A Comparison between Recursive Neural Networks and Graph Neural Networks

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Abstract—Recursive Neural Networks (RNNs) and Graph Neural Networks (GNNs) are two connectionist models that can directly process graphs. RNNs and GNNs exploit a similar processing framework, but they can be applied to different input domains. RNNs require the input graphs to be directed and acyclic, whereas GNNs can process any kind of graphs. The aim of this paper consists in understanding whether such a difference affects the behaviour of the models on a real application. An experimental comparison on an image classification problem is presented, showing that GNNs outperforms RNNs. Moreover the main differences between the models are also discussed w.r.t. their input domains, their approximation capabilities and their learning algorithms.

I. Introduction

In several application domains the data of interest can be naturally described by trees or graphs. In fact, nodes can be used to represent objects while edges intuitively define the relationships between them. These applications comprise problems from pattern recognition, natural language processing, Web page scoring and relational learning [1]–[6].

Classical graph theory includes several theoretical studies and ad-hoc methods that deal with graph classification (or regression), graph matching, link prediction, and similarity measures for non-flat data [7]–[9].

Standard machine learning approaches require to map graphs or trees into simpler representations, like vectors or sequences of real numbers. However, their performances vary largely with the application at hand. In fact, the preprocessing phase is rather problem dependent and its design usually results in a time—consuming trial and error procedure. Moreover, the topological information inherently contained in structural representations might be partially lost or difficult to be exploited.

More recently, new connectionist models, capable to directly elaborate graphs and trees without a preprocessing phase were proposed [10]. They extend support vector machines [11]–[14], neural networks [15]–[18] and SOMs [19] to structured data. Such techniques have been applied successfully on several problems as logical term classification [20], chemical compound classification [21], logo recognition [2], [22], Web page scoring [23], and face localization [24].

The main idea is to obtain, adaptively and automatically, a flat representation of the symbolic and subsymbolic information collected in the structures. While in kernels for graphs the internal encoding is defined by the selected kernel,

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in neural network models it is automatically learned by examples. Finally, SOMs-SD differ from the other methods since they implement an unsupervised learning framework, instead of a supervised one.

Graph Neural Networks have been recently proposed to process very general types of graphs and can be considered an extension of RNNs. Actually, RNNs require input graphs to be directed and acyclic, while cyclic or non–directed structures must undergo a preprocessing phase. However, GNNs have not been widely tested yet, and it is unknown whether the performance of GNNs and RNNs is different in practical applications. This paper presents an experimental comparison between GNNs and RNNs. The two models are evaluated on a real–world computer vision problem, which consists in classifying a set of images. Moreover, the theoretical differences between the two models are also investigated, paying attention to their admitted input domains, their approximation capabilities and their learning algorithms.

The paper is organized as follows. In the next section, the notation adopted in the paper is introduced. Sect. III reviews the RNN and the GNN model. Sect. IV illustrates the real—world computer vision task and shows the results obtained by RNNs and GNNs. Finally, the conclusions are drawn in Sect. V.

II. NOTATION

A labeled graph (or graph) G is a quadruple $(N, E, \mathcal{L}, \mathcal{E})$, where N is the set of nodes (or vertices), and E is the set of edges between nodes: $E \subseteq \{(u,v)|u,v\in N\}$. Nodes and edges constitute the skeleton of the graph. The last two items, that associate vectors of real numbers of dimension respectively \mathbb{R}^{l_N} and \mathbb{R}^{l_E} to each node and edge, are a node-labeling function $\mathcal{L}: N \to \mathbb{R}^{l_N}$ and an edge-labeling function $\mathcal{E}: E \to \mathbb{R}^{l_E}$. Node labels are represented by l_n and edge labels by $l_{(u,v)}$. In the following, the symbol l with no further specification is the vector obtained by stacking together all the labels of the graph. Finally, the symbol $|\cdot|$ denotes the cardinality or the absolute value, according to whether it is applied to a set or to a number.

If the graph is directed, an edge (u, v) is an *ordered* pair of nodes, where u is the *father* and v the *child*. If the graph is undirected, the ordering between u and v in (u, v) is not defined, i.e. (u, v) = (v, u). A graph is called *acyclic* if there is no path, i.e. a sequence of connected edges, that starts and ends in the same node. Given a node n, the nodes adjacent to n (or neighbors of n) are those connected to it by an edge, and are represented by ne[n]. If the graph is

directed, the neighbors of n either belong to the set of its children (ch[n]), or to the set of its parents (pa[n]). Moreover, a graph can be said positional or non-positional according to whether a function π_n is defined for each node n that assigns a different position $\pi_n(u)$ to each neighbor $u \in \text{ne}[n]$. Thus, combining the properties described so far it is possible to specify various graph categories: Directed Positional Acyclic Graphs (DPAGs), Directed Acyclic Graphs (DAGs) and so on.

The goal of the proposed models consists in approximating a target function $\tau(G,n) \in \mathbb{R}^m$ that maps a graph G and one of its nodes n into a vector of real numbers¹. In graph classification, τ does not depend on n, i.e. only one target is given for each graph. In node classification problems, each node in a given set has a target to be approximated.

III. RECURSIVE AND GRAPH NEURAL NETWORKS

A very general framework for graph processing must implement a function φ that computes an output $\varphi(G,n)$ for each pair (G,n). The main idea is to derive a flat description of the information related to each node n. A node represents an object of the domain of interest: its description is a vector of real numbers called *state* and denoted by $x_n \in R^s$, where the state dimension s is a predefined parameter. In order to obtain a distributed processing framework, the states are computed locally at each node. A reasonable choice is to design x_n as the output of a parametric function f_w (called *state transition function*), that depends on the node label l_n and on the relationships between n and its neighbors

$$\boldsymbol{x}_n = f_{\boldsymbol{w}}(\boldsymbol{l}_n, \boldsymbol{x}_{\text{ne}[n]}, \boldsymbol{l}_{\text{ne}[n]}, \boldsymbol{l}_{(n,\text{ne}[n])}), \quad n \in \boldsymbol{N}, \quad (1)$$

where $\operatorname{ne}[n]$ is the set of neighbors of node n, $x_{\operatorname{ne}[n]}$ and $l_{\operatorname{ne}[n]}$ are the sets containing the states and the labels of the nodes in $\operatorname{ne}[n]$ respectively, and $l_{(n,\operatorname{ne}[n])}$ is the set of the edge labels between n and its neighbors (Fig. 1). Once each node has a vectorial representation, it can also be assigned an output o_n , evaluated by another parametric function g_w , called *output function*

$$o_n = g_w(x_n, l_n), \quad n \in \mathbb{N}.$$
 (2)

Eqs. (1) and (2) define a method to produce an output $o_n = \varphi_w(G, n)$ for each node of the graph G. Moreover, the symbolic and subsymbolic information related to the nodes is indeed automatically encoded into a vector by the state transition function.

Given a graph G, the computation can be graphically visualized substituting all of the nodes with "units" that compute the function f_w and are connected according to graph topology. The obtained network is called the *encoding network* (Fig. 2). In this way, the encoding network has the same skeleton as the input graph. Each "f-unit" is also connected to another unit that implements the output function g_w . Since the same parametric functions are applied to all the nodes, the units of the same type share the same set of parameters.

¹More precisely, in classification problems, the codomain of τ is \mathbb{N}^m , whereas in regression problems the codomain is \mathbb{R}^m .

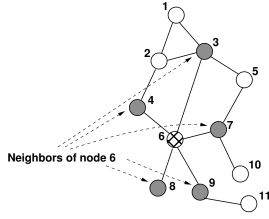


Fig. 1. The state x_n depends on the information in its neighborhood.

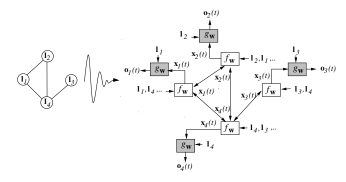


Fig. 2. Overview of the graph processing. Notice that the skeleton of the input graph defines the interconnection among the processing units.

Let F_w and G_w be the vectorial functions obtained stacking all the instances of f_w and g_w , respectively. Then Eqs. (1) and (2) can be rewritten as

$$x = F_{w}(x, l) \tag{3a}$$

$$o = G_w(x, l) \tag{3b}$$

where l represents the vector containing all the labels and x collects all the states. Equation (3a) defines the global state x, while Eq. (3b) computes the output. It is worth to mention that Eq. (3a) is recursive w.r.t. the state x, thus x is well defined only if Eq. (3a) has a unique solution. In conclusions, the viability of the method depends on the particular implementation of the transition function f_w .

In our supervised framework, for a subset $S \subseteq N$ of nodes, called supervised nodes, a target value t_n is defined. Thus an error signal can be specified,

$$e_{\mathbf{w}} = \sum_{i \in \mathbf{S}} (\mathbf{t}_i - \varphi_{\mathbf{w}}(\mathbf{G}, i))^2. \tag{4}$$

This signal drives an error backpropagation procedure that adapts the parameters of $f_{\boldsymbol{w}}$ and $g_{\boldsymbol{w}}$ so that the function realized by the network approximates the targets, i.e. $\varphi_{\boldsymbol{w}}(\boldsymbol{G},i) \approx \boldsymbol{t}_i, \ \forall i \in \boldsymbol{S}$.

In practice, Eq. (1) is well suited to process positional graphs, since each neighbor position can be associated to a specific input argument of function f_w . In non-positional graphs, this scheme introduces an unnecessary constraint, since neighbors should be artificially ordered. A reasonable solution consists in calculating the state x_n as a sum of "contributions", one for each of its neighbors. Thus, state transition function can be rewritten as

$$\boldsymbol{x}_{n} = \sum_{i=1}^{\left| \text{ne}[n] \right|} h_{\boldsymbol{w}} \left(\boldsymbol{l}_{n}, \boldsymbol{x}_{\text{ne}_{i}[n]}, \boldsymbol{l}_{\text{ne}_{i}[n]}, \boldsymbol{l}_{(n, \text{ne}_{i}[n])} \right), \quad n \in \boldsymbol{N}$$
(5)

where $ne_i[n]$ is the *i*-th neighbor and |ne[n]| is the number of neighbors of n.

Several possible implementations of the functions $f_{\boldsymbol{w}}$ (or $h_{\boldsymbol{w}}$) and $g_{\boldsymbol{w}}$ can be selected. RNNs and GNNs differ in the implementation of the state transition function $f_{\boldsymbol{w}}$ and in the class of graphs that can be processed.

A. Recursive Neural Networks

Recursive Neural Networks (RNNs) were originally proposed in [15], [16], [20]. In RNNs, the problem posed by the solution of Eq. (3) is faced enforcing the following two constraints:

- Input graphs are acyclic;
- The state x_n of node n depends only on its children, instead of its neighbours, i.e. Eq. (5) is replaced by

$$\boldsymbol{x}_{n} = \sum_{i=1}^{\left| \operatorname{ch}_{[n]} \right|} h_{\boldsymbol{w}} \left(\boldsymbol{l}_{n}, \boldsymbol{x}_{\operatorname{ch}_{i}[n]}, \boldsymbol{l}_{\operatorname{ch}_{i}[n]}, \boldsymbol{l}_{(n, \operatorname{ch}_{i}[n])} \right)$$
(6)

In this way, the definition of x is not recursive and Eq. (3) has a unique solution. In fact, the state x can be computed as follows: for each leaf node n, x_n can be computed directly, assuming that $x_{\operatorname{ch}[n]}$ is assigned a predefined value x_0 ; for any other node, x_n is computed by Eq. (6) as soon the states of the children $x_{\operatorname{ch}[n]}$ are available.

In graph classification problems, each graph must contains a node r, called the *supersource*, from which there is a path to all the other nodes². The supersource is the only supervised node, i.e. a target t_r is assigned to it. Actually, the supersource is the best candidate to collect all the information embedded in the structure and to compute a suitable output.

The original RNN model can only process Directed Positional Acyclic Graphs (DPAGs). More recently, an extended model was proposed that can cope with non–positional acyclic graphs with labeled edges (i.e. DAGs) [25]. Such a model uses the transition function defined in Eq. (6), where function h_w is implemented by the composition of two different parametric functions. The former function $\phi_w: \mathbb{R}^{(s+l_E)} \to \mathbb{R}^s$ computes the sum \overline{x}_n of the neighbor contributions to the state (Eq. (7)), while the latter

 $\rho_w : \mathbb{R}^{(s+l_N)} \to \mathbb{R}^s$ combines this value with the node label l_n to obtain the state of the node (Eq. (8))

$$\overline{\boldsymbol{x}}_{n} = \sum_{i=1}^{\left| \operatorname{ch}[n] \right|} \phi_{\boldsymbol{w}}(\boldsymbol{x}_{\operatorname{ch}_{i}[n]}, \boldsymbol{l}_{(n, \operatorname{ch}_{i}[n])})$$
 (7)

$$\boldsymbol{x}_n = \rho_{\boldsymbol{w}}(\overline{\boldsymbol{x}}_n, \boldsymbol{l}_n) \tag{8}$$

As mentioned before, the output is evaluated only at the supersource r,

$$o_r = g_{\boldsymbol{w}}(\boldsymbol{x}_r). \tag{9}$$

The parametric functions ϕ_w , ρ_w , and g_w can be implemented by a variety of neural network models (Fig. 3). As shown in Fig. 4, the corresponding encoding network is a feedforward network. Thus, the output o_r can be calculated starting from the leaves and proceeding toward the supersource node r, whereas the error is backpropagated from r to the leaves of the encoding network.

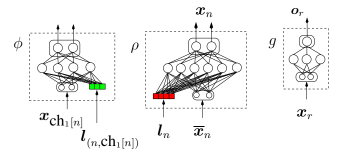


Fig. 3. Functions ϕ_w , ρ_w , and g_w may be implemented by a variety of neural network models.

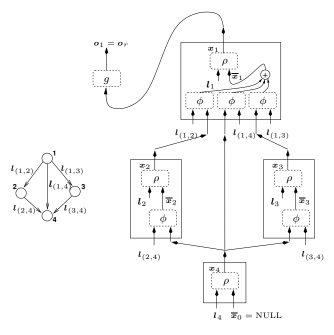


Fig. 4. An input graph (on the left) and its corresponding encoding network.

Finally, RNNs have been proved to be able to approximate in probability any function on positional trees [26] and positional and non-positional trees with labeled edges [25].

²If such a node does not exist it is always possible to add a dummy supersource node with the minimum number of edges [15].

In conclusions, the main limitations of RNNs w.r.t. the original framework are the restriction of the input domain to acyclic graphs and the dependence of a state x_n just on the children.

B. Graph Neural Networks

Graph Neural Networks (GNNs) have been recently proposed [17], [18] as an evolution of Recursive Neural Networks and have been conceived to fully implement the framework presented in Sect. III, in particular Eqs. (5) and (2). Thus, the input domain is the whole class of general graphs, including cyclic, acyclic, positional and non–positional, with labeled nodes and edges. In particular, this model can directly deal with undirected graphs.

Graph processing is performed associating to each graph G its encoding network, as shown in Fig. 2. As in RNNs, there could be many possible ways to implement the two functions f_w and g_w . In the following we assume to use three-layered feedforward neural network (i.e. with one hidden layer)³, since they have been proved to be universal approximators [27], [28]. However, this is not a sufficient condition for GNNs to perform universal approximation on graphs.

As already remarked, in case of cyclic graphs the system described by Eqs. (3) is well defined only if state updating has a unique solution and this depends on the properties of the transition function $f_{\boldsymbol{w}}$. The proposed solution defines $f_{\boldsymbol{w}}$ such that the global function $F_{\boldsymbol{w}}$ results to be a *contraction mapping* w.r.t. the state \boldsymbol{x} . A generic function $\rho: \mathbb{R}^n \to \mathbb{R}^n$ is a contraction mapping w.r.t. a vector norm $\|\cdot\|$, if it exists a real number μ , $0 \le \mu < 1$, such that for any $\boldsymbol{x}_1, \boldsymbol{x}_2 \in \mathbb{R}^n$, $\|\rho(\boldsymbol{x}_1) - \rho(\boldsymbol{x}_2)\| \le \mu \|\boldsymbol{x}_1 - \boldsymbol{x}_2\|$. If $F_{\boldsymbol{w}}$ is a contraction mapping, it is possible to prove that Eq. (3a) has a unique solution [29].

Since the function f_w is implemented using a feedforward neural network, this constraint can be satisfied limiting the range of values that weights w can assume. In fact it has been proved that if the norm of the jacobian matrix of a vectorial function is bounded by a number smaller than 1, then the function is a contraction mapping. In our implementation we used the one-norm, defined, for a generic matrix A, as $\|A\|_1 = max_j(\sum_i |A_{ij}|)$. Forcing the weights of the network to sufficiently small values surely satisfies the requirement. This approach, however, has one important drawback. If the weights are no more allowed to vary arbitrarily one can suspect that the global approximation property may be lost. For this reason it is preferable to add a penalty term to the error function,

$$e_{w} = \sum_{i \in S} (t_{i} - \varphi_{w}(G, i))^{2} + \beta L \left(\left\| \frac{\partial F_{w}}{\partial x} \right\|_{1} \right), \quad (10)$$

where the function L is specified as follows

$$L(y) = \begin{cases} (y - \mu)^2 & \text{if } y > \mu \\ 0 & \text{elsewhere} \end{cases}$$

and μ is the desired contraction constant. The value β in Eq. (10) is an user definable parameter balancing the relative importance of the error on supervised patterns and the error w.r.t. the chosen contraction constant μ .

It should be clear that the greater is β , the harder is the constraint on the norm of the jacobian matrix of F_w . At the same time, no guarantee is given that the norm remains always strictly under μ . This is not a major problem, as actually the risk of instability of Eq. (3) is very low even if the norm significantly overcomes the value of 1. The reason is that the bound on the norm of the jacobian matrix gives only a sufficient condition for F_w to be a contraction mapping.

1) The learning algorithm: The Banach fixed point theorem suggests a simple algorithm to compute the fixed point of Eq. (3a). It states that if ρ is a generic contraction mapping, then the dynamical system $x(t+1) = \rho(x(t))$, where x(t) denotes the t-th iterate of x, converges exponentially fast to the solution of the equation $x = \rho(x)$ for any initial state x(0). Thus x_n can be obtained iteratively as

$$\boldsymbol{x}_n(t+1) = f_{\boldsymbol{w}}(\boldsymbol{l}_n, \boldsymbol{x}_{\mathsf{ne}[n]}(t), \boldsymbol{l}_{\mathsf{ne}[n]}, \boldsymbol{l}_{(n,\mathsf{ne}[n])}).$$

The simultaneous and repeated activation of the "f-units" in the encoding network (Fig. 2) produces the behavior described by the previous equation. Notice that there is no learning in this phase. Simply, the state vector x is evolved until it reaches its fixed point x^* . Then the "g-units" evaluate all the outputs o_n , that can be compared with targets t_n to derive an error signal as in Eq. (10).

The goal of the learning procedure is to adapt the parameters of the transition and the output functions for a better approximation of the targets. This can be accomplished using a gradient descent strategy, as in canonical backpropagation algorithm. Thus, for each parameter w_k , the quantity $\frac{\partial e_w}{\partial w_k}$ needs to be evaluated. As g_w is a standard feedforward neural network, the gradient of the error w.r.t. each component of its weight vector w can be easily obtained, together with the gradient $\frac{\partial e_w}{\partial x^*}$ that expresses the sensibility of the overall error to a perturbation of the fixed point of the state vector x. We can suppose that the system has reached its fixed point x^* at time T, i.e. $x^* \simeq x(T)$. The gradient of the error w.r.t. the parameters of function f_w can be obtained as

$$\frac{\partial e_{w}}{\partial w} = \frac{\partial e_{w}}{\partial x(T)} \frac{\partial x(T)}{\partial w}.$$
 (11)

Unfortunately, $\boldsymbol{x}(T)$ has an implicit dependence on the parameters \boldsymbol{w} , so the second factor of the right part of Eq. (11) cannot be evaluated directly. Instead, the error should be backpropagated from one time instant to the previous, accumulating the gradient until convergence. In fact $\boldsymbol{x}(T)$ depends on \boldsymbol{w} directly, but also through the state \boldsymbol{x} at previous time instants. Remembering that

$$\boldsymbol{x}(T) = \boldsymbol{F}_w(\boldsymbol{x}(T-1), \boldsymbol{l})$$

and using the chain rule, we have the following recursive

³A different implementation of transition and output functions is presented in [18].

equation

$$\frac{\partial \boldsymbol{x}(T)}{\partial \boldsymbol{w}} = \frac{\partial F_{\boldsymbol{\psi}}(\boldsymbol{x}(T-1), \boldsymbol{l})}{\partial \boldsymbol{\psi}} \Big|_{\boldsymbol{\psi} = \boldsymbol{w}} + \frac{\partial F_{\boldsymbol{w}}(\boldsymbol{x}(T-1), \boldsymbol{l})}{\partial \boldsymbol{x}(T-1)} \frac{\partial \boldsymbol{x}(T-1)}{\partial \boldsymbol{w}} \tag{12}$$

where term $\frac{\partial F_w(x(T-1),l)}{\partial x(T-1)}$ is simply the jacobian matrix of F_w w.r.t. its second argument, evaluated at the point x(T-1). The notation in Eq. (12) is not unnecessarily burdensome. In fact the symbol $\frac{\partial F_w(x(T-1),l)}{\partial w}$ is ambiguous in this context as it means the overall sensitivity of the steady state x(T) to a small variation of w, but also the partial derivative of the function F_w w.r.t. to its first argument w, regardless of the fact that even the second argument x(T-1) depends on x. For this reason, it is better to resort to a more formal notation to ensure clarity. Thus the symbol $\frac{\partial F_w(x(T-1),l)}{\partial w}$ has the former meaning while we introduce a formal parameter y when we need the partial derivative only w.r.t. to the first argument of F_w .

Suppose now that in the forward phase the state evolved for a long time after convergence. Obviously

$$\boldsymbol{x}(T+\overline{t}\,)\simeq\boldsymbol{x}(T) \qquad \forall \overline{t}>0$$

Gradient accumulation needs, in principle, to evaluate Eq. (12) backward until time $t=-\infty$. From a practical point of view, however, the series of the gradient values is convergent, as the generic term results from the multiplication of increasingly more jacobian matrices. If all these matrices have norm smaller than one, gradient accumulation can be stopped after some steps backward, as contributions become progressively smaller. The conclusion is that gradient computation is indeed feasible, given that the condition on the jacobian matrix norm is satisfied. Thus we can stop gradient accumulation at time instant t_0 , that we can safely assume to be posterior to convergence time T, i.e. $t_0 > T$. This allows to consider only the terms at time instant T,

$$\begin{split} J_{\boldsymbol{w}} & \stackrel{\circ}{=} & J_{\boldsymbol{w}}(T) = \frac{\partial F_{\boldsymbol{\psi}}(\boldsymbol{x}(T), \boldsymbol{l})}{\partial \boldsymbol{\psi}} \, \bigg|_{\,\boldsymbol{\psi} = \boldsymbol{w}} \\ J_{\boldsymbol{x}} & \stackrel{\circ}{=} & J_{\boldsymbol{x}}(T) = \frac{\partial F_{\boldsymbol{w}}(\boldsymbol{x}(T), \boldsymbol{l})}{\partial \boldsymbol{x}(T)} \end{split}$$

and Eq. (12) becomes

$$\frac{\partial x(T)}{\partial w} = J_w \{ 1 + J_x + (J_x)^2 + (J_x)^3 + \dots + (J_x)^{t_0} \}.$$

This intuition is very useful also to contain memory requirements, as there is no need to keep memory of the gradient values calculated at each time instant before state convergence. Gradient evaluation is represented schematically in Fig. 5.

The proposed algorithm is not completely new. In fact it combines the backpropagation through structure algorithm, adopted to train Recursive Neural Networks, and the Almeida–Pineda algorithm [30]–[32] (also called *Recurrent Backpropagation*). The latter is a particular version of the backpropagation through time algorithm which can be used

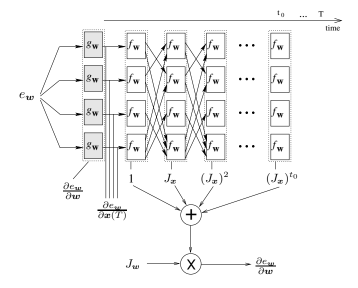


Fig. 5. Overall gradient is evaluated accumulating gradient contributions backward until time instant t_0 .

to train recurrent networks. In the Almeida–Pineda algorithm, input is kept constant and the network settles down to a fixed point for its dynamics. The aim is to move this fixed state toward a given target, backpropagating the error and accumulating the gradient exactly in the way seen before for GNNs. Moreover, as explained clearly in [32], Recurrent Backpropagation doesn't require the storage of any state history, assuming, as seen before, that the state equilibrium point was reached long before starting gradient backpropagation. In conclusion, our approach applies the Almeida-Pineda algorithm to the encoding network, where all instances of f_w and $g_{\boldsymbol{w}}$ are considered to be independent networks, but, as usual in Recursive Neural Networks, contributions for shared weights are combined. When the gradient is available, the weights can be updated using resilient backpropagation [33], a well-known variant of standard backpropagation that speeds up the convergence.

2) Some remarks on generalization capabilities: In this section we will report some results concerning the generalization capabilities of the GNN model. More details can be found in [34].

Given a graph G, the unfolding tree of depth $d T_n^d$ of one of its nodes n is the graph obtained "unfolding" G up to the depth d, using n as the starting point (Fig. 6).

Two nodes n, u are said to be unfolding equivalent $n \sim u$, if $T_n^d = T_u^d$ for every $d \geq 1$. In Fig. 6 the two nodes with label \boldsymbol{b} are unfolding equivalent, while the two nodes with label \boldsymbol{a} are not equivalent, as their trees already differ in the very first levels.

A function $p: \mathcal{D} \to \mathbb{R}^m$ is said to preserve the unfolding equivalence on \mathcal{D} , if $n \sim u$ implies $p(\mathbf{G}, n) = p(\mathbf{G}, u)$, for any patterns (\mathbf{G}, n) , (\mathbf{G}, u) that belong to \mathcal{D} .

The nodes of the two graphs on the left of Fig. 7 share the same unfolding tree. If a function assigns different values to them, f.i. the number of nodes of the graph to which they belong, it doesn't preserve the unfolding equivalence.

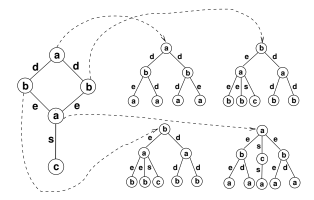


Fig. 6. A graph and the unfolding trees of depth 2 of its nodes.

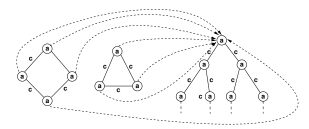


Fig. 7. The nodes of the two graphs on the left are all unfolding equivalent nodes.

Since GNNs adopt an inherently local computational scheme, they are able to approximate only functions that preserve unfolding equivalence. Noteworthy, also the converse is true. In fact it is possible to show that, given a function that preserves the unfolding equivalence, it always exists a GNN model capable of approximating it in probability to the desired degree of accuracy [34].

IV. EXPERIMENTAL RESULTS

RNNs and GNNs have been experimentally compared on an image classification problem which was based on images extracted from the Caltech benchmark database [35]. In the experiments, four classes have been considered, containing images of 'bottles', "camels", "guitars" and "houses". Two of the four categories, namely bottles and camels, are particularly difficult to be recognized, since they were originally collected using Google image search, without any kind of normalization.

For each class, a set of 350 random images were extracted from the original dataset. A half of those images consisted of positive examples of the category, the other half consisted of negative examples, i.e. images belonging to the other categories. Then, the set was subdivided into a training set, a validation set⁴ and a test set containing 150, 50 and 150 images, respectively.

A. From images to DAGs

Each image was preprocessed in order to obtain the Region Adjacency Graph (RAG) that represents it. A RAG is a

graph where nodes denote homogeneous regions of the image and edges stand for the adjacency relationships. Nodes and edges are labeled. Node labels contain geometric features of the corresponding regions (area, perimeter, orientation of the principal axes, and so on), while edge labels encode the mutual orientation of the two regions and the difference between their average colors.

RAGs were obtained by the following steps:

- The image was filtered using the Mean Shift algorithm [36]⁵.
- A k-means [37] color quantization procedure was performed to locate an initial big number of homogeneous regions.
- Some adjacent regions were merged to achieve the
 desired number of final regions. The procedure evaluated a dissimilarity function between all the pairs
 of regions and merged those that achieve the lowest
 values, updating the list at each fusion. The dissimilarity
 function has been chosen heuristically considering the
 distance between the average colors of the two regions
 and their dimension.

The RAGs obtained by the above procedure constituted the dataset used for the experiments with the GNNs. On the other hand, since RAGs are undirected graphs, RNNs are not able to process them. As in other applications of RNNs to image processing [24], [38], the RAGs were transformed into Directed Acyclic Graphs (DAGs). The transformation was accomplished by performing a breadth-first visit of the RAG, starting from the region which contained the central pixel of the image. Each time an edge was traversed in a certain direction, the undirected edge was replaced by an arc with that direction (Fig 8).

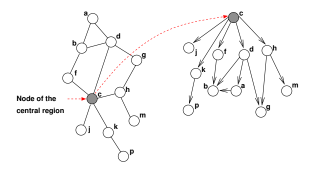


Fig. 8. The original RAG (on the left) and the corresponding DAG.

B. Network architectures and results

The experimentation consisted of a set of runs. Each experiment varies according whether a GNN or a RNN is trained, according to the category that is considered and to the network architecture that was used. In fact, every experiment concerns a category in 'bottles', "camels", "guitars"

⁴The validation set was exploited to choose the optimal network. In fact, the optimal network was the one that, during the training procedure, achieved the lowest error on the validation set.

⁵The main advantage of this technique is that the method was especially designed to preserve gradient discontinuities. Thus smallest textures, that are often over segmented by techniques that employ only color information, were heavily smoothed while significant edges remained almost crisp, easing region boundaries detection.

TABLE I The number of hidden units of the RNN and GNN functions, w.r.t. the three configurations small, medium and big

Model	Function	Configuration		
Model		small	medium	big
RNN	ϕ	10 hidden u.	11 hidden u.	12 hidden u.
	ρ	5 hidden u.	7 hidden u.	10 hidden u.
	g	5 hidden u.	7 hidden u.	10 hidden u.
GNN	f	5 hidden u.	7 hidden u.	10 hidden u.
	g	5 hidden u.	7 hidden u.	10 hidden u.

TABLE II

COMPARISON BETWEEN THE ACCURACIES OBTAINED BY RECURSIVE

NEURAL NETWORKS (RNN) AND GRAPH NEURAL NETWORKS (GNN)

IN THE IMAGE CLASSIFICATION PROBLEM

Class	Model	Configuration		
Ciass		small	medium	big
Bottles	RNN	69.33%	70.66%	68.66%
	GNN	76.67%	83.33%	84.67%
Camels	RNN	62.67%	62.67%	65.33%
Cameis	GNN	70.00%	73.33%	74.67%
Guitars	RNN	61.33%	62,67%	60.00%
Guitars	GNN	70.67%	70.00%	67.33%
Houses	RNN	80.67%	81.33%	79.33%
Houses	GNN	84.00%	83.33%	84.67%

and "houses" and the network is trained to decide whether the input graph belongs or not to the considered category.

Three different configurations, with an increasing number of weights, were experimented for each model. In order to make the comparison as fair as possible, the configurations were chosen so that GNNs and RNNs have the same number of weights (Tab. I). The three configurations will be denoted as *small*, *medium* and *big*.

Table II shows the accuracies achieved in the experiments. Here, the accuracy is defined as the percentage of correct predictions w.r.t. the total number of predictions. Table III shows the same results, taking also into account the different accuracies in the positive and negative examples. Finally, Table IV displays the *equal error rate* of the two models. The *equal error rate* is obtained plotting the accuracies on positive and negative examples of the test set varying the classification threshold (Fig. 9). The equal error rate is defined as the accuracy of the system when the number of errors in the two classes is equal.

The results show that GNNs almost always outperformed RNNs. Even if the two models are conceptually very similar, Graph Neural Networks have the advantage to process RAGs directly. In the transformation from RAGs to DAGs, the edges, which are naturally undirected, are replaced with directed ones, probably causing a loss of information. Moreover, the direction of the edges affects also the order of the computation which in RNNs must go from the leaves of the DAGs to the supersource. Those two factors have probably determined the better results obtained by GNNs.

 $\label{thm:comparison} \textbf{TABLE III}$ Comparison between the accuracies obtained by the models

W.R.T. NEGATIVE AND POSITIVE EXAMPLES. THE BEST CLASS ACCURACIES ARE SHOWN IN BOLD

Class	Model	Subclass	Configuration		
Class			small	medium	big
Bottles	RNN	positive	69.33%	81.33%	68.00%
		negative	69.33%	60.00%	69.33%
	GNN	positive	80.00%	69.33%	73.33%
		negative	73.33%	78.67%	80.00%
Camels	RNN	positive	82.67%	84.00%	92.00%
		negative	42.67%	41.33%	38.66%
	GNN	positive	73.33%	80.00%	81.33%
		negative	66.67%	66.67%	68.00%
	RNN	positive	82.66%	81.33%	74.67%
Guitars		negative	44.00%	44.00%	45.33%
Guitars	GNN	positive	62.67%	61.33%	60.00%
		negative	78.67%	78.67%	74.67%
Houses	RNN	positive	76.67%	85.33%	85.33%
		negative	84.67%	77.33%	73.33%
	GNN	positive	86.67%	85.33%	82.67%
		negative	81.33%	81.33%	86.67%

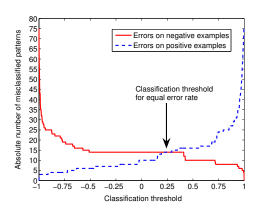


Fig. 9. Typical curves describing the absolute number of errors on the test set varying the classification threshold. Test set had 75 positive and 75 negative examples.

TABLE IV

COMPARISON BETWEEN THE EQUAL ERROR RATES OBTAINED BY

RECURSIVE NEURAL NETWORKS (RNN) AND GNNs IN THE IMAGE

CLASSIFICATION PROBLEM

Class	Model	Configuration		
Class		small	medium	big
Bottles	RNN	69.33%	68.00%	68.66%
	GNN	77.33%	77.33%	78.67%
Camels	RNN	58.67%	58.67%	58.67%
	GNN	69.33%	76.00%	73.33%
Guitars	RNN	58.67%	62.67%	60.00%
Guitars	GNN	68.00%	70.67%	69.33%
Houses	RNN	78.67%	78.67%	78.67%
Houses	GNN	81.33%	85.33%	82.67%

On the other hand, notice that GNNs require a longer training time due to need of iterating the computation of the states and of the gradients until a stable point is achieved.

V. Conclusions

This paper presented a comparison between two neural network models, RNNs and GNNs, capable of directly processing graphs with labeled nodes and edges. Their main differences and similarities w.r.t. input domains, approximation capabilities and learning algorithms have been discussed. A real—world image classification task showed that GNNs outperforms RNNs, both in terms of accuracies and equal error rates.

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