes single source query, choosing high (low) degree nodes uniformly. TU denotes single target query, choosing nodes uniformly. TH (TL) denotes single target query, choosing high (low) degree nodes uniformly.

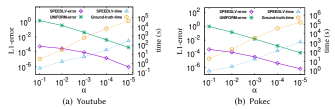


Figure 17: Results with very small  $\alpha$  (solid lines denote the  $L_1$ -error, and dashed lines represent the runtime)

vector which is indistinguishable for all nodes (1/2m for all nodes). However, as shown in [50], the degree-normalized PPR vector can still produce effective rankings and clusterings even when  $\alpha = 10^{-6}$ .

We vary  $\alpha$  from  $10^{-1}$  to  $10^{-5}$ , and use the state-of-the-art deterministic method in [49] to compute the ground-truth of the single source PPR vector to an L1-error bound  $10^{-9}$ . After that, we calculate the L1-error between SPEEDLV and the ground-truth PPR vector, and also compute the L1-error for the simple baseline. We randomly sample 50 nodes uniformly and take the average value the final result. The results on Youtube and Pokec are shown in Fig. 17. Similar results can also be observed on the other datasets. As can be seen, the L1-error of SPEEDLV is at least two orders of magnitude lower than that of the baseline method with varying  $\alpha$ . These results indicate that even for a very small  $\alpha$ , our algorithm can still produce much more accurate results than the baseline method. In addition, we can see that the L1-errors of both SPEEDLV and baseline decrease as  $\alpha$  decreases. The reason could be that the results of SPEEDLV, the baseline method, and the ground truth PPR converge to the degree-weighted uniform distribution when  $\alpha$  is very small, and thereby the L1-errors will be small. Fig. 17 also shows the time overheads for computing the ground truth and the time consumption by our SPEEDLV algorithm. We can see that the time costs by our algorithm are much lower than the time overheads for computing the ground truth. For a very small  $\alpha$  ( $\alpha \le 10^{-4}$ ), SPEEDLV is at least two orders of magnitude faster than the ground truth computation algorithm [49]. These results indicate that our SPEEDLV algorithm can achieve a very good trade-off between accuracy and runtime.

# 8 RELATED WORK

PageRank computation. Methods for computing personalized Page-Rank can be divided into two categories: deterministic algorithms and randomized approximate algorithms. For deterministic methods, there are many studies that focus on matrix-based power methods [25, 53]. Based on the power method, many different optimization techniques were proposed. [19, 20] applied the Cheyshev polynomials to accelerate the convergence rate. BEAR [39] preprocessed the adjacency matrix so that there is a large and easy-to-invert submatrix and also pre-computed several submatrix required to form an index. BePI [29] improved BEAR by using the power method instead of matrix inversion. TPA [52] was also an index-based iterative method which used PageRank value to approximate the nodes that are far from the source node. [34] developed a core-tree decomposition technique to further improve the efficiency of the power method. Also, there are a large number of local methods for computing personalized PageRank, notable examples including the forward push method [4, 10] and the backward push method [3, 28, 33]. Although

much progress has been made, deterministic methods are still slow for high-precision personalized PageRank computation.

For approximate methods, most of them are based on Monte Carlo simulation [7]. The idea of combining Monte Carlo and deterministic push method was first introduced in [32]. Much work follows this idea to improve different types of personalized PageRank queries. [44, 46] utilized the two-stage framework, which combines Monte Carlo and deterministic push, to answer the single source query. [31] and [49] further improved the single source query algorithm. [45] answered several new queries which aims to find heavy hitters in a graph based on the two-stage framework. However, in the Monte Carlo stage, all of the previous studies just simply simulate random walks. Unlike these studies, we propose an alternative method based on sampling of spanning forests which is shown to be more efficient than the random walk based sampling methods. Additionally, there also exist a number of algorithms to answer the top-k personalized PageRank query, which are also based on matrix operations [23, 25, 29], local methods [24, 27] and Monte Carlo techniques [8]. Specifically, matrix-based methods are based on the power method with a given absolute error bound  $\epsilon_a$ ; local methods conduct a local search from the source node while maintaining lower and upper bounds, and stops the search when the top-k results can be obtained by the lower and upper bounds; Monte Carlo techniques, including BiPPR [32], HubPPR [44] and FORA [46] are used for approximating the top-k PPR queries, which ensure a relative error  $\epsilon_r$  for any PPR value larger than 1/n, with probability at least 1 - 1/n. TopPPR [47] is the state-of-the-art algorithm which combines forward push, backward push and Monte Carlo together to answer the top-k query. Matrix forest theorem and spanning forest sampling. The Kirchhoff matrix tree theorem is perhaps the most classic result in spectral graph theory. Such a theorem has been generalized to spanning forest in early years [1, 11, 12, 38]. Most previous studies on the matrix forest theorem are based on the matrix L+qI where q is a constant [1]. Unlike the previous studies, we establish a new PageRank matrix forest theorem based on the  $\beta$ -Laplacian matrix  $(L_{\beta} = (\beta D)^{-1}(L + \beta D))$ . We note that Chung and Zhao also introduced a PageRank matrix forest theorem on undirected graphs [14, 15]. Their results are mainly based on the classic Cauchy-Binet formula which are hard to extend to directed graphs. Moreover, their matrix forest theorem is based

The algorithms for sampling spanning trees also have been heavily investigated [2, 26, 48]. The most well-known algorithms include (1) the Aldous-Broder algorithm [2] which simulates a random walk until the whole graph is covered; and (2) the Wilson algorithm [48] which simulates loop-erased random walks. Note that the concept of loop-erased random walk was also studied from the probability point of view [5, 6, 35]; and it was applied to generate spanning forests [5, 6] with an extended Wilson algorithm. Such an extended Wilson algorithm was also used for graph signal processing applications. [9, 37]. Unlike these work, we develop a loop-erased  $\alpha$ -random walk algorithm to sample spanning forests for personalized PageRank computation.

on the lazy random walk model, instead of the  $\beta$ -Laplacian matrix.

## 9 CONCLUSION

In this work, we develop several novel personalized PageRank matrix-forest theorems which connects the personalized PageRank values to the weights of spanning forests. Based on this connection, we propose a new personalized PageRank computation algorithm that samples spanning forests via simulating loop-erased  $\alpha$ -random walks on a graph. Compared to the previous algorithms, the proposed algorithm is shown to be more robust w.r.t. the parameter  $\alpha$ . This enable us to improve the efficiency of the state-of-the-art algorithms when  $\alpha$  is small. Specifically, by using our technique, we can significantly improve the efficiency of the state-of-the-art algorithms for answering two types of personalized PageRank queries, including single source and single target queries. Extensive experiments on 5 large real-life graphs demonstrate the efficiency of the proposed algorithms.

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