Message Passing Networks for Molecules with Tetrahedral Chirality

Abstract

Molecules with identical graph connectivity can exhibit different physical and biological properties if they exhibit stereochemistry—a spatial structural characteristic. However, modern neural architectures designed for learning structure-property relationships from molecular structures treat molecules as graph-structured data and therefore are invariant to stereochemistry. Here, we develop two custom aggregation functions for message passing neural networks to learn properties of molecules with tetrahedral chirality, one common form of stereochemistry. We evaluate performance on synthetic data as well as a newly-proposed protein-ligand docking dataset with relevance to drug discovery. Results show modest improvements over a baseline sum aggregator, highlighting opportunities for further architecture development.

1 Introduction

Recent advances in machine learning for chemistry that enable learning directly from molecular structures without preprocessing them into fixed-length vectors have yielded improvements in diverse settings including property prediction [19, 26, 7], predictive synthesis [8], and molecular optimization [12, 4]. However, stereochemistry has remained largely unexplored through the lens of deep learning despite being a significant aspect of molecular representation. Stereoisomers are molecular structures with the same graph connectivity but different spatial arrangements. Many different types of stereochemistry exist, ranging from common forms such as tetrahedral chirality and cis/trans isomerism to rarer forms such as atropisomerism and (the recently discovered) akamptisomerism [5]. Each of them influences the accessible conformations of a small molecule, which in turn can influence the molecule’s properties. Stereoisomers can have different physical properties (e.g., melting and boiling...
point), pharmacokinetics (e.g., absorption, distribution, metabolism, and excretion), and bioactivity (e.g., protein affinity) [3, 18, 27, 6].

This work focuses on tetrahedral chirality and its influence on property prediction. In principle, a molecule exhibiting tetrahedral chirality is neither a 2D graph nor a 3D structure, but somewhere in between. That is, the stereochemical designation of a chiral center limits the range of accessible conformers, but does not lock in a single 3D structure. Thus, we view using 3D representations to address tetrahedral chirality as too restrictive for conformationally-flexible molecules.

Message passing neural networks (MPNNs) operate on molecules as graphs by treating atoms as nodes and bonds as edges [11]. MPNNs operate by iteratively aggregating neighbor representations; traditional aggregation functions developed for graph-structured data such as sum, mean, and max are symmetric operators. Because stereoisomers have identical graph connectivities, symmetric aggregators operating on two different chiral centers will collapse their neighbors to an identical representation, regardless of chirality; that is, aggregation functions are the primary barrier to effectively realizing chirality in MPNN architectures. Equation 1 shows a generic MPNN update with $x^{(k)}_i \in \mathbb{R}^F$ denoting the representation of node $i$ at layer $k$ and $x_j, j \in \mathcal{N}(i)$ denoting its neighbors. $\gamma$ and $\phi$ represent differentiable functions while $e_{j,i} \in \mathbb{R}^{F_e}$ is the directed edge representation from node $j$ to $i$. $\text{AGG}$ refers to a generic aggregation scheme.

$$x^{(k)}_i = \gamma(k)\left(x^{(k-1)}_i, x^{(k)}_j\right), \quad x^{(k)}_j = \text{AGG}_{j \in \mathcal{N}(i)} \phi(k)\left(x^{(k-1)}_i, x^{(k-1)}_j, e^{(k-1)}_{j,i}\right)$$

Here, we focus on extending MPNNs to remove the unwanted invariance to tetrahedral chirality. We develop and evaluate our methods on a new dataset that exemplifies the influence of tetrahedral chirality on molecular properties. Our primary contributions are: (1) Establishing a customized graph representation where edges of each node are ordered, not merely sets; (2) Proposing two theory-motivated methods to distinguish molecules with different tetrahedral chiralities through asymmetric node aggregation while still achieving physically-meaningful invariances; (3) Proposing a benchmark dataset where the property of interest is a function of stereochemical configuration; and (4) Evaluating empirical performance on this and other datasets using the proposed aggregation schemes through multiple criteria. We use the PyTorch Geometric framework to implement our network [13] and make our code publicly available at https://github.com/PattanaikL/chiral_gnn

2 Related Work

Message passing on 2D molecular graphs MPNNs have gained relevance for their application to molecular property prediction [11, 20, 15] and are now a staple of many QSAR approaches [16]. Recent innovations include directional messages [25] and hierarchical messages [25]. Many of these 2D graph methods do not address stereochemistry at all; in fact, the recent work by Flam-Shepherd et al. [14], which includes 3D information, is one of the first MPNN architectures to explicitly address stereochemistry by performing message passing and aggregation over larger subgraphs.

Learning molecular invariances Achieving (in,equi)variances for molecules with neural networks has traditionally focused on rotational and translational symmetries, primarily for computed properties like electronic energy, which are identical between enantiomers. Architectures leading this effort include DTNNs [31], SchNet [30], Comorant [2], and the recent DimeNet [22]. However, all of these networks operate in $\mathbb{R}^3$, which does not reflect the conformational flexibility of molecules.

3 Approach

Preliminaries Ostensibly, MPNNs might capture stereochemistry through atom- or bond-level descriptors, such as an R/S atom feature to distinguish enantiomers (i.e., non-superimposable stereoisomers that are mirror images of each other). These features represent global measures of chirality, determined through CIP side-chain ranking conventions [9]. Figure 1 illustrates why global chirality descriptors as atom features are insufficient for MPNNs to meaningfully differentiate these structures.
With global descriptors, the two "R" structures would receive the same atom-level feature while the two "S" structures would receive a different feature. However, the local configurations at the tetrahedral center are different for each of the two “R” structures, despite sharing the same global descriptor.

An alternate method captures the relevant information through local chirality descriptors, denoted by clockwise/counter-clockwise labels (CW/CCW) in Figure 1, as used in SMILES notation [12]. Chiral centers receive a parity label based on the order of neighboring atoms, e.g. as provided in the SMILES string [10], relative to their spatial orientation, which assigns them to a local chiral group of 12 equivalent permutations of four neighbors out of 24 total (Figure 2). Directly using this parity bit as an atom feature would make the MPNN sensitive to arbitrary bookkeeping and SMILES canonicalization conventions. We therefore use this local parity bit in combination with the ordered set of neighbors to preserve physically-meaningful invariances.

Theory We now prove the model structure necessary to constitute a universal approximator of all aggregation functions defined over the same chiral group. For any set $X$, we define the quadruples set $X^4 = \{(x_1, x_2, x_3, x_4), x_i \in X, i = 1, 2, 3, 4\}$. We define the equivalence relation $\sim_C$ for a given chiral group $C$ as $(x_1, x_2, x_3, x_4) \sim_C (x'_1, x'_2, x'_3, x'_4)$ iff $\exists \pi \in \mathcal{P}(C)$ such that $(x_1, x_2, x_3, x_4) = (x'_{\pi(1)}, x'_{\pi(2)}, x'_{\pi(3)}, x'_{\pi(4)})$, where $\mathcal{P}(C)$ is the set of all 12 permutations of quadruples that are in the $C$ chiral group. Finally, we define the quotient set $\mathcal{C}_4(X) = X^4 \sim_C$. Then, the following proposition proves the universal approximation capability of our proposed architecture. See the Appendix for a proof of the proposition.

Proposition 1. Any function $f : \mathcal{C}_4(X) \to \mathbb{R}^n$ is a valid function on quadruples from a chiral group $C$, i.e. it is invariant to any permutation from $\mathcal{P}(C)$, iff it can be decomposed in the form

$$f(x_1, x_2, x_3, x_4) = h \left( \sum_{\pi \in \mathcal{P}(C)} g(x_{\pi(1)}, x_{\pi(2)}, x_{\pi(3)}, x_{\pi(4)}) \right)$$

for suitable functions $g$ and $h$.

Models Following Equation 2, we propose two novel aggregation functions for updating the representation of a tetrahedral center $x_i$ by adjusting the function $g$. Each aggregation function allows enantiomers to be distinguished while remaining invariant to arbitrary bookkeeping conventions.

Permutation (PERM). PERM introduces four separate weight matrices $\{W_{g_i}\}_{i=1}^4$ for each ordered neighbor in a given permutation (denoted by $\pi \in \mathcal{P}(i)$). Note that in Equation 3, $\tau_{g_1}$ and $\tau_{g_2}$ represent different nonlinear operators (tanh and ReLU, respectively, in our case).

$$x_i^{*}\left(k\right) = \frac{1}{3} W_{h_2} \tau_h \left( \frac{1}{3} W_{h_1} \sum_{\pi \in \mathcal{P}(i)} \tau_{g_2} \left( \tau_{g_1} \left( W_{g_1} x^{(k-1)}_{\pi(1)} \right) + \tau_{g_1} \left( W_{g_2} x^{(k-1)}_{\pi(2)} \right) + \tau_{g_1} \left( W_{g_3} x^{(k-1)}_{\pi(3)} \right) + \tau_{g_1} \left( W_{g_4} x^{(k-1)}_{\pi(4)} \right) \right) \right)$$
Permutation-concatenation (PERM_CAT). PERM_CAT is similar to the PERM architecture, but rather than using four separate weight matrices, we use a single weight matrix and concatenate atom representations in each permutation. PERM_CAT and PERM are equivalent when $\tau_g$ is linear.

$$x_i^{* (k)} = W_{h_2} \tau_h \left( \frac{1}{3} W_{h_1} \sum_{\pi \in P(i)} \tau_g \left( W_g(x_{\pi (1)}^{(k-1)}) \| x_{\pi (2)}^{(k-1)} \| x_{\pi (3)}^{(k-1)} \| x_{\pi (4)}^{(k-1)} \right) \right)$$

We test each aggregation function within several message passing frameworks to assess empirical performance. These include the standard graph convolutional network (GCN) [21] modified to include edge features, the graph isomorphism network (GIN) [34] modified to use edge features [17], and the directed message passing neural network (DMPNN) [35]. We also test the effect of including only atom-level local chiral features describing CW/CCW, only global chiral features describing R/S, and including both sets of features; we denote these runs by “LCF,” “GCF,” and “BCF,” respectively. A third aggregation function that achieves the correct invariances but is not motivated by Proposition 1, the Product of Differences, can be found in the Appendix.

4 Experiments

Datasets Most property prediction datasets lack clean stereochemical information on which to benchmark chirality-aware models. To this end, we extract a subset of data from the D4 dopamine receptor protein-ligand docking screen by Lyu et al. [24]. Stereoisomers may exhibit distinct interaction energies when complexing with a protein of interest due to the conformations/poses they can achieve. Our dataset (D4DCHP) narrows the original 138 million molecules to stereoisomer pairs for a single 1,3-dicyclohexylpropane skeletal scaffold. We further define two additional subsets: one where enantiomers exhibit $> 5$ kcal/mol differences in docking score (DIFF5) and one where molecules have a single tetrahedral center (CHIRAL1). We also evaluate performance on the “Lipo” dataset of log-scaled partition coefficients from Wu et al. [33] as 1127 of 4200 molecules in this dataset contain at least one tetrahedral center; unlike the D4DCHP dataset, however, we do not have complete pairs of enantiomers. See Appendix Section 6.5.3 for Lipo results. Additional dataset details can also be found in the Appendix.

Metrics We evaluate empirical performance of our custom aggregators on each dataset using root mean squared error (RMSE). However, as we are especially interested in how our methods separate stereoisomers in the D4DCHP dataset, we additionally evaluate the model’s ability to correctly order two enantiomers; predictions within 0.1 kcal/mol of each other are considered indistinguishable and treated as a random guess with 50% accuracy. We denote this task rank classification.

R/S Classification To ensure that our new aggregation methods understand tetrahedral chirality, we first evaluate them on the simple task of classifying molecules from CHIRAL1 as R or S. This tests if the MPNNs can (1) distinguish enantiomers and (2) learn CIP rules to assign an R/S label. Because the input atom/bonds features and graph connectivity are identical for a pair of stereoisomers, we expect a sum aggregator to achieve an accuracy of 50%. Empirical results (Table 1) support this hypothesis. Sum aggregators across MPNN architectures do not perform better than a random classifier, while both custom aggregators for all three architectures reach nearly perfect accuracy when classifying molecules as R or S.

| Aggregation method | Graph architecture |
|--------------------|--------------------|
| SUM                | GCN 50 50 50       |
| PERM               | GCN 100 100 98     |
| PERM_CAT           | GCN 100 100 98     |

4 D4DCHP We next evaluate empirical performance on the full D4DCHP dataset (Table 2). We shuffle the data randomly but ensure that stereoisomer pairs are kept together in the train/validation/test splits. In the absence of atom-level chiral features, the asymmetric aggregation methods of PERM and PERM_CAT provide a significant benefit over the SUM aggregator baseline.

The performance of the different aggregation methods in the presence of chiral features depends on the architecture. The custom aggregation functions show measurable improvements in both rank
classification and RMSE for the GCN architecture, but these improvements are not so clear for the GIN and DMPNN architectures. For the GIN, including both atom-level chiral features is sufficient to capture the trends picked up by the custom aggregators in the absence of such features. The custom aggregators yield improved performance for the DMPNN in terms of rank classification but not RMSE.

The strong influence of atom-level chiral features is rather surprising. By including just two additional atom-level features, the sum aggregators achieve approximately a 10% improvement in rank classification, regardless of graph architecture. The more expressive graph architectures can more effectively use this chiral information, such that the explicit equivariances offered by the custom aggregators no longer provide a significant benefit. Regardless, the highest observed rank classification accuracy of 62% provides clear room for improvement, motivating further research into representation learning methods that can better address tetrahedral chirality.

Table 2: Rank classification percent for test set (trained on full D4DCHP, results for DIFF5 subset) and full D4DCHP RMSE for test set

| Aggregation method | GCN rank classification accuracy (%) | D4DCHP error (RMSE) |
|--------------------|--------------------------------------|---------------------|
|                    | GCN | GIN | DMPNN | GCN | GIN | DMPNN |
| SUM                | 50.0 ± 0.0 | 50.0 ± 0.0 | 50.0 ± 0.0 | 6.77 ± 0.05 | 6.45 ± 0.07 | 6.45 ± 0.06 |
| SUM (LCF)          | 55.6 ± 0.1 | 57.9 ± 0.3 | 57.8 ± 0.3 | 6.76 ± 0.04 | 6.42 ± 0.06 | 6.41 ± 0.06 |
| SUM (GCF)          | 57.9 ± 0.5 | 61.5 ± 0.2 | 60.6 ± 0.0 | 6.73 ± 0.05 | 6.38 ± 0.06 | 6.39 ± 0.06 |
| SUM (BCF)          | 58.7 ± 0.7 | 62.1 ± 0.1 | 60.9 ± 0.4 | 6.95 ± 0.30 | 6.38 ± 0.06 | 6.37 ± 0.06 |
| PERM               | 58.3 ± 0.5 | 58.6 ± 0.3 | 59.7 ± 0.5 | 6.69 ± 0.06 | 6.44 ± 0.06 | 6.42 ± 0.05 |
| PERM (LCF)         | 60.4 ± 0.5 | 61.1 ± 0.1 | 61.5 ± 0.1 | 6.69 ± 0.07 | 6.41 ± 0.06 | 6.41 ± 0.05 |
| PERM (GCF)         | 59.8 ± 0.5 | 61.2 ± 0.4 | 61.5 ± 0.3 | 6.68 ± 0.06 | 6.40 ± 0.07 | 6.40 ± 0.05 |
| PERM (BCF)         | 60.7 ± 0.3 | 61.5 ± 0.2 | 61.3 ± 0.2 | 6.67 ± 0.06 | 6.39 ± 0.06 | 6.39 ± 0.06 |
| PERM_CAT           | 60.5 ± 0.2 | 60.5 ± 0.3 | 61.2 ± 0.3 | 6.65 ± 0.06 | 6.40 ± 0.05 | 6.40 ± 0.06 |
| PERM_CAT (LCF)     | 61.0 ± 0.1 | 62.0 ± 0.3 | 61.3 ± 0.3 | 6.62 ± 0.06 | 6.38 ± 0.06 | 6.40 ± 0.05 |
| PERM_CAT (GCF)     | 60.7 ± 0.2 | 61.8 ± 0.5 | 61.8 ± 0.2 | 6.64 ± 0.05 | 6.40 ± 0.07 | 6.38 ± 0.06 |
| PERM_CAT (BCF)     | 60.8 ± 0.1 | 62.2 ± 0.1 | 62.4 ± 0.2 | 6.65 ± 0.06 | 6.37 ± 0.07 | 6.39 ± 0.07 |

5 Conclusion and Outlook

We have developed two aggregation functions to learn properties for molecules with tetrahedral chirality, motivated by a general expression for a universal function approximator. The custom aggregators can fully distinguish enantiomers on a toy R/S classification problem and show comparable or modestly-improved performance over a baseline sum aggregator on a newly-proposed D4DCHP dataset depending on the MPNN architecture and inclusion of atom-level chiral features.

Empirical performance is a function of both (a) whether the network understand stereochemistry and (b) the extent to which stereochemistry matters for that dataset and property. Many existing datasets are inadequate; either stereochemistry is not relevant (e.g., QM9 energies [28]), or the data are not rich enough to convey the nuanced trends yielded by chirality (e.g., lipophilicity). Thus, until such datasets become available, we recommend that algorithm development occur in controlled environments like D4DCHP. Finally, we have only attempted to address tetrahedral chirality for MPNNs here; we leave stereoisomerism beyond tetrahedral chirality to future work.

Acknowledgments and Disclosure of Funding

We gratefully acknowledge the Machine Learning for Pharmaceutical Discovery and Synthesis consortium for funding this work. We also acknowledge the MIT SuperCloud and Lincoln Laboratory Supercomputing Center for providing HPC resources that have contributed to the research results reported within this paper [29].

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6 Appendix

6.1 Theory

6.1.1 Proof of proposition 1

Proof: First, we easily observe that the functional form in Eq. [3] is a valid function on \( C_4(\mathcal{X}) \). Conversely, we note that there is a bijection between \( C_4(\mathcal{X}) \) and the set of sets of C-chiral group permutations of a quadruple. This bijection is given by the following function:

\[
\phi(x_1, x_2, x_3, x_4) = \{(x_{\pi(1)}, x_{\pi(2)}, x_{\pi(3)}, x_{\pi(4)}) \mid \pi \in \mathcal{P}(C)\}
\]

Using [36], we know that all functions from sets \( \mathcal{Y} \subset \mathcal{X} \) to \( \mathbb{R}^\mathcal{N} \) have the form \( h(\sum_{y \in \mathcal{Y}} \psi(y)) \) for some functions \( h \) and \( \psi \). Taking \( \mathcal{Y} = \mathcal{X}^4 \) and \( g = \psi \circ \phi \), we obtain the desired result. \( \square \)

6.2 Models

6.2.1 Product of differences (PD)

Separate from the PERM and PERM_CAT aggregators, we develop an additional aggregator that achieves the desired equivariances between local chiral groups without using multiple permutations. This method takes pairwise differences between neighbor atom representations and then computes the signed geometric mean. Note that the inversion of the tetrahedral center (e.g., by exchanging two neighbors) will invert the sign of the resulting product.

\[
x_i^{(k)} = \tau_h \left( \frac{W_h \prod_{j \in \mathcal{N}(i)} \prod_{m \in \mathcal{N}(i), m < j} (x_j^{(k-1)} - x_m^{(k-1)})^{1/6}}{6} \right)
\]

\[
= \tau_h \left( W_h (x_1^{(k-1)} - x_2^{(k-1)})^{1/6} \times (x_1^{(k-1)} - x_3^{(k-1)})^{1/6} \times (x_1^{(k-1)} - x_4^{(k-1)})^{1/6} \times (x_2^{(k-1)} - x_3^{(k-1)})^{1/6} \times (x_2^{(k-1)} - x_4^{(k-1)})^{1/6} \times (x_3^{(k-1)} - x_4^{(k-1)})^{1/6} \right)
\]

6.2.2 Aggregation visualization

Figure [3] shows an example of the sum aggregator and all custom aggregators applied to a tetrahedral chiral center for a given initial ordering.

Figure 3: A. Two stereoisomers and their atom feature vectors representing different local chiral groups. B-E. SUM, PD, PERM, and PERM_CAT aggregation examples for first chiral group.
6.2.3 Chiral group permutations

Table 3 shows all permutations for both chiral groups. Any initial atom ordering falls into one of these two groups, which determines all permutations considered for the chiral message. Note that, in practice, we adjust all atom orderings to fall within a single group (Group 1 in this case), so we only need to apply a single set of MLP functions (corresponding to $g$ and $h$ in Equation 2). This adjustment only changes the bookkeeping used for tetrahedral centers, not the physical meaning. See Figure 2 for a visual explanation on how to obtain these permutation groups.

| Group 1                                      | Group 2                                      |
|----------------------------------------------|----------------------------------------------|
| {0, 1, 2, 3}, {0, 2, 3, 1}, {0, 3, 1, 2}   | {0, 1, 3, 2}, {0, 2, 1, 3}, {0, 3, 2, 1}   |
| {1, 0, 3, 2}, {1, 3, 2, 0}, {1, 2, 0, 3}   | {1, 0, 2, 3}, {1, 3, 0, 2}, {1, 2, 3, 0}   |
| {2, 0, 1, 3}, {2, 1, 3, 0}, {2, 3, 0, 1}   | {2, 0, 3, 1}, {2, 1, 0, 3}, {2, 3, 1, 0}   |
| {3, 0, 2, 1}, {3, 2, 1, 0}, {3, 1, 0, 2}   | {3, 0, 1, 2}, {3, 2, 0, 1}, {3, 1, 2, 0}   |

6.3 Datasets

6.3.1 D4DCHP dataset justification

Evaluating chiral interactions on an enzyme binding pocket in theory is a complex task. While some stereoisomers will interact differently with the enzyme, others will show few differences. That is, the chiral portion of the small molecule structure may not be relevant to the binding thermodynamics, such that two enantiomers may even interact with identical binding energies. This uncertainty of whether or not chirality is relevant presents an important challenge, and one that must be present when designing a benchmark for this task.

6.3.2 D4DCHP dataset construction

We filter the D4 docking study from \[24\] by the generalized Bemis-Murcko scaffold 1,3-dicyclohexylpropane (hence the dataset name D4DCHP). Additionally, we mandate that all molecules have at least one tetrahedral chiral center (i.e. we remove molecules with chiral centers with fewer than four neighbors\[^1\], such as with sulfoxides) and further filter the dataset to include only stereoisomer pairs (i.e., enantiomers or diastereomers), removing structures without such a complement in the dataset. The constitutes the full dataset. We also create two subsets of this full dataset: one retains pairs of enantiomers with differences in docking score greater than five kcal/mol (DIFF5) while the other retains molecules with only one tetrahedral center (CHIRAL1). Note that the provided d4_docking.csv file contains the full dataset, while the \{subset\}/split{x}.npy files define the subsets.

6.3.3 Synthetic R/S dataset generation

For the toy R/S classification task, we use the CHIRAL1 dataset and label each pair of enantiomers with a 0/1 denoting that the single stereocenter is R/S. The assignment is made using RDKit version 2020.03.2 \[23\]. We keep pairs of stereoisomers together in the training/validation/testing sets but determine each split randomly, and we only run one split. The dataset is provided as d4_docking_rs.csv and the split we used for the results is rs/split0.npy.

6.3.4 D4DCHP dataset statistics

Table 4 shows various statistics for the full dataset and its subsets. The top left of Figure 5 shows the data distribution for the full set, highlighting the large range of values present. The bottom left of Figure 5 shows the distribution obtained by taking the difference between all pairs on stereoisomers in the full dataset. This reinforces the earlier justification of the D4DCHP dataset; while some pairs show little to no relative difference in docking score, others show large differences. Thus, tetrahedral stereochemistry has a complex effect on this function, rendering this dataset a challenging but realistic challenge.

\[^1\] Chiral carbons with three heavy atom neighbors and one hydrogen are retained.
one for development of chiral representation models. The rest of Figure 4 shows the scaffold used to extract the data and example structures from D4DCHP.

| Statistics                        | FULL | DIFF5 | CHIRAL1 |
|-----------------------------------|------|-------|---------|
| Number datapoints                 | 287468 | 119166 | 204778  |
| Min (kcal/mol)                    | -71.29 | -71.29 | -67.54  |
| Max (kcal/mol)                    | 86.94  | 86.94  | 86.94   |
| Mean (kcal/mol)                   | -30.67 | -29.32 | -31.19  |
| Standard deviation (kcal/mol)     | 10.07  | 11.86  | 9.91    |

Figure 4: (Top left) Distribution of docking scores for full dataset. (Bottom left) Distribution of difference in docking score between pairs of stereoisomers. (Right) Generalized Bemis-Murcko scaffold used to subset data from original study and example structures with small and large differences in docking score.

6.4 Experimental details

6.4.1 Featurization

We convert each SMILES in the csv files to PyTorch Geometric data objects with node and edge features, detailed in Table 5, using RDKit [23]. For structures with multiple fragments, we only retain the largest fragment. This only affects one entry in the Lipo dataset and does not affect the D4DCHP dataset, as all molecules in the dataset consist of a single structure. We include only heavy atoms (i.e. non-hydrogen atoms) in the molecular graph. However, for chiral centers that include
a hydrogen atom as one of the four substituents, we make the attached hydrogen explicit; this is the case for a significant number of structures in the dataset. This construction facilitates use of the custom aggregation functions, but note that even for the sum aggregator, we make such hydrogens explicit in the molecular graph.

### Table 5: Atom and bond features

| Indices | Description | Options | Type   |
|---------|-------------|---------|--------|
| 0-11    | atom identity | H, C, N, O, F, Si, P, S, Cl, Br, I, other | one-hot |
| 12-18   | degree      | 0, 1, 2, 3, 4, 5, other | one-hot |
| 19-24   | formal charge | -2, -1, 0, 1, 2, other | one-hot |
| 25-30   | total number of Hs on atom | 0, 1, 2, 3, 4, other | one-hot |
| 31-36   | hybridization | sp, sp², sp³, sp³d, sp³d², other | one-hot |
| 37      | aromaticity | true, false | one-hot |
| 38      | atomic mass | R > 0 value * 0.1 | |
| 39-41   | local chirality (optional) | CW, CCW, other | one-hot |
| 42-44   | global chirality (optional) | R, S, other | one-hot |

| Indices | Description | Options | Type   |
|---------|-------------|---------|--------|
| 0       | bond presence | true, false | one-hot |
| 1-4     | bond type | single, double, triple, aromatic | one-hot |
| 5       | conjugation | true, false | one-hot |
| 6       | presence in ring | true, false | one-hot |

#### 6.4.2 Tetrahedral centers

We take advantage of RDKit’s internal bookkeeping to assign CW/CCW labels to chiral tetrahedral centers [23]. Before calculating the updated node representation for a custom aggregation function, we artificially change all tetrahedral centers to the same chiral grouping so they can be described by the same parity bit. That is, for all tetrahedral centers marked as CCW, we switch the order of the first and second neighbors as defined by RDKit before applying permutations. This allows us to feed all tetrahedral centers through consistently-parametrized matrices and ensures that the correct permutations are applied to all tetrahedral chiral centers.

#### 6.4.3 Training

We split the data into training/validation/testing sets using an 80/10/10 split. The reported statistics are averages of three runs, each using different splits, and the sample standard deviation. We run each method for a fixed 80 epochs and retain the model with the best validation performance. For the the D4DCHP dataset, we keep pairs of stereoisomers together in the training/validation/testing sets but determine each split randomly. All models use the same seed to initialize network weights, ensuring consistency between runs.

We use the Noam scheduler implementation from Yang et al. [35], which increases the learning rate linearly to the max rate for two warm-up epochs, and decreases it exponentially afterwards to the final rate. In our implementation, we ask the user to supply the maximum rate, setting the initial and final rates to be a tenth of the max rate.

For the D4DCHP dataset, we choose hyperparameters by using a simple grid scan for the DMPNN graph architecture with the sum aggregator. We use the final hyperparameters, which provide good performances across all base architectures, for all test runs with the other aggregators (Table 6).
Table 6: D4DCHP hyperparameter choices

| Hyperparameter     | Sample range       | Final choice |
|--------------------|--------------------|--------------|
| Hidden size        | 300, 600, 900, 1200| 300          |
| Depth              | 2, 3, 4, 5, 6      | 3            |
| Dropout            | 0, 0.2, 0.4, 0.6, 0.8, 1 | 0.2        |
| Max learning rate  | 1e-5, 1e-4, 1e-3   | 1e-4         |
| Batch size         | 25, 50, 100        | 50           |

Table 7: Classification of R/S stereocenters of CHIRAL1 test set for all custom aggregators (%)

| Aggregation method | Graph architecture | GCN | GIN | DMPNN |
|--------------------|--------------------|-----|-----|-------|
| SUM                | 50                 | 50  | 50  |
| PD                 | 100                | 100 | 100 |
| PERM               | 100                | 100 | 98  |
| PERM_CAT           | 100                | 100 | 98  |

6.5 Additional results

6.5.1 R/S Classification

Table 7 includes the R/S classification results for the product of differences aggregator, which shows that it too can perfectly distinguish between chiral centers.

6.5.2 D4DCHP

Table 8 shows the full validation and test RMSE values from the D4DCHP dataset. The validation results follow the same trends as the test results in that GCN architecture benefits from the custom aggregators, the DMPNN architecture shows improvements in rank classification but not RMSE with the custom aggregators, and the GIN architecture achieves good performance with only a sum aggregator and chiral features. RMSE results of the PD aggregation are consistently worse than others. While the method is constructed to achieve specific equivariances, a product of differences has little physical motivation relative to other aggregation schemes more akin to belief propagation.

Figure 5 shows rank classification results for different threshold values (such that a threshold of 5 kcal/mol corresponds to the results in Table 2). The PERM_CAT aggregator is consistently the best across graph architectures. With large differences in stereoisomers, all aggregators tend to increase rank classification accuracy for the GIN and DMPNN architectures. While the PD aggregator showed worse RMSE than the baseline SUM aggregator, it is able to rank stereoisomers correctly more than 50% of the time.

6.5.3 Lipophilicity

We also test our custom aggregation functions on the standard lipophilicity MoleculeNet dataset without pairs of stereoisomers present [33]. Over 25% of these data contain a chiral center, but because the dataset is so small and stereoisomer pairs are not present, we posit that the data are not rich enough for our aggregators to learn information about chirality. Table 9 shows these results, which agree with our hypothesis. Without a substantial number of chiral examples, the custom aggregators cannot learn nuanced trends and achieve similar performance to the baseline SUM aggregator. Note that the small size of the dataset allows us to perform a full hyperparameter optimization for each individual run, using the hyperparameters and sample ranges in Table 6 and a budget of 128 combinations. We perform this search using the Optuna framework [1].
Table 8: D4DCHP RMSE validation/test results for full dataset and rank classification results for DIFF5 subset

| Aggregation method | Graph architecture | RMSE (validation/test) | Rank classification (%) |
|--------------------|--------------------|-------------------------|--------------------------|
|                    | GCN                | GIN                     | DMPNN                    |
| SUM                | 6.75 ± 0.05 / 6.77 ± 0.05 | 6.43 ± 0.04 / 6.45 ± 0.07 | 6.41 ± 0.02 / 6.45 ± 0.06 |
| SUM (LCF)          | 6.74 ± 0.06 / 6.76 ± 0.04 | 6.39 ± 0.03 / 6.42 ± 0.06 | 6.39 ± 0.02 / 6.41 ± 0.06 |
| SUM (GCF)          | 6.72 ± 0.05 / 6.73 ± 0.05 | 6.36 ± 0.03 / 6.38 ± 0.06 | 6.36 ± 0.03 / 6.39 ± 0.06 |
| SUM (BCF)          | 6.72 ± 0.07 / 6.95 ± 0.30 | 6.35 ± 0.04 / 6.38 ± 0.06 | 6.35 ± 0.03 / 6.37 ± 0.06 |
| PD                 | 7.02 ± 0.06 / 7.03 ± 0.05 | 6.62 ± 0.06 / 6.63 ± 0.04 | 6.64 ± 0.07 / 6.64 ± 0.06 |
| PD (LCF)           | 7.02 ± 0.06 / 7.04 ± 0.06 | 6.64 ± 0.04 / 6.66 ± 0.05 | 6.64 ± 0.06 / 6.64 ± 0.06 |
| PD (GCF)           | 7.00 ± 0.04 / 7.03 ± 0.05 | 6.64 ± 0.05 / 6.66 ± 0.05 | 6.59 ± 0.04 / 6.58 ± 0.05 |
| PD (BCF)           | 7.01 ± 0.05 / 7.02 ± 0.05 | 6.65 ± 0.06 / 6.66 ± 0.06 | 6.57 ± 0.04 / 6.59 ± 0.05 |
| PERM               | 6.67 ± 0.05 / 6.69 ± 0.06 | 6.41 ± 0.04 / 6.44 ± 0.06 | 6.40 ± 0.04 / 6.42 ± 0.05 |
| PERM (LCF)         | 6.65 ± 0.05 / 6.69 ± 0.07 | 6.38 ± 0.04 / 6.41 ± 0.06 | 6.39 ± 0.04 / 6.41 ± 0.05 |
| PERM (GCF)         | 6.67 ± 0.03 / 6.68 ± 0.06 | 6.38 ± 0.04 / 6.40 ± 0.07 | 6.37 ± 0.04 / 6.40 ± 0.05 |
| PERM (BCF)         | 6.64 ± 0.04 / 6.67 ± 0.06 | 6.37 ± 0.04 / 6.39 ± 0.06 | 6.37 ± 0.04 / 6.39 ± 0.06 |
| PERM_CAT           | 6.61 ± 0.03 / 6.65 ± 0.06 | 6.37 ± 0.03 / 6.40 ± 0.05 | 6.38 ± 0.03 / 6.40 ± 0.06 |
| PERM_CAT (LCF)     | 6.59 ± 0.04 / 6.62 ± 0.06 | 6.36 ± 0.04 / 6.38 ± 0.06 | 6.37 ± 0.03 / 6.40 ± 0.05 |
| PERM_CAT (GCF)     | 6.61 ± 0.03 / 6.64 ± 0.05 | 6.36 ± 0.03 / 6.40 ± 0.06 | 6.36 ± 0.04 / 6.38 ± 0.06 |
| PERM_CAT (BCF)     | 6.63 ± 0.04 / 6.65 ± 0.06 | 6.35 ± 0.03 / 6.37 ± 0.07 | 6.35 ± 0.03 / 6.39 ± 0.07 |

| Aggregation method | Graph architecture | |
|--------------------|--------------------|------------------|
|                    | GCN                | GIN              |
| SUM                | 50.0 ± 0.0         | 50.0 ± 0.0       |
| SUM (LCF)          | 55.6 ± 0.1         | 57.9 ± 0.3       |
| SUM (GCF)          | 57.9 ± 0.5         | 61.5 ± 0.2       |
| SUM (BCF)          | 58.7 ± 0.7         | 62.1 ± 0.1       |
| PD                 | 57.3 ± 0.9         | 58.9 ± 0.4       |
| PD (LCF)           | 57.6 ± 0.4         | 59.0 ± 0.4       |
| PD (GCF)           | 57.5 ± 0.5         | 58.2 ± 0.3       |
| PD (BCF)           | 58.1 ± 0.4         | 58.9 ± 0.2       |
| PERM               | 58.3 ± 0.5         | 58.6 ± 0.3       |
| PERM (LCF)         | 60.4 ± 0.5         | 61.1 ± 0.1       |
| PERM (GCF)         | 59.8 ± 0.5         | 61.2 ± 0.4       |
| PERM (BCF)         | 60.7 ± 0.3         | 61.5 ± 0.2       |
| PERM_CAT           | 60.5 ± 0.2         | 60.5 ± 0.3       |
| PERM_CAT (LCF)     | 61.0 ± 0.1         | 62.0 ± 0.3       |
| PERM_CAT (GCF)     | 60.7 ± 0.2         | 61.8 ± 0.5       |
| PERM_CAT (BCF)     | 60.8 ± 0.1         | 62.2 ± 0.1       |
Figure 5: Rank classification accuracy for different test subsets selected based on a minimum threshold for the difference in ground truth docking score. Accuracy generally increases with larger ground truth differences between stereoisomers.

Table 9: Lipo test RMSE

| Aggregation method | GCN       | GIN       | DMPNN     |
|--------------------|-----------|-----------|-----------|
| SUM                | 0.63 ± 0.04 | 0.64 ± 0.02 | 0.60 ± 0.01 |
| SUM (LCF)          | 0.64 ± 0.03 | 0.64 ± 0.01 | 0.60 ± 0.01 |
| PD                 | 0.65 ± 0.01 | 0.67 ± 0.02 | 0.68 ± 0.02 |
| PD (LCF)           | 0.67 ± 0.05 | 0.69 ± 0.03 | 0.67 ± 0.01 |
| PERM               | 0.65 ± 0.05 | 0.66 ± 0.01 | 0.62 ± 0.01 |
| PERM (LCF)         | 0.65 ± 0.02 | 0.65 ± 0.02 | 0.62 ± 0.01 |
| PERM_CAT           | 0.66 ± 0.03 | 0.65 ± 0.03 | 0.62 ± 0.01 |
| PERM_CAT (LCF)     | 0.67 ± 0.04 | 0.65 ± 0.01 | 0.62 ± 0.02 |