Edge Dithering for Robust Adaptive Graph Convolutional Networks

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Abstract—Graph convolutional networks (GCNs) are vulnerable to perturbations of the graph structure that are either random, or, adversarially designed. The perturbed links modify the graph neighborhoods, which critically affects the performance of GCNs in semi-supervised learning (SSL) tasks. Aiming at robustifying GCNs conditioned on the perturbed graph, the present paper generates multiple auxiliary graphs that perform SSL. Robustness is enabled through learnable graph-combining weights along with suitable regularizers. Relative to GCN, the novel AGCN achieves markedly improved performance in tests with noisy inputs, graph perturbations, and state-of-the-art adversarial attacks. Further experiments with protein interaction networks showcase the competitive performance of AGCN for SSL over multiple graphs.

I. INTRODUCTION

A task of major importance at the cross-roads of machine learning and network science is semi-supervised learning (SSL) over graphs. SSL aims at predicting nodal labels given: i) the graph connections; ii) feature vectors at all nodes; and iii) labels only at a subset of nodes. This partial label availability may be attributed to privacy concerns (e.g., with medical data); energy considerations (e.g., with wireless sensor networks); or unrated items (e.g., with recommender systems).

Standard SSL schemes typically assume that the available labels and graph connections have certain properties such as smoothness, which asserts that connected nodes have similar attributes [1]. In various scenarios however, robustness issues arise. Powerful adversaries manipulate nodal attributes and connections to bias learning, and promote their malicious goals [2]. Further, human annotators or noisy data introduce errors during the graph construction that leads to perturbed edge weights [3]. Adversarially perturbed or simply anomalous graph data may degrade the performance of SSL algorithms with severe consequences. The recent era of misinformation and “fake” news calls for robust machine learning algorithms for network science [4], [5], [6]. In this context, a novel robust GCN framework is introduced here that utilizes edge-dithered auxiliary graphs, which are combined using learnable weights.

A. Related work

Graph-based SSL methods typically assume that the true labels are “smooth” with respect to the underlying network structure, which naturally motivates leveraging the topology of the network to propagate the labels and increase classification performance. Graph-induced smoothness may be captured by kernels on graphs [7], [1]; or Gaussian random fields [8]. Graph convolutional networks (GCNs) incorporate the graph structure to achieve state-of-the-art results in SSL tasks [9], [10], [11], [12].

With the success of GCNs on graph learning tasks granted, recent results indicate that perturbations of the graph topology or nodal features can severely deteriorate their classification performance [2], [13], [14]. Structural attacks target a subset of nodes and modify their links to promote mis-classification of targeted nodes [15]. The designed graph perturbations are “un-noticeable”, which is feasible so long as the degree distribution of the perturbed graphs are similar to the initial distribution [2]. GCNs learn nodal representations by extracting information within local neighborhoods. Adversaries poison the learned features by perturbing the node’s neighborhood. Hence, the vulnerability of GCNs challenges their deployment to critical applications dealing with security or healthcare, where robust learning is of major importance. Defending against adversaries may unleash the potential of GCNs and broaden the scope of machine learning applications altogether. Recent works robustify GCNs against structural perturbations by utilizing the nodal features [15], [16]. Gaussian regularizers are employed in [16] to protect the network from adversarial attacks. Jaccard similarity among features is utilized in [15] to prune perturbed edges. However, these methods are challenged in the absence of nodal feature vectors.

B. Contributions

The present paper develops a framework for robust deep learning over perturbed graphs. Specifically, the contribution of this work is threefold.

C1. Given the perturbed unweighted graph and aiming at robust SSL, multiple auxiliary graphs are drawn by dithering (adding or removing) edges with probabilities selected to boost robustness. The novel edge-dithering (ED) approach reconstructs the original neighborhood structure with high probability (whp) as the number of sampled graphs increases. ED can be applied even in the absence of nodal features.
C2. A weighted combination of the auxiliary ED graphs is employed across GCN layers. Per layer weights are adapted to promote those ED graphs that maximally avoid the adversarially perturbed edges. Further, a residual feed of the data is utilized to facilitate diffusion of the features across the graph. Robust graph-based regularizers are also included to prevent overfitting, and further account for the underlying graph topology.

C3. Numerical tests involving noisy features, noisy edge weights, and random as well as adversarial edge perturbations showcase the merits of the novel approach. The proposed ED-AGCN achieves also competitive results in predicting protein functions over multiple graphs.

II. MODELING AND PROBLEM FORMULATION

Consider a graph $G := (V, A)$ of $N$ nodes, with $V := \{v_1, \ldots, v_N\}$ denoting the vertex set, and $A$ the $N \times N$ adjacency matrix capturing edge connectivity through $A_{n,m}$ that is 1 if an edge connects $v_n$ and $v_m$, and 0 otherwise. The neighborhood of $v_n$ is

$$N_n := \{n' : A_{n,n'} \neq 0, \quad v'_n \in V\}. \quad (1)$$

The perturbed graph is $\tilde{G} := (V, \tilde{A})$ with corresponding adjacency $\tilde{A} := A + \Delta$ having entries perturbed by

$$\tilde{A}_{n,n'} = \begin{cases} 
1, & \text{if } A_{n,n'} = 0 \text{ and } \tilde{A}_{n,n'} = 1 \\
-1, & \text{if } A_{n,n'} = 1 \text{ and } \tilde{A}_{n,n'} = 0 \\
0, & \text{otherwise}
\end{cases} \quad (2)$$

where +1 corresponds to edge insertion, −1 to edge deletion, and 0 to no perturbation. Evidently, these links may drastically degrade the performance of SSL methods since the neighborhood is either adversarially or randomly modified [2]. The adversarial attacks aim at unnoticeable changes, a constraint that limits the number of perturbations. Hence, the number of perturbed links (nonzero elements in $\tilde{A}$) is small relative to the original number of edges in $G$.

Associated with the $n$-th node can be an $F \times 1$ feature vector $x_n$. These vectors are collected in the $N \times F$ feature matrix $X := [x_1^T, \ldots, x_N^T]^T$, where $X_{n,j}$ may denote, for example, the salary of the $n$-th individual in the LinkedIn social network. Let also $y_n \in \{0, 1, \ldots, K-1\}$ denote the label of node $n$, which may represent, for example, the education level of a person. The $N \times K$ matrix $Y$ is the “one-hot” representation of the nodal labels belonging to $K$ classes, that is, if $y_n = k$ then $Y_{n,k} = 1$ and $Y_{n,k'} = 0, \forall k' \neq k$.

Goal. Given the perturbed topology $A$, the features in $X$, and labels only at a subset $\mathcal{L}$ of nodes $\{y_n\}_{n \in \mathcal{L}}$ with $\mathcal{L} \subset V$, the goal of this paper is to design robust GCN architectures that are minimally affected by the perturbed edges.

III. EDGE DITHERING

The ever-expanding interconnection of social, email, and media service platforms presents an opportunity for adversaries manipulating networked data to launch malicious attacks [4], [5], [2]. Perturbed edges modify the graph neighborhoods, which leads to significant degradation in the performance of GCNs. Aiming to restore a node’s initial graph neighborhood an edge-dithering (ED) module is developed in this section, where auxiliary graphs are created with probabilities designed to enhance robustness. Dithering in visual and audio applications, refers to intentional injection of noise so that the quantization error is converted to random noise, which can be easily handled [17].

Permeating the benefits of dithering towards robustify GCNs, we generate ED graphs $\{G_i\}_{i=1}^I$, where $G_i := (V, A_i)$. Each auxiliary graph $G_i$ is a dithered version of the perturbed graph $\tilde{G}$, where the edges in $A_i$ are selected in a probabilistic fashion as follows

$$A_{n,n',i} = \begin{cases} 
1 \text{ wp. } q_1 \delta(\tilde{A}_{n,n'} = 1 - q_2)\delta(\tilde{A}_{n,n'} = 0), & \delta(\tilde{A}_{n,n'} = 1) \\
0 \text{ wp. } q_2 \delta(\tilde{A}_{n,n'} = 0)\delta(\tilde{A}_{n,n'} = 1), & \delta(\tilde{A}_{n,n'} = 0)
\end{cases} \quad (3)$$

where $\delta(\cdot)$ is the indicator function, $q_1 := \Pr(\tilde{A}_{n,n'} = 1|\tilde{A}_{n,n'} = 1)$ and $q_2 := \Pr(\tilde{A}_{n,n'} = 0|\tilde{A}_{n,n'} = 0)$. If $n$ and $n'$ are connected in $G$, the edge connecting $n$ with $n'$ is deleted with probability $1 - q_1$. Otherwise, if $n$ and $n'$ are not connected in $G$ i.e. $(\tilde{A}_{n,n'} = 0)$, an edge between $n$ and $n'$ is inserted with probability $1 - q_2$.

Hence, the $i$th ED graph neighborhood of $v_n$ is

$$\mathcal{N}_{n}^{(i)} := \{n' : A_{n,n',i} \neq 0, \quad v'_n \in V\}. \quad (4)$$

The ED graphs give rise to different neighborhoods $\mathcal{N}_{n}^{(i)}$, and the role of ED is to ensure that the unperturbed neighborhood of each node will be present with high probability (whp) in at least one of the $I$ graphs. The ensuing remarks assert that this will happen whp as $I$ increases.

Remark 1: With high probability, there exists $G_i$ such that a perturbed edge will be restored to its initial value. This means that there exists an ED graph $i$ such that $A_{n,n',i} = A_{n,n'}$.

Since, each $G_i$ is independently drawn, it holds that

$$\Pr\left(\bigcup_{i=1}^I (A_{n,n',i} = 0)\big| A_{n,n'} = 1, A_{n,n'} = 0\right) = 1 - q_1$$
$$\Pr\left(\bigcup_{i=1}^I (A_{n,n',i} = 1)\big| A_{n,n'} = 0, A_{n,n'} = 1\right) = 1 - q_2$$

Remark 2: Whp there exists $G_i$ which will recover the original neighborhood structure of a node, i.e. $\mathcal{N}_{n}^{(i)} = \mathcal{N}_{n}$. The proof of this remark is included in the Appendix.

The high probability claims asserted in Remarks 1 and 2 hold as $I$ increases. Nevertheless, experiments with adversarial attacks demonstrate that even with a small $I$ the use of ED significantly boosts classification performance. The operation of the ED module is detailed in Fig. 1. Note that the proposed ED does not require availability of nodal feature vectors. The generated graphs have to be processed by a dedicated architecture that promotes the learned features from unperturbed nodal neighborhoods.

IV. ADAPTIVE GCN WITH EDGE DITHERING

The ED module generates $\{G_i\}_{i=1}^I$, which along with the perturbed graph $\tilde{G}$ will be judiciously combined to obtain a robust learning architecture. Typically, deep or shallow learning over graphs considers that the relation among the
output features at our AGCN architecture and training are presented.

To the graph that supports the data [10], [9], [21], [22]. Next, linearly combined local features the idea is to progressively capture a specific form of social interaction, such as friendship, a conveniently parametrized linear transformation, a scalar L

A novel GCN that adapts to multiple relations and enhances combination of the auxiliary ED graphs, this section develops ˇ

While the entries of h(l) depend only on the one-hop neighbors of n (one-hop diffusion), successive application of this operation will increase the diffusion range, spreading the information across the network. Generalizing to neighborhoods with larger diameter, consider the kth power of the adjacency matrix A(k). Indeed, the vector A(k)x holds the linear combinations of the values of x in the k-hop neighborhood [9].

After defining the matrices A(k) for k = 1, . . . , K, i = 1, . . . , I, consider the following parametrized mapping

where the learnable coefficients {c(i,k) }Kk=1 weight the effect of the corresponding k-th hop neighbors for relation i. At the l-th layer, the coefficients {c(i,k) }Kk=1,l=1 are collected in the K × I matrix C(l). The proposed map in (6) aggregates the diffused features in the K-hop neighborhoods per i; see also Fig. 2.

The extracted feature vector h(l) captures the diffused features per ED graph i. Aiming at robustness, the learning algorithm should promote features originating from non-perturbed graph neighborhoods. Towards this end, a graph adaptive module is developed that combines h(l) across i′ as

where R(l)i,i′′,n mixes features across ED graphs. A key contribution of this paper is viewing {R(l)i,i′′,n}i,i′′,n as training parameters, which allows AGCN to learn how to combine the different relations encoded by the ED graphs. This characteristic enables the novel AGCN to navigate through the ED graphs {G(n)}n=1, and assign larger weights to features originating from non-perturbed neighborhoods.

The graph adaptive module in (7) allows for different R(l)i,i′′,n per n. Considering the same R for each n, that is R(l)i,i′′,n = R(l)i,i′′, results in a design with less parameters at the expense of reduced flexibility. On the other hand, the flexible design

Fig. 1: ED in operation on a perturbed social network among voters. Black solid edges are the true links and dashed red edges represent adversarially perturbed links.

A. Per layer operation

Consider a hidden layer (say the lth one), whose output is the N × I × P(l) tensor Z(1) that holds the P(l) × 1 feature vectors Z(l,i), ∀n,i, with P(l) being the number of output features at l. Similarly, let Z(l) = 1 represent the input to this lth layer.

The mapping from Z(l−1) to Z(l) can be split into two steps. A linear one designed to map the tensor Z(l−1) to the tensor Z(l). The latter is then processed elementwise to obtain ˇ

Of critical importance is the design of the linear map from Z(l−1) to Z(l) that is tailored to our ED-based setup. Convolutional (C)NNs typically consider a small number of trainable weights, and then generate the linear output by

Convolving the input with these weights [20]. The convolution combines values of close-by inputs (consecutive time instants, or neighboring pixels), and thus extracts information of local neighborhoods.

GCNs generalize CNNs to operate on graph data by replacing the convolution with a graph filter whose parameters are also learned [10], [9]. This preserves locality, reduces the degrees of freedom of the map, and leverages the graph structure.

Neighborhood aggregation module. First, a neighborhood aggregation module is considered that combines linearly the nodal features within a graph neighborhood. Since the neighborhood depends on the particular ED graph (4), the combined n-th feature of the i-th graph is

where

Aiming at multi-layer graphs [18].

Many works refer to these as multi-layer graphs [18].
Fig. 2: The neighborhood aggregation module (NAM) combines features \( \{x_n\} \) of the multiple ED graphs. For this example, the 2-hop neighborhood is considered \( (K = 2) \). Different graphs define different local neighborhoods.

in (7) allows large weights \( R_{i,n',n} \) for neighborhoods without corrupted edges, even if \( G_i \) is perturbed.

**Feature aggregation module.** Next, the graph adaptive features \( g_{n,i}^{(l)} \) are mixed using learnable vectors \( w_{n,i,p}^{(l)} \) to obtain

\[
Z_{n,i,p}^{(l)} := g_{n,i}^{(l)} w_{n,i,p}^{(l)}, \quad i = 1, \ldots, I, \ n = 1, \ldots, N, \ p = 1, \ldots, P^{(l)}.
\]

The \( P^{(l-1)} \times N \times I \times P^{(l)} \) tensor \( W^{(l)} \) collects the feature mixing weights \( \{w_{n,i,p}^{(l)}\} \), while the \( I \times I \times P^{(l)} \) tensor \( R^{(l)} \) collects the graph mixing weights \( \{R_{i,n',n}^{(l)}\} \). Upon collecting all the scalars \( \{Z_{n,i,p}^{(l)}\} \) in \( Z^{(l)} \) (6)-(8) reduce to

\[
Z^{(l)} := f(Z^{(l-1)}; \theta^{(l)}_z), \quad \theta^{(l)}_z := [\text{vec}(W^{(l)}); \text{vec}(R^{(l)}); \text{vec}(C^{(l)})]^T.
\]

**Residual GCN layer.** Concatenating \( L \) GCN layers diffuses the input \( X \) across the \( L \)-hop graph neighborhood, cf. (5). However, the exact size of the relevant neighborhood is not always known a priori. To endow our architecture with increased flexibility, a residual GCN layer is introduced that inputs \( X \) at each \( l \), and thus captures multiple types of diffusion\(^2\). As a result, the linear operation in (9) is replaced by the residual linear tensor mapping

\[
Z^{(l)} := f(Z^{(l-1)}; \theta^{(l)}_z) + f(X; \theta^{(l)}_z)
\]

where \( \theta^{(l)}_z \) encodes trainable parameters, cf. (10). When viewed as a map from \( X \) to \( Z^{(l)} \), the operator in (11) implements a broader class of graph diffusions than the one in (9). If \( l = 3 \) and \( K = 1 \) for example, the first summand in (11) is a one-hop diffusion of the input that corresponds to a two-hop (nonlinear) diffused version of \( X \), while the second summand diffuses \( X \) in one-hop. At a more intuitive level, the second summand also guarantees that the impact of \( X \) in the output does not vanish as the number of layers grows.

The output of our graph architecture is

\[
\hat{Y} = g(Z^{(L)}; \theta_y)
\]

where \( g(\cdot) \) is the normalized exponential function (softmax), \( \hat{Y} \) is an \( N \times K \) matrix, \( \hat{Y}_{n,k} \) represents the probability that \( y_n = k \), and \( \theta_y \) are trainable parameters.

**B. Training with graph-smooth regularizers**

The AGCN weights are estimated by minimizing the discrepancy between estimated labels and the given ones as

\[
\min_{\theta^{(l)}_i, \theta^{(l)}_g} \mathcal{L}_{tr}(\hat{Y}, Y) + \mu_1 \sum_{l=1}^L \text{Tr}(\hat{Y}^T A_l \hat{Y}) + \mu_2 \rho(\theta^{(L)}_y, \theta^{(L)}_g) + \lambda \sum_{l=1}^L \|R^{(l)}\|_1
\]

where \( \mathcal{L}_{tr}(\hat{Y}, Y) := - \sum_{n \in \mathcal{L}} \sum_{k=1}^K Y_{nk} \ln \hat{Y}_{nk} \) is the cross-entropy loss function over the labeled examples.

The first regularizer in (13) promotes smooth label estimates over the graphs [1], and \( \rho(\cdot) \) is an \( L_2 \) norm over the AGCN parameters that is used to avoid overfitting [20]. Finally, the \( L_1 \) norm in the third regularizer encourages learning sparse mixing coefficients, and hence it promotes activating only a subset of edge-dithered graphs per \( l \). ED graphs with a large number of perturbed edges will result to a higher cost function in (13). Hence, the learning algorithm will assign larger combining weights to non-perturbed topologies. The backpropagation algorithm is employed to minimize (13).

To recap, aiming at a robust GCN architecture, we introduce a novel edge-dithering module that generates probabilistically auxiliary graphs. These graphs are processed by a robust AGCN architecture that: combines features within neighborhoods originating from the different graphs; adapts to each graph by aggregating the learned features with \( \mathbf{R} \); uses a simple but versatile residual tensor mapping (11); and employs smoothness and sparsity promoting graph-based regularizers; see also Fig. 3.

**V. Experiments**

The AGCN is tested with \( L = 3, \ P^{(1)} = 64, \ P^{(2)} = 8, \) and \( P^{(w)} = K \). The regularization parameters \( \{\mu_1, \mu_2, \lambda\} \) are chosen based on the performance of the AGCN in the validation set for each experiment. For the training stage, an ADAM optimizer with learning rate 0.005 was employed [24], for 300 epochs with early stopping at 60 epochs. The goal here is to provide tangible answers to the following research questions.

**RQ1.** How does AGCN compare to state-of-the-art methods for SSL over multi-relational graphs?

**RQ2.** How robust is AGCN compared to GCN under noisy features, noisy edge weights, and random as well as adversarial edge perturbations?
**RQ3.** How sensitive is AGCN to the parameters of the ED module \(q_1, q_2\) and \(I\)?

**A. Predicting multi-relational protein functions**

This section assesses the performance of the proposed AGCN when predicting protein functions over multiple graphs. For this experiment, the given network is multi-relational and no perturbations were considered, hence the ED module is not used. Protein-to-protein interaction networks relate two proteins via multiple cell-dependent relations and protein classification seeks the unknown function of some proteins based on the functionality of a (small) subset of them [25].

Given a target function \(y_n\) that is known on a subset of proteins \(n \in \mathcal{L}\), known functions on all proteins summarized in \(X\), and the multi-relational protein networks \(A\), the goal is to predict whether proteins in the unlabeled set \(n \in \{V \setminus \mathcal{L}\}\) are associated with the target function or not. Hence, the number of target classes in this application is simply \(K = 2\). In this setting, \(A_i\) represents the protein connectivity in the \(i\)-th cell type. Examples of such cells include cerebellum, midbrain, or frontal lobe. Table I summarizes the dimensions of the three datasets used in our experiments.

Next, AGCN is compared with the GCN [9] that is the single-relational alternative, and the Mune [26] that is a state-of-the-art diffusion-based approach for SSL over multi-relational graphs. Since GCN only accounts for a single graph, GCN employs the graph \(i\) that achieves the best results in the validation set. Furthermore, Mune does not account for feature vectors in the nodes of the graph. Hence, to facilitate fair comparison, we also employ our AGCN without using the feature vectors, i.e. \(X = I_N\). Finally, since the classes are heavily unbalanced, we evaluate the performance of the various approaches using the macro F1 score for predicting the protein functions.\(^3\)

**Table I: Protein-to-protein interaction datasets**

| Dataset       | Nodes  | Features | Relations |
|---------------|--------|----------|-----------|
| Generic cells | 4,487  | 502      | 144       |
| Brain cells   | 2,702  | 81       | 9         |
| Circulation cells | 3,385 | 62       | 4         |

**Table II: Macro F1 for the brain cells dataset.**

| \(|\mathcal{L}|\) | 440 | 220 | 110 | 55  |
|------------------|-----|-----|-----|-----|
| AGCN             | 0.86| 0.79| 0.71| 0.69|
| GCN              | 0.49| 0.48| 0.48| 0.47|
| AGCN (No feat.)  | 0.41| 0.43| 0.41| 0.35|
| Mune (No feat.)  | 0.27| 0.27| 0.32| 0.14|

**Table III: Macro F1 for the circulation cells dataset.**

| \(|\mathcal{L}|\) | 440 | 220 | 110 | 55  |
|------------------|-----|-----|-----|-----|
| AGCN             | 0.77| 0.76| 0.70| 0.69|
| GCN              | 0.48| 0.48| 0.48| 0.47|
| AGCN (No feat.)  | 0.41| 0.42| 0.40| 0.35|
| Mune (No feat.)  | 0.28| 0.27| 0.26| 0.13|

**Table IV: Macro F1 for the generic cells dataset.**

| \(|\mathcal{L}|\) | 440 | 220 | 110 | 55  |
|------------------|-----|-----|-----|-----|
| AGCN             | 0.70| 0.66| 0.60| 0.58|
| GCN              | 0.49| 0.48| 0.48| 0.47|
| AGCN (No feat.)  | 0.40| 0.44| 0.41| 0.43|
| Mune (No feat.)  | 0.28| 0.25| 0.24| 0.13|

Tables II-IV report macro F1 values for the aforementioned approaches for varying number of observed (labeled) nodes \(|\mathcal{L}|\). The results for all datasets demonstrate that: i) the macro F1 score improves as \(|\mathcal{L}|\) increases; ii) AGCN, that judiciously combines the multiple-relations, outperforms the GCN by a large margin; and iii) for the case where nodal features are not used (bottom two rows of each table), AGCN outperforms the state-of-the-art Mune.

**B. Robustness to additive Gaussian input noise**

This section reports the performance of the proposed ED-AGCN architecture under noisy graphs or features. For this experiment, the ionosphere dataset is considered, which contains \(N = 351\) data points with \(F = 34\) features that belong

\(^3\)Accurate classifiers achieve macro F1 values close to 1.
to \( K = 2 \) classes [27]. In this case, the graphs \( G^\kappa \) are formed using \( \kappa \)-nearest neighbors graphs for different values of \( \kappa \) (i.e., different number of neighbors). This method computes the link between \( n \) and \( n' \) based on the Euclidean distance of their features \( \| x_n - x_{n'} \|^2 \).

Oftentimes, the available topology and feature vectors might be corrupted with noise. Capturing this noise, \( A_{\Delta} \) and \( X \) are obtained as \( A_{\Delta} = A_{\Delta_r} + O_{\Delta} \) and \( X = X_{tr} + O_X \), where \( A_{\Delta_r} \) and \( X_{tr} \) represent the true multi-relational topology and features and \( O_{\Delta} \) and \( O_X \) denote the corresponding additive perturbations. We draw \( O_{\Delta} \) and \( O_X \) from a zero mean white Gaussian distribution with specified signal to noise ratio (SNR). Since random additive noise is considered here, the ED module is not employed.

Fig. 4 reports the SSL classification performance for the ionosphere dataset of AGCNs. The AGCN is tested with different values of \( \kappa \), where \( \kappa = 5, 10 \) corresponds to a two relational graph, i.e. \( I = 2 \). We deduce that multiple \( \kappa \)-nearest neighbors graphs lead to learning more robust representations of the data, which testifies to the merits of proposed multi-relational architecture.

**C. Robustness to Bernoulli noise on edges**

This experiment tests our architecture with four network datasets [28]: “Cora” (\( N = 2708, K = 7, |\mathcal{L}| = 140 \)), “Citeseer” (\( N = 3327, K = 6, |\mathcal{L}| = 120 \)) and “Pubmed” (\( N = 19717, K = 3, |\mathcal{L}| = 30 \)) are citation graphs, while “Polblogs” (\( N = 1224, K = 2, |\mathcal{L}| = 24 \)) is a political blog network. To facilitate comparison, we reproduce the same experimental setup than in [9], i.e., the same split of the data in train, validation, and test sets. To study the effect of graph perturbations on the neural network architecture, the feature vectors of the citation datasets are not used. Notice that our robust GCN architecture can be applied even in the absence of nodal features, whereas existing approaches are not directly applicable [15]. For this experiment, the perturbed graph \( \hat{\mathbf{A}} \) is generated by inserting new edges in the original graphs between a random pair of nodes \( n, n' \) that are not connected in \( \mathbf{A} \), i.e. \( A_{n,n'} = 0 \). The added edges can be regarded as drawn from Bernoulli distribution. AGCN utilizes the multiple graphs generated via the ED module with \( I = 10 \), and \( q_1 = 0.9 \), and \( q_2 = 1 \) since no edge is deleted in \( \hat{\mathbf{A}} \).

Fig. 4 reports the SSL classification accuracy of AGCN with \( |\mathcal{L}| = 50 \) for noisy features (left) and noisy graphs (right).

**TABLE V: Classification accuracy in percent for nodes in \( \mathcal{I} \) for different numbers of attacked nodes.**

| Dataset      | Method | Number of attacked nodes \( |\mathcal{I}| \) | 20 | 30 | 40 | 50 | 60 |
|--------------|--------|---------------------------------|----|----|----|----|----|
| Citeseer     | GCN    |                                 | 60.49 & 56.00 & 61.49 & 56.39 & 58.99 |    |    |    |
|              | AGCN   |                                 | 70.99 & 56.00 & 61.49 & 61.20 & 58.66 |    |    |    |
| Cora         | GCN    |                                 | 76.00 & 74.66 & 76.00 & 62.39 & 73.66 |    |    |    |
|              | AGCN   |                                 | 78.00 & 82.00 & 84.00 & 73.59 & 74.99 |    |    |    |
| Pubmed       | GCN    |                                 | 74.00 & 71.33 & 68.99 & 66.40 & 69.66 |    |    |    |
|              | AGCN   |                                 | 72.00 & 75.36 & 71.44 & 68.50 & 74.43 |    |    |    |
| Polblogs     | GCN    |                                 | 85.03 & 86.00 & 84.99 & 78.79 & 86.91 |    |    |    |
|              | AGCN   |                                 | 84.00 & 88.00 & 91.99 & 78.79 & 92.00 |    |    |    |

**D. Robustness to adversarial attacks on edges**

The original graphs in Cora, Citeseer, Pubmed, and Polblogs were perturbed using the adversarial setup in [2], where structural attacks are effected on attributed graphs. These attacks perturb connections adjacent to \( \mathcal{I} \) set of targeted nodes by adding or deleting edges [2]. Our ED module uses \( I = 10 \) sampled graphs with \( q_1 = 0.9 \), and \( q_2 = 0.999 \). For this experiment, \( 30\% \) of the nodes are used for training, \( 30\% \) for validation and \( 40\% \) for testing.

Table V reports the classification accuracy of GCN and AGCN for different number of attacked nodes (\(|\mathcal{I}|\)). Different from Fig. 5 where the classification accuracy over the test set is reported, Table V reports the classification accuracy over the set of attacked nodes \( \mathcal{I} \). It is observed that the proposed AGCN is more robust relative to GCN under adversarial attacks [2]. This finding justifies the use of the novel ED in conjunction with the AGCN that judiciously selects extracted features originating from non-corrupted neighborhoods.

**Parameter sensitivity analysis**. Fig. 6 includes sensitivity of the AGCN to varying parameters of the ED module for the experiment in Table V with the Cora and \( |\mathcal{I}| = 30 \). It is observed that the AGCN’s performance is relative smooth for certain ranges of the parameters. In accordance with Remark 2, notice that even for small I AGCN’s performance is increased significantly.

**VI. CONCLUSIONS**

This work advocates a novel deep learning approach to robust SSL over perturbed graphs. It relies on random dithering applied to edges with probabilities selected to restore a node’s

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\(^4\)The nodes in \( \mathcal{T} \) are in the testing set.
original neighborhood with high probability. The auxiliary edge-dithered graphs are combined and jointly exploited by an adaptive GCN. The latter assigns larger combining weights to learned features extracted from graph neighborhoods without perturbed edges. Experiments demonstrate the performance gains of AGCN in the presence of noisy features, noisy edge weights, and random as well as adversarial edge perturbations.

Fig. 5: Classification accuracy for increasing number of perturbed edges. (Left) Cora, (Middle left) Pubmed, (Middle right) Citeseer, (Right) Polblogs.

Fig. 6: SSL classification accuracy of AGCN under varying edge creation prob. $q_1$, edge deletion prob. $q_2$, and number of samples $I$.

REFERENCES

[1] A. J. Smola and R. I. Kondor, “Kernels and regularization on graphs,” in Learning Theory and Kernel Machines. Springer, 2003, pp. 144–158.
[2] D. Zügner, A. Akbarnejad, and S. Günnemann, “Adversarial attacks on neural networks for graph data,” in Proc. Intl. Conf. on Knowledge Disc. and Data Mining (KDD), 2018, pp. 2847–2856.
[3] L. Akoglu, H. Tong, and D. Koutra, “Graph based anomaly detection and description: a survey,” Data mining and knowledge discovery, vol. 29, no. 3, pp. 626–688, 2015.
[4] I. J. Goodfellow, J. Shlens, and C. Szegedy, “Explaining and harnessing adversarial examples,” in Proc. Int. Conf. on Learn. Representations (ICLR), 2014.
[5] C. C. Aggarwal, “Outlier analysis,” in Data mining. Springer, 2015, pp. 237–263.
[6] Y. Yan, Z. Xu, J. W. Tsang, G. Long, and Y. Yang, “Robust semi-supervised learning through label aggregation,” in Thirtieth AAAI Conference on Artificial Intelligence, 2016.
[7] M. Belkin, P. Niyogi, and V. Sindhwani, “Manifold regularization: A geometric framework for learning from labeled and unlabeled examples,” J. Mach. Learn. Res., vol. 7, pp. 2399–2434, 2006.
[8] Z. Z. Ghahramani and J. D. Lafferty, “Semi-supervised learning using gaussian fields and harmonic functions,” in Proc. Intl. Conf. Mach. Learn. (ICML), Washington, USA, Jun. 2003, pp. 912–919.
[9] T. N. Kipf and M. Welling, “Semi-supervised classification with graph convolutional networks,” in Proc. Int. Conf. on Learn. Representations (ICLR), Toulon, France, Apr. 2017.
[10] M. M. Bronstein, J. Bruna, Y. LeCun, A. Szlam, and P. Vandergheynst, “Geometric deep learning: Going beyond euclidean data,” IEEE Sig. Process. Mag., vol. 34, no. 4, pp. 18–42, 2017.
[11] P. Veličković, G. Cucurull, A. Casanova, A. Romero, P. Lio, and Y. Bengio, “Graph attention networks,” in Proc. Int. Conf. on Learn. Representations (ICLR), 2018.
[12] K. Xu, W. Hu, J. Leskovec, and S. Jegelka, “How powerful are graph neural networks?” in Proc. Int. Conf. Learn. Representations (ICLR), 2019.
[13] K. Xu, H. Chen, S. Liu, P.-Y. Chen, T.-W. Weng, M. Hong, and X. Lin, “Topology attack and defense for graph neural networks: An optimization perspective,” in IJCAI, 2019.
[14] H. Dai, H. Li, T. Tian, X. Huang, L. Wang, J. Zhu, and L. Song, “Adversarial attack on graph structured data,” in Proc. Intl. Conf. Mach. Learn. (ICML), 2018.
[15] H. Wu, C. Wang, Y. Tsybikov, A. Docherty, K. Lu, and L. Zhu, “Adversarial examples on graph data: Deep insights into attack and defense,” in IJCAI, 2019.
[16] D. Zhu, Z. Zhang, P. Cui, and W. Zhu, “Robust graph convolutional networks against adversarial attacks,” in Proc. Intl. Conf. on Knowledge Disc. and Data Mining (KDD), 2019.
[17] R. A. Ulichney, “Dithering with blue noise,” Proceedings of the IEEE, vol. 76, no. 1, pp. 56–79, 1988.
[18] M. Kivelä, A. Arenas, M. Barthelemy, J. P. Gleeson, Y. Moreno, and M. A. Porter, “Multilayer networks,” Journal of Complex Networks, vol. 2, no. 3, pp. 203–271, 2014.
[19] S. Wasserman and K. Faust, Social Network Analysis: Methods and Applications. Cambridge university press, 1994, vol. 8.
[20] I. Goodfellow, Y. Bengio, A. Courville, and Y. Bengio, Deep Learning. MIT press Cambridge, 2016, vol. 1.
[21] X. Cao, W. Lu, and Q. Xu, “Deep neural networks for learning graph representations,” in Thirtieth AAAI Conference on Artificial Intelligence, 2016.
[22] R. Li, S. Wang, F. Zhu, and J. Huang, “Adaptive graph convolutional neural networks,” in Thirty-Second AAAI Conference on Artificial Intelligence, 2018.
[23] K. He, X. Zhang, S. Ren, and J. Sun, “Deep residual learning for image recognition,” in IEEE Conf on Comp. Vision and Pat. Rec., 2016, pp. 770–778.
[24] D. P. Kingma and J. L. Ba, “Adam: A method for stochastic optimization,” in Proc. Int. Conf. on Learn. Representations (ICLR), San Diego, CA, USA, May 2015.
[25] M. Zitnik and J. Leskovec, “Predicting multicellular function through multi-layer tissue networks,” Bioinformatics, vol. 33, no. 14, pp. 1190–1198, 2017.
[26] J. Ye and L. Akoglu, “Robust semi-supervised learning on multiple networks with noise,” in Pacific-Asia Conf. on Knowl. Disc. and Data Mining (PKDD). Springer, 2018, pp. 196–208.
APPENDIX

[27] D. Dheeru and E. Karra Taniskidou, “UCI machine learning repository,” 2017. [Online]. Available: http://archive.ics.uci.edu/ml

[28] P. Sen, G. Namata, M. Bilgic, L. Getoor, B. Galligher, and T. Eliassi-Rad, “Collective classification in network data,” AI magazine, vol. 29, no. 3, p. 93, 2008.
Remark 3: With high probability there exists $G_i$ such that a perturbed edge will be restored to its initial value. This means that there exist a graph $i$ such that $A_{n,n',i} = A_{n,n'}$. Since, each $G_i$ is independently drawn, it holds that

$$\Pr\left(\bigcup_{i=1}^{I} (A_{n,n',i} = 0) \mid \bar{A}_{n,n'} = 1, A_{n,n'} = 0\right) = 1 - q_1$$

(14)

$$\Pr\left(\bigcup_{i=1}^{I} (A_{n,n',i} = 1) \mid \bar{A}_{n,n'} = 0, A_{n,n'} = 1\right) = 1 - q_2$$

(15)

Remark 4: With high probability there exists $G_i$ which will recover the original neighborhood structure of a node, i.e. $N_n^{(i)} = N_n$.

Proof. The two neighborhood structures will be the same $N_n^{(i)} = N_n$ if and only if $A_{n,n',i} = A_{n,n'}; \forall n'$. For any edge $A_{n,n',i}$ there are 4 scenarios

1) $\bar{A}_{n,n'} = 1$ and $A_{n,n'} = 1$
2) $\bar{A}_{n,n'} = 1$ and $A_{n,n'} = 0$
3) $\bar{A}_{n,n'} = 0$ and $A_{n,n'} = 1$
4) $\bar{A}_{n,n'} = 0$ and $A_{n,n'} = 0$

It further holds that

$$\Pr\left(\bigcup_{i=1}^{I} (A_{n,n',i} = 0) \mid \bar{A}_{n,n'} = 1, A_{n,n'} = 0\right) = 1 - q_1^I$$

(16)

$$\Pr\left(\bigcup_{i=1}^{I} (A_{n,n',i} = 1) \mid \bar{A}_{n,n'} = 0, A_{n,n'} = 1\right) = 1 - q_2^I$$

(17)

$$\Pr\left(\bigcup_{i=1}^{I} (A_{n,n',i} = 0) \mid \bar{A}_{n,n'} = 1, A_{n,n'} = 1\right) = 1 - (1 - q_1)^I$$

(18)

$$\Pr\left(\bigcup_{i=1}^{I} (A_{n,n',i} = 1) \mid \bar{A}_{n,n'} = 0, A_{n,n'} = 0\right) = 1 - (1 - q_2)^I$$

(19)

Without loss of generality assume for the $N$ connections $\{A_{n,n'}\}_{n,n'=1}^{N}$ the events 1)-4) appear with the following frequency $\kappa, \lambda, \mu, \nu$. Since sampling each edge of the graph is done independently across edges and across draws we arrive to the following

$$\Pr\left(\bigcup_{i=1}^{I} (N_n^{(i)} = N_n)\right) = \left(1 - (1 - q_1)^I\right)^{\kappa} \left(1 - q_1^I\right)^{\lambda} \left(1 - q_2^I\right)^{\mu} \left(1 - (1 - q_2)^I\right)^{\nu}$$

(20)

Equations (k) with $k \leq 13$ correspond to the original manuscript while (k) with $k > 13$ correspond to the supplementary material).