Limitations of low dimensional graph embeddings

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Abstract

The learning of graph representations, also called graph embeddings, is a fundamental technique in machine learning (ML). The aim is to represent the vertices of a graph by low-dimensional real-valued vectors, which can be used for a variety of downstream ML tasks. The popular technique of Graph Neural Networks (GNNs) can be thought of a type of graph representation learning method. This article surveys some recent work on the limitations of graph embedding methods. These results provide mathematical theorems proving that low-dimensional embeddings cannot recreate “community-like” structure, through commonly used kernels. These theorems are supported by empirical results, showing that many classic graph embedding methods actually perform poorly on important machine learning tasks. Low-dimensional representations often lose much of the fine-grained community structure of real-world data. The results surveyed in this article provide an interesting counterpoint to the popularity of graph embeddings and GNNs for machine learning.

1 Introduction

Capturing the rich structure of graph is central for many machine learning tasks. The heterogeneous, non-local, and massive structure of modern graphs are a problem for many basic machine learning tasks, such as ranking, link prediction, and classification [14]. Graph representation learning provides a convenient solution to this problem. Each vertex of a graph is represented as a vector in low-dimensional space. These vectors form the (low-dimensional) embedding of the graph. The vectors can be used for a plethora of downstream machine learning tasks. The aim is to have these vectors combine the graph structure with vertex features into a compact geometric representation.

The study of low-dimensional graph embeddings is an incredibly popular research area, and has generated many exciting results over the past few years (see surveys [19, 8] and a Chapter 23 in [27]). The most successful methods for graph embeddings often use Deep Learning techniques, together with classic approaches like matrix factorization [29, 16, 28, 30, 31, 24]. Graph Neural Networks (GNNs) can be thought of as specific class of graph embeddings algorithms [18, 37, 40]. Despite the large variety of algorithms used for graph representation learning (and specifically GNNs), their output has a simple, consistent structure. Given a graph $G$ on $n$ vertices, these methods map each vertex to a vector in $\mathbb{R}^d$, where $d \ll n$. In a typical applications, $n$ is the order of millions or more, while $d$ is in the hundreds.

Much of the advances in this field are primarily empirical; there are numerous papers in major machine learning conferences on more complex GNN architectures or more involved graph embedding methods. Nonetheless, there is limited principled understanding of the power of low-dimensional graph embeddings. Small changes in training or input data can lead to major differences in the output [17]. Much of the research on graph embeddings and GNNs, for example, report significant success in prediction tasks on graphs [16, 18, 31]. On the other hand, some papers suggest that low-dimensional graph embeddings can be beaten by simpler hand-tuned methods [17, 22]. It is useful to have a rigorous mathematical framework to understand graph embeddings. Specially, we wish to address the following questions.

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• To what extent can relevant graph structure be captured by low-dimensional graph embeddings?
• How does low-dimensional geometry relate to downstream ML tasks on graphs?
• How does the choice of kernel functions (for prediction tasks) affect the task?

Contrary to most research in this area, our approach to investigating these questions is through limitations or lower bounds. Our aim is develop a theoretical framework that is agnostic to the specific algorithm performing the embedding. We first describe some basic terminology and central concepts.

The importance of matrix factorizations: A low-dimensional matrix factorization approximates gives an algorithm-independent view of how low-dimensional embeddings represent graphs. We think of $M$ as an adjacency matrix, or any matrix representation of our graph $G$. In many Deep Learning methods, $M$ is constructed by long walks with appropriate vertex-centric aggregation functions [29, 16, 30]. The matrix $V$ is the collection of embeddings, where each column vector corresponds to a vertex. Recent work has shown the many different graph embedding methods can be cast as matrix factorizations, with different choices of matrices $M$, and different notions of approximations [31]. Indeed, the simplest low-dimensional embedding is obtained by a low-rank SVD of the adjacency matrix, arguably the most basic of matrix factorizations.

The importance of dot products: Regardless of how the graph embedding vectors are obtained, the general strategy is to use “nearness” in geometric space as a proxy for the similarity of the corresponding vectors. Thus, the downstream ML tasks treat vertices $i$ and $j$ as similar if their corresponding vectors $\vec{v}_i$ and $\vec{v}_j$ are similar. Arguably, the most important measure of similarity is the dot product $\vec{v}_i \cdot \vec{v}_j$ or cosine similarity (which is a normalized dot product). This has special significance for matrix factorizations, since $V^TV$ is precisely the matrix of all pairs of dot products.

Even when the embedding vectors are not obtained by matrix factorizations, the Gram matrix $V^TV$ is commonly used for downstream prediction tasks. A natural strategy for any prediction task is to use nearest neighbor (k-NN) search according to the dot product/cosine similarity. There are many open course packages for k-NN, making it the default choice in industrial applications of graph embeddings [2].

We now can formalize our initial questions in matrix language. Observe how there is no reference to any embedding method or GNN; we directly analyze the behavior of the output representation.

Can a Gram matrix $V^TV$ for $V \in \mathbb{R}^{d \times n}$ ($d \ll n$) capture the structure of either real-world networks or the properties of prediction matrices arising from downstream ML tasks (on graphs)?

This framework captures the vast majority of graph embedding constructions and applications.

1.1 Limitations of graph embeddings

This article presents two results on the limitations of low-dimensional graph embeddings for prediction and machine learning on real-world graphs [34, 36]. The underlying theoretical results are algorithm agnostic, and the limitations hold for any graph embedding algorithm. These results come with empirical backing, performed on classic and important graph embedding methods.

Inability to recreate triangle structure [34]: This result focuses on the fundamental premise of low-dimensional embeddings, rather than a specific ML task. Algorithms to construct (or predict from) low-dimensional embeddings implicitly pose a low-dimensional graph/matrix model. This model represents a distribution of graphs that are created from a set of real vectors (where each vector represents a vertex). The embedding is constructed by fitting this model to an input graph, typically using dot product as a proximity measure.
This result mathematically proves that no low-dimensional model (using a dot product kernel) can simultaneously recreate two central properties of real-world graphs: sparsity and triangle density. It is well-known from the early days of network science that real-world graphs are sparse but have high clustering coefficients [39, 32, 33, 13]. Hence, typical models used to construct low-dimensional embeddings cannot generate realistic graphs. They miss the (crucial) triangle structure of real data.

The theory is supported with empirical results. These empirical results are demonstrated on other kernels beyond the dot product, thus suggesting that the limitations are fundamental. We discuss these results in §2.

There are notable counterpoints to these results. Firstly, Chanpuriya et al show that asymmetric embeddings avoid the rank lower bounds discussed above [9]. Another result of Chanpuriya et al shows that graphs can sometimes be reconstructed from their embeddings. In some cases, the community structure of the reconstruction is actually enhanced [10]. We give more detail in §2.

Weak performance on community labeling tasks [36]: This result takes a complementary angle, and focuses on a downstream ML task. Community labeling is a binary prediction problem, where we wish to predict if two vertices belong to a community. This work sets up a community labeling problem for both real and synthetic data sets. First, the authors create a simple benchmark algorithm using logistic regression on classic graph features (like Personalized PageRank and short path counts). This benchmark has fairly good performance, as measured by distributions of local precision. On the other hand, many well-established graph embedding methods have surprisingly poor performance on the same task.

These observations are backed up with theoretical proofs. The prediction matrices based on low-dimensional embeddings provably do not have the typical community structure of real data. This work also investigates alternate kernels, like the normalized softmax (used in results like DeepWalk and node2vec [29, 16]). These kernels have other limitations in that slight noise can destroy community structure in the corresponding prediction matrices.

These results are discussed in §3.

1.2 Broader context

A reader may ask: how are the limitations stated in this article consistent with large body of work on the effectiveness of graph embeddings and GNNs? Our answer is two-fold, backed up by [17, 22]. First, most research on graph embeddings compare various representation learning methods with each other, and do not consider alternative baselines. Secondly, we believe that many graph embeddings methods do not have good predictions with respect to other metrics. For example, almost all these result use the AUC metric to measure link prediction performance. On the other hand, AUC is a bad measure for sparse ground truth [20, 25]. Indeed, in §3, we show poor prediction performance (for graph embedding methods) on a local precision metric.

There has been compelling empirical work showing that GNNs and embeddings can be outperformed by simpler methods. We mention two results in detail because they highlight empirical weaknesses in graph embeddings, and reinforce the previous points.

Gurukar et al, the lack of good experimental design [17]: Gurukar et al do a detailed comparison of twelve different graph representation learning methods on a variety of ML tasks. They focus on two of the most important tasks of link prediction and node classification. Despite there being many newer methods, they observe that the M-NMF algorithm is best for link prediction [38] and NetMF algorithm [31] is best for node classification. No graph representation method outperforms on both metrics. This paper does an exceptional job of clearly specifying the experimental design and thoroughly investigating previous work.

Along the lines of the current article, simple task specific baselines are competitive with graph embedding methods. These baselines are formed by a simple model that uses basic graph features (like Common Neighbors, Adamic-Adar index [4], Jaccard similarity, etc.). These results are analogous to what we observe in [36].
Gurukar et al point out a major problem with evaluations in graph embedding papers. Quoting from there: “. . . a new method almost always compares its performance against a subset of other methods [and] datasets previously evaluated. While great care is taken to tune the new method, the same care is often not taken when evaluating baselines.”

Huang et al, easier methods beat GNNs [22]: Huang et al devise an alternate graph learning algorithm called Correct and Smooth (C&S). Consider the problem of node classification. This method starts with a “base predictor” that ignores the graph structure. Using only node features, one can train a basic model for classification. Then, the graph is introduced a post-processing step to “correct and smooth” the errors. The idea is that the errors should be correlated along edges. So this method tries to find a smooth error estimation on the graph, which is computed using standard label propagation methods. This smoothed error estimate is used to correct the base predictor.

The C&S method is shown to outperform GNNs on node classification tasks on the OGB leaderboard [1]. In some cases, a state-of-the-art GNN has slightly better performance, but the GNNs always have many orders of magnitude more parameters. Hence, it is much more expensive to train them than C&S.

We also mention another line of weaknesses, specific to message passing GNN architectures. These results show that certain graph-theoretic properties cannot be learned by common GNN architectures [40, 26, 15]. There are formal limits on what GNN-based representations can learn. Some of these results show that message passing GNNs are no more powerful than the classic Weisfeiler-Lehman (WL) isomorphism test [40].

The work presented in the current article is different from these results because of the (algorithm agnostic) focus on the geometry of graph representations.

2 Impossibility of capturing sparse, triangle-rich structure

The first result we present argues that low-dimensional embeddings with dot product geometries are not good representation of real-world graphs. Seshadhri, Sharma, Stolman, and Goel demonstrate mathematically and empirically that they lose local cluster structure, a central aspect of graphs that arise from real data [34].

Graph embeddings are often generated by assuming that the embeddings vectors lie in a hidden, latent space. We assume a model that generates graphs from these vectors, which can be thought of as the model parameters. The embeddings are constructed by optimizing for these parameters, given the input graph.

Consider the graph embedding vectors \( \vec{v}_1, \vec{v}_2, \ldots, \vec{v}_n \in \mathbb{R}^d \) (denoted by the \( d \times n \) matrix \( V \)). Let \( G_V \) denote the following distribution of graphs over the vertex set \( [n] \). For each index pair \( i, j \), independently insert (undirected) edge \( (i, j) \) with probability \( \max(0, \min(\vec{v}_i \cdot \vec{v}_j, 1)) \). (If \( \vec{v}_i \cdot \vec{v}_j \) is negative, \( (i, j) \) is never inserted. If \( \vec{v}_i \cdot \vec{v}_j \geq 1 \), \( (i, j) \) is always inserted.) This model subsumes the classic Stochastic Block Model [21] and Random Dot Product Model [42, 6]. Many graph embedding methods, including GNNs, effectively optimize over such a model to generate the embedding vectors.

Two hallmarks of real-world graphs are: (i) Sparsity: average degree is constant with respect to \( n \), and (ii) Triangle density: there are many triangles incident to low degree vertices [39, 32, 33, 13]. The large number of triangles is an important aspect of community structure.

**Definition 2.1:** For parameters \( c > 1 \) and \( \Delta > 0 \), a graph \( G \) with \( n \) vertices has a \((c, \Delta)\)-triangle foundation if there are at least \( \Delta n \) triangles contained among vertices of degree at most \( c \).

Typically, we think of both \( c \) and \( \Delta \) as constants. We emphasize that \( n \) is the total number of vertices in \( G \), not the number of vertices in \( S \). In Figure 1, we plot the value of \( c \) vs \( \Delta \) as the thick blue line. (Specifically, the \( y \) axis is the number of triangles divided by \( n \).) Observe that for all graphs, for \( c \in [10, 50] \), we get a value of \( \Delta > 1 \) (in many cases \( \Delta > 10 \)). As mentioned earlier, there is much work in network science showing that there are often a linear number of triangles among low degree vertices [33, 13].
Figure 1: Plots of degree $c$ vs $\Delta$: We plot results for a variety of real-world graphs: (i) **ca-HepPh**, a High Energy Physics coauthorship network (ii) **Facebook**, a small snapshot of a Facebook social network, (iii) **String_hs**, a Protein-protein interaction network. We plot $c$ versus the total number of triangles only involving vertices of degree at most $c$. We divide the latter by the total number of vertices $n$, so it corresponds to $\Delta$, as in Definition 2.1. We plot these both for the original graph (in thick blue), and for a variety of embeddings and kernel functions. For each embedding, we plot the maximum $\Delta$ in a set of 100 samples from a 100-dimensional embedding. The embedding analyzed by Theorem 2.1 (TDP) is given in thick red. Observe how the embeddings generate graphs with very few triangles among low degree vertices. The gap in $\Delta$ for low degree is 2-3 orders of magnitude. The other lines correspond to alternate embeddings, using the NODE2VEC vectors and/or different functions of the dot product.

Our main result is that any embedding of graphs that generates graphs with $(c, \Delta)$-triangle foundations, with constant $c$, $\Delta$, must have near linear rank. This contradicts the belief that low-dimensional embeddings capture the structure of real-world complex networks.

**Theorem 2.1**: Fix constant $c > 4$, $\Delta > 0$. Suppose the expected number of triangles in $G \sim \mathcal{G}_V$ that only involve vertices of expected degree $c$ is at least $\Delta n$. Then, the rank of $V$ is at least $\Omega(\sqrt{n \log^2 n})$.

Equivalently, graphs generated from low-dimensional embeddings cannot contain many triangles only on low-degree vertices. In all applications, $d$ is thought of as a constant, or at least much smaller than $n$. On the contrary, Theorem 2.1 implies that $d$ must be $\Omega(n/\log^2 n)$ to accurately model the low-degree triangle behavior. This lower bound holds regardless of how the vectors are constructed.

### 2.1 Empirical validation

We empirically validate the theory on a collection of complex networks. For each real-world graph, we compute a 100-dimensional embedding through SVD and an important Deep Learning method, node2vec [16]. We generate 100 samples of graphs from these embeddings, and compute their $c$ vs $\Delta$ plot. This is plotted with the true $c$ vs $\Delta$ plot. (To account for statistical variation, we plot the maximum value of $\Delta$ observed in the samples, over all graphs. The variation observed was negligible.) Fig. 1 shows such a plot for three different real-world networks.

In all cases, this plot is significantly off the mark at low degrees for the embedding. Around the lowest degree, the value of $\Delta$ (for the graphs generated by the embedding) is 2-3 orders of magnitude smaller than the original value. The local triangle structure is destroyed around low degree vertices. The total number of triangles is preserved well, as shown towards the right side of each plot. A nuanced view of the triangle distribution, as given in Definition 2.1, is required to see the shortcomings of low dimensional embeddings.

We note that several other functions of dot product have been proposed in the literature, such as the softmax function [29, 16] and linear models of the dot product [18]. Theorem 2.1 does not have direct implications for such models, but our empirical validation holds for them as well. The embedding in Theorem 2.1 uses the truncated dot product (TDP) function $\max(0, \min(\vec{v}_i \cdot \vec{v}_j, 1))$ to model edge probabilities. We construct other embeddings that compute edge probabilities using machine learning models with the dot product and Hadamard
product as features. This subsumes linear models as given in [18]. We also consider (scaled) softmax functions, as in [29], and standard machine learning models (LRDP, LRHP).

For each of these models, we perform the same experiment described above. Fig. 1 also shows the plots for these other models. Observe that none of them capture the low-degree triangle structure, and their $\Delta$ values are all 2-3 orders of magnitude lower than the original. Chanpuriya et al also validate these results on various other embedding methods [9].

### 2.2 Counterpoints

We discuss two important results of Chanpuriya, Musco, Sotiropoulos, and Tsourakakis [9, 10]. The first result show that the rank lower bound of Theorem 2.1 can be circumvented with an asymmetric embedding. They prove that one can construct two matrices $U, V \in \mathbb{R}^{d \times n}$ such that the graph distribution from $UV^T$ can generate realistic triangle structure. They show that any bounded degree graph can be embedded in this method with at most max-degree dimensions. This introduces a new technique for graph embeddings. We note, however, that such asymmetric embeddings lose the geometric structure of the standard embeddings. One wants geometric proximity of vectors to represent structural closeness. For vertices $i$ and $j$, an asymmetric embedding approximates the edge probability by $\vec{u}_i \cdot \vec{v}_j$, but the similarity of $i$ and $j$ would be measured by looking at (say) $\vec{u}_i$ and $\vec{u}_j$. It would be interesting to incorporate similarity in asymmetric embeddings.

Another result shows that, in some cases, community structure can be reconstructed from DeepWalk embeddings [10]. This shows that some structure is being retained by the embeddings. We note that these results mostly focus on SBMs with a constant (at most 5) blocks, or only the largest few communities in real data. Theorem 2.1 and the other results in this article focus on cases where the number of blocks/communities is large. We believe that low-dimensional embeddings can recover the top few communities, but fail to capture the rich structure of many small communities. In the next section, we discuss this point further.

### 3 Challenges for community labeling using graph embeddings

One central promise of unsupervised graph embedding methods is to preserve network structure in the geometry. To what extent do embedding methods capture graph structure relevant to downstream ML tasks?

This section is based on the paper of Stolman, Levy, Seshadhri, and Sharma [36]. We begin this discussion with the empirical results, because they highlight the core observation. After that, we will go into the mathematical explanations that relate to low-dimensional embeddings and factorization.

Consider the following well-defined pairwise community labeling problem. Given two vertices $i$ and $j$, the binary classification task is to determine whether they belong to the same community. We note that this community labeling problem is an instance of a broad range of community detection problems that have a long history of study in the graph mining literature [23].

#### 3.1 Empirical setup

We use a set of real-world datasets with ground truth community labels, an Amazon co-purchase graph of products and a DBLP citation network [41]. We also create a synthetic Stochastic Block model, with 100K vertices, and small blocks of size 20 each. There is a dense graph within each block, and a random sparse graph connecting all the blocks.

As explained earlier, the prediction task is to determine if an input pair $i, j$ of vertices belong to a community. (They may belong to multiple communities; to make the problem simpler, we do not require any community labels to be determined.)
Figure 2: Each point, \((x, y)\), on the curve represents the approximate fraction of vertices, \(y\), for which the given method produces a precision@10 score of at least \(x\). LR-Structural is plotted against the two best performing embedding methods. 1000 vertices are sampled and for each vertex \(v\) sampled, the vertices of the graph \(u_1, \ldots, u_n\), are ordered by decreasing score assigned by the given classifier. The precision@10 is the fraction of \(u_1, \ldots, u_{10}\) which share a community with \(v\). Across all instances, the simple baseline LR-Structural handily outperforms more complex graph embedding methods.

The setup for graph embeddings: We experiment with a set of important graph embedding methods based on factorizations and Deep Learning (GraRep, DeepWalk, node2vec, NetMF [29, 7, 28, 16, 31]). We note that the NetMF method was reported to be one of the best embedding methods for node classification [17]. Given the embedding for each vertex, we need to construct pairwise features for pairs in \((i, j) \in V \times V\), for the community prediction task. The standard approach is to either take the dot product \(\vec{v}_i \cdot \vec{v}_j\) or the Hadamard product \(\vec{v}_i \circ \vec{v}_j\). (Recall that the Hadamard product is a \(d\)-dimensional vector whose \(r\)th coordinate is the product of the \(r\)th coordinates of \(\vec{v}_i\) and \(\vec{v}_j\).) We finally train a logistic regression model on these features for the prediction problem. This is the standard pipeline used in prior work [11, 16, 8].

A simple baseline: We compute four basic structural graph features for pairs of vertices \((i, j)\). We look at the cosine similarity and cut size between neighborhoods, and the Personalized PageRank values. These are well-known classic features used in the literature [35, 5]. We train a simple logistic regression model on these four features; the model is denoted LR-Structural.

Performance metric: For both real and simulated data, the ground truth is sparse, i.e. the vast majority of node pairs do not belong to the same community. We do not use AUC because of its problems in measuring sparse data [20, 25]. Instead, it is appropriate to measure the prediction performance using precision-recall curves for this highly imbalanced label distribution [12].

The methods are evaluated by comparing the “precision@10” distributions. We sample 1000 random vertices. For each of 1000 vertices sampled, \(v\), we order the other vertices of the graph, \(u_1, \ldots, u_n\), in decreasing order of their prediction score. (The model predictors based on logistic regression on the embedding vectors or structural features assign a score in \([0, 1]\).) We compute the precision, per vertex, of the classifier among the top 10 scores, with respect to the ground truth. When the predictor is based on dot product, this is simply the top 10 neighbors in geometric space. In other words, we sample a vertex at random and report the fraction of its ten nearest neighbors with which it shares a community.

We represent the distribution of values of precision@10 scores as a reliability curve. This is the curve \((x, y)\) such that at least a \(y\) fraction of vertices sampled had a precision@10 score score of at least \(x\). Higher \(y\) values for a given \(x\) indicate better performance. Fig. 2 contains the curves for the best methods against the baseline (we leave out methods with poorer performance).
Main observation: Across all instances, the baseline LR-Structural method heavily outperforms the more complex graph embedding methods. The performance gap for the simple Stochastic Block Model instance is striking. The graph is practically learnable (just a collection of dense blocks with sparse connections), but the embedding methods fail to accurately predict pairs within blocks. For the LR-Structural method, in all instances, at least 90% of the vertices have a precision@10 of at least 0.5. This means, for at least 90% of the vertices, at least five of the top 10 scores are in the same community. By contrast, for the embedding methods, this fraction is less than 75%. When looking for a precision of more than 0.8, the embeddings methods have less than 20% of vertices achieving high scores. Overall, we observe that the embeddings methods do not perform the community labeling task well, despite the simple LR-Structural baseline having good precision values.

3.2 Theoretical explanation of limitations

As explained earlier, a large variety of graph embedding methods (including those using Deep Learning) implicitly factorize a matrix $M$ as $V^T V$. Here, $M$ denotes some matrix representing the input graph data or the final prediction, and $V \in \mathbb{R}^{d \times n}$ is the matrix of graph embeddings. Broadly speaking, we can classify these methods into two categories:

- **Direct factorizations**: Here, we set $V$ as \[ \arg\min_V \| V^T V - M \|_2, \] where $M$ is typically (some power of) the graph adjacency matrix. Methods such as Graph Factorization, GraRep [7], and HOPE [28] would fall under this category.

- **Softmax factorizations**: These methods factorize a stochastic matrix, such as (powers of) the random walk matrix. (A stochastic matrix has row sums equal to one.) Since $V^T V$ is not necessarily stochastic, these methods apply the softmax to generate a stochastic matrix. Notable examples are such methods are DeepWalk [29] and Node2vec [16]. Formally, consider the normalized softmax matrix $\text{nsm}(V)$ given by
\[
\text{nsm}(V)_{ij} = \frac{\exp(\vec{v}_i \cdot \vec{v}_j)}{\sum_k \exp(\vec{v}_i \cdot \vec{v}_k)}
\] (28)

Note that $\text{nsm}(V)$ is stochastic by construction.

The NetMF [31] method interpolates between these categories and shows that a number of existing methods can be expressed as factorization methods, especially of the above forms.

The notion of community pairs: We start with an abstraction of community structure from a matrix standpoint: many dense blocks in an overall sparse matrix. We quantify “how much” community structure can be present in a matrix $V^T V$ or $\text{nsm}(V)$, for any matrix $V \in \mathbb{R}^{d \times n}$ (for $d \ll n$). This formulation captures the fundamental notion of a low-dimensional embedding, without referring to any specific method to compute it.

Let us start with an $n \times n$ matrix $M$ that represents the “similarity” or likelihood of connection between vertices. This is the final prediction matrix for community labeling. For convenience, let us normalize so that the $\forall i \in [n], \sum_{j \leq n} M_{ij} \leq 1$. (So the sum of similarities of a vertex is at most 1.) A communities is essentially a dense block of entries, which motivates the following definition. We use $\varepsilon$ to denote a parameter for the threshold of community strength. One should think of $\varepsilon$ as a small constant, or something slowly decreasing in $n$ (like $1/\text{poly}(\log n)$).

**Definition 3.1:** A pair of vertices $(i, j)$ is a potential community pair if both $M_{ij}$ and $M_{ji}$ are at least $\varepsilon$.

Note that we do not expect all such pairs $(i, j)$ to truly be together in a community. Hence, we only consider such a pair a potential candidate. We expect community relationships to be mutual, even if the matrix $M$ is not. A community can be thought of as a submatrix where at least a constant fraction of pairs are potential community pairs. It is natural to expect that $\Theta(n)$ pairs are community pairs; indeed, most vertices should participate in communities, and will have at least a constant number of community neighbors. Our mathematical analyses shows that direct and softmax factorizations cannot produce these many potential community pairs.
**Lower bound for direct factorizations:** We prove that the number of potential community pairs in $V^T V$ is linear in the rank, and thus, a low-dimensional factorization cannot capture community structure. The proof uses the rotational invariance of Frobenius norms.

**Theorem 3.1:** Consider any matrix $V \in \mathbb{R}^{d \times n}$ such that row sums in $V^T V$ have absolute value at most 1. Then $V$ has at most $d/2\varepsilon^2$ potential community pairs.

**Proof:** Since $V^T V$ has row sums of absolute value at most 1, the largest absolute value of eigenvalue is also at most 1 (a consequence of the Gershgorin circle theorem [3]). The rank of $V^T V$ is at most $d$, so $V^T V$ has at most $d$ non-zero eigenvalues. We can express the Frobenius norm squared, $\|V^T V\|_F^2$, by the sums of squares of eigenvalues. By the arguments above, $\|V^T V\|_F^2 \leq d$.

But the Frobenius norm squared $\|V^T V\|_F^2$ is also the sums of squares of entries. Each potential community pair contributes at least $2\varepsilon^2$ to this sum. Hence, there can be at most $d/2\varepsilon^2$ potential community pairs.

**The instability of softmax factorizations:** The properties of softmax factorizations are more nuanced. Firstly, we can prove that softmax factorizations can represent community structure quite effectively.

**Theorem 3.2:** For $d = O(\log n)$, there exists $V \in \mathbb{R}^{d \times n}$ such that $\text{nsm}(V)$ exhibits community structure. Specifically, for any natural number $b \leq n$, there exists $V \in \mathbb{R}^{d \times n}$ such that $\text{nsm}(V)$ has $n/b$ blocks of size $b$, such that all entries within blocks are at least $1/2b$.

Indeed, this covers the various SBM settings we study, and demonstrates the superiority of softmax factorizations for modeling community structure. We note that a similar theorem, for asymmetric factorizations, was proved in [9].

On the other hand, these factorizations are highly unstable to small perturbations. Indeed, with a tiny amount of noise, any community pair can be destroyed with high probability. The noise model scales each vector with small $(1 \pm \delta)$ Gaussian noise to get the matrix $\text{nsm}(V^{(\delta)})$. (The formal definition is given in [36].)

**Theorem 3.3:** Let $c$ denote some absolute positive constant. Consider any $V \in \mathbb{R}^{d \times n}$. For any $\delta > c \ln(1/\varepsilon)/\ln n$, the following holds in $\text{nsm}(V^{(\delta)})$ (this is the matrix formed by $\text{nsm}(V)$ with $\delta$ Gaussian noise). For at least $0.98n$ vertices $i$, for any pair $(i, j)$, the pair is not a potential community pair with probability at least 0.99.

Thus, with overwhelming probability, any community structure in $\text{nsm}(V)$ is destroyed by adding $o(1)$ (asymptotic) noise. This is strong evidence that either noise in the input or numerical precision in the final optimization lead to destruction of community structure. These theorems give an explanation of the poor performance of the embeddings.

### 4 Conclusion

Instead of interpreting these limitations pessimistically, we reiterate the need for rigorous, foundational work in graph embeddings and GNNs. The work in this article merely scratches the surface. The limitations given in [34, 36] might not hold for all low-dimensional embedding methods, but they cover a large class of them. The limitations certainly hold for the most popular methods used, and is reinforced by the empirical results. The counterpoints of [9, 10] lead to a more nuanced picture for specialized embedding methods. We need a deeper understanding of how limitations can be avoided and how they relate to the downstream ML tasks.

The limitations question a purely empirical approach of designing better and better embedding methods and GNNs. As [17] correctly point out, each method comes with many hyperparameters, so it might be possible to tune one method to beat another and vice versa. Small improvements on some test datasets might not reveal the
complete picture. The theoretical and mathematical framework discussed in this article provide a more rigorous basis for research. If there are fundamental limitations from low-dimensional geometry for certain methods, we should not try to “tune” the problems away by experimenting with hyperparameters.

Overall, we believe that the work surveyed in this article provide an exciting new research perspective for graph embeddings and GNNs.

References


