Convolutional Kernel Networks for Graph-Structured Data

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Abstract

We introduce a family of multilayer graph kernels and establish new links between graph convolutional neural networks and kernel methods. Our approach generalizes convolutional kernel networks to graph-structured data, by representing graphs as a sequence of kernel feature maps, where each node carries information about local graph substructures. On the one hand, the kernel point of view offers an unsupervised, expressive, and easy-to-regularize data representation, which is useful when limited samples are available. On the other hand, our model can also be trained end-to-end on large-scale data, leading to new types of graph convolutional neural networks. We show that our method achieves competitive performance on several graph classification benchmarks, while offering simple model interpretation. Our code is freely available at https://github.com/claying/GCKN.

1. Introduction

Graph kernels are classical tools for representing graph-structured data (see Kriege et al., 2020, for a survey). Most successful examples represent graphs as very-high-dimensional feature vectors that enumerate and count occurrences of local graph sub-structures. In order to perform well, a graph kernel should be as expressive as possible, i.e., able to distinguish graphs with different topological properties (Kriege et al., 2018), while admitting polynomial-time algorithms for its evaluation. Common sub-structures include walks (Gärtner et al., 2003), shortest paths (Borgwardt & Kriege, 2005), subtrees (Shervashidze et al., 2011), or graphlets (Shervashidze et al., 2009).

Graph kernels have shown to be expressive enough to yield good empirical results, but decouple data representation and model learning. In order to obtain task-adaptive representations, another line of research based on neural networks has been developed recently (Niepert et al., 2016; Kipf & Welling, 2017; Xu et al., 2019; Verma et al., 2018). The resulting tools, called graph neural networks (GNNs), are conceptually similar to convolutional neural networks (CNNs) for images; they provide graph-structured multilayer models, where each layer operates on the previous layer by aggregating local neighbor information. Even though harder to regularize than kernel methods, these models are trained end-to-end and are able to extract features adapted to a specific task. In a recent work, Xu et al. (2019) have shown that the class of GNNs based on neighborhood aggregation is at most as powerful as the Weisfeiler-Lehman (WL) graph isomorphism test, on which the WL kernel is based (Shervashidze et al., 2011), and other types of network architectures than simple neighborhood aggregation are needed for more powerful features.

Since GNNs and kernel methods seem to benefit from different characteristics, several links have been drawn between both worlds in the context of graph modeling. For instance, Lei et al. (2017) introduce a class of GNNs whose output lives in the reproducing kernel Hilbert space (RKHS) of a WL kernel. In this line of research, the kernel framework is essentially used to design the architecture of the GNN since the final model is trained as a classical neural network. This is also the approach used by Zhang et al. (2018a) and Morris et al. (2019). By contrast, Du et al. (2019) adopt an opposite strategy and leverage a GNN architecture to design new graph kernels, which are equivalent to infinitely-wide GNNs initialized with random weights and trained with gradient descent. Other attempts to merge neural networks and graph kernels involve using the metric induced by graph kernels to initialize a GNN (Navarin et al., 2018), or using graph kernels to obtain continuous embeddings that are plugged to neural networks (Nikolentzos et al., 2018).

In this paper, we go a step further in bridging graph neural networks and kernel methods by proposing an explicit multilayer kernel representation, which can be used either as a traditional kernel method, or trained end-to-end as a GNN when enough labeled data are available. The multilayer construction allows to compute a series of maps
which account for local sub-structures (“receptive fields”) of increasing size. The graph representation is obtained by pooling the final representations of its nodes. The resulting kernel extends to graph-structured data the concept of convolutional kernel networks (CKNs), which was originally designed for images and sequences (Mairal, 2016; Chen et al., 2019a). As our representation of nodes is built by iteratively aggregating representations of their outgoing paths, our model can also be seen as a multilayer extension of path kernels. Relying on paths rather than neighbors for the aggregation step makes our approach more expressive than the GNNS considered in Xu et al. (2019), which implicitly rely on walks and whose power cannot exceed the Weisfeiler-Lehman (WL) graph isomorphism test. Even with medium/small path lengths (which leads to reasonable computational complexity in practice), we show that the resulting representation outperforms walk or WL kernels.

Our model called graph convolutional kernel network (GCKN) relies on the successive uses of the Nyström method (Williams & Seeger, 2001) to approximate the feature map at each layer, which makes our approach scalable. GCKNs can then be interpreted as a new type of graph neural network whose filters may be learned without supervision, by following kernel approximation principles. Such unsupervised graph representation is known to be particularly effective when small amounts of labeled data are available. Similar to CKNs, our model can also be trained end-to-end, as a GNN, leading to task-adaptive representations, with a computational complexity similar to that of a GNN when the path lengths are small enough.

**Notation.** A graph $G$ is defined as a triplet $(V, E, a)$, where $V$ is the set of vertices, $E$ is the set of edges, and $a : V \rightarrow \Sigma$ is a function that assigns attributes, either discrete or continuous, from a set $\Sigma$ to nodes in the graph. A path is a sequence of distinct vertices linked by edges and we denote by $P(G)$ the set of paths and paths of length $k$ in $G$, respectively. In particular, $P_0(G)$ is reduced to $V$. We also denote by $P_k(G, u) \subset P_k(G)$ the set of paths of length $k$ starting from $u$ in $V$. For any path $p$ in $P(G)$, we denote by $a(p)$ in $\Sigma^{1+k}$ the concatenation of node attributes in this path. We replace $P$ by $\mathcal{W}$ to denote the corresponding sets of walks by allowing repeated nodes.

**2. Related Work on Graph Kernels**

Graph kernels were originally introduced by Gärtner et al. (2003) and Kashima et al. (2003), and have been the subject of intense research during the last twenty years (see the reviews of Vishwanathan et al., 2010; Kriege et al., 2020).

In this paper, we consider graph kernels that represent a graph as a feature vector counting the number of occurrences of some local connected sub-structure. Enumerating common local sub-structures between two graphs is unfortunately often intractable; for instance, enumerating common subgraphs or common paths is known to be NP-hard (Gärtner et al., 2003). For this reason, the literature on graph kernels has focused on alternative structures allowing for polynomial-time algorithms, e.g., walks.

More specifically, we consider graph kernels that perform pairwise comparisons between local sub-structures centered at every node. Given two graphs $G = (V, E, a)$ and $G' = (V', E', a')$, we consider the kernel

$$K(G, G') = \sum_{u \in V} \sum_{u' \in V'} \kappa_{\text{base}}(l_G(u), l_{G'}(u')),$$

where the base kernel $\kappa_{\text{base}}$ compares a set of local patterns centered at nodes $u$ and $u'$, denoted by $l_G(u)$ and $l_{G'}(u')$, respectively. For simplicity, we will omit the notation $l_G(u)$ in the rest of the paper, and the base kernel will be simply written $\kappa_{\text{base}}(u, u')$ with an abuse of notation. As noted by Lei et al. (2017); Kriege et al. (2020), this class of kernels covers most of the examples mentioned in the introduction.

**Walks and path kernels.** Since computing all path co-occurrences between graphs is NP-hard, it is possible instead to consider paths of length $k$, which can be reasonably enumerated if $k$ is small enough, or the graphs are sparse. Then, we may define the kernel $\kappa_{\text{path}}^{(k)}$ as (1) with

$$\kappa_{\text{base}}(u, u') = \sum_{p \in P_k(G, u)} \sum_{p' \in P_k(G', u')} \delta(a(p), a'(p')),$$

where $a(p)$ represents the attributes for path $p$ in $G$, and $\delta$ is the Dirac kernel such that $\delta(a(p), a'(p')) = 1$ if $a(p) = a'(p')$ and 0 otherwise.

It is also possible to define a variant that enumerates all paths up to length $k$, by simply adding the kernels $\kappa_{\text{path}}^{(i)}$:

$$K_{\text{path}}(G, G') = \sum_{i=0}^{k} K_{\text{path}}^{(i)}(G, G').$$

Similarly, one may also consider using walks by simply replacing the notation $P$ by $\mathcal{W}$ in the previous definitions.

**Weisfeiler-Lehman subtree kernels.** A subtree is a subgraph with a tree structure. It can be extended to subtree patterns (Shervashidze et al., 2011; Bach, 2008) by allowing nodes to be repeated, just as the notion of walks extends that of paths. All previous subtree kernels compare subtree patterns instead of subtrees. Among them, the Weisfeiler-Lehman (WL) subtree kernel is one of the most widely used graph kernels to capture such patterns. It is essentially based on a mechanism to augment node attributes by iteratively aggregating and hashing the attributes of each node’s neighborhoods. After $i$ iterations, we denote by $a_i$ the new node
attributes for graph $G = (V, E, a)$, which is defined in Algorithm 1 of Shervashidze et al. (2011) and then the WL subtree kernel after $k$ iterations is defined, for two graphs $G = (V, E, a)$ and $G' = (V', E', a')$, as

$$K_{WL}(G, G') = \sum_{i=0}^{k} K_{\text{subtree}}^{(i)}(G, G'),$$

where

$$K_{\text{subtree}}^{(i)}(G, G') = \sum_{u \in V} \sum_{u' \in V'} K_{\text{subtree}}^{(i)}(u, u'),$$

with $K_{\text{subtree}}^{(i)}(u, u') = \delta(a_i(u), a'_i(u'))$ and the attributes $a_i(u)$ capture subtree patterns of depth $i$ rooted at node $u$.

3. Graph Convolutional Kernel Networks

In this section, we introduce our model, which builds upon the concept of graph-structured feature maps, following the terminology of convolutional neural networks.

**Definition 1** (Graph feature map). Given a graph $G = (V, E, a)$ and a RKHS $\mathcal{H}$, a graph feature map is a mapping $\varphi : V \rightarrow \mathcal{H}$, which associates to every node a point in $\mathcal{H}$ representing information about local graph substructures.

We note that the definition matches that of convolutional neural networks. The RKHS of $\mathcal{H}$ may admit a graph feature map $\varphi$ for graphs $G, G'$ with feature maps $\varphi_0, \varphi'_0$ of the form (1) with

$$\kappa_{\text{base}}(u, u') = \sum_{p \in P_k(G, u)} \sum_{p' \in P_k(G', u')} \kappa_1(\varphi(p), \varphi'(p')),$$

where $\varphi_0(p) = [\varphi_0(p_i)]_{i=0}^k$ denotes the concatenation of $k + 1$ attributes along path $p_i$, which is an element of $\mathcal{H}_0^{k+1}$. $p_i$ is the $i$-th node on path $p$ starting from index 0, and $\kappa_1$ is a Gaussian kernel comparing such attributes:

$$\kappa_1(\varphi(p), \varphi'(p')) = e^{-\frac{k}{\sigma^2} \sum_{i=0}^{k} ||\varphi(p_i) - \varphi'(p_i)||_2^2}. $$

This is an extension of the path kernel, obtained by replacing the hard matching function $\delta$ in (2) by $\kappa_1$, as done for instance by Togninalli et al. (2019) for the WL kernel. This replacement not only allows us to use continuous attributes, but also has important consequences in the discrete case since it allows to perform inexact matching between paths. For instance, when the graph is a chain with discrete attributes—in other words, a string—then, paths are simply $k$-mers, and the path kernel (with matching function $\delta$) becomes the spectrum kernel for sequences (Leslie et al., 2001). By using $\kappa_1$, instead, we obtain the single-layer CKN kernel of Chen et al. (2019a), which performs inexact matching, as the mismatch kernel does (Leslie et al., 2004), and leads to better performances in many tasks involving biological sequences.

3.1. Single-Layer Construction of the Feature Map

We propose a single-layer model corresponding to a continuous relaxation of the path kernel. We assume that the input attributes $a(u)$ live in $\mathbb{R}^{q_0}$, such that a graph $G = (V, E, a)$ admits a graph feature map $\varphi_0 : V \rightarrow \mathcal{H}_0$ with $\mathcal{H}_0 = \mathbb{R}^{q_0}$ and $\varphi_0(u) = a(u)$. Note that this assumption also allows us to handle discrete labels by using a one-hot encoding strategy—that is, e.g., four labels $\{A, B, C, D\}$ are represented by four-dimensional vectors $(1, 0, 0, 0), (0, 1, 0, 0), (0, 0, 1, 0), (0, 0, 0, 1)$, respectively.

**Continuous relaxation of the path kernel.** We rely on paths of length $k$, and introduce the kernel $K_1$ for graphs $G, G'$ with feature maps $\varphi_0, \varphi'_0$ of the form (1) with

$$K_1(\varphi_0, \varphi'_0) = \kappa_1(\varphi_0(p), \varphi'_0(p')),$$

with $\kappa_1(\varphi_0(p), \varphi'_0(p')) = e^{-\frac{k}{\sigma^2} \sum_{i=0}^{k} ||\varphi(p_i) - \varphi'(p'_i)||_2^2}. $$

The RKHS of $K_1$ can be characterized by using Theorem 2 in Appendix A. It is the space of functions $f_z : G \mapsto \langle z, \Phi(G) \rangle_{\mathcal{H}}$ for all $z$ in $\mathcal{H}$ endowed with a particular norm.

Note that even though graph feature maps $\varphi, \varphi'$ are graph-dependent, learning with $K_1$ is possible as long as they all map nodes to the same RKHS $\mathcal{H}$—as $\Phi$ will then also map all graphs to the same space $\mathcal{H}$. We now detail the full construction of the kernel, starting with a single layer.

From graph feature map $\varphi_0$ to graph feature map $\varphi_1$. The kernel $K_1$ acts on pairs of paths in potentially different graphs, but only through their mappings to the same space $\mathcal{H}^{k+1}$. Since $K_1$ is positive definite, we denote by $\mathcal{H}_1$ its RKHS and consider its mapping $\varphi_1^{\text{path}} : \mathcal{H}_0^{k+1} \rightarrow \mathcal{H}_1$ such that

$$K_1(\varphi_0(p), \varphi'_0(p')) = \langle \varphi_1^{\text{path}}(\varphi_0(p)), \varphi_1^{\text{path}}(\varphi'_0(p')) \rangle_{\mathcal{H}_1}. $$

For any graph $G$, we can now define a graph feature map $\varphi_1 : V \mapsto \mathcal{H}_1$, operating on nodes $u$ in $V$, as

$$\varphi_1(u) = \sum_{p \in P_k(G, u)} \varphi_1^{\text{path}}(\varphi_0(p)).$$

Then, the continuous relaxation of the path kernel, denoted by $K_1(G, G')$, can also be written as (6) with $\varphi = \varphi_1$, and its underlying kernel representation $\Phi_1$ is given by (7). The construction of $\varphi_1$ from $\varphi_0$ is illustrated in Figure 1.
3.2. Concrete Implementation and GCKNs

We now discuss algorithmic aspects, leading to the graph convolutional kernel network (GCKN) model, which consists in building a finite-dimensional embedding $\Psi(G)$ that may be used in various learning tasks without scalability issues. We start here with the single-layer case.

The Nyström method and the single-layer model. A naive computation of the path kernel $K_1$ requires comparing all pairs of paths in each pair of graphs. To gain scalability, a key component of the CKN model is the Nyström method (Williams & Seeger, 2001), which computes finite-dimensional approximate kernel embeddings. We discuss here the use of such a technique to define finite-dimensional maps $\phi_1 : \mathcal{V} \to \mathbb{R}^{n_1}$ and $\phi_1' : \mathcal{V}' \to \mathbb{R}^{n_1}$ for graphs $G, G'$ such that for all pairs of nodes $u, u'$ in $\mathcal{V}, \mathcal{V}'$, respectively,

$$\langle \phi_1(u), \phi_1'(u') \rangle_{\mathcal{H}_1} \approx \langle \psi_1(u), \psi_1'(u') \rangle_{\mathbb{R}^{n_1}}.$$

The consequence of such an approximation is that it provides a finite-dimensional approximation $\Psi_1$ of $\Phi_1$:

$$K_1(G, G') \approx \langle \Psi_1(G), \Psi_1(G') \rangle_{\mathbb{R}^{n_1}}$$

with $\Psi_1(G) = \sum_{u \in \mathcal{V}} \psi_1(u)$.

Then, a supervised learning problem with kernel $K_1$ on a dataset $(G_i, y_i)_{i=1,\ldots,n}$, where $y_i$ are labels in $\mathbb{R}$, can be solved by minimizing the regularized empirical risk

$$\min \sum_{i=1}^n L(y_i, \langle \Psi_1(G_i), w \rangle) + \lambda \|w\|^2,$$  \hspace{1cm} (11)

where $L$ is a convex loss function. Next, we show that using the Nyström method to approximate the kernel $K_1$ yields a new type of GNN, represented by $\Psi_1(G)$, whose filters can be obtained without supervision, or, as discussed later, with back-propagation in a task-adaptive manner.

Specifically, the Nyström method projects points from a given RKHS onto a finite-dimensional subspace and performs all subsequent operations within that subspace. In the context of $\kappa_1$, whose RKHS is $\mathcal{H}_1$ with mapping function $\phi_1$, we consider a collection $Z = \{z_1, \ldots, z_{q_1}\}$ of $q_1$ prototype paths represented by attributes in $\mathcal{H}_0^{k+1}$, and we define the subspace $\mathcal{E}_1 = \text{Span}(\phi_1(z_1), \ldots, \phi_1(z_{q_1}))$. Given a new path with attributes $z$, it is then possible to show (see Chen et al., 2019a) that the projection of path...
attributes \(z\) onto \(\mathcal{E}_1\) leads to the \(q_1\)-dimensional mapping

\[
\psi_1^{\text{path}}(z) = [\kappa_1(z_i, z_j)]_{i,j}^{-\frac{1}{2}}[\kappa_1(z_1, z), \ldots, \kappa_1(z_{q_1}, z)]^T,
\]

where \([\kappa_1(z_i, z_j)]_{i,j}\) is a \(q_1 \times q_1\) Gram matrix. Then, the approximate graph feature map \(\psi_1\) is obtained by pooling

\[
\psi_1(u) = \sum_{p \in P_k(G, u)} \psi_1^{\text{path}}(\psi_0(p)) \quad \text{for all } u \in V,
\]

where \(\psi_0 = \varphi_0\) and \(\psi_0(p) = [\psi_0(p_i)]_{i=0,\ldots,k}\) in \(\mathbb{R}^{q_0(k+1)}\) represents the attributes of path \(p\), with an abuse of notation.

**Interpretation as a GNN.** When input attributes \(\psi_0(u)\) have unit-norm, which is the case if we use one-hot encoding on discrete attributes, the Gaussian kernel \(\kappa_1\) between two path attributes \(z, z'\) in \(\mathbb{R}^{q_0(k+1)}\) may be written

\[
\kappa_1(z, z') = e^{-\frac{1}{2}\|z-z'\|^2} = e^{\alpha_1(z^T z' - k-1)} = \sigma_1(z^T z'),
\]

which is a dot-product kernel with a non-linear function \(\sigma_1\). Then, calling \(Z\) in \(\mathbb{R}^{q_0(k+1) \times q_1}\) the matrix of prototype path attributes, we have

\[
\psi_1(u) = \sum_{p \in P_k(G, u)} \sigma_1(Z^T Z)^{-\frac{1}{2}} \sigma_1(Z^T \psi_0(p)),
\]

where, with an abuse of notation, the non-linear function \(\sigma_1\) is applied pointwise. Then, the map \(\psi_1\) is build from \(\psi_0\) with the following steps: (i) feature aggregation along the paths, (ii) encoding of the paths with a linear operation followed by point-wise non-linearity, (iii) multiplication by the \(q_1 \times q_1\) matrix \(\sigma_1(Z^T Z)^{-\frac{1}{2}}\) and (iv) linear pooling. The major difference with a classical GNN is that the “filtering” operation may be interpreted as an orthogonal projection onto a linear subspace, due to the matrix \(\sigma_1(Z^T Z)^{-\frac{1}{2}}\). Unlike the Dirac function, the exponential function \(\sigma_1\) is differentiable. A useful consequence is the possibility of optimizing the filters \(Z\) with back-propagation as detailed below. Note that in practice we add a small regularization term to the diagonal for stability reason: \((\sigma_1(Z^T Z) + \varepsilon I)^{-\frac{1}{2}}\) with \(\varepsilon = 0.01\).

**Learning without supervision.** Learning the “filters” \(Z\) with Nyström can be achieved by simply running a K-means algorithm on path attributes extracted from training data (Zhang et al., 2008). This is the approach adopted for CKNs by Mairal (2016); Chen et al. (2019a), which proved to be very effective as shown in the experimental section.

**End-to-end learning with back-propagation.** While the previous unsupervised learning strategy consists in finding a good kernel approximation that is independent of labels, it is also possible to learn the parameters \(Z\) end-to-end, by minimizing (11) jointly with respect to \(Z\) and \(u\). The main observations from Chen et al. (2019a) in the context of biological sequences is that such a supervised learning approach may yield good models with much fewer filters \(q_1\) than with the unsupervised learning strategy. We refer the reader to Chen et al. (2019a,b) for how to perform back-propagation with the inverse square root matrix \(\sigma_1(Z^T Z)^{-\frac{1}{2}}\).

**Complexity.** The complexity for computing the feature map \(\psi_1\) is dominated by the complexity of finding all the paths of length \(k\) from each node. This can be done by simply using a depth first search algorithm, whose worst-case complexity for each graph is \(O(|V|d^k)\), where \(d\) is the maximum degree of each node, meaning that large \(k\) may be used only for sparse graphs. Then, each path is encoded in \(O(q_1q_0(k+1))\) operations; When learning with back-propagation, each gradient step requires computing the eigenvalue decomposition of \(\sigma_1(Z^T Z)^{-\frac{1}{2}}\) whose complexity is \(O(q_1^3)\), which is not a computational bottleneck when using mini-batches of order \(O(q_1)\), where typical practical values for \(q_1\) are reasonably small, e.g., less than 128.

### 3.3. Multilayer Extensions

The mechanism to build the feature map \(\varphi_1\) from \(\varphi_0\) can be iterated, as illustrated in Figure 1 which shows how to build a feature map \(\varphi_{j+1}\) from a previous one \(\varphi_j\). As discussed by Mairal (2016) for CKNs, the Nyström method may then be extended to build a sequence of finite-dimensional maps \(\psi_0, \ldots, \psi_J\), and the final graph representation is given by

\[
\Psi_j(G) = \sum_{u \in V} \psi_j(u).
\]

The computation of \(\Psi_j(G)\) is illustrated in Algorithm 1. Here we discuss two possible uses for these additional layers, either to account for more complex structures than paths, or to extend the receptive field of the representation without resorting to the enumeration of long paths. We will denote by \(k_j\) the path length used at layer \(j\).

#### A simple two-layer model to account for subtrees.

As emphasized in (7), GCKN relies on a representation \(\Phi(G)\) of graphs, which is a sum of node-level representations provided by a graph feature map \(\varphi\). If \(\varphi\) is a sum over

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**Algorithm 1 Forward pass for multilayer GCKN**

1: **Input:** graph \(G = (V, E, \psi_0 : V \to \mathbb{R}^{|V|})\), set of anchor points (filters) \(Z_j \in \mathbb{R}^{(k+1)q_1 \times q_1}\) for \(j = 1, \ldots, J\).
2: **for** \(j = 1, \ldots, J\) **do**
3: **for** \(u \in V\) **do**
4: \(\psi_j(u) = \sum_{p \in P_k(G, u)} \psi_1^{\text{path}}(\psi_{j-1}(p))\);
5: **end for**
6: **end for**
7: **Global pooling:** \(\Psi(G) = \sum_{u \in V} \psi_J(u)\);
path starting at the represented node, $\Phi(G)$ can simply be written as a sum over all paths in $G$, consistently with our observation that (6) recovers the path kernel when using a Dirac kernel to compare paths in $\kappa_2$. The path kernel often leads to good performances, but it is also blind to more complex structures. Figure 2 provides a simple example of this phenomenon, using $k = 1$: $G_1$ and $G_3$ differ by a single edge, while $G_4$ has a different set of nodes and a rather different structure. Yet $P_1(G_3) = P_1(G_4)$, making $K_1(G_1, G_3) = K_1(G_1, G_4)$ for the path kernel.

![Figure 2. Example cases using $\kappa_1 = \kappa_2 = \delta$, with path lengths $k_1 = 1$ and $k_2 = 0$; The one-layer kernel $K_1$ counts the number of common edges while the two-layer $K_2$ counts the number of nodes with the same set of outgoing edges. The figure suggests using $K_1 + K_2$ to gain expressiveness.](image)

Expressing more complex structures requires breaking the succession of linearity introduced in (7) and (10)—much like pointwise nonlinearities are used in neural networks. Concretely, this effect can simply be obtained by using a second layer with path length $k_2 = 0$—paths are then identified to vertices—which produces the feature map $\varphi_2(u) = \phi_2^{\text{path}}(\varphi_1(u))$, where $\phi_2^{\text{path}} : H_1 \to H_2$ is a non-linear kernel mapping. The resulting kernel is then

$$K_2(G, G') = \sum_{u \in V} \sum_{u' \in V'} \langle \varphi_2(u), \varphi_2(u') \rangle_{H_2}$$

$$= \sum_{u \in V} \sum_{u' \in V'} \kappa_2(\varphi_1(u), \varphi_1(u')).$$  

(15)

When $\kappa_1$ and $\kappa_2$ are both Dirac kernels, $K_2$ counts the number of nodes in $G$ and $G'$ with the exact same set of outgoing paths $P(G, u)$, as illustrated in Figure 2.

Theorem 1 further illustrates the effect of using a nonlinear $\phi_2^{\text{path}}$ on the feature map $\varphi_1$, by formally linking the walk and WL subtree kernel through our framework.

**Theorem 1.** Let $G = (V, E)$, $G' = (V', E')$, $\mathcal{M}$ be the set of exact matchings of subsets of the neighborhoods of two nodes, as defined in Shervashidze et al. (2011), and $\varphi$ defined as in (10) with $\kappa_1 = \delta$ and replacing paths by walks. For any $u \in V$ and $u' \in V'$ such that $|\mathcal{M}(u, u')| = 1$,

$$\delta(\varphi_1(u), \varphi_1(u')) = \kappa_{\text{subtree}}^{(k)}(u, u').$$  

(16)

Recall that when using (8) with walks instead of paths and a Dirac kernel for $\varphi_1$, the kernel (6) with $\varphi = \varphi_1$ is the walk kernel. The condition $|\mathcal{M}(u, u')| = 1$ indicates that $u$ and $u'$ have the same degrees and each of them has distinct neighbors. This can be always ensured by including degree information and adding noise to node attributes. For a large class of graphs, both the walk and WL subtree kernels can therefore be written as (6) with the same first layer $\varphi_1$ representing nodes by their walk histograms. While walk kernels use a single layer, WL subtree kernels rely on a second layer $\varphi_2$ mapping nodes to the indicator function of $\varphi_1(u)$.

Theorem 1 also shows that the kernel built in (15) is a path-based version of WL subtree kernels, therefore more expressive as it captures subtrees rather than subtree patterns. However, the Dirac kernel lacks flexibility, as it only accounts for pairs of nodes with identical $P(G, u)$. For example, in Figure 2, $K_2(G_1, G_2) = 0$ even though $G_1$ only differs from $G_2$ by two edges, because these two edges belong to the set $P(G, u)$ of all nodes in the graph. In order to retain the stratification by node of (15) while allowing for a softer comparison between sets of outgoing paths, we replace $\delta$ by the kernel $\kappa_2(\varphi_1(u), \varphi_1(u')) = e^{-\alpha_2 ||\varphi_1(u) - \varphi_1(u')||_2}$. Large values of $\alpha_2$ recover the behavior of the Dirac, while smaller values gives non-zero values for similar $\varphi_1(u)$.

A multilayer model to account for longer paths. In the previous paragraph, we have seen that adding a second layer could bring some benefits in terms of expressiveness, even when using path lengths $k_2 = 0$. Yet, a major limitation of this model is the exponential complexity of path enumeration, which is required to compute the feature map $\varphi_1$, preventing us to use large values of $k$ as soon as the graph is dense. Representing large receptive fields while relying on path enumerations with small $k$, e.g., $k \leq 3$, is nevertheless possible with a multilayer model. To account for a receptive field of size $k$, the previous model requires a path enumeration with complexity $O(|V|d^k)$, whereas the complexity of a multilayer model is linear in $k$.

3.4. Practical Variants

Summing the kernels for different $k$ and different scales. As noted in Section 2, summing the kernels corresponding to different values of $k$ provides a richer representation. We also adopt such a strategy, which corresponds to concatenating the feature vectors $\Psi(G)$ obtained for various path lengths $k$. When considering a multilayer model, it is also possible to concatenate the feature representations obtained at every layer $j$, allowing to obtain a multi-scale feature representation of the graph and gain expressiveness.

**Use of homogeneous dot-product kernel.** Instead of the Gaussian kernel (9), it is possible to use a homogeneous dot-
product kernel, as suggested by Mairal (2016) for CKNs:
\[ \kappa_{1}(z, z') = \|z\| \|z'\| \sigma_{1}\left(\frac{\langle z, z' \rangle}{\|z\| \|z'\|}\right), \]
where \( \sigma_{1} \) is defined in (12). Note that when \( z, z' \) have unit norm, we recover the Gaussian kernel (9). In our paper, we use such a kernel for upper layers, or for continuous input attributes when they do not have unit norm. For multilayer models, this homogenization is useful for preventing vanishing or exponentially growing representations. Note that ReLU is also a homogeneous non-linear mapping.

Other types of pooling operations. Another variant consists in replacing the sum pooling operation in (13) and (14) by a mean or a max pooling. While using max pooling as a heuristic seems to be effective on some datasets, it is hard to justify from a RKHS point of view since max operations typically do not yield positive definite kernels. Yet, such a heuristic is widely adopted in the kernel literature, e.g., for string alignment kernels (Saigo et al., 2004). In order to solve such a discrepancy between theory and practice, Chen et al. (2019b) propose to use the generalized max pooling operator of Murray & Perronnin (2014), which is compatible with the RKHS point of view. Applying the same ideas to GCKNs is straightforward.

Using walk kernel instead of path kernel. One can use a relaxed walk kernel instead of the path kernel in (8), at the cost of losing some expressiveness but gaining some time complexity. Indeed, there exists a very efficient recursive way to enumerate walks and thus to compute the resulting complexity. Indeed, there exists a very efficient recursive way to enumerate walks and thus to compute the resulting approximate feature map in (13) for the walk kernel. Specifically, if we denote the \( k \)-walk kernel by \( \kappa_{\text{walk}}^{(k)} \), then its value between two nodes can be decomposed as the product of the 0-walk kernel between the nodes and the sum of the \( (k-1) \)-walk kernel between their neighbors
\[ \kappa_{\text{walk}}^{(k)}(u, u') = \kappa_{\text{walk}}^{(0)}(u, u') \sum_{v \in \mathcal{N}(u)} \sum_{v' \in \mathcal{N}(u')} \kappa_{\text{walk}}^{(k-1)}(v, v'), \]
where \( \kappa_{\text{walk}}^{(0)}(u, u') = \kappa_{1}(\varphi_{0}(u), \varphi_{0}(u')) \). After applying the Nyström method, the approximate feature map of the walk kernel is written, similar to (13), as
\[ \psi_{1}(u) = \sigma_{1}(Z^{T}Z)^{-\frac{1}{2}} \sum_{p \in \mathcal{W}_{k}(G, u)} \sigma_{1}(Z^{T} \psi_{0}(p)) \cdot c_{k}(u) := \]

Based on the above observation and following similar induction arguments as Chen et al. (2019b), it is not hard to show that \( (c_{j}(u))_{j=1, \ldots, k} \) obeys the following recursion
\[ c_{j}(u) = b_{j}(u) \odot \sum_{v \in \mathcal{N}(u)} c_{j-1}(v), \]
where \( \odot \) denotes the element-wise product and \( b_{j}(u) \) is a vector in \( \mathbb{R}^{q_{1}} \) whose entry \( i \) in \( \{1, \ldots, q_{1}\} \) is \( \kappa_{1}(u, z_{i}^{(k+1-j)}) \) and \( z_{i}^{(k+1-j)} \) denotes the \( k + 1 - j \)-th column vector of \( z_{i} \) in \( \mathbb{R}^{q_{0}} \). More details can be found in Appendix C.

4. Model Interpretation

Ying et al. (2019) introduced an approach to interpret trained GNN models, by finding a subgraph of an input graph \( G \) maximizing the mutual information with its predicted label (note that this approach depends on a specific input graph). We show here how to adapt similar ideas to our framework.

Interpreting GCKN-path and GCKN-subtree. We call GCKN-path our model \( \Psi_{1} \) with a single layer, and GCKN-subtree our model \( \Psi_{2} \) with two layers but with \( k_{2} = 0 \), which is the first model presented in Section 3.3 that accounts for subtree structures. As these models are built upon path enumeration, we extend the method of Ying et al. (2019) by identifying a small subset of paths in an input graph \( G \) preserving the prediction. We then reconstruct a subgraph by merging the selected paths. For simplicity, let us consider a one-layer model. As \( \Psi_{1}(G) \) only depends on \( G \) through its set of paths \( \mathcal{P}_{k}(G) \), we note \( \Psi_{1}(\mathcal{P}) \) with an abuse of notation for any subset of \( \mathcal{P} \) of paths in \( G \), to emphasize the dependency in this set of paths. For a trained model \( \Psi_{1}(\mathcal{P}, w) \) and a graph \( G \), our objective is to solve
\[ \min_{\mathcal{P}' \subseteq \mathcal{P}_{k}(G)} L(\hat{y}, (\Psi_{1}(\mathcal{P}'), w)) + \mu |\mathcal{P}'|, \tag{17} \]
where \( \hat{y} \) is the predicted label of \( G \) and \( \mu \) a regularization parameter controlling the number of paths to select. This problem is combinatorial and can be computationally intractable when \( |\mathcal{P}(G)| \) is large. Following Ying et al. (2019), we relax it by using a mask \( M \) with values in \( \{0; 1\} \) over the set of paths, and replace the number of paths \( |\mathcal{P}'| \) by the \( \ell_{1} \)-norm of \( M \), which is known to have a sparsity-inducing effect (Tibshirani, 1996). The problem then becomes
\[ \min_{M \in \{0; 1\}^{\mathcal{P}_{k}(G)}} L(\hat{y}, (\Psi_{1}(\mathcal{P}_{k}(G) \odot M), w)) + \mu \|M\|_{1}, \tag{18} \]
where \( \mathcal{P}_{k}(G) \odot M \) denotes the use of \( M(p) \alpha(p) \) instead of \( \alpha(p) \) in the computation of \( \Psi_{1} \) for all \( p \) in \( \mathcal{P}_{k}(G) \). Even though the problem is non-convex due to the non-linear mapping \( \Psi_{1} \), it may still be solved approximately by using projected gradient-based optimization techniques.

Interpreting multilayer models. By noting that \( \Psi_{j}(G) \) only depends on the union of the set of paths \( \mathcal{P}_{k_{j}}(G) \), for all layers \( l \leq j \), we introduce a collection of masks \( M_{l} \) at each layer, and then optimize the same objective as (18) over all masks \( (M_{l})_{l=1, \ldots, j} \), with the regularization \( \sum_{l=1}^{j} \|M_{l}\|_{1} \).
5. Experiments

We evaluate GCKN and compare its variants to state-of-the-art methods, including GNNs and graph kernels, on several real-world graph classification datasets, involving either discrete or continuous attributes.

5.1. Implementation Details

We follow the same protocols as (Du et al., 2019; Xu et al., 2019), and report the average accuracy and standard deviation over a 10-fold cross validation on each dataset. We use the same data splits as Xu et al. (2019), using their code. Note that performing nested 10-fold cross validation would have provided better estimates of test accuracy for all models, but it would have unfortunately required 10 times more computation, which we could not afford for many of the baselines we considered.

Considered models. We consider two single-layer models called GCKN-walk and GCKN-path, corresponding to the continuous relaxation of the walk and path kernels respectively. We also consider the two-layer model GCKN-subtree introduced in Section 3.3 with path length \( k_2 = 0 \), which accounts for subtrees. Finally, we consider a 3-layer model GCKN-3layers with path length \( k_2 = 2 \) (which enumerates paths with three vertices for the second layer), and \( k_3 = 0 \), which introduces a non-linear mapping before global pooling, as in GCKN-subtree. We use the same parameters \( \alpha_j \) and \( q_j \) (number of filters) across layers. Our comparisons include state-of-the-art graph kernels such as WL kernel (Shervashidze et al., 2011), AWL (Ivanov & Buneaev, 2018), RetGK (Zhang et al., 2018b), GNTK (Du et al., 2019), WWL (Togninalli et al., 2019) and recent GNNs including GCN (Kipf & Welling, 2017), PatchySAN (Niepert et al., 2016) and GIN (Xu et al., 2019). We also include a simple baseline method LDP (Cai & Wang, 2018) based on node degree information and a Gaussian SVM.

Learning unsupervised models. Following Mairal (2016), we learn the anchor points \( Z_j \) for each layer by K-means over 300000 extracted paths from each training fold. The resulting graph representations are then mean-centered, standardized, and used within a linear SVM classifier (11) with squared hinge loss. In practice, we use the SVM implementation of the Cyanure toolbox (Mairal, 2019).\(^1\) For each 10-fold cross validation, we tune the bandwidth of the Gaussian kernel (identical for all layers), pooling operation (local (13) or global (14)), path size \( k_1 \) at the first layer, number of filters (identical for all layers) and regularization parameter \( \lambda \) in (11). More details are provided in Appendix B, as well as a study of the model robustness to hyperparameters.

Learning supervised models. Following Xu et al. (2019), we use an Adam optimizer (Kingma & Ba, 2015) with the initial learning rate equal to 0.01 and halved every 50 epochs, and fix the batch size to 32. We use the unsupervised model based described above for initialization. We select the best model based on the same hyperparameters as for unsupervised models, with the number of epochs as an additional hyperparameter as used in Xu et al. (2019). Note that we do not use DropOut or batch normalization, which are typically used in GNNs such as Xu et al. (2019). Importantly, the number of filters needed for supervised models is always much smaller (e.g., 32 vs 512) than that for unsupervised models to achieve comparable performance.

5.2. Results

Graphs with categorical node labels. We use the same benchmark datasets as in Du et al. (2019), including 4 biochemical datasets MUTAG, PROTEINS, PTC and NCI1 and 3 social network datasets IMDB-B, IMDB-MULTI and COLLAB. All the biochemical datasets have categorical node labels while none of the social network datasets has node features. We use degrees as node labels for these datasets, following the protocols of previous works (Du et al., 2019; Xu et al., 2019; Togninalli et al., 2019). Similarly, we also transform all the categorical node labels to one-hot representations. The results are reported in Table 1. With a few exceptions, GCKN-walk has a small edge on graph kernels and GNNs—both implicitly relying on walks too—probably because of the soft structure comparison allowed by the Gaussian kernel. GCKN-path often brings some further improvement, which can be explained by its increasing the expressivity. Both multilayer GCKNs bring a stronger increase, whereas supervising the filter learning of GCKN-subtree does not help. Yet, the number of filters selected by GCKN-subtree-sup is smaller than GCKN-subtree-unsup (see Appendix B), allowing for faster classification at test time. GCKN-3layers-unsup performs in the same ballpark as GCKN-subtree-unsup, but benefits from lower complexity due to smaller path length \( k_1 \).

Graphs with continuous node attributes. We use 4 real-world graph classification datasets with continuous node attributes: ENZYMES, PROTEINS_full, BZR, COX2. All datasets and size information about the graphs can be found in Kersting et al. (2016). The node attributes are preprocessed with standardization as in Togninalli et al. (2019). To make a fair comparison, we follow the same protocol as used in Togninalli et al. (2019). Specifically, we perform 10 different 10-fold cross validations, using the same hyperparameters that give the best average validation accuracy. The hyperparameter search grids remain the same as for training graphs with categorical node labels. The results are shown in Table 2. They are comparable to the

\(^1\)http://julien.mairal.org/cyanure/
We train a supervised GCKN-subtree model on the Mutagenicity dataset (Kersting et al., 2016), and use our method described in Section 4 to identify important subgraphs. Figure 3 shows examples of detected subgraphs. Our method is able to identify chemical groups known for their mutagenicity such as polycyclic aromatic hydrocarbon (top row left), Diphenyl ether (top row middle) or NO₂ (top row right), thus admitting simple model interpretation. We also find some groups whose mutagenicity is not known, such as polyphenylene sulfide (bottom row middle) and 2-chloroethyl- (bottom row right). More details and additional results are provided in Appendix B.

5.3. Model Interpretation

We train a supervised GCKN-subtree model on the Mutagenicity dataset (Kersting et al., 2016), and use our method described in Section 4 to identify important subgraphs. Figure 3 shows examples of detected subgraphs. Our method is able to identify chemical groups known for their mutagenicity such as polycyclic aromatic hydrocarbon (top row left), Diphenyl ether (top row middle) or NO₂ (top row right), thus admitting simple model interpretation. We also find some groups whose mutagenicity is not known, such as polyphenylene sulfide (bottom row middle) and 2-chloroethyl- (bottom row right). More details and additional results are provided in Appendix B.

Table 1. Classification accuracies on graphs with discrete node attributes. The accuracies of other models are taken from Du et al. (2019) except LDP, which we evaluate on our splits and for which we tune bin size, the regularization parameter in the SVM and Gaussian kernel bandwidth. Note that RetGK uses a different protocol, performing 10-fold cross-validation 10 times and reporting the average accuracy.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>MUTAG</th>
<th>PROTEINS</th>
<th>PTC</th>
<th>NCI1</th>
<th>IMDB-B</th>
<th>IMDB-M</th>
<th>COLLAB</th>
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<td>1113</td>
<td>344</td>
<td>4110</td>
<td>1000</td>
<td>1500</td>
<td>5000</td>
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<td>2</td>
<td>3</td>
<td>3</td>
<td>3</td>
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<tr>
<td>avg ťnodes</td>
<td>18</td>
<td>39</td>
<td>26</td>
<td>30</td>
<td>20</td>
<td>13</td>
<td>74</td>
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<tr>
<td>avg ťedges</td>
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<td>73</td>
<td>51</td>
<td>32</td>
<td>97</td>
<td>66</td>
<td>2458</td>
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<tr>
<td>LDP</td>
<td>88.9 ± 9.6</td>
<td>73.3 ± 5.7</td>
<td>63.8 ± 6.6</td>
<td>72.0 ± 2.0</td>
<td>68.5 ± 4.0</td>
<td>42.9 ± 3.7</td>
<td>76.1 ± 1.4</td>
</tr>
<tr>
<td>WL subtree</td>
<td>90.4 ± 5.7</td>
<td>75.0 ± 3.1</td>
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<td>86.0 ± 1.8</td>
<td>73.8 ± 3.9</td>
<td>50.9 ± 3.8</td>
<td>78.9 ± 1.9</td>
</tr>
<tr>
<td>AWL</td>
<td>87.9 ± 9.8</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>74.5 ± 5.9</td>
<td>51.5 ± 3.6</td>
</tr>
<tr>
<td>RetGK</td>
<td>90.3 ± 1.1</td>
<td>75.8 ± 0.6</td>
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<td>84.5 ± 0.2</td>
<td>71.9 ± 1.0</td>
<td>47.3 ± 0.7</td>
<td>81.0 ± 0.3</td>
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<tr>
<td>GNTK</td>
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<td>84.2 ± 1.5</td>
<td>76.9 ± 3.6</td>
<td>52.8 ± 4.6</td>
<td>83.6 ± 1.0</td>
</tr>
<tr>
<td>GCN</td>
<td>85.6 ± 5.8</td>
<td>76.0 ± 3.2</td>
<td>64.2 ± 4.3</td>
<td>80.2 ± 2.0</td>
<td>74.0 ± 3.4</td>
<td>51.9 ± 3.8</td>
<td>79.0 ± 1.8</td>
</tr>
<tr>
<td>PatchySAN</td>
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<td>75.9 ± 2.8</td>
<td>60.0 ± 4.8</td>
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<td>71.0 ± 2.2</td>
<td>45.2 ± 2.8</td>
<td>72.6 ± 2.2</td>
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<td>GIN</td>
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<td>80.2 ± 1.9</td>
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<tr>
<td>GCKN-walk-unsup</td>
<td>92.8 ± 6.1</td>
<td>75.7 ± 4.0</td>
<td>65.9 ± 2.0</td>
<td>80.1 ± 1.8</td>
<td>75.9 ± 3.7</td>
<td>53.4 ± 4.7</td>
<td>81.7 ± 1.4</td>
</tr>
<tr>
<td>GCKN-path-unsup</td>
<td>92.8 ± 6.1</td>
<td>76.0 ± 3.4</td>
<td>67.3 ± 5.0</td>
<td>81.4 ± 1.6</td>
<td>75.9 ± 3.7</td>
<td>53.0 ± 3.1</td>
<td>82.3 ± 1.1</td>
</tr>
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<td>GCKN-subtree-unsup</td>
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<td>76.4 ± 3.9</td>
<td>70.8 ± 4.6</td>
<td>83.9 ± 1.6</td>
<td>77.8 ± 2.6</td>
<td>53.5 ± 4.1</td>
<td>83.2 ± 1.1</td>
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<tr>
<td>GCKN-3layer-unsup</td>
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<td>77.2 ± 3.8</td>
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<td>83.4 ± 1.5</td>
</tr>
<tr>
<td>GCKN-subtree-sup</td>
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<td>76.2 ± 2.5</td>
<td>68.4 ± 7.4</td>
<td>82.0 ± 1.2</td>
<td>76.5 ± 5.7</td>
<td>53.3 ± 3.9</td>
<td>82.9 ± 1.6</td>
</tr>
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</table>

Table 2. Classification accuracies on graphs with continuous attributes. The accuracies of other models except GNTK are taken from Togninalli et al. (2019). The accuracies of GNTK are obtained by running the code of Du et al. (2019) on a similar setting.

<table>
<thead>
<tr>
<th>Dataset</th>
<th>ENZYMES</th>
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<td>3</td>
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<tr>
<td>avg ťedges</td>
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<td>72.8</td>
<td>38.3</td>
<td>43.5</td>
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<td>RBF-WL</td>
<td>68.4 ± 1.5</td>
<td>75.4 ± 0.3</td>
<td>81.0 ± 1.7</td>
<td>75.5 ± 1.5</td>
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<tr>
<td>HKG-WL</td>
<td>63.0 ± 0.7</td>
<td>75.9 ± 0.2</td>
<td>78.6 ± 0.6</td>
<td>78.1 ± 0.5</td>
</tr>
<tr>
<td>HKG-SP</td>
<td>66.4 ± 0.4</td>
<td>75.8 ± 0.2</td>
<td>76.4 ± 0.7</td>
<td>72.8 ± 1.2</td>
</tr>
<tr>
<td>WWL</td>
<td>73.3 ± 0.9</td>
<td>77.9 ± 0.8</td>
<td>84.4 ± 2.0</td>
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<tr>
<td>GNTK</td>
<td>69.6 ± 0.9</td>
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<td>GCKN-subtree-unsup</td>
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<td>GCKN-subtree-sup</td>
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<td>77.6 ± 0.4</td>
<td>86.4 ± 0.5</td>
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</tr>
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Acknowledgements

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