Generalization and Representational Limits of Graph Neural Networks

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Abstract

We address two fundamental questions about graph neural networks (GNNs). First, we prove that several important graph properties, e.g., shortest/longest cycle, diameter, or certain motifs, cannot be computed by GNNs that rely entirely on local information. Such GNNs include the standard message passing models, and more powerful variants that exploit local graph structure (e.g., via relative orientation of messages, or local port ordering) to distinguish neighbors of each node. Our treatment includes a novel graph-theoretic formalism.

Second, we provide the first data dependent generalization bounds for message passing GNNs. This analysis explicitly accounts for the local permutation invariance of GNNs. Our bounds are much tighter than existing VC-dimension based guarantees for GNNs, and are comparable to Rademacher bounds for recurrent neural networks.

1. Introduction

Graph neural networks (Scarselli et al., 2009; Gori et al., 2005), in their various incarnations, have emerged as models of choice for embedding graph-structured data from a diverse set of domains, including molecular structures, knowledge graphs, biological networks, social networks, and n-body problems (Duvenaud et al., 2015; Defferrard et al., 2016; Battaglia et al., 2016; Zhang et al., 2018b; Santoro et al., 2018; Yun et al., 2019; Jin et al., 2019).

The working of a graph neural network (GNN) on an input graph, with a feature vector associated with each node, can be outlined as follows. Layer \( \ell \) of the GNN updates the embedding of each node \( v \) by aggregating the feature vectors, or node and/or edge embeddings, of \( v \)’s neighbors from layer \( \ell - 1 \) via a non-linear transformation, possibly combining this with \( v \)’s embedding. The exact form of the aggregate and combine steps varies across architectures, and empirical success has been demonstrated for several variants. These include Graph Convolutional Networks (GCN) (Kipf and Welling, 2017), Graph Attention Networks (GAT) (Veličković et al., 2018), Graph Isomorphism Network (GIN) (Xu et al., 2019), and GraphSAGE (Hamilton et al., 2017). GNNs are known to have fundamental connections to message passing (Dai et al., 2016; Gilmer et al., 2017), the Weisfeiler-Leman (WL) graph isomorphism test (Xu et al., 2019; Morris et al., 2019), and local algorithms (Sato et al., 2019; Loukas 2020).

In this work, we investigate the representational limitations and generalization properties of GNNs. That is, we examine the performance of GNNs from a learning perspective: (a) how well they can discriminate graphs that differ in a specific graph property (represented by assigning different labels to such graphs), and (b) how well they can predict labels, e.g., a graph property, for unseen graphs. Specifically, we focus on classification: (1) a GNN with learnable parameters embeds the nodes of each input graph, (2) the node embeddings are combined into a single graph vector via a readout function such as sum, average, or element-wise maximum, and (3) a parameterized classifier either makes a binary prediction on the resulting graph vector (our setting for representation limits), or local binary predictions, one per node, the majority of which determines the graph label (our setting for generalization bounds).

Our contributions. (1) We show that GNNs that generate node embeddings solely based on local information cannot distinguish some simple non-isomorphic graphs. As a result, these GNNs cannot compute important graph properties such as longest or shortest cycle, diameter, etc. This limitation holds for popular models such as GraphSAGE, GCN, GIN, and GAT. Our impossibility results also extend to more powerful variants that provide to each node information about the layout of its neighbors, e.g., via port numbering, like CPNGNN (Sato et al., 2019), or geometric information, like DimeNet (Klicpera et al., 2020).

We introduce a novel graph-theoretic formalism for analyzing CPNGNNs, and our constructions provide insights that may facilitate the design of more effective GNNs.
Among other works, Barceló et al. (2020) proved results for message passing GNNs. Our guarantees are significantly tighter than the VC bounds established by Scarselli et al. (2018) for a class of GNNs. Interestingly, the dependence of our bounds on parameters is comparable to Rademacher bounds for recurrent neural networks (RNNs). Our results also hold for folding networks (Hammer, 2001) that operate on tree-structured inputs.

Our generalization analysis specifically accounts for local permutation invariance of the GNN aggregation function. This relies on a specific sum form that extends to aggregating port-numbered messages, and therefore opens avenues for analyzing generalization of CPNGNNs.

The first part of the paper is dedicated to analyzing what GNNs cannot learn. The second part investigates how well GNNs learn when they can: we establish new generalization bounds for GNNs. We outline some key proof steps for our results in the main text, and defer the detailed proofs to the Supplementary material.

2. Related Work

GNNs continue to generate much interest from both theoretical and practical perspectives. An important theoretical focus has been on understanding the expressivity of existing architectures, and thereby introducing richer (invariant) models that can generate more nuanced embeddings. But, much less is known about the generalization ability of GNNs. We briefly review some of these works.

Expressivity. Scarselli et al. (2009) extended the universal approximation property of feed-forward networks (FFNs) to GNNs using the notion of unfolding equivalence. Recurrent neural operations for graphs were introduced with their associated kernel spaces (Lei et al., 2017). Dai et al. (2016) performed a sequence of mappings inspired by mean field and belief propagation procedures from graphical models, and Gilmer et al. (2017) showed that common graph neural net models models may be studied as Message Passing Neural Networks (MPNNs). It is known (Xu et al., 2019) that GNN variants such as GCNs (Kipf and Welling, 2017) and GraphSAGE (Hamilton et al., 2017) are no more discriminative than the Weisfeiler-Leeman (WL) test. In order to match the power of the WL test, Xu et al. (2019) also proposed GINs. Showing GNNs are not powerful enough to represent probabilistic logic inference, Zhang et al. (2020) introduced Express-GNN.

Among other works, Barceló et al. (2020) proved results in the context of first order logic, and Dehmamy et al. (2019) investigated GCNs through the lens of graph moments underscoring the importance of depth compared to width in learning higher order moments. The inability of some graph kernels to distinguish properties such as planarity has also been established (Kriege et al., 2018, 2020).

Spatial, hierarchical, and higher order GNN variants have also been explored. Notably, Sato et al. (2019) exploited a local port ordering of nodes to introduce the Consistent Port Numbering GNN (CPNGNN), which they proved to be strictly more powerful than WL. They and Loukas (2020) also established connections to distributed local algorithms. Higher order generalizations have been studied by Morris et al. (2019), Murphy et al. (2019), Maron et al. (2019c), in particular, Maron et al. (2019a) introduced models that are more powerful than WL. Hella et al. (2015) investigated models weaker than port numbering. Several other works exploit spatial information to obtain more nuanced embeddings (Ying et al., 2018, You et al., 2019, Ingraham et al., 2019, Klicpera et al., 2020, Chen et al., 2019). Xu et al. (2018) learned locally adaptive structure-aware representations by adaptively aggregating information over extended neighborhoods. Veličković et al. (2018) introduced GATs that obviate specifying the graph structure in advance.

Concurrent to our work, Chen et al. (2020b) established results regarding the ability of MPNNs and Invariant Graph Networks (IGNs) to count substructures in graphs. Sub-graph counts and related properties have also been studied in Arvind et al. (2020). Sato et al. (2020) advocated adding random features to nodes in order to better approximate some problems. We refer the reader to Sato (2020) for a survey about expressivity of GNNs.

Invariance. An important consideration in the design of GNNs is their ability to produce output embeddings that are equivariant or permutation-invariant to the input feature vectors. Maron et al. (2019b) constructed permutation-invariant and equivariant linear layers, and showed that their model can approximate any GNN that can be cast as a MPNN in the framework of (Gilmer et al., 2017). Murphy et al. (2019) constructed new permutation-invariant functions for variable-size inputs, and suggested some approximations. Maron et al. (2019c). Keriven and Peyré (2019) proved universality theorems for a specific class of invariant and equivariant networks, respectively.

Generalization. Several works have established generalization guarantees for FFNs (Bartlett et al., 2017, Golowich et al., 2018, Neyshabur et al., 2018, Zhang et al., 2018a) and RNNs (Chen et al., 2020a, Allen-Zhu and Li, 2019). GNNs differ in some key aspects from those models. Unlike RNNs that process sequences, GNNs operate on graph-structured data: sharing of recurrent weights takes place along both the depth and width of a GNN. Unlike FFNs, GNNs deal with irregular local structure. Moreover, at each node, GNNs typically employ permutation-invariant aggregations, in contrast to global permutation invariance (Sokolic et al., 2017). Scarselli et al. (2018) proved VC-
When computing node embeddings, the embedding of node $v$ is updated by processing the information from its neighbors as an ordered set, ordered by the port numbering, i.e., the aggregation function is generally not permutation invariant. In addition to a neighbor node $u$’s current embedding, $v$ receives the port number that connects $u$ to $v$.

Another model, DimeNet (Klicpera et al. 2020), is a directional message passing algorithm introduced in the context of molecular graphs. Specifically, DimeNet embeds atoms via a set of messages (i.e., edge embeddings) and leverages the directional information by transforming messages based on the angle between them. For each node $v$, the embedding for an incoming message from neighbor $u$ is computed as

$$m_{uv}^{(t)} = f_1 (m_{uv}^{(t-1)}, \tilde{m}_{uv}^{(t-1)})$$

$$\tilde{m}_{uv}^{(t-1)} = \sum_{w \in N(u) \setminus \{v\}} f_2 (m_{uw}^{(t-1)}, e_{uw}, a_{uw}, a_{uw})$$

and $e_{uw}$ is a representation of the distance from $u$ to $v$, $a_{uw}$ combines $\angle uvw$ with the distance from $w$ to $u$, and $f_1$ and $f_2$ are update functions similar to AGG and COMBINE. The node embedding $h_v^{(t)}$ is simply the sum of message embeddings $m_{uv}^{(t)}$.

For the purposes of this paper, message passing GNNs consist of LU-GNNs, CPNGNN, and DimeNet. We establish representational limits of message passing GNNs with respect to several important graph properties in this paper. Namely, (a) girth (length of the shortest cycle), (b) circumference (length of the longest cycle), (c) diameter (maximum distance, in terms of shortest path, between any pair of nodes in the graph), (d) radius (minimum node eccentricity, where eccentricity of a node $u$ is defined as the maximum distance from $u$ to other vertices), (e) conjunct cycle (two cycles that share an edge), (f) total number of cycles, and (g) $k$-clique (a subgraph of at least $k \geq 3$ vertices such that each vertex in the subgraph is connected by an edge to any other vertex in the subgraph).

We will appeal to the following definition to formalize when GNNs cannot distinguish non-isomorphic graphs, that is, when two distinct graphs are represented with the same embedding regardless of GNN parameters.

**Definition 1.** For a given graph property $P$ and readout function $f$, we say that a GNN $Q$ decides $P$, if for any pair of graphs $(G_1, G_2)$ such that $G_1$ and $G_2$ differ on $P$, we have $f(g_Q(G_1)) \neq f(g_Q(G_2))$. Here, $g_Q(G)$ denotes the collection of embeddings of nodes in $G$ when $G$ is provided as input to $Q$. By extension, a class $\mathcal{Q}$ of GNNs cannot decide $P$ if there does not exist any $Q \in \mathcal{Q}$ that decides $P$.

We can compute the embedding of any node $v$ using a local computation tree, rooted at $v$ and obtained by unrolling...
4. Representational limits of GNNs

We now sketch novel constructions to illustrate the limits of LU-GNNs, CPNGNNs, and DimeNets. First, we show that in some cases, CPNGNNs can be more discriminative than LU-GNNs, depending on the port numbering. Then, we demonstrate that still, LU-GNNs, CPNGNNs, and DimeNets cannot compute certain graph properties. Our proofs build examples of graphs that (1) differ in important graph properties, but that (2) these models cannot distinguish. As a consequence, these models will not be able to compute such graph properties in general.

To formalize this framework, we introduce a condition of local isomorphism for a pair of graphs. This condition implies that CPNGNNs and LU-GNNs cannot distinguish the two graphs. A similar framework applies to DimeNet. Finally, our insights point to a new GNN variant that leverages additional geometric features to circumvent our constructions for CPNGNNs and DimeNets.

Limitations of LU-GNNs.

**Proposition 1.** There exist non-isomorphic graphs that LU-GNNs cannot distinguish, but CPNGNN can distinguish with some consistent port ordering.

Fig. 1 shows two graphs, \( G \) (consisting of two triangles) and \( \Gamma \). Nodes with the same color (or, equivalently, same uppercase symbol without the subscripts and underline) have identical feature vectors. The port numbers for each node are shown next to the node on the respective edges; the numbering is consistent. Moreover, for LU-GNNs, edges on nodes with same color have identical edge feature vectors; for CPNGNNs, edge features are the same if, in addition, the local ports for nodes that have the same color are identical. As explained in Fig. 1, CPNGNN can distinguish between the two graphs by exploiting the port information. However, LU-GNNs do not leverage such information, and fail to find distinct representations.

We note that Theorem 2 in Sato et al. (2019) also establishes the discrepancy in expressivity of CPNGNNs and LU-GNNs, by exploiting a connection between GNNs and local algorithms from distributed computing. In contrast to their approach, our construction in Proposition 1 may also be invoked to argue that DimeNets are strictly more expressive than LU-GNNs.

**Limitations of CPNGNNs.** Port orderings can help CPNGNNs distinguish graphs that LU-GNNs cannot. But, port orderings are not unique, and not all orderings distinguish the same set of graphs.

**Proposition 2.** There exist pairs of non-isomorphic graphs and consistent port numberings \( p \) and \( q \) such that CPNGNN can distinguish the graphs with \( p \) but not \( q \).

Fig. 2 shows the same pair of graphs \( G \) and \( \Gamma \), but with a different ordering. CPNGNN can no longer distinguish the two non-isomorphic graphs with this new ordering. Thus, it may be useful to try multiple random orderings, or even parameterize and learn one along with GNN parameters.

Henceforth, we assume that an ordering is given with the input graph. We now demonstrate the inability of CPNGNNs to decide several graph properties. Toward this goal, note that in Fig. 1, we constructed an explicit bijection...
between nodes in $G$ and $\mathcal{G}$ to reason about permutation-invariant readouts. We now introduce a graph-theoretic formalism for CPNGNNs that obviates the need for an explicit bijection and is easier to check.

We define a pair of surjective mappings between two graphs in question, and impose additional conditions that guarantee the existence of a bijection. This bijection implies that corresponding nodes in the graphs receive identical embeddings, and hence both graphs obtain the same set of node embeddings, making them indistinguishable.

The main idea is that a node $v_1$ in graph $G_1$ is locally indistinguishable from $v_2$ in $G_2$ if (1) the node features agree: $x_{v_1} = x_{v_2}$, and (2) the port-ordered local neighborhoods of $v_1$ and $v_2$ cannot be told apart. That is, if port $i$ of $v_1$ connects to port $k$ of $v$, then a locality preserving bijection connects the nodes corresponding to images of $v_1$ and $v$ via the same ports. In the notation here, we include the port numbers $(i, j)$ associated with each edge $(u, v)$ in the edge notation, i.e., $((u, i), (v, j))$.

**Definition 2.** We say that graph $G_1(V_1, E_1, p)$ port-covers $G_2(V_2, E_2, q)$ if the following conditions are satisfied: (a) there exists a surjection $f : V_1 \rightarrow V_2$ such that $x_v = x_{f(v)}$ for all $v \in V_1$, (b) $p$ and $q$ are consistent, and (c) for all $v \in V_1$ there exists a local bijection $g_{v_1}$ such that for all $i \in \text{degree}(v_1)$ and $(v, k) = p(v_1, i)$, we have

$$g_{v_1}(((v_1, i), (v, k))) = (q(f(v), k), q(f(v_1), i)),$$

such that $q(f(v), k) = (f(v_1), i); q(f(v_1), i) = (f(v), k); ((v_1, i), (v, k)) \in E_1$ and $(q(f(v), k), q(f(v_1), i)) \in E_2$. Moreover, we say that $G_1(V_1, E_1, p)$ and $G_2(V_2, E_2, q)$ are port-locally isomorphic if they both cover each other.

Definition 2 does not preclude the possibility that $f$ maps multiple nodes in $G_1$ to the same node in $G_2$, or the other way round. Hence, the claim that $G_1$ and $G_2$ cannot be distinguished by CPNGNN might not hold. Fortunately, the following result comes to our rescue.

**Proposition 3.** If $G_1(V_1, E_1, p)$ and $G_2(V_2, E_2, q)$ are port-locally isomorphic, there exists a bijection $h$ that satisfies (a)-(c) in Definition 2 (with $h$ replacing $f$). As a result, CPNGNNs produce identical embeddings for the corresponding nodes in $G_1$ and $G_2$, so CPNGNNs cannot separate $G_1$ and $G_2$ with permutation-invariant readout.

We may characterize port-local isomorphism in alternative ways, e.g., in terms of an identical multiset of port-numbered computation trees (i.e., computation trees with associated port numbers on edges). Our formalism allows us to construct an explicit port-cover bijection from only a pair of surjective mappings, one in each direction (see the proof of Proposition 3 for details). Thus, it provides insight into the role played by locally isomorphic neighborhoods in unraveling a port-cover bijection whenever it exists.

We now proceed to establish that CPNGNNs are limited in that they fail to decide important graph properties. We can invoke conditions of Proposition 3 or define a bijection, to show the following result (see Fig. 3 for our constructions).

**Proposition 4.** There exist consistent port orderings such that CPNGNNs with permutation-invariant readout cannot decide several graph properties: girth, circumference, diameter, radius, conjoint cycle, total number of cycles, and $k$-clique.

Clearly, these impossibility results apply to LU-GNNs as well (see Fig. 4). However, as described in Fig. 4, our constructions for CPNGNNs do not work for DimeNets. This immediately leads us to the question whether DimeNets are expressive enough to decide the graph properties.

**Limitations of DimeNets.** Unfortunately, as we show in Fig. 4, there exists an example of two graphs that differ in several of these properties but cannot be distinguished by DimeNets.

**Proposition 5.** DimeNet with permutation-invariant readout cannot decide several graph properties: girth, circumference, diameter, radius, or total number of cycles.
In fact, as we argue in Fig. 4, augmenting DimeNet with port-numbering would still not be sufficient. Therefore, a natural question that arises is whether we can obtain a more expressive model than both CPNGNN and DimeNet. Leveraging insights from our constructions, we now introduce one such variant, H-DCPN (short for Hierarchical Directional Message Passing Consistent Port Numbering Networks), that generalizes both CPNGNN and DimeNet.

More powerful GNNs. The main idea is to augment DimeNet not just with port ordering, but also additional spatial information. Observe that the construction in Fig. 4 will fail if for each edge \((u, v)\), we additionally model the set of angles \(\alpha_{uv}\) between planes \(P(w, u, v)\) and \(P(u, v, z)\) due to neighbors \(w \) of \(u\) and neighbors \(z \) of \(v\). Similarly, we could use the distances between these planes. We denote by \(\Phi_{uv}\) all such features derived from these planes. Denote by \(m_{uv}(l)\) the message from neighbor \(u\) of \(v\) at time \(l\), and by \(m_{uv}(l) = f(m_{uv}(l), \Phi_{uv})\) a refined message that encapsulates the effect of geometric features.

We can incorporate salient aspects of CPNGNN as well. Specifically, we first fix a consistent port numbering, as in CPNGNN. Denote the degree of \(v\) by \(d(v)\). Let \(c_e(j)\) be the neighbor of \(v\) that connects to port \(j\) of \(v\) via port \(t_j\), for \(j \in [d(v)]\). We suggest to update the embedding of \(v\) as

\[
h_v^{(l)} = f(h_v^{(l-1)}, m_{uv}^{(l-1)}, t_1, v, \ldots, m_{uv}^{(l-1)}, t_{d(v)}, v),
\]

where \(f\) can potentially take into account the ordering of its arguments. The update resembles CPNGNN when we define \(m_{uv}^{(0)} = h_v^{(0)}\); and DimeNet when \(f\) ignores \(h_v^{(l-1)}\) (and ports) and we define \(m_{uv}^{(l)}\) using (1) in section 3. H-DCPN derives its additional discriminative power from the features \(\Phi_{uv}\) encoded in messages \(m_{uv}^{(l)}\). For instance, the nodes labeled \(A_1, B_1, C_1, D_1\) lie on the same plane in \(G_3\). In contrast, the plane defined by nodes with labels \(A_1, B_1, C_1\) in \(G_4\) is orthogonal to that defined by nodes with labels \(D_2, A_1, B_1\); thus allowing H-DCPN to distinguish the node labeled \(A_1\) from the node labeled \(A_1\) (Fig. 4).

5. Generalization bounds for GNNs

Next, we study the generalization ability of GNNs via Rademacher bounds, focusing on binary classification. We generalize the previous results on the complexity of feedforward networks (Bartlett et al., 2017; Neyshabur et al., 2018) and RNNs (Chen et al., 2020a) in mainly three ways.

First, we process graphs, unlike sequences in RNNs, or instances restricted to the input layer in feedforward networks. In particular, we show that the complexity of GNNs that combine predictions from individual nodes may be analyzed by focusing on local node-wise computation trees. Second, we share weights across all nodes in these computation trees (i.e., both along the depth and the width of the tree). Third, we model local permutation-invariance in the aggregation function at each node in the tree. Curiously, our resulting bounds are comparable to Rademacher bounds for RNNs.

We consider locally permutation invariant GNNs, where in each layer \(\ell\), the embedding \(h_v^\ell \in \mathbb{R}^r\) of node \(v\) of a given input graph is updated by aggregating the embeddings of its neighbors, \(u \in N(v)\), via an aggregation function \(\rho : \mathbb{R}^r \rightarrow \mathbb{R}^r\). Different types of updates are possible; we focus on a mean field update (Dai et al., 2016; Jin et al., 2018; 2019)

\[
h_v^\ell = \phi(W_1 x_v + W_2 \rho(\sum_{u \in N(v)} g(h_u^{\ell-1}))),
\]

where \(\phi\) and \(g\) are nonlinear transformations and \(x_v \in \mathbb{R}^r\) is the feature vector for \(v\). We assume \(\rho(0) = 0, ||x_v||_2 \leq B_2\) for all \(v, ||\phi(x)||_\infty \leq b < \infty\) for all \(x \in \mathbb{R}^r\), \(\phi(0) = 0\), and \(g(0) = 0\). One possible choice of \(\phi\) is a squashing function such as tanh. We also assume that \(\phi, \rho, g\) have Lipschitz constants \(C_\phi, C_\rho, C_g\) respectively; and that \(W_1\) and \(W_2\) have bounded norms: \(||W_1||_2 \leq B_1, \ ||W_2||_2 \leq B_2\). The weights \(W_1, W_2\) and functions \(\phi, \rho, g\) are shared across nodes and layers.

The graph label is generated by a readout function that aggregates node embeddings of the final layer \(L\). Here, we assume this function applies a local binary classifier of the form \(f_c(h_v^L) = \psi(\beta^T h_v^L)\) from a family \(\mathcal{F}_\beta\) parameterized by \(\beta\) such that \(||\beta||_2 \leq B_\beta\), with sigmoid function \(\psi\), to each node representation \(h_v^L\), and then averages the binary predictions of all nodes, i.e., \(f(G) = \sum_{v \in V} f_c(h_v^L)\). We predict label 1 if \(f(G) > 0.5\), else 0. Such networks implement permutation invariance locally in each neighborhood, and globally when aggregating the node embeddings. This invariance will play an important role in the analysis.

Let \(f(G)\) be the output of the entire GNN for input graph \(G\) with true label \(y \in \{0, 1\}\). Our loss is a margin loss applied
to the difference in probability between true and incorrect label:
\[
\tau(f(G), y) = y(2f(G) - 1) + (1 - y)(1 - 2f(G)),
\]
with \(\tau(f(G), y) < 0\) if and only if there is a classification error. The margin loss is then, with \(a = -\tau(f(G), y)\) and indicator function \(1[\cdot]\):
\[
\text{loss}_\gamma(a) = 1[a > 0] + (1 + a/\gamma)1[a \in [-\gamma, 0]].
\]

A standard result in learning theory relates the population risk \(P[\tau(f(G), y) \leq 0]\) to the empirical risk for training examples \(\{(G_j, y_j)\}_{j=1}^m\):
\[
\hat{R}_\gamma(f) = \frac{1}{m} \sum_{j=1}^m \text{loss}_\gamma(\tau(f(G_j), y_j))
\]
and to the empirical Rademacher complexity \(\hat{R}_\mathcal{S}(\mathcal{J}_\gamma)\) of the class \(\mathcal{J}_\gamma\) of functions concatenating the loss with the GNN prediction function \(f\).

**Lemma 1 [Mohri et al. (2012)].** For any margin \(\gamma > 0\), any prediction function \(f\) in a class \(\mathcal{F}\) and \(\mathcal{J}_\gamma \in \{(G, y) \mapsto \text{loss}_\gamma(\tau(f(G), y))/f \in \mathcal{F}\}\), given \(m\) samples \((G_j, y_j) \sim \mathcal{D}\), with probability \(1 - \delta\), the population risk for \(\mathcal{D}\) and \(f\) is bounded as
\[
\mathbb{P}(\tau(f(G), y) \leq 0) \leq \hat{R}_\gamma(f) + 2\hat{R}_\mathcal{S}(\mathcal{J}_\gamma) + 3\sqrt{\frac{\log(2/\delta)}{2m}}.
\]

Hence, we need to bound the empirical Rademacher complexity \(\hat{R}_\mathcal{S}(\mathcal{J}_\gamma)\) for GNNs. We do this in two steps: (1) we show that it is sufficient to bound the Rademacher complexity of local node-wise computation trees; (2) we bound the complexity for a single tree via recursive spectral bounds, taking into account permutation invariance.

### 5.1. From Graphs to Trees

We begin by relating the Rademacher complexity of \(\mathcal{J}_\gamma\) to the complexity of each node classification. The node embedding \(h^L_t\) is equal to a function applied to the local computation tree of depth \(L\), rooted at \(v\), that we obtain when unrolling the \(L\) neighborhood aggregations. That is, the tree represents the structured \(L\)-hop neighborhood of \(v\), where the children of any node \(u\) in the tree are the nodes in \(N(u)\). Hence, if \(t\) is the tree at \(v\), we may write, with a slight abuse of notation, \(f_c(h^L_t) = f_c(t; \Theta)\), where \(\Theta\) represents the parameters \(W_1, W_2\) of the embedding and \(\beta\) of the node classifier.

With this notation, we rewrite \(f(G; \Theta)\) as an expectation over functions applied to trees. Let \(T_1, \ldots, T_n\) be the set of all possible computation trees of depth \(L\), and \(w_i(G)\) the number of times \(T_i\) occurs in \(G\). Then, note that we may write \(f(G; \Theta)\) as the sum
\[
\sum_{i=1}^n \frac{w_i(G)}{w(G)} f_c(T_i; \Theta) = \mathbb{E}_{T \sim w(G)} f_c(T; \Theta).
\]

This perspective implies a key insight of our analysis: the complexity of the GNN may be bounded by the complexity of the computation trees.

**Proposition 6.** Let \(\mathcal{G} = \{G_1, \ldots, G_m\}\) be a set of i.i.d. graphs, and let \(\mathcal{T} = \{t_1, \ldots, t_m\}\) be such that \(t_j \sim w'(G_j), j \in [m]\). Denote by \(\hat{R}_\mathcal{G}\) and \(\hat{R}_\mathcal{T}\) the empirical Rademacher complexity of GNNs for graphs \(\mathcal{G}\) and trees \(\mathcal{T}\), respectively. Then \(\hat{R}_\mathcal{G} \leq \mathbb{E}_{t_1, \ldots, t_m} \hat{R}_\mathcal{T}\).

Therefore, to apply Lemma 1 it is sufficient to bound the Rademacher complexity of classifying single node-wise computation trees. Before addressing this next step in detail, we state and discuss our main result for this section.

### 5.2. Generalization Bound for GNNs

We define the percolation complexity of our GNNs to be the product \(\mathbb{C} \triangleq C_\rho C_\phi C_\beta B_2\) of Lipschitz constants and weight norm bounds. We now bound \(\hat{R}_\mathcal{T}(\mathcal{J}_\gamma)\), when each tree \(t_j \in \mathcal{T}\) has a branching factor \(i.e.,\) maximum number of neighbors for any node) at most \(d\), and \(\mathcal{J}_\gamma\) maps each \((t, y)\) pair to \(\text{loss}_\gamma(-p(f_c(t; \Theta), y))\).

**Proposition 7.** The empirical Rademacher complexity of \(\mathcal{J}_\gamma\) with respect to \(\mathcal{T}\) is
\[
\hat{R}_\mathcal{T}(\mathcal{J}_\gamma) \leq \frac{4}{\gamma m} + \frac{24rB_\beta Z}{\gamma \sqrt{m}} \sqrt{3\log Q}, \quad \text{where}
\]
\[
Q = 2AB_\beta \sqrt{m} \max\{Z, M\sqrt{r} \max\{B_x B_1, B_x R\}\},
\]
\[
M = C_\phi \frac{(Cd)^{L}}{Cd - 1}, \quad Z = C_\phi B_1 B_x + C_\phi B_2 R, \quad \overline{R} \leq C_\rho C_\phi d \min\{b \sqrt{r}, B_1 B_x M\}.
\]

Note that we do not need to prespecify \(B_1\) and \(B_2\); we can simply take these values to be the spectral norm, respectively, of the learned weights \(W_1\) and \(W_2\). Before proceeding with the proof, we discuss some important implications of this result in the wake of Lemma 1 and Proposition 6.

**Comparison with RNN.** Table 1 summarizes the dependence of our generalization error on the embedding dimension \(r\), branching factor \(d\), depth \(L\) and sample size \(m\) for different \(C\) up to log factors (denoted by notation \(\mathcal{O}\)). Importantly, our bounds increase mostly linearly, at most with power 1.5, as a function of \(r, d\) and \(L\). Curiously, these dependencies are analogous to bounds for RNNs with sequence of length \(L\) (Chen et al. 2020a), when the spectral
norm of recurrent weights in RNN plays the role of $C$. The additional dependence on the branching factor $d$ is due to processing trees in contrast to sequences in RNNs.

**Comparison with VC-bounds for GNNs.** Scarselli et al. (2018) established generalization bounds based on VC-dimension. These bounds, for tanh and logistic sigmoid activations, depend with fourth-order on the number of hidden units $H$, and quadratically on $r$ and the maximum number of nodes $N$ in any input graph. Note that $N$ is at least $d$, and possibly much larger than $d$. Since $H = r$ in our setting, this amounts to having the VC-dimension scale as $\mathcal{O}(r^6N^2)$, and consequently, generalization error as $\tilde{O}(r^3N/\sqrt{m})$. Thus, our generalization bounds are significantly tighter even if $L = \mathcal{O}(r)$.

**Local permutation invariance.** Previous works focused on permutation-invariance at a global level (San-nai and Imaizumi (2019), Sokolic et al. (2017)). In contrast, GNNs are a composition of permutation-invariant transformations, applied to the neighborhoods of single nodes. We exploit local permutation-invariance via sum-decomposability. In the absence of local permutation-invariance, we would need to address the ordering of messages at each node and the complexity would depend on the worst case ordering.

**Extension to other GNN variants.** Note that we define $C_q$ and $C_p$ with respect to the aggregation function (e.g., unweighted sum or mean) that acts prior to transformation by $W_2$. This allows us to disentangle the role of shared weights from aggregation. Our analysis easily extends to the setting where messages are weighted (e.g., based on the edge embeddings) prior to aggregation. While we considered message passing with *mean field updates* (Dai et al. 2016), other updates, such as *embedded loopy belief propagation*, may be analyzed similarly in our framework.

**Analysis.** Our proof proceeds in multiple steps. We first quantify how changing shared weights affects the embedding of the root node of a fixed tree. To do so, we recursively bound the effect on each subtree of the root by the maximum effect across these subtrees. Since both the non-linear activation function and the permutation-invariant aggregation function are Lipschitz-continuous, and the feature vector at the root and the shared weights have bounded norm, the embedding at the root of the tree adapts (i.e., changes gradually in a controlled way) to the embeddings from the subtrees. We then quantify the effect of changing not only the shared weights but also changing the classifier parameters. Since the classifier parameters are chosen from a bounded norm family, we can bound the change in prediction probability. This allows us to use a covering number argument to approximate the predictions, and thus bound the empirical Rademacher complexity via Dudley’s entropy integral.

Fix the feature vectors for the computation tree of depth $L$ with degree of each internal node equal to $d$. Let the feature vector associated with the root (assumed to be at level $L$) of the tree be given by $x_L$. We denote the feature vector associated with node $j$ at level $\ell \in [L-1] \triangleq \{1, 2, \ldots, L-1\}$ by $x_{\ell,j}$. Denote the embedding produced by the subtree rooted at node $j$ on level $\ell \in [L-1]$ by $T_{\ell,j}(W_a, W_b)$ when $W_a$ and $W_b$ are the parameters of the model. Consider two sets of parameters $\{W_1, W_2\}$ and $\{W'_1, W'_2\}$. We will denote the embedding vector produced by the GNN after processing the entire tree by $T_L(W_1, W_2)$ as a shorthand for $T_{L,1}(W_1, W_2)$. Denote the set of subtrees of node with feature vector $x$ by $C(x)$.

We first quantify the changes in the root embedding when changing the shared weight parameters.

**Lemma 2.** The $l_2$-distance between embedding vectors produced by $(W_1, W_2)$ and $(W'_1, W'_2)$ after they process the tree from the leaf level to the root can be bounded recursively as

$$\Delta_L \triangleq || T_L(W_1, W_2) - T_L(W'_1, W'_2) ||_2 \leq C_b B_x ||(W_1 - W'_1)||_2 + C_d \max_{j \in C(x_L)} \Delta_{L-1,j} + C_b ||(W_2 - W'_2)R(W_1, W_2, x_L)||_2,$$

where

$$R(W_1, W_2, x_L) = \rho \left( \sum_{j \in C(x_L)} g(T_{L-1,j}(W_1, W_2)) \right)$$

is the permutation-invariant aggregation of the embeddings of the subtrees rooted at level $L-1$ under $(W_1, W_2)$.

We therefore proceed to bounding $||R(W_1, W_2, x_L)||_2$.

**Lemma 3.**

$$||R(W_1, W_2, x_L)||_2 \leq C_p C_g d \min \left\{ b \sqrt{r}, C_g B_1 B_2 \frac{(Cd)^L - 1}{Cd - 1} \right\}$$

Next, we quantify how changing the shared weights and classifier parameters affects the probability of outputting tree label 1.
The change in probability $\Lambda_L$ due to change in parameters from $(W_1, W_2, \beta)$ to $(W_1', W_2', \beta')$ is
\[
\Lambda_L = |\psi(\beta^T T_L(W_1, W_2)) - \psi(\beta'^T T_L(W_1', W_2'))| \\
\leq ||\beta - \beta'||_2 Z + B_\beta \Delta_L ,
\]
where $Z$ is an upper bound on $||T_L(W_1, W_2)||_2$. Moreover, we can bound $\Delta_L$ non-recursively:
\[
\Delta_L \leq MB_x ||W_1 - W_1'||_2 \\
+ M||R(W_1, W_2, x_L)||_2 ||W_2 - W_2'||_2 .
\]

Lemma 4 allows us to ensure that $\Lambda_L$ is small via a sufficiently large covering. Specifically, the change in probability $\Lambda_L$ can be bounded by $\epsilon$, using a covering of size $P$, where $P$ depends on $\epsilon$. Here, $\log P$ grows as $O(\log (1/\epsilon))$ for sufficiently small values of $\epsilon$. The stability of prediction translates into good generalization guarantees.

Lemma 5. The change in probability $\Lambda_L$ can be bounded by $\epsilon$ using a covering of size $P$, where $\log P$ is at most
\[
3\epsilon^2 \log \left(1 + \frac{6B_\beta \max\{Z, M\sqrt{\max\{B_x B_1, R B_2\}}\}}{\epsilon}\right).
\]

Moreover, if the cover parameter (radius)
\[
\epsilon < 6B_\beta \max\{Z, M\sqrt{\max\{B_x B_1, R B_2\}}\},
\]
then a covering of size $P$ such that $\log P$ is at most
\[
3\epsilon^2 \log \left(\frac{12B_\beta \max\{Z, M\sqrt{\max\{B_x B_1, R B_2\}}\}}{\epsilon}\right)
\]
suffices to ensure $\Lambda_L \leq \epsilon$. Here, $R \triangleq ||R(W_1, W_2, x_L)||_2$, and $Z, M$ are as defined in the statement of Proposition 4.

The remaining steps for Proposition 7 are straightforward and deferred to the Supplementary. We now outline some steps for understanding generalization of CPNGNNs.

5.3. Toward a generalization analysis for CPNGNNs

Two parts were integral to our analysis: (a) bounding complexity via local computation trees, and (b) the sum decomposition property of permutation-invariant functions. We now provide their counterparts for CPNGNNs.

Like before, we start with a vertex $v$, and unroll the $L$ neighborhood aggregations to obtain a computation tree of depth $L$, rooted at $v$. However, now we additionally label each edge in the computation tree with the respective ports of the nodes incident on the edge (enabled by consistent ordering). Thus, we may analyze a input port-numbered graph using its node-wise port-numbered trees.

Since permutation-invariance applies only to multisets of messages, but not port-numbered messages, we cannot express the aggregation in CPNGNNs as in Equation (2). Instead, we provide an injective function for aggregating a collection of port-numbered messages. The function takes a general sum-form that decouples the dependence on each message and its corresponding port number.

**Proposition 8.** Assume $\mathcal{X}$ is countable. There exist functions $f$ and $g$ such that $h((x_1, p_1), \ldots, (x_P, p_P)) = \sum_{i \in |P|} g(p_i) f(x_i)$ is unique for each collection of $(x_i, p_i)$ pairs, where $P \subset \mathbb{N}$, $X = \{x_1, x_2, \ldots, x_P\} \subset \mathcal{X}$ is a multiset of bounded size, and $p_i$ are all distinct numbers from $|P|$.

The result in Proposition 8 holds particular significance, since it is known (Hella et al., 2015) that port-numbered messages provide a strictly richer class than sets and multisets. The generalization bound for CPNGNN will be likely worse than the result in Proposition 7 since each port appears as an exponent in our generalized decomposition so the complexity of aggregation grows rapidly in the neighborhood size. We leave a detailed analysis for future work.

**Discussion**

We discuss some limitations and implications of this work.

**Tradeoffs between expressivity and generalization.** We introduced H-DCPN primarily to illustrate some ideas that could be used to improve the existing models. In particular, we suggested incorporating geometric information to circumvent some issues with DimeNet. While reasonable for settings where graphs represent actual spatial structures, such models are clearly inapplicable for graphs without any underlying geometric interpretation. Incorporating higher order information via a hypergraph or factor graph might be more natural in some scenarios. Beyond the nature of the graph, a good tradeoff can often be guided by application-specific considerations such as size of the training data, computation budget, and constraints (e.g., on training and inference time). Often, additional expressivity from a complex model might be offset by factors such as computational intractability and lack of generalization.

**Theory and practice.** Our results shed some light on the observed empirical success and limitations of message passing GNNs. For example, LU-GNNs have been observed to perform suboptimally on molecular graphs, where models that exploit the spatial structure (e.g., DimeNet) perform significantly better (Klicpera et al., 2020). Additional features can avoid some pitfalls of LU-GNNs and have improved performance elsewhere, e.g., Position-aware GNNs (You et al., 2019) and Structured Transformers (Ingraham et al., 2019). Likewise, Proposition 2 elucidates that choosing a good port-numbering can have a crucial impact on the performance of CPNGNNs. Despite addressing complex graphs instead of sequences, generalization bounds for GNNs and RNNs are comparable, so the empirical success of GNNs is not surprising.
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References


