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Computational capabilities of graph neural networks

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Abstract
In this paper, we will consider the approximation properties of a recently introduced neural network model called graph neural network (GNN), which can be used to process-structured data inputs, e.g., acyclic graphs, cyclic graphs, and directed or undirected graphs. This class of neural networks implements a function \( \tau(G, n) \in \mathbb{R}^m \) that maps a graph \( G \) and one of its nodes \( n \) onto an \( m \)-dimensional Euclidean space. We characterize the functions that can be approximated by GNNs, in probability, up to any prescribed degree of precision. This set contains the maps that satisfy a property called preservation of the unfolding equivalence, and includes most of the practically useful functions on graphs; the only known exception is when the input graph contains particular patterns of symmetries when unfolding equivalence may not be preserved. The result can be considered an extension of the universal approximation property established for the classic feedforward neural networks (FNNs). Some experimental examples are used to show the computational capabilities of the proposed model.

Keywords
Computational, capabilities, graph, neural, networks

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Abstract—In this paper, we will consider the approximation properties of a recently introduced neural network model called graph neural network (GNN), which can be used to process-structured data inputs, e.g., acyclic graphs, cyclic graphs, and directed or undirected graphs. This class of neural networks implements a function \( \tau(G, \eta) \in \mathbb{R}^m \) that maps a graph \( G \) and one of its nodes \( \eta \) onto an \( m \)-dimensional Euclidean space. We characterize the functions that can be approximated by GNNs, in probability, up to any prescribed degree of precision. This set contains the maps that satisfy a property called preservation of the unfolding equivalence, and includes most of the practically useful functions on graphs; the only known exception is when the input graph contains particular patterns of symmetries when unfolding equivalence may not be preserved. The result can be considered an extension of the universal approximation property established for the classic feedforward neural networks (FNNs). Some experimental examples are used to show the computational capabilities of the proposed model.

Index Terms—Approximation theory, graphical domains, graph neural networks (GNNs), universal approximators.

I. INTRODUCTION

In a large number of practical and engineering applications, the underlying data are often more conveniently represented in terms of graphs. In fact, a graph naturally represents a set of objects (nodes) and their relationships (edges). For example, in an image, it is natural to represent as nodes regions of the image that have similar intensity or color, and to represent the relationship among these regions by edges. This is often known as a region adjacency graph. As another example, it is convenient to model the individual web pages as nodes of a graph, and the hyperlink connections among the web pages as edges of the graph.

Traditionally, to process graph-structured input data, one first “squashes” the graph structure into a vector, and then uses neural network models that accept vectorial inputs, e.g., multilayer perceptrons and self-organizing maps, to process such resulting data [1]. Such “squashing” of the graph-structured input may lose most of the topological relationships among the nodes of the graph. An alternative approach is to preserve the topological relationships among the data items in a graph-structured input data, and to follow the graph structure in a node-by-node processing of the input data [2]–[4]. This general approach underpins a number of proposed neural network models, e.g., recursive neural networks (RNNs) [2], [4] and self-organizing maps for structured data [3]. The advantages of this approach include: the topological relationship among the data items are preserved, and taken into account in the data processing steps; and less data processing is required for each node. However, at least in the ways in which the RNN models or the self-organizing maps for structured data are formulated [3], [4], they can process limited types of graphs, e.g., acyclic and directed graphs. While RNNs or self-organizing maps for structured data can be extended to handle more general graph structures, e.g., cyclic graphs or undirected graphs or to adopt a more sophisticated processing scheme, e.g., taking into account the ancestors as well as descendants of a node in the processing, they tend to become relatively complicated.

Recently, the supervised approaches of this class of methods have been unified in a novel neural network model called graph neural networks (GNNs) [5]. GNNs can handle acyclic and cyclic graphs, directed and undirected graphs, and graphs with locally neighborhood dependency. A GNN handles such complexity by deploying two functions in the model: a transition function \( f \), which defines the relationship between the nodes of the graph, and an output function \( g \), which specifies an output for each node. By using these functions, a GNN implements a mapping \( \varphi(G, \eta) \in \mathbb{R}^m \), where \( G \) is a graph, \( \eta \) denotes a node in \( G \), and \( \mathbb{R}^m \) is the \( m \)-dimensional Euclidean space. It was shown empirically that GNNs can be used to model graph-structured data, and that trained GNNs can generalize to unforeseen data [6].

However, the approximation capabilities of this model have not been investigated yet and it has not been defined which functions on graphs the GNNs are able to realize. In other words, an interesting question arises: given a generic function \( \tau(G, \eta) \in \mathbb{R}^m \), can it be realized or approximated by a function \( \varphi \) implemented by a GNN model?

In this paper, we will seek to answer this question. In particular, we will show that under mild generic conditions, most of the practically useful functions on graphs can be approximated in probability by GNNs up to any prescribed degree of accuracy. Such a result can be considered an extension of the uni-
resolution approximation property that was proved for feedforward neural networks (FNNs) [7]–[10]. It also extends the universal approximation property of RNNs [11], [12].

The structure of this paper is as follows. After the introduction of some notations used in this paper as well as some preliminary definitions, Section II briefly presents the concept of a graph neural network model. A universal approximation theorem is shown in Section III and the proof of the theorem together with its auxiliary lemmas are given in the Appendix, while Section IV collects some experimental results on a number of examples used to illustrate the demonstrated property. Finally, conclusions are drawn in Section V.

II. GRAPH NEURAL NETWORKS

The GNN model was first introduced in [5] and [13]. In this section, we briefly introduce the model and the notation needed in this paper. Readers are referred to [5] for more details on the GNN model.

A. Notation

A graph $G$ is a pair $(N, E)$, where $N$ is a set of nodes and $E$ is a set of edges (or arcs) between nodes in $N$. Graphs are assumed to be undirected, i.e., for each arc $(u, v)$, the equality $(u, v) = (v, u)$ holds. The set $\mathcal{N}[n]$ collects the neighbors of $n$, i.e., the nodes connected to $n$ by an arc, while $\mathcal{E}[n]$ denotes the set of arcs having $n$ as a vertex. Nodes and edges may have labels, which are assumed to be real vectors. The labels attached to node $n$ and edge $(n, m)$ are represented by $l_n \in \mathbb{R}^N$ and $l_{(n,m)} \in \mathbb{R}^E$, respectively, and $l$ is the vector obtained by stacking together all the labels of the graph. The notation adopted for the labels follows a more general scheme. If $y$ is a vector that contains data from a graph and $S$ is a subset of its nodes (edges), then $y_S$ is the vector obtained by selecting from $y$ only the components related to the nodes (edges) in $S$. Thus, for example, $l_{\mathcal{N}[n]}$ is the vector containing the labels of all the neighbors of $n$.

Graphs may be either positional or nonpositional. The latter are those described so far, while positional graphs differ since a unique integer identifier is assigned to each neighbor of a node $n$ to indicate its logical position. Formally, for each node $n$ in a positional graph, there exists an injective function $\nu_n : \mathcal{N}[n] \rightarrow \{1, \ldots, |N|\}$, which assigns to each neighbor $u$ of $n$ a position $\nu_n(u)$. The position of the neighbor may be important in certain practical applications, e.g., object locations [12].

The graphical domain considered in this paper is the set $\mathcal{D}$ of pairs of a graph and a node, i.e., $\mathcal{D} = G \times N$ where $G$ is a set of graphs and $N$ is a subset of their nodes. We assume a supervised learning framework with the learning set $\mathcal{L} = \{(G_i, n_{ij}, t_{ij}) | G_i = (\mathcal{N}_i, \mathcal{E}_i) \in G, n_{ij} \in \mathcal{N}_i, t_{ij} \in \mathbb{R}^m, 1 \leq i \leq p, 1 \leq j \leq q_i\}$, where $n_{ij}$ denotes the $j$th node in the graph $G_i$ and $t_{ij}$ is the desired target associated to $n_{ij}$. Finally, $p \leq |G|$ and $q_i \leq |N_i|$. Interestingly, a set of graphs can be seen as one large graph that contains disconnected components. Hence, one can refer to a learning set as the pair $\mathcal{L} = (G, T)$ where $G = (N, E)$ is a graph and $T$ is a set of pairs $\{(n_i, t_i) | n_i \in N, t_i \in \mathbb{R}^m, 1 \leq i \leq q\}$.

B. The Model

The intuitive idea underlying the proposed approach is that nodes in a graph represent objects or concepts, and edges represent their relationships. Each concept is naturally defined by its features and the related concepts. Thus, we can attach a state $x_n \in \mathbb{R}^n$ to each node $n$ that is based on the information contained in the neighborhood of $n$ (see Fig. 1). The variable $x_n$ contains a representation of the concept embodied in node $n$ and can be used to produce an output $o_n \in \mathbb{R}^m$, i.e., a decision about the concept.

Let $f_w$ be a parametric function, called local transition function, that expresses the dependence of a node $n$ on its neighborhood and let $g_u$ be the local output function that describes how the output is produced. Then, $x_n$ and $o_n$ are defined as follows:

$$x_n = f_w(l_n, l_{\mathcal{E}[n]}, x_{\mathcal{N}[n]}, l_{\mathcal{E}[n]})$$
$$o_n = g_u(x_n, l_n)$$

where $l_n, l_{\mathcal{E}[n]}, x_{\mathcal{N}[n]}$, and $l_{\mathcal{E}[n]}$ are the label of $n$, the labels of its edges, the states, and the labels of the nodes in the neighborhood of $n$, respectively. In GNNs, the transition and the output functions are implemented by multilayer FNNs [5].

Remark 1: For the sake of simplicity, only the case of undirected graphs is studied, but the results can be easily extended to directed graphs and even to graphs with mixed directed and undirected arcs. In fact, with minor modifications, GNNs can process general types of graphs. For example, when dealing with directed graphs, the function $f$ must also accept as an input the direction of each arc, coded, for instance, as an additional parameter $d_k$ for each arc $k \in \mathcal{E}[n]$ such that $d_k = 1$, if $k$ is directed towards $n$ and $d_k = 0$, if $k$ comes from $n$. Moreover, when different kinds of edges coexist in the same data set, the label should be designed to distinguish between them.

Note that (1) makes it possible to process both positional and nonpositional graphs. For positional graphs, $f_w$ needs to receive as additional input the positions of the neighbors. In practice, this can be easily achieved provided that the information contained in $x_{\mathcal{N}[n]}, l_{\mathcal{E}[n]},$ and $l_{\mathcal{E}[n]}$ is sorted according to neighbor positions and is properly padded with special null values in positions corresponding to nonexisting neighbors. For example, $x_{\mathcal{N}[n]} = [x_{y_1}, \ldots, x_{y_{M}}]$, where $y_{i} = x_{n_{ij}}$, if $u$ is the $i$th neighbor of $n$ ($\nu_n(u) = i$), and $y_{i} = x_{0}$, for some predefined null state $x_{0}$, if there is no $i$th neighbor, and $M = \max_{u \in N} \nu_n(u)$ is the maximum number of neighbors of the node $n$.

For nonpositional graphs, on the contrary, it is useful to replace function $f_w$ of (1) with

$$x_n = \sum_{n \in \mathcal{N}[n]} h_w(l_n, l_{(n,u)}, x_n, l_n), \quad n \in N$$

where $h_w$ is a parametric function. In the following, (2) is referred to as the nonpositional form, while (1) is called the positional form. It is worth mentioning that the same structure of (2) can also be applied to positional graphs provided that the parameters of $h_w$ are extended to include a description of the position $\nu_n(u)$ of each neighbor $u$ of $n$. Formally, positional graphs can
be processed when $h_{\mathbf{w}}$ takes the position of the neighbors as input, i.e.,

$$
x_n = \sum_{u \in \pi(n)} h_{\mathbf{w}}(l_n, l_{t(n u)}, \mathbf{x}_u, l_u, \nu_n(u)), \quad n \in \mathcal{N}. \quad (3)
$$

In practical implementations of GNNs and RNNs, the form defined in (1) is preferred to (3). However, (3) is a special case of (1) and will be particularly useful for proving our results.

Let $\mathbf{x}, \mathbf{o}, \mathbf{l}$, and $\mathbf{l}_N$ be the vectors constructed by stacking all the states, all the outputs, and all the node labels, respectively. Then, (1) can be written in a vectorial form as follows:

$$
\mathbf{x} = F_{\mathbf{w}}(\mathbf{x}, \mathbf{l})
$$

$$
\mathbf{o} = G_{\mathbf{w}}(\mathbf{x}, \mathbf{l}_N)
$$

(4)

where $F_{\mathbf{w}}$ and $G_{\mathbf{w}}$ are the composition of $|\mathcal{N}|$ instances of $f_{\mathbf{w}}$ and $g_{\mathbf{w}}$, respectively. In GNNs, $F_{\mathbf{w}}$ is called the global transition function while $G_{\mathbf{w}}$ is the global output function. Note that in order to ensure that $\mathbf{x}$ is correctly defined, (4) must have a unique solution. The Banach fixed point theorem [14] provides a sufficient condition for the existence and uniqueness of the solution of such a system of equations. According to Banach’s theorem [14], (4) has a unique solution provided that $F_{\mathbf{w}}$ is a contraction map with respect to the state, i.e., there exists a real number $\mu$, $0 \leq \mu < 1$, such that $\|F_{\mathbf{w}}(\mathbf{x}, \mathbf{l}) - F_{\mathbf{w}}(\mathbf{y}, \mathbf{l})\| \leq \mu \|\mathbf{x} - \mathbf{y}\|$ holds for any $\mathbf{x}, \mathbf{y}$, where $\| \cdot \|$ is any vectorial norm. In GNNs, $f_{\mathbf{w}}$ is designed so that $F_{\mathbf{w}}$ is a contraction map.

Thus, (1) provides a method to realize a function $\varphi$ that returns an output $\varphi(\mathbf{G}, n) = \mathbf{o}_n$ for each graph $\mathbf{G}$ and each node $n$.

Definition 1—Harmolodic Functions: Let $F_{\mathbf{w}}$ be a contraction map with respect to (w.r.t.) $\mathbf{x}$. Then, any function $\varphi : \mathcal{D} \rightarrow \mathbb{R}^n$ generated by $\varphi(\mathbf{G}, n) = \mathbf{o}_n$ is referred to as a harmolodic function. The class of harmolodic functions on $\mathcal{D}$ will be denoted by $\mathcal{H}(\mathcal{D})$.

Banach’s fixed point theorem suggests also the following classic iterative scheme for computing the value of the stable state:

$$
x(t + 1) = F_{\mathbf{w}}(\mathbf{x}(t), \mathbf{l})
$$

(5)

where $\mathbf{x}(t)$ denotes the $t$th iteration of $\mathbf{x}$. This equation converges exponentially fast to the solution of (4) for any initial value $\mathbf{x}(0)$. In fact, (5) implements the Jacobi iterative method for the solution of nonlinear systems [15].

Learning phase in GNN model aims at adapting the parameter set $\mathbf{w}$ such that $\varphi_{\mathbf{w}}$ approximates the learning set $\mathcal{L} = \{(\mathbf{G}_i, \mathbf{n}_{i,j}, \mathbf{t}_{i,j})\}$, $G_i = (\mathbf{N}_i, \mathbf{E}_i) \in \mathcal{G}$, $n_{i,j} \in \mathbf{N}_i$, $t_{i,j} \in \mathbb{R}^n$, $1 \leq i \leq p, 1 \leq j \leq q_i$. This learning task can be posed as the minimization of a quadratic error function

$$
e_{\mathbf{w}} = \sum_{i=1}^{p} \sum_{j=1}^{q_i} (t_{i,j} - \varphi_{\mathbf{w}}(\mathbf{G}_i, \mathbf{n}_{i,j}))^2.
$$

(6)

The name “harmolodic function” is inspired by the harmolodic philosophy that is behind jazz music of saxophonist Ornette Coleman.
In GNNs, the minimization is achieved by a new learning algorithm [5] that combines backpropagation-through-structure algorithm [4], which is used in RNNs, with the Almeida–Pineda algorithm [16], [17]. In order to ensure that the global transition function $F_w$ remains a contraction map during learning phase, a penalty term $L(||(\partial F_w)/(\partial x)||)$ may be added to the error function (6), where $L(y)$ is $(y - \mu)^2$ if $y > \mu$ and 0 otherwise, and the parameter $\mu \in (0, 1)$ defines the desired contraction constant of $F_w$.

### III. Computational Capabilities of GNNs

FNNs have been proved to be universal approximators [7]–[9] for functions having Euclidean domain and codomain, i.e., they can approximate any map $\tau: \mathbb{R}^m \rightarrow \mathbb{R}^b$. Several versions of the result have been proposed, which adopt different classes of functions, different measures of the approximation, and different network architectures [10]. Recently, also RNNs have been shown to approximate in probability any function on trees up to any degree of precision [11], [12]. More precisely, it has been proved that for any probability measure $P$, any reals $\varepsilon, \lambda$, and any real function $\tau$ defined on trees, there exists a function $\varphi$ implemented by a RNN such that $P(||\tau(T) - \varphi(T)|| \geq \varepsilon) \leq 1 - \lambda$ holds. In the following, the approximation capabilities of GNN model are investigated. The analysis presented here concerns the undirected graphs the labels of which are expressed as a vector of reals, i.e., graphs where node labels belong to $\mathbb{R}^{|V|}$ and edge labels belong to $\mathbb{R}^{|E|}$. Both positional and nonpositional GNNs are studied.

In order to discuss the results, some new concepts will be introduced. First, we will define an equivalence $\sim$ on nodes, called unfolding equivalence, that aims to specify which concepts, among those represented by a graph, can or cannot be distinguished using only the information contained in the graph. Then, we will demonstrate that the class of functions that can be approximated by GNNs consists of maps $\tau: \mathcal{D} \rightarrow \mathbb{R}^m$, which are generic except for the fact that $\tau$ is constrained to produce the same output on nodes that are unfolding equivalent i.e., $n \sim u$ implies $\tau(G, n) = \tau(G, u)$. The equivalence $\sim$ will be formally defined using another concept, the unfolding tree, that is defined in the following.

An unfolding tree $T^d_n$ is the graph obtained by unfolding $G$ up to the depth $d$, using the node $n$ as the starting point (see Fig. 2).

**Definition 2—Unfolding Tree:** An unfolding tree $T^d_n$ having depth $d$ of a node $n$ is recursively defined as

$$
T^d_n = \begin{cases} 
\text{Tree}(l_n), & \text{if } d = 1 \\
\text{Tree}(l_n, l_{\varnothing \cup \{n\}}; T^{|d|-1}_{\varnothing \cup \{n\}}), & \text{if } d > 1.
\end{cases}
$$

Here, $T^{|d|-1}_{\varnothing \cup \{n\}} = [T^{d-1}_{u_1}, T^{d-1}_{u_2}, \ldots]$ is the vector containing the unfolding trees having depth $d - 1$ of the neighbors $u_1, u_2, \ldots$ of $n$. The operator Tree constructs a tree from the label of the root, the labels of the edges entering into the root, and a set of subtrees. Moreover, the possibly infinite tree $T_n = \lim_{d \rightarrow \infty} T^d_n$ that can be constructed by merging all the unfolding trees $T^d_n$ for any $d$ will simply be called the unfolding tree of $n$.

An example of construction of the unfolding tree is shown in Fig. 2. Unfolding trees naturally induce an equivalence relationship on the nodes of $G$.

**Definition 3—Unfolding Equivalence:** Let $G = (N, E)$ be an undirected graph. The nodes $n, u \in N$ are said to be unfolding equivalent, $n \sim u$, if $T_n = T_u$.

For example, Fig. 2 shows a graph with two unfolding nonequivalent nodes, two unfolding equivalent nodes, and their respective unfolding trees of depth 3. In this particular example, nonequivalent nodes can be immediately distinguished at the first level of the trees, since they have a different number of children.

Functions that do not distinguish nodes which are unfolding equivalent are said to preserve the unfolding equivalence.

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2For the sake of simplicity, only the case of undirected graphs is studied. The results can be easily extended to directed graphs.

3If no subtree is given, as in Tree($l_n$), the constructed tree contains only one node.

---

**Fig. 2.** Graph and four unfolding trees of depth 3. Dashed lines specify the correspondence between a node and its unfolding tree. The two nodes with label $b$ are not unfolding equivalent because their unfolding trees are different, whereas the two nodes with label $a$ are unfolding equivalent.
**Definition 4—Functions Preserving the Unfolding Equivalence:** A function \( l : \mathcal{D} \rightarrow \mathbb{R}^m \) is said to preserve the unfolding equivalence on \( \mathcal{D} \), if
\[
\exists \ u \implies l(\mathcal{G}, n) = l(\mathcal{G}, u).
\]
The class of functions that preserves the unfolding equivalence on \( \mathcal{D} \) is denoted by \( \mathcal{F}(\mathcal{D}) \).

For example, let us apply a given function \( l : \mathcal{D} \rightarrow \mathbb{R}^m \) to the graph in Fig. 2. If \( l \) preserves the unfolding equivalence, then \( l \) is constrained to produce the same output for the two nodes \( n_1 \) and \( n_2 \) having label \( a \), i.e., \( l(\mathcal{G}, n_1) = l(\mathcal{G}, n_2) \).

**Remark 2:** The exact meaning of the given definitions is slightly different according to whether positional or nonpositional graphs are to be considered. If the graphs are positional, the unfolding trees should take into account also the original neighbors’ positions. Moreover, equation \( T_n = T_u \) in Definition 3 uses the equality embedded in positional trees. For nonpositional graphs, the unfolding trees and the equality are both nonpositional.

The following theorem states that functions preserving the unfolding equivalence compute the outputs at a node \( \mathcal{G} \) considering only the information contained in the unfolding tree \( T_n \).

**Theorem 1—Functions of Unfolding Trees:** A function \( l \) belongs to \( \mathcal{F}(\mathcal{D}) \) if and only if there exists a function \( k \) defined on trees such that \( l(\mathcal{G}, n) = k(T_n) \) for any node \( n \) of the domain \( \mathcal{D} \).

The proofs of all theorems and corollaries presented in this section have been moved to the Appendix to improve paper’s readability.

The following corollary, which is an immediate consequence of Theorem 1, suggests that \( \mathcal{F}(\mathcal{D}) \) is a large class of functions. It can be applied, for example, to all the real-life domains where the labels contain real numbers.

**Corollary 1—Graphs With Distinct Labels:** Let \( \mathcal{G} \) be the set of the graphs of \( \mathcal{D} \) and assume that all the nodes have distinct labels, i.e., \( \forall n \neq u \implies l_n \neq l_u \) for any nodes \( n, u \) of \( \mathcal{G} \). Then, any function defined on \( \mathcal{D} \) preserves the unfolding equivalence.

In the following, we assume that \( \mathcal{D} \) is equipped with a probability measure \( \mathcal{P} \) and an integral operator is defined on the functions from \( \mathcal{D} \) onto \( \mathbb{R}^m \). In order to clarify how these concepts can be formally defined, note that a graph is specified by its structure and its labels. Since node labels and the possible structures of a graph are enumerable, there exists an enumerable partition \( \mathcal{D}_1, \mathcal{D}_2, \ldots \) of the domain \( \mathcal{D} \) such that \( \mathcal{D}_i = \{ \mathcal{G} \times \{ i \} \} \) and each set \( \mathcal{G}_i \) contains graphs having the same structure. For each \( i \), a graph \( \mathcal{G} \in \mathcal{G}_i \) is completely defined by the vector \( \mathbf{l}_G \) formed by stacking all its labels and the set \( \mathcal{G}_i = \{ \mathbf{l}_G | \mathcal{G} \in \mathcal{G}_i \} \), obtained by collecting all those vectors, is a subset of an Euclidean space, i.e., \( \mathcal{G}_i \subseteq \mathbb{R}^n \). Thus, any measure \( \mathcal{P} \) on \( \mathcal{D} \), when restricted to \( \mathcal{G}_i \), is equivalent to a measure \( m_i \) defined on the linear space \( \mathbb{R}^n \). As a consequence, \( \mathcal{P} \) can be formally defined, for each \( \mathcal{D}_i \subseteq \mathcal{D} \), as
\[
\mathcal{P}(\mathcal{D}) = \sum_i \alpha_i \cdot \mathcal{P}(\mathcal{D} \cap \mathcal{D}_i) = \sum_i \alpha_i \cdot m_i(\mathcal{G}_i)
\]
where \( \mathcal{G}_i \) is specified by the equality \( \mathcal{G}_i \times \{ i \} = \mathcal{D}_i \cap \mathcal{D}_i \) and the \( \alpha_i \) are positive numbers such that \( \sum_i \alpha_i = 1 \).

Moreover, we will define the integral of a function \( l \) on \( \mathcal{D} \) as
\[
\int_{\mathcal{D}} l(\mathcal{G}, n) d\mathcal{G} dn = \sum_i \int_{\mathcal{D}_i} l(\mathcal{G}, n) d\mathcal{G},
\]
where each \( \int_{\mathcal{D}_i} l(\mathcal{G}, n) d\mathcal{G} \) is computed using the Lebesgue measure theory [18].

The set \( \mathcal{F}(\mathcal{D}) \) plays an important role in our analysis. In fact, it will be proved that any measurable function \( \tau \in \mathcal{F}(\mathcal{D}) \) can be approximated by a GNN in probability. Moreover, the converse holds: all the functions implemented by a GNN preserve the unfolding equivalence. First, the result is proved for positional GNNs.

**Theorem 2—Approximation by Positional GNNs:** Let \( \mathcal{D} \) be a domain that contains positional graphs. For any measurable function \( \tau \in \mathcal{F}(\mathcal{D}) \) preserving the unfolding equivalence, any norm \( ||\cdot|| \) on \( \mathbb{R}^m \), any probability measure \( P \) on \( \mathcal{D} \), and any reals \( \varepsilon_1, \varepsilon_2 > 0 \), there exist two continuously differentiable functions \( f \) and \( g \) such that, for the GNN defined by
\[
x_n = f(l_n, l_{\alpha_0}[n], l_{\beta_1}[n], l_{\gamma_2}[n])
\]
\[
o_n = g(x_n, l_n), \quad n \in \mathcal{N}
\]
the global transition function \( F \) is a contraction map with a contracting constant \( \mu \), the state dimension is \( s = 1 \), the stable state is uniformly bounded, and the corresponding harmonic function defined by \( \varphi(\mathcal{G}, n) \) satisfies the condition
\[
P( ||\tau(\mathcal{G}, n) - \varphi(\mathcal{G}, n)|| \geq \varepsilon ) \leq 1 - \lambda.
\]

Commonly used FNNs are universal approximators [7]–[10] and, obviously, they can also approximate the functions \( f \) and \( g \) of Theorem 2. However, to perfectly simulate the GNN dynamics, we must consider a restricted class of network architectures that can approximate any function and its derivatives at the same time.

**Definition 5—FNNs Suitable to Implement GNNs:** A class \( \mathcal{Q} \) of FNNs is said to be suitable to implement GNNs, if for any positive integers \( a, b \), any continuously differentiable function \( l \in \mathbb{R}^a \rightarrow \mathbb{R}^b \) with a bounded support, and any real numbers \( \varepsilon_1 > 0, \varepsilon_2 > 0 \), there exist a function \( k_{\mathcal{G}} \) implemented by a network in \( \mathcal{Q} \) and a set of parameters \( \mathbf{w} \), such that \( ||l(x) - k_{\mathcal{G}}(\mathbf{w})|| \leq \varepsilon_1 \) and \( ||(\partial l(\mathbf{x}))/\partial \mathbf{x} - (\partial k_{\mathcal{G}}(\mathbf{w})/\partial \mathbf{x})|| \leq \varepsilon_2 \) hold for any \( x \in \mathbb{R}^b \).

In [19], it is proved that the class of three-layered neural networks with activation function \( \sigma \) in the hidden neurons and a linear activation function in the output neurons can approximate any function \( l \) and its derivatives on \( \mathbb{R}^a \), provided that there exists a linear combination \( k \) of scaled shifted rotations of \( \sigma \) such that \((\partial l(\mathbf{x}))/\partial \mathbf{x})\) is a square integrable function of uniformly locally bounded variation. It can be easily proved that three-layered neural networks using common differentiable activation functions, e.g., \( \sigma(x) = 1/(1+e^{-x}), \sigma(x) = (1-e^{-x})/(1+e^{-x}), \)

\footnote{It is worth mentioning that also the converse holds: in fact, any measure on \( \mathcal{D} \) can be represented as in (7) where \( \alpha_i = \mathcal{P}(\mathcal{D}_i) \).}

\footnote{This is stated in Theorem 4.}

\footnote{Notice that since all the norms on the Euclidean space are equivalent, the definition is not affected by considered norm \( ||\cdot|| \).}
or $\sigma(x) = \text{atan}(x)$, satisfy the above property and are suitable to implement GNNs.

Corollary 2 proves that $f$ and $g$ can be replaced by networks suitable to implement GNNs without losing the property stated in Theorem 2.

Corollary 2—Connectionist Implementation of Positional GNNs: Let us assume that the hypotheses of Theorem 2 holds, that the nodes of the graph in $\mathcal{D}$ have a bounded number of neighbors, and that $\mathcal{Q}$ is a class of networks suitable to implement GNNs. Then, there exist a parameter set $\mathcal{W}$ and two functions $f_{\mathcal{W}}$ (transition function) and $g_{\mathcal{W}}$ (output function) implemented by networks in $\mathcal{Q}$ such that the thesis of Theorem 2 is true.

The hypothesis on the boundedness of the number of neighbors is needed because $f$, without such a constraint, can have any number of inputs, whereas an FNN can only have a predefined number of inputs. It is worth mentioning that the hypothesis could be removed by adopting the form defined in (3) in place of the one expressed in (1). In this case, we can prove that $h_{\mathcal{W}}$ can be implemented by a multilayered FNN.\(^7\)

The definition of network class suitable to implement GNNs can be weakened, if we admit that the GNN state $x(t)$ remains bounded during the computation of the fixed point. Such an assumption is reasonable in a real application and can be guaranteed by using a fixed initial state, e.g., $x(0) = 0$. In fact, the proofs of Theorem 2 and Corollary 2 demonstrate that if the states are bounded, $g$ and $f$ have to be approximated only on compact subsets of their domains, instead of the whole domains. With such a simplification, the universal approximation literature provides several other results about the approximation of a function along with its derivatives [10], [20], [21]. For example, in [10], it is proved that three-layered networks with nonpolynomial analytic activation functions can implement any polynomial on compact sets. Since polynomials are dense in continuous functions also with respect to derivatives, three-layered networks with nonpolynomial analytic activations are suitable to implement GNNs.

The transition function defined in (2) is less general than the one in (1). For this reason, one may wonder whether nonpositional GNN based on (2) has narrower approximation capabilities than the GNN of (1). Theorem 3 states that both models have the same computational power.

Theorem 3—Approximation by Nonpositional GNNs: Let $\mathcal{D}$ be a domain that contains nonpositional graphs. For any measurable function $\tau \in \mathcal{F}(\mathcal{D})$ that preserves the unfolding equivalence, any norm $\|\cdot\|$ on $\mathcal{F}^{\mathcal{D}}$, any probability measure $P$ on $\mathcal{D}$, and any reals $\varepsilon,\mu,\lambda$, where $\varepsilon > 0, 0 < \lambda < 1$, and $0 < \mu < 1$, there exist two continuously differentiable functions $h$ and $g$ such that, for the GNN defined by

$$x_n = \sum_{u \in \mathcal{W}[n]} h(l_n, l_{(n,u)}, x_u, l_u),$$

$$o_n = g(x_n, l_n),$$

the global transition function $F$ is a contraction map with contraction constant $\mu$, the state dimension is $s = 1$, the stable state

is uniformly bounded, and the corresponding hamologic function defined by $\varphi(\mathcal{G}, n) = \alpha_n$ satisfies the condition

$$P(\|\tau(\mathcal{G}, n) - \varphi(\mathcal{G}, n)\| \geq \varepsilon) \leq 1 - \lambda.$$  

\[\Box\]

In addition, we have the following corollary.

Corollary 3—Connectionist Implementation of Nonpositional GNNs: Let us assume that the hypothesis of Theorem 2 holds and $\mathcal{Q}$ is a class of network suitable to implement GNNs. Then, there exists a parameter set $\mathcal{W}$ and two functions $f_{\mathcal{W}}$ (transition function) and $g_{\mathcal{W}}$ (output function) implemented by networks in $\mathcal{Q}$, such that the thesis of Theorem 3 holds.

Finally, the following theorem proves that a GNN can implement only functions that preserve the unfolding equivalence. Hence, the functions realizable by the proposed model are exactly those described in Theorems 2 and 3 respectively.

Theorem 4—$\mathcal{H}(\mathcal{D}) \subseteq \mathcal{F}(\mathcal{D})$: Let $\varphi$ be the function implemented by a GNN. If the GNN is positional, then $\varphi$ preserves the unfolding equivalence on positional graphs, while if the GNN is nonpositional, then $\varphi$ preserves the unfolding equivalence on nonpositional graphs.

Theorems 2–4 can be provided with intuitive explanations. GNNs use a local computational framework, i.e., the processing consists of “small jobs” operated on each single node. There is no global activity and two “small jobs” can communicate only if the corresponding nodes are neighbors. The output $o_n = \varphi(\mathcal{G}, n_1)$ of node $n_1$ depends only on the information contained in its neighbors, and recursively, in all the connected nodes. In other words, $\varphi(\mathcal{G}, n_1)$ is a function of the unfolding tree $T_n$, which, according to Theorem 1, implies that $\varphi$ preserves the unfolding equivalence.

What the GNNs cannot do is described by the following two cases. Theorems 2–4 ensure that GNNs do not suffer from other limitations except for those mentioned here. If two nodes $n_1$ and $n_2$ are “completely symmetric” (recursively equivalent) and cannot be distinguished on the basis of information contained in the connected nodes, then a GNN will produce the same output for those nodes. In the example depicted in Fig. 3, every node has the same label $\mathbf{a}$ and graphs $\mathbf{A}$ and $\mathbf{B}$ are regular, i.e., each node has exactly the same number of edges. Thus, all the nodes of graph $\mathbf{A}$ (graph $\mathbf{B}$) are “symmetric” and will have the same output, i.e., $\varphi(\mathcal{G}, n_1) = \varphi(\mathcal{G}, n_2)$ if both $n_1$ and $n_2$ belong to $\mathbf{A}$ (or both $n_1$ and $n_2$ belong to $\mathbf{B}$). Moreover, GNNs cannot compute general functions on disconnected graphs. If $\mathbf{G}$ is composed of disconnected graphs, the information contained in a subgraph cannot influence the output of a node, which is not reachable from that subgraph. For example, if $n$ is a node of graph $\mathbf{A}$ in Fig. 3, then $\varphi(\mathcal{G}, n)$ cannot be influenced by $\mathbf{B}$, e.g., $\varphi(\mathcal{G}, n)$ cannot count the number of edges of graph $\mathbf{B}$.

It is worth mentioning that in common graph theory all the nodes of a graph are considered different entities. On the contrary, in GNNs, two nodes are equal unless the available information suggests otherwise. Such a property is not necessarily a limitation, for two different reasons. 1) It may capture an intuitive idea of the information contained in a graph. In fact, the unfolding tree $T_n$ contains all the data that can be reached by surfing the graph from $n$. If we assume that the graph defines
all available information about the domain objects and their relationships, then it is reasonable to think that $T_n$ describes all our knowledge about $n$. In addition, the definition of function preserving the unfolding equivalence captures all the reasonable functions on a graphical domain. 2) If the considered application requires that some nodes are distinct, then the goal can be practically obtained by inserting into the data set the appropriate information. Let us consider again the examples depicted in Fig. 3. If $n$ is a node of graph $A$ and $\varphi(G, n)$ should depend on the information contained in $B$, then there must be some hidden relationship between the object represented by $n$ and the objects represented by the nodes of $B$. By explicitly representing this relationship with appropriate edges, $A$ and $B$ become a connected graph and the GNN model can produce the desired function. Similarly, if some nodes are unfolding equivalent, but $\varphi$ should produce different outputs, then there exists some information that distinguishes among the equivalent nodes and is not represented in the graph. Including such information into the labels (or, in general, into the graph) will solve the problem.

The presented theory also extends all the currently known results on approximation capabilities of RNNs. In fact, it has been proved that RNNs can approximate in probability any function on trees [11], [12]. On the other hand, when processing a tree, an RNN acts as a GNN where the neighborhood of a node only contains its children, i.e., the father is not included (see [5] for a more detailed comparison). It can be easily observed that under this definition of neighborhood, any function on trees that satisfies the unfolding equivalence and Theorems 2 and 3 reproduces those presented in [11] and [12].

Moreover, the concept of unfolding tree has been introduced in [22], where it is used to implement a procedure that allows to process cyclic graphs by RNNs. Such an approach extracts, from the input graph, the unfolding trees of all the nodes: then, those trees are processed by an RNN. It is proved that such a method allows to approximate in probability any function on cyclic graphs with distinct labels. Such a result can now be deduced by using Corollary 1.

The intuition delivered by these results is that a wide class of maps on graphs is implementable by a diffusion mechanism based on a transition function and an output function. Here, we also proved that the global transition function can be restricted to be a contraction map. Such result is crucial for the applications of the GNN model to practical problems using generic forms of graphs (because the functions that cannot be approximated by the proposed GNNs are pathological in nature). These universal approximation results thus recommend the GNNs as suitable practical models for processing of most classes of graph-structured input data, e.g., cyclic or acyclic and directed or undirected.

IV. EXPERIMENTAL RESULTS

This section presents four experiments designed to demonstrate peculiarities of the GNN model that can be observed in its practical applications and are related to its approximation properties. In the first example, it is shown that by adding noise to the node labels of a data set, we can transform a function that does not preserve the unfolding equivalence to a function that preserves the unfolding equivalence. The experiment demonstrates that such a function, which in theory is approximable by a GNN, can be, even if only partially, learned. The other three experiments face problems with different levels of difficulties. Here, the difficulty depends on the complexity of the coding that must be stored in the states. Even if in theory a GNN can realize most of the functions on graphs, in practice, the learnability may be limited by the architecture adopted for the transition function $f$ and the output function $g$, and by the presence of local minima in the error function. We will observe that the accuracy of the learned function decreases while the coding becomes more complex. Other experiments, whose goal is to assess the performance and the properties of the GNN model on wider and real-life applications, can be found in [5], [6], and [23]–[27]. The following facts hold for each experiment, unless otherwise specified. The functions involved in the GNN model $g_{\varphi}, h_{\varphi}$ were implemented by three-layered (one hidden layer) FNNs with sigmoidal activation functions. The presented results were averaged on five different runs. In each run, the data set was a collection of random graphs constructed by the following procedure: each pair of nodes was connected with a certain probability $\delta$: the resulting graph was checked to verify whether it was connected and, finally, if it was not, random edges were inserted until the condition was satisfied. The data set was split into a training set, a validation set, and a test set and the validation set was used to avoid possible issues with overfitting. In every trial, the training procedure performed at most 5000 epochs and every 20 epochs the GNN was evaluated on the validation set. The GNN that achieved the lowest error on the validation set was considered the best model, which was then applied to the test set.

The performance of the model is measured by the accuracy in classification problems (when $t_{i,j}$ can take only the values $-1$ or $1$) and by the relative error in regression problems (when $t_{i,j}$ may be any real number). More precisely, in classification problems, a pattern is considered correctly classified if $\varphi_{\varphi}(G, n_{i,j}) > 0$ and $t_{i,j} = 1$ or if $\varphi_{\varphi}(G, n_{i,j}) < 0$ and $t_{i,j} = -1$. Thus, the accuracy is defined as the percentage of patterns correctly classified by the GNN on the test set. On the other hand, in regression problems, the relative error on a pattern is given by $|t_{i,j} - \varphi_{\varphi}(G, n_{i,j})|/t_{i,j}$.

A. Half-Hot on Uniform Graphs

This problem consists of learning by examples a relation $\tau$ that, given a graph $G_i$, returns $\tau(G_i, n_{i,j}) = 1$ for half of the
nodes of $G_i$ and $\tau(G_i, n_{i,j}) = -1$ for the other half. Fig. 3(b) shows an example of $\tau$.

The data set contained connected regular graphs, i.e., graphs where each node has the same number of connections. As discussed in Section III, if all the labels of the nodes are equal and the graphs are regular, then $\tau$ does not preserve the unfolding equivalence and cannot be realized by a GNN. In practice, when a GNN is applied on a regular graph, it produces the same output on each node. However, the labels can be made distinct by extending them with a random component. With this extension, according to Corollary 1, $\tau$ can be realized by a GNN.

The purpose of this experiment is to check the above theoretical results and to verify whether the extension of the labels with random vectors can actually increase the computational power of GNNs. In this experiment, 300 uniform graphs with random labels and random connectivity were equally subdivided into training set, validation set, and test set. Each graph $G_i$ was generated by the following three-step procedure.

Step 1) An even random number of nodes $d$ in the range $[4, 10]$ and a random integer number of links $c$ in the range $[3, 20]$ were generated. The numbers are produced by uniform probability distributions.

Step 2) A random undirected regular graph with $d$ nodes and $c$ connections for each node was generated. The graph was produced by recursively inserting random edges between nodes that did not reach the maximal number of connections. The construction procedure may be stopped either because a regular graph was obtained or because a configuration was reached where no more edges could be inserted. The construction procedure was repeated until a regular graph was generated.

Step 3) A random node label $I_n$ was attached to each node $n$. Each label is a five–dimensional vector containing integers in the range $[0, 5]$.

Note that given a graph $G_i = (N_i, E_i)$, there are many different functions $\tau$ solving the task. However, for our purposes, no particular one is preferable. Such a concept can be expressed applying the following error function:

$$
\epsilon_w = \sum_{j, k: j \neq k} \varphi_w(G_i, n_{i,j}) \cdot \varphi_w(G_i, n_{i,k})
$$

to each graph $G_i$. It can be easily proved that if $G_i$ contains an even number of nodes and $\varphi_w$ produces values in the range $[-1, 1]$, then $\epsilon_w$ reaches a minima when for half of the nodes $\varphi_w(G_i, n_{i,j}) = -1$ and $\varphi_w(G_i, n_{i,j}) = 1$ for the other half.

For this experiment, a GNN was employed where both the transition function $h_w$ and the output function $g_w$ were implemented by three-layered FNNs with five hidden neurons. The constraint $\alpha_n \in [-1, 1]$ was enforced using a hyperbolic tangent activation in the output layer of the FNN that implements $g_w$.

For each graph $G_i$ of the data set, the test procedure computed the difference between the desired result and the achieved one as $\delta_i = (|G_i|/2) - b_i$, where $b_i$ was the number of “hot” nodes. A node $n$ was considered hot if $\alpha_n > 0$. The GNN predicted the
correct result, i.e., $\delta_i = 0$, in 38% of the cases. Moreover, for only 2% of the total number of patterns, the differences were larger than 2 ($|\delta_i| > 2$). The dotted lines in Fig. 4 show the results achieved for each possible value of $\delta_i$ on the test set and the training set, respectively.

One may argue that the results achieved by GNNs cannot be correctly evaluated without a statistical analysis of the data set. In fact, even a simple procedure that assigns to each output a random value may often produce the right result, because the case $\delta_i = 0$ is the most probable one. On the other hand, the expected behavior of such a procedure can be easily computed\(^8\) and is depicted in Fig. 4 (continuous line). Interestingly, the GNN used the random labels to distinguish nodes and outperformed the random procedure. Moreover, the results have been compared also with a three-layer FNN (dashed line in Fig. 4). The FNN was fed only by node labels and did not use graph connectivity. The results obtained by such a network were very similar to those expected for the random procedure. In fact, the experiments have shown that the FNN just learns to produce a balanced number of hot and nonhot nodes in the whole data set.

### B. The Clique Problem

A clique of size $k$ is a complete subgraph with $k$ nodes\(^9\) in a larger graph (see Fig. 5). The goal of this experiment was to detect cliques of size 5 in the input graphs. More precisely, the GNN was trained to approximate the function defined by $\tau(G_i, n_{i,j}) = 1$, if $n_{i,j}$ belongs to a clique of size 5, and $\tau(G_i, n_{i,j}) = -1$, otherwise. The data set contained 2000 random graphs of 20 nodes each: 300 graphs in the training set, 300 in the validation set, and the rest in the test set. After the construction procedure described at the beginning of this section, a clique of size 5 was inserted into each graph of the data set. Thus, each graph had at least one clique, but it could have more cliques, due to the random data set construction. The graph density used in the construction ($\delta = 0.3$) was heuristically selected so as to build a small but not negligible number of graphs with two or more cliques. In fact, only about 65% of the graphs had only five nodes belonging to a clique (the graph contains just one clique), while in some particular cases more than half the nodes of a graph were involved in a clique (Fig. 6).

The overall percentage of nodes belonging to a clique was 28.2%. All the nodes were supervised and the desired outputs $t_{i,j} = \tau(G_i, n_{i,j})$ were generated by a brute force algorithm that localized all the cliques of the graphs.

Table I shows the accuracies achieved on this problem by a set of GNNs obtained by varying the number of hidden neurons of the FNNs that compose the GNN. The performance is measured as the percentage of nodes that have been correctly classified.

\begin{table}[ht]
\centering
\begin{tabular}{|c|c|c|c|c|}
\hline
Hiddens & test & train \\
\hline
2 & 83.73% & 83.45% \\
5 & 86.95% & 86.60% \\
10 & 90.74% & 90.33% \\
20 & 90.20% & 89.72% \\
30 & 90.32% & 89.82% \\
\hline
\end{tabular}
\caption{Results on the Clique Problem. The table displays the performance achieved on test and training sets with different numbers of hidden nodes in the FNNs that compose the GNN. The performance is measured as the percentage of nodes that have been correctly classified.}
\end{table}

\(^8\)Note that the most useful random procedure is the process that sets $\alpha_i$ to a value in $[-1, 1]$ with uniform probability. In this case, the probability of producing $\delta_i$, hot nodes in a graph with $d$ nodes is $\binom{d}{5}/2^d$, where $\binom{d}{5}$ is the binomial coefficient.

\(^9\)A graph is complete if there is an edge between each pair of nodes.
both FNNs. Finally, the dimension of the state was set to \( s = 2 \). Some experiments with larger states have shown only a marginal improvement of the performance.

The accuracy achieved on the test set is very close to the accuracy on training set, with any number of hidden units. This proves that the GNN model did not suffer from overfitting problems on this experiment and that the accuracy is satisfactory even with a reduced number of hidden neurons.

Finally, one may wonder whether the clique problem can be solved by a simpler approach, for example, by an FNN that takes in as input only the number of neighbors \( |\text{ne}[n_{i,j}]| \) of each node \( n_{i,j} \). The number of neighbors is informative on the nature of the data; this can be statistically closely correlated with the target \( t_{i,j} \). For instance, it is obvious that if \( |\text{ne}[n_{i,j}]| < 5 \), then \( n_{i,j} \) cannot belong to any clique of size five. Thus, an FNN with one input, 20 hidden neurons, and one output neuron was trained to predict \( t_{i,j} \) from \( |\text{ne}[n_{i,j}]| \). The accuracy reached by FNN averaged on five runs was 81.56%. As a consequence, GNNs always outperform FNNs, suggesting that GNNs are able to exploit more information from the graph topology than just the number of neighbors.

However, the difference between the performances of the two models, GNNs and FNNs, was not large. The clique task is a difficult problem for GNNs. In fact, in GNN model, the computation is localized on the nodes of the graph [see (1)], while the detection of a clique requires the simultaneous knowledge of the properties of all the nodes involved in the clique. Learning procedure should adapt the parameters so that the transition function \( h_m \) accumulates the needed information into the node states, while the output function \( g_m \) decodes the states and produces the right answer. Thus, as suggested by the proofs of Theorems 2 and 3, those functions may be very complex and the learning may be difficult.\(^{11}\)

\( ^{11} \)It is difficult to make a deeper analysis of the reasons for which a given function that can be realized in theory cannot be learned in practice. It is worth noticing, however, that similar problems can be encountered also in common recurrent neural networks, e.g., when a long sequence of inputs is processed (these problems are usually referred to as long time dependencies problems [28]).

C. The Neighbors Problem

This simple task consists of computing the number of neighbors \( |\text{ne}[n]| \) of each node \( n \). Since the information required to compute the desired output is directly available by counting the arcs entering to each node, GNNs are expected to perform much better on this problem than on the clique problem. On the other hand, the peculiarity of this experiment lies in the fact that the data set consisted of only one single large graph \( G \).

In each run of this experiment, one random graph \( G \) with 500 nodes was built. The data set \( L \) contained a pattern \((G, t_{i,j}, t_j)\), where \( t_j = |\text{ne}[n_j]| \), for each node \( n_j \) of the graph. The data set was randomly split into a training set (125 patterns), a validation set (125 patterns), and a test set (250 patterns). The performance was measured by the percentage of the patterns where GNNs achieved a relative error \( e_r \) lower than 0.05 and 0.1, respectively. Table II shows that GNNs solve this problem. As the number of hidden neurons in the FNNs becomes larger, so does the percentage of the patterns whose prediction is very close to the desired output \( t_j \). For a large number of hidden neurons, most of the patterns are correctly predicted.

\( ^{10} \)Increasing the number of hidden neurons did not improve the result significantly.

### Table II

<table>
<thead>
<tr>
<th>Hiddens</th>
<th>Test</th>
<th>Train</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( e_r &lt; 0.05 )</td>
<td>( e_r &lt; 0.1 )</td>
</tr>
<tr>
<td>2</td>
<td>73.64%</td>
<td>77.40%</td>
</tr>
<tr>
<td>5</td>
<td>89.56%</td>
<td>89.76%</td>
</tr>
<tr>
<td>10</td>
<td>90.64%</td>
<td>91.44%</td>
</tr>
<tr>
<td>20</td>
<td>99.04%</td>
<td>99.72%</td>
</tr>
</tbody>
</table>

D. The Second-Order Neighbors Problem

For this experiment, the graph was constructed as in the neighbors problem. Here, the goal is to compute, for each node \( n_j \), the number of distinct neighbors’ neighbors. In other words, the GNN should predict the number of nodes \( |\text{ne}[\text{ne}[n_j]]| \) that are reachable from \( n_j \) by a path containing two edges; the nodes that are connected to \( n_j \) by several paths must be counted only once and \( n_j \) itself should not be counted.\(^{12}\)

For this reason, this problem is more difficult to learn than the neighbors problem. Table III shows the obtained results. As in the neighbors problem, the error decreases for larger numbers of hidden units. However, in this case, the GNNs can solve the problem only partially and the percentage of patterns with small relative error \( e_r < 0.1 \) never exceeds 89%.

### Table III

<table>
<thead>
<tr>
<th>Hiddens</th>
<th>Test</th>
<th>Train</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( e_r &lt; 0.05 )</td>
<td>( e_r &lt; 0.1 )</td>
</tr>
<tr>
<td>2</td>
<td>65.96%</td>
<td>81.88%</td>
</tr>
<tr>
<td>5</td>
<td>66.00%</td>
<td>81.64%</td>
</tr>
<tr>
<td>10</td>
<td>66.04%</td>
<td>80.00%</td>
</tr>
<tr>
<td>20</td>
<td>72.48%</td>
<td>88.48%</td>
</tr>
<tr>
<td>30</td>
<td>70.40%</td>
<td>81.08%</td>
</tr>
</tbody>
</table>

E. The Tree Depth Problem

The goal of the task was to compute the depth \( d(n_j) \) of each node \( n_j \) in a tree, i.e., the length of the path from the root of the tree to node \( n_j \). In each run, the data set contained one large tree \( T \), with 10 000 nodes. The tree was built starting from the root and attaching to each node a number of children randomly chosen between 0 and 5. Then, the procedure was applied recursively to each leaf until \( T \) contained the given number of nodes. If the final tree had less than 10,000 nodes (this could

\( ^{12} \)More precisely, the desired output \( t_{i,j} \) was normalized so that it belongs to \([0, 1]\), i.e., \( t_{i,j} = |\text{ne}[\text{ne}[n_j]]|/M \), where \( M \) is the maximum number of neighbors’ neighbors \( M = \max_j |\text{ne}[\text{ne}[n_j]]| \).
have happened as nodes may have no children), the construction was repeated. The depth of the trees, measured after the completion of the construction process, usually belonged to the interval [10, 25].

Thus, each data set consisted of 10,000 patterns \((T, n_j, t_j)\), where \(t_j = d(n_j) / D\) and \(D\) is the maximum depth of the tree, i.e., \(D = \max_j d(n_j)\). Training set and validation set collected 2000 random patterns from the data set; the remaining 6000 patterns constituted the test set.

Intuitively, this task appears to be more difficult than the neighbors problem, but less difficult than neighbors’ neighbors problem. In fact, the depth cannot be computed using only the local information as in the neighbors problem. On the other hand, the depth of a node depends on the depth of the parent and such a dependence is expressed by a simpler function than in neighbors’ neighbors problem. The results achieved in the experiments seem to confirm such an intuitive idea.

V. CONCLUSION

In this paper, we studied the approximation properties of graph neural networks, a recently introduced connectionist model for graph processing. First, we defined the class of functions preserving the unfolding equivalence. Such a class contains most of the practically useful maps on graphs. In fact, only when the input graph contains symmetries, the unfolding equivalence may not be preserved. Then, we proved that GNNs can approximate, in probability, up to any degree of precision any function that preserves the unfolding equivalence and that, vice versa, any function implemented by GNNs preserves the unfolding equivalence. The presented results extend and include those already obtained for RNNs, the predecessor model of GNNs, and prove that the GNN model can be applied to more general classes of applications. Some experimental examples shed some light on the computational capability of the model and have been discussed w.r.t. the developed theory.

As a topic of future research, it may be useful to consider theoretical issues that have been considered for common connectionist models, but have not been studied for GNNs. For example, the investigation of the generalization properties of GNNs may require the extension of the concepts of Vapnik–Chervonenkis dimension [29] and minimum description length [30]. Moreover, conditions under which the error function does not have any local minima have been considered for FNNs [31–33], but not yet for GNNs. Similarly, there are no studies, analogous to those in [34], on the closure of the class of functions that can be implemented by GNNs.

### APPENDIX

#### PROOFS

The proofs of the main results can be found in this appendix.

A. Proof of Theorem 1—Functions of Unfolding Trees

If there exists \(\kappa\) such that \(\varphi(G, n) = \kappa(T_n)\), then \(n_1 \sim n_2\) implies

\[
l(G, n_1) = \kappa(T_{n_1}) = \kappa(T_{n_2}) = l(G, n_2).
\]

On the other hand, if \(l\) preserves the unfolding equivalence, then we can define \(\kappa\) as \(\kappa(T_n) = l(G, n_1)\). Note that the above equality is a correct specification for a function. In fact, if \(T_{n_1}\) and \(T_{n_2}\) are two unfolding trees, then \(T_{n_1} = T_{n_2}\) implies \(l(G, n_1) = l(G, n_2)\), such that \(\kappa\) is uniquely defined.

B. Proof of Theorem 2—Approximation by Positional GNNs

For the sake of simplicity, the theorem will be proved assuming \(m = 1\), i.e., \(\tau(G, n) \in \mathbb{R}\). However, the result is easily extended to the general case when \(\tau(G, n) \in \mathbb{R}^m\) is a vector. The GNN that satisfies the theorem can be defined by composition of \(m\) GNNs, each one approximating a component of \(\tau(G, n)\).

According to Theorem 1, there exists a function \(\kappa\) such that \(\tau(G, n) = \kappa(T_n)\). Thus, the main idea of the proof consists of designing a GNN that is able to encode the unfolding trees into the node states. The stable state of a node will be \(x_n = \nabla(T_n)\), where \(\nabla\) is an encoding function that maps trees to real numbers. In this way, the output function \(g\) will obtain a representation of \(T_n\) by decoding the state and will produce the desired output using \(\kappa\). Said differently, the recursive activation of \(f\) will implement \(\nabla\), and \(g\) will implement \(\kappa \circ \nabla^{-1}\), where \(\nabla^{-1}\) is the inverse function of \(\nabla\) and \(\circ\) is the function composition operator.

The proof of the theorem is organized into three sections. In the next section, some preliminary lemmas are proved, which allow to restate the theorem in a simpler form. Then, the coding function \(\nabla\) is defined. Finally, it is proved that \(\nabla\) can be implemented by a transition function \(f\) and that the corresponding global transition function \(F\) is a contraction map.

1) Preliminary Results: Theorem 2 requires \(\tau\) to be approximated in probability on the whole \(D\), i.e.,

\[
P(\|\tau(G, n) - \varphi(G, n)\| \leq \varepsilon) \geq 1 - \lambda.
\]

The first step of the proof consists of two lemmas, which simplify this problem by showing that the theorem can be reduced to a simpler form where the approximation \(\|\tau(G, n) - \varphi(G, n)\| \leq \varepsilon\) is achieved just on finite sets of patterns \((G, n_i)\) where \(1 \leq i \leq v\).

Moreover, it is also proved that it is sufficient to consider graphs with integer labels only. Formally, Theorem 2 will be reduced to the following theorem.

**Theorem 5:** For any finite set of patterns \((G, n_i)\) where the graphs have integer labels, any function: \(\tau : D \to \mathbb{R}^m\), which preserves the unfolding equivalence, any real: \(\varepsilon, \mu\), where \(\varepsilon > 0\) and \(0 < \mu < 1\), there exist two continuously differentiable functions \(f\) and \(g\) such that for the GNN defined by

\[
x_n = f(l_n, l_{\text{in}}[n], x_{\text{in}}[n], l_{\text{in}}[n]),
\]

\[
o_n = g(x_n, l_n), \quad n \in \mathbb{N}
\]
the global transition function \( F \) is a contraction map with contracting constant \( \mu \), the dimension of the state is \( s = 1 \), the stable state is uniformly bounded, and

\[
\| \tau(G_i, n_i) - \varphi(G_i, n_i) \| \leq \varepsilon
\]

holds for any \( i, 1 \leq i \leq \nu \), where \( \varphi \) is the function implemented by the GNN.

The reduction is carried out by proving two lemmas. The first lemma proves that the domain can be divided into small subsets \( D_1, D_2, \ldots \), such that the graphs in each subset have the same structure and have similar labels (see Fig. 7). A finite number of \( D_i \) is sufficient to cover a subset of the domain whose probability is larger than \( 1 - \lambda \).

**Lemma 1:** For any probability measure \( P \) on \( D \), and any reals \( \lambda \) and \( \delta \), where \( 0 < \lambda \leq 1 \) and \( \delta > 0 \), there exist a real \( \eta > 0 \), which is independent of \( \delta \), a set \( D \subseteq D \), and a finite number of partitions \( \bar{D}_1, \ldots, \bar{D}_v \) of \( D \), where \( \bar{D}_i = G_i \times \{ n_{i} \} \) for a graph \( G_i \subseteq G \), and a node \( n_{i} \), such that:

1) \( P(\bar{D}) \geq 1 - \lambda \) holds;
2) for each \( i \), all the graphs in \( G_i \) have the same structure, i.e., they differ only in the values of their labels;
3) for each set \( \bar{D}_i \), there exists a hypercube \( H_i \in \mathbb{R}^\nu \) such that \( l_G \notin H_i \) holds for any graph \( G \in G_i \), where \( l_G \) denotes the vector obtained by stacking all the labels of \( G \);
4) for any different sets \( G_i, G_j, i \neq j \), their graphs have different structures or their hypercubes \( H_i, H_j \) have a null intersection \( H_i \cap H_j = \emptyset \);
5) for each \( i \) and each pair of graphs \( G_1, G_2 \in G_i \), the inequality \( \| l_{G_1} - l_{G_2} \|_\infty \leq \delta \) holds;
6) for each graph \( G \) in \( \mathcal{D} \), the inequality \( \| l_G \|_\infty \leq \bar{b} \) holds.

**Proof:** Two graphs may differ either because of their different structures or because of the different values of their labels. Since the set of the possible structures is enumerable, the set of graphs \( G \) can be partitioned into a sequence of disjoint subsets \( A_1, A_2, A_3, \ldots \), where each \( A_i \) contains graphs having the same structure (they differ only for their label values). Moreover, since there is a finite number of nodes in a graph, also \( \mathcal{D} \) can be partitioned into a sequence \( B_1, B_2, \ldots \), where, for each \( i \), \( B_j = C_j \times \{ v_{ij} \} \), where \( v_{ij} \) is equal to an \( A_i \), for some \( i \), and \( v_{ij} \) is a node of the corresponding graph (structure).

Let \( \eta \) be a real number, \( 0 < \eta \leq \delta \), \( \delta \) be defined by \( b = \delta \eta \) for some integer \( \delta \), and \( l_i^\eta \) be the interval \( l_i^\eta = [-b + (i - 1)\delta, -b + i\delta - \eta] \), where \( 1 \leq i \leq 2d \). Moreover, consider all the hypercubes \( H_i^\eta \) that can be constructed by taking values in the \( l_i^\eta \), e.g., \( H_i^\eta = l_1^\eta \times l_2^\eta \times l_3^\eta \times l_1^\eta \) is a four-dimensional hypercube. In the following, we will denote these hypercubes as \( H_i^\eta, H_j^\eta, \ldots \). Note that each \( H_i^\eta \) is contained in \([-b, b]^\nu \), for some \( a_\nu \), and their union \( H_i^\eta = \bigcup_{i=1}^{d} H_i^\eta \) approximates \( \bigcup_{i=1}^{d} [b, 2b]^\nu \), when \( \eta \to 0 \). Moreover, for any points \( l_1, l_2 \in H_j^\eta \), we have \( |l_1 - l_2|_\infty \leq \delta \), since each interval is shorter than \( \delta \).

Let \( B_i^\eta \) be the subset of \( B_i \) containing only the graphs the labels of which belong to \( H_i^\eta \). Since

\[
\lim_{b \to \infty} P\left( \bigcup_{i,j} B_i^\eta_{i,j} \right) = P\left( \lim_{b \to \infty} \left( \bigcup_{i,j} B_i^\eta_{i,j} \right) \right) = P\left( \mathcal{D} \right) = 1
\]

there exists \( \bar{b} \) such that

\[
P\left( \bigcup_{i,j} B_i^\eta_{i,j} \right) \geq 1 - \lambda/2
\]

Moreover, since

\[
\lim_{k \to \infty} \eta = 0 \left( \bigcup_{i,j,k} B_i^\eta_{i,j,k} \right) = \bigcup_{i,j} B_i^\eta_{i,j}
\]

there exist \( \bar{b} \) and \( \eta \) such that

\[
P\left( \bigcup_{i,j,k} B_i^\eta_{i,j,k} \right) \geq P\left( \bigcup_{i,j,k} B_i^\eta_{i,j,k} \right) - \lambda/2 \geq 1 - \lambda.
\]

The sets \( B_i^\eta_{i,j} \) involved in (9) satisfy the properties expected of the sets \( D_1, \ldots, D_v \) of the theorem and the \( H_i^\eta \) are the corresponding hypercubes. In fact, (9) implies point 1 of the theorem. Points 2–4 of the theorem follow by definition of the sets \( B_i^\eta_{i,j} \). Moreover, point 5 of the theorem holds because the labels of the graphs in \( B_i^\eta_{i,j} \) belong to the same hypercube \( H_i^\eta \).
Finally, since the labels of the graphs in $L_d^G$ are vectors with components in $[-T, T]$, also point 6 of the theorem holds.

The following lemma proves the proof of the equivalence between Theorem 2 and Theorem 5. The intuitive idea behind the proof of the theorem is that of constructing a GNN, which produces a constant output on each subset $D_i$. Since there is only a finite number of subsets $D_i$, Theorem 5 ensures that the construction is possible. Since the $D_i$ are small and $\tau$ is continuous, such a GNN will also satisfy the hypothesis of Theorem 2.

**Lemma 2:** Theorem 2 holds if and only if Theorem 5 holds.

**Proof:** Theorem 2 is more general than Theorem 5, so one direction of the implication is straightforward. On the other hand, let us assume that Theorem 5 holds and we have to show that this implies Theorem 2.

Let us apply Lemma 1 setting the values $P$ and $\lambda$ of the hypothesis equal to the corresponding values of Theorem 2 and $\delta$ being any positive real number. It follows that there is a real $\delta$ and a subset $\delta$ of $D$ such that $P(\delta) > 1 - \lambda$. Let $M$ be the subset of $D$ that contains only the graphs $G$ satisfying $\|G\|_\infty \leq \delta$. Note that since $\delta$ is independent of $\delta$, then $\delta \subset M$ for any $\delta$.

Since $\tau$ is integrable, there exists a continuous function$^{14}$ that approximates $\tau$ up to any degree of precision in probability. Thus, without loss of generality, we can assume that $\tau$ is continuous w.r.t. the labels. Moreover, since $M$ is bounded, $\tau$ is equicontinuous on $M$. By definition of equicontinuity, a real $\delta > 0$ exists such that

$$|\tau(G_1, n) - \tau(G_2, n)| \leq \frac{\varepsilon}{2}$$

holds for any node $n$ and for any pair of graphs $G_1, G_2$ having the same structure and satisfying $\|G_1 - G_2\|_\infty \leq \delta$.

Let us apply Lemma 1 again, where, now, the $\delta$ of the hypothesis is set to $\delta$, i.e., $\delta = \delta$. In the following, $D_i = G_i \times \{n\}$, $1 \leq i \leq v$, represents the sets obtained by the new application of the intervals defined in the proof of Lemma 1.

Let $\theta : \mathbb{R}^l \rightarrow \mathbb{R}^l$ be a function that encodes reals into integers as follows: for any $i$ and any $z \in \Omega_i$, $\theta(z) = i$. Thus, $\theta$ assigns to all the values of an interval $I_i^\Omega_i$ the index $i$ of the interval itself. Since the intervals do not overlap (see Fig. 7) and are not contiguous, $\theta$ can be continuously extended to the entire $\mathbb{R}^l$. Moreover, $\theta$ can be extended also to vectors: let $\Theta(Z)$ denote the vector of integers obtained by coding all the components of $Z$. Finally, let $\Theta : G \rightarrow \mathbb{G}$ represent the function that transforms each graph by replacing all the labels with their coding, i.e., $L_\Theta(G) = \theta(L_G)$.

Let $G_1, \ldots, G_v$ be graphs, each one extracted from a different set $G_i$. Note that, according to points 3–5 of Lemma 1, $\Theta$ produces an encoding of the sets $G_i$. More precisely, for any two graphs $G_1, G_2$ of $\overline{D}$, we have $\Theta(G_1) = \Theta(G_2)$, if the graphs belong to the same set, i.e., $G_1, G_2 \in G_i$; and we have $\Theta(G_1) \neq \Theta(G_2)$, otherwise. Thus, we can define a function $\Gamma$ such that $\Gamma(\Theta(G_i), n_i) = (G_i, n_i)$, $1 \leq i \leq v$.

Consider the problem of approximating $\tau$ on $\overline{D}$ on the set $(\Theta(G_1), n_1), \ldots, (\Theta(G_v), n_v)$. Theorem 5 can be applied to such a set, because the set contains a finite number of graphs with integer labels. It follows that there exists a GNN that implements a function $\varphi$ such that, for each $i$

$$|\tau(\Gamma(\Theta(G_i), n_i)) - \varphi(\Theta(G_i), n_i)| \leq \frac{\varepsilon}{2},$$

(11)

Let $f$ and $g$ be the encoding function and the output function, respectively, that realize the above GNN. Consider the GNN described by

$$\begin{align*}
x_n &= f(\theta(l_n), \hat{\theta}(l_{i[X]}), x_{\hat{\theta}(l_{i[X]})}) \\
o_n &= g(x_n, n_n)
\end{align*}$$

(12)

and let $\varphi$ be the function implemented by this GNN. It is easily shown that for any $i$ and $G \in G_i$

$$\varphi(G, n_i) = \varphi(\Theta(G_i), n_i)$$

holds. Putting together the above equality with (10) and (11), it immediately follows, for any $(G, n) \in D_i$

$$|\tau(G, n) - \varphi(G, n)| = |\tau(G, n) - \tau(G_i, n) + \tau(G_i, n) - \varphi(G, n)|$$

$$\leq |\tau(G_i, n) - \varphi(G, n)| + \frac{\varepsilon}{2}$$

$$= |\tau(\Gamma(\Theta(G_i), n)) - \varphi(\Theta(G_i), n)| + \frac{\varepsilon}{2} \leq \varepsilon.$$

Thus, the GNN described by (12) satisfies $|\tau(G, n) - \varphi(G, n)| \leq \varepsilon$ in the restricted domain $D$. Since $P(\delta) \geq 1 - \lambda$, we have

$$P(|\tau(G, n) - \varphi(G, n)| \leq \varepsilon) \geq 1 - \lambda$$

and the lemma has been shown to be true.

2) The Coding Function: The main idea of the proof is that of designing a transition function $f$, which is able to encode the input graph into the node states. In this way, the output function $g$ has to only decode the state and produce the desired outputs. Of course, the transition function $f$ cannot access directly the whole input graph, but has to read it using the information stored in the states of the neighbor nodes. On the other hand, the target function $\tau$ preserves the unfolding equivalence by hypothesis and there exists a function $\kappa$ such that $\tau(G, n) = \kappa(T_n)$. Thus, an obvious solution will be to store directly the unfolding $T_n$ of node $n$ into the state $x_n$. More precisely, in place of $T_n$, which is infinite and cannot be directly memorized, it is sufficient to store the unfolding up to a depth $r$, where $\tau$ is the total number of nodes contained in the graphs $G_1, \ldots, G_v$. In fact, the following lemma shows that $T_n$ is sufficient to define the unfolding equivalence.

**Lemma 3:** Let us consider the unfolding equivalence defined on a set of graphs $G_1, \ldots, G_v$. For any two nodes $u$ and $v$, $u \sim n$ holds if and only if $T_u^v = T_u^{v'}$ holds, where $r = \sum_{i=1}^{n} |N_i|$, and $G_i = (N_i, E_i)$.

**Proof:** The “only if” part of the proof is straightforward. In fact, by definition, $u \sim v$ implies $T_u^v = T_u^{v'}$ for each $k$. Thus, $T_u^v = T_u^{v'}$ follows. For the “if part,” let us assume $T_u^v = T_u^{v'}$. Note that, for any integer $k \geq 1$, $T_u^v = T_u^{v'}$ implies $T_u^{v'} = T_u^{v'} - 1 = T_u^{v'} - 1$, because $T_u^{v'} - 1$ and $T_u^{v'} - 1$ are subtrees of $T_u^v$ and $T_u^{v'}$, respectively. Thus, there are only three possible cases: 1) $T_u^v = T_u^{v'}$ for any $k$; 2) $T_u^v = T_u^{v'}$, for any $k$; and 3) there exists a $k$ such that $T_u^v = T_u^{v'}$, for $k < \kappa$ and $T_u^{v'} = T_u^{v'}$, for $k \geq \kappa$.
1) immediately supports our theorem, and case 2) is absurd by the assumption that $T^{k}_n = T^{k}_u$ of Theorem 5. Hence, case 2) cannot be true. Let us discuss case 3): we will show that $k < r$. If $n$ and $u$ have different (node or edge) labels, their unfolding trees are immediately different at depth 1, i.e., $T^{k}_n \neq T^{k}_u$. On the other hand, if two nodes $n$ and $u$ have the same labels and are connected to the neighbors by edges having the same labels, then $T^{k}_n \neq T^{k}_u$ may happen only because they have different subtrees, which implies that the set of the unfolding trees of the neighbors are different. Putting together the above reasoning with the assumption of case 3), we deduce the following inference rule:

If $T^{k}_n \neq T^{k}_u$ and $T^{k-1}_n = T^{k-1}_u$, then there are two neighbors $n_2, n_3$ of $n, u$, respectively, for which $T^{k-1}_n \neq T^{k-1}_u$ and $T^{k-2}_n = T^{k-2}_u$ hold.

Let us consider the equivalence $\sim_k$ defined by $u \sim_k u$ if and only if $T^{k}_n = T^{k}_u$ and let us denote by $\equiv$ the equality for equivalences. The largest equivalence, i.e., $u \sim_1 v$ for each $u, v$ having the same label. Then, while $k$ increases, $\sim_k$ becomes more and more refined until $\sim_k$ becomes constant and equals the unfolding equivalence $\equiv$. The above inference rule suggests that if $\sim_{k-1} \neq \sim_{k-2}$ then $\sim_{k-1} \neq \sim_{k-2}$, i.e., $\sim_{k-1} \equiv \sim_{k-2}$ implies $\sim_{k} \equiv \sim_{k-1}$. Thus, all the steps $k$ where $\sim_k$ is refined are consecutive. Since at each refining step at least a class of the equivalence defined by $\sim_k$ is split and the number of equivalences classes cannot be larger than the number of nodes, then there exist at most $r - 1$ refining steps. As a consequence, $k < r$ holds.

In the following, we describe a representation that will encode trees by real numbers. Such a representation will be used to store the unfolding trees into the states. More precisely, let $G_1, \ldots, G_r$ be the graphs considered in Theorem 5. We will restrict our attention only to the trees up to depth $r$ that can be built from the graphs $G_1, \ldots, G_r$, i.e., the trees $\mathcal{U} = \{ T^{d}_n | 1 \leq d \leq r, n$ is a node of $G_i, 1 \leq i \leq v\}$. Our purpose is that of designing an encoding $\nabla(T^{d}_n)$, which maps the tree $T^{d}_n$ to a real number and is defined for any $T^{d}_n \in \mathcal{U}$. The function $\nabla$ will be specified in two steps.

Step 1) A map $\nabla$ will be defined, which assigns a different integer number to each quintuple $(i, l, n_1, l_1, n_2, l_2)$, where $l_2$ is the $i$th neighbor of $n_2$. Moreover, the coding $\nabla$ will be defined as

$$\nabla(T^{d+1}_n) = \sum_{i=1}^{[\text{ne}(n)]} \gamma^0(i, l, n_1, n_2, l_1, l_2)$$

where $\gamma$ is any positive real number smaller than $Q/(2r(1+Q))$. Here, $Q$ is given by $Q = Q_1/Q_2$, where $\mu$ is the contraction constant of Theorem 2 (which we are proving), and $Q_1, Q_2$ are two real numbers such that $Q_1 |z| \leq \|z\| \leq Q_2 |z|$ holds for any $z$, the 1-norm $\|z\|_1 = \sum_i |z_i|$, and the norm $\|z\|$ of the hypothesis of Theorem 2.\(^{15}\)

Step 2) It will be proved that $\nabla$ is injective on $\mathcal{U}$ and there exists a decoding function $\nabla^{-1}$ such that $T^{d}_n = \nabla^{-1}(\nabla(T^{d}_n))$.

The two steps are discussed with more details in the following.

Step 1) Function $\nabla$: Since $\mathcal{U}$ contains a finite number of trees, only a finite number of quintuples $(i, l, n_1, n_2, l_1, l_2)$ exists. So, we can enumerate all the possible quintuples and define the coding $\nabla$ that assigns a different integer to each quintuple. Among the possible assignments, we select a $\nabla$ that is monotonically increasing w.r.t. $d$. More precisely, we assume that for any $d$ and any nodes $n, u, n_1, n_2, n_3$ and $\nabla_j$

$$\nabla \left( i, l, n_1, n_2, l_1, l_2 \right) > \nabla \left( j, l, n_1, n_2, l_1, l_2 \right)$$

holds.

Step 2) The Decoding Function $\nabla^{-1}$: Let us consider the function $\xi : \mathcal{U} \rightarrow \mathcal{P}$ that takes in as input an unfolding tree $T^{d}_n$ and returns the polynomial of the variable $\gamma$ that is represented on the right-hand side of (13). Notice that the function $\xi$ is injective on $\mathcal{U}$, because the polynomial $\xi(T^{d}_n)$ contains a term for each quintuple $(i, l, n_1, n_2, l_1, l_2)$. In fact, a quintuple contains all the information related to a neighbor of $n$ and is uniquely described by $\nabla$.

We will show that $\nabla$ is also injective by using a reduction to absurdity argument. Let us assume that $\nabla(T^{d+1}_n) = \nabla(T^{d+1}_u)$ holds, for some $n_1, n_2, d_1, d_2$, and that $T^{d_1}_n = T^{d_2}_u$ does not hold. By definition, we have $\xi(T^{d_1}_n)(\gamma) = \nabla(T^{d_1}_n) = \nabla(T^{d_2}_u) = \xi(T^{d_2}_u)(\gamma)$. On the other hand, the polynomial function $\xi(T^{d_1}_n)$ is different from $\xi(T^{d_2}_u)$ because $\xi$ is injective. Thus, $\gamma$ is a root of the nonnull polynomial $\xi(T^{d_1}_n) - \xi(T^{d_2}_u)$. Such a conclusion cannot be true by the following lemma, which shows that if $\gamma$ is a positive real number, sufficiently close to 0, then $\gamma$ cannot be a root of $\xi(T^{d_1}_n) - \xi(T^{d_2}_u)$.

Lemma 4: Let $p(x) = \sum_{i=1}^{k} a_i x^i$ be a polynomial in $x$ with integer coefficients and let $B$ be the maximal magnitude of the coefficients, i.e., $B = \max_{i=1}^{k} |a_i|$. Then, $p$ has no root in the open interval $(0, 1/2B)$.

Proof: Let $a_j$ be the first nonnull coefficient, i.e., $a_j = 0, 1 \leq i \leq j - 1, a_j \neq 0$. Moreover, let us assume $a_j > 0$: the proof when $a_j < 0$ follows by a similar reasoning as shown here. By using simple algebra

$$p(x) = \sum_{i=1}^{k} a_i x^i \geq a_j x^j - \sum_{i=j+1}^{k} a_i x^i \geq a_j x^j - \sum_{i=j+1}^{k} B x^i$$

$$= a_j x^j - B x^{j+1} \frac{1 - x^{k-j}}{1 - x}$$

$$= \frac{x^j}{1 - x} (1 - x - B x (1 - x^{k-j}))$$

$$= \frac{x^{j+1} (1 - x^{k-j})}{1 - x}$$

$$= \frac{x^{j+1} (1 - x^{k-j})}{1 - x} (\frac{1 - x}{1 - x^{k-j}})$$

$$= \frac{2B}{2 - B} = 0$$

\(^{15}\)Such a definition is made possible by the fact that all norms on a finite-dimensional space over $R$ are equivalent.
where the last inequality follows by the assumption $x \in (0,1/(2\varepsilon))$, which implies $1-x > 1/2$, and $1/(x(1-x^{r-1})) > 2B$. Hence the lemma is true. □

More precisely, note that the coefficients of the polynomial $\xi(T_{n,t_1}^d) - \xi(T_{n,t_2}^d)$ can assume only three numerical values $-1,0,1$. Thus, we can apply Lemma 4 to $\xi(T_{n,t_1}^d) - \xi(T_{n,t_2}^d)$ with $B = 1$. It follows that provided that $0 < \gamma < Q/(2r(1+Q)) < 1/(2r) \leq 1/(2B)$ holds, $\triangledown$ is injective on $U$ and there exists a decoding function such that $T_{n,t_1}^d = \triangledown^{-1}(\triangledown(T_{n,t_1}^d))$.

3) Implementation of $\triangledown$: In this section, we will show how a GNN can implement the coding $\triangledown$ and store $\triangledown(T_{n,t_1}^d)$ in the state of a node $n$. In fact, a GNN can construct the coding $\triangledown$ recursively storing in the states larger and larger unfolding trees. At the beginning, the states are set to a predefined initial value, which represents a void tree $(x(n_0) = \triangledown(T_0))$. Then, the transition function $f$ constructs the representation of a deeper unfolding tree each time the node is activated. In fact, $f$ builds $\triangledown(T_{n,t_1}^d)$, using the set $\{\triangledown(T_{n,t_1}^d), \ldots, \triangledown(T_{n,t_1}^{d+1})\}$ of the representations stored in the states of the neighbors. The construction process is stopped when the depth $r$ is reached: $f$ is defined so that $x(n,d) = \triangledown(T_{n,t_1}^d)$ for each $n$ and $d \geq r$. Thus, our goal is to implement the following transition function:

$$f\left(l_n, \triangledown(T_{n,t_1}^d), l_{\text{col}}[n], l_{\text{ne}}[n]\right) = \begin{cases} \triangledown(T_{454}^{d+1}), & \text{if } 0 \leq d \leq r - 1 \\ \triangledown(T_{n,t_1}^d), & \text{if } d \geq r. \end{cases} \quad (15)$$

Such a goal is reached by defining $f$ as

$$f(l_n, y_{\text{col}}[n], l_{\text{ne}}[n]) = \sum_{i=1}^{[x(n)]} h(i, l_n, l_{(n,t_i)}), y_i, l_{u_i} \quad (16)$$

where $y$ is the representation of any set of unfolding trees and $y_i$ is the representation of the $i$th tree contained in $y$. Moreover, $h$ is the function

$$h(i, l_n, l_{(n,t_i)}), z, l_{u_i}) = \begin{cases} \gamma^i(l_i, l_{(n,t_i)}, \triangledown^{-1}(z), l_{u_i}), & \text{if } 0 \leq d \leq r - 1 \\ \gamma^i(l_i, l_{(n,t_i)}, \triangledown^{-1}(z), l_{u_i}), & \text{if } d \geq r. \end{cases} \quad (17)$$

where $\gamma$ is the real number in the definition of the coding function $\triangledown$ [see (13)], $z$ is a representation of an unfolding tree, and $\zeta$ is defined as

$$\zeta(T_{u_i}^d) = T_{u_i}^{d-1} \quad (18)$$

i.e., $\zeta$ is a function that extracts from the unfolding tree $T_{u_i}^d$ the tree $T_{u_i}^{d-1}$, which is related to the same node $u_i$ but has a shallower depth.\footnote{Note that such a definition is made possible by the fact that an unfolding tree of a given depth $d$ contains the unfolding tree of a shallower depth $d - 1$.}

It is easily observed that such a function $f$ satisfies (15) and realizes the construction of the coding $\triangledown(T_{n,t_1}^d)$ as desired. In fact, from (13), it follows:

$$f\left(l_n, \triangledown(T_{n,t_1}^d), l_{\text{col}}[n], l_{\text{ne}}[n]\right) = \sum_{i=1}^{[x(n)]} h(i, l_n, l_{(n,t_i)}), \triangledown(T_{n,t_1}^d), l_{u_i} \quad (19)$$

On the other hand, $f$ and $g$ are still defined only on a finite set of points, e.g., $f$ is not defined when the first input parameter does not contain a label of a node or the second input parameter is not the coding of a tree. Since we are looking for a differentiable function, $f$ and $g$ must be extended to accept any vector of reals. Any continuously differentiable extension of $g$ works, because $g$ will operate only on the final stable state. On the other hand, the extension of $f$ must be carefully designed to ensure that the corresponding global transition function $F$ is a contraction map. Lemma 5 produces the needed results to achieve this goal.

**Lemma 5:** For any positive real $\vartheta$, there exists a continuously differentiable function $h : \mathbb{R}^{d+2N+r+1} \to \mathbb{R}$ such that if $f$ is defined as in (16) and $F$ is the global transition function corresponding to $f$, then:

1) equation (15) holds for any unfolding tree $T_{n,t_1}^d \in U$;
2) the inequality

$$||F(x,l) - F(y,l)||_1 < r \left(\frac{\gamma}{1-r\gamma} + \vartheta\right)||x-y||_1$$

holds for any $x$, $y$ and any $l$.

**Proof:** The proof of this lemma is more involved. In order to preserve the flow of the proof of Theorem 2, we will defer the proof until Section B4 of the Appendix. □

In fact, since $0 < \gamma < Q/(2r(1+Q))$ by definition of $\gamma$, then

$$r \left(\frac{\gamma}{1-r\gamma} + \vartheta\right) = \frac{Q}{1+Q} + \vartheta < Q$$

holds for a sufficiently small $\vartheta$. As a consequence, the second point of Lemma 5 and the definition of $Q, Q_1, Q_2$ (see definition of $\triangledown$ in step 1 in Section B2 of the Appendix) implies

$$||F(x,l) - F(y,l)||_1 \leq \frac{1}{Q_1}||F(x,l) - F(y,l)||_1$$

Thus, $F$ is a contraction map with contraction constant smaller than $\mu$ and Theorem 2 has been proved.

4) **Proof of Lemma 5:** In order to carry out the proof, some properties of the function $h$ and of the coding $\triangledown$ must be considered. The following lemma shows that $h$ behaves as a contraction map with respect to the domain of the trees in $U$. The proof is straightforward.
Lemma 6: Let \( h \) be defined as in (17). For any node \( n \) and any integers: \( d_1 \geq 1, d_2 \geq 1 \), and \( 1 \leq i \leq r \), the inequality
\[
\left| h \left( i, l_n, l_{(n,u_i)}, \nabla (T^h_{u_i}), l_{u_i} \right) - h \left( i, l_n, l_{(n,u_i)}, \nabla (T^d_{u_i}), l_{u_i} \right) \right| \leq \frac{\gamma}{1 - \nu \gamma} \left| \nabla (T^h_{u_i}) - \nabla (T^d_{u_i}) \right|
\]
holds, where \( u_i \) is the \( i \)th neighbor of \( n \).

Proof: Without loss of generality, we can assume \( r \geq d_2 > d_1 \geq 1 \). In fact, the proof of case \( r \geq d_1 > d_2 \geq 1 \) follows the same reasoning, and, in the case \( d_1 = d_2 \), it is straightforward. Moreover, by definition of \( h \), the cases \( d_1 > r \) and \( d_2 > r \) can be reduced to \( d_1 = r \) and \( d_2 = r \), respectively.

In the following, let \( m_d \) and \( M_d \) be, respectively
\[
m_d = \min_{n \in X_d} \left( \rho \left( i, l_n, l_{(n,u_i)}, T^d_{u_i}, l_{u_i} \right) \right)
\]
\[
M_d = \max_{n \in X_d} \left( \rho \left( i, l_n, l_{(n,u_i)}, T^d_{u_i}, l_{u_i} \right) \right)
\]
where \( \rho \) is the tuple coding function used in \( h \). Since, by definition, \( \rho \) is monotonically increasing w.r.t. \( d \) [see (14)], then
\[
m_{d-1} < M_{d-1} < m_d < M_d
\]
and, since \( 0 < \gamma < 1 \)
\[
(18)
\]
holds for any \( n, i, \) and \( d > 1 \). Using (18), it follows:
\[
\left| \nabla (T^h_{u_i}) - \nabla (T^d_{u_i}) \right| \leq \frac{\gamma M_d}{1 - \nu \gamma} \leq \frac{\gamma M_{d-1}}{1 - \nu \gamma} \leq \frac{\gamma m_{d-1}}{1 - \nu \gamma}
\]

Moreover, an upper bound on \( \gamma m_{d1} \) is established as
\[
\gamma m_{d1} = \frac{\gamma M_{d1}}{1 - \nu \gamma} \leq \frac{\gamma m_{d1} - \nu \gamma m_{d2-1}}{1 - \nu \gamma} \leq \frac{\gamma M_{d1} - \nu \gamma m_{d2-1}}{1 - \nu \gamma}.
\]

Lemma 7 shows that if a function is defined and if it is a contraction map only on a finite set of points, it can be extended to a contraction map on the entire input domain.

Lemma 7: Let \( \eta \) be a positive real number, \( l : A \rightarrow B \) be a function, and \( A \subset B \times B^d \) be a finite set of vectors. Assume that
\[
\| l(x, z) - l(y, z) \| \leq \eta \| x - y \| \tag{21}
\]
holds for any vectors \( [x, z], [y, z] \) that belong to \( A \), where \( x, y \in B \), \( z \in B^d \), and \( [\cdot, \cdot] \) denotes the operator that stacks two vectors. Then, for any positive real \( \theta \), \( l \) can be extended to the entire \( B \times B^d \). The resulting function \( \overline{l} \) equals \( l \) on \( A \), is infinitely differentiable, and satisfies
\[
\| \overline{l}(x, z) - l(y, z) \| \leq (\eta + \theta) \| x - y \| \tag{22}
\]
on the entire domain, i.e., for any vectors \( [x, z], [y, z] \) that belong to \( B \times B^d \).

Proof: The proof is carried out in five steps. Each step defines a new function \( l_i \) using the previous one: \( l_{i-1} \). The first function is the function \( l \) defined by the hypothesis; the last will be the function that satisfies the lemma.

Step 1—Extending \( l(x, z) \) to Some Large and Small Values \( x \): Let \( M = \{ z_1, \ldots, z_l \} \) be the set obtained by removing the first component from each vector in \( A \). For each \( z_i \), \( A_i = \{ [x_i, z_i] \} \) denotes the subset of \( A \) that includes all the vectors containing \( z_i \). Moreover, for each \( i \), let \( x_{iki}, x_{iki+1} \) be two real numbers that fulfill
\[
x_{iki} < \min_{j \neq i} \left( x_{iki}, \frac{l(x_{iki}, z_i)}{\eta} \right)
\]
\[
x_{iki+1} > \max_{j \neq i} \left( l(x_{iki}, z_i) \frac{x_{iki} + x_{iki+1}}{\eta} \right).
\]

In the following, \( B_i \) represents the superset of \( A_i \) defined by \( B_i = A_i \cup \{ [x, z_i] \} \). The function \( l_1 \) is a simple extension of \( l \) to \( B = \bigcup B_i \) and is defined by \( l_1(x, z) = l(x, z) \), if \( [x, z] \in A \), and \( l_1(x, z) = 0 \), otherwise.

We will prove that \( l_1 \) satisfies inequality (21) on \( B \). In fact, this claim holds in a straightforward manner if both \( [x, z] \) and \( [y, z] \)

belong to \( A \). On the other hand, if \( [x, z] = [x_{i,j}, z_i] \) and \([y, z] = [x_{i,j}, z_i]\) for some \( i,j \), then

\[
\eta|x_{i,j} - x_{i,j}| = \eta|v_{i,j} - v_{i,0}|
\]

\[
= \eta|v_{i,j} - \eta\min_{t \geq 0} \left(x_{t,i} - \frac{\|l(x_{t,i}, z_i)\|}{\eta}\right)
\]

\[
\geq \|l(x_{i,j}, z_i) - l(x_{i,j}, z_i)\|
\]

The proof of the claim follows a similar reasoning for the other cases, i.e., \([x, z] = [x_{i,j}, z_i]\), \([y, z] = [x_{i,j}, z_i]\), and \([y, z] = [x_{i,j}, z_i]\).

Step 2—Extending \( l_1(x, z) \) to any \( x \): Without loss of generality, let us assume that, for each \( i, x_{i,0}, \ldots, x_{i,k+1} \) are sorted according to their values, i.e., \( x_{i,j} < x_{i,j+1}, 0 \leq j \leq k \). Moreover, let \( C \) be defined as \( C = R \times M \).

The function \( l_2 \) generalizes \( l_1 \) to the set \( C \). More precisely, \( l_2 \) is

\[
l_2(x, z) = \begin{cases} 
1 \frac{x-x_{i,j}}{x_{i,j}-x_{i,j-1}}(l_{1}(x_{i,j+1}, z_i) - l_{1}(x_{i,j-1}, z_i)) & \text{if } x_{i,j} \leq x \leq x_{i,j+1}, 0 \leq j \leq k \\[0, \]
if \[x \leq x_{i,0} \] or \[x \geq x_{i,k+1} \].
\end{cases}
\]

Actually, \( l_2 \) is a piecewise linear function on \( C \) and it equals \( l_1 \) on \( B \). Moreover, \( \|l_2(x, z) - l_2(x_{i,j}, z_i)\| \leq \eta|x - x_{i,j}| \) holds, because if \( j = k + 1 \), then \( \|l_2(x, z) - l_2(x_{i,j}, z_i)\| = 0 \) by definition of \( l_2 \), and if \( j \leq k \), then

\[
\|l_2(x, z) - l_2(x_{i,j}, z_i)\| = \left|l_{1}(x_{i,j-1}, z_i) - l_{1}(x_{i,j-1}, z_i)\right|
\]

\[
\frac{x - x_{i,j-1}}{x_{i,j} - x_{i,j-1}}(l_1(x_{i,j-1}, z_i) - l_1(x_{i,j-1}, z_i))
\]

\[
\leq \eta|x - x_{i,j}|.
\]

A similar reasoning can be used to prove \( \|l_2(x_{i,t}, z_i) - l_2(y, z_i)\| \leq \eta|y - y| \).

Let \( [x, z], [y, z] \) be vectors in \( C \), and without loss of generality, assume that \( x \geq y \) holds. Let \( j \) be the largest index satisfying \( x \geq x_{i,j} \) and let \( t \) be the smallest index satisfying \( y \leq x_{i,t} \). Using (21) and the inequality \( \|l_2(x, z_i) - l_2(x_{i,j}, z_i)\| \leq \eta|x - x_{i,j}| \), it follows:

\[
\|l_2(x, z_i) - l_2(y, z_i)\|
\]

\[
\leq \|l_2(x, z_i) - l_2(x_{i,j}, z_i)\|
\]

\[
+ \|l_2(x_{i,j}, z_i) - l_2(x_{i,t}, z_i)\|
\]

\[
+ \|l_2(x_{i,t}, z_i) - l_2(y, z_i)\|
\]

\[
\leq \eta|x - x_{i,j}| + \left|\|l_1(x_{i,j}, z_i) - l_1(x_{i,t}, z_i)\| + \eta|x_{i,t}-y|\right|
\]

\[
\leq \eta|x - x_{i,j}| + \eta|x - x_{i,j}| - |x_{i,t}-y| = \eta|x - y|
\]

which implies that \( l_2 \) satisfies (21) on \( C \).

Step 3—Extending \( l_2(x, z) \) to the Entire \( R \times R^b \): Let \( V \) be the vertices of a hypercube \( H \) in \( R^b \) that contains the vectors in \( M \) as interior points. By some results shown in \([35]\), \( H \) can be partitioned, by a process called triangulation, into \( b \)-simplices having \( M \cup V \) as vertices and such that no vector of \( M \cup V \) is an interior point of a simplex. A \( b \)-simplex is a geometric figure having \( b + 1 \) vertices and it is a generalization of a triangle in the domain \( R^b \). Each point of a simplex can be obtained as a linear combination of its vertices. Thus, for any \( z \in H \), let us denote by \( S_z \subseteq M \cup V \) the set of the \( b + 1 \) vertices of the simplex where \( z \) is included. Since, a simplex is the convex hull of its vertices, there exist \( b + 1 \) positive reals \( \alpha_{z, \nu} \in R \), such that

\[
z = \sum_{v \in S_z} \alpha_{z, \nu} \cdot v \sum_{v \in S_z} \alpha_{z, \nu} = 1.
\]

The function \( l_2 \) is defined on the entire \( R \times R^b \) as

\[
l_3(x, z) = \begin{cases} \sum_{v \in S_z} \alpha_{z, \nu} \cdot l_2(x, \nu) & \text{if } z \in H \\[0, \]
if \[z \notin H \].
\end{cases}
\]

Note that \( l_3 \) is a linear function on each simplex and interpolates \( l_2 \) on the vertices. Thus, \( l_3 \) is piecwise continuous on \( H \). Moreover, \( l_3 \) is 0 on the faces of \( H \) and it is 0 outside \( H \). Thus, \( l_3 \) is piecwise continuous on \( R \times R^b \). Finally, by simple algebra

\[
\|l_3(x, z) - l_3(y, y)\|
\]

\[
= \sum_{v \in S_z} \alpha_{z, \nu} \cdot \left[l_2(x, \nu) - l_2(y, \nu)\right]
\]

\[
\leq \sum_{v \in S_z} \alpha_{z, \nu} \cdot \left[l_2(x, \nu) - l_2(y, \nu)\right]
\]

\[
\leq \sum_{v \in S_z} \alpha_{z, \nu} \cdot \eta|x - y|
\]

\[
= \eta|x - y|
\]

which implies that \( l_3 \) satisfies (21) for any \( [x, z] \in R \times R^b \).

Step 4—Approximating \( l_3 \) by a Differentiable Function: In the following, \( \xi \) will denote an infinitely differentiable probability distribution. We further assume that the support of \( \xi \) is inside the unit ball, i.e., \( \xi(x, z) = 0 \) if \( \|x[z]\| \geq 1 \) and \( \xi \) is not null in \((0,0)\). Finally, the constants \( L \) and \( P \) are specified as follows:

\[
L = \max_{x \in R^b, \|z\| \leq \delta} \frac{\|l_3(x, z) - \xi(x, z)\|}{\|x - y\|}
\]

\[
P = 2 \max_{x \in R^b, \|z_1, z_2, z_3, z_4 \| \leq \delta} \frac{\|x_{i,j} - x_{k,\nu}, z_{i} - z_{k}\|}{\|x - y\|}
\]

Function \( l_4 \) will be an infinitely differentiable function that approximates \( l_3 \). Let us consider a smoothing operation on \( l_3 \) as follows:

\[
l_3(x, z) = \int l_3(x - \mu, z - \mu) \sigma_\delta(\mu, z) d\mu
\]

where \( \delta \) is a positive real and the smoothing function \( \sigma_\delta \) is defined as \( \sigma_\delta(x, z) = \delta^{b+1} \xi(x, z/\delta) \). According to well-known results on convolutions \([18]\), \( l_3 \) is an infinitely differentiable
function and \( \lim_{x \to \infty} b_i = l_3 \) uniformly. Since the convergence is uniform, there exists \( \delta \) such that

\[
\max_{x, z} |l_i^3(x, z) - l_3(x, z)| \leq \frac{\eta P \cdot \xi(0, 0)}{2L}.
\]

(26)

Thus, we define \( l_4 = l_i^3 \). Finally, note that by (23)

\[
|l_i(x, z) - l_i(y, z)|
= \int \left( l_3(x - \xi, z - \xi) - l_3(y - \xi, z - \xi) \right) \sigma(z, \xi) d\xi d\xi \\
\leq \eta |x - y| \cdot \int \sigma(z, \xi) d\xi d\xi \\
= \eta |x - y|
\]

holds, so that \( l_4 \) fulfills (21) on \( \mathbb{R} \times \mathbb{R}^6 \).

Step 5—Adjust the Function on \( A \) for an Interpolation: Note that \( l_4 \) is differentiable, but it does not interpolate \( l \) on \( A \) anymore. Function \( l_5 \) will be an infinitely differentiable map that interpolates \( l \) on \( A \). More precisely, \( l_5 \) is built by slightly changing \( l_4 \) in the neighborhood of the points of \( A \)

\[
l_5(x, z) = l_4(x, z) + \sum_{[x_i, z_i] \in A} \frac{\xi(0, 0)}{\xi(0, 0)} \cdot \xi \left( \frac{x - x_i, z - z_i}{P} \right)
\]

Note that, since \( \xi \) is null outside the unit ball and \( P \) is twice the maximal distance of the points in \( A \) [see (25) and (24)], then \( \xi((x - x_i)/P, (z - z_i)/P) > 0 \) holds only if \([x_i, z_i]\) is the point of \( A \) closest to \([x, z]\). Thus, for any \([x, z]\), at most one term of those involved in the sum of (28) is nonnull. Since \([x_i, z_i] \in A\) is the closest point to itself, then

\[
l_5(x_i, z_i) = l_4(x_i, z_i) + \sum_{[x_i, z_i] \in A} \frac{\xi(0, 0)}{\xi(0, 0)} \cdot \xi \left( \frac{x - x_i, z - z_i}{P} \right)
\]

holds.

Finally, let \([x, z]\), and \([y, z]\) be vectors in \( \mathbb{R} \times \mathbb{R}^6 \). Then, by definition of \( l_5 \) and (27) and (26)

\[
|l_5(x, z) - l_5(y, z)|
= \left| l_4(x, z) - l_4(y, z) \right|
+ \sum_{[x_i, z_i] \in A} \left( \frac{\xi(0, 0)}{\xi(0, 0)} \cdot \xi \left( \frac{x - x_i, z - z_i}{P} \right) \right)
+ \sum_{[x_i, z_i] \in A} \left( \frac{\xi(0, 0)}{\xi(0, 0)} \cdot \xi \left( \frac{y - x_i, z - z_i}{P} \right) \right)
\]

Again, since \( \xi \) is null outside the unit ball and \( P \) is twice the maximal distance of the points in \( A \), there are at most two \([x_i, z_i] \in A \) for which \( \xi((x - x_i)/P, (z - z_i)/P) - \xi((y - x_i)/P, (z - z_i)/P) \neq 0 \) holds. Moreover, the definition of \( L \) implies \( \xi((x - x_i)/P, (z - z_i)/P) - \xi((y - x_i)/P, (z - z_i)/P) \leq \left| \frac{L}{P} \right| |x - y| \). Thus

\[
|l_5(x, z) - l_5(y, z)| \leq \eta |x - y| + \eta |x - y| = (\eta + \vartheta) |x - y|
\]

and Lemma 7 has been proved with \( \overline{l} = l_5 \).

Proof of Lemma 5: Now, we can proceed with the proof of Lemma 5. To avoid confusion, let us use an alternative notation to represent the function \( h \) in (16): \( l \) is

\[
l(y, l_{n-1-u}) = h(i, l_{n}, l_{n}, l_{i}, l_{n-1-u}, y, l_{n-1-u})
\]

where \( l_{n-1-u} \) collects into a vector the values \( i, l_{n-1-u}, l_{n-1-u}, l_{n-1-u}, y, l_{n-1-u} \). Note that according to the specification of \( h \), function \( l \) is defined only for the labels and the unfolding tree of a node of the graphs \( G_1, \ldots, G_v \) of Section B2 of the Appendix. By Lemma 6

\[
|l(x, l_{n-1-u}) - l(y, l_{n-1-u})| \leq \frac{\gamma}{1 - \gamma} |x - y|
\]

holds for any \( x = \nabla(T_{u_i}^{l_{n}}), y = \nabla(T_{u_i}^{l_{n}}) \), \( d_1, d_2 \in \mathbb{R} \). Moreover, by Lemma 7, \( l \) can be extended to an infinitely differentiable function \( \overline{l} \) that satisfies

\[
|\overline{l}(x, l_{n-1-u}) - \overline{l}(y, l_{n-1-u})| \leq \left( \frac{\gamma}{1 - \gamma} + \vartheta \right) |x - y|
\]

for any positive real \( \vartheta \), any \( x, y \in \mathbb{R} \), and any \( l_{n-1-u} \in \mathbb{R}^b \), \( b = 2|N| + 2E \). Thus, let \( f \) be defined as in (16), with its parameters being any value in the corresponding Euclidean spaces, i.e.,

\[
f \left( l_{n}, y_{n}[n], l_{n}[n], y_{n}[n], l_{n}[n], \right.
\]

for any \( y \in \mathbb{R}^a, a = s|N|, \) and any \( l \in \mathbb{R}^b, b = 2|N| + 2E \). Here, \( \overline{h} \) is the extension of \( h \) represented by \( \overline{l} \).

It is clear that function \( f \) fulfills point 1) of Lemma 5 by definition of \( \overline{l} \). On the other hand, by (28)

\[
|f \left( l_{n}, l_{n}[n], y_{n}[n], l_{n}[n], l_{n}[n], \right. - f \left( l_{n}, l_{n}[n], y_{n}[n], l_{n}[n], l_{n}[n], \right. |
\]

for any \( y \in \mathbb{R}^a, a = s|N|, \) and any \( l \in \mathbb{R}^b, b = 2|N| + 2E \). Here, \( \overline{h} \) is the extension of \( h \) represented by \( \overline{l} \).
Thus, if \( F \) is the global transition function corresponding to \( f \), then

\[
\|F(x, I) - F(y, I)\|_1 \\
\leq \sum_{n} |f(l_n, l_{x[n]}, x_{x[n]}, l_{x[n]}) - f(l_n, l_{y[n]}, y_{x[n]}, l_{x[n]})| \\
\leq \frac{\gamma}{1 - r^\gamma} \|x - y\|_1
\]

holds for any \( x, y \in \mathbb{R}^n \). Thus, if \( F \) is the global transition function corresponding to \( f \), then

\[
\|F(x, I) - F(y, I)\|_1 \\
\leq \sum_{n} |f(l_n, l_{x[n]}, x_{x[n]}, l_{x[n]}) - f(l_n, l_{y[n]}, y_{x[n]}, l_{x[n]})| \\
\leq \frac{\gamma}{1 - r^\gamma} \|x - y\|_1
\]

holds, and hence point 2) of Lemma 5 has been proved.

\[\square\]

C. Proof of Corollary 2: Connectionist Implementation of Positional GNNs

Let \( \phi_{f, g} \) denote the function realized by an GNN, where \( f \) and \( g \) are the local transition and local output functions, respectively. Moreover, for any function \( I \), let \( \|I\|_\infty \) represent the superior norm, i.e., \( \|I\|_\infty = \sup_{x \in \mathbb{R}} |I(x)| \). Lemma 8 proves that \( \phi_{f, g} \) depends continuously on \( f \) and \( g \) w.r.t. the superior norm.

Lemma 8: Let \( \phi_{f, g} \) be the function realized by an GNN. Suppose that \( f \) and \( g \) are continuously differentiable, \( g \) has a bounded support, and the global transition function \( F \) is a contraction map. Then, for any real \( \zeta > 0 \), there exist two reals \( \delta_f, \delta_g > 0 \) such that

\[
\|\phi_{f, g} - \phi_{\tilde{f}, \tilde{g}}\|_\infty \leq \zeta
\]

holds for any \( \phi_{\tilde{f}, \tilde{g}} \) implemented by a GNN, provided that the corresponding global transition function \( \tilde{F} \) is a contraction, and the local transition and local output functions fulfill

\[
\|g - \tilde{g}\|_\infty \leq \delta_g \quad \text{and} \quad \|f - \tilde{f}\|_\infty \leq \delta_f
\]

respectively.

Proof: Since \( g \) is continuous and has a bounded support, then it is equicontinuous. Moreover, also \( G \) is equicontinuous, because it is built by stacking copies of \( G \). Thus, there exists a real \( \Delta > 0 \) such that \( |x - \tilde{x}| \leq \Delta \) implies \( |G(x, I_N) - G(\tilde{x}, I_N)| \leq \zeta/2 \), for any \( x, \tilde{x} \).

Let us define \( \delta_f = ((1 - \eta) \Delta)/\|M\|_\infty \). This implies \( |f(x, I) - \tilde{f}(x, I)| \leq \zeta/2 \), for any \( x \).

Moreover, assume that \( F \) and \( \tilde{F} \) consist of stacking copies of \( f \) and \( \tilde{f} \), respectively, then \( \|F - \tilde{F}\|_\infty \leq \Delta \).

\[\square\]

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Moreover, since, in our setting, all the norms are equivalent, there exists a constant $Q$ such that
\[
\left\| \frac{\partial F}{\partial x}(G, l_N) \right\| \leq \left\| \frac{\partial F}{\partial x} + \frac{\partial F}{\partial x} - \frac{\partial F}{\partial x} \right\| + \frac{\eta}{2} + Q \left\| \frac{\partial F}{\partial x} - \frac{\partial F}{\partial x} \right\|_1 \leq \frac{\eta}{2} + Q \vartheta_2,
\]
As a consequence, it is sufficient to set $\vartheta \leq \eta/(2Q)$ in order to ensure that $\bar{F}$ is a contraction map (with contraction constant smaller than $\eta$). Thus, the corollary is shown to be true.

D. Proof of Theorem 3 and Corollary 3: Approximation by Nonpositional GNNs and the Connectionist Implementation

The proof of Theorem 3 follows the same reasoning as the proof of Theorem 2 with few minor differences in the definition of the function $\tilde{\phi}$ (Step 1 in Section B2 of the Appendix) and in the demonstration of the existence of a decoding function $\tilde{\nabla}^{-1}$ (Step 2 in Section B2 of the Appendix). In fact, in the definition of $\tilde{\phi}$, we must take into account that the processed graphs are nonpositional. Such a difference can be overcome by discarding the neighbor position from the input parameters of $\tilde{\phi}$.19 Thus, $\tilde{\phi}$ will be defined as a function that is monotonically increasing w.r.t. $d$ and produces a different integer $\tilde{\phi}(L_n, l_{(n,u)}, T^d_n, l_u)$ for each different value of $L_n, l_{(n,u)}, T^d_n$, and $l_u$.

Moreover, also the proof of the existence of a decoding function $\tilde{\nabla}^{-1}$ must be changed due to the different definition of $\tilde{\phi}$, and, as a consequence, of $\nabla$. However, an inspection of the proof indicates that the new definition of $\tilde{\nabla}$ affects only the maximum coefficient $B$ of the polynomial $\xi(T^d_n) - \xi(T^d_{n+1})$. In fact, $B$ was equal to 1 in Theorem 2, whereas it will be shown that $B \leq r$ in the current case. On the other hand, $B$ affects only Lemma 4, which still holds if $B \leq r$, because for the lemma to be true, it is sufficient that $0 < \gamma \leq 1/(2B)$ holds and, in this case, we have $0 < \gamma = Q/(2r(1 + Q)) < 1/(2r) = 1/(2B)$.

Thus, in order to prove Theorem 3, we have only to demonstrate $B \leq r$. Note that each neighbor of $n$ is represented by a term of the polynomial $\xi(T^d_n)$. In this case, it is different from Theorem 2 in that several children may be represented by the same term since the position of the child is not considered. More precisely, this happens when two neighbors $n_l$ and $n_j$ of $n$ have the same unfolding tree, i.e., $T^d_{n_l} = T^d_{n_j}$. Intuitively, such an occurrence is not a problem, since the coefficient corresponding to each term of $\xi(T^d_n)$ will count the number of subtrees of a given “type” and such information is sufficient to reconstruct the original nonpositional tree $T^d_n$. Formally, since $B$ is the maximum coefficient of the polynomial $\xi(T^d_n) - \xi(T^d_{n+1})$, $B$ cannot be larger than the maximum number of possible trees $T^d_{n+1}$, which is smaller than the number of neighbors of $n$. As a consequence, $B \leq \max[n] \leq r$ holds.

Finally, Corollary 3 can be demonstrated using the same argument used in the proof of Corollary 2. In fact, the proof of Corollary 3 shows that a GNN can approximate another GNN, provided that we can approximate up to any degree of precision the transition function $f$ and its derivatives by a network in $Q$. Similarly, in nonpositional GNNs, the function $h$ is approximated by a network in $Q$. It turns out that, for each $n$
\[
\left\| \tilde{f}(L_n, l_{(n,u)}, x_{(n,u)}, l_{(n,u)}) - f(L_n, l_{(n,u)}, x_{(n,u)}, l_{(n,u)}) \right\|
\]
holds, where $\tilde{h}$ is the function implemented by the neural network, $\tilde{f}$ the corresponding transition function, and $\vartheta_1$ is a bound on the achievable accuracy. Since the accuracy is proportional to the number of neighbors, it may appear that $\tilde{f}$ cannot be approximated up to any desired accuracy. On the contrary, we can observe that the function implemented by the GNN does not actually approximate the target function $\tau$ on the whole domain $D$, but only on graphs having a finite set of structures as defined by Theorem 5. Thus, we can concentrate our attention only on those graphs and we can assume that $\max[n]$ is bounded. As a consequence, $\tilde{f}$ can be approximated up to any degree of precision by implementing $h$ with a network in $Q$ and a similar reasoning applies also to the approximation of the Jacobian of $f$.

E. Proof of Theorem 4: $\mathcal{H}(\mathcal{D}) \subseteq \mathcal{F}(\mathcal{D})$

This theorem is proved for positional GNNs. The demonstration of the other cases follows the same reasoning. Let $f$ and $g$ be, respectively, the local transition and output functions of the GNN, and consider the following:
\[
x_n(t + 1) = f(L_n, l_{(n,u)}, x_{(n,u)}(t), l_{(n,u)}), \\
o_n(t + 1) = g(x_n(t + 1), l_n), \quad n \in \mathcal{N}
\]
where $x_n(0) = 0$ holds, for each $n$. In the following, it is shown by an induction argument on $t$ that there exists a function $\mathcal{H}$ such that $x(t) = \mathcal{H}(T^d_t)$ for $t \geq 1$. Note that this immediately implies that the theorem is true, since we can define a function $\kappa$ such that
\[
\kappa(T_n) = \lim_{t \to \infty} g(\mathcal{H}(T^d_n), l_n) \\
= \lim_{t \to \infty} g(x_n(t - 1), l_n) = \varphi(G, n)
\]
that satisfies the hypothesis of Theorem 1.

The induction argument goes as follows.

Base: $t = 1$.

The state $x_n(1)$ is computed by applying $f$ on $L_n, l_{(n,u)}, x_{(n,u)}, l_{(n,u)}$. All this data belong to $T^2_n$, so that we can define a function $\mathcal{H}$ such that
\[
\mathcal{H}(T^2_n) = f(L_n, l_{(n,u)}, 0, l_{(n,u)}),
\]
holds.

Induction: $t > 1$.19
Note that $x_u(t)$ is calculated from $l_{\omega t}, x_{\nu t}[n](t)$, $l_{\nu t \omega} l_{\nu t n}$, $r(t)$. By using the induction argument, there exists $r_u$ such that $x_u(t-1) = r(T_u)$ holds, for each $u \in \Omega$. Thus, $x_u(t)$ depends on $l_{\omega t}, l_{\nu t \omega}, l_{\nu t n}$ and all the $T_u$. Since such information is contained in $T_n$, we can define

$$r(T_n) = f\left(l_{\omega t}, l_{\nu t \omega}, l_{\nu t n}, T_n, T_n, T_n, \ldots, T_n\right)$$

where $r(T_n)$ is a vector obtained by stacking all the $r(T_u), u \in \Omega$. $\blacksquare$

REFERENCES


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