EEGNN: Edge Enhanced Graph Neural Networks

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Abstract

Training deep graph neural networks (GNNs) poses a challenging task, as the performance of GNNs may suffer from the number of hidden message-passing layers. The literature has focused on the proposals of over-smoothing and under-reaching to explain the performance deterioration of deep GNNs. In this paper, we propose a new explanation for such deteriorated performance phenomenon, mis-simplification, that is, mistakenly simplifying graphs by preventing self-loops and forcing edges to be unweighted. We show that such simplifying can reduce the potential of message-passing layers to capture the structural information of graphs. In view of this, we propose a new framework, edge enhanced graph neural network (EEGNN). EEGNN uses the structural information extracted from the proposed Dirichlet mixture Poisson graph model, a Bayesian nonparametric model for graphs, to improve the performance of various deep message-passing GNNs. Experiments over different datasets show that our method achieves considerable performance increase compared to baselines.

1 Introduction

Graph neural networks (GNN) (Zhou et al., 2020; Wu et al., 2020) is an important tool for analyzing graph data, such as social network (You et al., 2020), transportation network (Chen et al., 2021a), molecular graph (Huang et al., 2020), biological network (Zhang et al., 2021), financial transaction network (Wang et al., 2021), academic citation graph (Xu et al., 2021), and knowledge graph (Ji et al., 2021). GNNs have become popular with the state-of-the-art performance by applying deep learning methodologies to graphs. Among them, message passing neural networks (MPNN) (Gilmer et al., 2017) use message-passing layers to compute node embeddings. Examples of MPNNs include graph convolutional neural networks (GCN) (Kipf and Welling, 2017), GraphSAGE (Hamilton et al., 2017), graph attention networks (GAT) (Veličković et al., 2018), and gated graph neural networks (GGNN) (Li et al., 2016). Similar to standard multi-layer perceptron (MLP) in deep learning, the message passing layer in a GNN framework aggregates information from the local neighbors of each node, and then transforms the information via an activation function into the embedding
A node embedding can aggregate information over $N$ hop neighbors, in the form of $N$ hidden message-passing layers, thus incorporating further reaches of the graph.

Although deeper layers in a non-graph neural network often achieve better performance (Krizhevsky et al., 2012; He et al., 2016), GNNs typically perform best with only 2 to 4 hop neighbors, that is, 2 to 4 hidden layers. In contrast, using a larger number of layers, termed as deep stacking, may lead to a substantial drop in the performance for GNNs (Klicpera et al., 2019; Rong et al., 2019; Li et al., 2020; Chen et al., 2020b). One explanation for this phenomenon is the over-smoothing. By applying graph convolution repeatedly over many hidden layers, the representation of the nodes will be indistinguishable. As a result, the over-smoothing can jeopardize the performance of deep GNNs. Another explanation is the under-reaching. When GNNs aggregate messages over long paths, the information propagation across distant nodes in the graph becomes difficult because it is susceptible to bottlenecks (Alon and Yahav, 2020). This causes GNNs to perform poorly in predicting tasks that require the remote interaction (Singh et al., 2021; Hwang et al., 2021).

Many efforts have been devoted to addressing these limitations. To handle the over-smoothing, DropEdge (Rong et al., 2019) and DropNode (Huang et al., 2021) were proposed to randomly remove a certain number of edges or nodes from the input graph at each training epoch. These methods are likened to Dropout (Srivastava et al., 2014), which randomly drops hidden neurons in neural networks to prevent overfitting. On the contrary, to address the under-reaching, virtual edges (Gilmer et al., 2017), super nodes (Scarselli et al., 2009; Hwang et al., 2021), or short-cut edges (Allamanis et al., 2018) can be added to the original graph. However, none of aforementioned methods consider adding or removing based on the structural information of the graph. Instead, the pattern of deciding which nodes or edges to be added or removed comes from an arbitrarily random selection.

Different from the over-smoothing and under-reaching, we propose a new explanation for performance deterioration of deeper GNNs from the perspective of misusing edge structural information, mis-simplification, explained as follows. Most observed graphs are recorded as simple graphs, where self-loops are not allowed, and all edges are unweighted and undirected (Shafie, 2015). In a natural way, GNNs are designed for learning such simple graphs that can be constructed by collapsing multiple edges into a single edge as well as removing self-loops. This approach, however, discards the information inherent in the original network. Take one example, for a source node connected to many neighboring target nodes (see node 1 in Figure 1a), its self-loop has an equal weight to neighboring non-loop edge, which may under-weigh the importance of this node. Take another example, no matter how similar the two nodes are (see nodes 1 and 2 in Figure 1a), only one edge is allowed to connect the pair of nodes, and as a result, the information passing between both nodes is restricted. Furthermore, edge (1, 2) should play a more important role in message passing than edge (1, 3), because node 2 is a key node. However, typical GNNs treat both edges indifferently as they are equally weighted in the simple graph. Therefore, such mis-simplification can reduce the potential of message-passing layers to capture structural information in GNNs.

To solve this issue, we propose an edge-enhanced graph neural network (EEGNN), which incorporates edge structural information in the message-passing layer. First, we assume that there is an underlying virtual multigraph, allowing for self-loops and for multiple edges between pairs of nodes, and the observed graph model can be viewed as a transformation of the virtual multigraph. As illustrated in Figure 1, the left is the
original observed simple graph, while the right is the corresponding virtual graph. Second, to build the virtual multigraph that can capture the edge structural information, we propose the Dirichlet mixture Poisson graph model, a Bayesian nonparametric model. Following Caron and Fox (2017), the interactions between nodes are modelled by assigning a sociability parameter to each node. Then, the counts of edges are generated from a Poisson distribution, where the Poisson rate is the product of sociability parameters of the nodes in two ends. Finally, in the framework of EEGNN, we can then replace the observed graph in a GNN with the virtual multigraph. In this architecture, message-passing layers can then assign weights proportionally to the importance of the edges, thus passing the information from nodes to nodes in a more reasonable manner.

The main contribution of our paper is fourfold.

• We outline a new explanation for the poor performance of deep GNNs;
• We propose a new way to enhance existing GNN methods by utilizing the structural information of graphs;
• We propose a Bayesian nonparametric graph model and its Monte Carlo Markov chain (MCMC) inference procedure;
• We demonstrate the superior sample performance of our proposal over existing methods through the experiments on six real datasets.

2 Preliminaries

2.1 GNN and Message Passing Layer

We begin by introducing some notation. Let $G = (V, E)$ be a graph with node set $V = \{v_1, \ldots, v_{|V|}\}$ and edge set $E = \{e_1, \ldots, e_{|E|}\}$, where $|V|$ and $|E|$ denote the number of nodes and edges in $G$, respectively. The adjacency matrix $A \in \mathbb{R}^{|V| \times |V|}$ is defined as $A_{ij} = 1$ if $(v_i, v_j) \in E$ and 0 otherwise. The corresponding degree matrix $D = \text{diag}(D_1, \ldots, D_{|V|})$, where $D_i = \sum_{j=1}^{|V|} A_{ij}$. We denote the data matrix by $X \in \mathbb{R}^{m \times |V|}$, whose $j$th column corresponds to the node $j$ with $m$ features.
GNN is a neural network model to process graphs for node classification, edge prediction and graph classification (Gori et al., 2005; Zhou et al., 2020; Wang et al., 2021). Within various GNNs, information is exchanged between nodes and is updated by neural networks via message passing layers (Gilmer et al., 2017). Specifically, the initial representation, \( h_i^0 \) for node \( i \), is generated by a function of this node’s features. Then, the message passing layers update the representation based on this node’s neighbors. The message passing contains two steps: the aggregation step and the update step. Denote the representation for node \( i \) in layer \( l \) by \( h_i^l \). A message passing layer in GNN can be expressed as

\[
h_i^{l+1} = \text{UPDATE}(h_i^l, \text{AGGREGATE}(h_j^l \mid j \in \mathcal{N}_i)),
\]

where AGGREGATE(\( \cdot \)) denotes a permutation-invariant function, such as the sum, mean, and maximum, to send information from one node to another through edges, and UPDATE(\( \cdot \)) denotes linear or nonlinear differentiable functions such as MLP. For example, the vanilla GCN uses

\[
h_i^{l+1} = \sigma(\sum_{j=1}^{\vert V \vert} \tilde{P}_{ij} h_j^l W^l \mid j \in \mathcal{N}_i \cup \{i\}),
\]

or in matrix form, \( H^{l+1} = \sigma(\tilde{P} H^l W^l) \) (Kipf and Welling, 2017), APPNP uses \( H^{l+1} = (1-\alpha)\tilde{P} H^l + \alpha H^0 \) (Klicpera et al., 2019), and GCNII uses \( H^{l+1} = \sigma((1-\alpha)\tilde{P} H^l + \alpha H^0)((1-\beta)I_n + \beta W^l) \) (Chen et al., 2020b), where \( \tilde{P} = (D+I)^{-1/2}(A+I)(D+I)^{-1/2} \), \( H^l \) is the representation for all nodes in layer \( l \), \( \sigma \) and \( W^l \) are respectively the activation function and the corresponding weight in a neural network layer, and \( \alpha \) and \( \beta \) are hyper-parameters. The formulas above show that the GNN treats each edge with equal weight and hence leads to mis-simplification. In order to solve this issue, we adopt a Bayesian nonparametric sparse graph model to generate the virtual edges and virtual multigraph.

### 2.2 Bayesian Nonparametric Sparse Graph Model

In contrast with other graph models that are based on node feature embeddings, Caron and Fox (2017) represent the observed graph \( G \) as a point process on \( \mathbb{R}^2 \), \( A = \sum_{i,j} A_{ij} \delta(\theta_i, \theta_j) \), where \( \theta_i \) is a proxy for node \( i \) on the real axis, as illustrate in Figure 2a. Noting that the same definition for node \( i \) is also applied to node \( j \), but we omit explanation for node \( j \) to avoid redundancy. This representation specifies the source and target nodes for each edge. To model the possibility for two nodes constructing an edge, a sociability parameter \( w_i > 0 \) is assigned to node \( i \) for each \( i = 1, \ldots, \vert V \vert \). Following Aldous (1997), the graph model can be factorized as

\[
p(A_{ij} = 1) = 1 - \exp(-2w_i w_j) \quad \text{for } i \neq j \\
p(A_{ii} = 1) = 1 - \exp(-w_i^2)
\]

This is equivalent to modelling an unobserved integer-valued multigraph as \( z_{ij} \sim \text{Poisson}(w_i w_j) \) and setting \( A_{ij} = 1_{z_{ij} + z_{ji} > 0} \). To model the sparsity property in real graphs, that is, \( \vert E \vert = O(\vert V \vert^2) \), the sociability is generated from a completely random measure. See also Appendix A for a short review. This model allows for self-loop and multi-edges, and thus can be used to build a virtual multigraph. However, as the Poisson intensity is factorized as the outer product of a vector with itself, only one feature for each node is considered, which restricts the capability of this model. To address this issue, we propose a novel model in Section 3.1 below.
3 Methodology

3.1 Dirichlet Mixture Poisson Graph Model

To adopt the latent community information among nodes in the graph, mixed-membership stochastic block model (Airoldi et al., 2008) associates each node with latent cluster distributions. In an analogy, we add cluster-membership features to each pair of edges instead of nodes. Specifically, we extend the graph model in Caron and Fox (2017) by proposing the following Dirichlet mixture Poisson graph model (DMPGM),

\[
\pi = (\pi_1, \pi_2, \ldots) \sim \text{GEM}(\alpha),
\]

\[
W_0 = \sum_{i=1}^{\infty} w_{0,i} \delta_{\theta_i} \sim \text{CRM}(\nu, \lambda), \quad W_k = \sum_{i=1}^{\infty} w_{k,i} \delta_{\theta_i} \sim \text{GP}(W_0),
\]

\[
z_{ij} \sim \text{PP}(\sum_{k=1}^{\infty} \pi_k w_{k,i} \times w_{k,j}), \quad A_{ij} = \min(z_{ij} + z_{ji}, 1)\mathbb{1}_{i \neq j},
\]

for \(i, j, k \in \mathbb{N}^+\), where GEM(\(\alpha\)) is the distribution for atom sizes of a Dirichlet process DP(\(\alpha\)) and each atom corresponds to a distinct node. Moreover, CRM(\(\rho, \lambda\)) denotes a completely random measure with \(v^C(dw, d\theta) = v(dw)\lambda(d\theta)\) as its Levy measure, and GP(\(H\)) denotes gamma process with base measure \(H\). We summarize the probabilistic generative steps as follows. First, the cluster distribution \(\pi\) is assigned with a prior GEM(\(\alpha\)), which allows for infinitely many clusters. Second, we use a hierarchical structure to generate values for the node sociability parameter in each cluster, \(w_{k,n}\). Then \(W_0\), sampled from a completely random measure, is used as the base measure in GP(\(W_0\)) for \(W_k\) such that \(w_{k,n}\) belongs to gamma distribution \(\Gamma(w_{0,i})\). This hierarchical setting is designed to ensure the components in \(W_k\) share atom locations (Teh et al., 2006; Liu et al., 2022). Finally, following Caron and Fox (2017), an undirected multigraph is generated from a Poisson process, and a simple graph is transformed from the multigraph by aggregating multiple edges to a single edge for each pair of nodes and removing self-loops. The corresponding adjacent matrix \(A\) to the observable simple graph can then be generated.
DMPGM can be equivalently expressed from a mixture model perspective. Specifically, a set of edges in each cluster is sampled from Poisson($\pi_k \tilde{w}_k^2$), where $\tilde{w}_k = \sum_{i=1}^{\infty} w_{k,i}$. As a consequence, this is equivalent to sampling the total number of edge from Poisson($\sum_{k=1}^{\infty} \tilde{w}_k$), and then assigning each edge a cluster membership from Categorical($\pi_1 \tilde{w}_1^2, \pi_2 \tilde{w}_2^2, \ldots$). Following the same methodology, for each edge, a pair of nodes is then sampled from Categorical($w_{k,1}, w_{k,2}, \ldots$) in the cluster $k$. Hence, a relationship between edges and nodes is constructed. In Appendix C, we show that DMPGM enjoys similar properties as the model in Caron and Fox (2017) in the following theorem.

**Theorem 1** The graph constructed by DMPGM is sparse if CRM in (2) has infinite activity.

It is worth noting that DMPGM extends the model in Caron and Fox (2017) by assuming that edges can belong to different clusters. As a result, DMPGM is more flexible and applicable in modellling real data. We also note that DMPGM is distinct from the overlapping communities graph model (Todeschini et al., 2020) and graph Poisson factorization (Zhou, 2015), because we assign a Dirichlet prior for the clustering step 1 nonparametric manner in the following steps.

In Appendix C, we show that DMPGM enjoys similar properties as the model in Caron and Fox (2017) by assuming that edges are restricted to the weights $w_{k,i}$ because the locations $\{\theta_i\}$ of both observed and unobserved nodes are not likelihood identifiable, thus being ignored. Moreover, given the observed nodes set $V$, the weights for each $W_k$ are truncated to a $(|V| + 1)$-dimensional vector, $w_k = (w_{k,0}, w_{k,1}, \ldots, w_{k,|V|})^t$, where $w_{k,i}$ corresponds to the weight on an observed node $i$ for $1 \leq i \leq |V|$, and $w_{k,0}$ is the sum of weights for all unobserved nodes. Similarly, the posterior distribution for $\pi$ is truncated to a $(K + 1)$-dimensional vector, $\pi = (\pi_0, \pi_1, \ldots, \pi_K)^t$, where $K$ is the truncated number of clusters and is inferred adaptively in Step 4 below, and $\pi_0$ corresponds to the cluster without any observation. Consequently, given the truncation levels $|V|$ and $K$ for $W_0$ and $\pi$, respectively, DMPGM contains the following parameters to infer: $\pi$, $\{w_k\}_{k \geq 0}$, $\mathbf{c} = \{c_{ij}\}_{A_{ij} = 1, 1 \leq i \leq z_{ij}}$, and $z = \{z_{ij}\}_{A_{ij} = 1}$.

We next propose a MCMC inference framework that can infer the number of edge clusters in a Bayesian nonparametric manner in the following steps.

**Step 1** Update $w_{0,1}, \ldots, w_{0,|V|} | \bar{w}_0, z, \mathbf{c}$ using Hamilton Monte Carlo (Kroese et al., 2011), where $\bar{w}_0 = \sum_{i=0}^{|V|} w_{0,i}$ with the log-posterior and its gradient provided in Appendix B.1.

**Step 2** Update $w_{k,1}, \ldots, w_{k,|V|} | \bar{w}_k, w_0, z, \mathbf{c}$ for $k = 1, \ldots, K$ given the conjugacy, where $\bar{w}_k = \sum_{i=0}^{|V|} w_{k,i}$.

We sample $\bar{w}_{k,i}$ ~ Dirichlet($\nu_0, \nu_1, \ldots, \nu_{|V|}$), where $\nu_i = w_{0,i} + \sum_{j=1}^{|V|} n_{k,i}$, $n_{k,i} = \sum_{j=1}^{|V|} 1_{c_{ij}=k}$, and then compute $w_{k,i} = \bar{w}_k \bar{w}_{k,i}$.
**Step 3** Update $\pi_0, \pi_1, \ldots, \pi_K | \mathbf{z}, \mathbf{c}$ using the conjugacy. Analogous to Step 2, we sample $\pi_k \sim \text{Dirichlet}(n_0, n_1, \ldots, n_k)$, where $n_k = \sum_{i=1}^{||V||} \sum_{j=1}^{||V||} c_{ij} - 1$ for $k > 0$ and $n_0 = \alpha$.

**Step 4** Update the latent edge cluster membership $c_{ij} | \{w_k\}_{k \geq 0}, \pi$ for each pair $(i, j)$ such that $A_{ij} = 1$. For each edge we sample from the multinomial distribution $p(c_{ij} = k) \propto \pi_k w_{k,i} w_{k,j}$ for $k = 0, 1, \ldots, K$. In this step, if $k = 0$ is sampled, we add a new cluster (Teh et al., 2006; Liu et al., 2022), and increase the truncated number of clusters from $K$ to $K + 1$.

**Step 5** Update the unobserved $z_{ij} | \pi, w_k \sim \text{Truncated–Poisson}(\sum_{k=0}^{K} \pi_k w_{k,i} w_{k,j})$ for each pair $(i, j)$ such that $A_{ij} = 1$, where truncated Poisson is a conditional probability distribution of a Poisson-distributed random variable with strictly positive counts (Cohen, 1960).

**Step 6** Update the $\bar{w}_k$ and $\bar{w}_0$ using Metropolis–Hastings (Kroese et al., 2011) algorithm based on the log-posterior provided in Appendix B.2.

We iterate over **Steps 1-6** until convergence. For the MCMC algorithm, the global variables are updated in linear time, and the Monte Carlo step iteratively samples from $K$ clusters. Therefore, the computational complexity is dominated by $O(K \max(||V||, ||E||))$.

### 3.3 Edge Enhanced Message Passing

In conventional message passing layers built from a simple graph, information for node $i$ is obtained from edges linked to its neighboring nodes $\mathcal{N}_i$ and from its self-loop. In these layers, each edge $(i, j)$ for $j \in \mathcal{N}_i \cup \{i\}$ has equal weight, resulting in *mis-simplification* of the more complex structural information for the GNN, as described in Section 1. To overcome this *mis-simplification*, we sample artificial edges given the estimated DMPGM, from which we construct a virtual multigraph

$$G^* = (V, E, r), \quad r((i, j)) = z_{ij},$$

(3)

where the multiplicity-map $r : E \rightarrow \mathbb{N}^+$ assigns to each edge an integer to represent its multiplicity, and $z_{ij}$ in DMPGM is defined in (2) and is inferred from Step 5 in Section 3.2. In this way, we can extract the edge structural information, via the inferred multiplicity for each edge, using the DMPGM model to build the virtual multigraph. For example, as illustrated in Figure 1, two artificial self-loops are added to nodes 1, one artificial self-loop is added to nodes 2, and the edge $(1, 2)$ is assigned with multiplicity 2, where the multiplicity is decided by $z_{11} = 2$, $z_{22} = 1$ and $z_{12} + z_{21} = 2$, respectively. We then replace the original simple graph in the message passing layers by the generated virtual multigraph, that is,

$$h_{i}^{l+1} = \text{UPDATE}^l(h_{i}^l, r((i, i), \text{AGGREGATE}^l(h_{j}^l, r((i, j) \mid j \in \mathcal{N}_i))).$$

(4)

For example, for GCN, APPNP and GCNII, we replace $\hat{P}$ by $\hat{P}$, where $\hat{P}$ is defined as

$$\hat{P} = \hat{D}^{-\frac{1}{2}} \hat{A} \hat{D}^{-\frac{1}{2}}, \quad \hat{A} = (A_{ij} = z_{ij}), \quad \hat{D} = \text{diag}(\hat{D}_i = \sum_j z_{ij}).$$

(5)

In addition, as the virtual multigraph already contains self-loops, there is no need to add the self-loops again to the message passing layers. This is different from conventional GNNs, where the self-loops are often added
and forced to be a single edge. Though GIN (Xu et al., 2019) and JKNet (Xu et al., 2018) also assign different weight for self-loops empirically, we are the first to propose a method to systematically decide the relative weights for self-loops and other edges.

Finally, conditional on the updated parameters of DMPGM in each iteration, we sample the multiplicity of each edge. Then a set of multiedges and self-loops are generated from DMPGM, which can be used to build virtual graph and update GNN trainable parameters. We summarize the algorithm in Algorithm 1.

**Algorithm 1: EEGNN Algorithm**

Iterate Step 1 to Step 6 in Section 3.2 till the MCMC chains converge.

Set up initialization of trainable parameters in EEGNN.

repeat
1. Build the virtual graph and sample $\hat{P}$ according to (5),
2. Use $\hat{P}$ to replace $\tilde{P}$,
3. Update GNN parameters using the gradient descent,
4. Obtain a new sampling for the parameters in DMPGM by implementing Step 1 to Step 6,
until the convergence of the loss function of EEGNN

4 Experiments

4.1 Datasets and Bayesian Estimation

In this section, we demonstrate through real data examples that EEGNN can effectively use the edge structural information to improve the performance for various GNNs. We conduct empirical experiments to compare EEGNN with representative baselines across six well-established network datasets. First, *Cora*, *Citeseer*, and *PubMed* are standard benchmark datasets for citation networks (Yang et al., 2016). In these networks, nodes represent papers, and edges indicate cross citations between papers. Node features are the bag-of-words embedding of the contents, and node labels are academic subjects. Second, *Texas*, *Cornell*, and *Wisconsin* are webpage cross-link networks (Pei et al., 2020). Their nodes represent web pages of universities, and edges represent hyperlinks between them. Node features are bag-of-words embedding of the websites. Node labels contain five categories for the webs including students, projects, courses, staff, and faculty. Statistics for these datasets are summarized in Appendix D.

Our experiments are implemented by using a gamma process as the completely random measure in (2). Following Section 3.2, we infer the parameters of DMPGM using MCMC in the following way. We use population based training (Jaderberg et al., 2017) to tune the hyperparameters in DMPGM. For each dataset, we grow the MCMC chain up to 50,000 epochs. Figures 3a and 3b display the log-likelihood and number of clusters with respect to training epochs for the *Texas* dataset. (The training results for the other datasets are shown in Appendix E.) Figure 3a shows that the log-likelihood of *Texas* for the DMPGM converges after 10,000 epochs. Moreover, benefiting from the Bayesian nonparametric model, we can infer the number of edge clusters in a data adaptive manner (Liu et al., 2022). Figure 3b shows that the inferred number of edges
per node (termed as *multiplicity* of virtual edges per node) rises from an initial value of 10 to 50 at the start of training, then converges to around 35. The inferred edge multiplicity is displayed in the histograms in Figure 3c. These histograms show that a large proportion of the edges in the observed graph have underlying multi-edges, suggesting the *mis-simplification* in the original observed graph.

### 4.2 Comparison with Baselines

With the inference results of DMPGM, following Section 3.3, we implement the experiments to compare baseline GNNs and their edge enhanced versions. For the baseline GNNs, we chose SGC (Wu et al., 2019) and its variant, including APPNP (Klicpera et al., 2019) and GCNII (Chen et al., 2020a), and hence name their edge enhanced versions as EE-SGC, EE-APPNP and EE-GCNII, respectively. To make a fair comparison, we follow the settings of ‘sweet point’ GNN hyperparameter configuration in Chen et al. (2021b) for all datasets. The details of these hyperparameter settings are collected in Appendix F. For all experiments, the GNNS are trained with a maximum of 1000 epochs and an early stopping patience of 100 epochs. To analyze the effect of EEGNN with different number of layers, we run the experiments for 2, 16, 32 and 64 layers. We randomly split each dataset into training and test sets, train the baseline GNNs and the edge enhanced versions using the same training set, and then compute the node clustering accuracy on the test set. We repeat this procedure 50 times for each model and dataset. The mean predictive accuracy and the corresponding standard deviation are reported in Table 1.

We observe a few apparent patterns. First, EEGNN can improve the performance of the baseline models in most of the cases. For example, SGC, the backbone GNN for various models, performs poorly with 32 layers (see Table 1c). However, with the aid of our EEGNN framework, the accuracy of the SGC model is increased by more than 6% for Cora, and by approximately 2% across other candidate datasets. Moreover, SGC performs even worse with 64 layers for Cora and Pubmed (see Table 1d). EEGNN largely improve the prediction accuracy by 10% and 15%, respectively, in both cases. It is worth noting that the improvements are attained without changing any other settings. As using virtual multigraph or observed simple graph brings in the only difference, this provides a strong evidence to reveal that EEGNN can be used as a tool to enhance baseline GNNs by alleviating the *mis-simplification* problem.
Second, for APPNP and GCNII, EEGNNs achieve similar accuracies on the Cora, Citeseer and PubMed datasets, but significantly improves the performance on Texas, Wisconsin and Cornell. Especially, with 64 layers, EE-GCNII for Texas leads to more than 6% improvement, and EE-APPNP for Citeseer results in more than 10% increase in the predictive accuracy. On the other hand, as APPNP and GCNII have already reached relatively high accuracy (approximately 70% – 80%) on the Cora, Citeseer and PubMed, further enhancement to higher accuracy tends to be difficult.

Finally, we observe that EEGNN has larger impact on the performance of deeper SGC on the Cora, Citeseer and PubMed. With only 2 layers, edge enhanced versions behave slightly worse than baseline models. However, with 32 or 64 layers, EEGNNs achieve considerable improvements. This is because the mis-simplification applies to all layers. Therefore, the distortion of edge structural information is accumulated from the first to the last layer, resulting in severe performance deterioration.

5 Conclusion

This paper presents a novel explanation for the performance deterioration of deeper GNNs: mis-simplification. We propose DMPGM, a Bayesian nonparametric graph model and its MCMC inference framework. Using the information obtained from DMPGM, we replace the original simple graph by the virtual graph, and use the virtual graph to aggregate the information in the graph. The experiments over various real datasets demonstrate that EEGNN can improve the performance of baseline GNN methods. Our paper paves a new way to use information extracted by statistical graph modelling to improve the performance of GNNs. One limitation of our proposal is that EEGNN only adds the virtual edges to the observed graph, but does not remove edges according to the structural information. It is left for future work to develop a framework that allows to simultaneously add and remove edges with structural information.

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A Completely Random Measure

Completely random measures (Ghosal and Van der Vaart, 2017), including gamma process, inverse Gaussian process and stable process, are commonly used as priors for infinite-dimensional latent variables in Bayesian nonparametric models, because their realizations are atomic measures with countable-dimensional supports. Suppose that $(\Omega, \mathcal{F})$ is a Polish sample space, $\Theta$ is the set of all bounded measures on $(\Omega, \mathcal{F})$ and $\mathcal{M}$ is a $\sigma$-algebra on $\Theta$. A complete random measure from $(\Theta, \mathcal{M})$ into $(\Omega, \mathcal{F})$ can be characterized by its Laplace transform (Kingman, 1993),

$$E[e^{-tP(A)}] = \exp \left\{ - \int_A \int_{(0, \infty]} (1 - e^{-t\pi}) \nu(c, dx, ds) \right\},$$

where $A$ is any measurable subset of $\Omega$ and $\nu(c, dx, ds)$ is called the intensity measure. If $\nu(c, dx, ds) = \kappa(dx)\nu(ds)$, where $\kappa(\cdot)$ and $\nu(\cdot)$ are measures on $\Omega$ and $(0, \infty]$, respectively, the completely random measure is homogeneous and $\nu(\cdot)$ is called the Lévy measure. We can view this completely random measure as a Poisson process on the product space $\Omega \times (0, \infty]$ using the intensity measure and denote this completely random measure as $\text{CRM}(\kappa, \nu)$. For example, the gamma process $\Gamma(\kappa)$ has Lévy measure $\nu(ds) = s^{-1} e^{-s} ds$ such that $Q(A) \sim \Gamma(\kappa(A), 1)$ if $Q \sim \Gamma(\kappa)$, where $\Gamma(\alpha, \beta)$ is a gamma distribution with density $\frac{\beta^\alpha}{\Gamma(\alpha)} e^{-\beta x}$. Therefore, its normalization, Dirichlet process $P \sim DP(\kappa)$ (Ferguson, 1973) satisfies

$$(P(A_1), \ldots, P(A_n)) \sim \text{Dirichlet}(\kappa(A_1), \ldots, \kappa(A_n))$$

for any partition $\Omega = (A_1, \ldots, A_n)$, where $\bigcup_{i=1}^n A_i = \Omega$ and $A_i \cap A_j = \emptyset$ for any $i$ and $j$. Griffiths–Engen–McCloskey (GEM) distribution, which is the distribution of the weights in a Dirichlet process. For $(\pi_1, \pi_2, \cdots) \sim \text{GEM}(\alpha)$, it can be sampled by $\pi_i = g_i \prod_{i=1}^{i-1} g_i$, where $g_i \sim \text{Beta}(1, \alpha)$ independently (Ghosal and Van der Vaart, 2017).

B MCMC technical details and derivations

B.1 Derivations for Step 1

With the setup of DMPGM in (2) and the formula of moments for Dirichlet-multinomial distribution, we obtain that

$$p(w_0,1, \ldots, w_0,|V| \mid \bar{w}_0, z, c) \propto \prod_{i=1}^K \frac{\Gamma(\bar{w}_0)}{\Gamma(\bar{w}_0 + n_k)} \prod_{i=0}^N \frac{\Gamma(w_0,i + n_{k,i})}{\Gamma(w_0,i)} \prod_{i=1}^N v(w_0,i) \cdot u(\bar{w}_0 - \sum_{i=1}^{|V|} w_0,i),$$

where $n_{k,i} = \sum_{j=1}^{|V|} \sum_{j=1}^{|V|} 1_{c_{ji} = k} + 1_{c_{ji} = k}$, $v(\cdot)$ is the weight intensity measure for the complete random measure of $W_0$, and $u(\cdot)$ is the density function for $W_0(\Omega)$ that can be derived using its Laplace transform. To infer the posterior distributions for these parameters, we present the gradient of the log-posterior with respect to $w_0$, which will be used in Hamiltonian Monte Carlo,

$$\nabla_{w_0,i} \log p(w_0,1, \ldots, w_0,|V| \mid \bar{w}_0, z, c) = \sum_{i=1}^{|V|} \sum_{k=1}^K \left\{ \Phi(n_{k,i} + w_0,i) - \Phi(w_0,i) \right\}$$

$$+ \sum_{i=1}^{|V|} \nabla_{w_0,i} \log v(w_0,i) + \nabla_{w_0,i} \log u(\bar{w}_0 - \sum_{i=1}^{|V|} w_0,i),$$

15
where $\Phi$ is the digamma function.

### B.2 Derivations for Step 6

By the formulas of the densities for Poisson distribution and gamma distribution, we have that

$$
p(w_k | \bar{w}_0, \pi, c, z) \propto \left(\frac{\pi_k \bar{w}_k^2}{n_k!} e^{-\pi_k \bar{w}_k} \right) \cdot \frac{1}{\Gamma(\bar{w}_0)} \bar{w}_k^{\bar{w}_0-1} e^{-\bar{w}_k}.
$$

Therefore, the log-posterior with respect to $\bar{w}_k$ is

$$
\log p(w_k | \bar{w}_0, \pi, c, z) = (2n_k + \bar{w}_0 - 1) \log \bar{w}_k - \bar{w}_k - \bar{w}_k^2 \pi_k + \text{constant}.
$$

Similarly, following Caron and Fox (2017) and Liu et al. (2022), we obtain that

$$
p(w_0 | \bar{w}_k, \pi, c, z) \propto \prod_{k=1}^{K} \frac{1}{\Gamma(w_0)} \bar{w}_k^{w_0-1} e^{-\bar{w}_k} \cdot u(\bar{w}_0).
$$

and hence the corresponding log-posterior is

$$
\log p(w_0 | \bar{w}_k, \pi, c, z) = \log u(\bar{w}_0) + w_0 \sum_{k=1}^{K} \log(\bar{w}_k) - K \log \Gamma(\bar{w}_0) + \text{constant}.
$$

### C Technical Proofs and Derivations

#### C.1 Proof of Theorem 1

The proof for Theorems 3 and 4 in Caron and Fox (2017) can be directly adapted to DMPGM, Therefore, we only provide a sketch of the proof. First, we show that Theorem 3 in Caron and Fox (2017) also holds for DMPGM. We use

$$
\hat{D}_{ij} | \{W_k\} \sim \text{Poisson}\left(\sum_k \pi_k W_k([i-1, i]) W([j-1, j])\right),
$$

to replace (54) in Appendix C.1 of Caron and Fox (2017). Consequently, (55) holds because for any $k$ we have

$$
W_k([0, \alpha]) / W_0([0, \alpha]) \rightarrow 1 \quad \text{almost surely as } \alpha \rightarrow \infty. \quad (B.1)
$$

Second, we show that Theorem 4 in Caron and Fox (2017) also holds for DMPGM. Specifically, (59) becomes

$$
X_n | \{W_k^{(2)}\} = \text{Poisson}\left[\frac{1}{2} \phi\{W(s_n^{(2)})\}\right],
$$

so that (62) in (Caron and Fox, 2017) can be achieved by (B.1). Finally, with others part of the proof unchanged, Theorem 1 for DMPGM is proved.

### D Dataset Details

We present the following summary statistics for the datasets in Table 2.
E Inference results for DMPGM

The log likelihood and the number of edge clusters in the training process are shown in Figure 4 and Figure 5, respectively. The inferred edge multiplicity is shown in Figure 6.

F Hyperparameters

We list the hyperparameters used in our experiments in Table 3 below.

G Data and Code

We obtained the datasets from the publically available source https://pytorch-geometric.readthedocs.io/en/latest/modules/datasets.html. All data do not contain personally identifiable information or offensive content. We conducted our experiments on a c5d.4xlarge instance on the AWS EC2 platform, with 16 vCPUs and 32 GB RAM. The codes for training conventional GNNs are from https://github.com/VITA-Group/Deep_GCN_Benchmarking under MIT license.
Figure 5: Number of clusters inferred by DMPGM over the course of the MCMC chain.

Figure 6: Histograms of the expected multiplicity of virtual edges formed in the EEGNN framework using each dataset.
Table 1: Results on real datasets: mean accuracy (%) ± standard deviation (%)

(a) Number of layers: 2

|        | Cora       | Citeseer   | PubMed     | Texas      | Wisconsin  | Cornell   |
|--------|------------|------------|------------|------------|------------|-----------|
| SGC    | 77.01±0.34 | 69.18±0.35 | 75.46±0.28 | 56.16±4.99 | 48.59±6.59 | 57.84±2.76 |
| EE-SGC | 76.78±0.29 | 69.60±0.37 | 75.80±0.21 | 61.24±6.48 | 53.45±8.95 | 58.92±3.55 |
| APPNP  | 82.22±0.39 | 71.73±0.76 | 79.41±0.48 | 61.41±5.27 | 52.55±7.44 | 57.73±2.74 |
| EE-APPNP | 81.48±0.47 | 71.45±0.54 | 78.90±0.52 | 66.80±3.74 | 66.23±2.93 | 60.17±6.00 |
| GCNII  | 82.21±0.67 | 67.65±0.96 | 77.91±1.71 | 61.35±8.18 | 72.51±4.91 | 74.22±8.75 |
| EE-GCNII | 81.94±0.51 | 81.48±0.59 | 77.30±0.97 | 64.22±9.02 | 70.94±6.10 | 75.68±9.78 |

(b) Number of layers: 16

|        | Cora       | Citeseer   | PubMed     | Texas      | Wisconsin  | Cornell   |
|--------|------------|------------|------------|------------|------------|-----------|
| SGC    | 73.11±0.43 | 67.79±0.56 | 70.45±0.17 | 56.27±4.92 | 48.63±6.62 | 57.84±2.76 |
| EE-SGC | 73.07±0.34 | 68.55±0.35 | 70.60±0.25 | 59.96±6.46 | 50.59±8.72 | 57.84±2.76 |
| APPNP  | 83.70±0.20 | 72.51±0.52 | 80.42±0.30 | 60.76±5.05 | 53.29±7.09 | 57.68±2.79 |
| EE-APPNP | 83.47±0.66 | 73.20±0.92 | 77.90±0.36 | 66.08±4.62 | 66.08±3.17 | 61.30±7.18 |
| GCNII  | 84.77±0.37 | 72.30±0.80 | 78.60±0.52 | 66.38±8.69 | 70.71±2.40 | 74.49±8.98 |
| EE-GCNII | 84.10±0.57 | 72.50±1.40 | 78.81±0.66 | 73.30±3.85 | 78.94±4.90 | 75.24±8.08 |

(c) Number of layers: 32

|        | Cora       | Citeseer   | PubMed     | Texas      | Wisconsin  | Cornell   |
|--------|------------|------------|------------|------------|------------|-----------|
| SGC    | 59.94±0.56 | 66.17±0.50 | 68.97±0.19 | 56.16±4.99 | 48.59±6.59 | 57.84±2.76 |
| EE-SGC | 66.46±0.83 | 67.68±0.44 | 70.68±0.68 | 59.46±6.16 | 50.59±8.72 | 58.92±3.15 |
| APPNP  | 83.55±0.50 | 72.11±0.64 | 80.22±0.34 | 61.57±5.28 | 52.71±7.34 | 57.78±2.75 |
| EE-APPNP | 83.79±0.39 | 72.47±0.53 | 79.23±0.27 | 66.00±4.33 | 66.39±3.09 | 74.86±7.84 |
| GCNII  | 85.34±0.53 | 73.26±0.86 | 79.89±0.33 | 70.49±5.48 | 69.06±2.70 | 74.05±8.56 |
| EE-GCNII | 85.70±0.41 | 73.45±1.40 | 79.72±0.43 | 75.24±3.72 | 79.37±0.43 | 74.86±7.84 |

(d) Number of layers: 64

|        | Cora       | Citeseer   | PubMed     | Texas      | Wisconsin  | Cornell   |
|--------|------------|------------|------------|------------|------------|-----------|
| SGC    | 25.65±1.93 | 63.08±0.52 | 40.98±1.73 | 56.16±4.99 | 48.55±6.58 | 57.84±2.76 |
| EE-SGC | 35.54±1.36 | 65.42±0.17 | 64.28±0.82 | 59.46±6.16 | 50.31±8.42 | 58.92±3.15 |
| APPNP  | 83.58±0.49 | 72.10±0.48 | 80.42±0.42 | 61.19±5.29 | 53.06±7.10 | 57.68±2.74 |
| EE-APPNP | 83.76±0.41 | 72.16±0.65 | 79.94±0.22 | 66.00±4.08 | 66.63±3.12 | 61.75±7.43 |
| GCNII  | 85.46±0.31 | 73.44±1.00 | 80.08±0.37 | 69.57±5.70 | 68.63±1.05 | 73.19±8.83 |
| EE-GCNII | 85.54±0.59 | 72.24±1.26 | 79.93±0.46 | 75.62±3.65 | 76.57±3.89 | 73.26±7.39 |
Table 2: Graph datasets statistics.

| Dataset  | Cora  | Citeseer | PubMed | Texas | Wisconsin | Cornell |
|----------|-------|----------|--------|-------|-----------|---------|
| Nodes    | 2,708 | 3,327    | 19,717 | 183   | 183       | 183     |
| Edges    | 5,429 | 4,732    | 44,338 | 309   | 499       | 295     |
| Ave. Degree | 3.88 | 2.84     | 4.50   | 3.38  | 5.45      | 3.22    |
| Features | 1,433 | 3703     | 500    | 1703  | 1703      | 1703    |
| Classes  | 7     | 6        | 3      | 5     | 5         | 5       |

(a) Hyperparameters for SGC and EE-SGC.

| Dataset  | Cora  | Citeseer | PubMed | Texas | Wisconsin | Cornell |
|----------|-------|----------|--------|-------|-----------|---------|
| dim_hidden | 64   | 256     | 256    | 64    | 64        | 64      |
| alpha    | 0.1  | 0.1     | 0.1    | 0.1   | 0.1       | 0.1     |
| weight_decay | 0.0005 | 0.0005 | 0.0005 | 0.0001 | 0.0005   | 0.0005 |
| lr       | 0.01 | 0.01    | 0.01   | 0.01  | 0.01      | 0.01    |
| dropout  | 0.6  | 0.7     | 0.6    | 0.5   | 0.5       | 0.5     |

(b) Hyperparameters for APPNP and EE-APPNP.

| Dataset  | Cora  | Citeseer | PubMed | Texas | Wisconsin | Cornell |
|----------|-------|----------|--------|-------|-----------|---------|
| dim_hidden | 64   | 64       | 64     | 64    | 64        | 64      |
| alpha    | 0.1  | 0.1      | 0.1    | 0.1   | 0.1       | 0.1     |
| lr       | 0.01 | 0.01     | 0.01   | 0.01  | 0.01      | 0.01    |
| dropout  | 0.   | 0.       | 0.     | 0.    | 0.        | 0.      |
| weight_decay1 | 0.005 | 0.005 | 0.005 | 0.005 | 0.005   | 0.005 |
| weight_decay2 | 0.   | 0.       | 0.     | 0.    | 0.        | 0.      |
| embedding_dropout | 0.5  | 0.5      | 0.5    | 0.5   | 0.5       | 0.5     |

(c) Hyperparameters for GCNII and EE-GCNII.

Table 3: Hyperparameters in the training.
Acknowledgments

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