Computing Effective Resistances on Large Graphs Based on Approximate Inverse of Cholesky Factor

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Abstract-Effective resistance, which originates from the field of circuits analysis, is an important graph distance in spectral graph theory. It has found numerous applications in various areas, such as graph data mining, spectral graph sparsification, circuits simulation, etc. However, computing effective resistances accurately can be intractable and we still lack efficient methods for

ITERN STATE OF THE STATE OF THE ADART OF THE STATE OF is of two-dimensional grid structure. Despite of the importance of effective resistances, we still lack efficient methods for computing effective resistances on general large graphs.

The goal of power grid (PG) reduction is to reduce the original large power grid to a smaller one which can preserve the electrical behavior of port nodes. There are mainly three types of PG reduction methods in the literature: moment matching based methods [11], [12], node elimination based methods [13],

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[14] and multigrid-like methods [15], [16]. Moment matching based methods [11], [12] cannot reduce power grids efficiently because there are hundreds of thousands of port nodes in typical power grids. Node elimination based methods [13], [14] have been successful for reducing tree-like RC networks, but they tend to generate much denser models with even more edges than original models when dealing with mesh-like power grids. The multigrid-like methods [15], [16] can generate realizable and sparse reduced models, but their error is hard to control.

Effective resistances have been utilized for developing more efficient PG reduction methods [8], [9]. They employ the effective resistances based sampling approach [4] to sparsify the dense reduced models. The method proposed in [9] does not scale well to large problems due to the high computational complexity, as admitted in [9]. The method proposed in [8] is more scalable since it leverages the techniques of graph partitioning and effective resistances based port merging. However, it computes effective resistances accurately, which still takes a large amount of time for large-scale power grids.

In this paper, we aim to develop an efficient algorithm for computing effective resistances on large graphs and also a fast PG reduction method. Our main contributions are summarized as follows.

1) We propose an efficient algorithm for computing effective resistances on general weighted graphs, which is based on a sparse approximate inverse technique for the Cholesky factor of Laplacian matrix.

2) Incorporating the proposed algorithm for computing effective resistances with the PG reduction framework proposed in [8], we develop a fast PG reduction method.

Extensive experiments have been conducted to validate the efficiency and the accuracy of the proposed algorithm for computing effective resistances, which shows an average 168X speedup and also significant reduction in errors over the recent competitor [1]. The proposed algorithm is highly scalable. For a graph with 6.0E7 nodes and 1.0E8 edges, effective resistances of all edges can be computed within about 8 minutes, with an estimated average relative error of only 0.17%. It also derives a fast PG reduction method, which achieves an average 6.4X speedups over the method based on accurate effective resistances, with no increase in reduction errors. The resulted fast PG reduction method can be utilized in many downstream applications. For example, it brings 1.7X and 2.5X speedups of overall time for power grid transient analysis and DC incremental analysis, respectively.

II. BACKGROUND

A. Effective Resistances and Their Applications

Suppose G = (V, E, w) denotes a weighted undirected graph, where V and E are the sets of vertices (nodes) and edges, w is a positive weight function. Let n = |V| and m = |E|. The incidence matrix B is defined to be an $m \times n$ matrix such that each row of B corresponds to an edge in E and each column of B corresponds to a node in V. The entries of B satisfy:

$$B(e,v) = \begin{cases} 1, & v \text{ is } e's \text{ head} \\ -1, & v \text{ is } e's \text{ tail} \\ 0, & \text{otherwise} \end{cases}$$
(1)

Let W denote an $m \times m$ diagnoal matrix with W(e, e) = w(e). Then the Laplacian matrix $L_G \in \mathbb{R}^{n \times n}$ is defined as:

$$L_G = B^T W B {.} (2)$$

Given two nodes p and q, the *effective resistance* $R_G(p,q)$ across p and q refers to the voltage difference between p and q when a unit current flows into G through p and leaves through q. Formally, it can be defined as:

$$R_G(p,q) = e_{p,q}^T L_G^{\dagger} e_{p,q} . \tag{3}$$

Here L_G^{\dagger} denotes pseudo-inverse of L_G and $e_{p,q} = e_p - e_q$, where e_p is the *p*-th column of the identy matrix. L_G is singular as the smallest eigenvalue is 0. To handle the singularity of L_G , we introduce a ground node and connect it to a randomly selected node in each connected component of G. It corresponds to adding some small positive values to the diagonal elements of L_G . To simplify the notations, we still use L_G to denote the resulted symmetric diagonally dominant (SDD) matrix.

Below we briefly review the power grid reduction method proposed in [8], which employs effective resistances based graph sparsification.

Power grid reduction [17] aims to reduce the original large power grid to a small one which contains all the port nodes. A port node is defined to be a node which is connected to a voltage or current source. The voltage drops of port nodes can be obtained by analyzing the reduced model, enabling more efficient analysis for large power grids. Schur complement based method [14] is adopted to eliminate the non-port nodes without loss of accuracy but tends to generate much denser models. To address this issue, a graph sparsification based power grid reduction approach was introduced [8], which consists of circuits partitioning, Schur complement based reduction, effective resistances based port merging and effective resistances based graph sparsification. We remark that all the port nodes have important physical information and should be preserved. Although at least half of the port nodes were eliminated in [8], the algorithm can be easily extended to the case where all the port nodes should be kept. We summarize the modified version as Alg. 1.

Among all these steps, computing effective resistances is the most time-consuming one. So, to obtain a fast power

Algorithm 1 Power Grid Reduction via Effective Resistances Based Graph Sparsification [8]

Input: The original power grid.

Output: The reduced power grid.

- 1: Partition the original power grid into many blocks. The nodes are classified into three types: port nodes, non-port interface nodes and non-port interior nodes.
- 2: For each block, eliminate the non-port interior nodes using Schur complement based method.
- 3: For each reduced block, compute effective resistances exactly using (3) for all the edges.
- 4: For each reduced block, merge the nodes based on effective resistances and then sparsify the reduced grid using effective resistances based sampling approach.
- 5: Stitch all the reduced and sparsified models together to form the final reduced power grid.

grid reduction algorithm, more efficient methods for computing effective resistances are demanded.

B. Methods for Computing Effective Resistances

Computing effective resistances accurately is computational challenging. For each query (p,q), computing $R_G(p,q)$ requires solving linear equations whose coefficient matrix is L_G . It should be noted that many applications require computing effective resistances for every edge $(p,q) \in E$. Solving |E| linear equations takes at least $\Omega(|E|^2)$ time, which can be prohibitive for large-scale problems.

To compute effective resistances more efficiently, a random projection based method was proposed in [4]. It is based on the fact that the effective resistance $R_G(p,q)$ can be written as the distance between two vectors:

$$R_{G}(p,q) = e_{p,q}^{T} L_{G}^{\dagger} e_{p,q} = e_{p,q}^{T} L_{G}^{\dagger} L_{G} L_{G}^{\dagger} L_{G}^{\dagger} e_{p,q}$$

$$= e_{p,q}^{T} L_{G}^{\dagger} B^{T} W B L_{G}^{\dagger} e_{p,q}$$

$$= \| W^{1/2} B L_{G}^{\dagger} e_{p} - W^{1/2} B L_{G}^{\dagger} e_{q} \|_{2}^{2} .$$
(4)

For each node p, $W^{1/2}BL_G^{\dagger}e_p$ is an *m*-dimensional vector. With the Johnson-Lindenstraus Lemma, these vectors can be projected into a *k*-dimensional space, such that:

$$R_G(p,q) \approx \|QW^{1/2}BL_G^{\dagger}e_p - QW^{1/2}BL_G^{\dagger}e_q\|_2^2 , \quad (5)$$

where k = O(logm) and $Q \in \mathbb{R}^{k \times m}$ is a random matrix whose elements are uniformly sampled from $\pm \frac{1}{\sqrt{k}}$. Note that using (5), only k linear equations need to be solved to construct the matrix $QW^{1/2}BL_G^{\dagger}$ and then each effective resistance query can be answered in O(k) = O(logm) time. A more practical version of this algorithm was presented in [1], which incorporates the random projection based method and fast linear equation solver [18]. However, due to the big hidden constants, it still takes a huge amount of time to compute effective resistances on large graphs with reasonably high accuracy.

III. EFFICIENTLY COMPUTING EFFECTIVE RESISTANCES BASED ON APPROXIMATE INVERSE OF CHOLESKY FACTOR

A. The Idea

Suppose L_G is factorized with Cholesky factorization:

$$L_G = L L^T , (6)$$

where L is a lower triangular matrix. Then the effective resistance of query (p,q) can be written as:

$$R_G(p,q) = e_{p,q}^T L_G^{-1} e_{p,q} = e_{p,q}^T L^{-T} L^{-1} e_{p,q}$$

= $||L^{-1} e_{p,q}||_2^2 = ||L^{-1} e_p - L^{-1} e_q||_2^2$. (7)

Eq. (7) shows that the effective resistance $R_G(p,q)$ can be written as the distance between the p-th column and the q-th column of L^{-1} . If L^{-1} is available, the effective resistances can be computed efficiently, but computing and storing the dense L^{-1} explicitly can be prohibitive for large-scale problems. So, our idea is to derive a sparse and approximate inverse of L.

B. Sparse Approximate Inverse of Cholesky Factor

Based on the observation that most elements in L^{-1} are very small, we develop a method for computing a sparse approximation of L^{-1} . Let $Z = L^{-1}$ and z_j be the *j*-th column of Z. Recall that L is the Cholesky factor of L_G , it can be shown that all the diagonal elements in L are positive and all the off-diagonal elements in L are nonpositive [19].

The matrix Z has some useful structural properties which are summarized as the following lemma. Here a matrix is nonnegative means that all the elements in it are nonnegative.

Lemma 1. Suppose $Z = L^{-1}$, where L is the Cholesky factor of Laplacian matrix. Then, Z is nonnegative and the columns of Z satisfy:

$$z_j = \frac{1}{L_{j,j}} e_j + \sum_{i>j \& L_{i,j} \neq 0} \frac{-L_{i,j}}{L_{j,j}} z_i , \qquad (8)$$

Proof. Eq. (8) can be derived by multiplying L on both sides. Suppose z_i is nonnegative. Since $L_{i,j} \leq 0$ and $L_{j,j} > 0$, Eq. (8) implies z_j is nonnegative. So it can be proved by a simple mathematical induction that Z is nonnegative.

Eq. (8) suggests that we can first compute the *n*-th column of L^{-1} and then compute the (n-1)-th, (n-2)-th, \cdots , and the 1st columns one by one. Suppose we have some sparse approximations to z_i , denoted by \tilde{z}_i . Then z_i can be computed approximately by:

$$z_j \approx z_j^* = \frac{1}{L_{j,j}} e_j + \sum_{i>j \& L_{i,j} \neq 0} \frac{-L_{i,j}}{L_{j,j}} \tilde{z}_i$$
 (9)

The z_i^* can be computed efficiently because the \tilde{z}_i s are sparse. Note that many elements in z_i^* are very small, which can be set to 0. Here we adopt a prunning strategy to control the error in the sense of 1-norm. Note that for a vector x, the 1-norm of x, denoted by $||x||_1$, is defined as the sum of absolute value of each element. Let $trunc_k(x)$ denote the vector obtained by setting the k smallest elements (in the sense of absolute values) in x to 0. Our strategy is to find the largest k and to set $\tilde{z}_j =$ $trunc_k(z_i^*)$, such that:

$$\frac{\|\tilde{z}_j - z_j^*\|_1}{\|z_j^*\|_1} \le \epsilon , \qquad (10)$$

where ϵ is a user-defined threshold. It can be implemented by first sorting the nonzero elements in z_i^* and then finding the largest k which satisfies the above constraint. We summarize the algorithm for computing sparse approximate inverse of Cholesky factor as Alg. 2.

Algorithm 2 Sparse Approximate Inverse of Cholesky Factor **Input:** Cholesky factor of L_S : L, a user-defined threshold ϵ . **Output:** A sparse approximation to L^{-1} : Z.

1: **for** j = n to 1 **do** Compute $z_j^* = \frac{1}{L_{j,j}} e_j + \sum_{i>j \& L_{i,j} \neq 0} \frac{-L_{i,j}}{L_{j,j}} \tilde{z}_i$. if $nnz(z_j^*) \le \log n$ then 2: 3: $\tilde{z}_j = z_j^*.$ 4: 5: Continue. end if 6: Find the largest k such that $\frac{\|trunc_k(z_j^*)-z_j^*\|_1}{\|z_i^*\|_1} \leq \epsilon$. Set 7: $\tilde{z}_j = trunc_k(z_j^*).$ 8: end for

Now we analyze the errors caused by approximating L^{-1} with Z. Formally, we give a theorem below, which relates the approximation error $||z_p - \tilde{z}_p||_1$ to the depth of node p in the filled graph. The filled graph of G refers to the undirected graph corresponding to the matrix L [19]. Let $G_L = (V, F)$ denote the filled graph, where $F = \{(i, j) | i \neq j \text{ and } L_{i,j} \neq 0\}$. The depth of node p, denoted by depth(p), is defined as:

$$depth(p) = \begin{cases} 0, \quad L(p+1:n,p) = \mathbf{0} \\ 1 + \max_{i > p \& L(i,p) \neq 0} depth(i), \quad \text{otherwise} \ . \end{cases}$$
(11)

Theorem 1. Suppose L and $\tilde{Z} = [\tilde{z}_1, \tilde{z}_2, ..., \tilde{z}_n]$ are the input and output of Alg. 2. $Z = L^{-1} = [z_1, z_2, ..., z_n]$. The depth(p) is defined as (11). Then, for any node p:

$$\frac{\|z_p - \tilde{z}_p\|_1}{\|z_p\|_1} \le depth(p) \times \epsilon .$$
(12)

Proof. We prove the theorem by induction. First note that, for node q with L(q+1:n,q) = 0 or depth(q) = 0, we have:

$$z_q = z_q^* = \tilde{z}_q = \frac{1}{L_{q,q}} e_q ,$$
 (13)

so $||z_q - \tilde{z}_q||_1 = 0$, which satisfies (12). Now we consider node p with $L(p+1:n,p) \neq 0$. It is hypothesized that for node i with i > p and $L(i, p) \neq 0$, we have $\frac{\|z_i - \tilde{z}_i\|_1}{\|z_i\|_1} \leq depth(i) \times \epsilon$.

The error $||z_p - \tilde{z}_p||_1$ can be divided into two parts:

$$\begin{aligned} \|z_p - \tilde{z}_p\|_1 &= \|(z_p - z_p^*) + (z_p^* - \tilde{z}_p)\|_1 \le \|z_p - z_p^*\|_1 + \|z_p^* - \tilde{z}_p\|_1 . \end{aligned} \tag{14}$$

For the second part, Alg. 2 ensures that $\|z_p^* - \tilde{z}_p\|_1 \le \epsilon \times \|z_p^*\|_1.$
Recall that z_p is a vector with all elements nonnegative and z_p^* is a truncated version of z_p , it is easy to show $\|z_p^*\|_1 \le \|z_p\|_1$, so we have:

$$||z_p^* - \tilde{z}_p||_1 \le \epsilon \times ||z_p||_1 .$$
(15)

Now consider the first part. Recall that:

2

$$z_p^* = \frac{1}{L_{p,p}} e_p + \sum_{i > p \& L_{i,p} \neq 0} \frac{-L_{i,p}}{L_{p,p}} \tilde{z}_i .$$
 (16)

Here $L_{i,p} < 0$, $L_{p,p} > 0$ and $\sum_{i} \frac{-L_{i,p}}{L_{p,p}} \leq 1$. In the rest of this proof, we use \sum_{i} to replace $\sum_{i>p\&L_{i,p}\neq 0}$. Then

$$\|z_p - z_p^*\|_1 = \|\sum_i \frac{-L_{i,p}}{L_{p,p}} (z_i - \tilde{z}_i)\|_1 \le \sum_i \frac{-L_{i,p}}{L_{p,p}} \|(z_i - \tilde{z}_i)\|_1.$$
(17)

By the inductive hypothesis, we have

$$||z_p - z_p^*||_1 \le \sum_i \frac{-L_{i,p}}{L_{p,p}} ||z_i||_1 \times depth(i) \times \epsilon$$

$$\le \max_i depth(i) \times \epsilon \times \sum_i \frac{-L_{i,p}}{L_{p,p}} ||z_i||_1 .$$
(18)

Note that for nonnegative vectors z_i s, we have

$$\sum_{i} \frac{-L_{i,p}}{L_{p,p}} \|z_i\|_1 = \|\sum_{i} \frac{-L_{i,p}}{L_{p,p}} z_i\|_1 \le \|z_p\|_1 .$$
(19)

Substituting (19) into (18), we obtain

$$||z_p - z_p^*||_1 \le \max_i depth(i) \times \epsilon \times ||z_p||_1 .$$
 (20)

So

$$\frac{\|z_p - \tilde{z}_p\|_1}{\|z_p\|_1} \le (\max_i depth(i) + 1) \times \epsilon = depth(p) \times \epsilon .$$
(21)

By mathematical induction, Eq. (12) is true for all nodes. This ends the proof. $\hfill \Box$

Note that for most graphs that stem from real world, the maximum node depth in the filled graph is not very large, as reported in the next section. By setting a sufficiently small ϵ , \tilde{Z} approximates L^{-1} very well.

C. The Overall Algorithm and Discussion

Based on the approximate inverse technique, the effective resistance $R_G(p,q)$ can be computed as:

$$R_G(p,q) = \|L^{-1}e_p - L^{-1}e_q\|_2^2 \approx \|\tilde{z}_p - \tilde{z}_q\|_2^2 .$$
 (22)

Now we analyze the errors of effective resistances. Let $z_{p,q} = z_p - z_q$, $\tilde{z}_{p,q} = \tilde{z}_p - \tilde{z}_q$, $\Delta z_p = \tilde{z}_p - z_p$, $\Delta z_q = \tilde{z}_q - z_q$ and $\Delta z_{p,q} = \tilde{z}_{p,q} - z_{p,q}$. Eq. (12) indicates that:

$$\begin{aligned} \|\Delta z_{p,q}\|_{1} &\leq \|\Delta z_{p}\|_{1} + \|\Delta z_{q}\|_{1} \\ &\leq (\|z_{p}\|_{1} depth(p) + \|z_{q}\|_{1} depth(q))\epsilon \end{aligned}$$
(23)

We can assume that $\|\Delta z_{p,q}\|_1$ is very small and $\Delta z_{p,q}$ is close to **0**. Then by ignoring the second-order terms, we have:

$$\|\tilde{z}_{p,q}\|_{2}^{2} = \|z_{p,q} + \Delta z_{p,q}\|_{2}^{2} \approx \|z_{p,q}\|_{2}^{2} + 2z_{p,q}^{T}\Delta z_{p,q} .$$
(24)

Let $\hat{R}_{p,q} = \|\tilde{z}_{p,q}\|_2^2$ denote the approximate effective resistance, then we obtain:

$$\begin{aligned} |\frac{\tilde{R}_{p,q}}{R_{p,q}} - 1| &\approx |\frac{2z_{p,q}^{T}\Delta z_{p,q}}{\|z_{p,q}\|_{2}^{2}}| \leq \frac{2\|z_{p,q}\|_{1}\|\Delta z_{p,q}\|_{1}}{\|z_{p,q}\|_{2}^{2}} \\ &\leq \frac{2\|z_{p,q}\|_{1}(\|z_{p}\|_{1}depth(p) + \|z_{q}\|_{1}depth(q))}{\|z_{p,q}\|_{2}^{2}}\epsilon . \end{aligned}$$

If we denote $\alpha_{p,q} = \frac{2\|z_{p,q}\|_1(\|z_p\|_1 depth(p) + \|z_q\|_1 depth(q))}{\|z_{p,q}\|_2^2}$, then

$$1 - \alpha_{p,q}\epsilon \le \frac{\dot{R}_{p,q}}{R_{p,q}} \le 1 + \alpha_{p,q}\epsilon .$$
⁽²⁶⁾

This implies that the relative error of effective resistance scales linearly with the parameter ϵ . The smaller ϵ is, the closer $\tilde{R}_{p,q}$ is to $R_{p,q}$.

Computing effective resistances using (22) requires Cholesky factorization on L_G . However, for large-scale graphs, especially for graphs that stem from social networks, Cholesky factorization on L_G can be very expensive. In this work, we propose to use incomplete Cholesky factorization instead. In incomplete Cholesky factorization, some fill-ins with very small absolute values are dropped, which corresponds to set some branches with large resistances to open and does not introduce large errors to effective resistances. We summarize the overall algorithm for computing effective resistances as follows.

Algorithm 3 Computing Effective Resistances Based on Sparse Approximate Inverse of Cholesky Factor

Input: A weighted undirected graph: G, a set of effective resistance queries Q_r .

Output: Effective resistances for each query in Q_r .

- 1: Run incomplete Cholesky factorization on L_G to obtain: $L_G \approx LL^T$.
- 2: Compute the sparse and approximate inverse with Alg. 2 for L: $\tilde{Z} \approx L^{-1}$.
- 3: for each query (p,q) in Q_r do
- 4: Compute effective resistance: $R_G(p,q) \approx \|\tilde{z}_p \tilde{z}_q\|_2^2$.

The time complexity of the proposed algorithm is closely related to the number of nonzeros in \tilde{Z} . In our experiments, the number of nonzeros in \tilde{Z} is about Cnlogn where C is a small constant (e.g. C < 20). The reason behind this phenomenon may be the decay property of L^{-1} [20]. Here we just assume the average number of nonzeros in one column of \tilde{Z} is O(logn). Incomplete Cholesky factorization takes O(n) time and the number of nonzeros in L is O(n). Each iteration of Alg. 2 takes $O(\frac{nnz(L)}{n}logn + logn \cdot loglogn) = O(logn \cdot loglogn)$ time and the time complexity of Alg. 2 is $O(nlogn \cdot loglogn)$. So the overall time complexity of Alg. 3 is $O(nlogn \cdot loglogn) + O(|Q_r|logn)$.

IV. NUMERICAL RESULTS

We first compare the proposed algorithm for computing effective resistances (Alg. 3) with the algorithm proposed in [1]. Because the codes shared by the authors of [1] are written in MATLAB, we also implement our Alg. 3 in MATLAB. Then we implement the graph sparsification based power grid reduction (Alg. 1) and compare the scenarios where effective resistances are computed accurately, approximately using the random projection based method [1] and using the proposed Alg. 3. Finally, the resulted fast power grid reduction algorithm is leveraged to solve problems of DC incremental analysis and transient analysis. These programs are written in C++. All experiments are conducted using a single CPU core of a computer with Intel Xeon E5-2630 CPU @2.40 GHz and 256 GB RAM.

A. Results on Computing Effective Resistances

In this subsection, we compare the proposed algorithm (Alg. 3) with the random projection based method [1], whose

results are obtained by running the codes shared on [21]. We do not compare the algorithms in [2], [3] because these algorithms can only handle unweighted graphs. The results are listed in Table I. The test cases cover a great variety of graphs obtained from social networks, finite element analysis and circuit simulation problems [22]–[24]. For each case, effective resistances of all edges are computed, i.e. the set of effective resistance queries $Q_r = E$. T denotes the runtime for computing effective

 TABLE I

 Results for Computing Effective Resistances on Large Graphs.

Case	V (E)	dpt	WWW15 [1]				Alg. 3			
Case	V (<i>L</i>)	upt	$\overline{T(s)}$	E_a	E_m	$\frac{nnz(Q)}{nlogn}$	T(s)	E_a	E_m	$\frac{nnz(Z)}{nlogn}$
com-DBLP	3.2E5(1.0E6)	464	517	2.6E-2	1.4E-1	108	4.14	7.1E-5	1.9E-3	5.40
com-Amaz	3.3E5(9.3E5)	590	719	2.2E-2	1.4E-1	149	4.71	8.0E-5	3.9E-3	7.47
com-Yout	1.1E6(3.0E6)	1370	926	3.5E-2	2.1E-1	32.6	21.0	1.5E-4	2.1E-2	1.63
coAuDBLP	3.0E5(1.0E6)	1040	513	2.5E-2	1.1E-1	108	3.87	7.1E-5	4.0E-3	5.43
coAuCite	2.3E5(8.1E5)	774	414	2.4E-2	1.0E-1	129	2.32	5.6E-5	7.9E-3	6.45
fe_tooth	7.8E4(4.5E5)	1892	322	1.8E-2	7.4E-2	304	1.73	8.6E-4	1.1E-2	15.2
fe_rotor	1.0E5(7.6E5)	2448	488	1.7E-2	7.0E-2	344	2.84	8.3E-4	2.1E-2	17.2
NACA0015	1.0E6(3.1E6)	543	2447	2.2E-2	7.5E-2	163	12.1	1.0E-3	3.6E-3	8.17
ibmpg5	1.1E6(1.6E6)	513	691	2.2E-2	1.2E-1	123	3.16	1.7E-3	2.7E-2	6.17
ibmpg6	1.7E6(2.5E6)	602	934	2.3E-2	1.2E-1	109	4.64	1.8E-3	2.2E-2	5.44
thupg1	5.0E6(8.2E6)	1097	7158	1.8E-2	8.1E-2	122	36.6	1.7E-3	1.4E-2	6.10
G2_circuit	1.5E5(2.9E5)	720	214	2.0E-2	1.2E-1	166	1.15	1.3E-3	4.4E-2	8.30
G3_circuit	1.6E6(3.1E6)	1237	2388	2.0E-2	9.8E-2	140	13.3	3.1E-3	4.0E-2	7.02
thupg10	6.0E7(1.0E8)	3725	-	-	-	-	481	1.7E-3	1.7E-2	1.24

resistances. E_a and E_m denote the average and the maximum relative errors, which are estimated by randomly selecting 1000 edges, computing accurate effective resistances for these edges and then computing the errors. For Alg. 3, T includes the time for incomplete Cholesky factorization, computing approximate inverse and computing effective resistances. The drop tolerance in incomplete Cholesky factorization is set to 1E-3 and the parameter ϵ in Alg. 2 is also set to 1E-3. We also record the maximum depth of the filled graph, which is defined in the last section and denoted by dpt, and the number of nonzeros in the random projection matrix Q and in the appproximate inverse matrix \tilde{Z} , both of which are divided by nlogn. "-" means that it takes more than 10 hours.

From the results we see that, the proposed Alg. 3 achieves an average **168X** speedup over the random projection based method [1]. Although the number of nonzeros in the random projection matrix Q is much larger than that in the approximate inverse matrix \tilde{Z} , the proposed algorithm shows one to two orders of magnitude improvement in the avarage relative error and also significant reduction in the maximum relative error. For the largest case named "thupg10", with 6.0E7 nodes and 1.0E8 edges, effective resistances for up to 1.0E8 node pairs can be computed within just 481 seconds, while the average relative error is about 0.17%, which demonstrates the extremely high scalability of the proposed algorithm.

B. Results on Graph Sparsification Based PG Reduction

Combining the proposed algorithm for computing effective resistances (Alg. 3) with the graph sparsification based power grid reduction framework (Alg. 1), we obtain a fast power grid reduction algorithm. Test cases are from the well-known IBM power grid benchmarks [23]. For graph partitioning, we use the widely adopted graph partitioner METIS [25] and the number of blocks are set to $\frac{\#ports}{50}$. For transient analysis, each case is simulated for 1000 fixed-size time steps and both original

models and reduced models are analyzed with the direct solver CHOLMOD [26] (performing just once matrix factorization). For DC incremental analysis, 10% blocks of each benchmark are modified to mimick the design process where the initial power grid is modified to fix violations. We compare three power grid reduction methods, which only differ in computing effective resistances. The first method computes effective resistances accurately, while the other two compute effective resistances approximately, using the random projection based approach [1] and the proposed approach (Alg. 3) respectively.

The results are listed in Table II. |V| and |E| denote the number of nodes and resistors. T_{red} denotes the time for power grid reduction. Note that in the application of DC incremental analysis, only the modified blocks need to be reduced, so the T_{red} in incremental analysis is just about 10% of that in transient analysis. T_{tr} and T_{inc} are the time for transient analysis and DC incremental analysis, respectively. Err is the average absolute error and Rel is the relative error, which is computed by dividing the Err by the maximum voltage drop.

From the results we see that, with the proposed algorithm for computing effective resistances (Alg. 3), the time for power grid reduction can be reduced largely, showing an average 6.4X speedup over the power grid reduction approach based on accurate effective resistances. In terms of the reduction accuracy, there is almost no increase in reduction errors. This validates the efficiency and the effectiveness of the proposed power grid reduction approach. On the other hand, with the random projection based method, it still takes a large amount of time to generate reduced models. Meanwhile, the reduction accuracy is also deteriorated, which is due to the large errors of effective resistances.

As for the total time of transient analysis, which includes the time for power grid reduction and the time for analyzing the reduced power grid, the proposed method achieves 1.7X speedup averagely over the power grid reduction approach based on accurate effective resistances. Compared with analyzing the original power grid, the proposed method shows an average 2.9X speedup, with the relative error below 1.51% for all cases. The transient waveforms of node n0_20706300_8937900 and node n1_29561400_9521100 in case "ibmpg3t" are plotted in Fig. 1. They validate the accuracy of transient simulation using the proposed fast power grid reduction method.

In the application of DC incremental analysis, the proposed method shows significant improvement in the incremental power grid reduction stage, thus achieves an average 2.5X speedup for the total time over the power grid reduction method based on accurate effective resistances. Compared with analyzing the modified power grid directly using CHOLMOD ("Original" in Table II), the proposed method shows 4.2X speedups on average, while the relative error is below 1.34% for all cases.

V. CONCLUSIONS

In this paper, we propose an efficient algorithm for computing effective resistances on massive weighted graphs. The proposed algorithm is based on a sparse approximate inverse technique for Cholesky factors and achieves 168X speedups on

 TABLE II

 Results on Graph Sparsification Based Power Grid Reduction for Transient Analysis (upper) and DC Incremental Analysis (lower).

Case Original	w/ Acc. Eff. Res.	W	/ App. Eff. Res. ([1])	w/ App. Eff. Res. (Alg. 3)		
Case $ V (E) = T_{tr}$	$ V (E)$ T_{red} T_{tr} Err(mV)	Rel(%) $V (E)$	T_{red} T_{tr} Err(mV) Rel($\frac{(W)}{W} = \frac{ V (E) - T_{red} - T_{tr} - \text{Err}(\text{mV}) \text{ Rel}(\%)}{V(E) - T_{red} - T_{tr} - \text{Err}(\text{mV}) - \text{Rel}(\%)}$		
ibmpg2t 1.3E5(2.08E5) 13.3	5.1E4(1.49E5) 6.55 4.33 0.801	1.52 5.1E4(1.47)	E5) 3.51 4.23 2.300 4.2	8 5.1E4(1.49E5) 0.951 4.20 0.796 1.51		
ibmpg3t 8.5E5(1.40E6) 201	3.0E5(8.59E5) 67.2 35.1 0.153	0.78 3.0E4(8.54)	E5) 29.4 34.7 0.253 1.2	9 3.0E4(8.59E5) 7.70 35.8 0.162 0.83		
ibmpg4t 9.5E5(1.55E6) 310	4.0E5(1.35E6) 81.9 132 0.006	0.93 4.0E5(1.35)	E6) 36.3 133 0.034 4.8	5 4.0E5(1.35E6) 10.6 129 0.006 0.93		
ibmpg5t 1.1E6(1.62E6) 141	5.1E5(1.04E6) 24.1 48.0 0.124	0.87 5.1E5(1.03	E6) 21.5 47.4 0.137 0.9	6 5.1E5(1.04E6) 5.59 48.8 0.125 0.87		
ibmpg6t 1.7E6(2.48E6) 179	7.7E5(1.64E6) 39.4 67.7 0.173	1.02 7.7E5(1.66)	E5) 33.5 67.0 0.334 1.9	7 7.7E5(1.64E6) 8.76 67.5 0.173 1.02		
Case Original	w/ Acc. Eff. Res.	W	/ App. Eff. Res. ([1])	w/ App. Eff. Res. (Alg. 3)		
Case $ V (E) = T_{inc}$	$- \frac{ V (E)}{ V_{red} } T_{red} T_{inc} \operatorname{Err}(\mathrm{mV})$) Rel(%) $V (E)$	T_{red} T_{inc} Err(mV) Rel	$(\%) V (E) T_{red} T_{inc} \text{Err(mV) Rel(\%)}$		
ibmpg2 1.3E5(2.08E5) 0.627	5.1E4(1.49E5) 0.784 0.159 6.052	1.21 5.1E4(1.47E	(5) 0.381 0.149 14.10 2.	31 5.1E4(1.49E5) 0.115 0.158 6.020 1.20		
ibmpg3 8.5E5(1.40E6) 20.4	3.0E5(8.59E5) 6.73 1.54 1.612	0.66 3.0E4(8.54E	25) 2.66 1.46 3.004 1.	22 3.0E4(8.59E5) 0.769 1.57 1.734 0.71		
ibmpg4 9.5E5(1.55E6) 36.3	4.0E5(1.35E6) 8.85 11.7 0.093	0.76 4.0E5(1.35E	6) 3.94 11.5 1.159 9.	45 4.0E5(1.35E6) 1.04 11.6 0.089 0.73		
ibmpg5 1.1E6(1.62E6) 10.5		1.34 5.1E5(1.03E	6) 2.11 1.74 5.257 7.			
ibmpg6 1.7E6(2.48E6) 8.68	7.7E5(1.64E6) 3.98 2.42 1.503	0.73 7.7E5(1.66E	5) 3.53 2.45 3.181 1.	54 7.7E5(1.64E6) 0.888 2.41 1.512 0.73		

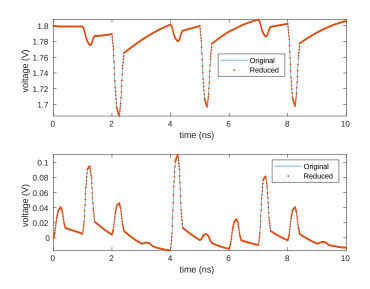


Fig. 1. The transient simulation results of a VDD node (up) and a GND node (down) in case "ibmpg3t", obtained by analyzing the original power grid and the reduced power grid.

average and also significant reduction in errors, compared with the random projection based method. Combining the proposed algorithm with the graph sparsification based power grid reduction framework, we obtain a fast power grid reduction method, which shows 6.4X speedups averagely and almost the same reduction accuracy. Utilized for the downstream applications of power grid transient analysis and incremental analysis, the proposed fast power grid reduction method brings 1.7X and 2.5X speedups of overall time, compared to using the reduction method based on accurate effective resistances. Given the wide range of applications of effective resistances, we hope that this work can promote progress in more fields in the future.

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