SpeqNets: Sparsity-aware Permutation-equivariant
Graph Networks

Christopher Morris\(^1,2\), Gaurav Rattan\(^2\), Sandra Kiefer\(^3\), and Siamak Ravanbakhsh\(^1\)

\(^1\)McGill University and Mila Quebec AI Institute  
\(^2\)RWTH Aachen University  
\(^3\)Max Planck Institute for Software Systems

While message-passing graph neural networks have clear limitations in approximating permutation-equivariant functions over graphs or general relational data, more expressive, higher-order graph neural networks do not scale to large graphs. They either operate on \(k\)-order tensors or consider all \(k\)-node subgraphs, implying an exponential dependence on \(k\) in memory requirements, and do not adapt to the sparsity of the graph. By introducing new heuristics for the graph isomorphism problem, we devise a class of universal, permutation-equivariant graph networks, which, unlike previous architectures, offer a fine-grained control between expressivity and scalability and adapt to the sparsity of the graph. These architectures lead to vastly reduced computation times compared to standard higher-order graph networks in the supervised node- and graph-level classification and regression regime while significantly improving standard graph neural network and graph kernel architectures in terms of predictive performance.

1 Introduction

Graph-structured data is ubiquitous across application domains ranging from chemo- and bioinformatics [Barabasi and Oltvai, 2004, Jumper et al., 2021, Stokes et al., 2020] to image [Simonovsky and Komodakis, 2017] and social-network analysis [Easley and Kleinberg, 2010]. To develop successful machine-learning models in these domains, we need techniques that exploit the rich information inherent in the graph structure and the feature information within nodes and edges. In recent years, numerous approaches have been proposed for machine learning with graphs—most notably, approaches based on graph kernels [Borgwardt et al., 2020, Kriege et al., 2020] or using graph neural networks (GNNs) [Chami et al., 2020, Gilmer et al., 2017, Grohe, 2021, Morris et al., 2021]. Here, graph kernels based on the 1-dimensional Weisfeiler–Leman algorithm (1-WL) [Weisfeiler and Leman, 1968], a simple heuristic for the graph isomorphism problem, and corresponding GNNs [Morris et al., 2019, Xu et al., 2019] have recently advanced the state-of-the-art in supervised node- and graph-level learning. However, the 1-WL operates via simple neighborhood aggregation, and the purely local nature of the

\(^*\)Email: chris@christophermorris.info
related approaches misses important patterns in the given data. Moreover, they are only applicable to binary structures and therefore cannot deal with general structures containing relations of higher arity, e.g., hypergraphs. A more powerful algorithm for graph isomorphism testing is the \( k \)-dimensional Weisfeiler–Leman algorithm (\( k \)-WL) \cite{babai1979automorphism,Bai}. The algorithm captures more global, higher-order patterns by iteratively computing a coloring or labeling for \( k \)-tuples defined over the set of nodes of a given graph based on a certain notion of adjacency between tuples. See \cite{kief} for a survey and more background. However, since the algorithm considers all \( n^k \) many \( k \)-tuples of an \( n \)-node graph, it does not scale to large real-world graphs. Moreover, the cardinality of the considered neighborhood is always \( k \cdot n \). Hence, a potential sparsity of the input graph does not reduce the running time.

New neural architectures that possess the same power as the \( k \)-WL in terms of separating non-isomorphic graphs \cite{azizian2020higherProduct,geerts2020higherOrderGraphs,maron2019higherOrderGraphs} suffer from the same drawbacks, i.e., their memory requirement is lower-bounded by \( n^k \) for an \( n \)-node graph, and they have to resort to dense matrix multiplication. Recently, \cite{mmp} introduced the local variant (\( \delta \)-\( k \)-LWL) of the \( k \)-WL considering only a subset of the neighborhoods in \( k \)-WL. However, like the original algorithm, the local variant operates on the set of all possible \( k \)-tuples, again resulting in the same (exponential) memory requirements, rendering the algorithm not practical for large, real-world graphs.

Present work To address the described drawbacks, we introduce a new set of heuristics for the graph isomorphism problem, denoted \((k,s)\text{-LWL}\), which only considers a subset of all \( k \)-tuples, namely those inducing subgraphs with at most \( s \) connected components. We study the effect of \( k \) and \( s \) on the expressive power of the heuristics. Specifically, we show that the \((k,1)\text{-LWL}\) induces a hierarchy of provably expressive heuristics for the graph isomorphism problem, i.e., with increasing \( k \), the algorithm becomes strictly more expressive. Additionally, we prove that the \((k,2)\text{-LWL}\) is strictly more expressive than the \((k,1)\text{-LWL}\). Further, we separate the \((k,2)\text{-LWL}\) and \((k,k)\text{-LWL}\) by showing that the \((k,k)\text{-LWL}\) is strictly more expressive than the \((k,2)\text{-LWL}\). Building on these combinatorial insights, we derive corresponding provably expressive, permutation-equivariant neural architectures, denoted \((k,s)\text{-SpeqNets}\), which offer a more fine-grained trade-off between scalability and expressivity compared to previous architectures based on the \( k \)-WL, see Figure 1 for a high-level overview of the theoretical results. Empirically, we show how our architectures offer vastly reduced computation times while beating baseline GNNs and other higher-order graph networks in terms of predictive performance on well-known node- and graph-level prediction benchmark datasets.

1.1 Related work

In the following, we review related work and background on graph kernels, GNNs, and graph theory.

Graph kernels Historically, kernel methods—which implicitly or explicitly map graphs to elements of a Hilbert space—have been the dominant approach for supervised learning on graphs. Important early work in this area includes random-walk based kernels \cite{gartner2003kernel,kashima2003kernels,kriege2017wasserdistanz} and kernels based on shortest paths \cite{borgwardt2005shortest}. More recently, developments in the field have emphasized scalability, focusing

\footnote{In \cite{babai2016graph}, László Babai mentions that he introduced the algorithm in 1979 together with Rudolf Mathon.}
on techniques that bypass expensive Gram matrix computations by using explicit feature maps, see, e.g., [Shervashidze et al., 2011]. Morris et al. [2017] devised a local, set-based variant of the $k$-WL and a corresponding kernel. However, the approach is weaker than the tuple-based algorithm. Further, Morris et al. [2020] proposed kernels based on the $\delta$-$k$-LWL.

Yanardag and Vishwanathan [2015] successfully employed Graphlet [Shervashidze et al., 2009] and Weisfeiler–Leman kernels within frameworks for smoothed [Yanardag and Vishwanathan, 2015] and deep graph kernels [Yanardag and Vishwanathan, 2015]. Other recent work focuses on assignment-based [Johansson and Dubhashi, 2015, Kriege et al., 2016, Nikolentzos et al., 2017], spectral [Kondor and Pan, 2016, Verma and Zhang, 2017], graph decomposition-based [Nikolentzos et al., 2018], randomized binning approaches [Heimann et al., 2019], and the extension of kernels based on the 1-WL [Morris et al., 2016, Rieck et al., 2019, Togninalli et al., 2019]. For a theoretical investigation of graph kernels, see [Kriege et al., 2018], and for a thorough survey of graph kernels, see [Borgwardt et al., 2020, Kriege et al., 2020].

**GNNs** Recently, GNNs [Gilmer et al., 2017, Scarselli et al., 2009] emerged as an alternative to graph kernels. Instances of this architecture are described in, e.g., [Duvenaud et al., 2015, Hamilton et al., 2017, Velickovic et al., 2018], which can be subsumed under the message-passing framework introduced in [Gilmer et al., 2017]. In parallel, approaches based on spectral information were introduced in, e.g., [Defferrard et al., 2016, Bruna et al., 2014, Kipf and Welling, 2017, Monti et al., 2017]—all of which descend from early work in [Kireev, 1995, Baskin et al., 1997, Sperduti and Starita, 1997, Micheli and Sestito, 2005, Merkwirth and Lengauer, 2005, Micheli, 2009, Scarselli et al., 2009]. Recent extensions and improvements to the GNN framework include approaches to incorporate different local structures (around subgraphs), e.g., [Abu-El-Haija et al., 2019, Flam-Shepherd et al., 2020, Jin et al., 2019, Niepert et al., 2016, Xu et al., 2018], novel techniques for pooling node representations in order to perform graph classification, e.g., [Cangea et al., 2018, Gao and Ji, 2019, Ying et al., 2018, Zhang et al., 2018], incorporating distance information [You et al., 2019], and non-Euclidean geometry approaches [Chami et al., 2019]. Moreover, recently, empirical studies on neighborhood-aggregation functions for continuous node features [Corso et al., 2020], edge-based GNNs leveraging physical knowledge [Anderson et al., 2019, J. Klicpera, 2020, Klicpera et al., 2021], and sparsification methods [Rong et al., 2020] emerged. Surveys of recent advancements in GNN techniques can be found, e.g., in [Chami et al., 2020, Wu et al., 2019, Zhou et al., 2018].
Limits of GNNs and more expressive architectures

Recently, connections of GNNs to Weisfeiler–Leman type algorithms have been shown [Azizian and Lelarge, 2020, Barceló et al., 2020, Chen et al., 2019, Geerts et al., 2020, Geerts, 2020, Maehara and NT, 2019, Maron et al., 2019, Morris et al., 2019, Xu et al., 2019]. Specifically, [Morris et al., 2019, Xu et al., 2019] showed that the expressive power of any possible GNN architecture is limited by the 1-WL in terms of distinguishing non-isomorphic graphs.

Triggered by the above results, a large set of papers proposed architectures to overcome the expressivity limitations of the 1-WL. Morris et al. [2019] introduced $k$-dimensional GNNs ($k$-GNN) which rely on a message-passing scheme between subgraphs of cardinality $k$. Similarly to [Morris et al., 2017], the paper employed a local, set-based (neural) variant of the $k$-WL. Later, this was refined in [Maron et al., 2019, Azizian and Lelarge, 2020] by introducing $k$-order folklore graph neural networks ($k$-FGNN), which are equivalent to the folklore or oblivious variant of the $k$-WL [Grohe, 2021, Morris et al., 2021] in terms of distinguishing non-isomorphic graphs. Subsequently, Morris et al. [2020] introduced neural architectures based on the $\delta$-$k$-LWL, which only considers a subset of the neighborhood from the $k$-WL, taking the potential sparsity of the underlying graph (to some extent) into account. Although more scalable, the algorithm reaches computational exhaustion on large graphs since it considers all $n^k$ tuples of size $k$. Chen et al. [2019] connected the theory of universal approximations of permutation-invariant functions and the graph isomorphism viewpoint and introduced a variation of the 2-WL. See [Morris et al., 2021] for an in-depth survey on this topic.

Recent work has extended the expressive power of GNNs, e.g., by encoding node identifiers [Murphy et al., 2019, Vignac et al., 2020], leveraging random features [Abboud et al., 2020, Dasoulas et al., 2020, Sato et al., 2020], subgraph information [Bevilacqua et al., 2021, Bouritsas et al., 2020, Cotta et al., 2021, Papp et al., 2021, Thiede et al., 2021, You et al., 2021, Zhang and Li, 2021, Zhao et al., 2021], homomorphism counts [Barceló et al., 2021, NT and Maehara, 2020], spectral information [Balcilar et al., 2021], simplicial and cellular complexes [Bodnar et al., 2021], random walks [Tönshoff et al., 2021], graph decompositions [Talak et al., 2021], distance [Li et al., 2020] and directional information [Beaini et al., 2020].

However, all of the above approaches mentioned in the previous paragraph only overcome limitations of the 1-WL, 2-WL, or 3-WL, and do not induce a hierarchy of provably powerful, permutation-equivariant neural architectures aligned with the $k$-WL hierarchy.

Theory

The Weisfeiler–Leman algorithm constitutes one of the earliest and most natural approaches to isomorphism testing [Weisfeiler, 1976, Weisfeiler and Leman, 1968], having been heavily investigated by the theory community over the last few decades [Grohe, 2017]. Moreover, the fundamental nature of the $k$-WL is evident from a variety of connections to other fields such as logic, optimization, counting complexity, and quantum computing. The power and limitations of the $k$-WL can be neatly characterized in terms of logic and descriptive complexity [Babai, 1979, Immerman and Lander, 1990], Schr"{o}dinger-Adams relaxations of the natural integer linear program for the graph isomorphism problem [Atserias and Maneva, 2013, Grohe and Otto, 2015, Malkin, 2014], homomorphism counts [Dell et al., 2018], and quantum isomorphism games [Atserias et al., 2019]. In their seminal paper, Cai et al. [1992] showed that, for each $k$, there exists a pair of non-isomorphic graphs of size $\mathcal{O}(k)$ that are not distinguished by the $k$-WL. [Kiefer, 2020] gives a thorough survey of more background and related results concerning the expressive power of the $k$-WL. For $k = 1$, the power of the algorithm has been completely characterized [Arvind et al., 2015, Kiefer et al., 2015]. Moreover, upper bounds on the running
time [Berkholz et al., 2017] and the number of iterations for \( k = 1 \) [Kiefer and McKay, 2020] and for the folklore \( k = 2 \) [Kiefer and Schweitzer, 2016, Lichter et al., 2019] have been shown. For \( k \) in \( \{1, 2\} \), Arvind et al. [2019] studied the abilities of the (folklore) \( k\)-WL to detect and count fixed subgraphs, extending the work of F"urer [2017]. The former was refined in [Chen et al., 2020]. Kiefer et al. [2019] showed that the folklore 3-WL completely captures the structure of planar graphs. The algorithm (for logarithmic \( k \)) plays a prominent role in the recent result of Babai [2016] improving the best known running time for the graph isomorphism problem. Recently, Grohe et al. [2020] introduced the framework of Deep Weisfeiler–Lemans algorithms, which allow the design of a more powerful graph isomorphism test than Weisfeiler–Lemans type algorithms. Finally, the emerging connections between the Weisfeiler–Lemans paradigm and graph learning are described in two recent surveys [Grohe, 2020, Morris et al., 2021].

2 Preliminaries

As usual, for \( n \geq 1 \), let \([n] := \{1, \ldots, n\} \subset \mathbb{N}\). We use \( \{\ldots\} \) to denote multisets, i.e., the generalization of sets allowing for multiple instances for each of its elements.

**Graphs** A graph \( G \) is a pair \((V(G), E(G))\) with finite sets of nodes \( V(G) \) and edges \( E(G) \subseteq \{(u, v) \subseteq V(G) \mid u \neq v\} \). If not otherwise stated, we set \( n := |V(G)| \). For ease of notation, we denote the edge \( (u, v) \) in \( E(G) \) by \((u, v) \) or \((v, u) \). In the case of directed graphs, \( E \subseteq \{(u, v) \in V \times V \mid u \neq v\} \). A labeled graph \( G \) is a triple \((V, E, \ell)\) with a label function \( \ell : V(G) \cup E(G) \to \mathbb{N} \). Then \( \ell(v) \) is a label of \( v \) for \( v \) in \( V(G) \cup E(G) \). The neighborhood of \( v \) in \( V(G) \) is denoted by \( \delta(v) = \{u \in V(G) \mid \{u, v\} \in E(G)\} \) and the degree of a node \( v \) is \( |\delta(v)| \). For \( S \subseteq V(G) \), the graph \( G[S] = (S, E_S) \) is the subgraph induced by \( S \), where \( E_S = \{(u, v) \in E(G) \mid u, v \in S\} \). A connected component of a graph \( G \) is an inclusion-wise maximal subgraph of \( G \) in which every two nodes are connected by paths. A tree is a connected graph without cycles. A rooted tree is an oriented tree with a designated node called root, in which the edges are directed away from the root. Let \( p \) be a node in a rooted tree. Then we call its out-neighbors children with parent \( p \). We denote an undirected cycle on \( k \) nodes by \( C_k \). Given two graphs \( G \) and \( H \) with disjoint node sets, we denote their disjoint union by \( G \cup H \).

Two graphs \( G \) and \( H \) are isomorphic and we write \( G \simeq H \) if there exists a bijection \( \varphi : V(G) \to V(H) \) preserving the adjacency relation, i.e., \((u, v) \) is in \( E(G) \) if and only if \((\varphi(u), \varphi(v)) \) is in \( E(H) \). Then \( \varphi \) is an isomorphism between \( G \) and \( H \). Moreover, we call the equivalence classes induced by \( \simeq \) isomorphism types, and denote the isomorphism type of \( G \) by \( \tau(G) \). In the case of labeled graphs, we additionally require that \( \ell(v) = \ell(\varphi(v)) \) for \( v \) in \( V(G) \) and \( \ell((u, v)) = \ell((\varphi(u), \varphi(v))) \) for \( (u, v) \) in \( E(G) \). Let \( v \) be a tuple in \( V(G)^k \) for \( k > 0 \), then \( G[v] \) is the subgraph induced by the multiset of elements of \( v \), where the nodes are labeled with integers from \( \{1, \ldots, k\} \) corresponding to their positions in \( v \).

**Equivariance** For \( n > 0 \), let \( S_n \) denote the set of all permutations of \([n]\), i.e., the set of all bijections from \([n]\) to itself. For \( \sigma \) in \( S_n \) and a graph \( G \) such that \( V(G) = [n] \), let \( \sigma \cdot G \) be the graph such that \( V(\sigma \cdot G) = \{v_{\sigma^{-1}(1)}, \ldots, v_{\sigma^{-1}(n)}\} \) and \( E(\sigma \cdot G) = \{(v_{\sigma^{-1}(i)}, v_{\sigma^{-1}(j)}) \mid (v_i, v_j) \in E(G)\} \). That is, applying the permutation \( \sigma \) reorders the nodes. Hence, for two isomorphic graphs \( G \) and \( H \) on the same vertex set, i.e., \( G \simeq H \), there exists \( \sigma \) in \( S_n \) such that \( \sigma \cdot G = H \). Let \( \mathcal{G}_n \) denote the set of all graphs on \( n \) nodes. A function \( f : \mathcal{G} \to \mathbb{R} \) is invariant if for every \( n > 0 \) and every \( \sigma \) in \( S_n \) and every graph \( G \) of order \( n \), it
holds that \( f(\sigma \cdot G) = f(G) \). A function \( f : \mathcal{G} \to \mathcal{G} \) is equivariant if for every \( n > 0 \), \( f(\mathcal{G}_n) \subseteq \mathcal{G}_n \) and for every \( \sigma \) in \( S_n \), \( f(\sigma \cdot G) = \sigma \cdot f(G) \).

**Kernels**  A kernel on a non-empty set \( \mathcal{X} \) is a symmetric, positive semidefinite function \( k : \mathcal{X} \times \mathcal{X} \to \mathbb{R} \). Equivalently, a function \( k : \mathcal{X} \times \mathcal{X} \to \mathbb{R} \) is a kernel if there is a feature map \( \phi : \mathcal{X} \to \mathcal{H} \) to a Hilbert space \( \mathcal{H} \) with inner product \( \langle \cdot, \cdot \rangle \) such that \( k(x,y) = \langle \phi(x), \phi(y) \rangle \) for all \( x \) and \( y \) in \( \mathcal{X} \). A graph kernel is a kernel on the set \( \mathcal{G} \) of all graphs.

### 2.1 Node-refinement algorithms

In the following, we briefly describe the Weisfeiler–Leman algorithm and related variants [Morris et al., 2020]. Let \( k \) be a fixed positive integer. There exist two definitions of the \( k\text{-WL} \), the so-called oblivious \( k\text{-WL} \) and the folklore or non-oblivious \( k\text{-WL} \), see, e.g., [Grohe, 2021]. There is a subtle difference in how they aggregate neighborhood information. Within the graph learning community, it is customary to abbreviate the oblivious \( k\text{-WL} \) as \( k\text{-WL} \), a convention that we follow in this paper.

We proceed to the definition of the \( k\text{-WL} \). Let \( V(G)^k \) denote the set of \( k\)-tuples of nodes of the graph \( G \). A coloring of \( V(G)^k \) is a mapping \( C : V(G)^k \to \mathbb{N} \), i.e., we assign a number (color) to every tuple in \( V(G)^k \). The initial coloring \( C_0 \) of \( V(G)^k \) is specified by the atomic types of the tuples. So two tuples \( \mathbf{v} \) and \( \mathbf{w} \) in \( V(G)^k \) have the same initial color if the mapping \( v_i \mapsto w_i \) induces an isomorphism between the labeled subgraphs \( G[\mathbf{v}] \) and \( G[\mathbf{w}] \). Note that, given a tuple \( \mathbf{v} \) in \( V(G)^k \), we can upper-bound the running time of the computation of this initial coloring for \( \mathbf{v} \) by \( O(k^2) \). A color class corresponding to a color \( c \) is the set of all tuples colored \( c \), i.e., the set \( C^{-1}(c) \).

For \( j \) in \([k]\) and \( w \) in \( V(G) \), let \( \phi_j(\mathbf{v}, w) \) be the \( k\)-tuple obtained by replacing the \( j\)th component of \( \mathbf{v} \) with the node \( w \). That is, \( \phi_j(\mathbf{v}, w) = (v_1, \ldots, v_{j-1}, w, v_{j+1}, \ldots, v_k) \). If \( \mathbf{w} = \phi_j(\mathbf{v}, w) \) for some \( w \) in \( V(G) \), call \( \mathbf{w} \) a \( j\)-neighbor of \( \mathbf{v} \). The neighborhood of \( \mathbf{v} \) is the set of all \( \mathbf{w} \) such that \( \mathbf{w} = \phi_j(\mathbf{v}, w) \) for some \( j \) in \([k]\) and a \( w \) in \( V(G) \).

The refinement of a coloring \( C : V(G)^k \to \mathbb{N} \), denoted by \( \hat{C} \), is the coloring \( \hat{C} : V(G)^k \to \mathbb{N} \) defined as follows. For each \( j \) in \([k]\), collect the colors of the \( j\)-neighbors of \( \mathbf{v} \) in a multiset \( S_j := \{ C(\phi_j(\mathbf{v}, w)) \mid w \in V(G) \} \). Then, for a tuple \( \mathbf{v} \), define

\[
\hat{C}(\mathbf{v}) := (C(\mathbf{v}), M(\mathbf{v})),
\]

where \( M(\mathbf{v}) \) is the \( k\)-tuple \((S_1, \ldots, S_k)\). For consistency, the obtained strings \( \hat{C}(\mathbf{v}) \) are lexicographically sorted and renamed as integers, not used in previous iterations. Observe that the new color \( \hat{C}(\mathbf{v}) \) is solely dictated by the color histogram of the neighborhood of \( \mathbf{v} \). In general, a different mapping \( M(\cdot) \) could be used, depending on the neighborhood information that we would like to aggregate. We will refer to a mapping \( M(\cdot) \) as an aggregation map.

**\( k\)-dimensional Weisfeiler–Leman**  For \( k \geq 2 \), the \( k\text{-WL} \) computes a coloring \( C_{\infty} : V(G)^k \to \mathbb{N} \) of a given graph \( G \), as described next.\(^2\) To begin with, the initial coloring \( C_0 \) is computed. Then, starting with \( C_0 \), successive refinements \( C_{i+1} = \hat{C}_i \) are computed until convergence. More precisely,

\[
C_{i+1}(\mathbf{v}) := (C_i(\mathbf{v}), M_i(\mathbf{v})),
\]

\(^2\)We define the 1-WL in the next subsection.
We define the successive refinement steps are also called *rounds* or *iterations*. Since the color classes form a partition of $V(G)^k$, there must exist a finite $\ell \leq |V(G)|^k$ such that $C_\ell$ and $\tilde{C}_\ell$ induce the same partition on the vertex tuples, i.e., the partition induced by $C_\ell$ cannot be refined further. The $k$-WL outputs this $C_\ell$ as the stable coloring $C_\infty$.

The $k$-WL distinguishes two graphs $G$ and $H$ if, upon running the $k$-WL on their disjoint union $G \cup H$, there exists a color $c$ in $\mathbb{N}$ in the stable coloring such that the corresponding color class $S_c$ satisfies

$$|V(G)^k \cap S_c| \neq |V(H)^k \cap S_c|,$$

i.e., the numbers of $c$-colored tuples in $V(G)^k$ and $V(H)^k$ differ. Two graphs that are distinguished by the $k$-WL must be non-isomorphic, because the algorithm is defined in an isomorphism-invariant way.

Finally, the application of different aggregation maps $M$ yields related versions of $k$-WL. For example, setting $M(\cdot)$ to be

$$M^F(v) = \{(C(\phi_1(v, w)), \ldots, C(\phi_k(v, w))) \mid w \in V(G)\},$$

yields the so-called folklore version of $k$-WL (see e.g., [Cai et al., 1992]). It is known that the oblivious version of the $k$-WL is as powerful as the folklore version of the $(k-1)$-WL [Grohe, 2021].

**Local $\delta$-$k$-dimensional Weisfeiler–Leman algorithm** Morris et al. [2020] introduced a more efficient variant of the $k$-WL, the local $\delta$-$k$-dimensional Weisfeiler–Leman algorithm ($\delta$-$k$-LWL). In contrast to the $k$-WL, the $\delta$-$k$-LWL considers only a subset of the entire neighborhood of a node tuple. Let the tuple $w = \phi_j(v, w)$ be a $j$-neighbor of $v$. We say that $w$ is a local $j$-neighbor of $v$ if $w$ is adjacent to the replaced node $v_j$. Otherwise, the tuple $w$ is a global $j$-neighbor of $v$. The $\delta$-$k$-LWL considers only local neighbors during the neighborhood aggregation process, and discards any information about the global neighbors. Formally, the $\delta$-$k$-LWL refines a coloring $C^k(v)$ (obtained after $i$ rounds of the $\delta$-$k$-LWL) via the aggregation function

$$M^\delta_i(v) = \{\{C^k_i(\phi_1(v, w)) \mid w \in \delta(v_1)\}, \ldots, \{C^k_i(\phi_k(v, w)) \mid w \in \delta(v_k)\}\},$$

hence considering only the local $j$-neighbors of the tuple $v$ in each iteration. The coloring functions for the iterations of the $\delta$-$k$-LWL are then defined by

$$C^k_{i+1}(v) = (C^k_i(v), M^\delta_i(v)).$$

We define the 1-WL to be the $\delta$-1-LWL, which is commonly known as Color Refinement or Naïve Node Classification.

\[\text{for a node } v \text{ in } V(G).\]

\[\text{Strictly speaking, the 1-WL and Color Refinement are two different algorithms. That is, the 1-WL considers neighbors and non-neighbors to update the coloring, resulting in a slightly higher expressivity when distinguishing nodes in a given graph, see [Grohe, 2021] for details.}\]
Morris et al. [2020] also defined the $\delta$-$k$-LWL$^+$, a minor variation of the $\delta$-$k$-LWL. Formally, the $\delta$-$k$-LWL$^+$ refines a coloring $C_i$ (obtained after $i$ rounds) via the aggregation function

$$M^{\delta+}(v) = \{ (c_i^{k,\delta}(\phi_1(v, w)), \#(v, \phi_1(v, w))) | w \in \delta(v_1) \}, \ldots,$$

$$\{ (c_i^{k,\delta}(\phi_k(v, w)), \#(v, \phi_k(v, w))) | w \in \delta(v_k) \},$$

instead of the $\delta$-k-LWL aggregation defined in Equation (2). Here, we set

$$\#_i^j(v, x) := |\{w : w \sim_j v, C_i^{k,\delta}(w) = C_i^{k,\delta}(x)\}|,$$

where $w \sim_j v$ denotes that $w$ is a $j$-neighbor of $v$, for $j$ in $[k]$. Essentially, $\#_i^j(v, x)$ counts the number of (local or global) $j$-neighbors of $v$ which have the same color as $x$ under the coloring $C_i$ (i.e., after $i$ rounds). Morris et al. [2020] showed that the $\delta$-$k$-LWL$^+$ is slightly more powerful than the $k$-WL in distinguishing non-isomorphic graphs.

### The Weisfeiler–Leman hierarchy and permutation-invariant function approximation

The Weisfeiler–Leman hierarchy is a purely combinatorial algorithm for testing graph isomorphism. However, the graph isomorphism function, mapping non-isomorphic graphs to different values, is the hardest to approximate permutation-invariant function. Hence, the Weisfeiler–Leman hierarchy has strong ties to GNNs’ capabilities to approximate permutation-invariant or equivariant functions over graphs. For example, Morris et al. [2019], Xu et al. [2019] showed that the expressive power of any possible GNN architecture is limited by 1-WL in terms of distinguishing non-isomorphic graphs. Azizian and Lelarge [2020] refined these results by showing that if an architecture is capable of simulating $k$-WL and allows the application of universal neural networks on vertex features, it will be able to approximate any permutation-equivariant function below the expressive power of $k$-WL; see also [Chen et al., 2019]. Hence, if one shows that one architecture distinguishes more graphs than another, it follows that the corresponding GNN can approximate more functions. These results were refined in [Geerts and Reutter, 2022] for Color Refinement and taking into account the number of iterations of $k$-WL.

### 3 The $(k, s)$-LWL algorithm

Since both $k$-WL and its local variant $\delta$-$k$-LWL consider all $k$-tuples of a graph, they do not scale to large graphs for larger $k$. Specifically, for an $n$-node graph, the memory requirement is in $\Omega(n^k)$. Further, since the $k$-WL considers the graph structure only at initialization, it does not adapt to its sparsity, i.e., it does not run faster on sparser graphs. To address this issue, we introduce the $(k, s)$-LWL. The algorithm offers more fine-grained control over the trade-off between expressivity and scalability by only considering a subset of all $k$-tuples, namely those inducing subgraphs with at most $s$ connected components. This combinatorial algorithm will be the basis of the permutation-equivariant neural architectures of Section 4.

Let $G$ be a graph. Then $\#\text{com}(G)$ denotes the number of (connected) components of $G$. Further, let $k \geq 1$ and $1 \leq s \leq k$, then

$$V(G)^k_s := \{ v \in V(G)^k | \#\text{com}(G[v]) \leq s \}$$

is the set of $(k, s)$-tuples of nodes, i.e, $k$-tuples which induce (sub-)graphs with at most $s$ (connected) components.
In contrast to the algorithms of Section 2.1, the \((k, s)\)-LWL colors tuples from \(V(G)^k\) instead of the entire \(V(G)^k\). Hence, analogously to Section 2.1, a coloring of \(V(G)^k\) is a mapping \(C^{k,s}: V(G)^k \rightarrow \mathbb{N}\), assigning a number (color) to every tuple in \(V(G)^k\). The initial coloring \(C_0^{k,s}\) of \(V(G)^k\) is defined in the same way as before, i.e., specified by the isomorphism types of the tuples, see Section 2.1. Subsequently, the coloring is updated using the \(\delta\)-k-LWL aggregation map, see Equation (2). Hence, the \((k, s)\)-LWL is a variant of the \(\delta\)-k-LWL considering only \((k, s)\)-tuples, i.e., Equation (2) is replaced with

\[
M_i^{\delta,k,s}(\mathbf{v}) := \{\{C_i^{k,s}(\phi_1(\mathbf{v}, w)) | w \in \delta(v_1) \text{ and } \phi_1(\mathbf{v}, w) \in V(G)^k_s\}, \ldots ,
\{C_i^{k,s}(\phi_k(\mathbf{v}, w)) | w \in \delta(v_k) \text{ and } \phi_k(\mathbf{v}, w) \in V(G)^k_s\}\},
\]

i.e., \(M_i^\delta(\mathbf{v})\) restricted to colors of \((k, s)\)-tuples. The stable coloring \(C^{k,s}_\infty\) is defined analogously to the stable coloring \(C^{k}_\infty\). In the following two subsections, we investigate the properties of the algorithm in detail.

Analogously to the \(\delta\)-k-LWL\(^+\), we also define the \((k, s)\)-LWL\(^+\) using

\[
M_i^{k,+}(\mathbf{v}) = \{\{C_i^{k,s}(\phi_1(\mathbf{v}, w)), \#_{i,s}(\mathbf{v}, \phi_1(\mathbf{v}, w))) | w \in \delta(v_1) \text{ and } \phi_1(\mathbf{v}, w) \in V(G)^k_s\}, \ldots ,
\{C_i^{k,s}(\phi_k(\mathbf{v}, w)), \#_{i,s}(\mathbf{v}, \phi_k(\mathbf{v}, w))) | w \in \delta(v_k) \text{ and } \phi_k(\mathbf{v}, w) \in V(G)^k_s\}\),
\]

where the function

\[
\#_{i,s}(\mathbf{v}, \mathbf{x}) = |\{\mathbf{w}: \mathbf{w} \sim_j \mathbf{v}, C_i^{k,s}(\mathbf{w}) = C_i^{k,s}(\mathbf{x}) \text{ and } \mathbf{w} \in V(G)^k_s\}|,
\]

restricts \(\#_{i,s}(\mathbf{v}, \mathbf{x})\) to \((k, s)\)-tuples.

### 3.1 Expressivity

Here, we investigate the expressivity of the \((k, s)\)-LWL, i.e., its ability to distinguish non-isomorphic graphs, for different choices of \(k\) and \(s\). In Section 4, we will leverage these results to devise universal, permutation-equivariant graph networks. We start off with the following simple observation. Since the \((k, k)\)-LWL colors all \(k\)-tuples, it is equal to the \(\delta\)-k-LWL.

**Observation 1.** Let \(k \geq 1\), then

\((k, k)\)-LWL \(\equiv\) \(\delta\)-k-LWL and \((1, 1)\)-LWL \(\equiv\) \(\delta\)-k-LWL \(\equiv\) 1-WL.

The following result shows that the \((k, 1)\)-LWL form a hierarchy, i.e., the algorithm becomes more expressive as \(k\) increases.

**Theorem 1.** Let \(k \geq 1\), then

\((k + 1, 1)\)-LWL \(\subset\) \((k, 1)\)-LWL.

Moreover, we also show that the \((k, 2)\)-LWL is more expressive than the \((k, 1)\)-LWL.

**Proposition 2.** For \(k \geq 2\), it holds that

\((k, 2)\)-LWL \(\subset\) \((k, 1)\)-LWL.
Further, the following theorem yields that increasing the parameter $s$ results in higher expressivity. Formally, we show that the $(k,k)$-LWL is strictly more expressive than the $(k,2)$-LWL.

**Theorem 3.** For $k \geq 2$, it holds that

$$(k,k)\text{-LWL} \subsetneq (k,2)\text{-LWL}.$$ 

See Section 3.2 for an analysis of the asymptotic running time of the $(k,s)$-LWL, showing that it only depends on $k$, $s$, and the sparsity of the graph. In particular, the running time of the $(k,s)$-LWL on an $n$-vertex graph of bounded degree is $\tilde{O}(n^s)$ instead of the usual $\tilde{O}(n^k)$ for the $k$-WL, for fixed $k$ and $s$.

### 3.1.1 Proofs of Theorems 1 and 3 and Proposition 2

To prove Theorem 1, we introduce the $(k,s)$-tuple graph. It essentially contains all $(k,s)$-tuples as nodes, where each node $v_t$ is labeled by the isomorphism type of the $(k,s)$-tuple $t$. We join two nodes by an edge, labeled $j$, if the underlying $(k,s)$-tuples are $j$-neighbors. The formal definition of the $(k,s)$-tuple graph is as follows. Recall that $\tau$ denotes an isomorphism type.

**Definition 4.** Let $G$ be a graph and let $k \geq 1$, and $s$ in $[k]$. Further, let $s$ and $t$ be tuples in $V(G)^k_s$. Then the directed, labeled $(k,s)$-tuple graph $T^k_s(G) = (V_T, E_T, \ell_T)$ has node set $V_T = \{v_t | t \in V(G)^k_s\}$, and

$$(v_s, v_t) \in E_T \iff t = \phi_j(s, w) \text{ holds for some } j \in [k] \text{ and some } w \in V(G).$$

We set $\ell_T((v_s, v_t)) := j$ if $t$ is a local $j$-neighbor of $s$, and let $\ell_T(v_s) := \tau_{G[s]}$.

Given a graph $G$ and the corresponding $(k,s)$-tuple graph $T^k_s(G)$, we define a variant of the 1-WL, which takes into account edge labels. Namely, for $v_t$ in $V_T$, the new algorithm uses the colorings $C^1_0,\delta,s(v_t) = \tau_{G[t]}$ and

$$C^1_{i+1}(v_t) = (C^1_{i-1}(v_t), \{C^1_{i-1}(v_s), \ell(v_t, v_s)) | v_s \in \delta(v_t)\}$$

for $i > 0$. Note that the 1-WL, see Equation (4), and the variant defined via Equation (9) have the same asymptotic running time. The following lemma states that the $(k,s)$-LWL can be simulated on the $(k,s)$-tuple graph using the above variant of the 1-WL.

**Lemma 5.** Let $G$ be a graph, $k \geq 1$, and $s$ in $[k]$. Then

$$C^k_s(t) = C^k_s(u) \iff C^1_{i,s}(v_t) = C^1_{i,s}(v_u),$$

for all $i \geq 0$, and all $(k,s)$-tuples $t$ and $u$ in $V(G)^k_s$.

**Proof sketch.** Induction on the number of iterations using Definition 4. □

The unrolling of a neighborhood around a node of a given graph to a tree is defined as follows, see Figure 2 for an illustration.
Figure 2: Illustration of the unrolling operation around the node \( a \) for \( i = 2 \).

**Definition 6.** Let \( G = (V, E, \ell) \) be a labeled (directed) graph and let \( v \) be in \( V \). Then \( U^i_G,v = (W^i, F^i, l^i) \) for \( i \geq 0 \) denotes the unrolled tree \( G \) around \( v \) at depth \( i \), where

\[
W^i = \begin{cases} \{v(0,v)\} & \text{if } i = 0 \\ W^{i-1} \cup \{u(i_{(i-1,p)},w(i_{(i-1,p)}) \mid u \in \delta(w) \text{ for } w(i_{(i-1,p)}) \in W^{i-1}\} & \text{otherwise,} \end{cases}
\]

and

\[
F^i = \begin{cases} \emptyset & \text{if } i = 0 \\ F^{i-1} \cup \{(w(i_{(i-1,p)}) \mid u \in \delta(w) \text{ for } w(i_{(i-1,p)}) \in W^{i-1}\} & \text{otherwise.} \end{cases}
\]

The label function is defined as \( l_i(u(j,p)) = \ell(u) \) for \( u \) in \( V \), and \( l_i(u(j,w)) = \ell((w,u)) \). For notational convenience, we usually omit the subscript \( i \).

In the following, we use the unrolled tree for the above defined \((k,s)\)-tuple graph. For \( k \geq 2 \) and \( s \) in \([k]\), we denote the directed, unrolled tree of the \((k,s)\)-tuple graph of \( G \) around the node \( v_t \) at depth \( i \) for the tuple \( t \) in \( V(G)^k_s \) by \( U^i_{T^k_s(G),v_t} \). For notational convenience, we write \( U^i_{T,v_t} \) for \( U^i_{T^k_s(G),v_t} \). Further, for two \((k,s)\)-tuples \( t \) and \( u \), we write

\[
U^i_{T,v_t} \simeq_{v_t \rightarrow v_u} U^i_{T,v_u} \tag{10}
\]

if there exists a (labeled) isomorphism \( \varphi \) between the two unrolled trees, mapping the (root) node \( v_t \) to \( v_u \). Moreover, we need the following two results.

**Theorem 7** ([Busacker and Saaty, 1965, Valiente, 2002]). The 1-WL distinguishes any two directed, labeled non-isomorphic trees.

Using the first result, the second one states that the \((k,s)\)-LWL can be simulated by the variant of the 1-WL of Equation (9) on the unrolled tree of the \((k,s)\)-tuple graph, and hence can be reduced to a tree-isomorphism problem.

**Lemma 8.** Let \( G \) be a connected graph, then the \((k,s)\)-LWL colors the tuples \( t \) and \( u \) in \( V(G)^k_s \) equally if and only if the corresponding unrolled \((k,s)\)-tuple trees are isomorphic, i.e.,

\[
C^i_{k,s}(t) = C^i_{k,s}(u) \iff U^i_{T,v_t} \simeq_{v_t \rightarrow v_u} U^i_{T,v_u},
\]

for all \( i \geq 0 \).
Figure 3: The graphs $A_{k+2}$ and $B_{k+2}$ for $k = 4.$

**Proof sketch.** First, by Lemma 5, we can simulate the $(k, s)$-WL for the graph $G$ using the $(k, s)$-tuple graph $T^k_s(G).$ Secondly, consider a node $v_t$ in the $(k, s)$-tuple graph $T^k_s(G)$ and a corresponding node in the unrolled tree around $v_t.$ Observe that the neighborhoods for both nodes are identical. By definition, this holds for all nodes (excluding the leaves) in the unrolled tree. Hence, by Lemma 5, we can simulate the $(k, s)$-WL for each tuple $t$ by running the 1-WL in the unrolled tree around $v_t$ in the $(k, s)$-tuple graph. Since the 1-WL decides isomorphism for trees, see Theorem 7, the result follows.

The following lemma shows that the $(k + 1, 1)$-WL is strictly more expressive than the $(k, 1)$-WL for every $k \geq 2.$

**Lemma 9.** Let $k \geq 2.$ Let $G := C_{2(k+2)}$ and $H := C_{(k+2)} \cup C_{(k+2)}.$ Then, the graphs $G$ and $H$ are distinguished by $(k + 1, 1)$-WL, but they are not distinguished by $(k, 1)$-WL.

**Proof.** We first show that the $(k + 1, 1)$-WL distinguishes the graphs $G$ and $H.$ Let $v$ in $V(G)^{k+1}_1$ be a tuple $(v_1, \ldots, v_{k+1})$ such that $v_1, \ldots, v_{k+1}$ is a path of length $k$ in $G.$ Let $w$ in $V(H)^{k+1}_1$ be a tuple $(w_1, \ldots, w_{k+1})$ such that $w_1, \ldots, w_{k+1}$ is a path of length $k$ in $H.$ By the structure of $H,$ there exists a vertex $w_{k+2}$ in $V(H)$ such that $w_1, \ldots, w_{k+2}$ forms a cycle of length $k + 2.$ We claim that $v$ does not have any local 1-neighbor $x$ in $V(G)^{k+1}_1$ such that $x$ is non-repeating, i.e., every vertex in $x$ is distinct. This holds because replacing the first vertex of $v$ with any other vertex of $G$ will yield a disconnected tuple. On the other hand, $w$ admits a non-repeating, local 1-neighbor, obtained by replacing the first vertex $w_1$ by $w_{k+2}.$ Hence, the $(k + 1, 1)$-WL distinguishes $G$ and $H.$

Next, we show that the $(k, 1)$-WL does not distinguish the graphs $G$ and $H.$ Indeed, for every $j$ in $[k],$ every $k$-tuple $x$ in $V(G)^k_j$ or $V(H)^k_j$ has exactly two local $j$-neighbors, corresponding to the two neighbors $y, z$ of the vertex $x_j.$ The exact number of local $j$-neighbors of $k$ which additionally lie in $V(G)^k_j$ or $V(H)^k_j) depends only on the atomic type of $x,$ since the length of cycles in $G$ and $H$ is at least $k + 2.$ Hence, the $(k, 1)$-WL neighborhood of every tuple in $G$ or $H$ depends only on its atomic type. This implies that the $(k, 1)$-WL does not refine the initial coloring for $G$ as well as $H,$ and hence it does not distinguish $G$ and $H.$

Although Lemma 9 already implies Theorem 1, the construction hinges on the fact that the graphs $G$ and $H$ are not connected. To address this, for $k \geq 2,$ we introduce two connected graphs $A_{k+2}$ and $B_{k+2}$ defined as follows. The graph $A_{k+2}$ has $2(k + 2)$ nodes and $2(k + 2) + 1$ edges, and consists of two disjoint cycles on $k + 2$ nodes connected by a single edge. The graph $B_{k+2}$ also has the same number of nodes and edges, and consists of two cycles on $k + 3$ nodes, each, sharing exactly two adjacent nodes. See Figure 3 for an illustration of the graphs $A_{k+2}$ and $B_{k+2}$ for $k = 4.$ We obtain the following result for the two graphs.

**Lemma 10.** For $k \geq 2,$ the $(k, 1)$-WL does not distinguish the graphs $A_{k+2}$ and $B_{k+2},$ while the $(k + 1, 1)$-WL does.
Proof. We first show the second part, i.e., that the \((k + 1, 1)\)-LWL distinguishes the graphs \(A_{k+2}\) and \(B_{k+2}\). Without loss of generality, assume that \(V(A_{k+2}) = \{a_1, \ldots, a_{2(k+2)}\}\) and that \(E(A_{k+2})\) consists of the edges \((a_i, a_{i+1})\) for \(1 \leq i \leq k + 1\), \((a_i, a_{(k+2)})\), \((a_i, a_{+1})\) for \((k + 3) \leq i \leq 2(k+2) - 1\), and \((a_{k+3}, a_{2(k+2)})\). The two cycles are connected by the edge \((a_{(k+2)}, a_{(k+3)})\) in \(E(A_{k+2})\). Further, assume \(V(B_{k+2}) = \{b_1, \ldots, b_{2(k+2)}\}\) where \((b_i, b_{i+1})\) in \(E(B_{k+2})\) for \(1 \leq i \leq 2(k+2) - 1\) and \((b_1, b_{2(k+2)+2})\) in \(E(B_{k+2})\). Finally, the edge \((b_1, b_{k+3})\) in \(E(B_{k+2})\) is shared by the two cycles of length \(k + 3\) each.

Now, let \(t = (a_1, \ldots, a_{k+1})\) in \(V(A_{k+2})^k_{1+1}\). Observe that the tuple \((a_1, \ldots, a_{k+2})\) is a \((k + 1)\)-neighbor of the tuple \(t\), inducing a graph on \(k + 1\) nodes. Further, since the two cycles in the graph \(B_{k+2}\) have length \(k+3\), there is no tuple without repeated nodes that has a \((k + 1)\)-neighbor without repeated nodes. Hence, the two graphs are distinguished by \((k + 1, 1)\)-LWL.

We now show that the \((k, 1)\)-LWL does not distinguish the graphs \(A_{k+2}\) and \(B_{k+2}\). First, we construct a bijection \(\theta: V(A_{k+2}) \rightarrow V(B_{k+2})\) as induced by the following coloring:

![Diagram](image)

Based on the bijection \(\theta\), we define the bijection \(\theta^k: V(A_{k+2})^k_{1} \rightarrow (A_{B_{k+2}})^k_{1}\), by applying \(\theta\) component-wise to \((k, s)\)-tuples. Observe that \(G[s] \simeq G[\theta^k(s)]\) for \(s\) in \(V(A_{k+2})^k_{1}\).

Claim 11. Let \(s\) be a tuple in \(V(G)^k_{1}\) and \(t = \theta^k(s)\) in \(V(H)^k_{1}\). Let \(N_j(s)\) and \(N_j(t)\) be the \(j\)-neighbors of the tuple \(s\) and \(t\), respectively, for \(j \in [k]\). Then \(\theta^k\) yields a one-to-one correspondence between \(N_j(s)\) and \(N_j(t)\). Consequently, \(G[u] \simeq G[\theta^k(u)]\) for \(u\) in \(N_j(s)\) and \(\theta^k(u)\) in \(N_j(t)\).

Proof. The desired claim follows by observing that the bijective map \(\theta: V(A_{k+2}) \rightarrow V(B_{k+2})\) preserves neighborhoods, i.e. for every \(x\) in \(V(A_{k+2})\), \(\theta(N_F(x)) = N_K(\theta(x))\).

We now again leverage the above claim to show that \(C^k,s_i(s) = C^k,s_i(\theta^k(s))\) for \(i \geq 0\), implying the required result. By a straightforward inductive argument, using Claim 11, we can inductively construct a tree isomorphism between the unrolled trees around the node \(v_s\) and \(v_t\) in the corresponding \((k, s)\)-tuple graph such that \(U^k_{1,v_s} \simeq v_s \rightarrow v_t U^k_{1,v_t}\). By Lemma 8, this implies \(C^k,s_i(s) = C^k,s_i(\theta^k(s))\) for \(i \geq 0\). This shows that the \((k, s)\)-LWL does not distinguish \(A_{k+2}\) and \(B_{k+2}\).

Hence, Theorem 1 directly follows from Lemma 10. Moreover, we also show that the \((k, 2)\)-LWL is more expressive than the \((k, 1)\)-LWL.

Proposition 12. Let \(k \geq 2\). Then

\[(k, 2)\text{-LWL} \subseteq (k, 1)\text{-LWL}.

Proof. As in Lemma 9, let \(G := C_{2(k+2)}\) and \(H := C_{(k+2)} \cup C_{(k+2)}\). By Lemma 9, \(G\) and \(H\) are not distinguished by the \((k, 1)\)-LWL for \(k \geq 2\). We claim that the \((k, 2)\)-LWL distinguishes \(G\) and \(H\) for \(k = 2\) already. Since the \((k, 2)\)-LWL is at least as powerful as the \((2, 2)\)-LWL, this yields the desired claim.

With respect to the \((2, 2)\)-LWL, observe that the \((2, 2)\)-tuple graph \(T^2_2(H)\) consists of four connected components while the \((2, 2)\)-tuple graph \(T^2_2(G)\) consists of a single connected component.
More precisely, there exist two connected components of $T_2^2(H)$ that consist only of 2-tuples containing two non-adjacent nodes which are in the same connected component of the graph $H$. Note that none of these 2-tuples is adjacent to any 2-tuples in $V(H)^2_1$. Moreover, there exists no such connected component in $T_2^2(G)$. Also, note that the number of neighbors of each 2-tuple of the graphs is exactly 4, excluding self loops. Hence, the (2, 2)-LWL will distinguish the two graphs. 

Moreover, the following result shows that increasing the parameter $s$ results in higher expressivity. Formally, we show that the $(k, k)$-LWL is strictly more expressive than the $(k, 2)$-LWL. Note that we use vertex-colored graphs (rather than simple undirected graphs) in our proofs.

**Theorem 13.** Let $k \geq 2$, then

$$(k, k)\text{-LWL} \subsetneq (k, 2)\text{-LWL}.$$ 

For the proof of Theorem 13, we modify the construction employed in [Morris et al., 2020], Appendix C.1.1., where they provide an infinite family of graphs $(G_k, H_k)_{k \in \mathbb{N}}$ such that (a) $k$-WL does not distinguish $G_k$ and $H_k$, although (b) $\delta$-LWL distinguishes $G_k$ and $H_k$. Since our proof closely follows theirs, let us recall some relevant definitions from their paper.

**Construction of $G_k$ and $H_k$.** Let $K$ denote the complete graph on $k + 1$ vertices (without any self-loops). The vertices of $K$ are indexed from 0 to $k$. Let $E(v)$ denote the set of edges incident to $v$ in $K$: clearly, $|E(v)| = k$ for all $v$ in $V(K)$. We call the elements of $V(K)$ base vertices, and the elements of $E(K)$ base edges. Define the graph $G_k$ as follows:

1. For the vertex set $V(G_k)$, we add
   (a) $(v, S)$ for each $v$ in $V(K)$ and for each even subset $S$ of $E(v)$,
   (b) two vertices $e^1, e^0$ for each edge $e$ in $E(K)$.
2. For the edge set $E(G_k)$, we add
   (a) an edge $\{e^0, e^1\}$ for each $e$ in $E(K)$,
   (b) an edge between $(v, S)$ and $e^1$ if $v$ in $e$ and $e$ in $S$,
   (c) an edge between $(v, S)$ and $e^0$ if $v$ in $e$ and $e$ not in $S$,

For every $v$ in $K$, the set of vertices of the form $(v, S)$ is called the vertex cloud for $v$. Similarly, for every edge $e$ in $E(K)$, the set of vertices of the form $\{e^0, e^1\}$ is called the edge cloud for $e$.

Define a companion graph $H_k$, in a similar manner to $G_k$, with the following exception: in Step 1(a), for the vertex 0 in $V(K)$, we choose all odd subsets of $E(0)$. Counting vertices, we find that $|V(G)| = |V(H)| = (k + 1) \cdot 2^{k-1} + (k+1) \cdot 2$. This finishes the construction of the graphs $G$ and $H$. We set $G_k := G$ and $H_k := H$.

**Distance-two-cliques.** A set $S$ of vertices is said to form a distance-two-clique if the distance between any two vertices in $S$ is exactly 2. The following results were shown in [Morris et al., 2020].

**Lemma 14** ([Morris et al., 2020]). The following holds for the graphs $G_k$ and $H_k$ defined above.

- There exists a distance-two-clique of size $(k + 1)$ inside $G_k$.
- There does not exist a distance-two-clique of size $(k + 1)$ inside $H_k$. 

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Hence, $G_k$ and $H_k$ are non-isomorphic.

**Lemma 15** ([Morris et al., 2020]). The $\delta$-$k$-LWL distinguishes $G_k$ and $H_k$. On the other hand, $k$-WL does not distinguish $G_k$ and $H_k$.

We are ready to present the proof of Theorem 13.

**Proof of Theorem 13.** Observe that the $(k, k)$-LWL is the same as the $\delta$-$k$-LWL. Hence, it suffices to show an infinite family of graphs $(X_k, Y_k)$, $k \in \mathbb{N}$, such that (a) $(k, 2)$-LWL does not distinguish $X_k$ and $Y_k$, although (b) $\delta$-$k$-LWL distinguishes $X_k$ and $Y_k$.

Let $X_k$ be the graph obtained from the graph $G_k$ as follows. First, for every base vertex $v$ in $V(K)$, every vertex of $V(G_k)$ in the vertex cloud for $v$ receives a color Red. Hence, vertex clouds form color classes, where each such class has a distinct color. Similarly, for every base edge $e$ in $E(K)$, every vertex of $V(G_k)$ in the edge cloud for $e$ receives a color Blue. Finally, let $\Delta > 3k$. Then, we replace every edge $e$ in $G_k$ by a path of length $\Delta$, such that every vertex on this path is colored with the color $\{c, c'\}$, where $c$ and $c'$ are the colors of the endpoints of $e$ in $G_k$. We call such path vertices auxiliary vertices. The graph $Y_k$ is obtained from $H_k$ by an identical construction.

First, we show that the $(k, 2)$-LWL does not distinguish the graphs $X_k$ and $Y_k$. We use a modified version of the bijective $k$-pebble game [Grohe, 2017]: (a) we enforce the $k$ pebbles to form at most two components at any point during the game, and (b) when the Spoiler and Duplicator pick the $i^{th}$ pebble from each graph, the Duplicator is required to exhibit a bijection only between the position-$i$ local neighbourhoods of the two pebbling configuration tuples $x \in X_k^i$ and $y \in Y_k^i$, instead of a bijection between the vertex sets of $X_k$ and $Y_k$.

Observe that there can be at most two vertices out of $k + 1$ vertices in $V(K)$ such that the corresponding vertex clouds contain a tupled vertex, by our choice of $\Delta$. Hence, in the usual parlance of Cai-Fürer-Immerman games [Cai et al., 1992], the twisted edge can always be hidden among the remaining $(k - 1)$ vertices of $K$. This ensures that for all $i \in [k]$, a partial isomorphism between $x \setminus x_i$ and $y \setminus y_i$ can always be extended to a bijective mapping between the $i$-local-neighborhoods of $x$ and $y$. Hence, the Duplicator cannot win this pebble game and therefore, $(k, 2)$-LWL cannot distinguish the graphs $X_k$ and $Y_k$.

Next, we show that the $\delta$-$k$-LWL distinguishes the graphs $X_k$ and $Y_k$. Our proof closely follows the corresponding proof in [Morris et al., 2020]. Instead of showing a discrepancy in the number of distance-two-cliques, we instead use colored-distance-$(2\Delta + 1)$-cliques defined as follows. Let $S$ be a set of vertices belonging to the vertex clouds. The set $S$ is said to form a colored-distance-$(2\Delta + 1)$-clique if any two vertices in $S$ are connected by a path of exactly $2\Delta + 1$ vertices, of which $2\Delta$ are auxiliary vertices and one vertex is a vertex from an edge cloud. Analogously to their proof, it can be shown that (a) there exists a colored-distance-$(2\Delta + 1)$-clique of size $(k + 1)$ inside $X_k$, and (b) there does not exist a colored-distance-$(2\Delta + 1)$-clique of size $(k + 1)$ inside $Y_k$, and hence, (c) $X_k$ and $Y_k$ are non-isomorphic. Finally, we claim that the $\delta$-$k$-LWL is powerful enough to detect colored-distance-$(2\Delta + 1)$-cliques. The proof is analogous to [Morris et al., 2020, Appendix C.1.1, Proof of Lemma 9]. This yields that the $\delta$-$k$-LWL distinguishes the graphs $X_k$ and $Y_k$.  

\[\square\]

### 3.2 Asymptotic running time

In the following, we bound the asymptotic running time of the $(k, s)$-LWL. Due to Lemma 5, we can upper-bound the running time of the $(k, s)$-LWL for a given graph by upper-bounding the
time to construct the \((k, s)\)-tuple graph and running the 1-WL variant of Equation (9) on top. Proposition 17 establishes an upper bound on the asymptotic running time for constructing the \((k, s)\)-tuple graph from a given graph. Thereto, we assume a \(d\)-bounded-degree graph \(G\), for \(d \geq 1\), i.e., each node has at most \(d\) neighbors.

To prove the proposition, we define \((k, s)\)-multisets. Let \(G\) be a graph, \(k \geq 1\), and \(s\) in \([k]\), then the set of \((k, s)\)-multisets

\[
S(G)^k_s = \{ \{v_1, \ldots, v_k\} \mid v \in V(G)^k_s \}
\]

contains the set of multisets inducing subgraphs of \(G\) on at most \(k\) nodes with at most \(s\) components. The following results upper-bounds the running time for the construction of \(S(G)^k_s\).

Algorithm 1 Generate \((k, s)\)-multisets

Input: Graph \(G\), \(k\), \(s\), and \(S(G)^s_s\)
Output: \((k, s)\)-multiset \(S(G)^k_s\)

1: Let \(R\) be an empty set data structure
2: for \(M \in S(G)^s_s\) do
3: \hspace{1em} Let \(S\) be a queue data structure containing only \((M, s)\)
4: \hspace{1em} while \(S\) not empty do
5: \hspace{2em} Pop \((T, c)\) from queue \(S\)
6: \hspace{2em} if \(c + 1 \leq k\) then
7: \hspace{3em} for \(t \in T\) do
8: \hspace{4em} for \(u \in \delta(t) \cup \{t\}\) do
9: \hspace{5em} Add \((T \cup \{u\}, c + 1)\) to \(S\)
10: \hspace{2em} \hspace{1em} else
11: \hspace{2em} \hspace{1em} Add \(T\) to \(R\)
12: \hspace{1em} \hspace{1em} return \(R\)

Proposition 16. Let \(G\) be a \(d\)-bounded-degree graph, \(k \geq 2\), and \(s\) in \([k − 1]\). Then Algorithm 1 computes \(S^k_s(G)\) in time \(\tilde{O}(n^s \cdot k^{k−s}(d + 1)^{k−s})\).

Proof. Let \(c < k\) and let \(T'\) be an element in \(S(G)^{c+1}_s\). By definition of \(S(G)^{c+1}_s\) and \(S(G)^{c+1}_s\), there exists a \(c\)-element multiset \(T\) in \(S(G)^c_s\) such that \(T' = T \cup \{v\}\) is in \(S(G)^{c+1}_s\) for a node \(v\) in \(V(G)\). Since \(s\) is fixed, \(v\) is either in the neighborhood \(\delta(w)\) for \(w\) in \(T\) or \(v = w'\) for \(w'\) in \(T\). Hence, lines 7 to 9 in Algorithm 1 generate \(S(G)^{c+1}_s\) from \(S(G)^c_s\). The set data structure \(R\) makes sure that the final solution will not contain duplicates. The running time follows directly when using, e.g., a red–black tree, to represent the set \(R\).

Based on the above result, we can easily construct \(T^k_s(G)\) from \(S^k_s(G)\), implying the following result.

Proposition 17. Let \(G\) be a \(d\)-bounded-degree graph, \(k \geq 3\), and \(s\) in \([k − 1]\). Then we can compute \(T^k_s(G)\) in time \(O(n^s \cdot k^{k−s}(d + 1)^{k−s+1} \cdot k! \cdot k)\).

Proof. The running time follows directly from Proposition 16. That is, from \(S^k_s(G)\), we can generate the set \(V^k_s(G)\) by generating all permutations of each element in the former. By iterating over each resulting \((k, s)\)-tuple and each component of such \((k, s)\)-tuple, we can construct the needed adjacency information.
Hence, unlike for the $k$-WL, the running time of the $(k, s)$-LWL does not depend on $n^k$ for an $n$-node graph and is solely dictated by $s$, $k$, and the sparsity of the graph.

Moreover, observe that the upper bound given in Proposition 17, by leveraging Lemma 5, also upper-bounds the asymptotic running time for one iteration of the $(k, s)$-LWL.

4 SpeqNets: Sparse, permutation-equivariant graph networks

We can now leverage the above combinatorial insights to derive sparsity-aware, permutation-equivariant graph networks, denoted $(k, s)$-SpeqNet. Given a labeled graph $G$, let each $(k, s)$-tuple $v$ in $V(G)^k_s$ be annotated with an initial feature $f^{(0)}(v)$ determined by its (labeled) isomorphism type, e.g., a one-hot encoding of $\tau_{G[v]}$. Alternatively, we can also use some application-specific, real-valued feature. In each layer $t > 0$, we compute a new feature $f^{(t)}(v)$ as

$$f^{(t-1)}_{\text{mrg}}(f^{(t-1)}(v), f^{(t-1)}_{\text{agg}}(f^{(t-1)}(v_1, w) \mid w \in \delta(v_1) \text{ and } \phi_1(v, w) \in V(G)^k_s), \ldots, f^{(t-1)}(v_k, w) \mid w \in \delta(v_k) \text{ and } \phi_k(v, w) \in V(G)^k_s)), \tag{11}$$

in $\mathbb{R}^{1 \times e}$, where $W^{(t)}_1$ and $W^{(t)}_2$ are learnable parameter matrices from $\mathbb{R}^{d \times e}$ for some $d, e > 0$. Here, $f^{(t)}_{\text{mrg}}$ and $f^{(t)}_{\text{agg}}$ are arbitrary differentiable functions, responsible for merging and aggregating the relevant feature information, respectively. Note that we can naturally handle discrete node and edge labels as well as directed graphs. The following result demonstrates the expressive power of the $(k, s)$-SpeqNet, in terms of distinguishing non-isomorphic graphs.

Theorem 18. Let $(V, E, \ell)$ be a labeled graph, and let $k \geq 1$ and $s$ in $[k]$. Then for all $t \geq 0$, there exists weights $W^{(t)}_1$ and $W^{(t)}_2$ such that

$$C^{k,s}_t(v) = C^{k,s}_t(w) \iff f^{(t)}(v) = f^{(t)}(w).$$

Hence, the following holds for all $k \geq 1$:

$$(k, s)$$-SpeqNet $\equiv (k, s)$-LWL.

Proof sketch. First, observe that the $(k, s)$-LWL can be simulated on an appropriate node- and edge-labeled graph, see Lemma 5. Secondly, following the proof of [Morris et al., 2019, Theorem 2], there exists a parameter matrix $W^{(t)}_2$ such that we can injectively map each multiset in Equation (11), representing the local $j$-neighbors for $j$ in $[k]$, to a $d$-dimensional vector. Moreover, we concatenate $j$ to each such vector to distinguish between different neighborhoods. Again, by [Morris et al., 2019, Theorem 2], there exists a parameter matrix $W^{(t)}_1$ such that we can injectively map the set of resulting $k$ vectors to a unique vector representation. Alternatively, one can concatenate the resulting $k$ vectors and use a multi-layer perceptron to learn a joint lower-dimensional representation.

It is not possible to come up with an architecture, i.e., instantiations of $f^{(t)}_{\text{mrg}}$ and $f^{(t)}_{\text{agg}}$ such that it becomes more powerful than the $(k, s)$-LWL, see [Morris et al., 2019]. However, all results from the previous section can be lifted to the neural setting, see also Section 2.1. Analogously to GNNs, the above architecture can naturally handle continuous node and edge labels. By using
the tools developed in [Azizian and Lelarge, 2020], it is straightforward to show that the above
architecture is universal, i.e., it can approximate any possible permutation-invariant function
over graphs up to an arbitrarily small additive error.

4.1 Node-, edge-, and subgraph-level learning tasks

The above architecture computes representation for $k$-tuples, rendering it mostly suitable for
graph-level learning tasks, e.g., graph classification or regression. However, it is also possible to
derive neural architectures based on the $(k,s)$-LWL for node- and edge-level learning tasks, e.g.,
node or link prediction. Given a graph $G$, to learn a node feature for node $v$, we can simply
pool over the feature learned for $(k,s)$-tuples containing the node $v$ as a component. That is, let $t \geq 0$, then we consider the multisets

$$m^t(v)_i = \{ f^{t-1}(t) \mid t \in V(G)^{k}_s \text{ and } t_i = v \}$$

for $i$ in $[k]$. Hence, to compute a vectorial representation of the node $v$, we compute a vectorial
representation of $m^t(v)_i$ for $i$ in $[k]$, e.g., using a neural architecture for multi-sets, see [Wagstaff
et al., 2021], followed by learning a joint vectorial representation for the node $v$. Again,
by [Azizian and Lelarge, 2020], it is straightforward to show that the above architecture is
universal, i.e., it can approximate any possible permutation-equivariant function over graphs
up to an arbitrarily small additive error. Also, the above approach can be straightforwardly
generalized to learn subgraph representations on an arbitrary number of vertices.

5 Experimental evaluation

Here, we aim to empirically investigate the learning performance of the kernel, see Section 2.1,
and neural architectures, see Section 4, based on the $(k,s)$-LWL, compared with standard kernel
and (higher-order) GNN baselines. Concretely, we aim to answer the following questions.

Q1 Do the $(k,s)$-LWL-based algorithms, both kernel and neural architectures, lead to improved
classification and regression scores on real-world, graph-level benchmark datasets compared
with dense algorithms and standard baselines?

Q2 How does the $(k,s)$-SpeqNet architecture compare to standard GNN baselines on node-
classification tasks?

Q3 To what extent does the $(k,s)$-LWL reduce computation times compared with architectures
induced by the $k$-WL?

Q4 What is the effect of $k$ and $s$ with respect to computation times and predictive performance?

The source code of all methods and evaluation procedures is available at https://www.
github.com/chrsmrrs/speqnets.

Datasets To compare the $(k,s)$-LWL-based kernels, we used the well-known graph-classification
benchmark datasets from [Morris et al., 2020], see Table 1 for dataset statistics and properties.4
To compare the $(k,s)$-SpeqNet architecture with GNN baselines, we used the ALCHEMY [Chen
et al., 2019] and the Qm9 [Ramakrishnan et al., 2014, Wu et al., 2018] graph regression datasets,

4All datasets are publicly available at www.graphlearning.io.
Table 1: Dataset statistics and properties for graph-level prediction tasks, †—Continuous vertex labels following Gilmer et al. [2017], the last three components encode 3D coordinates.

| Dataset   | Number of graphs | Number of classes/targets | Number of nodes | Number of edges | Node labels | Edge labels |
|-----------|------------------|---------------------------|----------------|----------------|-------------|-------------|
| ENZYMES   | 600              | 6                         | 32.6           | 62.1           | ✓           | ✓           |
| IMDB-BINARY | 1000            | 2                         | 19.8           | 96.5           | ×           | ×           |
| IMDB-MULTI | 1500            | 3                         | 13.0           | 65.9           | ×           | ×           |
| MUTAG     | 188              | 2                         | 17.9           | 19.8           | ✓           | ×           |
| NC11      | 4110             | 2                         | 29.9           | 32.3           | ✓           | ×           |
| PTC_FM    | 349              | 2                         | 14.1           | 14.5           | ✓           | ×           |
| PROTEINS  | 1113             | 2                         | 39.1           | 72.8           | ✓           | ×           |
| REDDIT-BINARY | 2000        | 2                         | 429.6          | 497.8          | ×           | ×           |
| ALCHEMY   | 202579           | 12                        | 10.1           | 10.4           | ✓           | ✓           |
| QM9       | 129433           | 12                        | 18.0           | 18.6           | ✓(13+3D)† | ✓(4)        |

Table 2: Dataset statistics and properties for node-level prediction tasks.

| Dataset   | Number of nodes | Number of edges | Number of node features |
|-----------|-----------------|-----------------|-------------------------|
| CORNELL   | 183             | 295             | 1703                    |
| TEXAS     | 183             | 309             | 1703                    |
| WISCONSIN | 251             | 490             | 1703                    |

again see Table 3 for dataset statistics and properties. Following Morris et al. [2020], we opted for not using the 3D-coordinates of the ALCHEMY dataset to solely show the benefits of the (sparse) higher-order structures concerning graph structure and discrete labels. To investigate the performance of the architecture for node classification, we used the WebKB datasets [Pei et al., 2020], see Table 2 for dataset statistics and properties.

Kernels We implemented the \((k, s)\)-LWL and \((k, s)\)-LWL\(^+\) for \(k \in \{2, 3\}\) and \(s \in \{1, 2\}\). We compared our kernels to the Weisfeiler–Leman subtree kernel (1-WL) [Shervashidze et al., 2011], the Weisfeiler–Leman Optimal Assignment kernel (WLOA) [Kriege et al., 2016], the graphlet kernel (GR) [Shervashidze et al., 2009], and the shortest-path kernel [Borgwardt and Kriegel, 2005] (SP). Further, we implemented the higher-order kernels \(\delta-k\)-LWL, \(\delta-k\)-LWL\(^+\), \(\delta-k\)-WL, and \(k\)-WL kernel for \(k \in \{2, 3\}\) as outlined in [Morris et al., 2020]. All kernels were (re-)implemented in C++11. For the graphlet kernel, we counted (labeled) connected subgraphs of size 3. We followed the evaluation guidelines outlined in [Morris et al., 2020].

Neural architectures We used the GIN-\(\varepsilon\) and Gin-\(\varepsilon\)-JK architectures [Xu et al., 2019] as neural baselines. For data with (continuous) edge features, we used a 2-layer MLP to map them to the same number of components as the node features and combined them using summation (GINE and GINE-\(\varepsilon\)). For the evaluation of the \((k, s)\)-SpeqNet neural architectures of Section 4, we implemented them using PYTORCH GEOMETRIC [Fey and Lenssen, 2019], using a Python-wrapped C++11 preprocessing routine to compute the computational graphs for the higher-order GNNs. We used the GIN-\(\varepsilon\) layer to express \(f_{\text{agg}}^W_1\) and \(f_{\text{agg}}^W_2\) of Equation (11). For the GNN baseline for the Qm9 dataset, following [Gilmer et al., 2017], we used a complete graph, computed
pairwise $\ell_2$ distances based on the 3D coordinates, and concatenated them to the edge features. We note here that our intent is not the beat state-of-the-art, physical knowledge-incorporating architectures, e.g., DimeNet [J. Klicpera, 2020] or Cormorant [Anderson et al., 2019], but to solely show the benefits of the local, sparse higher-order architectures compared to the corresponding (1-dimensional) GNN. For the $(k, s)$-SpeqNet architectures, in the case of the QM9 dataset, to compute the initial features, for each $(k, s)$-tuple, we concatenated the node and edge features, computed pairwise $\ell_2$ distances based on the 3D coordinates, and a one-hot encoding of the (labeled) isomorphism type. Finally, we used a 2-layer MLP to learn a joint, initial vectorial representation. For the node-classification experiments, we used mean pooling to implement Equation (12) and a standard GCN or GIN layer for all experiments, including the $(k, s)$-SpeqNet architectures. Further, we used the architectures (SDRF) outlined in [Topping et al., 2021] as baselines.

For the kernel experiments, we computed the (cosine) normalized Gram matrix for each kernel. We computed the classification accuracies using the $C$-SVM implementation of LibSVM [Chang and Lin, 2011], using 10-fold cross-validation. We repeated each 10-fold cross-validation ten times with different random folds and report average accuracies and standard deviations.

Following the evaluation method proposed in [Morris et al., 2020], the $C$-parameter was selected from $\{10^{-3}, 10^{-2}, \ldots, 10^{2}, 10^{3}\}$ using a validation set sampled uniformly at random from the training fold (using 10% of the training fold). Similarly, the numbers of iterations of the $(k, s)$-LWL, $(k, s)$-LWL$^+$, 1-WL, WLOA, $\delta$-$k$-LWL, $\delta$-$k$-LWL$^+$, and $k$-WL were selected from $\{0, \ldots, 5\}$ using the validation set. Moreover, for the $(k, s)$-LWL$^+$ and $\delta$-$k$-LWL$^+$, we only added the label function # on the last iteration to prevent overfitting. We report computation times for the $(k, s)$-LWL, $(k, s)$-LWL$^+$, WLOA, $\delta$-$k$-LWL, $\delta$-$k$-LWL$^+$, and $k$-WL with five refinement steps.

All kernel experiments were conducted on a workstation with 791GB of RAM using a single core. Moreover, we used the GNU C++ Compiler 4.8.5 with the flag -02.

For comparing the kernel approaches to GNN baselines, we used 10-fold cross-validation and again used the approach outlined in [Morris et al., 2020]. The number of components of the (hidden) node features in $\{32, 64, 128\}$ and the number of layers in $\{1, 2, 3, 4, 5\}$ of the GIN and GIN-$\varepsilon$ layer were again selected using a validation set sampled uniformly at random from the training fold (using 10% of the training fold). We used mean pooling to pool the learned node embeddings to a graph embedding and used a 2-layer MLP for the final classification, using a dropout layer with $p = 0.5$ after the first layer of the MLP. We repeated each 10-fold cross-validation ten times with different random folds and report the average accuracies and standard deviations. Due to the different training methods, we do not provide computation times for the GNN baselines.

For the larger molecular regression tasks ALCHEMY and QM9, we closely followed the hyperparameters found in [Chen et al., 2019] and [Gilmer et al., 2017], respectively, for the GINE-$\varepsilon$ layers. That is, we used six layers with 64 (hidden) node features and a set2seq layer [Vinyals et al., 2016] for graph-level pooling, followed by a 2-layer MLP for the joint regression of the twelve targets. We used the same architecture details and hyperparameters for the $(k, s)$-SpeqNet. For the ALCHEMY, we used the subset of 12000 graphs from [Morris et al., 2020]. For both datasets, we uniformly and at random sampled 80% of the graphs for training, and 10% for validation and testing, respectively. Moreover, following [Chen et al., 2019, Gilmer et al., 2017], we normalized the targets of the training split to zero mean and unit variance. We used a single model to predict all targets. Following [J. Klicpera, 2020, Appendix C], we report mean standardized MAE and mean standardized logMAE. We repeated each experiment five
Table 3: Classification accuracies in percent and standard deviations, OOT—Computation did not finish within one day, OOM—Out of memory. The (2, 2)-LWL and (2, 2)-LWL+ are omitted as they are equivalent to the δ-2-LWL and δ-2-LWL+.

| Method | Dataset | ENZYMES | IMDB-BINARY | IMDB-MULTI | MUTAG | NCI1 | PROTEINS | PTC_MR | REDDIT-BINARY |
|--------|---------|---------|-------------|------------|-------|------|----------|--------|---------------|
| GR     |         | 29.9 ± 0.8 | 59.3 ± 0.9 | 39.2 ± 0.6 | 72.5 ± 1.7 | 66.2 ± 0.2 | 71.5 ± 0.5 | 56.6 ± 1.3 |
| SP     |         | 40.3 ± 0.9 | 58.7 ± 0.6 | 39.7 ± 0.3 | 81.7 ± 1.5 | 74.1 ± 0.2 | 75.8 ± 0.7 | 59.6 ± 1.5 |
| 1-WL   |         | 50.6 ± 1.2 | 72.5 ± 0.8 | 50.0 ± 0.8 | 75.9 ± 2.0 | 84.4 ± 0.3 | 73.1 ± 0.6 | 59.3 ± 2.1 |
| WLOA   |         | 57.1 ± 0.8 | 73.2 ± 0.4 | 49.8 ± 0.4 | 84.2 ± 1.2 | 85.2 ± 0.2 | 73.0 ± 0.9 | 60.3 ± 1.9 |
| GIN-ε  |         | 38.7 ± 1.5 | 72.9 ± 0.7 | 49.7 ± 0.7 | 84.1 ± 1.4 | 77.7 ± 0.8 | 72.2 ± 0.6 | 55.2 ± 1.7 |
| GIN-c-JK |       | 39.3 ± 1.6 | 73.0 ± 1.1 | 49.6 ± 0.7 | 83.4 ± 2.0 | 78.3 ± 0.3 | 72.2 ± 0.7 | 56.0 ± 3.2 |
| 2-WL   |         | 37.0 ± 1.0 | 68.1 ± 1.7 | 47.5 ± 0.7 | 85.7 ± 1.6 | 66.9 ± 0.3 | 75.2 ± 0.4 | 60.5 ± 1.1 |
| 3-WL   |         | 42.3 ± 1.1 | 67.1 ± 1.5 | 46.8 ± 0.8 | 85.4 ± 1.5 | 67.0 ± 0.4 | 75.4 ± 0.4 | 61.5 ± 1.3 |

Table 4: Additional experimental results for graph regression and node classification.

| Method     | Dataset   | ALCHEMY (10k) | Qm9         |
|------------|-----------|---------------|-------------|
| GINE-ε     |           | 0.180 ± 0.000 | -1.958 ± 0.047 | 0.079 ± 0.001 | -3.439 ± 0.000 |
| (2, 1)-LWL |           | 0.160 ± 0.001 | -2.010 ± 0.056 | 0.078 ± 0.007 | -2.947 ± 0.171 |
| (2, 2)-LWL |           | 0.115 ± 0.001 | -2.722 ± 0.064 | 0.029 ± 0.001 | -4.881 ± 0.008 |
| (3, 1)-LWL |           | 0.115 ± 0.001 | -1.914 ± 0.079 | 0.008 ± 0.003 | -3.397 ± 0.000 |
| (3, 2)-LWL |           | 0.115 ± 0.002 | -2.767 ± 0.079 | OOM         |             |

(a) Mean MAE (mean std. MAE, logMAE) on large-scale (multi-target) molecular regression tasks.

(b) Classification accuracies and standard deviations for node classification.

Times and report average scores and standard deviations. We used the provided ten training, validation, and test splits for the node-classification datasets. All neural experiments were conducted on a workstation with one GPU card with 32GB of GPU memory.

To compare training and testing times between the (2, 1)-SpeqNet, (2, 2)-SpeqNet, GINE-ε architectures, we trained all three models on ALCHEMY (10k) and Qm9 to convergence, divided by the number of epochs, and calculated the ratio with respect to the average epoch computation time of the (2, 1)-SpeqNet (average computation time of dense or baseline layer divided by average computation time of the (2, 1)-SpeqNet). Contrary to the kernel timing experiments, we did not take into account the time of the preprocessing routine to compute the computational graphs to focus purely on the neural component of the architecture. Clearly, the time for the preprocessing of (k, s)-SpeqNet with small s is much smaller than that of, e.g., the δ-k-WL.
Table 6: Overall computation times for selected datasets in seconds (Number of iterations for WL-based methods: 5), Oot—Computation did not finish within one day (24h), Oom—Out of memory.

| Graph Kernel | Dataset       | ENZYMES | IMDB-Binary | IMDB-Multi | MUTAG | NCI1 | PTC_MR |
|--------------|---------------|---------|-------------|------------|-------|------|--------|
| 1-WL         |               | <1      | <1          | <1         | <1    | 1.9  | <1     |
| 2-WL         |               | 225.9   | 91.2        | 38.3       | 4.3   | 1127.8 | 10.7   |
| 3-WL         |               | 55242.7 | 17565.2     | 4977.1     | 259.8 | Oot  | 1324.2 |
| δ-2-LWL      |               | 25.2    | 27.3        | 19.8       | <1    | 82.2 | 1.1    |
| δ-2-LWL+     |               | 25.6    | 26.1        | 18.5       | <1    | 108.3 | 1.2    |
| δ-3-LWL      |               | 3519.0  | 3560.3      | 1957.5     | 36.5  | 15207.3 | 89.9  |
| δ-3-LWL+     |               | 3674.9  | 3636.5      | 2162.3     | 43.6  | 15945.6 | 111.1 |
| (k,s)-LWL+   | 2               | 1.6     | 12.0        | 11.0       | <1    | 5.8  | <1     |
| (k,s)-LWL+   | 3               | 1.7     | 12.6        | 11.0       | <1    | 6.7  | <1     |
| (k,s)-LWL+   | 4               | 51.1    | 1049.4      | 1238.7     | 1.6   | 120.1 | 2.0    |
| (k,s)-LWL+   | 5               | 937.9   | 2571.1      | 2252.6     | 19.0  | 3502.6 | 29.6   |
| (k,s)-LWL+   | 6               | 1046.1  | 2937.8      | 2572.2     | 22.4  | 3888.7 | 34.4   |

Table 7: Average speed-up ratios over all epochs (training and testing).

| Method       | Dataset       | Alchemy (10k) | Qm9 |
|--------------|---------------|---------------|-----|
| GINE-ε       |               | 0.5           | 1.3 |
| (2,1)-SpeqNet|               | 1.0           | 1.0 |
| (2,2)-SpeqNet|               | 1.3           | 3.4 |

5.1 Results and discussion

In the following, we answer questions Q1 to Q4.

A1 Kernels See Table 3. The (k, s)-LWL for k, s in {2, 3} significantly improves the classification accuracy compared to the k-WL and the δ-k-WL, and the other kernel baselines, while being on par with or better than the δ-2-LWL and δ-3-LWL. The (k, s)-LWL and (k, s)-LWL+ achieve a new state-of-the-art on five out of eight datasets. Our algorithms also perform vastly better than the neural baselines.

Neural architectures See Table 4. On both datasets, all (k, s)-SpeqNet architectures beat the GNN baseline. On the Alchemy dataset, the (2, 2)-SpeqNet and (3, 1)-SpeqNet perform best, while on the Qm9 dataset, the (2, 2)-SpeqNet performs best by a large margin.

A2 See Table 4. Over all three datasets, the (k, s)-SpeqNet architectures improve over the GCN baseline. Specifically, over all datasets, the (2, 1)-SpeqNet and the (2, 2)-SpeqNet lead to an increase of at least 7% in accuracy. For example, both architectures beat the GCN baseline by at least 17% on the Wisconsin dataset. Further, the (k, s)-SpeqNet architectures lead to better accuracies compared to the SDRF architecture, e.g., improving on it by more than 10% on the Cornell dataset. Hence, node-level tasks also benefit from higher-order information. However, increasing k more does not result in increased accuracies.
A3 Kernels See Table 6. Clearly, for the same $k$ and $s < k$, the $(k, s)$-WL improves over the $k$-WL and its (local) variants. For example, on the ENZYMES dataset, the $(2, 1)$-WL is more than 20 times faster in terms of computation times compared to the $\delta$-2-WL. The speed-up is even more significant for the (non-local) 2-WL. This speed-up factor increases more as $k$ increases, e.g., the $(3, 1)$-WL is more than 1700 times faster compared to the 3-WL, whereas the $(3, 2)$-WL is still more than 87 times faster, while giving better accuracies. Similar speed-up factors can be observed over all datasets.

Neural architectures See Table 7. The $(2, 1)$-SpeqNet severely speeds up the computation time across both datasets. Specifically, on the ALCHEMY dataset, the $(2, 1)$-SpeqNet is 1.3 times faster compared to the $(2, 2)$-SpeqNet, while requiring twice the computation time of the GINE-$\varepsilon$ but achieving a lower MAE. More interestingly, on the QM9 dataset, the $(2, 1)$-SpeqNet is 3.4 times faster compared to the $(2, 2)$-SpeqNet, while also being 1.3 times faster compared to the GINE-$\varepsilon$. The speed-up over GINE-$\varepsilon$ is most likely due to the latter considering the complete graph to compute all pairwise $\ell_2$ distances, whereas the $(2, 1)$-SpeqNet only considers connected node pairs.

A4 See Tables 3 and 4. The $(2, 1)$-LWL, $(2, 1)$-LWL$^+$, and $(3, 1)$-LWL$^+$ beat the 1-WL on six out of eight datasets. Going from the $(3, 1)$-LWL to the $(3, 2)$-LWL often leads to a slight increase in accuracy, e.g., on the ENZYMES and MUTAG datasets, while sometimes leading to a drop in accuracy, e.g., on the IMDB-BINARY dataset. Hence, a better understanding of the model’s generalization performance with respect to $s$ needs to be investigated in future work. The effect is less pronounced for the neural architectures; however, all higher-order models beat the GNN baseline. Reducing $s$ leads to a vast reduction in computation time. For example, on the ENZYMES dataset, going from the $(3, 2)$-LWL to the $(3, 1)$-LWL leads to a speed-up factor of more than 18, while only inducing a small drop in terms of accuracy, whereas the $(3, 1)$-LWL$^+$ beats the $(3, 2)$-LWL while only increasing the computation time by one second. Similar observations can be made across all datasets.

Increasing $s$ often leads to a better performance on the graph regression tasks. For example, on the ALCHEMY dataset, going from a $(2, 1)$-SpeqNet to a $(2, 2)$-SpeqNet architecture reduces the MAE by 0.054. Similar effects can be observed for the QM9 dataset, and when going from a $(3, 1)$-SpeqNet to a $(3, 2)$-SpeqNet architecture. On the node-classification datasets, reducing $s$ leads to a slight drop in accuracy, between 0.5 and 4%, while increasing $k$ beyond 2 often results in a drop in accuracy.

6 Conclusion

To circumvent the exponential running time requirements of the $k$-WL, we introduced a new heuristic for the graph isomorphism problem, namely the $(k, s)$-WL. By varying the parameters $k$ and $s$, the $(k, s)$-WL offers a trade-off between scalability and expressivity and, unlike the $k$-WL, takes into account the potential sparsity of the graph. Based on these combinatorial insights, we designed provably expressive machine-learning architectures, $(k, s)$-SpeqNets, suitable for node-, subgraph-, and graph-level prediction tasks. Empirically, we showed that such architectures lead to state-of-the-art results in node- and graph-level classification regimes while obtaining promising results on graph-level regression tasks. We believe that this principled approach paves the way for designing new permutation-equivariant architectures, overcoming the limitations of current graph neural networks.
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