# A Review of Methods for Explainable and Interpretable Graph Neural Networks

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# 1 Introduction

Graphs provide an incredibly flexible structure for modeling complex data. Data can naturally appear as graphs, like molecules. We can reduce data to a graph, such as the key points of an image. We can even use graphs to add structure, such as grammatical relationships. Graph Neural Networks (GNNs) have become a popular choice for prediction and inference on graph data. At their core, GNNs work by iteratively updating node embeddings based on information from neighboring nodes. The idea is to use the graph's structure to engineer better features. This message passing scheme allows GNNs to capture complex dependencies and patterns present within the graph structure. GNN architectures typically consist of multiple layers, each performing message passing and aggregation operations to refine the embeddings. These layers are often followed by pooling and dense prediction layers to produce the final output. Some important applications of graph classification include predicting chemical toxicity (Bai et al. 2019), classifying proteins (Gallicchio and Micheli 2019), and even detecting cancer from pathology slides (Xiao et al. 2023). While GNNs achieve remarkable predictive power, their complexity prevents the exaction of the scientific rationale. Like many deep learning models, the black box nature of GNNs prevents wide adoption. Without strong methods for understanding their predictions, GNNs are more susceptible to adversarial attacks and undetected discrimination. Inferential methods also serve to direct modeling efforts by highlighting common structures or features that may be predictive in other applications. These reasons underscore the critical importance of methods for explaining and interpreting GNN models.

Although explainability and interpretability are sometimes used interchangeably in the literature, we will adopt the distinction expressed in Yuan et al. (2022). In this article the authors say that a model is explainable if the models predictions can be reasoned post-hoc (Yuan et al. 2022). For example, the effect of a specific input variable can be estimated by either dropping or randomly permuting the variable and accessing the effect on the output (Breiman 2001). This paradigm could be directly applied to GNNs; however, researchers are often not interested in the statistical significance of tabular node and edge-level features. Most scientific questions focus on the importance of specific graph-level substructures. The challenge with GNN explanations is that in addition to computing the importance of features, the high-level features themselves must first be identified. On the other hand, a model is interpretable if the model's decision process can be readily understood by humans (Yuan et al. 2022). For example, a linear regression model is interpretable because each coefficient clearly defines the relationship between any input and output. An interpretable model is typically superior to an explainable one because, similar to a partial F-test, explanation methods might identify that certain variables enhance the model, but they cannot explain how these variables contribute. On the flip side, interpretable models may not reach the performance levels of black-box models. GNN models are generally not interpretable because the relation expressed by the parameters tends to exceed human understanding. A direct translation from traditional statistics would be a circuit type analysis, which has work for image processing convolution neural networks (Olah et al. 2020). For graphs, this would involve using coefficients on subgraphs to produce predictions. Similar to the difficulties of explanation methods, these subgraphs would have to be identified, which can be computational complex and expensive given the combinatorial nature of graphs. Overcoming these challenges has substantial scientific impacts. In applications where GNNs demonstrate strong predictive power, it enables the formulation of testable scientific hypotheses about the nature of the classification. Conversely, in cases where GNNs exhibit weak predictive power, it helps identify and understand potential misunderstandings within the model.

### 1.1 General Notation

• Any graph G can be describe by X, A, E. The node feature, matrix, edge feature matrix, and adjacency matrix respectively.

- Let  $X = [X_c, X_d]$ , where  $X_c$  is the subset of continuous node features and  $X_d$  is the subset of one-hot discrete node features. Let  $E = [E_c, E_d]$ , denoted in the same manner.
- Let n represent the number of nodes in a graph.
- For any graph, let  $\nu$  denote the set of node and  $\mathcal{E}$  denote the set of edges.
- A is a binary  $n \times n$  matrix, where A[i, j] = 1 indicates that an edge exists between nodes labeled i and j.
- Let  $\operatorname{explainee}(G;\ \Omega) = h_G^{(1)}, \dots h_G^{(L)}, \rho_G$  be an L layer GNN model with parameters  $\Omega$  that we would like to explain. Here  $h_G^{(l)}$  denotes the graph embeddings outputted by the  $l^th$  layer, and  $\rho_G$  denotes the final prediction vector.
- Let arg max  $\rho_G = \hat{Y}_G$  be the predicted class label for graph G predicted from explainee().
- Unless otherwise noted, assume any GNN model discussed is a graph-level classification model.

# 2 Analysis of Core Papers

# 2.1 GNNInterpreter (X. Wang and Shen 2024)

GNNInterpreter (X. Wang and Shen 2024) is a method for generating model-level explanations of GNN graph classification models. In general, explanation methods serve to elucidate which features within the data influence disparate predictions. These methods typically fall into two categories: instance-level and model-level. Instance-level explanations aim to unveil the model's rationale behind a particular prediction. In domains such as image and text analysis, a prevalent approach involves masking or perturbing the instance and assessing the impact on the model's prediction. On the other hand, model-level explanations seek to understand how a model generally distinguishes between classes. In image and text analysis, for instance, one common technique involves treating the input as a trainable parameter and optimizing the model's prediction towards a specific class. Consequently, the resulting optimized input comprises a set of features strongly associated with the targeted class. GNNInterpreter provides model level explanations in this manner. Formally, GNNInterpreter tries to learn the graph generating distribution for each class. GNNInterpreter works by optimizing the parameters of a generic graph generating distribution to produce samples

that closely match the explainee's understanding of the targeted class.

Graph generating distributions are hard to specify because there can be discrete and continuous elements of X, E and A. Furthermore, the interactions between these matrices can be complex. The authors tackle these issues by making two simplify assumptions. First, they assume that every graph is a Gilbert random graph (Gilbert 1959), where every possible edge as an independent fixed probability of occurring. Second, the author assume that the features of every node and edge are independently distributed. The justification of the first assumption is that the other common types of random graphs are not suitable for this application. Erdo-Renyi random graphs (ERDdS and R&wi 1959) have a fixed number of edges, which limits the diversity of explanations, Rado random graphs (Rado 1964) are infinite in size, and the random dot-product graph model is just a generalization of Gilbert's model (X. Wang and Shen 2024). The second assumption is justified by the fact that the parameters of the independent distributions will be updated jointly using the explainee model. Therefore, the explainee's understanding of the latent correlation structure should be contained in the final estimates.

Although the graphs discuss in this paper only contain discrete features,  $X_c$  and  $E_c$  can be sampled from any continuous distribution that can be expressed as a location-scale family. Separating the stochastic and systematic components is necessary for gradient based optimization. It is commonly known as the reparametrization trick. The discrete feature matrices,  $(X_d, E_d, A)$ , need to be sampled from a continuous distribution for gradient based optimization, but the distribution has to have sampling properties close to a discrete distribution. The author assume that the true underlying distribution for every discrete node and edge feature is categorical. The categorical distribution is also know as the multi-bernoulli, where every sample has a fixed probability of being in one of the discrete categories. Let  $\theta_{\rm cat}$  represent the associated vector of un-normalized or relative probabilities, where each entries is > 0. Then,

$$\operatorname{argmax} \log \theta_{\operatorname{cat}} + \operatorname{Gumbel}(0, 1) \sim \operatorname{Cat}(\theta_{\operatorname{cat}}) \tag{1}$$

The intuition is that the Gumbel distribution, which is the density of the maximum order statistic of i.i.d. standard normals, makes it a good candidate for modeling the winning or maximum

probability category. Adding Gumbel noise to the logits should maintain the true relative proportions, but introduce enough skewness such that every category has some probability of having the maximum noised logit. A proof of equation 1 is provide in section Section 4.3. Approximating the argmax function with the Softmax function allows for approximate categorical sampling that is differentiable w.r.t.  $\theta_{\text{cat}}$ . The associate inverse uniform CDF sampling formula below is referred to as the Gumbel-Softmax trick or the concrete distribution (Maddison, Mnih, and Teh 2017),

Softmax 
$$\left(\frac{\log \theta_{\text{Cat}} - \log(-\log \epsilon)}{\tau}\right), \quad \epsilon \sim U[0, 1].$$
 (2)

 $\tau$  is a hyperparameter that controls the degree of relaxation. Smaller value of  $\tau$  approximate the discrete sampling better, but can result in numerical issues. The adjacency matrix can be sampled in a similar manner since the Bernoulli is just a special case of the categorical. The sampling formula is as follows:

sigmoid 
$$\left(\frac{\log(\theta_A/(1-\theta_A)) + \log \epsilon - \log(1-\log \epsilon)}{\tau}\right), \quad \epsilon \sim U[0,1],$$
 (3)

where  $\theta_A$  is an  $n \times n$  matrix with each [i, j] entry representing the relative probability that the ij edge exists. Equation 3 samples from the binary concrete distribution (Maddison, Mnih, and Teh 2017). Taken together, let

$$G_{\rm gen} \sim {\rm gen}(\Theta)$$

notate the combined graph generating distribution, where  $\Theta$  is the set of all parameters.

An obvious objective is to maximize the likelihood that the *explainee* model predicts a sampled graph to be a member of the target class. Let  $\tilde{\rho}$  denote the desired prediction. Then the above objective can be expressed as:

$$\mathcal{L}_{\text{pred}}(\Theta \mid G_{\text{gen}}) = \mathbb{E}_{G_{\text{gen}}} \text{ CrsEnt}(\text{explainee}(G_{\text{gen}}), \ \tilde{\rho})$$
(4)

While the above objective enforces a desirable property, it fails to be restrictive enough to generate

realistic graphs. This is because the final prediction,  $\rho_{G_{\rm gen}}$  is compute using only final embeddings,  $h_{G_{\rm gen}}^{(L)}$ . Normally  $h_{G_{\rm gen}}^{(L)}$  contains all the needed information from the graphs structure; however, the generation scheme allows the feature distributions to be optimized directly. This means that an explanation can ignore the graph structure and optimize towards the desired final embeddings. It is dangerously common for nonsensical features and structures at the graph level to end up on the desired side of the decision boundary. Empirically, the authors found that even completely random graphs can produce confidence and consistent predictions.

Dataset [Method]		Predicted Class Probability by GNN	Training Time Per Class
Is_Acyclic [XGNN]	Cyclic $0.076 \pm 0.000$	Acyclic $0.927 \pm 0.000$	45 s
Is_Acyclic [Ours]	Cyclic 0.999 ± 0.001	Acyclic $1.000 \pm 0.000$	20 s
Is_Acyclic [Random]	Cyclic $0.143 \pm 0.155$	Acyclic $0.857 \pm 0.155$	-
MUTAG [XGNN]	Mutagen 0.986 ± 0.057	Nonmutagen $0.991 \pm 0.083$	128 s
MUTAG [Ours]	Mutagen 1.000 ± 0.000	Nonmutagen $1.000 \pm 0.000$	12 s
MUTAG [Random]	Mutagen 0.068 ± 0.251	Nonmutagen $0.932 \pm 0.251$	-
ColorConsistency [Ours]	Consistent 0.968 ± 0.110	Inconsistent $1.000 \pm 0.000$	58 s
ColorConsistency [Random]	Consistent $0.017 \pm 0.118$	Inconsistent $0.983 \pm 0.118$	-

Figure 1: An abridged copy of Table 5 from X. Wang and Shen (2024).

For example, Figure 1 shows that random graphs had an average predicted probability of being non-mutagenic, in the MUTAG dataset (Debnath et al. 1991), of 93.2%. In order to mitigate this issue, the authors proposed additionally minimizing the cosine distance between the average embedding of all the observed graphs from the targeted class,  $\bar{h}_{G_c}^{(L)}$ , and the embedding of the generated explanation:

$$\mathcal{L}_{\text{embed}}(\Theta \mid G_{\text{gen}}) = \mathbb{E}_{G_{\text{gen}}} \text{ CosDist}\left(\bar{h}_{G_c}^{(L)}, h_{G_{\text{gen}}}^{(L)}\right). \tag{5}$$

Additionally, the author wanted to encourage sparsity for ease of interpretation. This was done by employing an  $L_1$ ,  $L_2$ , and a budget penalty on the edge probabilities:

$$\mathcal{L}_{\text{spars}}(\theta_A) = ||\theta_A||_1 + ||\theta_A||_2 + \text{softplus}(\text{sigmoid}||\theta_A||_1 - B)^2, \tag{6}$$

where B is the expected maximum number of edges for generated explanation graphs. Connectivity is another desirable property as it ensures cohesive explanations. To encourage connectivity the author minimize the KL-Divergence between edge probabilities that share a common node.

$$\mathcal{L}_{\text{conn}}(\theta_A) = \sum_{i \in \nu} \sum_{j,k \in \mathcal{E}(i)} D_{KL}(\operatorname{sigmoid}(\theta_A[i,\ j])\ ||\ \operatorname{sigmoid}(\theta_A[i,\ k])), \tag{7}$$

where  $\mathcal{E}(i)$  is the set of edges that connect to node i.

The final generator model is trained by sampling  $G_{\text{gen}} \sim \text{gen}(\Theta)$  and then iteratively updating  $\Theta$  via gradient descent on the full loss:

$$\mathcal{L}_{\text{GNNInterpreter}}(\Theta \mid G_{\text{gen}}) = \lambda_{1} \mathcal{L}_{\text{pred}}(\Theta \mid G_{\text{gen}}) + \lambda_{2} \mathcal{L}_{\text{embed}}(\Theta \mid G_{\text{gen}}) + \lambda_{3} \mathcal{L}_{\text{spars}}(\Theta \mid G_{\text{gen}}) + \lambda_{4} \mathcal{L}_{\text{conn}}(\Theta \mid G_{\text{gen}})$$
(8)

GNNInterpreter achieves remarkable accuracy on most target classes, with most intervals in Figure 2 being tight and very close to 1. This indicates that the generated examples are almost always classified as the targeted class. The explanations for the house motif and the lollipop shape perform worse in terms of predictions. Although the authors critique the use of predictions as the sole objective, they do not employ any other quantitative metrics to evaluate the validity of their explanations.

Dataset [Method]		Predicted Class Pr	robability by GNN		Training Time Per Class
MUTAG [XGNN]	Mutagen	Nonmutagen			128 s
MOTAG [AGIVIV]	$0.986 \pm 0.057$	$0.991 \pm 0.083$			120 3
MUTAG [Ours]	Mutagen	Nonmutagen			12 s
MCTAG [Ours]	$1.000 \pm 0.000$	$1.000 \pm 0.000$			12.8
Cyclicity [Ours]	Red Cyclic	Green Cyclic	Acyclic		49 s
Cyclicity [Ours]	$1.000 \pm 0.000$	$1.000 \pm 0.000$	$1.000 \pm 0.000$		498
Motif [Ours]	House	House-X	Complete-4	Complete-5	83 s
wour [Ours]	$0.918 \pm 0.268$	$0.999 \pm 0.032$	$1.000 \pm 0.000$	$0.998 \pm 0.045$	038
Shape [Ours]	Lollipop	Wheel	Grid	Star	24 s
Shape [Ours]	$0.742 \pm 0.360$	$0.989 \pm 0.100$	$0.996 \pm 0.032$	$1.000 \pm 0.000$	24 S

Figure 2: A copy of the quantitative modeling results from Table 2 in X. Wang and Shen (2024).

Qualitatively, Figure 3 displays limitations in terms of realism. For instance, for the mutagen class, the explanation correctly identifies the importance of the  $NO_2$  group; however, the generated graph is unrealistic and might not even be chemically feasible. Additionally, the non-mutagen example

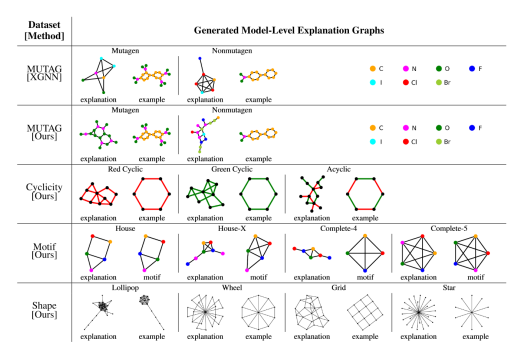


Figure 3: A copy of the qualitative modeling results from Figure 1 in X. Wang and Shen (2024).

lacks clear patterns or identifiable structures. Furthermore, the explanations for the Cyclicity dataset, as well as the Wheel and Grid classes, do not seem to be from their respective underlying data distributions. In summary, GNNInterpreter offers a method for generating example graphs that would be classified as a targeted class by a GNN model, without requiring domain-specific rules. However, optimizing for predictions and even embeddings alone does not seem to be a sufficient objective for producing *in-distribution* graphs. While the authors correctly note that focusing on predictions can lead to unrealistic graphs, they also inadvertently show that optimizing embeddings alone is not necessarily sufficient either.

#### 2.2 D4Explainer (Chen et al. 2023)

D4Explainer (Chen et al. 2023), or in-Distribution GNN Explanations via Discrete Denoising Diffusion, directly addresses the realism of generated graphs in model-level explanations by using a diffusion based generator model. In the image domain, diffusion models have proven to produce more realistic images than other generative AI methods, such as Generative Adversarial Networks (GANs). These models work by iteratively adding noise to an observation until it becomes pure noise. A denoising model is then trained to predict the noise added at any step. New observations

can be generated by reversing this process, passing pure noise through the denoising model, a technique known as reverse sampling. Additional objectives, such as a desired class label, can be added to the loss function to generate samples with certain properties.

The authors here are focused on discrete structural diffusion. D4Explainer generate example graph by noising and denoising the adjacency matrices of observed graphs. The sampled graphs have the same features, but different structures. The process of gradually adding noise to the input data is called forward diffusion. Let  $t \in [0, T]$  denote the current iteration. Let  $\beta_t$  be the common probability that any edge changes state at time step t.  $(\beta_1, \dots, \beta_T)$  is known as the variance schedule and is a set hyperparameter. Let  $A_t$  be a one-hot encoded version of the  $t^{th}$  noised adjacency. Then the forward diffusion process can be expressed as:

$$A_t[i,j] \sim q(A_t[i,j] \mid A_{t-1}[i,j]) = \operatorname{Cat}(A_{t-1}[i,j] \cdot Q_t),$$
 (9)

or

$$A_t[i,j] \sim q(A_t[i,\ j] \mid A_0[i,\ j]) = \text{Cat}\left(A_0[i,\ j] \prod_{i=1}^t Q_i\right), \tag{10}$$

where

$$Q_t = \begin{bmatrix} 1 - \beta_t & \beta_t \\ \beta_t & 1 - \beta_t \end{bmatrix},$$

the  $t^{th}$  element-wise transition matrix. The data need to approach pure noise during the final iteration. This ensures that the denoising model can start with pure noise. In the context of this problem, a completely uninformative adjacency would be a Gilbert random graph where the probability of any edge is 0.5. A proof of this property for the above diffusion process is provide in Section 4.3.

Even though only the adjacency matrix is being noised, the authors still wanted to use all of the available information during the *backwards diffusion* process. Thus the denoising model was parameterized as:

$$p(\hat{A}_0 \mid A_t, t, X_0, E_0; \Omega) \tag{11}$$

where  $\Omega$  is the set of trainable parameters. The authors employed a *Provably Powerful Graph Network* (PPGN) (Maron et al. 2020) as their denoising architecture; however, they have added an additional neural network to learn the time or noise level effect. Like most diffusion models, the primary objective is to minimize the distance between the predicted denoised observation and the original. The authors have added an additional weight term to focus the model on noisier or more difficult training examples. The core loss function is expressed as:

$$\mathcal{L}_{dist}(\Omega \mid A_0) = \mathbb{E}_{\hat{A}_0} \sum_{t=1}^{T} \left( 1 - 2\bar{\beta}_t + \frac{1}{T} \right) \text{CrsEnt} \left( A_0, \hat{A}_0 \right), \tag{12}$$

where  $\hat{A}_0 = p(\hat{A}_0 \mid A_t, t, X_0, E_0; \ \hat{\Omega}), \ A_t[i,j] \sim q(A_t[i,\ j] \mid A_0[i,\ j]),$  and

$$\bar{\beta}_t = \frac{1}{2} - \frac{1}{2} \prod_{i=1}^t (1 - 2\beta_i),$$

the cumulative transition probability. After the denoising model has been trained, model-level explanations are produced by iterative sampling candidate adjacency structures for an observed set of graph features and selecting the candidate with the highest probability of being from the targeted class. This is a multi-step procedures, where at each step  $A_t$  is denoised to k candidates  $\hat{A}_{0,1...k}$ . The candidate with the best predicted probability is then noised to a level of t-1 and the process is repeated. The pseudocode is reproduced below.

#### Algorithm 1 D4Explainer Model-level Explanation Reverse Sampling Algorithm

**Require:**  $\hat{\Omega}$ : trained denoising parameters;  $q(A_t \mid A_0)$ : forward diffusion process.

**Input:** N: maximum number of nodes; T: maximum noise level; K: number of candidates per iteration;  $\tilde{\rho}$  targeted prediction vector; (X, E): node and edge features.

**Output:** A: adjacency matrix for model-level explanation.

```
Sample A_T[1:n,1:n] Bernoulli(0.5) for t in T to 1 do Sample candidates \{\hat{A}_{0,k} \sim p(A_t,t,X,E;\ \hat{\Omega}): k \in 1,\dots,K\} Select the best candidate argmin \operatorname{CrsEnt}(\operatorname{explainee}(G=(X,\hat{A}_{0,j},E)),\ \tilde{\rho}) Sample A_{t-1}[1:n,1:n] \sim q(A_{t-1},\hat{A}_{0,j}) end for return \hat{A}_0
```

To assess how well a sampled explanation matches the observed graph distribution, the authors

compute various maximum mean discrepancy (MMD) statistics. MMD is a general technique used to measure the distance between two data distributions based solely on observed data. The distributions are estimated using kernel density estimation, and then the distance between their means is compared. As long as the chosen kernel is characteristic, an MMD statistics defined as the  $L_2$  distance between kernel mean embeddings, is representative of the distance between the true distributions (Gretton et al. 2012). The Gaussian Earth Mover's Distance kernel was selected here. The authors use MMD statistics to compare the degree, clustering, and spectrum distributions of the generated explanation against the observed data. The degree distribution of a graph indicates how frequently nodes have different numbers of connections, reflecting the overall connectivity pattern. The clustering coefficient of a node measures the proportion of the node's neighbors that are also connected to each other, representing local clustering. The spectrum distribution, which is the distribution of the eigenvalues of the adjacency matrix or the Laplacian matrix, provides insights into the graph's structural characteristics and dynamic properties. Similar to GNNInterpreter, the author of D4Explainer also value the sparsity of their generate examples. They measure the density of a graph as the number of present edges divided by the number of possible edges or

Density = 
$$|\mathcal{E}|/|\nu|^2$$
. (13)

In additional to model-level explanations, D4Explainer can provide counterfactual explanations by adding a term to the loss function. The authors define a counterfactual explanation for a particular observed graph to be  $G^c$  such that  $\hat{Y}_{G^c} \neq \hat{Y}_G$ , but the difference between  $G^c$  and G is minimal.  $\mathcal{L}_{dist}$  (12) ensures that the  $G^c$  generated does not stray too far from the observed graph distribution, and adding  $\mathcal{L}_{CF}$  minimize the probability that the generated graph is classified the same as the input.

$$\mathcal{L}_{CF}(\Omega \mid A_0) = \mathbb{E}_{\hat{A}_0} - \log(1 - \rho_{G^c}[\hat{Y}_G]), \tag{14}$$

where  $G = (X, A_0, E)$  is an observed graph,  $G^c = (X, \hat{A}_0, E)$ , and  $\rho_{G^c}[\hat{Y}_G]$  denote the probability that  $G^c$  belongs to the same class as G. To quantify the quality of generated counterfactual explanations, the author report *Counterfactual Accuracy*, *Fidelity*, and *modification ratio*. CF-ACC measure the proportion of  $G^c$ s that have different predicted labels than there associated

observed graph:

$$CF-ACC = \mathbb{E}_{\hat{A}_0} I(\hat{Y}_{G^c} \neq \hat{Y}_G). \tag{15}$$

Fidelity measures the difference in the predicted probability of G and  $G^c$  with respected to the original class label:

$$\text{Fidelity} = \mathbb{E}_{\hat{A}_0} \ \rho_G[\hat{Y}_G] - \rho_{G^c}[\hat{Y}_G]. \tag{16}$$

Finally, modification ratio measures difference in edges between  $G^c$  and G relative to the size of the original graph:

$$MR = \mathbb{E}_{\hat{A}_0} \frac{|\sum_{i,j} A[i,j] - \hat{A}_0(i,j)|}{\sum_{i,j} A[i,j]}.$$
 (17)

Similar to GNNInterpreter, the authors report the mean predicted probability for the targeted class. Using a maximum of 6 nodes, the author were able to achieve a mean of 0.832 on the MUTAG dataset. Since the counterfactual explanation are generated on a per observation basis, the authors compared their method to other popular instance-level explanation methods.

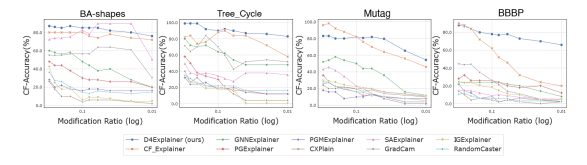


Figure 4: Comparison of the CF-ACC relative to MR of various methods and datasets. A copy of Figure 4 from Chen et al. (2023).

Figure 4 displays that D4Explainer can flip the prediction on roughly 80% of observed graphs at most modification levels. Missing from the plot is a variance measure. It is possible that the counterfactual accuracy varies more between different data splits or hyperparameter than the other methods. Figure 5 indicates that the D4Explainer generated graphs tend to have feature distributions that are closer to the original dataset; however, there might be a hidden class bias. The distributions might only be close for the majority class, skewing the averages.

		Mu	ıtag			BB	BP			NO	CI1	
Models	Deg.	Clus.	Spec.	Sum.	Deg.	Clus.	Spec.	Sum.	Deg.	Clus.	Spec.	Sum.
RamdomCaster	0.1593	0.0247	0.0417	0.2257	0.1693	0.0072	0.0397	0.2162	0.1847	1.9769	0.0404	2.2020
GNNExplainer	0.1614	0.0002	0.0409	0.2025	0.1615	0.0002	0.0395	0.2012	0.1577	0.0005	0.0405	0.1987
SAExplainer	0.0940	0.0032	0.0412	0.1384	0.1594	0.0032	0.0402	0.2028	0.189	0.0002	0.0408	0.2300
GradCam	0.1122	0.0083	0.0416	0.1621	0.0699	0.0026	0.0384	0.1109	0.1638	0.0003	0.0404	0.2045
IGExplainer	0.1292	0.0000	0.0411	0.1703	0.0908	0.0000	0.0394	0.1302	0.4288	0.0002	0.0398	0.4688
PGExplainer	0.1475	0.0002	0.0418	0.1895	0.2014	0.0018	0.0403	0.2435	0.1937	0.0000	0.0396	0.2333
PGMExplainer	0.1800	0.0002	0.0419	0.2221	0.1916	0.0003	0.0403	0.2322	0.2199	0.0000	0.0404	0.2603
CXPlain	0.1734	1.2706	0.0417	1.4857	0.1768	0.0001	0.0394	0.2163	0.1629	0.0001	0.0404	0.2034
CF-GNNExplainer	0.1172	0.0000	0.0380	0.1552	0.0870	0.0001	0.0393	0.1264	0.1224	0.0001	0.0404	0.1629
D4Explainer	0.1172	0.0000	0.0244	<u>0.1416</u>	0.0530	$\overline{0.0000}$	0.0331	0.0861	0.1006	$\overline{0.0000}$	0.0353	0.1359

Figure 5: MMD statistic for various methods and datasets copied from Table 2 in Chen et al. (2023).

#### 2.3 ProtGNN (Zhang et al. 2021)

Up until now, we have been discussing post-hoc explanation methods; however, Rudin (2019) raises two important concerns with this approach. First, parity in the predictions does not guarantee parity in the reasoning. Explanations may bear strong predictions, but for distinct and sometimes nonsensical reasons. Recall that Figure 1 showed that completely random graphs can illicit confident predictions. Second, feature importance indicates that a feature is significant to the model, but it does not reveal how the model uses the feature. For instance, a substructure might lead to a prediction due to a strong positive relation with the target class, or it could be important because of a strong negative relation with the other classes. ProtGNN (Zhang et al. 2021) uses the prototype modeling paradigm to create a GNN that is *interpretable*. Prototype based models make prediction by comparing new inputs to exemplar cases or *learned prototypes* of each class. This results in an interpretable model because, as long as the prediction function applied to the prototype similarity scores is simple, the reasoning for any given prediction will be clear. Intuitively, this can be thought of as using a black-box model to engineer features, which are then passed to a straightforward white-box model.

ProtGNN initializes m random prototype vectors for each of the C classes. Let  $p_k$  denote a prototype vector and  $P_{Y_G}$  denote the set of prototypes vectors assigned to the ground truth label of G,  $Y_G$ . Now, let  $f(G; \Omega_f) = \hat{p}_G$  be a graph encoder function that maps any graph into the prototype vector space, where  $\Omega_f$  is the set of trainable parameters. f is generally one of the standard GNN architectures. Since ProtGNN was primarily designed to be a graph classification model, the author optimized for predictive power using:

$$\mathcal{L}_{\text{prot}}(\Omega_f, \ \Omega_{\psi}, \ p_1, \dots, p_{mC} \mid G) = \text{CrsEnt}\left[\psi\left(sim\left(\hat{p}_G, p_1\right), \dots, sim\left(\hat{p}_G, p_{mC}\right); \ \Omega_{\psi}\right), \ Y_G\right], \quad (18)$$

where  $\psi$  is any simple prediction model with trainable parameters  $\Omega_{\psi}$  and sim is any vector similarity function. ProtGNN uses a full connected layer with a softmax output activation for  $\psi$ , which is similar to a multinomial regression model. To address similar out-of-distribution concerns discussed earlier, the authors include three additional loss terms as constraints. The cluster loss ensure that each embedding is close to at least one prototype assigned to its ground truth class:

$$\mathcal{L}_{\text{clst}}(\Omega_f, \ \Omega_{\psi}, \ p_1, \dots, p_{mC} \mid G) = \min_{p_k \in P_{Y_G}} ||\hat{p}_G - p_k||_2^2. \tag{19}$$

A separation loss ensures that embeddings are far from the prototypes of other classes:

$$\mathcal{L}_{\text{sep}}(\Omega_f, \ \Omega_{\psi}, \ p_1, \dots, p_{mC} \mid G) = - \min_{p_k \notin P_{Y_G}} ||\hat{p}_G - p_k||_2^2. \tag{20}$$

Finally, the diversity loss encourages each prototype within a class to learn different information:

$$\mathcal{L}_{\mathrm{div}}(p_1,\ldots,p_{mC}) = \sum_{k=1}^{C} \sum_{p_i \neq p_j \in P_k} \; \max(0,\cos(p_i,p_j) - s_{\mathrm{max}}) \tag{21} \label{eq:linear_loss}$$

where  $s_{\text{max}}$  is a set similarity threshold.

The learned prototypes are latent embedding vectors that are not directly interpretable. To address this issue, the authors used a Monte Carlo Tree Search (MCTS) algorithm (Coulom 2006) to identify the subgraph within the observed graphs of the prototype's class that has an embedding closest to the given prototype. Specifically, the author use the MCTS to select an optimal sequence of pruning actions. Let  $\mathcal{G}_{0,\ i,\dots,j}$  represent the subgraph resulting from the sequence of pruning actions  $\operatorname{act}_i,\dots,\operatorname{act}_j$ , where  $\mathcal{G}_0$ , the root of the search tree, is an observed graph. The MCTS algorithm proceeds in 4 core steps. First, an existing node in the search tree,  $\mathcal{G}_{0,\ i,\dots,j}$  is selected. Second, the chosen node is expanded by adding a child node  $\mathcal{G}_{0,\ i,\dots,j,k}$  resulting from action  $\operatorname{act}_k$ . To limit the search space, the authors restrict  $\operatorname{act}_k$  to be the removal of a peripheral node with minimum degree. Third, the reward for the sequence of actions  $i,\dots,j,k$  is estimated via a Monte

Carlo simulation where additional random actions are taken until a subgraph of the desired size is produced. The sampled embeddings are then compared to the prototype vector and the similarities are averaged. Finally, the child node's estimated score is then backpropagated to update the estimated scores of all its parent nodes. After a fixed number of iterations through the observed graphs from the prototype's class, the decision path with the highest resulting score is selected to produce the ultimate prototype projection. In addition to finding the closest subgraph in the observed class, the authors also suggest a method for finding similar subgraphs within each input. This is done by training a neural network to predict an edge mask conditional on a prototype vector:

$$\hat{A}_k = \operatorname{nn}(G, p_k; \ \Omega_{\operatorname{nn}}). \tag{22}$$

The resulting induced subgraph's embedding,  $\hat{p}_{G_k}$ , are then optimized to be as close as possible to the given prototype:

$$\mathcal{L}(\Omega_{\rm nn}, \ \Omega_f, \ p_k \mid G) = -sim(\hat{p}_{G_k}, p_k). \tag{23}$$

The authors denote the version of the model that utilizes this *conditional subgraph sampling module* as **ProtGNN+**.

ProtGNN achieves similar or better graph classification accuracy than state of the art models across 5 standard benchmark datasets (see Figure 6).

Datasets		GCN			GIN			GAT	
2 4	Original	ProtGNN	ProtGNN+	Original	ProtGNN	ProtGNN+	Original	ProtGNN	ProtGNN+
MUTAG	73.3±5.8	76.7±6.4	$73.3 \pm 2.9$	93.3±2.9	90.7±3.2	91.7±2.9	$75.0\pm5.0$	$78.3 \pm 4.2$	81.7±2.9
BBBP	$84.6 \pm 3.4$	$89.4 \pm 4.1$	$88.0 \pm 4.6$	$86.2 \pm 1.1$	$86.5 \pm 1.6$	$85.9 \pm 4.0$	$83.0 \pm 2.6$	$85.9 \!\pm\! 2.5$	$85.5 \pm 0.8$
Graph-SST2	$89.7 \pm 0.5$	$89.9 \pm 2.4$	$89.0 \pm 3.0$	$92.2 \pm 0.3$	$92.0 \pm 0.2$	$92.3 \pm 0.4$	$88.1 \pm 0.8$	$89.1 {\pm} 1.2$	$88.7 \pm 0.9$
Graph-Twitter	$67.5 \pm 1.9$	$68.9 \pm 5.9$	$66.1 \pm 6.5$	$66.2 \pm 1.3$	$75.2 \pm 2.8$	<b>76.5</b> $\pm$ <b>3.4</b>	$69.6 \pm 6.5$	$64.8 \pm 4.0$	$66.4 \pm 3.3$
BA-Shape	$91.9 \pm 1.7$	$\textbf{95.7} \!\pm\! 1.4$	$94.3 \pm 3.7$	$92.9 {\pm} 0.5$	$95.2 \pm 1.3$	$\textbf{95.5} \pm \textbf{2.4}$	$92.9 {\pm} 1.2$	$93.4 {\pm} 3.4$	$93.2 \pm 2.0$

Figure 6: A copy of the quantitative modeling results report in Table 1 of Zhang et al. (2021).

Figure 6 also indicates that ProtGNN is robust to the choice of embedding model as there seems to be benefits across various GNN architectures. The authors do note that the computational complexity of ProtGNN can be significantly higher than a standard GNN model, taking roughly 5 times longer to train. Most of this complexity is attributable to the use of the MCTS. On the other hand, ProtGNN allows for easier and clearer inference. Figure 7 demonstrates that the output of

ProtGNN is similar to a simple regression model. The prototypes can be thought of as features, the similar subgraphs and similarity scores as the observed feature values, and the class connections as the marginal effects on the prediction.

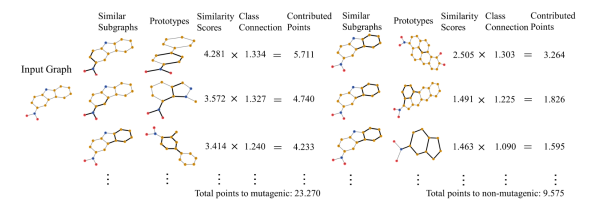


Figure 7: An example ProtGNN+ output for the MUTAG dataset taken from Figure 3 in Zhang et al. (2021).

Since prototypes from every class are used for the classification, ProtGNN can also be used for counterfactual type analysis. For any given predictions, researcher can determine why a certain classification was made and why the other classes were not selected. Figure 7 display both a strong similarity to the mutagenic prototypes as well as a weaker relation to the non-mutagenic prototypes.

# 3 Synthesis of Core Papers

GNNInterpreter and D4Explainer are both post-hoc generative explanation methods. Each defines a model-level explanation for a given class c as an estimated graph generating distribution:

$$G_{\rm Gen} \sim f(G \mid \Theta) \text{ subject to argmin } \mathbb{E}_{G_{\rm Gen}} \text{ CrsEnt [explainee}(G_{\rm Gen}), \ \tilde{\rho_c}] \,. \tag{24}$$

In other words, an explanation is the distribution that maximizes the probability that a sampled graph will be classified by the *explainee* as the desired class c. The key difference between the methods is the form of  $f(G \mid \Theta)$ . This is akin to the parametric or distributional assumption for classical statistical methods. GNNInterpreter assumes that each graph matrix has a separate distribution with possibly dependent parameters. On the other hand, D4Explainer assumes that the only random variation between graphs is in the adjacency structure. Both models assume

that every entry of the adjacency matrix follows a Bernoulli distribution, but diverge in terms of estimating the probability. The methods also utilize different sampling procedures. GNNInterpreter clearly estimates the distributional parameters and samples traditionally. In contrast, D4Explainer employs a black-box denoising model. D4Explainer makes use of the observed graphs directly, whereas GNNInterpreter only injects latent embeddings. This makes GNNInterpreter much more sensitive to the underlying explainee model.

Both sets of authors highlight the out-of-distribution problem that plagues generative explanations. GNN models are not robust to out-of-distribution or unrealistic graphs, which is to say that explainee models can still return a confident classification for a complete random graph (see Figure 1). D4Explainer partial resolves this issue by restricting generated node and edge features to the observed values. This ensures that the features of any generated graph will not be out-of-distribution; however, this limits inference. GNNInterpreter tackles the problem by optimizing the similarity to the average class embedding; however, embeddings can also suffer from out-of-distribution problems (Steck, Ekanadham, and Kallus 2024). Just as an unrealistic or random graph can yield similar predictions, it can also have similar embeddings due to information loss in the lower-dimensional space (see Table 2). While GNNInterpreter and D4Explainer place differing restrictions on their respective likelihoods, the terms are interchangeable. For example, there is no reason why  $\mathcal{L}_{\text{spars}}$  (6) could not be included in the loss for D4Explainer. Similarly, the MMD statistics from D4Explainer could be used to evaluate graphs generated from GNNInterpreter (see Table 1).

ProtGNN, unlike the first two methods, is a self interpretable GNN model. Rather than attempting to decipher a predictive model, ProtGNN itself serves as the predictive model. Prototype similarity is closely related to GNNInterpreter's  $\mathcal{L}_{\text{embed}}$  (5) as they both measure the similarity between graphs in a latent vector space; however, unlike GNNInterpreter, the embedding function in ProtGNN is trainable. Furthermore, ProtGNN also uses dissimilarity to the non-target classes, but this could be easily added to GNNInterpreter's loss. On the other hand, similar to D4Explainer, the graph projections created from the prototypes are restricted to the observed data. It is also worth distinguishing that the similar subgraphs created by ProtGNN+ are instance-level explanations, while the class connection weights and prototype projections are model-level. D4Explainer

produces counterfactual examples by directly computing possible input changes that could flip the prediction, while ProtGNN demonstrates how the prediction could have changed by display the similarity to structures indicative of other classes. While not the traditional direct counterfactual analysis, ProtGNN provides model-level insights.

Experts in the field seem to agree that a class-conditional graph-generating distribution would help illuminate the black-box nature of GNNs. However, there is no consensus on the ideal form or likelihood of such a distribution. While diffusion has proven to be a strong generative method in other applications, D4Explainer's feature restriction and convoluted sampling algorithm leave much to be desired. The results from GNNInterpreter suggest that conditional denoising might be superior. Specifically, optimizing for prediction accuracy tends to produce better intervals, with GNNInterpreter achieving 1.0 (see Figure 2) compared to D4Explainer's 0.832, on the MUTAG dataset. Explicit prototype-based models are fairly uncommon in the literature, but we believe most researchers would agree that Figure 7 provides very clear inference. On the other hand, completely refactoring the predictive model is not always feasible, so both explainable and interpretable GNN methods remain important research avenues.

#### 4 Technical Details

#### 4.1 Methodology

Since GNNInterpreter only makes qualitative assessments of generated structures, we applied the metrics describe in D4Explainer to quantify the properties of the generating distribution. We reimplemented GNNInterpreter using an identical generation scheme and loss function. We trained the reimplemented model on MUTAG (Debnath et al. 1991), a standard benchmark graph classification dataset. MUTAG consists of nitroaromatic compounds, with the goal being to predict their mutagenicity on Salmonella typhimurium. Each observation is a graphs representing a chemical structure, where vertices denote atoms and edges represent bonds. For our explainee model, we also implemented a 3-layer GCN model. There are slight variations between the architectures and training hyperparameters, but the accuracies are comparable, 86% (ours) and 92% (theirs). We additionally reimplement D4Explainer's counterfactual generator. Although not discussed in the

original paper, this model can also be used to produce model level explanations. We simply randomly sample a graph from the opposite class and take the generated counterfactual graph as our explanation.

#### 4.2 Results

Table 1: Comparison of Original and Reimplement Model Results.

	Predictions	Density	Deg.	Clus.	Spec.
GNNInterpro	${f eter~Original}^1$				
Class 0	1.0 + / - 0.0	NA	NA	NA	NA
Class 1	1.0 +/- 0.0	NA	NA	NA	NA
GNNInterpro	eter Reimplen	nented			
Class 0	0.93 +/- 0.01	0.38 +/- 0.002	1.77	1.53	0.07
Class 1	0.98 +/- 0.01	0.32 +/- 0.003	1.41	1.43	0.04
D4Explainer	Original				
Aggregated	$0.92^{2}$	$0.315^2$	$0.12^{3}$	$0.00^{3}$	$0.02^{3}$
D4Explainer	Reimplement	ed			
Class 0	0.63 +/- 0.23	0.13 +/- 0.03	0.27	0.57	0.05
Class 1	0.66 +/- 0.20	0.13 +/- 0.03	0.08	0.25	0.04

 $<sup>^*</sup>$  +/- 1 standard deviation; 1,000 sample graphs.

We were unable to replicate GNNInterpreter's 0 variance predictive accuracy, but the values are comparable given the lack of extensive hyperparameter tuning (see Table 1). The predictions are similarity superior to the original D4Explainer results, 0.93 and 0.98 v. 0.92. Our explainee model has a bias towards the mutagenic class (class 1), which can explain the greater deviation

<sup>\*</sup> The code that produced the reimplemented values can be found here.

<sup>&</sup>lt;sup>1</sup> From Table 2 in Wang and Shen (2024).

 $<sup>^{2}</sup>$  From Table 3 in Chen et al. (2023).

<sup>&</sup>lt;sup>3</sup> From Table 2 in Chen et al. (2023).

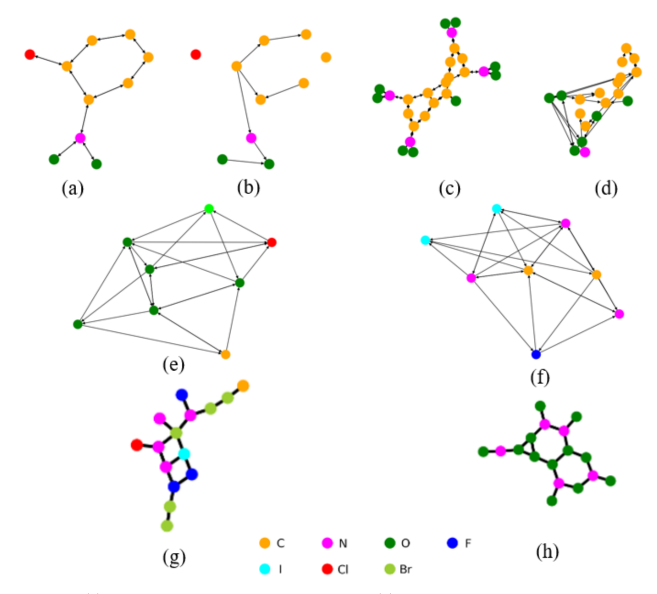


Figure 8: (a) Observed graph from class 0, Pr = 0.92; (b) Generated counterfactual explanation of (a) from D4Explainer reimplemented, Pr = 0.67; (c) Observed graph from class 1, Pr = 0.96; (d) Generated counterfactual explanation of (b) from D4Explainer reimplemented, Pr = 0.96; (e) Generated explanation for class 0 from GNNInterpreter reimplemented, Pr = 0.94; (f) Generated explanation for class 1 from GNNInterpreter reimplemented, Pr = 1.0; (g) Generated explanation for class 0 from GNNInterpreter (X. Wang and Shen (2024), Figure 1, pg.7), Pr = 1.0; (h) Generated explanation for class 1 from GNNInterpreter (X. Wang and Shen (2024), Figure 1, pg.7), Pr = 1.0. The code that produced the reimplemented figures can be found here.

between the class 0 models. Another limiting factor is a lower set maximum number of nodes. We set it to 8 to make the results more comparable to the model-level explanation reported by D4Explainer. Surprisingly, D4Explainer produces sparser graphs than GNNInterpreter in terms of average density, 0.38 and 0.32 v. 0.315. This is despite the fact that D4Explainer has no explicit penalty in the loss function. Based on the MMD statistics, D4Explainer produces graphs with better in-distribution properties (see Table 1). This might be due to the explicit regularization in GNNInterpreter, as penalizing the edge probabilities could lead to a different degree distribution. Additionally, the clustering coefficient might be affected, since fewer edges result in fewer triangle clusters. The spectrum distributions are similar for both models, as both enforce similarity in the explainee embedding space, which aligns with the nature of the spectrum distribution. However, one could argue that having a similar degree distribution and clustering coefficient may not always be desirable. For larger or more densely connected graphs, lower MMD statistics might result in explanations that are too complex to be useful.

Similar to the original examples display in GNNInterpreter, the generated non-mutagenic graphs do not have clear or realistic structures (see Figure 8). Graph (f) highlights that nitrogen in key. We believe that this result is consistent with the original findings about the  $NO_2$  groups, but because of the more restrictive node limit, our reproduction focuses on nitrogen since multiple oxygens can be found in non-mutagenic chemicals. The reimplemented D4Explainer has a CF-ACC in line with the original paper, 76.32% at 0.41 log MR (see Figure 4 for the original metrics). The reimplemented D4Explainer produces the least dense graphs, but has the least discriminative predictions. The latter makes sense, since the goal of the counterfactual loss is to flip the prediction, not necessarily maximize confidence. One the other hand, this model demonstrates that it is possible to generate sparse graphs with similar degree and clustering distributions. Visually the counterfactual explanation make sense. Graph (b) eliminated the red or chlorine atom which is typically associated with non-mutagenic compounds (Yuan et al. 2020). Graph (d) corrupted 3 of the 4  $NO_2$  groups, which is in line with GNNInterpreter's findings that multiple  $NO_2$  groups is a strong indicator of mutagenicity.

These results point to two main conclusions. First, the generator model should make use of graph level data. GNNInterpreter uses information about the graph structure through the explainee, but

never on the raw scale. This lack of graph level information can result in unrealistic distributional properties. One way of including this information would be to use the observed graphs to help predict the distributional parameters. Alternatively, the generator could be replaced with a diffusion model without any modification to the loss. Second, the generator model should use class or prediction information. D4Explainer's model-level sampling algorithm is less efficient and less accuracate than GNNInterpreter's direct optimization. The modified sampling algorithm demonstrates that single-step denoising can still maintain low MMD statistics. Adding  $\mathcal{L}_{\text{pred}}$  (4) and/or  $\mathcal{L}_{\text{embed}}$  (5) could further improve this type of sampling. Alternatively, a class label embedding, similar to the time embedding, could be included to establish a conditional diffusion model, which has worked extremely well in other applications. Additionally, it would be easy to explore diffusing the node and edge features as these types of models already exist (Vignac et al. 2023).

#### 4.3 Proofs

**Proof of Equation 1:** WLOG assume we want to sample for a categorical distribution  $cat(\theta_{cat})$  with a set of W categories each with a probability

$$\pi_{\omega} = \frac{\theta_{\mathrm{cat}}[\omega]}{\sum_{i \in W} \theta_{\mathrm{cat}}[i]}.$$

Define random variable

$$D = \underset{i \in W}{\operatorname{argmax}} \ \log \theta_{\operatorname{cat}} + \operatorname{Gumbel}(0,1) = \underset{i \in W}{\operatorname{argmax}} \ \log \theta_{\operatorname{cat}}[i] + \operatorname{Gumbel}(0,1)[i],$$

here  $\operatorname{Gumbel}(0,1)[i]$  denotes the  $i^{th}$  i.i.d. sample from the associated Gumbel. In order for D to be a true categorical random variable,  $\Pr[D=\omega]$  needs to be  $\pi_{\omega}$ .  $D=\omega$  if and only if  $\log \theta_{\mathrm{cat}}[\omega] + \operatorname{Gumbel}(0,1)[\omega] > \log \theta_{\mathrm{cat}}[i] + \operatorname{Gumbel}(0,1)[i] \ \forall i \in W \setminus \omega$ . Now, let  $M_i$  denote a

random variable that follows a Gumbel (log  $\theta_{\rm cat}[i],1)$  distribution. Then,

$$\begin{split} Pr[D = \omega] &= \mathbb{E}_{M_{\omega}} \prod_{i \in W \backslash \omega} Pr(M_i < m_{\omega}) \text{ i.i.d location shifted Gumbel distributions.} \\ &= \mathbb{E}_{M_{\omega}} \prod_{i \in W \backslash \omega} \exp\left(-e^{\log \theta_{\text{cat}}[i] - m_{\omega}}\right) \text{ Gumbel CDF.} \\ &= \mathbb{E}_{M_{\omega}} \exp\left(-\sum_{i \in W \backslash \omega} e^{\log \theta_{\text{cat}}[i] - m_{\omega}}\right) \\ &= \int_{-\infty}^{\infty} \exp\left(\log \theta_{\text{cat}}[\omega] - m_{\omega}\right) \exp\left(-e^{\log \theta_{\text{cat}}[\omega] - m_{\omega}}\right) \cdot \exp\left(-\sum_{i \in W \backslash \omega} e^{\log \theta_{\text{cat}}[i] - m_{\omega}}\right) \, dm \\ &= \int_{-\infty}^{\infty} \exp\left(\log \theta_{\text{cat}}[\omega] - m_{\omega}\right) \exp\left(-\sum_{i \in W} e^{\log \theta_{\text{cat}}[i] - m_{\omega}}\right) \, dm \\ &= \int_{-\infty}^{\infty} \theta_{\text{cat}}[\omega] \exp\left(-m_{\omega}\right) \exp\left(-e^{-m_{\omega}} \sum_{i \in W} \theta_{\text{cat}}[i]\right) \, dm \\ &= \pi_{\omega} \sum_{i \in W} \theta_{\text{cat}}[i] \int_{-\infty}^{\infty} \exp\left(-m_{\omega}\right) \exp\left(-e^{-m_{\omega}} \sum_{i \in W} \theta_{\text{cat}}[i]\right) \, dm \text{ From the above definition of } \pi_{\omega} \\ &= \pi_{\omega} \sum_{i \in W} \theta_{\text{cat}}[i] \frac{\exp\left(-e^{-m_{\omega}} \sum_{i \in W} \theta_{\text{cat}}[i]\right)}{\sum_{i \in W} \theta_{\text{cat}}[i]} \bigg|_{-\infty}^{\infty} \\ &= \pi_{\omega} \sum_{i \in W} \theta_{\text{cat}}[i] \frac{1}{\sum_{i \in W} \theta_{\text{cat}}[i]} = \pi_{\omega}. \end{split}$$

Reference: Huijben et al. (2022)

#### Proof of the limiting property of 10:

$$\begin{split} \lim_{t \to \infty} q(A_t[i,\ j] \mid A_{t-1}[i,\ j]) &= \lim_{t \to \infty} \operatorname{Cat} \left( A_0[i,\ j] \prod_{i=1}^t Q_i \right) \\ &= \lim_{t \to \infty} \operatorname{Cat} \left( A_0[i,\ j] \prod_{i=1}^t \begin{bmatrix} 1 & -1 \\ 1 & 1 \end{bmatrix} \begin{bmatrix} 1 & 0 \\ 0 & 1 - 2\beta_i \end{bmatrix} \begin{bmatrix} 0.5 & 0.5 \\ -0.5 & 0.5 \end{bmatrix} \right) \end{split}$$

Eigen decomposition.

$$\begin{split} &= \lim_{t \to \infty} \operatorname{Cat} \left( A_0[i, \ j] \begin{bmatrix} 1 & -1 \\ 1 & 1 \end{bmatrix} \begin{bmatrix} 1 & 0 \\ 0 & \prod_{i=1}^t 1 - 2\beta_i \end{bmatrix} \begin{bmatrix} 0.5 & 0.5 \\ -0.5 & 0.5 \end{bmatrix} \right) \\ &= \operatorname{Cat} \left( A_0[i, \ j] \begin{bmatrix} 1 & -1 \\ 1 & 1 \end{bmatrix} \begin{bmatrix} 1 & 0 \\ 0 & 0 \end{bmatrix} \begin{bmatrix} 0.5 & 0.5 \\ -0.5 & 0.5 \end{bmatrix} \right) \end{split}$$

Set  $\beta < 0.5$ .

$$\begin{split} &= \operatorname{Cat}\left(A_0[i,\ j] \begin{bmatrix} 0.5 & 0.5 \\ 0.5 & 0.5 \end{bmatrix}\right) \\ &= \operatorname{Cat}([0.5,0.5]) = \operatorname{Bernoulli}(0.5) \end{split}$$

#### 5 Future Directions

#### 5.1 Graph Edit Distance

GNNInterpreter generates graphs with predictions very close to the target class, but lack good in-distributions properties. D4Explainer, on the other hand, generates explanations with more consistent features distributions, but the predicted probabilities display higher variance. Furthermore, while the model parameters of GNNInterpreter's generator can be interpreted, D4Explainer introduces yet another black-box model. We propose using a differentiable approximation to the graph edit distance (GED) as a way of generically enforcing structural consistency. GED measures the similarity between two graphs by counting the minimum number of edits required to make the graphs isomorphic or subgraph isomorphic. GED is similar to the modification ratio for counterfactual explanations (see Equation 17). Edit distance has the key advantage of being human interpretable. For example, the percentage of atoms and bonds that need to be changed

for two molecules to match is more relatable than the cosine distance between their embeddings. Finding the exact optimal GED is a known NP-complete problem, and most solutions are not differentiable (Bougleux et al. 2015). Fortunately, approximating GED as a quadratic assignment problem (QAP) solves both the time complexity and derivative issue (R. Wang et al. 2024).

Let  $G_1$  and  $G_2$  be graphs we wish to compute the GED between. QAP approximates this be solving the following optimization problem:

$$\mathcal{L}_{\text{GED}}(G_1, G_2) = \max_{M} \text{ vec}(M) \ K(G_1, G_2) \ \text{vec}(M)^T. \tag{25} \label{eq:Loss}$$

Here M is a  $(n_1 \times n_2)$  binary node matching matrix and K is a  $(n_1 n_2 \times n_1 n_2)$  quadratic affinity matrix that encode both node and edge similarities. The efficiency of QAP comes from the fact that when any two nodes are matched together, the above objective will account for the possible edges that could connect the nodes. In order for the solution to be interpreted as an edit distance, each entry of K should range from [-1, 0], where -1 signals the need for a complete edit and 0 means no edits are required (R. Wang et al. 2024). We suggest the following similarity metric for every pair of nodes and edges:

$$\begin{split} &-\lambda_1 \left[1-\phi(\text{Continuous Features})\right] \ + \\ &-\lambda_2 \left[1-\phi(\text{Discrete Features})\right], \end{split} \tag{26}$$

where  $\phi(.)$  is the normalized cosine distance.  $1 - \phi(.)$  will return 2 is the vectors are in opposite directions, 1 if the vectors are orthogonal, and 0 if they are pointing in the same direction; however, one-hot encoded vectors will never be in opposite directions, so the different types of feature must be weighted accordingly. Computing affinity this way allows for K to be differentiable w.r.t. generated graph features. Furthermore we suggest using the reweighted random walks algorithm (Cho, Lee, and Lee 2010) to solve the objective (25), which will generate an M that is differentiable w.r.t. to K.

We conducted a simulation study to compare graph level and latent similarity, or  $\mathcal{L}_{GED}$  (25) and  $\mathcal{L}_{embed}$  (5). We generated 4 clusters of graph data. Every graph had one continuous edge

feature sampled from a standard normal, one discrete node and edge feature sampled from a Bernoulli, and an adjacency matrix, also Bernoulli. The only variation between the clusters was the maximum number of nodes as well as the mean of the continuous node feature distribution. Cluster 0 had 75-100 nodes, cluster 1 had 50-75 nodes, and cluster 2 as well as 3 had 10-35 nodes. This was designed to mimic a classic binary graph classification dataset where cluster 0 and 1 are the observed classes and clusters 2 and 3 are generated subgraph explanations. The embedding model was a 3 layer GNN with a neural network type convolution that utilized every feature. During each of the 50 trials, the GNN was trained using 50 samples (each) from only clusters 0 and 1. We then sampled 50 test graphs from each of the 4 cluster and computed 95% t intervals for GED and embedding cosine distance as percentages of their respective upper bounds. Since  $(\mu_0 = -1, \mu_1 = 1, \mu_2 = 0, \mu_3 = -5)$  we would expect an unbiased distance metric to maintain the following relationship:

$$dist(1, 2) \approx dist(0, 2) < dist(0, 1) < dist(0, 3) < dist(1, 3), \tag{27}$$

based on the distance between cluster means.

Table 2: Simulation Results

	Cluster 0	Cluster 1
Embedding	SS	
Within	(0.0008,  0.0015)	(0.0005,0.0009)
Between	(0.5662, 0.6264)	(0.5689,0.6289)
Cluster 2	(0.2894, 0.3589)	(0.1282, 0.209)
Cluster 3	(0.018,  0.0253)	(0.6521,0.7097)
GED		
Within	(0.4147, 0.419)	(0.462, 0.4674)
Between	(0.6589, 0.6639)	(0.6586, 0.6646)
Cluster 2	(0.7843, 0.7947)	(0.7571, 0.7715)
Cluster 3	(0.6164, 0.6286)	(0.8256,  0.8352)

Expected: 
$$\operatorname{dist}(1, 2) \approx \operatorname{dist}(0, 2) < \operatorname{dist}(0, 1) <$$

$$\operatorname{dist}(0, 3) < \operatorname{dist}(1, 3)$$
GED:  $(0.75, 0.77) \approx (0.78, 0.79) > (0.65, 0.66) >$ 

$$(0.61, 0.62) < (0.82, 0.83)$$
Embedding:  $(0.12, 0.20) < (0.28, 0.35) < (0.56, 0.62) >$ 

$$(0.018, 0.0253) < (0.65, 0.7)$$

The most worrisome bias arises between the distances from cluster 0 to 1 and from cluster 0 to 3 (see Table 2). Despite cluster 0's mean being twice as close to cluster 1, the embedding distance indicates that cluster 0 is (53, 60) percentage points farther away from cluster 1. Although GED also displayed this bias, it was considerably less pronounced, registering at (1, 5) percentage points. This implies that a generator initialized with a mean of -5 would produce samples with embeddings confidently assigning them to cluster 0, despite having a feature distribution far from the truth. Relying solely on embedding distance for training also increases the likelihood of the generator becoming trapped in a misleading local optimum due to the substantial bias. Although GED exhibits a statistically significant bias, its magnitude is considerably smaller, making it less likely to disrupt training. Another troubling bias observed is that the embedding distance between clusters 1 and 2 is approximately 10 percentage points less than the embedding distance between clusters 0 and 2, even though the mean of cluster 2 is equidistant from the means of clusters 1 and 0. This could stem from cluster 1 having fewer nodes making samples more similar in terms of size to cluster 2. However, since researcher typically cap the size of examples for interpretability, an objective with a degree bias could be problematic. On the contrary, GED introduces a bias absent in the embedding space. The edit distance between clusters 0 and 2 is actually greater than

 $<sup>^{1}</sup>$  Node Means:  $\mu_{0}=-1,\,\mu_{1}=1,\,\mu_{out1}=0,\,\mu_{out2}=$  -5.

 $<sup>^2</sup>$  95% t intervals on % difference.

<sup>\*</sup> The code for these simulation results can be found here.

the distance between clusters 0 and 1. While biased concerning mean locations, this direction is aligned with the maximum node size. Despite this, the differing directions of the biases of GED and embedding distance imply that they capture unique structural nuances, potentially complementing each other.

#### 5.2 Graph Protypes

ProtGNN's prototype projections are computationally expensive and cannot produce out-of-sample insights. These two issue can be resolve by using graph protypes as opposed to vectors. Redefine the prototype vectors,  $p_k$ , to be:

$$p_k = f(\tilde{G}_k; \Omega_f), \text{ where } \tilde{G}_k = (\tilde{X}_k, \tilde{A}_k, \tilde{E}_k)$$
 (29)

Redefining the prototypes in this manner maintains compatability with ProtGNN's loss terms as well as enabling the use of GED. The most complex loss function could be:

$$\mathcal{L}(\Omega_f, \Omega_{\psi}, \Omega_{\text{nn}}, \tilde{G}_1 \dots \tilde{G}_{mC} \mid G) = \mathcal{L}_{\text{prot}} + \mathcal{L}_{\text{clust}} + \mathcal{L}_{\text{sep}} + \mathcal{L}_{\text{div}} + \sum_{i=1}^{mC} \mathcal{L}_{\text{GED}}(G_i, \tilde{G}_i) + \sum_{i=1}^{mC} \mathcal{L}_{\text{GED}}(G_i, \tilde{G}_i),$$
(30)

where  $G_1 \dots G_{\mathrm{mC}}$  are the subgraphs induced by the predicted masks. Figure 9 depicts the general flow of such a network. Using graph prototypes also eliminates the computational bottleneck posed by the MCTS projection step. Furthermore, since  $X_{p_k}, A_{p_k}, E_{p_k}$  are free trainable parameters, they could contain structure not observed in the original dataset. We expect sample size to be a limiting factor. Although ProtGNN uses fairly high-dimensional vectors for its prototypes, graph prototypes might still introduce exponentially more parameters. The size and sparsity of the graph prototypes could be enforced, but this would introduce additional hyperparameters. If the dataset is extremely large, then the additional parameters should be manageable.

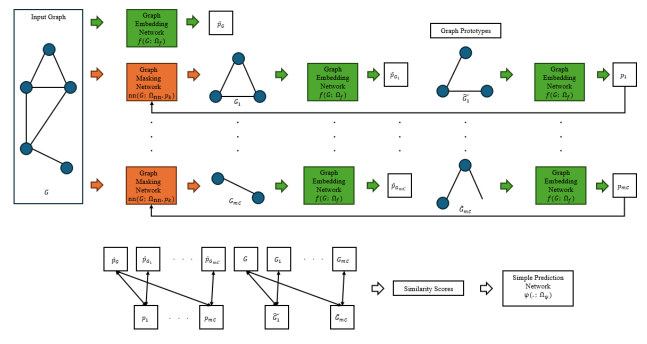


Figure 9: Potential GNN based architecture using graph protoypes.

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