Maximally Expressive GNNs for Outerplanar Graphs

Franka Bause∗
Faculty of Computer Science
UniVie Doctoral School Computer Science
University of Vienna, Vienna, Austria
franka.bause@univie.ac.at

Fabian Jogl∗
Center for Artificial Intelligence and Machine Learning
Machine Learning Research Unit
TU Wien, Vienna, Austria
fabian.jogl@tuwien.ac.at

Pascal Welke
Machine Learning Research Unit
TU Wien, Vienna, Austria
pascal.welke@tuwien.ac.at

Maximilian Thiessen
Machine Learning Research Unit
TU Wien, Vienna, Austria
maximilian.thiessen@tuwien.ac.at

Abstract
Most pharmaceutical molecules can be represented as outerplanar graphs. We propose a graph transformation that makes the Weisfeiler-Leman (WL) test and message passing graph neural networks maximally expressive on outerplanar graphs. While existing research predominantly focuses on enhancing expressivity of graph neural networks beyond the WL test on arbitrary graphs, our goal is to distinguish pharmaceutical graphs specifically. Our approach applies a linear time transformation, building on the fact that biconnected outerplanar graphs can be uniquely identified by their Hamiltonian adjacency list sequences. This pre-processing step can then be followed by any graph neural network. We achieve promising results on molecular benchmark datasets while keeping the pre-processing time low, in the order of seconds for common benchmarks.

1 Introduction
We study graph neural networks (GNNs) for the family of outerplanar graphs and devise a model that can distinguish all non-isomorphic outerplanar graphs. Most previous work relates the expressivity of GNNs to some \( k \)-dimensional Weisfeiler-Leman isomorphism test (\( k \)-WL), most famously showing that some GNNs can distinguish exactly all those graphs that \( k \)-WL can distinguish [Maron et al., 2019b; Morris et al., 2020; Xu et al., 2019]. In contrast, here we present an expressivity result for a well established class of graphs not defined by (generalized) message passing. We have chosen outerplanar graphs, since most pharmaceutical compounds are outerplanar [Droschinsky et al., 2017; Horváth and Ramon, 2010; Horváth et al., 2006]. In fact, most well-known benchmark datasets for graph-level tasks contain over 92% outerplanar graphs (see Table 2 in the appendix).

Relying on an isomorphism test for biconnected outerplanar graphs by Colbourn and Booth [1981], we develop a linear-time graph transformation [Jogl et al., 2023; Veličković, 2022]. This enables 1-WL, and thus message passing neural networks (MPNNs), to distinguish such graphs. We rely on Hamiltonian adjacency lists (HALs), which encode the structure of each biconnected component, and the fact that 1-WL can distinguish labeled trees. We discuss related work in Appendix D.

2 Preliminaries
A graph \( G = (V, E, \mu, \nu) \) consists of a set of nodes \( V \), a set of edges \( E \subseteq V \times V \) and \( \mu: V \to X \) and \( \nu: E \to X \) arbitrary attributes for the nodes and edges, respectively. We refer to an edge from \( u \) to \( v \) by \( uv \), and in case of undirected graphs \( uv = vu \). The in-neighbors of a node \( u \in V \) are denoted by \( N_i(u) = \{ v \mid vu \in E \} \). The out-neighbors of a node \( u \in V \) are denoted by \( N_o(u) = \{ v \mid wv \in E \} \)

*Equal contribution. For an extended version, see [Bause et al., 2023].

and in case of undirected graphs, \( N_i = N_o \). We focus on undirected input graphs, and will transform them into directed ones. A (directed) cycle \((v_1, \ldots, v_k)\) is a sequence of \( k \) distinct nodes, with \( \forall i \in \{1, \ldots, k-1\} : v_i v_{i+1} \in E \) and \( v_n v_1 \in E \). Given a graph \( G \), we denote the shortest path distance between \( v_i \) and \( v_j \) by \( d_G(v_i, v_j) \). A graph is outerplanar if it can be drawn in the plane without edge crossings and with all nodes belonging to the exterior face (for more details, see, e.g., Felsner [2012]). We call an undirected graph with at least three vertices biconnected if the removal of any single node does not disconnect the graph. A biconnected component is a maximal biconnected subgraph and we call biconnected outerplanar components of a graph blocks.

Identifying biconnected outerplanar graphs using Weisfeiler-Leman. We first present a graph transformation called \( \text{CAT} \), that allows the Weisfeiler-Leman algorithm to distinguish any two biconnected outerplanar graphs, and then extend it in \( \text{CAT} \) to all outerplanar graphs.

**Definition 1.** The \( \text{CAT}^* \) transformation takes a biconnected outerplanar graph \( G = (V, E, \mu, \nu) \) and yields a modified graph \( G' = \text{CAT}^*(G) = (V', E', \mu', \nu') \) by performing the steps below.

1. Let \( C = (v_1, \ldots, v_n) \) be a (directed) Hamiltonian cycle of \( G \) and \( \overline{C} \) be its reverse.
Two biconnected outerplanar graphs \( G \) and \( H \) are isomorphic, if and only if \( \text{WL}(\text{CAT}^*(G)) = \text{WL}(\text{CAT}^*(H)) \).

**Proof.** Two graphs are distinguished by WL iff the multisets of node colors of their stable colorings differ. Trivially, \( |V(G)| \neq |V(H)| \Rightarrow |V(\text{CAT}^*(G))| \neq |V(\text{CAT}^*(H))| \Rightarrow \text{WL}(\text{CAT}^*(G)) \neq \text{WL}(\text{CAT}^*(H)) \), so we only focus on graphs with \( |V(G)| = |V(H)| \). Two nodes only get the same color, if their unfolding trees are isomorphic. The first number in the HAL of each node is always 1, so it can be ignored, and the last number is always \( |V(G)| - 1 \), so this can simply be reconstructed by \( |V(\text{CAT}^*(G))| \). The rest of the HAL sequence and the node labels of \( G \) can be reconstructed from the unfolding tree of any node in \( \text{CAT}^*(G) \): Of course, each node has two direct neighbors in the Hamiltonian cycle. In the unfolding tree these are the parent and the single child with the 1-annotated edge. All other neighbors in the HAL can be reconstructed by looking at the weights of the edges that do not have weight 1. Figure 4 shows an example. Looking at any two biconnected outerplanar graphs with \( n \) nodes, Weisfeiler-Leman will be able to distinguish them after at most \( n \) iterations, if they are non-isomorphic: Since the HAL sequence is encoded in the unfolding trees from all starting points (cyclic shift) and, because of the reverse copy, in both directions (reverse direction), this identifies isomorphism by Lemma 1.

The **CAT transformation.** We define the CAT transformation by applying \( \text{CAT}^* \) to the blocks of the graphs and adding nodes and edges to make outerplanar graphs distinguishable by WL.

**Definition 2.** The \( \text{CAT}(G) = G' \) transformation, maps a graph \( G \) to a new graph \( G' \) as follows:

1. Let \( B_1, \ldots, B_t \) be the blocks of \( G \) and let \( F \) be the graph induced by the edges of \( G \) that are not in any block plus the nodes that are present in more than one block. Let \( \{\square, \odot, \otimes, \ast, \triangle\} \) be distinct node labels not in \( X \).

2. Add \( F \) to \( G' \) with labels \( \mu'(v) = (\perp, \mu(v)) \) for \( v \in F \).

3. For each block \( B_i \) in \( G' \):
   
   3.1. Add \( (B_i', B_i) = \text{CAT}^*(B_i) \) to \( G' \) (with labels).
   
   3.2. Let \( A_i = V(B_i) \cap V(F) \) be the nodes of \( B_i \) in \( F \).
   
   3.3. Let \( \gamma_i : A_i \to V(B_i') \) map nodes of \( F \) to their copy in \( B_i' \) and let \( \gamma_i \) to the copy in \( B_i' \).
   
   3.4. Add a node \( b_i \) and (undirected) edges \( \{b_i, v\} \) for all \( v \in V(B_i) \cup V(B_i') \) to \( G' \).
   
   3.5. For each \( a \in A_i \) add a node \( p \) and edges \( \{p, b_i\}, \{p, a\}, \{p, \gamma_i(a)\}, \{p, \gamma_i(a)\} \) to \( G' \).
   
   3.6. Let \( \mu'(b_i) = \square \), and for each \( a \in A_i \) let \( \mu'(a) = (\perp, \mu(a)) \) and \( \mu'(p) = (\perp, \mu(a)) \) for the corresponding \( p \).
4. Add a node \( g \) with \( \mu'(g) = \triangle \) to \( G' \) and for all nodes \( b_i \), add an edge \( \{g, b_i\} \) to \( G' \).

5. Let \( \text{CAT}(G) = G' \).

An example for a graph \( G \) and its corresponding graph \( G' = \text{CAT}(G) \) can be seen in Figure 1.

**Theorem 2.** Two outerplanar graphs \( G \) and \( H \) are isomorphic, iff \( \text{WL}(\text{CAT}(G)) = \text{WL}(\text{CAT}(H)) \).

**Proof.** Following Theorem 1, each block will be uniquely identified by WL. Since the additional nodes have distinct labels, they will not cause WL to falsely report two blocks as isomorphic when they are not. The information about the entire HAL sequence of each block is stored in the \( b \) nodes after some iteration. The \( p \) nodes connect the block and \( b \) nodes to the rest of the graph, determining the orientation of the block. Note that the graph returned by CAT without the CAT\(^*\) blocks and the node \( g \) is a tree. Relying on the labels of the \( p \) and \( b \) nodes, we can reconstruct the original graph from this tree. As WL can distinguish labeled trees [Arvind et al., 2015; Kiefer, 2020], it can thus distinguish non-isomorphic outerplanar graphs using CAT. For the other direction, note that CAT is permutation-invariant: for two isomorphic graphs \( G \) and \( H \), the graphs \( \text{CAT}(G) \) and \( \text{CAT}(H) \) are isomorphic and WL will give the same output for both.

Importantly, we can compute \( \text{CAT}(G) \) in linear time. The computational complexity is dominated by the computation of the blocks [Tarjan, 1972] and their Hamiltonian cycles [Mitchell, 1979], which both require linear time. Note that we only add a linear number of nodes and edges. From Xu et al. [2019] it follows, that MPNNs that are as expressive as 1-WL can distinguish \( \text{CAT}(G) \) and \( \text{CAT}(H) \) for non-isomorphic outerplanar graphs \( G \) and \( H \). Thus, we propose to transform the input graphs using CAT and then use any MPNN on them. We can also apply CAT to non-outrplanar graphs. In this case, the biconnected outerplanar components are identified and the steps described in Definition 2 are performed. The remaining graph will simply be copied without modification. This never reduces expressivity but also is not guaranteed to distinguish general non-outrplanar graphs.

### 4 Experimental Evaluation

We investigate whether our proposed CAT can improve the predictive performance of the MPNN GIN [Xu et al., 2019] on two commonly used molecular benchmarks: ZINC [Gómez-Bombarelli et al., 2018; Sterling and Irwin, 2015] and ogbg-molhiv [Hu et al., 2020]. For this, we compare CAT+GIN against GIN on both datasets. We tune the hyperparameters on the validation sets and evaluate the best performing hyperparameters 10 times. For each dataset we track a commonly used evaluation metric for this dataset and report the mean and standard deviation of this metric in the epoch with the highest validation performance. More details can be found in Appendix C and in our code repository\(^2\). Table 1 shows the results of our experiments and we can see that CAT+GIN convincingly outperforms GIN on both datasets. To measure the speed of CAT we measure its runtime on the training splits of ZINC and MOLHIV averaged over 10 trials. CAT requires 38 ± 1 s for ZINC and 133 ± 1 s for MOLHIV.

**Table 1:** Test performance of different GNNs over 10 random seeds. Arrows indicate whether smaller (↓) or bigger (↑) results are better. **Bold** indicates the best performing model for a dataset.

<table>
<thead>
<tr>
<th>Model</th>
<th>ZINC MAE ↓</th>
<th>MOLHIV ROC-AUC ↑</th>
</tr>
</thead>
<tbody>
<tr>
<td>GIN</td>
<td>0.177 ± 0.006</td>
<td>76.5 ± 1.2</td>
</tr>
<tr>
<td>CAT+GIN</td>
<td><strong>0.125 ± 0.006</strong></td>
<td><strong>78.4 ± 0.6</strong></td>
</tr>
</tbody>
</table>

### 5 Conclusion

We proposed a graph transformation, which enables the Weisfeiler-Leman test to be maximally expressive on outerplanar graphs by building on the fact that biconnected outerplanar graphs can be uniquely identified by their Hamiltonian adjacency list sequences. Our transformation encodes these HAL sequences in unfolding trees. We achieve promising first empirical results on molecular benchmark datasets, while keeping the pre-processing time very low. Interesting further directions would be extending the transformation to non-outerplanar graphs and investigating the effect of the transformation on the issues of oversquashing and oversmoothing.

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\(^2\)https://github.com/ocatias/OuterplanarGNNs_LoG
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### A Outerplanarity of Graph Datasets

Table 2 shows the percentage of outerplanar graphs in popular molecular datasets.

**Table 2: Common benchmark datasets and the percentage of outerplanar graphs in them: NCI (https://cactus.nci.nih.gov/) ZINC [Gómez-Bombarelli et al., 2018; Sterling and Irwin, 2015] and all other datasets are from ogb [Hu et al., 2020].**

<table>
<thead>
<tr>
<th>Dataset</th>
<th>#Graphs</th>
<th>Outerplanar</th>
</tr>
</thead>
<tbody>
<tr>
<td>NCI</td>
<td>250251</td>
<td>94%</td>
</tr>
<tr>
<td>ZINC</td>
<td>12000</td>
<td>98%</td>
</tr>
<tr>
<td>molhiv</td>
<td>41127</td>
<td>92%</td>
</tr>
<tr>
<td>moltox21</td>
<td>7831</td>
<td>96%</td>
</tr>
<tr>
<td>molesol</td>
<td>1128</td>
<td>97%</td>
</tr>
<tr>
<td>molbace</td>
<td>1513</td>
<td>93%</td>
</tr>
<tr>
<td>molclintox</td>
<td>1477</td>
<td>94%</td>
</tr>
<tr>
<td>molbbbp</td>
<td>2039</td>
<td>92%</td>
</tr>
<tr>
<td>molsider</td>
<td>1427</td>
<td>92%</td>
</tr>
<tr>
<td>moltoxcast</td>
<td>8576</td>
<td>96%</td>
</tr>
<tr>
<td>mollipo</td>
<td>4200</td>
<td>96%</td>
</tr>
</tbody>
</table>

### B Example for Hamiltonian Adjacency Lists

Figure 3 shows two graphs and their directed Hamiltonian cycles. The nodes are annotated with their HALs, the list of distances on the Hamiltonian cycle to their neighbors.

### C Experimental Evaluation

Our models are implemented in PyTorch-Geometric [Fey and Lenssen, 2019] and trained on a single NVIDIA GeForce RTX 3080 GPU. We use WandB [Biewald, 2020] for tracking. The used server has 64 GB of RAM, has an 11th Gen Intel(R) Core(TM) i9-11900KF CPU running at 3.50GHz and uses Fedora 38. For ZINC we train with a batch size of 128 and an initial learning rate of $10^{-3}$ that is halved whenever the validation metric does not improve for 20 epochs. Training stops after 500 epochs or after the learning rate dips below $10^{-5}$. For MOLHIV we train with a batch size of 128 and a fixed learning rate of $10^{-3}$ for 100 epochs. Table 3 shows the hyperparameters for GIN and CAT+GIN. We used the same hyperparameters for both models. We used a smaller hyperparameter grid for MOLHIV than for ZINC, as MOLHIV is larger than ZINC meaning that training takes much longer.

**More details on CAT.** CAT adds an additional feature to each node which encodes the type of that node i.e., nodes from Hamiltonian cycles, block nodes, pooling nodes, articulation nodes and or global block nodes. Furthermore, we create additional edge features encoding the types of nodes incident to this edge i.e., an edge between two different nodes in a Hamiltonian cycle has a different type than an edge from a pooling node to the block node. For newly created nodes and edges we set their remaining features to the feature of the node / edge they are based on; for example, a pooling node will have the features of the node they are performing the pooling operation for.

![Figure 3: Two graphs and their directed Hamiltonian cycles. Nodes are annotated with their HALs, the distances on the Hamiltonian cycle to their neighbors [Colbourn and Booth, 1981].](image)
have no natural representation in the graph (block and block pooling nodes) we set these features
to 0. To ensure that only these nodes get assigned 0 features, we shift the values of these features
for all other nodes by 1. Note that our GNN treats the distance on edges in blocks as a categorical
feature. We are not sure whether this is advantageous and will experiment with treating these features
as ordinals in the future.

Table 3: Hyperparamter grids for different datasets.

<table>
<thead>
<tr>
<th>Parameter</th>
<th>GIN, CAT+GIN On ZINC</th>
<th>GIN, CAT+GIN On MOLHIV</th>
</tr>
</thead>
<tbody>
<tr>
<td>Message passing layers</td>
<td>2, 3, 4, 5</td>
<td>4, 5</td>
</tr>
<tr>
<td>Final MLP layers</td>
<td>2</td>
<td>2</td>
</tr>
<tr>
<td>Pooling operation</td>
<td>mean, sum</td>
<td>mean, sum</td>
</tr>
<tr>
<td>Embedding dimension</td>
<td>64, 128, 256</td>
<td>64, 128</td>
</tr>
<tr>
<td>Jumping knowledge</td>
<td>last</td>
<td>concat</td>
</tr>
<tr>
<td>Dropout rate</td>
<td>0, 0.5</td>
<td>0.5</td>
</tr>
</tbody>
</table>

D Discussion and Related Work

It is well known that the expressiveness of MPNNs is bounded by the 1-WL test [Morris et al.,
2019; Xu et al., 2019]. The unlabelled graphs corresponding to decalin and bicyclooctyl cannot be
distinguished by any MPNN. As these two graphs are outerplanar this shows that MPNNs are not
expressive enough for outerplanar graphs. Even some biconnected outerplanar graphs cannot be
distinguished by MPNNs, see Fig. 3. The importance of biconnectivity in the context of GNNs
was recently discussed by Zhang et al. [2023]. The fact that many pharmaceutical molecules are
outerplanar is well known in the graph mining community [Horváth and Ramon, 2010; Horváth et al.,
2006]. Cotta et al. [2021] discussed outerplanar graphs in the context of reconstruction with GNNs.

It is known that 3-WL is sufficient and necessary to distinguish all outerplanar graphs [Kiefer,
2020]. Hence, any GNN matching the expressivity of 3-WL, such as 3-IGN [Maron et al., 2019a] or
3-GNN [Morris et al., 2019], is capable of solving our main goal of distinguishing all outerplanar
graphs. The 3-WL test, however, runs in \( O(n^3 \log n) \) time [Immerman and Lander, 1990; Kiefer,
2020], which is practically inefficient already for medium-sized graphs. Similarly, 3-GNN and 3-IGN
run in roughly \( O(n^3) \) time, see Maron et al. [2019a]; Morris et al. [2019]. Contrary, we perform a
linear-time pre-processing of the graph and run standard 1-WL, or an MPNN, with a much more
efficient runtime of \( O(n \log n) \). This is the same asymptotic runtime as running 1-WL on the original
graphs, as we only add a linear number of nodes and edges in our graph transformation.

Finally, there are other approaches towards building more expressive GNNs such as methods that
extract subgraphs [Bevilacqua et al., 2021; Maron et al., 2019a] or lift the graph to regular cell
complexes Bodnar et al. [2021]. The idea of using additional node labels to increase expressivity is
also well known [Abboud et al., 2021; Dasoulas et al., 2020].

E Additional Figures

Fig 5 demonstrates the difference between two blocks overlapping in the same articulation node (top)
or in different articulation nodes (bottom). Fig 6 demonstrates CAT on real-life molecular graphs. We
would like to point out that one of the graphs looks like a frog.
Figure 4: One part of the CAT$^*$ transformation of the graph from Figure 2 and an example unfolding tree of one of its nodes from which the HAL sequence of the original graph can be reconstructed.

Figure 5: Left: example graphs; Right: Result of applying CAT to these graphs. Colors indicate the type of node: red nodes are from Hamiltonian cycles, blue nodes correspond to blocks, yellow nodes pool nodes from the Hamiltonian cycles, orange nodes correspond to articulation nodes and the grey node pools block nodes.
Figure 6: Left: example graphs from MOLHIV (top) and ZINC (bottom); Right: Result of applying CAT to these graphs. Colors indicate the type of node: red nodes are from Hamiltonian cycles, blue nodes correspond to blocks, yellow nodes pool nodes from the Hamiltonian cycles, orange nodes correspond to articulation nodes and the grey node pools block nodes.