NetKI: A kirchhoff index based statistical graph embedding in nearly linear time

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A B S T R A C T
Recent advancements in learning from graph-structured data have shown promising results on the graph classification task. However, due to their high time complexities, making them scalable on large graphs, with millions of nodes and edges, remains a challenge. In this paper, we propose NetKI, an algorithm to extract sparse representation from a given graph with $n$ nodes and $m$ edges in $O(m + \log n)$ time. Our approach follows the notion of Kirchhoff index that encodes the structure of the graph by estimating effective resistance - relying on this approach yields nearly linear time graph representation method that allows scalability on sufficiently large graphs. Through extensive experiments, we show that NetKI provides improved results in terms of running time on large networks and the classification accuracy is within range 2% from the state-of-the-art results.

1. Introduction

Many real-world systems involve interactions between pairs of entities. Examples of such systems include social networks, financial systems, molecular graph structures, resistor networks, recommender systems, and protein–protein interaction networks. All of these systems and many more can be represented as graphs, that encode real-world objects (i.e., nodes) and the pair-wise interactions (i.e., edges) between them. More recently, graphs have also been used to model grid-structured data, such as images, to make the computations faster [1]. Encoding graphs into low-dimensional embeddings enables one to employ the traditional machine learning and data mining algorithms on graph-structured data. This approach facilitates researchers to solve hard problems on graphs such as graph classification. The graph classification problem refers to the understanding of complex graph structures among different classes, and it has a variety of real-world applications ranging from text classification to molecules toxicity prediction and classifying community structures in a social network [2]. However, graph classification poses several challenges such as permutation invariance, scalability, and, the runtime efficiency of the encoding. Generally, the same graph can be represented by many adjacency matrices due to a lack order on nodes of a graph, therefore, the ideal encoding procedure for graph classification should be invariant under permutations of the nodes. Moreover, it should preserve the atomic structure of the graph based on the local and global positions of pairs of nodes. Unfortunately, existing graph classification approaches often require a pairwise comparison among the graphs or are based solely on a statistical and spectral representations, which is hard to compute [3–6]. Therefore, appropriate representation methods are required to encode the atomic structure of the graph succinctly that are efficient. To address the aforementioned challenges, in this paper, we propose using a well-known measure called the Kirchhoff index for extracting graph representations. The Kirchhoff index encapsulates the atomic structure of the graph because it incorporates the information regarding the number of paths and quality of paths. It features in many other contexts, such as Markov chains (the average commute time of a Markov chain on the graph), experiment design, and Euclidean distance embeddings [7,8]. Recently, it has also been employed to quantify the resilience of networks based on noisy data in distributed networked control systems [9]. In fact, it has more equivalent descriptions including the Laplacian spectrum and the effective resistance which makes it suitable for extracting graph representations. Typically, Kirchhoff index is defined in terms of effective resistance using the analogy of graphs in an electrical circuit. A graph can be transformed into an electrical circuit by replacing each edge with a unit resistance.
The effective resistance between any two nodes in such a network refers to the equivalent electrical resistance between those two nodes. The sum of the effective resistance between all pair of nodes is referred to as the Kirchhoff index of a graph. We note that a higher value of effective resistance between nodes indicates an inferior or less robust overall connection between the nodes (as measured by the number and quality of paths between them) [8,10,11,7,12]. Since there is always a path between two nodes in a connected network (that is, any two nodes are always connected, maybe via some other nodes), the effective resistance between any two nodes is well defined. Consequently, the effective resistance of a graph, which is the sum of all pair-wise effective resistances between nodes, is also well defined. To illustrate this, consider the example in Fig. 1.

In (b), there are four paths between nodes 1 and 2 with lengths 1, 3, 4, and 5, and the effective resistance between the nodes is 0.72. In (c), there are three paths between nodes 1 and 2 with lengths 1, 3, and 5, and the effective resistance between is 0.73. We see only a slight increase in effective resistance (indicating a slightly less robust connection) as the path of length four does not exist anymore. The effect is minimal because the paths of smaller lengths, which are more important, are still there. In (d), there are only two paths between nodes 1 and 2 with lengths 3, and 5. The edge between nodes 1 and 2, hence the path of length 1, is removed. The effective resistance between the nodes increases significantly as the path with the shortest length having the maximum effect is deleted. In (e), there is only one path of length 3 between nodes 1 and 2, which increase the effective resistance from 2.75 to 3.0. Thus, we observe that effective resistance between two nodes (adjacent or non-adjacent) measure robustness in terms of the number and quality (length) of paths between nodes. Since Kirchhoff index measures the number as well as the quality of paths, we argue in this paper that this is a suitable measure to generate meaningful and expressive graph representations.

We empirically show that proposed method provides improved results in terms of running time on large networks while the classification accuracy is close to the state-of-the-art methods that are, otherwise, impractical for large graphs.

We are making our implementation publicly available for other researchers to use and improve upon.

2. Related work

Our related work section is divided into four categories: direct methods, graph kernel methods, statistical and spectral representations, and graph-theoretic approaches. The methods in the first three categories deal with graph representations, while the last category surveys the relevant graph-theoretic approaches used or referred in this work. Table 1 summarizes the related work’s necessary properties and time complexity.

2.1. Direct methods

Direct methods seek to extract features from graphs directly [3,4,15,16,5,17]. The Graph Edit Distance (GED) [3] approach is built upon the minimal number of modification needed to transform one graph into another. However, the problem is NP-hard even when the goal is to approximate the distance. A group-theoretic approach, the graphlet spectrum [4] characterizes graphs in terms of the position or frequency of common graphlets or motifs. Some other well-known approaches include propagation models [18], family of tractable distances [19] and heuristics-based approaches [20,21]. However, all of these methods require extensive computational power to run on any decent size graph dataset.

2.2. Kernel methods

Graph kernel methods are one of the most popular categories of graph classification. In general, these methods define a distance between pairs of elementary substructures of two graphs, and some function of the resulting matrix is the representation of distance between the two given graphs. [15,22–25,3,19,26,18,20]. Deep graph kernels [15] compute substructures or motifs of a graph, and use a word-embedding model to extract the underlying features. In order to extract both local and global-scale graph features, Kondor et al., [16] introduced the Multiscale Laplacian Graph Kernel (MLG) which captures nodes’ topological information.
through graph Laplacian. Other popular graph kernels include the Weisfeiler-Lehman kernel (WL) [24], shortest path kernel (SP) [22], edge histogram kernel (EHK) [27], neighbourhood hash kernel (NHK) [28], graphlet sampling kernel (GK) [29] and graph invariant kernel [25]. However, these methods suffer from higher time complexity. In order to capture the distribution of an individual component of the graph, recently [30] proposed a distance measure for graphs: the graph Wasserstein distance and Weisfeiler-Lehman-inspired embedding scheme derived from optimal transport theory. The proposed measure uses node feature representations and weighted edges to enhance the quality of the graph embeddings. Similarly, [31] introduced indefinite graph kernel for time series analysis by using the power of optimal transport theory.

The computational overhead of these methods makes them impractical for large-scale graph classification. The proposed measure uses node feature representations and weighted edges to enhance the quality of the graph embeddings. Similarly, [31] introduced indefinite graph kernel for time series analysis by using the power of optimal transport theory. The computational overhead of these methods makes them impractical for large-scale graph classification.

2.3. Statistical and spectral representations

Statistical and spectral representation methods use a graph’s statistical properties and graph spectrum to generate graph feature vectors [32,33,5,6]. Initial works in this area [32,33] use summary graph-statistics and handcrafted features such as nodes’ degrees to extract local graph properties. Recently, Verma and Zhang [5] introduced a histogram-based representation method, known as the Family of Graph Spectral Distances (FGSD). This approach based on computing multisets of spectral distances between all pairs of nodes using eigendecomposition. The authors in [6] introduced a spectral representation method, based on wave- and heat trace signatures at various times, and consider the associated heat diffusion process on the graph rather than use the Laplacian spectrum to obtain graph representations. Recent approaches [5,6] have shown promising results on graph comparison. However, their time complexities make them impractical for reasonably large graphs.

Under the assumption of the same number of nodes in the graphs, [34] recently propose a graph comparison framework that uses Wasserstein distance is mean to compare graphs. The measure uses smooth graph signal distributions associated with graphs and compares them using the Wasserstein distance. Similarly, [35] uses optimal transport theory and introduces a graph comparison framework based on empirical distribution with a graph-based regularization. The main idea of the work was to compute an optimal transportation plan by limiting the displacement of a pair of nodes. Graph-based regularization [36] approach finds the similarity between adjacent nodes by their position or displacements of the transported sample. Instead of matching graph feature vectors, [37] propose to use a Wasserstein distance to compare between them. [38] uses separable functions to approximate discrete graph matching in continuous domain. Likewise, [39,40] use Gumbelsinkhorn network to extract permutations from data. Closer to our work, the authors in [41] propose a graph streaming algorithm that doesn’t require the whole graph in memory and approximate feature vector by estimating counts of sub-graph. Some other distance measures on graphs include [42–44,40]. Many of these approaches have shown promising results on the graph classification task, however, due to their computational complexities, scaling them on sufficiently large graphs is still a challenging task.

With the advancement in deep learning approaches especially the convolutional neural networks and recurrent neural networks, recently there has been a surge in neural models for learning graph representations [45–48,1]. Graph neural models are end-to-end learning approaches based on neighborhood aggregation and recurrent neural networks [49–51].

2.4. Graph-theoretic methods

Li and Zhang [14] introduced nearly-linear time algorithms for finding nodes’ and edges’ importance in a given network. To find edge importance, the authors deployed the Sherman-Morrison formula and used two graph-theoretic approaches to attain the required goal. These methods will be discussed in details in forthcoming section, as our method is based on this idea. Spielman and Srivastava constructed spectral graph sparsifier in O(m) and proved the existence of sparse graph with O(n log n/c²) edges [54]. To construct the sparsifier, this method uses the notion of graph effective resistance approximated by solving graph Laplacian. The breakthrough results by Spielman and Teng show that it can be solved in O(mlog⁴n log log(1/δ)) time [55], Kyng and Sachdeva [13] introduced a nearly-linear time algorithm that approximates cholesky factorization for Laplacian matrices [56]. This algorithm is based purely on random sampling and Gaussian elimination to solve linear systems in graph Laplacians. Recently, Cohen et al.[57] introduced a sub-sampling iterative method (currently the best-known Laplacian solver) based on recursive preconditioning, and this aims to reduce graphs to trees. Many other follow-up works [58–61] have extended the field in the past decade, and, currently the best known algorithm has O(mlog⁴n log log n log(log(1/δ))) time complexity. Although [57] is the best-known Laplacian solver, we employed Kyng and Sachdeva’s [13] method in our experimental setup, since

<table>
<thead>
<tr>
<th>Method</th>
<th>Type</th>
<th>Labels</th>
<th>Attr.</th>
<th>Perm. Invar.</th>
<th>Complexity</th>
</tr>
</thead>
<tbody>
<tr>
<td>GED [3]</td>
<td>direct</td>
<td>+</td>
<td>–</td>
<td>+</td>
<td>NP-complete</td>
</tr>
<tr>
<td>FTD [19]</td>
<td>direct</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>O(n²)</td>
</tr>
<tr>
<td>RW [52]</td>
<td>kernel</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>O(n³)</td>
</tr>
<tr>
<td>SP [22]</td>
<td>kernel</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>O(dhD)</td>
</tr>
<tr>
<td>NHK [28]</td>
<td>kernel</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>O(hm)</td>
</tr>
<tr>
<td>WL [24]</td>
<td>kernel</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>O(k³)</td>
</tr>
<tr>
<td>MLG [16]</td>
<td>kernel</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>FGSD [5]</td>
<td>stat. rep.</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>O(n²)</td>
</tr>
<tr>
<td>NetKI</td>
<td>stat. rep.</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>NetLS1D [6]</td>
<td>spec. rep.</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>O(km ± k²n)</td>
</tr>
<tr>
<td>GCN [47]</td>
<td>GCNs</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>GraphSAGE [53]</td>
<td>GCNs</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>GIN [49]</td>
<td>GCNs</td>
<td>–</td>
<td>+</td>
<td>+</td>
<td>–</td>
</tr>
<tr>
<td>DGI [50]</td>
<td>GCNs</td>
<td>–</td>
<td>+</td>
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<td>–</td>
</tr>
</tbody>
</table>
it is simple and requires no graph theoretic constructions. This enabled us to solve a much broader family of linear systems. Additionally, the code of the solver is freely available\(^1\), and we used it in our experimental setup.

### 3. Preliminaries and basic notations

#### 3.1. Graph Basics

Let \( G = (V, E) \) denote an undirected connected graph where \( V = \{v_1, v_2, \ldots, v_n\} \) is the set of \( n \) nodes, and \( E \subseteq V \times V \) is the set of \( m \) edges. Throughout this study, we consider \( G \) as a resistor network where nodes represent junctions while edges represent resistors between the junctions. Let \( r_e \) denote a unit resistance over the edge \( e \). We write \( G(e) \) to represent the graph obtained from \( G \) by deactivating edge \( e \) by decreasing \( r_e \) to \( 0 \) for some small \( 0 < \theta < 1/2 \). We denote this process by \( \theta \)-deletion. Let \( L \) denote the \( n \times n \) Laplacian matrix for \( G \) where,

\[
L_{i,j} = \begin{cases} 
  k_u & \text{if } u = v \\
  -1 & \text{if } u \sim v \\
  0 & \text{otherwise}.
\end{cases}
\]  

(1)

Where \( k_u \) is the degree of node \( u \), and \( (u \sim v) \) indicate that node \( u \) and \( v \) are neighbors. Let \( L^{0i} \) denote the Laplacian of the graph after the deactivation of edge \( e \). Let \( W \) be an \( m \times m \) diagonal matrix with \( W[i,j] = w(e_{ij}) \) (\( e_{ij} \) indicates an edge from the edge set \( \{e_1, e_2, \ldots, e_m\} \subseteq E \)). Similarly, let \( B_{m \times n} \) denotes a signed edge-vertex incident matrix where each entry is

\[
B_{j,i} = \begin{cases} 
  1 & \text{if } v \in e_i \text{ is head} \\
  -1 & \text{if } v \in e_i \text{ is tail} \\
  0 & \text{otherwise}.
\end{cases}
\]  

(2)

Then the Laplacian matrix \( L = B^{T}WB \) [62]. Let \( L^{+} \) represents the pseudo-inverse of the Laplacian and \( e \) denote the \( i^{th} \) standard basis vector or one-hot encoding of a node \( v_i \) where only the corresponding \( i^{th} \) index is 1, and 0 otherwise, and \( b_{i,v} = e_i - e_v \). For each edge \( e \in E \), we define \( b_e = b_{i,v} \) where \( u \) and \( v \) are the end points of the edge \( e \) respectively. This implies that \( b_e \) contains only two non-zero entries: 1 (on the \( i^{th} \) index) and \( -1 \) (on \( j^{th} \) index). Using the standard basis vector \( b_e \), it can be shown that \( L = \sum_{e \in E} b_e b_e^T \). Similarly, for \( G(e) \), we have \( L^+ = L - (1-\theta)b_e b_e^T \) [14].

#### 3.2. Network Kirchhoff Index

Extracting features from graphs is hard and involves many challenges such as the permutation invariance, scalability, and efficiency of the algorithm. To tackle these challenges, here we leverage a resistor network metaphor. The electrical flow on the resistor network can be described as is follow.

Let \( I(u) \) represent currents injected at the graph node \( u \), and \( I(e) \) indicate the currents induced in the edge \( e \). Let \( \phi(u) \) be the potential induced at the node \( u \). Then the Kirchhoff current law states that the sum of total current injected to the node must be equal to the sum of current flowing out of it:

\[
B^{T}I = I_{inj}
\]  

(3)

By Ohm's law, the flow of current in an edge follows

\[
I = WB\phi
\]  

(4)

Combining (3) and (4), we get

\[
I_{inj} = B^{T}(WB\phi) = L\phi
\]  

(5)

If the total amount of current passed to a node equals the total current flowing out of it, then we can write it as follows.

\[
\phi = L^{+}I_{inj}
\]  

(6)

Now using \( L^{+} \), the effective resistance and Kirchhoff index can be defined as below.

**Effective Resistance:** The effective resistance (\( R_{eff} \)) of an edge \( e \) can be defined as the potential difference between endpoints of the edge when a unit current is passed at the head of the edge and extracted at the tail. In terms of Laplacian matrix, it can be defined as

\[
R_{eff}(e) = b_e L^{+} b_e
\]  

(7)

Generally, the effective resistance between a pair of nodes can be calculated through well-known manipulations: series, parallel. Edges \( (e_1, \ldots, e_k) \) correspond to resistors with resistance \( (r_1, \ldots, r_k) \). Ohm in series can be replaced by a single edge with resistance \( \sum r_i \). In case of parallel connectivity, edges can be replaced by one edge with resistance \( (\sum r_i)^{-1} \). The definitions of these series and parallel manipulations on resistor networks follow that the effective resistance captures both the number of paths, quality of paths and their length between pair of nodes which intuitively quantifies edges based on their electrical importance.

**Rayleigh’s Monotonicity Law:** “If the resistance of a circuit is increased, the effective resistance \( R_{eff} \) between any two points can only increase. If they are decreased, it can only decrease”.

By law of Rayleigh’s Monotonicity, we define edges’ centrality scores as follow [14].

**Definition 3.1.** The centrality score \( C_{i}(e) \) of an edge \( e \in E \) using the Kirchhoff index is defined as the increase of the Kirchhoff index of the graph obtained from \( G \) by \( \theta \)-deleting \( e \).

**Definition 3.2.** Kirchhoff Index: The Kirchhoff index of a graph \( G \) can be defined as the sum of the effective resistance over all pairs of nodes.

\[
\text{Kirchhoff index}(G) = \sum_{v_i \in V} R_{eff}(v_i, v_j)
\]  

(8)

The Kirchhoff index of a given graph quantifies graph connectivity. The less connected a graph is, the larger the Kirchhoff index will be. Similarly, when the graph is densely connected, the corresponding Kirchhoff index is smaller in general. Based on the definition of Kirchhoff index and Rayleigh’s Monotonicity Law, we have [14]:

**Fact 1.1:** The Kirchhoff index of a graph does not decrease by \( \theta \)-deleting \( e \).

**Fact 1.2:** “Let \( \lambda_1, \lambda_2, \ldots, \lambda_{n-1} \) be the nonzero eigenvalues of \( L \). The Kirchhoff index of graph \( G \) satisfies Kirchhoff index(G) = \( n \sum_{i=1}^{n-1} \frac{1}{\lambda_i} - n\text{Tr}(L^{+}) \)”. Based on Fact 1.1, edges scores \( C_{i}(e) \) using Kirchhoff index can be computed via \( \theta \)-deletion as Kirchhoff index encapsulates information regarding the number of paths and quality of paths. By Definition 3.1, the difference of the Kirchhoff index between \( G \) and \( G(e) \) reflects \( C_{i}(e) \) and it can be computed for all the edges to encode the structure of the graph. In the forthcoming sections, we present a graph embedding algorithm using the computation of exact Kirchhoff index of the graph. And then we show linear time approximation of Kirchhoff index for encoding sufficiently large graphs.

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\(^1\) http://danspielman.github.io/Laplacians.jl/latest/usingSolvers/
4. Methodology

4.1. Graph embedding algorithm using exact Kirchhoff index

Here we present Kirchhoff index based graph representation algorithm which encodes graph structures using network Kirchhoff index. The algorithm describes the procedure for extracting graph representation by computing exact network Kirchhoff index.

**Algorithm 1:** Compute \( \mathcal{F} \) using exact Kirchhoff index

**Input:** Graph \( G = (V, E, w) \), Laplacian \( L \), bin width \( h \), number of bins \( b \)

**Output:** \( \mathcal{F} \)

1. Let \( L^+ \) indicate pseudo-inverse of the Laplacian matrix
2. \( K = nTr(L^+) \)
3. initialize sparse matrix \( \mathcal{F}_{v,i} \) with zeros
4. for \( (u, v) \in E \)
5. \( G'(e) \rightarrow \theta - deletion \)
6. \( L^\theta = \text{Laplacian of } G'(e) \)
7. \( K^\theta = nTr(L^\theta(e)) \)
8. \( C_v(e) = K^\theta - K \)
9. \( \mathcal{F}_{uv} = C_v(e) \)
10. end for
11. Set \( \text{Flt} \leftarrow \{ \mathcal{F}_{uv} \mid (u, v) \in |V| \} \)
12. \( \mathcal{F} \leftarrow \text{histogram(Flt, b, h)} \)
13. return \( \mathcal{F} \)

The core idea of Algorithm 1 is to perform \( \theta - deletion \) and compute the Kirchhoff index of the graph. Initially, the algorithm computes pseudo-inverse in step-1 and by using Fact 1.2, it computes Kirchhoff index \( K \) (step-2) of the \( G \). In step 5, it performs \( \theta - deletion \) and computes the Laplacian \( L^\theta \), pseudo-inverse \( L^\theta(e) \) and \( K^\theta \) (step-6-8) of the new graph \( G'(e) \). Using Definition 3.1, edge centrality score is computed in step 9. This procedure is repeated for all edges and the corresponding scores \( C_v(e) \) are stored in matrix \( \mathcal{F} \). Finally, we flatten \( \mathcal{F} \) and compute the histogram \( \mathcal{F} \) (step-12-13).

From step 4–11, the algorithm computes Laplacian pseudo-inverse \( L^\theta \) and \( L(e) \) for each edge \( mn \) times which has \( O(n^2) \) running time complexity each. Thus, the total cost of this algorithm is \( O(mn^2) \).

4.2. NetKI: Algorithm for graph representation using the approximate Kirchhoff index

Here, we show an approximation algorithm that extracts feature vector from graphs in nearly linear time. Our proposed method is based on the approximation procedure proposed in [14]. Li and Zhang [14] introduced two alternate \( \epsilon - approximation \) algorithms to estimate the relative importance of graph edges and an algorithm for computing nodes’ importance in a given graph. To compute edge relative importance, the authors used the notion of Kirchhoff index and approximate it in two different ways. We follow the EDCGECENTCOMP2(Theorem 1.2) approximation algorithm for estimating the edge scores which is derived through Sherman-Morrison formula [63]. For an edge \( e \in E \) and a scalar \( 0 < \theta < 1 \), we have

\[
(L^\theta)^+ = (L - (1-\theta)b_v b_v^T)^+ \\
\]

By Sherman-Morrison formula, we have

\[
= L^+ + (1-\theta) \frac{b_v b_v^T L^+}{1 - \theta b_v L^+ b_v} \\
\]

Thus by the definition of the edge score, we have

\[
\varphi_{ie} = n(1-\theta) \frac{Tr(L^+ b_v b_i L^+)}{1 - (1-\theta)b_v L^+ b_v} \\
\]

Where \( (L^\theta)^+ \) is the Laplacian of the graph \( G'(e) \) which is obtained by \( \theta - deletion \) in the original graph \( G \). \( \varphi_{ie} \) is the score we aim to compute as discussed in Algorithm 1. The numerator of Eq. (10) is the trace of the implicit matrix \( Tr(L^+ b_v b_i L^+) \) which is approximated through Hutchinson’s Monte-Carlo method [64,65]. Further to apply \( L \), the author used nearly-linear time Laplacian solver [57]. The denominator of the Eq. (10) is just \( 1 - (1-\theta)b_v L^+ b_v \) derived in Eq. (7), and the authors used the random projection in [54] to approximate the effective resistance.

EDGECENTCOMP2 algorithm is based on two main procedures: solving linear systems in graph Laplacian and approximating the effective resistance. Due to the various real-world applications of the graph Laplacian including the graph partitioning, resistor networks and the theory of random walks, there has been a surge in approaches to solve it efficiently. In particular, the breakthrough results by Spielman and Teng [55] showed that linear systems in graph Laplacian could be solved in \( O(m\log^2 n\log |\lambda|) \) time. Various follow up works have improved the bound in the last decade. One notable Laplacian solver was proposed by Kyng and Sachdeva [13] which is purely based on random sampling and runs in expected time \( O(m\log^2 n) \). The algorithm does not use any graph theoretic constructions and performs Gaussian elimination based on Cholesky factorization to solve \( x = L^b \) in graph Laplacian. This allows us to recursively solve a much broader family of linear systems. In EDCGECENTCOMP2 algorithm, the Laplacian solver proposed in [57] was used which runs in \( O(m\log^2 n\log^2 n\log |\lambda|) \) expected time.

To approximate the effective resistance, EDCGECENTCOMP2 uses the algorithm proposed in [54]. The authors in [54] used vector random projection in a subspace \( \{Q\mathcal{W}^{1/2}LX \} \) where \( Q \) is a random Bernoulli matrix and \( X_n \) indicates standard basis vector of node \( n \). Adapted from [54], we show the procedure for approximating the effective resistance in Algorithm 2.

**Algorithm 2:** Compute approximation matrix \( Z \)

**Input:** Graph \( G = (V, E) \) and \( \epsilon > 0 \)

**Output:** \( R_{\text{eff}}(u, v) \)

1. Let \( Q_{\theta, n} \) be a random Bernoulli matrix, where \( k = 24\log n/\epsilon^2 \)
2. Compute \( \hat{\mathcal{W}} = Q\mathcal{W}^{1/2}B \)
3. Let \( Z_{k, n} \) be an approximate resistance matrix
4. Let \( \gamma_i \) denote the rows of \( \hat{\mathcal{W}} \)
5. for \( i = 1 \) to \( k \)
6. \( U^* D^* \rightarrow \text{SPARSEOCHOLESKY}(L, \epsilon) \)
7. \( Z_i \leftarrow (U^* \gamma_i) \epsilon (D^*) (U^* \gamma_i) \)
8. end for
9. return \( Z \)

The Algorithm 2 uses Johnson-Lindenstrauss Lemma and Kyng and Sachdeva’s Laplacian solver to estimate the effective resistance. Using the Johnson-Lindenstrauss lemma, the algorithm randomly generates \( Q_{\theta, n} \) Bernoulli matrix and considers it as a \( k \)-dimensional projection in \( \mathbb{R}^d \) [66,13]. Then the algorithm computes \( \hat{\mathcal{W}} \) matrix which takes \( 2m \times 24\log n/\epsilon^2 + m = O(m/\epsilon^2) \) time, since \( W \) is diagonal and \( B \) has \( 2m \) entries. Finally, the algorithm solves the Laplacian for each row of \( \hat{\mathcal{W}} \) matrix to form the resistance.
matrix Z which has $O(m \log^2 n)$ time complexity. After computing the resistance matrix, the effective resistance of each edge $(u, v)$ can be computed as follows [54].

$$R_{eff}(u, v) \sim |Z(X_u - X_v)|^2$$

(11)

Where $X_u$ and $X_v$ represent the corresponding columns of nodes $u$ and $v$ in the resistance matrix $Z$.

**Algorithm 3: Compute $\mathcal{X}$ using NetKI**

**Input:** Graph $G = (V, E, w)$, Laplacian $L$, bin width $h$, number of bins $b, \theta$ and $\epsilon : (0, 1/2]$.

**Output:** $\mathcal{X}$

1: Let $z_1, \ldots, z_M$ be independent random $\pm 1$ vectors, where $M = \lfloor e^{-\epsilon^2 \log n} \rfloor$
2: Let $A_{M \times b}$ be a solver matrix
3: Let $F_{b \times n}$ be a score matrix
4: for $i = 1$ to $M$
5: $U^T D U \leftarrow$ SPARSECHOLESKY($L, \epsilon$)
6: $x_i = (U^T)^T (D^T)^T (U)^T z_i$
7: foreach $e \in E$ do
8: $A_e \leftarrow x_i^T b_a b_e^T x_i$
9: end for
10: end for
11: $Z \leftarrow \text{Compute approximation matrix}(G, \theta e/9)$
12: for each $e(u, v) \in E$ do
13: $F_{u} \leftarrow (1 - \theta) \sum_{i=1}^{M} A_{e}/(1 - (1 - \theta)Z|X_u - X_v|)^2$
14: end for
15: Set $\text{Flt} \leftarrow \{\mathcal{X} \forall (u, v) \in |V|\}$
16: compute $\mathcal{X} \leftarrow \text{histogram}($Flt, $b, h$)
17: return $\mathcal{X}$

Using EDGECENTCOMP2, we present the approximation algorithm for generating graph representations in Algorithm 3. The core idea of the algorithm is to approximate Kirchhoff index based on Eq. (10). To apply Laplacian pseudo-inverse, we use nearly-linear time solver [13] to generate sparse Cholesky factorization (step 5). In step 6, the algorithm computes the solution vector and calculates Kirchhoff index score for each edge using Eq. (10). This process is repeated $M$ times which compute $M$ scores for each edge. Further, the algorithm computes approximate matrix $Z$ using Algorithm 2 to estimate the effective resistance (step-11). Following Monte-Carlo method which approximates trace of a matrix $L$ by $\frac{1}{n} \sum_{i=1}^{n} A_{e} L z_i$, where $E[z_i^T L z_i] = \text{Tr}(L)$, the algorithm computes score matrix $F$. Note that $z_i$ represents random $\pm 1$ vectors consisting of independent Bernoulli entries. Finally, we flatten the score matrix and generate histogram $\mathcal{X}$ to perform graph classification task using a standard classification algorithm.

### 4.3. Modification to the EDGECENTCOMP2 algorithm

EDGECENTCOMP2 algorithm uses Cohen et al. [57] Laplacian solver, which is currently the best-known method for solving linear systems in graph Laplacian with running time $O(m \log^2 n \log \log n \log(1/\epsilon))$. However, it requires many graph-theoretic constructions such as the recursive preconditioning, leverage score, super-sparsest, and low-stretch spanning trees, and thus, is difficult to implement. Instead, we use Kng and Sachdeva (step 4 in Algorithm 2) - a popular nearly linear time Laplacian solver which runs in $O(m \log^2 n)$ in our experimental setup. This method is quite simple and based purely on random sampling. We refer the reader to [13] for more details. Further, we extend the algorithm to extract graph representation using the approximated Kirchhoff index based edge scores.

### 4.4. Analysis of the algorithms

The NetKI method is based on two main procedures: SPARSECHOLESKY and the “Compute approximation matrix”. The running time of SPARSECHOLESKY method is $O(m \log^2 n)$ . The “Compute approximation matrix” makes a total of $O(\log n)$ calls to SPARSECHOLESKY each of which takes $O(m \log^2 n)$ time. Therefore the total running time is $O(m \log^2 n)$.

The NetKI algorithm makes $O(\epsilon^{-2} \log n)$ calls to SPARSECHOLESKY each of which takes $O(m \log^2 n)$ time, makes a call to “Compute approximation matrix” which runs in $O(m \log^2 n)$ time, and computes the histogram which runs in $O(n + b)$, where $b$ is the number of bins. Therefore, the running time of NetKI algorithm is bounded by $O(\epsilon^{-2} \log^2 n) + O(m \log^2 n) + O(n + b) = O(\epsilon^{-2} \log^2 n)$.

### 5. Evaluation

We performed a different number of experiments to evaluate NetKI in terms of scalability, feature sparsity, running time, classification accuracy and features comparison. The forthcoming sections describe the details of all the experimentation we performed for the evaluation of NetKI.

#### 5.1. Scaling to large graphs

Scalability on large graphs is an essential requirement for classification of graph-structured data using machine learning. Although, recent approaches have shown promising results on various benchmark datasets, making them scalable on large graphs remains a challenge. To corroborate the scalability power of NetKI, here we present its performance analysis in terms of time on large random graphs with millions of nodes and edges.

We generate five large random graphs of size $10^5, 10^4, 10^3, 10^5, 10^6$ using Erdős-Rényi model with probabilities $0.01, 0.001, 0.0001, 0.00003$ respectively. The combination of these parameters generates large densely connected graphs up to 15 millions of edges. The time complexity comparison against other state-of-the-art methods including NetLSD [6] (which is currently the best known spectral representation method in terms of complexity) is shown in Fig. 2. We can see that NetKI took only 9 min on a graph with $10^5$ nodes and took 30 min on a graph with $10^6$ nodes. On the other hand, NetLSD ($h_x$) executed in 22 min on graph of size $10^5$ nodes and took 3 h on $10^6$ nodes. Although, FGSD performs well on small graphs, however, its performance worsens sharply as the size of the graphs increase. On the other hand, NetKI performs quite well on large graphs and outperforms all other approaches. These results illustrate that NetKI is a suitable graph embedding and classification method for large-scale graphs.

#### 5.2. NetKI sparse representations

Representation sparsity is desirable for the task of machine learning and can be defined as the number of zero entries present in the feature vector. Sparse representations have received considerable attention in the last few years, and have found applications in various fields such as signal processing, image processing, and deep learning [67]. The main aspect in which sparse representations are superior is the adaptation on varying levels of informa-
tion and its closeness to the linearity, as dense representations are difficult to learn. On the other hand, it provides an intermediate form between one-hot representation and pure distributed representation. Thus, the more sparse representation, the easier it is to generalize and learn [67,5].

We empirically show NetKI’s representation sparsity and also provide comparison with FGSD in Fig. 3 (left). The NetKI shown minimum 80% representation sparsity on D&D dataset and maximum 98% on PTC dataset. To better motivate the sparse representations for the classification task, we show the classification accuracy of NetKI and FGSD on sparse and dense Erdős–Rényi random graphs in Fig. 3 (right). The classification accuracy of both the algorithms is observed better on the representations of sparse graphs as compared to the dense representations.

To create random datasets for the experiment shown in Fig. 3 (right), we form binary classification problem and generate Erdős–Rényi random graphs with different densities for each class. We consider two cases: sparse and dense graphs with 10 datasets for each case and generate 100 graphs for each class where each graph consists of 100 nodes. For sparse graphs, in each iteration $i$, we set $p = 0.3 + (i \times c)$ for positive class and $p = 0.3 - (i \times c)$ for negative class. For dense datasets, we set $p = 0.7 + (i \times c)$ for one class and $p = 0.7 - (i \times c)$ for the other class. The value of $c$ is set to 0.002. We observe that with gradual increase in value of $i$, the difference between the graph densities of two classes widens as it makes one class denser and the other sparser and we see an improvement in the performance of the classifier.

5.3. Numerical results

We compare NetKI for classification accuracy and running time against the state-of-the-art methods, namely FGSD [5], NetLSD’s wave $w(g)$ and heat $h(g)$ kernels [6], NetSimile [32], Random Walk kernel (RW) [52], Shortest Path kernel [22], Edge Histogram kernel (EHK) [27], Neighborhood Hash kernel (NHK) [28] and Graphlet sampling kernel (GK) [29]. FGSD, NetLSD and NetSimile are recent statistical and spectral representation methods that show promising results while the rest are well-known graph kernel methods.

Datasets: We used five bioinformatics [68] and six social networks benchmark datasets to evaluate NetKI and other methods on. The bioinformatics datasets include MUTAG, PROTEINS, NCI1 and D&D. The social networks include COLLAB, IMDB-B, IMDB-M, REDDIT-B, REDDIT-5 K, and REDDIT-L. We report their characteristics in Table 2.

Experimental Setup: All experiments were performed on Intel Core i7-6800 K CPU@3.40 GHz machine with 128 GB RAM. We implemented NetKI in Python 3.7 with Networkx2 library to interact with graphs and Laplacians.jl3 library to solve linear systems in graph Laplacian. The NetKI implementation has been made publicly available 4. In NetKI setting, we set the cand $\beta$ to 0.5 throughout all the experiments. In the preprocessing step, we apply five-number summary and replace scores above the third quartile $(Q3) + 2.5 \times IQR$ with the mean value by considering the multisets of the whole dataset. In case of disconnected graph, we considered largely connected component of the graph. For computing the histogram, we set its range from $(0$–maxval) where maxval indicates the maximum score of the edge seen in the entire collection of graphs in the given dataset. After experimentation, the number of bins is chosen from the set $(20, 50, 200, 500, 1000, 2000)$ independently for different datasets. We attest this setup to be a good choice in terms of classification accuracy tradeoff. For experimenting with graph kernel methods, we used Grake1 library and used the default parameters setting. Moreover, we use the default parameters setting for NetSimile and NetLSD presented in their papers. NetLSD provides two functions: heat $h_s$ and wave $w_s$, therefore we report results on both the functions. In FGSD setting, we performed experiments using all the provided bin width: $(0.001, 0.0001, 0.00001)$ as reported in the actual paper and show the best results achieved among them.

For graph classification, we employed Random Forest (RF) Classifier with 500 estimators. To make a fair comparison among all the methods, the same classifier with the same parameter setting and 10-fold cross-validation is employed in the experimental setup. The results comparison on bioinformatics datasets with graph kernels, NetSimile, FGSD and NetLSD is reported in Table 3 while Table 4 reports results comparison on social network datasets.

On bioinformatics datasets (Table 3), we found that NetKI out-performs all other methods on PTC dataset while the classification accuracy on MUTAG, PROTEINS and D&D is within range 2.0 from the best accuracy. On NCI1 dataset, FGSD performed well and out-

![Fig. 2. Performance comparison in terms of time with state-of-the-art approaches on large random graphs. NetKI efficiently performs on large-scale graphs.](image-url)
performed all other methods. On social network datasets (Table 4), NetSimile performed quite well and outperformed all other methods on COLLAB and IMDB-B datasets. FGSD outperformed on IMDB-M and REDDIT-B while NetKI has shown highest accuracy on REDDIT-M-5 K and REDDIT-L social networks. The classification accuracy of NetKI on IMDB-M and REDDIT-B is within range 2.0 from the best result. We also report the running times of the algorithms (excluding any data loading or classification time for all algorithms) in brackets in Tables 3 and 4. We can see from the results that on small datasets such as MUTAG, and PTC, etc., the performance of FGSD is quite well as compared to the other algorithms, although it has quadratic time complexity. The main reason of this is that the FGSD implementation simply involves matrix multiplication which can be done quite fast when the size of the

### Table 2

Datasets characteristics.

<table>
<thead>
<tr>
<th>Dataset</th>
<th></th>
<th>Classes</th>
<th>avg</th>
<th></th>
<th></th>
<th></th>
<th>min</th>
<th>max</th>
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<tr>
<td>MUTAG</td>
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<td>2</td>
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<td>19.79</td>
<td>10</td>
<td>28</td>
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<tr>
<td>PROTEINS</td>
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<td>14.69</td>
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<tr>
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<td>32.3</td>
<td>3</td>
<td>111</td>
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<tr>
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<td>284.3</td>
<td>715.65</td>
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<tr>
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<tr>
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<td>97.53</td>
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<td>136</td>
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<td></td>
</tr>
<tr>
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<td>3</td>
<td>13.00</td>
<td>65.93</td>
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<tr>
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</tbody>
</table>

### Table 3

Classification accuracy comparison of NetKI against well-known graph kernels and recent methods is presented. Bold results indicate accuracy within range 2.0 from the best results of all the methods, and the blue color indicates best results. The expression ‘‘D’’ indicates computations exceed 24 h while std indicates the standard deviation of the performance of the algorithms. Running time in seconds of each algorithm is shown in brackets with the accuracy where the minimum running time is highlighted bold.

### Table 4

Classification accuracy and running times on social network datasets. ‘OMR’ indicates out of memory error.

![Fig. 3.](image-url) (left) NetKI representations sparsity comparison with FGSD on bioinformatics datasets. (right) NetKI and FGSD performance comparison in terms of classification accuracy on sparse and dense graph datasets generated using Erős-Rényi model with 30% and 70% densities. Note that ‘‘S’’ in legends indicates performance on sparse networks while ‘‘D’’ represents performance on dense network datasets.
matrices is small. However, as the size of the matrices increase such as in D&D and REDDIT datasets, the performance of FGSD worsens sharply. In contrast, NetKI performs quite well in terms of running time on large datasets as well as compared to other approaches. However, due to the approximation factor involved and the nature of the datasets, the performance of NetKI in terms of classification accuracy is also worse on NCI1 and COLLAB datasets. Recall that the main objective of NetKI was the scalability on sufficiently large datasets. We show in Tables 3 and 4 that the performance of NetKI is within range 2.0 from the best accuracy on most of the datasets. On the other hand, we can see from the tables and also from Fig. 2 that the running times of NetKI reduces sharply when the size of the datasets increases. To evaluate NetKI in terms of time and accuracy on very large real networks, we also reported classification accuracy and running time on REDDIT-L dataset, which is a very large social network dataset having 97491 average nodes and more than 1 million edges on average (see Table 2). We can see that NetKI outperformed NetLSD and shown quality results with 3.24% improvements in terms of accuracy and 127% improvements in terms of running time. Overall, we can see that NetKI has shown improved results in terms of running time and the classification accuracy is close to the state-of-the-art approaches that are, otherwise, impractical for large datasets. These results demonstrate that NetKI is a suitable graph embedding and classification method for large networks. NetSimile and FGSD could not able to scale on such large graphs therefore, we do not report on them.

5.4. Features comparison to NetLSD

Apart from the methodology perspective, NetKI and NetLSD are directly linked with the graph spectrum and can be observed closely related to each other. Thus, to provide more crisper analysis among them, here we present feature comparison to analyze the information extracted by both the algorithms from the same dataset. We hypothesized that high overlap in clustering on the embeddings generated through both the algorithms will reflect similar feature extraction. To test the hypothesis, we generate feature vectors from all small non-isomorphic graphs up to size 8. The total of them are 12111 number of graphs. For NetKI, we set the length of feature vector to 50 and use the same parameter setup shown above for other experiments. For NetLSD, we use default parameter setting and generate feature vectors from the same dataset. We perform k-means clustering with k = 4 on the resultant feature vectors of both the algorithms and find overlap between the clusters. For finding the overlap, we compute the intersection between NetKI each cluster with every cluster of NetLSD. The highest overlap was found 56% between NetKI cluster 1 and NetLSD cluster 0. The overlap between all other cluster was found < 50%. We have also shown tSNE visualization of the resultant clustering of both the algorithm in Fig. 4. We can see that the distributions of both the clustering are different, while the ratio of the overlap is also small. Thus, we conclude that NetKI and NetLSD capture different information from the graph.

5.5. Discussion

Currently, large graphs with millions of nodes and edges have become very common. Examples include online social networks, collaboration networks, and biomedical networks. Scalability on such large networks especially for the encoding methods is one of the key requirements to apply state-of-the-art machine learning methods [1]. There has been tremendous growth in graph representation methods, however, very few efforts have been devoted to enhance the performance of these methods and make them scalable on sufficiently large networks. The extensive experiments on well-known state-of-the-art methods on several networks shown in Section 5.3 reveal that the existing methods are still incapable of dealing with sufficiently large networks. And thus, there is a necessity of the encoding methods that are scalable on large networks.

On large real-world datasets such as REDDIT, D&D, and other large random networks, we show that NetKI runs very fast as compared to the existing approaches while attaining quality results in terms of classification accuracy. This demonstrates the effectiveness of NetKI on different kinds of networks, in particular, when the networks are sufficiently large and well-connected. In addition to the application of such efficient approaches in various domains, recently, the graph encoding methods have also been applied with GCNs collectively to improve the performance of the algorithms [69]. Such approaches help GCNs to generalize on multiple domains data. Due to the efficiency and quality results of NetKI, we believe that it is a suitable method that can be combined with end-to-end deep learning models to design efficient and universal graph embedding models.

6. Conclusion

We introduced NetKI, a novel method for graphs classification. NetKI is purely based on graph statistical representation that can be computed efficiently in nearly linear time. We proposed using the network Kirchhoff index as a function of graph representation f(gi) and show its approximation in nearly-linear time. NetKI does not require any graph summary statistics or node attributes and
relies only on graph structure. We performed extensive experiments on various real-world benchmark datasets and synthetic networks with millions of nodes and edges, and show that NetKI processes huge graphs and produce sparse representations which lead to attain good quality results.

**CRediT authorship contribution statement**

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**Declaration of Competing Interest**

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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