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Abstract  GNNs are powerful models based on node representation learning that perform particularly well in many machine learning problems related to graphs. The major obstacle to the deployment of GNNs is mostly a problem of societal acceptability and trustworthiness, properties which require making explicit the internal functioning of such models. Here, we propose to mine activation rules in the hidden layers to understand how the GNNs perceive the world. The problem is not to discover activation rules that are individually highly discriminating for an output of the model. Instead, the challenge is to provide a small set of rules that cover all input graphs. To this end, we introduce the subjective activation pattern domain. We define an effective and principled algorithm to enumerate activations rules in each hidden layer. The proposed approach for quantifying the interest of these rules is rooted in information theory and is able to account for background knowledge on the input graph data. The activation rules can then be redescribed thanks to pattern languages involving interpretable features. We show that the activation rules provide insights on the characteristics used by the GNN to classify the graphs. Especially, this allows to identify the hidden features built by the GNN through its different layers. Also, these rules can subsequently be used for explaining GNN decisions. Experiments on both synthetic and real-life datasets show highly competitive performance, with up to 200% improvement in fidelity on explaining graph classification over the SOTA methods.

1 Introduction

Graphs are a powerful and widespread data structure used to represent relational data. One of their specificity is that their underlying structure is not in a Euclidean space and has not a grid-like structure (Bronstein et al., 2017), characteristics facilitating the direct use of generic machine learning techniques. Indeed, each node of a graph is characterized by its features, its neighboring nodes, and recursively their properties. Such intrinsically discrete information cannot be easily used by

1 Univ Lyon, INSA Lyon, CNRS, UCBL, LIRIS, UMR5205, F-69621 Villeurbanne, France
2 Laboratoire de Recherche de l'EPITA (LRE), Le Kremlin-Bicêtre, 94276, France
standard machine learning methods to either predict a label associated with the
graph or a label associated with each node of the graph. To overcome this diffi-
culty, Graph Neural Networks (GNNs) learn embedding vectors to represent each
node \( v \) in a metric space and ease comparison between nodes. GNN methods (Def-
ferrard et al., 2016; Wu et al., 2021) employ a message propagation strategy that
recursively aggregates information from nodes to neighboring nodes. This method
produces vectors that represent the ego-graphs centered at each node, in such a
way that the classification task based on these vectors is optimized. These ego-
graphs are induced by nodes that are less than a certain distance from the central
node. These distances are equal to the recursion index and correspond to the layer
indices in the GNN.

Although GNNs have achieved outstanding performance in many tasks, a ma-
jor drawback is their lack of interpretability. The last five years have witnessed a
huge growth in the definition of techniques for explaining deep neural networks
(Burkart and Huber, 2021; Molnar, 2020), particularly for image and text data.
However, the explainability of GNNs has been much less explored. Two types of ap-
proaches have recently been proposed and have gained certain visibility. Methods
based on perturbation (Luo et al., 2020; Ying et al., 2019) aim to learn a mask seen
as an explanation of the model decision for a graph instance. They obtain the best
performance for instance explanation. It appears that such masks can lead to unre-
liable explanations, and most importantly, can lead to misleading interpretations
for the end-user. One can be tempted to interpret all the nodes or features of the
mask as responsible for the prediction leading to wrong assumptions. An example
of misleading interpretations is when a node feature is perceived as important for
the GNN prediction, whereas there is no difference between its distribution within
and outside the mask. XGNN (Yuan et al., 2020a) aims at providing model-level
explanations by generating a graph pattern that maximizes a GNN output label.
Yet, this method assumes that there is a single pattern for each target which is not
the case in practice when dealing with complex phenomena. Moreover, these two
types of methods query the GNN with perturbed input graphs to evaluate their
impact on the GNN decision and build their masks from the model output. They
do not study the internal mechanisms of the GNNs, especially the different em-
bedding spaces produced by the graph convolutions, while we are convinced that
the study of GNN activation vectors may provide new insights on the information
used by GNN to achieve the classification of graphs.

In this paper, we consider GNNs for graph binary classification. We introduce
a new method, called INSIDE-GNN, that aims at discovering activation rules in each
hidden layer of the GNN. An activation rule captures a specific configuration in the
embedding space of a given layer that is considered important in the GNN decision,
i.e., discriminant for an output label. The problem is therefore not only to discover
highly discriminant activation rules but also to provide a pattern set that covers
all GNN decisions on the input graphs. To this end, we define a measure, rooted in
the FORSIED framework (De Bie, 2011) to quantify the information provided by a
rule relative to that supplied by the rules already extracted. The activation pattern
set can then support instance-level explanations as well as providing insights about
the hidden features captured and exploited by the GNN.

Fig. 1 illustrates the main steps of the proposed method. From a trained GNN
model and a set of graphs (ideally following the same distribution as the train-
ing set), (1) a binary activation matrix is derived to encode the activation by the
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Fig. 1 Overview INSIDE-GNN: For each layer, (1) a binary matrix encodes the activation by nodes of embedding vector components. (2) A background model synthesizes the knowledge we have of these data: at the beginning, the probabilities of the components to be activated are independent to the nodes of the graphs. (3) The most informative activation rule (with respect to the background knowledge) is extracted by INSIDE-GNN. (4) This rule is integrated into the background knowledge which gradually makes the marginal distributions of the margins of the background model less and less independent. It is then added to the pattern set (5). Steps (2-5) are repeated until no rule brings significant information about the data in the table. Then, the activation rules are used (6) to support instance level explanations or (7) to provide insights on the model.

Graph nodes of the vector components of the GNN. The decision of the GNN is also associated with the nodes. (2) A background model represents the knowledge we have of the matrix data. At the initialization, we have no particular knowledge and we assume that the activations are independent to the nodes of the graphs. (3) INSIDE-GNN discovers the most informative activation rule based on the activation matrix and the background model. (4) The background model is updated to reflect the latest discovered rule that is added to the pattern set (5). Steps (2-5) are repeated until no more informative rules are obtained or early termination conditions are reached. (6) The activation pattern set is then used to provide instance-level explanations. To this end, several mask strategies involving nodes that support activation rules are devised. (7) For each activation rule, we use exploratory analysis techniques (e.g., subgroup discovery on graph propositionalization, subgraph mining) to characterize the nodes supporting the rules and provide interpretable insights on what the GNN really captures.
Our main contributions are as follows. After discussing the most important related work in Section 2 and introducing the novel problem of mining activation rule sets in Section 3, we devise a branch-and-bound algorithm that exploits upper-bound-based pruning properties to discover such rules. We explain how we characterize the activation rules with graph properties in Section 4. We report an empirical evaluation in Section 5 which studies the performance and the potential of the proposed approach for providing instance-level explanations or insights about the model. INSIDE-GNN is compared against SOTA explanation methods and outperforms them by up to 200%. We also study the characterization of activation rules thanks to interpretable pattern languages. We demonstrate that this allows to obtain good summaries of the hidden features captured by the GNN. Based on this, we eventually compare our approach against a model-level explanation method.

2 Related work

GNNs are attracting widespread interest due to their performance in several tasks as node classification, link prediction, and graph classification (Wu et al., 2020). Numerous sophisticated techniques allow to improve the performance of such models as graph convolution (Kipf and Welling, 2017), graph attention (Velickovic et al., 2018), and graph pooling (Wang and Ji, 2020). However, few researchers have addressed the problem of the GNN explainability compared to image and text domains where a plethora of methods have been proposed (Burkart and Huber, 2021; Molnar, 2020). As stated in (Yuan et al., 2020b), existing methods for image classification models explanation cannot be directly applied to not grid-like data; the ones based on the computation of abstract images via back-propagation (Simonyan et al., 2014) would not provide meaningful results on discrete adjacency matrices; those that learn soft masks to capture important image regions (Olah et al., 2017) will destroy the discreteness property when applied to a graph.

Nevertheless, there have been some attempts to propose methods for explaining GNNs in the last three years. Given an input graph, the instance-level methods aim at providing input-dependent explanations by identifying the important input features on which the model builds its prediction. One can identify four different families of methods. (1) The gradient/feature-based methods – widely applied in image and text data – use the gradients or hidden feature map values to compute the importance of the input features (Baldassarre and Azizpour, 2019; Pope et al., 2019). (2) The perturbation-based methods aim at learning a graph mask by investigating the prediction changes when perturbing the input graphs. GNNExplainer (Ying et al., 2019) is the seminal perturbation based method for GNNs. It learns a soft mask by maximizing the mutual information between the original prediction and the predictions of the perturbed graphs. Similarly, PGExplainer (Luo et al., 2020) uses a generative probabilistic model to learn succinct underlying structures from the input graph data as explanations. (3) The surrogate methods explain an input graph by sampling its neighborhood and learning an interpretable model. GrapheLime (Huang et al., 2020) thus extends the LIME algorithm (Ribeiro et al., 2016) to GNN in the context of node classification. It uses a Hilbert-Schmidt Independence Criterion Lasso as a surrogate model. However, it does not take into account the graph structure and cannot be applied to graph classification models.
PGM-Explainer (Vu and Thai, 2020) builds a probabilistic graphical model for explaining node or graph classification models. Yet, it does not allow to take into consideration edges in its explanations. These surrogate models can be misleading because the user tends to generalize beyond its neighbourhood an explanation related to a local model. Furthermore, the identification of relevant neighborhood in graphs remains challenging. Finally, (4) the decomposition-based methods (Pope et al., 2019; Schnake et al., 2020) start by decomposing the prediction score to the neurons in the last hidden layer. Then, they back-propagate these scores layer by layer until reaching the input space. XGNN (Yuan et al., 2020a) proposes to provide a model-Level explanation of GNNs by training a graph generator so that the generated graph patterns maximize the prediction of the model for a given label. However, it relies on a strong assumption: each label is related to only one graph generator which is not realistic when considering complex phenomena. This is further discussed in Section 5 based on some empirical evidence.

GNNExplainer, PGExplainer, and PGM-Explainer are the methods that report the best performance on many datasets. We will compare our contribution against these methods in the experimental study. Nevertheless, these methods have some flaws when used in practice. Discretizing the soft mask (i.e., selecting the most important edges) requires choosing a parameter $k$ which is not trivial to set. Besides, based on such a mask, the explanation may be misleading because the user is tempted to interpret what is retained in the mask as responsible for the decision, and this, even if a node label appears both inside and outside the mask.

Our method aims to mine some activation patterns in the hidden layers of GNNs. There exists in the literature some rule extraction methods for DNNs (Tran and d’Avila Garcez, 2018), but not for GNNs. For example, (Tran and d’Avila Garcez, 2018) mine association rules from Deep Belief Networks. Still, their approach suffers from an explosion of the number of patterns, which makes the results of frequency-based rule mining mostly unusable in practice. Also, with its focus on DBNs, the method is not directly applicable to standard GNNs.

3 INSIDE-GNN method

3.1 Graph Neural Networks and activation matrix

We consider a set of graphs $G$ with labels: $G = (V, E, M)$ with $V$ a set of nodes, $E$ a set of edges in $V \times V$, and $M$ a mapping between the nodes and the labels, $M \subseteq V \times T$, with $T$ the set of labels. The graphs of $G$ are classified in two categories $\{c^0, c^1\}$ by a GNN: GNN: $G \rightarrow \{c^0, c^1\}$. The GNN takes decisions at the level of each graph on the basis of vectors computed at the level of the nodes. These vectors embed nodes into a metric space to ease comparisons. More precisely, we consider Graph Convolutional Networks (GCN) (Kipf and Welling, 2017) with $L$ layers. GCNs compute vectors $h^\ell_v$, $\ell = 1 \ldots L$ of dimension $K$, an hyperparameter of the method. $h^\ell_v$ represents the ego-graph centered at node $v$ with radius $\ell$. This ego-graph is induced by the nodes that are less than a distance $\ell$ (in number of edges) from $v$. Such vectors are recursively computed by the following formula:

$$h^\ell_v = ReLU \left( W^{\ell} \cdot \sum_{w \in \mathcal{N}(v)} \frac{e_{w,v}}{\sqrt{d_v d_w}} h^{\ell-1}_w \right), \text{ with } d_v = \sum_{w \in \mathcal{N}(v)} e_{v,w}. $$
$e_{v,w}$ is the weight of the edge between nodes $v$ and $w$, $\mathcal{N}(v)$ is the set of neighboring nodes of $v$ including $v$, ReLU is the rectified linear activation function, and $W_{\ell}$ are the parameters learned during the model training phase. Finally, $h^0_v$ is the initial vector for node $v$ with the one-hot encoding of its labels from set $T$.

Each vector is of size $K$ and $\ell$ ranges from 0 to $L$ (the maximum number of layers in the GNN), two hyperparameters of the GNN. For a trained GNN, the vectors $h^0_v$ capture the key characteristics of the corresponding ego-graphs on which the classification is made. When one of the vector components is of high value, it plays a role in the decision process. More precisely, activated components of the vectors — those for which $(h^0_v)_k > 0$ — are combined by the neural network in a path leading to the decision. We are therefore going to construct the activation matrix corresponding to the activated vector components.

**Definition 1 (Activation matrix)** The activation matrix $\hat{H}^\ell$ has dimensions $(n \times K)$, with $n = \sum_{G \in \mathcal{G}} |V_i|$.

$$\hat{H}^\ell[v,k] = \begin{cases} 1 \text{ if } (h^0_v)_k > 0 \\ 0 \text{ otherwise} \end{cases}$$

For a given layer $\ell$, the activated components of $h^0_v$ correspond to the part of the ego-graph centered at $v$ and of radius $\ell$ that triggers the decision. Therefore, we propose in what follows to identify the sets of components that are activated together for one of the two decisions made by the GNN.

### 3.2 Activation rules discovery

We propose to adopt a subgroup discovery approach to identify sets of vector components that are mostly activated in the graphs having the same GNN decision.

**Definition 2 (Activation rule and support)** An activation rule $A^\ell \rightarrow c$ is composed of a binary vector $A^\ell$ of size $K$ and $c \in \{c^0, c^1\}$ a decision class of the GNN. A graph $G_i = (V_i, E_i, L_i) \in \mathcal{G}$ activates the rule if there is a node $v$ in $V_i$ such that $\hat{H}^\ell[v,k] = (A^\ell)_k$, for $(A^\ell)_k = 1$ and $k \in [1,K]$. It is denoted $\text{Activate}(A^\ell \rightarrow c, v)$.

The activated graphs with GNN decision $c$ form the support of the rule:

$$\text{Supp}(A^\ell \rightarrow c, \mathcal{G}) = \{G_i \in \mathcal{G} \mid \exists v \in V_i, \text{Activate}(A^\ell \rightarrow c, v) \text{ and } GNN(G_i) = c\}$$

**Example 1** Fig. 2 presents the internal GNN representation ($h^0_v$) of 4 graphs on the third layer, where $K = 6$. Non-null components (grey cells) are considered as activated and encoded by a ‘1’ value in the binary activation matrix $\hat{H}^3$. The pattern $A^3 = (1,0,0,0,0,1)$ is activated by nodes 1 and 3 of $G_1$ and node 2 of $G_2$. Thus, $\text{Activate}(A^3 \rightarrow 1, v_1) = True$, $\text{Activate}(A^3 \rightarrow 1, v_3) = True$ for $G_1$ and $\text{Activate}(A^3 \rightarrow 1, v_2) = True$ for $G_2$. $\text{Supp}(A^3 \rightarrow 1, \mathcal{G}) = \{G_1, G_2\}$.

Hence, activated rules are more interesting if their supports are largely homogeneous in term of GNN decisions, i.e. the graphs of the support are mainly classified either in class $c^0$ or in class $c^1$. We propose to measure the interestingness of these rules in a subjective manner. It makes possible to take into account a priori knowledge on activation components, but also to perform an iterative extraction of the rules and thus limiting the redundancy between them. These notions are explained below.
### 3.2.1 Measuring the interest of an activation rule

The question now is how to evaluate the interest of the activation rules so as to obtain a set of non-redundant rules. One way to achieve this is to model the knowledge extracted from the activation matrix into a background model and to evaluate the interest of a rule by the knowledge it brings in relation to it. This is what the FORSIED framework (De Bie, 2011) does. It proposes an operational way to define the background model and to evaluate the subjective interest of a pattern by using information theory to quantify both its informativeness and complexity.

We consider the discrete random variable $H[v,k]$ associated to the activation matrix $\hat{H}[v,k]$ and we model the background knowledge by the probability $P(H[v,k] = 1)$. Intuitively, the information content (IC) of an activation rule should increase when its components are unusually activated for the nodes in the graphs of its support (it is unlikely that these components are activated when considering a random node, while this probability increases when considering graphs supporting the pattern).

---

1. Subjective means relative to the background knowledge model.
2. We use hats to signify the empirical values.
Thus, given the probabilities $P(H^ℓ[v,k] = 1)$ and with the assumption that all $H^ℓ[v,k]$ are independent of each other, we can evaluate the interest of a rule by the product of $P(H^ℓ[v,k] = 1)$ for $v$ activated by the rule and $k$ such that $(A^ℓ)_k = 1$. Equivalently, we use the negative log-probability. The more probable the pattern — and therefore the less interesting — the smaller this value. As there may exist several nodes activated in a single graph, we choose the one maximizing the measure. This is formalized in the definition below.

**Definition 3 (Rule information content)** Given a probabilistic background model $P$, the information provided by a rule $A^ℓ → c$ to characterize a set of graphs $G$ is measured by

$$IC(A^ℓ → c, G) = \sum_{G_i \in \text{Supp}(A^ℓ → c, G)} \max_{v \in V_i} \sum_{k : (A^ℓ)_k = 1} \log(P(H^ℓ[v,k] = 1))$$

**Example 2** Considering the example of Fig. 2 and the rule $A^3 → 1$ with $A^3 = (1, 0, 0, 0, 0, 1)$ and the probabilistic background model $P$ given in Table 1, $IC(A^3 → 1, G) = - \log(0.72) - \log(0.34) - \log(0.99) - \log(0.47) = 3.13.$

A pattern with a large IC is more informative, but it may be more difficult for the user to assimilate it, especially when its description is complex. To avoid this drawback, the IC value is contrasted by its description length which measures the complexity of communicating the pattern to the user. The higher the number of components in $A^ℓ$, the more difficult to communicate it to the user.

**Definition 4 (Description length of a rule)** The description length of a rule is evaluated by

$$DL(A^ℓ → c) = |A^ℓ| \times \alpha + \beta$$

with $|A^ℓ|$ the L1 norm of $A^ℓ$, $\alpha$ the cost for the user to assimilate each component and $\beta$ a fixed cost for the pattern. We set $\beta = 1$ and $\alpha = 0.6$, as the constant parameter $\beta$ does not influence the relative ranking of the patterns, and with a value of 1, it ensures that the DL value is greater than 1. With $\alpha = 0.6$, we express a slight preference toward shorter patterns.

**Example 3** $DL(A^3 → 1) = 2\alpha + \beta = 2.2.$

The subjective interestingness measure is defined as the trade-off between IC and DL.

**Definition 5 (Subjective interestingness of a rule)** The subjective interestingness of a rule on the whole set of graphs $G$ is defined by

$$SI(A^ℓ → c, G) = \frac{IC(A^ℓ → c, G)}{DL(A^ℓ → c)}$$

However, in order to identify rules specific to a GNN decision, we consider the difference of subjective interestingness of the measure evaluated on the two groups of graphs.
**Definition 6 (Differential measure of subjective interest)** If we denote by $G^0$ (resp. $G^1$) the graphs $G_i \in \mathcal{G}$ such that $\text{GNN}(G_i) = c^0$ (resp. $\text{GNN}(G_i) = c^1$), the subjective interest of the rule $A^\ell \rightarrow c$ with respect to the classes is evaluated by

$$SI_{SG}(A^\ell \rightarrow c) = \omega_c SI(A^\ell \rightarrow c, G^c) - \omega_{1-c} SI(A^\ell \rightarrow c, G^{1-c}).$$

The weights $\omega_0$ and $\omega_1$ are used to counterbalance the measure in unbalanced decision problems. The rational is to reduce the SI values of the majority class. We set $\omega_0 = \max(1, \frac{|G^1|}{|G^0|})$ and $\omega_1 = \max(1, \frac{|G^0|}{|G^1|})$.

**Example 4** $SI_{SG}(A^3 \rightarrow 1) = SI(A^3 \rightarrow 1, G^1) - SI(A^3 \rightarrow 1, G^0) = \frac{IC(A^3 \rightarrow 1, G^1)}{DL(A^3 \rightarrow 1)} - \frac{IC(A^3 \rightarrow 1, G^0)}{DL(A^3 \rightarrow 1)} = 3.13 - 0.22 = 2.91$.

### 3.2.2 Computing the background model

The background model is initialized with basic assumptions about the activation matrix and updated as rules are extracted.

**Definition 7 (Initial background model)** Some components can be activated more than others on all the graphs, or some nodes can activate a variable number of components. We assume that this information is known and use it to constrain the initial background distribution $P$:

$$\sum_v P(H^{\ell,v}[v, k] = 1) = \sum_v P(\hat{H}^{\ell,v}[v, k] = 1),$$

$$\sum_k P(H^{\ell,v}[v, k] = 1) = \sum_k P(\hat{H}^{\ell,v}[v, k] = 1).$$

However, these constraints do not completely specify the probability matrix. Among all the probability distributions satisfying these constraints, we choose the one with the maximum entropy. Indeed, any distribution $P$ with an entropy lower than the maximum entropy distribution effectively injects additional knowledge, reducing uncertainty unduly. The explicit mathematical MaxEnt model solution can be found in [De Bie, 2009].

The corresponding initial background model of example of Fig. 2 is given in Table 2.

Once a rule $A^\ell \rightarrow c$ has been extracted, it brings some information about the activation matrix that can be integrated into $P$. The model must integrate the knowledge carried by this rule, that is to say that all the components with value 1 of $A^\ell$ are activated by the vertices activating the rule.

**Definition 8 (Updating the background model)** The model $P$ integrates the rule $A^\ell \rightarrow c$ as follows: $\forall k$ such that $(A^\ell)_k = 1$ and $v$ such that $\hat{H}^{\ell,v}[v, k] = (A^\ell)_k$, $P(H^{\ell,v}[v, k] = 1)$ is set to 1.
### 3.2.3 Iterative extraction of subjective activation subgroups

We propose to compute the subjective activation rules with an enumerate-and-rank approach. It consists to compute the rule $A^\ell \rightarrow c$ with the largest $SI_{SG}$ value and to integrate it in the background distribution $P$ to take into account this newly learnt piece of information.

Algorithm 1 sketches the method. First, it sets the output set equal to the empty set (line 1) and the minSI value to the largest value (line 2). minSI corresponds to the smallest $SI_{SG}$ value of the extracted patterns. A stack of size $K$ is initialized line 3. The first considered rule $A^\ell$ is initialized as a bit-vector of size $K$ containing only 0’s. It corresponds to the rule with no activated components. It has an associate attribute $Pot$ that encodes the components that could still be activated for $A^\ell$, as it leads a yet unconsidered combination of activated components. Rule $A^\ell$ is then staked in Stack (lines 4 to 6). Line 7, it computes the background model $P$ from the activation matrix $H^\ell$ as defined in Definition 7. Then, in a loop (lines 8 to 11), it computes iteratively the rule $A^\ell \rightarrow c$ having the best $SI_{SG}(A^\ell \rightarrow c)$ value. Then, this best rule is used to update the model $P$ (line 11). Indeed, once the rule $A^\ell$ is known, its subjective interest falls down to 0. This consists in setting the corresponding probabilities to 1.

Algorithm 2 presents INSIDE-SI that computes the best rule with as activated components the one’s values of the vector stored in Stack[depth], and even more, depending on the recursive process. It considers a pattern $A$ stored in the stack at depth depth. $A$ has 5 attributes:

<table>
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<th>Nodes</th>
<th>Component 1</th>
<th>Component 2</th>
<th>Component 3</th>
<th>Component 4</th>
<th>Component 5</th>
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<td>0.556</td>
<td>0.507</td>
<td>0.346</td>
<td>0.346</td>
</tr>
<tr>
<td>$G_4$</td>
<td>1</td>
<td>0.527</td>
<td>0.402</td>
<td>0.366</td>
<td>0.250</td>
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<td>2</td>
<td>0.729</td>
<td>0.556</td>
<td>0.507</td>
<td>0.346</td>
<td>0.346</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>0.527</td>
<td>0.402</td>
<td>0.366</td>
<td>0.250</td>
<td>0.250</td>
</tr>
<tr>
<td></td>
<td>4</td>
<td>0.527</td>
<td>0.402</td>
<td>0.366</td>
<td>0.250</td>
<td>0.250</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>0.527</td>
<td>0.402</td>
<td>0.366</td>
<td>0.250</td>
<td>0.250</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>0.374</td>
<td>0.285</td>
<td>0.285</td>
<td>0.177</td>
<td>0.177</td>
</tr>
</tbody>
</table>

Table 1 Initial background model $P(H^\ell[v,k] = 1)$ of example of Fig.2
Algorithm 1: inside-GNN($\hat{H}^t$, $c$, nbPatt)

**Input:** $\hat{H}^t$ the activation matrix (see Definition 1), $c$ the class to be characterized and nbPatt the number of patterns to extract.

**Output:** output, up to nbPatt best activation rules $A^t$ w.r.t. $SI_{SG}$.

1. output $\leftarrow \emptyset$
2. minSI $\leftarrow 0$
3. Stack.maxsize $\leftarrow K$
4. $A^t$ $\leftarrow$ a size $K$ bit-vector initialized at 0
5. $A^t.Pot$ $\leftarrow$ a size $K$ bit-vector full of 1’s
6. $Stack[0]$ $\leftarrow A^t$
7. $P$ $\leftarrow$ Compute_Model($\hat{H}^t$)
8. repeat
9. $A^t, minSI \leftarrow$ INSIDE-SI($Stack$, $P$, $c$, minSI , 0, $\emptyset$)
10. output $\leftarrow$ output $\cup A^t$
11. Update_Model($P, A^t$)
12. until ((|output| $\geq$ nbPatt) or (minSI < 10));

Algorithm 2: INSIDE-SI($Stack$, $P$, $c$, minSI, depth, Best)

**Input:** Stack a stack of recursively enumerated patterns at depth depth, $P$ the background distribution, $c$ the class to be characterized, minSI a dynamic threshold on $SI_{SG}$.

**Output:** Best, the best rule w.r.t. $SI_{SG}$.

1. $A$ $\leftarrow$ Stack[depth]
2. $Best$ $\leftarrow$ $A$
3. if ($\phi(A) = False$) or ($UB_{SI}(A, P, c) < minSI$) then
4. return
5. if ($A.Pot = \emptyset$) then
6. if $Best = \emptyset$ then
7. if $SI_{SG}(A \rightarrow c) > minSI$ then
8. $Best$ $\leftarrow$ $A$
9. else
10. if ($SI_{SG}(A \rightarrow c) > SI_{SG}(Best \rightarrow c)$) then
11. $Best$ $\leftarrow$ $A$
12. minSI $\leftarrow$ $SI_{SG}(Best \rightarrow c)$
13. else
14. $x$ $\leftarrow$ first bit of $A.Pot$ set to 1
15. $A.Pot[x]$ $\leftarrow$ 0
16. $A[x]$ $\leftarrow$ 1
17. $Stack[depth + 1]$ $\leftarrow$ $A$
18. INSIDE-SI($Stack$, $P$, $c$, minSI, depth+1)
19. $A[x]$ $\leftarrow$ 0
20. $Stack[depth + 1]$ $\leftarrow$ $A$
21. INSIDE-SI($Stack$, $P$, $c$, minSI, depth+1)
22. return $Best$, minSI

- $A.Pot$, a vector whose one’s values represent the activated components that can be further added to $A$ during the enumeration process,
- $AG^c$ (resp. $AG^{1-c}$) the set of graphs from $G^c$ (resp. $G^{1-c}$) that support $A$,
- and $ATG^c$ (resp. $ATG^{1-c}$) the set of graphs that are supporting $A$ and all its descendants in the enumeration process (there is a node in these graphs that activates all the components of $A$ and $A.Pot$).
The algorithm computes the closure of \( A \) using the function \( \phi \). It consists in adding activated components to \( A \) (set some components of \( A \) to 1) as long as the set \( A.G^c \) of supporting graphs stays unchanged. Furthermore, if a component has been removed from \( A \) (on line 14) but can be added later to \( A \) (i.e. \( \phi(A) \& \& A.pot \neq A \) with \( \& \) the bitwise and operation), \( A \) is not closed, the function returns False and the recursion stops.

Line 2, a second criterion based on an upper bound \( UB_{SI} \) makes the recursion stop if its value is less than the one of the current best found rule. It relies on the following property.

**Property 1** Let \( A \) and \( B \) be two binary vectors of size \( K \). The components that are activated for \( A \) are also activated for \( B \) (i.e., \( A \& B = A \), with \( \& \) the bitwise and operation). We can upper bound the value \( SI_{SG}(B \to c) \) and have

\[
SI_{SG}(B \to c) \leq UB_{SI}(A, P, c) \text{ with }
\]

\[
UB_{SI}(A, P, c) = w_c \frac{\sum_{g \in A.G^c} \max_{v \in V_g} \sum_{k \text{ s.t. } (A \& A.Pot)_k = 1} \log(P(H^f[v, k] = 1))}{\alpha(|A|) + \beta}
\]

\[
- w_{1-c} \frac{\sum_{g \in A.TG^{1-c}} \max_{v \in V_g} \sum_{k \text{ s.t. } (A)_k = 1} \log(P(H^f[v, k] = 1))}{\alpha(|A \& A.Pot|) + \beta}
\]

with \(|A|\) the L1 norm of \( A \).

**Proof** To upper bound the measure \( SI_{SG}(B \to c) \), we follow the strategy explained in [Cerf et al., 2009]. Let

\[
SI_{SG}(B \to c) = w_c X - w_{1-c} Z
\]

with

\[
X = IC(B, G^c) = \sum_{G \in \text{Supp}(B, G^c)} \max_{v \in V_g} \sum_{k \text{ s.t. } (B)_k = 1} \log(P(H^f[v, k] = 1))
\]

\[
Z = IC(B, G^{1-c}) = \sum_{G \in \text{Supp}(B, G^{1-c})} \max_{v \in V_g} \sum_{k \text{ s.t. } (B)_k = 1} \log(P(H^f[v, k] = 1))
\]

\[
Y_1 = Y_2 = DL(B) = \alpha(|B|) + \beta
\]

Similarly, we denote the upper bound function by

\[
UB_{SG}(A, P, c) = w_c \gamma \frac{1}{\beta} - w_{1-c} \frac{\epsilon}{\eta}
\]

Therefore, the largest value of \( SI_{SG}(B \to c) \) is obtained if:

- \( X \) has the maximal possible value, that is to say \( B = A \& A.Pot \) and all the graphs of \( G^c \) that support \( A \), also support \( A \& A.Pot \). In that case, we have

\[
\gamma = \sum_{g \in A.G^c} \max_{v \in V_g} \sum_{k \text{ s.t. } (A \& A.Pot)_k = 1} \log(P(H^f[v, k] = 1))
\]
\( Y_1 \) has the smallest possible value \( \alpha(|A|) + \beta \) (more elements in \( B \) will decrease the value of the fraction)

\[
\delta = \alpha(|A|) + \beta
\]

\( Z \) has the smallest possible value and is computed over \( A \), and on the graphs from \( \mathcal{G}^c \) that support \( A \) and all its descendants (\( A.TG^{1-c} \))

\[
\epsilon = \sum_{g \in A.TG^{1-c}} \max_{v \in V_g} - \sum_{k \text{ s.t. } (A)_k = 1} \log(P(H^f[v, k] = 1))
\]

\( Y_2 \) has the value \( \alpha(|A \& A.Pot|) + \beta \) (less elements in \( B \) will decrease the value of the function)

\[
\eta = \alpha(|A \& A.Pot|) + \beta
\]

It results in the upper bound definition.

Line 4 of Algorithm \([2]\) Best is updated as well as minSI if there are no more component to enumerate, and if the SI-SG value of the current rule is better than the one already found. Otherwise (lines 9 to 16), the enumeration continues either 1) by adding a component from \( A.Pot \) to \( A \) (lines 9-12) and recursively call the function (line 13), or 2) by not adding the component while still removing form \( A.Pot \) (line 14) and recursively continue the process (line 16).

### 4 Characterization of activation rules with subgroups

Once the activation patterns are found, we aim to describe them in an intelligible and accurate way. We believe that each activation pattern can be linked to hidden features of the graphs, that are captured by the model as being related to the class to be predicted. The objective here is to make these features explicit. For this, we seek to characterize the nodes that support the activation pattern, and more precisely to describe the singular elements of their neighborhoods. Many pattern domains can be used to that end. In the following, we consider two of them: one based on numerical descriptions and the other one based on common subgraphs. In order to characterize the subgraphs centered on the nodes of the activation pattern support (called ego-graphs) in a discriminating way compared to the other subgraphs, we extend the well-known GSPAN algorithm \([3]\) so that it takes into account subgroup discovery quality measure.

#### 4.1 Numerical subgroups

In this approach, we propose to describe each node that supports a given activation pattern by some topological properties. We choose to consider its degree, its betweenness centrality value, its clustering-coefficient measure, and the number of triangles it is involved in, as characteristic features. These properties can be extended to the whole ego-graph by aggregating the values of the neighbors. We

\[3\] These attributes are computed with Networkx Python Library [https://networkx.org/](https://networkx.org/)
consider two aggregation functions: the sum and the mean. Thanks to these properties, we make a propositionalization of the nodes of the graphs and we consider as target value the fact that the node belongs to the support of the activation rule (labeled as a positive example) or not (labeled as a negative example). Therefore, we have a matrix $D$ whose rows denote graph nodes and columns correspond to numerical attributes describing the position of the node in its graph: $D[v] \in \mathbb{R}^p$, with $p$ the number of attributes. $D$ is split in two parts $D^0$ and $D^1$ with $v \in D^c$ iff $\text{Activate}(A' \rightarrow c, v)$.

To identify the specific descriptions of the support nodes, we propose to use a subgroup discovery method in numerical data. It makes it possible to find restrictions on numerical attributes (less or greater than a numerical value) that characterize the presence of a node within the support of the activation rule. A numerical pattern has the form $\times_{i=1}^p [a_i, b_i]$ (i.e. the pattern language) and a graph node supports the pattern if $\forall i = 1 \ldots p, a_i \leq D[v, i] \leq b_i$.

To discover such subgroups, we use the pysubgroup library [Lemmerich and Becker 2018].

4.2 Graph subgroups

Another approach consists to characterize activation rules by subgraphs that are common among positive examples in contrast to the negative ones. To this end, we consider as positive examples the ego-graphs (with a radius equal to the layer) of nodes that support the activation pattern of interest. By taking the radius into account, we are not going beyond what the model can actually capture at this layer. The negative examples are the graphs in $G$ for which none of their vertices support the activation pattern. Hence, $D$ is a set of graph nodes $v$ associated to ego-graphs $E_g = (V_g, E_g, L_g)$. $D$ is split into $D^0$ and $D^1$ with $v \in D^c$ iff $\text{Activate}(A' \rightarrow c, v)$.

A graph pattern has the form $G = (V, E, L)$ (i.e. the pattern language) and a graph node supports the pattern if there exists a graph isomorphism between $E_g = (V_g, E_g, L_g)$ and its ego-graph $E_g = (V_g, E_g, L_g)$.

4.3 Quality measure and algorithms

As for the identification of activation patterns, we could have used subjective interestingness measure to characterize the supporting ego-graphs of the activation patterns. However, we opt for a more usual measure, the Weighted Relative Accuracy [Lavrač et al. 1999]. Given a pattern $P$ of a given language, a dataset $D$ split into $D^0$ and $D^1$ and a $\text{Supp}(P, D)$ measure that gives all the graph nodes supporting the pattern $P$ in the data $D$, the WRAcc measure

$$WRAcc(P, c) = \frac{|\text{Supp}(P, D)|}{|D|} \left( \frac{|\text{Supp}(P, D^c)|}{|\text{Supp}(P, D)|} - \frac{|D^c|}{|D|} \right)$$

gives high values to patterns that are mainly supported by nodes of $D^c$ compared to the whole dataset $D$. Then, we use off-the-shelf algorithms to discover the best subgroups. We compute patterns $P$ such that

$$WRAcc(P, c) \geq \min \_WRAcc \text{ and } |\text{Supp}(P, D)| \geq \min \_sup$$  \hspace{1cm} (1)
or just the subgroup with the highest \(WRAcc\) value.

For the numerical subgroups, we use PySubgroup library [Lemmerich and Becker, 2018]. For graph subgroup discovery, we integrate the \(WRAcc\) measure into the GSPAN algorithm [Yan and Han, 2002]. As \(WRAcc\) measure is not anti-monotone, we use the following upper-bound instead of the \(WRAcc\) for pruning:

\[
UB(P,c) = \frac{|\text{Supp}(P,D)|}{|\mathcal{D}|} \left(1 - \max \left(\frac{\text{min}_\text{sup}}{|\mathcal{D}|}, |\mathcal{D}'|\right)\right)
\]

If \(\text{min}_\text{sup} < |\mathcal{D}'|\), then we have \(UB(P,c) = \frac{|\text{Supp}(P,D)|}{|\mathcal{D}|} \left(1 - \frac{|\mathcal{D}'|}{|\mathcal{D}|}\right)\). Since \(\frac{|\text{Supp}(P,D')|}{|\text{Supp}(P,D)|} \leq 1\), \(WRAcc(P,c) \leq UB(P,c)\). In the other case, we have:

\[
\frac{|\text{Supp}(P,D')|}{|\text{Supp}(P,D)|} - \frac{|\mathcal{D}'|}{|\mathcal{D}|} \leq \frac{|\text{Supp}(P,D')|}{|\text{Supp}(P,D)|} - \frac{|\text{Supp}(P,D)|}{|\mathcal{D}|} \iff \\
\frac{\text{min}_\text{sup}}{|\mathcal{D}|} \leq \frac{|\text{Supp}(P,D)|}{|\mathcal{D}|} - \frac{|\text{Supp}(P,D')|}{|\text{Supp}(P,D)|} \iff \\
\frac{1}{|\mathcal{D}|} (\text{min}_\text{sup} - |\mathcal{D}'|) \leq \frac{1}{|\text{Supp}(P,D)|} (|\text{Supp}(P,D)| - |\text{Supp}(P,D')|)
\]

The last inequality holds since \(\frac{1}{|\mathcal{D}|} \leq \frac{1}{|\text{Supp}(P,D)|}\), \(\text{min}_\text{sup} \leq |\text{Supp}(P,D)|\), and finally \(|\mathcal{D}'| \geq |\text{Supp}(P,D')|\).

Since \(UB\) is not dependent to the \(\text{Supp}(P,D')\), when \(|\mathcal{D}'|\) is much lower than the \(|\mathcal{D}|\), this upper bound is not tight. We can use another upper bound which is dependent to the \(|\mathcal{D}'|\). Let us call this upper bound \(UB2\):

\[
UB2(P,c) = \frac{|\text{Supp}(P,D')|}{|\mathcal{D}|} - \frac{\text{min}_\text{sup}}{|\mathcal{D}|} \times \frac{|\mathcal{D}'|}{|\mathcal{D}|}
\]

Since except \(\text{Supp}(P,D')\) everything is constant, and \(\text{Supp}(P,D')\) is anti-monotone, \(UB2\) is anti-monotone too. To show that \(UB2\) is an upper bound for \(WRAcc\), note that \(\frac{\text{min}_\text{sup}}{|\mathcal{D}|} \times |\mathcal{D}'| \leq \frac{|\text{Supp}(P,D)|}{|\mathcal{D}|} \times |\mathcal{D}'|\) and the first terms of \(WRAcc\) and \(UB2\) are equal. In our algorithm we use \(UB3(P,c) = \min\{UB2(P,c), UB(P,c)\}\) as upper bound for the \(WRAcc\).

5 Experimental study

In this section, we evaluate \textsc{inside-gnn} through several experiments. We first describe synthetic and real-world datasets and the experimental setup. Then we present a quantitative study of the patterns provided by \textsc{inside-gnn}. Next, we show the experimental results on explanations of graph classification against several SOTA methods. Finally, we report results on the characterization of activation rules by human understandable descriptions of what GNN models capture. \textsc{inside-gnn} has been implemented in Python and the experiments have been performed on a machine equipped with 8 Intel(R) Xeon(R) W-2125 CPU @ 4.00GHz cores 126GB main memory, running Debian GNU/Linux. The code and the data are available.

\[4\] https://www.dropbox.com/sh/jsri7jbhmku6c8b/AACKHwcM3GmaPC8iBMlFehCa?dl=0
5.1 Datasets and experimental setup

Experiments are performed on six graph classification datasets whose main characteristics are given in Table 2. BA2 (Ying et al., 2019) is a synthetic dataset generated with Barabasi-Albert graphs and hiding either a 5-cycle (negative class) or a “house” motif (positive class). The other datasets (Aids (Morris et al., 2020), BBBP (Wu et al., 2017), Mutagen (Morris et al., 2020), DD (Dobson and Doig, 2003), Proteins (Borgwardt et al., 2005)) depict real molecules and the class identifies important properties in Chemistry or Drug Discovery (i.e., possible activity against HIV, permeability and mutagenicity).

A 3-convolutional layer GNN (with $K = 20$) is trained on each dataset using 80% of the data (train set). The hyperparameters are chosen using a grid-search on other 10% of the data (validation set). The learned GNN are tested on the remaining 10% of the data (test set). The corresponding accuracy values are reported in Table 2.

### Table 2: Main characteristics of the datasets.

<table>
<thead>
<tr>
<th></th>
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<th></th>
<th></th>
<th></th>
<th></th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td>BA2(syn)</td>
<td>1000</td>
<td>(500, 500)</td>
<td>25</td>
<td>50.92</td>
<td>0.995</td>
<td>0.97</td>
<td>1.0</td>
</tr>
<tr>
<td>Aids</td>
<td>2000</td>
<td>(400, 1600)</td>
<td>15.69</td>
<td>32.39</td>
<td>0.989</td>
<td>0.99</td>
<td>0.975</td>
</tr>
<tr>
<td>BBBP</td>
<td>1640</td>
<td>(389, 1251)</td>
<td>24.08</td>
<td>51.96</td>
<td>0.855</td>
<td>0.787</td>
<td>0.848</td>
</tr>
<tr>
<td>Mutagen</td>
<td>4357</td>
<td>(2401, 1936)</td>
<td>30.32</td>
<td>61.54</td>
<td>0.815</td>
<td>0.786</td>
<td>0.804</td>
</tr>
<tr>
<td>DD</td>
<td>1168</td>
<td>(681, 487)</td>
<td>268</td>
<td>1352</td>
<td>0.932</td>
<td>0.692</td>
<td>0.760</td>
</tr>
<tr>
<td>Proteins</td>
<td>1113</td>
<td>(663, 450)</td>
<td>39</td>
<td>145</td>
<td>0.754</td>
<td>0.768</td>
<td>0.784</td>
</tr>
</tbody>
</table>

5.2 Quantitative study of activation rules

Table 3 reports general indicators about the discovery of activation rules by INSIDE-GNN. The execution time ranges from few minutes for simple task (i.e., synthetic graphs) to two days for more complex ones (i.e., DD). It shows the feasibility of the proposed method. Notice that this process is performed only once for each model.

To assess whether the set of extracted rules represent the GNN well and in its entirety, we used the rules to describe the input graphs (i.e. the graphs (in row) are described by the rules (in columns) and the data matrix contains the number of graph nodes supporting the corresponding rule). We then learned the simple and interpretable model that is the decision tree. Thus, from only the knowledge of the number of nodes of a graph supporting each of the rules, we can see, in Table 3 last column, that the decision tree can mimic the GNN decision output with high accuracy. Obviously, we do not provide an interpretable model yet, since the decision tree is based on the patterns that capture sets of activated components of the GNN. Nevertheless, the results demonstrate that the pattern set returned by INSIDE-GNN captures the inner workings of GNNs well.
Table 3 Execution time, number of discovered patterns by INSIDE-GNN and the ability of the pattern set to mimic GNN: the accuracy of a decision tree with activation rules as features and measured on a test set. The class variable is the GNN output $y_i$. The closer $\text{Acc}(\text{DT}^\mathbf{P}, y_i)$ to 1, the better the mimicry.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>Time (s)</th>
<th># Act. Rules</th>
<th>$\text{Acc}(\text{DT}^\mathbf{P}, y_i)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>BA2(syn)</td>
<td>180</td>
<td>20</td>
<td>0.98</td>
</tr>
<tr>
<td>Aids</td>
<td>5160</td>
<td>60</td>
<td>0.96</td>
</tr>
<tr>
<td>BBBP</td>
<td>6000</td>
<td>60</td>
<td>0.89</td>
</tr>
<tr>
<td>Mutagen</td>
<td>41940</td>
<td>60</td>
<td>0.87</td>
</tr>
<tr>
<td>DD</td>
<td>212400</td>
<td>47</td>
<td>0.86</td>
</tr>
<tr>
<td>Proteins</td>
<td>8220</td>
<td>29</td>
<td>0.87</td>
</tr>
</tbody>
</table>

The general characteristics of the activation rules for each dataset are provided in Figs. 3–6. One can observe – in Fig. 3 – that a rule is usually supported by more than one node within a graph. Rules from the first layer of the GNN tend to involve a higher number of vertices than those in the following layers. It may be due to the fact that the first layer captures some hidden common features about the direct neighborhood of the vertices. The features captured by the GNN become more discriminant with layer indexes, as evidenced by the increasing SI_SG score with layers in Fig. 5. For some datasets (e.g., BA2, AIDS, DD, Proteins), some rules have high discriminative power for the positive class (bottom right corner in Fig. 6) or the negative class (top left corner). Their discriminative power is less effective for Mutagen and BBBP datasets. The most discriminant rules come from the last layer of the GNN. Some rules are not discriminant (i.e., around the diagonal) but remains subjectively interesting. These rules uncover activated
components that capture general properties of the studied graphs. It is important to note that we study here the discriminative power of a rule according to its presence in graphs. These rules can be more discriminant if we take into account the number of occurrences of the rules in the graphs. For instance, a rule that is not discriminant can become highly discriminant if we add a condition on its number of occurrences in graph, as we did when learning the decision trees in Table 3.

5.3 Comparison with competitors for explainability of GNN output

We now assess the ability of activation rules to provide good explanations for the GNN decisions. According to the literature, the best competitors are GNNExplainer (Ying et al., 2019), PGExplainer (Luo et al., 2020) and PGM-Explainer (Vu and Thai, 2020). We consider all of them as baseline methods. Furthermore, we also consider a gradient-based method (Pope et al., 2019), denoted Grad, even if it has been shown that such method is outperformed by the three others. Therefore, we compare INSIDE-GNN against these 4 single-instance-explanation methods in our experiments.

Evaluating the reliability of an explanation is not trivial due to the lack of ground truths. In our case, only BA2 is provided with ground truths by construction. When we have ground truths, we expect a good explanation to match it perfectly, but sometimes the model captures a different explanation that is just as discriminating. Moreover, if fully present, ground truths contain only simple relationships (e.g., BA2) which are not sufficient for a full assessment. Therefore,
to be able to consider synthetic and real-world datasets, we consider a ground truth free metric. We opt for Fidelity [Pope et al., 2019] which is defined as the difference of accuracy (or predicted probability) between the predictions on the original graph and the one obtained when masking part of the graph based on the explanations:

\[
Fid_{acc} = \frac{1}{N} \sum_{i=1}^{N} (1 - \delta_{(y_i \setminus m_i = y_i)}),
\]

where \(y_i\) is the original prediction of graph \(g_i\), \(m_i\) is the mask and \(g_i \setminus m_i\) is the complementary mask, \(\hat{y}_i \setminus m_i\) is the prediction for the complementary mask and \(\delta_{(y_i \setminus m_i = y_i)}\) equals 1 if both predictions are equal.

The fidelity can also be measured by studying the raw probability score given by the model for each class instead of the accuracy:

\[
Fid_{prob} = \frac{1}{N} \sum_{i=1}^{N} (f(g_i)_{y_i} - f(g_i \setminus m_i)_{y_i}),
\]

with \(f(g)_{y_i}\) is the prediction score for class \(y_i\).

Similarly, we can study the prediction change by keeping important features (i.e., the mask) and removing the others as Infidelity measures do:

\[
Infid_{acc} = \frac{1}{N} \sum_{i=1}^{N} (1 - \delta_{(y_i = y_i)}),
\]

\[
Infid_{prob} = \frac{1}{N} \sum_{i=1}^{N} (f(g_i)_{y_i} - f(m_i)_{y_i}).
\]
Fig. 6 Coverage of positive and negative classes, coloured according to the layers. A “perfect”
discriminating pattern for the positive class (resp. negative class) would be projected to the
lower right corner (resp. upper left corner).
The higher the fidelity, the lower the infidelity, the better the explainer.

Obviously, masking all the input graph would have important impact to the model prediction. Therefore, the former measures should not be studied without considering the Sparsity metric that aims to measure the fraction of graph selected as mask by the explainer:

$$Sparsity = \frac{1}{N} \sum_{i=1}^{N} \left( 1 - \frac{|m_i|}{|g_i|} \right),$$

where $|m_i|$ denotes the size of the mask $m_i$ and $|g_i|$ is the size of $g_i$ (the size includes the number of nodes, of edges and the attributes associated to them). Based on these measures, a better explainability method achieves higher fidelity, lower infidelity while keeping a sparsity close to 1.

We devise four policies to build a mask from an activation rules:

1. **node**: the simplest policy which takes only the nodes that are covered by the activation rule and the edges adjacent to these nodes.
2. **ego**: the ego-graphs of radius $\ell$ centered on activated nodes, with $\ell$ the layer associated to the pattern.
3. **decay**: a continuous mask with a weight associated to the edges that depends on the distance of its end-points to the activated nodes:

$$w_{v,a} = \sum_{a \in V_A} \frac{1}{2^{1+d(v,a)}} \text{ if } d(v,a) \leq \ell, 0 \text{ otherwise}$$

with $V_A$ the set of activated nodes, $d(v,a)$ the geodesic distance between nodes $v$ and $a$ and $w_{(u,v)} = w_u + w_v$.
4. **top k**: a discrete mask containing only the $k$ edges from decay mask with the highest weights ($k = 5$ or $k = 10$ in our experiments).

For each policy, we select the mask (and the related pattern) that maximises the fidelity. As GNNExplainer and PGExplainer provide continuous masks, we report, for fair comparisons, the performance with both continuous and discrete masks built with the $k$ best edges. Note that the average time to provide an explanation ranges from 8ms to 84ms for INSIDE-GNN. This is faster than PGM-Explainer (about 5s), GNNExplainer (80ms to 240ms) and Grad (300ms). It remains slightly slower than PGExplainer (6ms to 20ms). Table 4(a) summarises the performance of the explainers based on the Fidelity measures. Results show that INSIDE-GNN outperforms the baselines regardless of policy. On average, the gain of our method against the best baseline is 231% for $Fid_{prob}$ and 207% for $Fid_{acc}$. These results must be analysed while considering the sparsity (see Table 4(c)). In most of the cases, INSIDE-GNN provides sparser explanation than the baselines. Furthermore, at equal sparsity (top $k$), INSIDE-GNN obtains higher fidelity values than both competitors. Notice that PGM-Explainer fails on BA2 because this dataset does not have labeled nodes and this method investigate only the nodes of the graphs.

We provide additional information on the Fidelity in Table 5. The Fidelity aims to measure the percentage of times that a model decision is changed when the input graphs is obfuscated by the mask $m$. In Table 5 we report a polarized version of the Fidelity for which we count the number of changes between the two possible decisions of the model. For instance, $F^{a \rightarrow b}$ measures the percentage of graphs initially classified as 'false' by the model that become classified as 'true'
when obfuscating the graph with a mask. We can observe a dissymmetry between the class changes. As an example, inside-GNN has a perfect fidelity on BA2 and DD when considering only the positive examples, i.e., the mask provided by inside-GNN makes the model change its decision. When dealing with the negative examples, we obtain much lower score. Intuitively, some class changes cannot be done by only removing some vertices or edges. Regarding BA2, it is impossible to obtain a house motif from a cycle without adding an edge to form a triangle.

The quality of the explanations are also assessed with the Infidelity metrics in Table 4(b). inside-GNN achieves excellent performance on BA2. On the other datasets, inside-GNN is outperformed by GNNExplainer. inside-GNN obtain similar scores or outperforms the other competitors (i.e., PGExplainer, PGM-Explainer, Grad) at equal sparsity on most of the datasets. Notice that, in these experiments, we made the choice to build mask based on a single activation rule which is not enough to obtain fully discriminant mask for complex datasets. This is in agreement with what we observed in Fig. 6. We have no fully discriminant activation rule for the positive and negative classes. Hence, it would be necessary to combine activation rules to build a more discriminant mask and thus better optimise the Infidelity.

5.4 Model insights via the (re)description of activation patterns

We argue that activation rules also help provide insight into the model, especially what the GNN model captures. As discussed in Section 4, this requires characterizing the nodes (and their neighborhood) that support a given activation rule. In this experimental study, we investigate the obtained numerical subgroups for BA2 and the subgraph characterizing the activation rules retrieved for Mutagen, BBBP and Aids datasets.

5.4.1 Numerical subgroups

Each node can be easily described with some topological properties (e.g., its degree, the number of triangles it is involved in). Similarly, we can describe its neighborhood by aggregating the values of the neighbors. Thanks to such properties, we make a propositionalization of the nodes of the graphs. Considering the two most discriminant activation rules\(^5\), we use the subgroup discovery algorithm from pysubgroup library [Lemmerich and Becker 2018]\(^\text{1}\) to find the discriminating conditions of the nodes supporting these two patterns. Fig. 7 reports a visualisation of two graphs with activated nodes in red. The best description based on WRAcc measure of pattern \(p^1\) (Fig. 7 left) and \(p^0\) (Fig. 7 right) are given below.

For the House motif (positive class of BA2), the nodes that support activation rules are almost perfectly described (the WRacc equals to 0.24 while maximum value is 0.25) with the following conditions: **Nodes connected to two neighbors (degree≥2) that are not connected between them (clustering coefficient=0), not involved in a triangle and one of its neighbors is involved in a triangle (triangle2=1).** In other

\(^5\) \(p^1 = \{a_3, a_6, a_7, a_9, a_{10}, a_{15}\} (\text{where } a_i \text{ are the activated components of the rule and } p \text{ is the set representation of the bitset } A^p \text{ of Definition 2}), |\text{Supp}(p^1, G^1)| = 474, |\text{Supp}(p^1, G^0)| = 16 \text{ and } p^0 = \{a_0, a_1, a_2, a_4, a_5, a_8, a_{11}, a_{17}, a_{18}, a_{19}\}, |\text{Supp}(p^0, G^1)| = 137, |\text{Supp}(p^0, G^0)| = 506.
### (a) Fidelity

<table>
<thead>
<tr>
<th>Model</th>
<th>DD</th>
<th>Proteins</th>
<th>BA2</th>
<th>Acts</th>
<th>BBP</th>
<th>Mitogen</th>
</tr>
</thead>
<tbody>
<tr>
<td>INSIDE-GNN (ego)</td>
<td>0.980</td>
<td>0.975</td>
<td>0.990</td>
<td>0.955</td>
<td>0.905</td>
<td>0.705</td>
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<td>INSIDE-GNN (node)</td>
<td>0.975</td>
<td>0.940</td>
<td>0.780</td>
<td>0.705</td>
<td>0.605</td>
<td>0.405</td>
</tr>
<tr>
<td>INSIDE-GNN (decay)</td>
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<td>0.940</td>
<td>0.780</td>
<td>0.705</td>
<td>0.605</td>
<td>0.405</td>
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<tr>
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<td>0.940</td>
<td>0.780</td>
<td>0.705</td>
<td>0.605</td>
<td>0.405</td>
</tr>
<tr>
<td>INSIDE-GNN (top 10)</td>
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<td>0.940</td>
<td>0.780</td>
<td>0.705</td>
<td>0.605</td>
<td>0.405</td>
</tr>
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<td>0.975</td>
<td>0.990</td>
<td>0.955</td>
<td>0.905</td>
<td>0.705</td>
</tr>
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### (b) Infidelity

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<th>Acts</th>
<th>BBP</th>
<th>Mitogen</th>
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<td>0.185</td>
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<td>0.165</td>
<td>0.175</td>
<td>0.185</td>
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<tr>
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<td>0.150</td>
<td>0.160</td>
<td>0.170</td>
<td>0.180</td>
<td>0.190</td>
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<td>0.145</td>
<td>0.155</td>
<td>0.165</td>
<td>0.175</td>
<td>0.185</td>
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### (c) Sparsity

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<tr>
<th>Model</th>
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<th>BA2</th>
<th>Acts</th>
<th>BBP</th>
<th>Mitogen</th>
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<tr>
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words, the activation rule captures one node of the floor of the "house motif". We have similar conditions to identify some nodes of the 5-node cycle (negative class of BA2) nodes without triangle in their direct neighborhood ($\text{clustering}^2=0$) and whose sum of neighbors' degree (including itself) equal 7 ($\text{degree}^2=7$).

Table 5 Polarized fidelity.

<table>
<thead>
<tr>
<th>(a) Fidelity</th>
<th>Model</th>
<th>DD</th>
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<th>Aids</th>
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<td>0.000</td>
<td>1.000</td>
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</tr>
</tbody>
</table>

Luca Verri-Ferrer et al.
On GNN explainability with activation rules

clustering=0.0 AND degree=2
AND triangle2=1 AND triangle=0
WRAcc=0.02

clustering2=0.0 AND degree=2
WRAcc=0.24

Fig. 7 Nodes (in red) in the support of two activation rules that are discriminant for \( p^1 \) support, related to the positive target (left), and for \( p^0 \) support, related to the negative target (right).

Table 6 Characterization of activation rules with numerical subgroups on BA2. We only report the subgroup whose WRAcc value is greater than 0.1.

<table>
<thead>
<tr>
<th>Layer</th>
<th>Class</th>
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<td>0.2475</td>
</tr>
<tr>
<td>2</td>
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<td>clustering2=0 AND degree=2 AND triangle2_avg=0</td>
<td>0.207</td>
</tr>
<tr>
<td>3</td>
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<td>0.101</td>
</tr>
<tr>
<td>3</td>
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<td>betweenness2: [0.37:0.38] AND betweenness2_avg: [0.19:0.20] AND clustering2=0.0</td>
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</tr>
<tr>
<td>3</td>
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<tr>
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</tr>
<tr>
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<td>0.162</td>
</tr>
<tr>
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<td>1</td>
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<td>0.227</td>
</tr>
<tr>
<td>3</td>
<td>1</td>
<td>degree2: [7.8] AND degree2_avg: [3.50:3.60] AND degree=2 AND triangle=0</td>
<td>0.224</td>
</tr>
<tr>
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<td>1</td>
<td>degree=2 AND triangle2=1</td>
<td>0.238</td>
</tr>
<tr>
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<td>1</td>
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<td>0.240</td>
</tr>
<tr>
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<td>1</td>
<td>degree=2</td>
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</tr>
<tr>
<td>3</td>
<td>1</td>
<td>clustering=0.0 AND degree=2 AND triangle2=1 AND triangle2_avg=0.5</td>
<td>0.232</td>
</tr>
</tbody>
</table>

We report the description in terms of numerical subgroups of the activation rules in Table 6. It is important to note that even if some activation rules were found as subjectively interesting according to a specific output of the model, they may capture some general properties of the BA2 graph that are not so specific of one of the classes. For instance, the second subgroup is related to the positive class (i.e., house motif) but what it captured is not specific to house motif (degree=2, absence of triangle).
5.4.2 Graph subgroups

Similarly, we can characterize activation rules with graph subgroups. We investigate the interest of such pattern language for three datasets: Aids, BBBP and Mutagen. For each activation rule, we compute the graph that has the maximum WRAcc value, using min\_sup = 10 (see Equation 1). In other words, this graph has an important number of isomorphisms with ego-graphs that support the rule and that correspond to the class of the target of the rule. In Fig. 8, we report the WRAcc values of the discovered graphs that aim to characterize the activation rules. We can observe that the WRAcc values are rather high (WRAcc ranges from -1 to 0.25) which demonstrates that these graphs well describe the parts of the GNN identified by the activation rules.

The subgraphs obtained for Mutagen dataset are summarised in Fig. 9. For each layer and decision, we display the subgraphs whose WRAcc is greater than 0.1 layer by layer. The negative class is related to mutagenic molecules. Several things can be observed from this figure. First, some subgraphs are known as toxicophores or fragment of toxicophores in the literature (Kazius et al., 2005). For instance, the subgraph with two hydrogen and one azote atoms is a part of an aromatic amine. Similarly, the subgraph with one azote and two oxygen atoms is an aromatic nitro. The subgraph involving 6 carbon atoms is a fragment of a bay region or a k-region. Second, some subgraphs appear several times. It means that several activation rules are described with the same subgraphs. This can be explained in several ways. Neural networks are known to have a lot of redundant information, as evidenced by the numerous papers in the domain that aim to compress or simplify deep neural networks (Chen et al., 2018; Pan et al., 2016; Pasandi et al., 2020; Xu et al., 2018). Accordingly, this is not surprising to have several parts of the GNN that are similar and described by the same subgraphs. Notice that this problem could be an interesting perspective for our work. Another explanation is that the subgraphs well describe the hidden features captured by the GNN but from different perspective, i.e., the center is different. For instance, for a simple chemical bond C-N, one may have the same graph with one centered in C and the other in N. A last explanation could be that the subgraph language is not enough powerful to capture the subtle differences between the activation rules. Once again, the definition of more sophisticated and appropriate languages to describe the hidden features captured by the GNN is a promising perspective of research.

These latter experiments show that inside-GNN represents a valuable alternative to GNN explainability methods. In addition to providing single instance explanations, inside-GNN can provide insights about what the GNN perceives. Especially, it allows to build a summary of the hidden features captured by the model (e.g., Fig. 8). In relation to this, our method is quite analogous to model explanation methods such as XGNN (Yuan et al., 2020a). This deserves a discussion and a comparison with XGNN.

5.4.3 Comparison to XGNN

XGNN (Yuan et al., 2020a) is a method rooted in reinforcement learning that generates graphs that maximise the model decision for a given class. For Mutagen, we generate 20 graphs for each class with a maximum size equal to 6. Considering
Fig. 8 Boxplot of the WRAcc values of graph subgroups related to activation rules by layer (left column) or by both layer and model decision (right column) for Aids (first row), BBBP (second row) and Mutagen (third row).

the 40 generated graphs, we observe that only one of them is a subgraph of at least one graph of the dataset. The other graphs have on average 60% of partial inclusion: the maximum common subgraph with molecules from Mutagen uncovers 60% of a generated graph. Therefore, we can conclude that XGNN generates graphs that are not enough realistic. The only graph that appears within the dataset involves a carbon atom bonded to 2 others carbon atoms and one hydrogen atom. With INSIDE-GNN, we obtained two subgraphs characterizing some activation rules that are super-graphs of this one (see Fig. 9). Notice that, we also found this
Fig. 9 Characterization of activation rules for Mutagen with discriminant subgraphs. We retain only the subgraphs with a WRAcc value greater than 0.1. Mutagenic chemicals are classified as False.

subgraph for some activation rules. We did not report it in Fig. 9 because its WRAcc value is lower than 0.1. Nevertheless, this graph appears in 21100 ego-graps in the dataset. It describes a fragment of molecule that is very common. One can wonder if such a fragment can be mutagenic or if XGNN has just captured it a biased of the GNN. Furthermore, XGNN has generated graphs that are not planar, which is not common in Chemistry. Based on these evidences, we argue that XGNN does not return realistic graphs while our approach – by construction – provides subgraphs from the dataset.

We search for each pattern produced by inside-GNN the closest pattern in XGNN according to the Graph Edit Distance (GED) and vice versa. We note that
the previously described prototype graph (i.e., 3 carbons and 1 hydrogen) is found
in most of the cases as being the closest to the patterns produced by INSIDE-GNN.
In average, the distance between each XGNN prototype and the closest pattern of
INSIDE-GNN is 4.6 while the mean distance between INSIDE-GNN subgraphs and the
closest from XGNN is 3.7. This is rather important since the graphs provided by
XGNN have at most 6 nodes.

We believe that a model decision for a class cannot be summarized into a single
prototype. Several different phenomena can lead to the same class. Furthermore,
as we observed, this can lead to unrealistic prototype even if domain knowledge is
integrated within the graph generation. INSIDE-GNN allows to have deeper insights
from the GNN by considering each hidden feature separately.

6 Discussion and Conclusion

We have introduced a novel method for the explainability of GNNs. INSIDE-GNN
is based on the discovery of relevant activation rules in each hidden layer of the
GNN. Prior beliefs are used to assess how contrastive a rule is. We have proposed
an algorithm that efficiently and iteratively builds a set of activation rules, limiting
the redundancy between them. Extensive empirical results on several real-world
datasets confirm that the activation rules capture interesting insights about how
the internal representations are built by the GNN. Based on these rules, INSIDE-
GNN outperforms the SOTA methods for GNN explainability when considering
Fidelity metric. Furthermore, the consideration of pattern languages involving
interpretable features (e.g., numerical subgroups on node topological properties,
graph subgroups) is promising since it makes possible to summarise the hidden
features built by the GNN through its different layers.

We believe that such method can support knowledge discovery from powerful
GNNs and provide insights on object of study for scientists or more generally
for any user. However, a number of potential limitations need to be considered for
future research to make this knowledge discovery from GNNs effective in practice.

First, assessing explanations without ground truth is not trivial. Our experi-
mental evaluation relies on Fidelity, Infidelity and Sparsity metrics. Fidelity as-
sumes that the GNN decision would change if key part of the graphs are removed.
However, it is not always the case in practice. For instance, it is difficult to obtain
a toxic molecule from a non-toxic one by only removing some atoms. That would
be interesting to investigate other evaluation measures that take into account the
negation (i.e., absence of important features) and evaluation measures based on
the addition of subgraphs.

In this paper, we have devised an exhaustive algorithm for discovering the acti-
vation rules. Even if pruning based on upper bound is featured, the execution time
remains a problem. It ranges from few minutes to two days. This shows only the
feasibility of the proposed method, not its practical application. To overcome this
limitation, the completeness must be relaxed and some heuristic-based algorithms
have to be defined. Beam-search or Monte-Carlo Tree Search-based algorithms are
good alternatives to the one we propose.

Activation rule patterns are the simplest pattern language to deal with activa-
tion matrices since such patterns involve only conjunction of activated components.
Even simple, these activation rules are able to capture the hidden features built
by the GNN as witnessed by the experiments. We believe that more sophisticated
pattern languages are possible for GNNs. For instance, we observed that taking
into account the number of occurrences within a graph leads to better characteri-
sations. This can be integrated to the pattern language. Considering the negation
(i.e., the absence of activations) is also promising and would offer a deeper de-
scription of the internal mechanism of the GNNs.

With INSIDE-GNN, the activation rules are mined for each layer independently.
As a consequence, the relations between layers are not taken into account in the
discovery of activation rules. This may lead to redundant results when considering
all the layers. To avoid such redundancy, it is necessary to take into account as
prior knowledge the previous layers of a given layer.

Finally, activation rules capture specific configurations in the embedding space
of a given layer that is discriminant for the GNN decision. Experiments demon-
strate that these rules can be directly used to support instance-level model ex-
planation. However, activation rules cannot be easily interpreted by human beings
because of the pattern language itself (i.e., conjunction of activated components
of the hidden layers). The consideration of pattern languages with interpretable
features makes it possible to characterize them. However, this second step can be
improved by query the model itself. Indeed, the current characterization methods
investigate a dataset generated from the support of the activation rules. The model
should be considered in this step to have guarantee that the interpretable pattern
that describes a rule well embeds in the subspace related to this rule.

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