
Similarity-Quantized Relative Difference Learning for Improved Molecular Activity Prediction

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Abstract

Accurate prediction of molecular activities is crucial for efficient drug discovery, yet remains challenging due to limited and noisy datasets. We introduce Similarity-Quantized Relative Learning (SQRL), a learning framework that reformulates molecular activity prediction as relative difference learning between structurally similar pairs of compounds. SQRL uses precomputed molecular similarities to enhance training of graph neural networks and other architectures, and significantly improves accuracy and generalization in low-data regimes common in drug discovery. We demonstrate its broad applicability and real-world potential through benchmarking on public datasets as well as proprietary industry data. Our findings demonstrate that leveraging similarity-aware relative differences provides an effective paradigm for molecular activity prediction.

1 Introduction and Background

The ability to predict molecular activity is critical for small molecule drug discovery. However, experimental data for training machine learning models are often limited and noisy, making robust generalization challenging. While deep learning approaches have enabled rich representations from chemical structures [1–13], most focus on predicting absolute property values, ignoring valuable information in relationships between structurally similar molecules.

Drawing inspiration from medicinal chemists [14], who examine how specific structural modifications influence properties relative to a parent compound or a matched molecular pair [15], we introduce **Similarity-Quantized Relative Learning (SQRL)**—a training and evaluation framework that reformulates property prediction as learning relative differences between nearby compounds.

The key contributions of our work are:

- We introduce a robust **similarity-thresholded learning approach** that significantly enhances model performance by focusing on **predicting property differences between the most informative compound pairs**. This allows effective learning from limited and noisy data.
- Our analysis across molecular distance metrics and thresholds demonstrates that **similarity-aware dataset matching outperforms indiscriminate pairing of all inputs**, as it more effectively leverages local structural information.
- Extensive **benchmarking** demonstrating the benefits of SQRL **across diverse state-of-the-art network architectures** and multiple activity prediction datasets, including publicly available activity cliff prediction tasks and real-world industry datasets.

2 Related Work

Molecular property prediction. Significant research has focused on methods for molecular property and activity prediction, including recent work on graph neural networks (GNNs) [1–7] along with chemical language models [8–12] that learn meaningful molecular representations from molecular data [13]. Despite these advances, simpler tree-based models often outperform more complex neural approaches due to limited data availability and inherent modeling challenges [16, 17].

Activity cliff prediction. Activity cliffs refer to pairs of molecules with high structural similarity but significantly different activity levels. Previous approaches have addressed this challenging problem by representing molecular graphs as images [18, 19], applying graph convolutional networks (GCNs) to matched molecular pairs [20], or using chemical reaction information [21]. Many of these methods have formulated the problem as a classification task, aiming to identify whether a given molecular pair exhibits an activity cliff [22] or as a standard regression task that relies on learning the discontinuous chemical space directly from the data [23]. Our approach focuses on predicting the difference in potency values between any pair of similar molecules, providing a more versatile solution that can also be applied to standard potency prediction.

Metric, similarity, and few-shot learning. Pairwise learning approaches have been widely adopted in fields like ranking, metric, and similarity learning [24, 25]. Notably, few-shot learning approaches have been developed for molecular property prediction for improving generalization in low-data regimes [26–28]. Additionally, pairwise data matching has proven effective in implicit guidance of generative models for drug design. [29].

Relative prediction. Recently, pairwise learning has been applied to regression tasks. [Wetzel et al.](#) used Siamese networks to predict differences between all data points in both supervised and unsupervised settings as a way of producing ensembles of predictions and uncertainty estimates [30, 31]. This approach has been extended to classification tasks for tree-based models [32]. [Tynes et al.](#) applied the concept of pairwise learning to computational chemistry, analyzing the performance of random forest models trained on all pairs of inputs points [33]. Similarly, [Fralish et al.](#) trained a D-MPNN model on paired compounds and observed improvements in ADME property predictions [34] and molecule selection in an active learning setting [35].

Unlike previous approaches that primarily focus on absolute property predictions or indiscriminate pairwise learning, our work introduces a similarity-thresholded framework that emphasizes learning from the most informative compound pairs.

3 Similarity-Thresholded Relative Representation

Problem formulation. We formulate the relative prediction task as follows. Given a dataset of molecular structures $\mathcal{D} = \{(x_i, y_i)\}_{i=1}^N$, where x_i represents molecule i and $y_i \in \mathbb{R}$ denotes its corresponding property value, our goal is to learn a function $f : \mathcal{X} \times \mathcal{X} \rightarrow \mathbb{R}$ that predicts the relative difference in property values between two molecules. Formally, for any pair of molecules (x_i, x_j) , we aim to predict the relative difference $\Delta y_{ij} = y_i - y_j$.

Dataset matching. To train models on the relative prediction task, we construct a new dataset \mathcal{D}_{rel} by considering pairs of molecules in the original dataset \mathcal{D} . To focus on the most informative comparisons, we restrict the pairs to those within a certain threshold of structural similarity, as measured by a predefined similarity metric (e.g., Tanimoto similarity). This allows models to learn from local differences in chemical space where relative changes in property values are most meaningful. We define \mathcal{D}_{rel} as:

$$\mathcal{D}_{\text{rel}} = \{((x_i, x_j), \Delta y_{ij}) \mid x_i, x_j \in \mathcal{D}, d(x_i, x_j) \leq \alpha\} \quad (1)$$

where $d : \mathcal{X} \times \mathcal{X} \rightarrow \mathbb{R}_{\geq 0}$ is a distance function in the input space \mathcal{X} and $\alpha \in \mathbb{R}_{>0}$ is a distance threshold. See Appendix A.4 for examples of paired structures.

The choice of α is critical, as it involves a trade-off between the quantity and relevance of generated pairs. We propose selecting α based on the distribution of distances in the training data, specifically by choosing a threshold smaller than the average pairwise distance. This approach can help form more informative pairs by focusing on molecules with greater structural similarity and relevance.

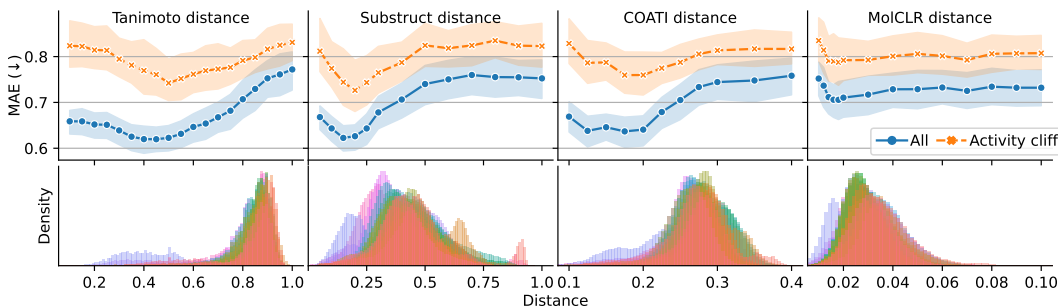


Figure 1: **Leveraging local structural information enhances predictive performance.** *Top:* Incorporating neighbors only up to a certain distance threshold α improves MAE (\downarrow). *Bottom:* Pairwise distance distributions of training data (overlaid for all 30 MoleculeACE tasks) with greater skewness and kurtosis yield the best performance and a wider range of acceptable values of α .

Relative representation. We define $g : \mathcal{X} \rightarrow \mathbb{R}^d$ as a mapping function that converts a molecular compound from the input space \mathcal{X} into a d -dimensional real-valued vector. This can be either a learnable model (such as a graph neural network), a pre-trained model, or a fixed molecular fingerprinting algorithm. Next, a machine learning model $f : \mathbb{R}^d \rightarrow \mathbb{R}$ uses the difference between molecular representations generated by g to predict the relative differences in properties.

We optimize parameters θ of f and g (if learnable) by minimizing:

$$\min_{\theta} \mathcal{L}(\theta) = \min_{\theta} \sum_{((x_i, x_j), \Delta y_{ij}) \in \mathcal{D}_{\text{rel}}} \ell(f(g(x_i) - g(x_j)), \Delta y_{ij}) \quad (2)$$

where ℓ is mean squared error loss. For a new molecule x_{new} , we compute the prediction \hat{y}_{new} as:

$$\hat{y}_{\text{new}} = \frac{1}{n} \sum_{x_i \in \text{NN}_n(x_{\text{new}})} y_i + f(g(x_i) - g(x_{\text{new}})) \quad (3)$$

where $\text{NN}_n(x_{\text{new}})$ denotes the set of n molecules from the training data \mathcal{D} that are nearest to x_{new} as determined by the distance function $d(x_i, x_{\text{new}})$. Unless otherwise specified, we set $n = 1$.

4 Experimental Results

4.1 Experimental setup

To understand the generality of SQRL, we conducted extensive evaluations across a diverse set of models and molecular activity datasets. Each model was trained to predict absolute property values directly (Standard) or to predict relative differences between selected molecule pairs (SQRL). More details about models, hyperparameter selection, and datasets can be found in Appendices A.1–A.4.

Models. We benchmarked baselines (RF [36], XGBoost [37], KNN) using Morgan fingerprints [38] with RDKit features [39], MLP with Morgan fingerprints, GNNs (AttentiveFP [4], GINE [40], PNA [41], MolCLR [42]), and transformer models (COATI [43], SAFE-GPT [44], and Uni-Mol [45]).

Distance metrics. Several distance metrics were used to create training data pairs and obtain the closest training molecules for inference. We evaluated Tanimoto distances between Morgan fingerprints [38], Tanimoto distances between substructure count vectors [46], as well as Euclidean distances using COATI [43], Uni-Mol, and MolCLR embeddings [42] (see A.5).

Datasets. We evaluated our approach on 30 activity prediction tasks ($\text{pEC}_{50}/\text{p}K_i$) for ChEMBL targets using the MoleculeACE dataset [23]. This dataset includes challenging activity cliff molecules (MoleculeACE-Cliff)—structurally similar compounds with large activity differences—providing a crucial test of local generalizability (see A.4). Additionally, we evaluated models on 5 proprietary drug discovery projects (Internal Targets) to assess the real-world applicability of our approach. Models are trained on single tasks and results are reported aggregated across tasks.

Table 1: **Spearman’s ρ (\uparrow) comparison of predictive performance.** Spearman’s ρ with standard deviation across tasks for Standard and SQRL methods using Tanimoto distance with $\alpha = 0.7$. Uni-Mol was evaluated on a subset of tasks due to computational constraints. See Appendix A.1 for more details and MAE scores (Table 3).

Model	MoleculeACE		MoleculeACE-Cliff		Internal Targets	
	Standard	SQRL	Standard	SQRL	Standard	SQRL
<i>Baselines</i>						
XGBoost	0.79 \pm 0.10	0.76 \pm 0.08	0.72 \pm 0.18	0.64 \pm 0.15	0.73 \pm 0.14	0.65 \pm 0.16
RF	0.80 \pm 0.09	0.77 \pm 0.07	0.72 \pm 0.18	0.68 \pm 0.14	0.70 \pm 0.13	0.68 \pm 0.14
KNN	0.66 \pm 0.10	0.67 \pm 0.13	0.57 \pm 0.19	0.59 \pm 0.23	0.52 \pm 0.14	0.52 \pm 0.14
MLP	0.32 \pm 0.17	0.73 \pm 0.09	0.23 \pm 0.16	0.62 \pm 0.16	0.39 \pm 0.14	0.59 \pm 0.20
<i>GNNs</i>						
AttentiveFP	0.52 \pm 0.20	0.77 \pm 0.09	0.43 \pm 0.19	0.67 \pm 0.17	0.49 \pm 0.13	0.66 \pm 0.19
GINE	0.33 \pm 0.19	0.76 \pm 0.09	0.29 \pm 0.21	0.68 \pm 0.18	0.40 \pm 0.22	0.61 \pm 0.17
PNA	0.51 \pm 0.18	0.72 \pm 0.08	0.41 \pm 0.17	0.61 \pm 0.18	0.52 \pm 0.12	0.64 \pm 0.15
MolCLR	0.35 \pm 0.22	0.77 \pm 0.09	0.28 \pm 0.20	0.66 \pm 0.18	0.39 \pm 0.27	0.68 \pm 0.17
<i>Transformers</i>						
COATI	0.69 \pm 0.11	0.74 \pm 0.09	0.59 \pm 0.15	0.64 \pm 0.16	0.52 \pm 0.26	0.61 \pm 0.18
Uni-Mol	0.26 \pm 0.19	0.69 \pm 0.10	0.19 \pm 0.20	0.57 \pm 0.18	0.40 \pm 0.16	0.51 \pm 0.14
SAFE-GPT	0.61 \pm 0.20	0.71 \pm 0.08	0.56 \pm 0.22	0.61 \pm 0.14	0.58 \pm 0.14	0.59 \pm 0.15

4.2 Leveraging local structural information improves learning

A key hypothesis of SQRL is that not all possible relative pairs are equally informative, and training on all pairwise comparisons as done previously [32–34] may overemphasize global relationships at the expense of local consistency. We analyzed pairwise distance distributions using various distance metrics and found that some distributions exhibit significant left-skew and/or high kurtosis (Figure 1 and Appendix Figure 5). We hypothesize that these highly similar compounds contain the most informative signal. To test this assumption, we trained MLP models on top of Morgan fingerprint features across a range of similarity thresholds for each of the metrics (Figure 1). We observed that incorporating neighbors up to a certain threshold provides a pronounced improvement in performance, but beyond this point, including more dissimilar pairs degrades performance. Moreover, performance is best for distance metrics with the desired distribution characteristics (e.g., Tanimoto and COATI). These results demonstrate the benefit of our similarity-thresholded approach: a smaller distance threshold α yields fewer but potentially more informative pairs, while a larger threshold increases pair count but may introduce less relevant comparisons. Our findings support the hypothesis that focusing on the most similar pairs effectively leverages local structural information (Appendix Figure 6), leading to improved predictive performance across various model architectures and datasets.

4.3 SQRL consistently improves predictive performance across all neural network architectures

To understand which model types benefit from this approach, we performed extensive benchmarking of molecular property prediction models trained using both the standard absolute prediction objective and SQRL. While the baseline models did not significantly benefit from relative training, all deep learning architectures exhibit consistent improvement of the Spearman rank correlation across all datasets when trained with SQRL (Table 1). We observe significant improvements for all GNN architectures and especially for the pre-trained transformer-based models—0.57 and 0.43 point improvement for COATI and Uni-Mol, respectively. Notably, substantial improvements are observed on the MoleculeACE-Cliff subset, highlighting SQRL’s ability to capture fine-grained structural differences that significantly impact molecular properties. Evaluation on internal targets shows that the observed benefits are transferable to real-world scenarios and underscore the robustness and broad applicability of this approach.

The lack of improvement for XGBoost, RF, and KNN with SQRL is notable. This may indicate that the simple difference fingerprint representation is not sufficient for these models as it forces these models to only learn from substructures that differ between a pair of molecules without taking into account the rest of the molecular structure of both molecules. A higher fidelity representation, such as concatenating the full molecular fingerprints to the difference representation, may overcome this limitation. Even so, most modern deep learning methods do not outperform conventional baselines (with or without SQRL). The small size of the training data for each task likely contributes to this result. However, we expect that neural network approaches may outperform baselines with additional tuning (e.g., more rigorous hyperparameter selection, choosing a more optimal distance threshold, or redesigning the training objective).

5 Conclusions and Future Directions

Overall, our work demonstrates consistent improvements for neural networks in molecular activity prediction using SQRL, particularly in capturing molecular activity cliffs. Our method shows promise in learning from pairwise differences, potentially offering a more nuanced understanding of structure-activity relationships. The limitations of the current work include the assumption that meaningful distance metrics are available. Future research could focus on refining SQRL for applications where similarity measures are less well-defined or more challenging to establish.

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A Appendix

A.1 Models

We used the following models to evaluate the effectiveness of the SQRL approach:

Baselines. All baseline models were trained on top of Morgan count fingerprints of size 2048, radius 2, and including chirality. For RF, XGBoost, and KNN, RDKit features from [39] were concatenated to Morgan fingerprints.

- **Random Forest (RF):** An ensemble learning method using decision trees. Scikit-learn implementation was used [47] with default parameters.
- **XGBoost:** A gradient boosting framework, optimized for efficiency and performance. Implementation from Ref. 37 was used with default parameters.
- **k-Nearest Neighbors (KNN):** A non-parametric method based on the similarity between data points. Scikit-learn implementation was used [47] with $k = 1$ to compare with our relative setting where we evaluate with respect to the closest training data point.
- **Multi-Layer Perceptron (MLP):** A standard feedforward neural network.

Graph neural networks. GNNs were trained end-to-end, with their learned representations followed by MLP layers for standard or relative predictions.

- **AttentiveFP:** A graph neural network that uses graph attention mechanisms to capture atomic interactions [4].
- **GINE:** Graph Isomorphism Network with Edge features, enhancing the model’s ability to capture bond information [40].
- **PNA:** Principal Neighbourhood Aggregation, a GNN architecture designed to be stable under permutations [41]. The implementation used in this paper follows Ref. 28.
- **MolCLR:** A pre-trained GNN using contrastive learning for molecular representation. In our experiments, we did not freeze the MolCLR layers, allowing them to be fine-tuned along with the rest of the model, according to the procedure of the original authors [42].

Transformer-based models. For transformer-based models, we utilized the pre-trained embeddings as fixed feature extractors, followed by MLP for task-specific predictions.

- **SAFE-GPT:** A chemical language model trained on linear molecular notation which has been adapted for autoregressive tasks [44].

- **Uni-Mol**: A 3D-aware transformer model trained on molecular conformations using SE(3) equivariant operations [45]. Due to computationally expensive conformation generation,
- **COATI**: A multi-modal generative model combining 2D and 3D molecular information through contrastive learning [43]. To obtain embeddings for our applications, the text encoder part of the model was used (Barlow_Closed).

Due to the computationally expensive conformer generation step, Uni-Mol-SQRL was evaluated on the following subset of MoleculeACE tasks (23/30): CHEMBL2971_Ki, CHEMBL2835_Ki, CHEMBL219_Ki, CHEMBL228_Ki, CHEMBL238_Ki, CHEMBL1862_Ki, CHEMBL218_EC50, CHEMBL231_Ki, CHEMBL235_EC50, CHEMBL287_Ki, CHEMBL2147_Ki, CHEMBL2047_EC50, CHEMBL4203_Ki, CHEMBL2034_Ki, CHEMBL1871_Ki, CHEMBL4792_Ki, CHEMBL244_Ki, CHEMBL234_Ki, CHEMBL239_EC50, CHEMBL262_Ki, CHEMBL4616_EC50, CHEMBL3979_EC50, CHEMBL4005_Ki.

A.2 Hyperparameter selection

See Table 2 for optimized hyperparameters used for each model in a standard and SQRL setting.

Table 2: Model hyperparameters.

Model	Linear Sizes	GNN Layers	Dropout	Learning Rate	Batch Size	Other
MLP	[256, 256]	-	0.0	1e-4	128	-
MLP-SQRL	[512, 256]	-	0.2	1e-5	64	-
AttentiveFP	[256]	3	0.2	1e-4	128	timesteps: 2
AttentiveFP-SQRL	[128]	3	0.0	1e-3	256	timesteps: 4
GINE	[128]	4	0.0	1e-4	64	-
GINE-SQRL	[256, 128]	5	0.0	1e-3	64	-
PNA	[128]	4	0.0	1e-4	64	-
PNA-SQRL	[128]	8	0.0	1e-5	256	-
MolCLR	[512]	-	0.0	1e-4	128	-
MolCLR-SQRL	[128]	-	0.0	1e-4	128	-
COATI	[256]	-	0.0	1e-5	64	-
COATI-SQRL	[256, 128]	-	0.0	1e-3	128	-
SAFE-GPT	[256, 128]	-	0.0	1e-3	32	-
SAFE-GPT-SQRL	[256, 128]	-	0.0	1e-4	128	-
Uni-Mol	[256, 128]	-	0.0	1e-3	128	-
Uni-Mol-SQRL	[128]	-	0.0	1e-4	128	-

A.3 MoleculeACE Dataset

The MoleculeACE dataset provided by [van Tilborg et al.](#) is curated from ChEMBL v29 and contains potency measurements for 30 targets. Activity cliff molecules were defined as pairs of molecules with greater than 90% substructure, scaffold, or SMILES similarity and greater than 10-fold activity difference. Figure 2 shows the number of training samples for each task in the MoleculeACE dataset along with the number of activity cliff molecules present in each task. The train-test split was performed by clustering molecules into 5 clusters and stratified splitting using the activity cliff label.

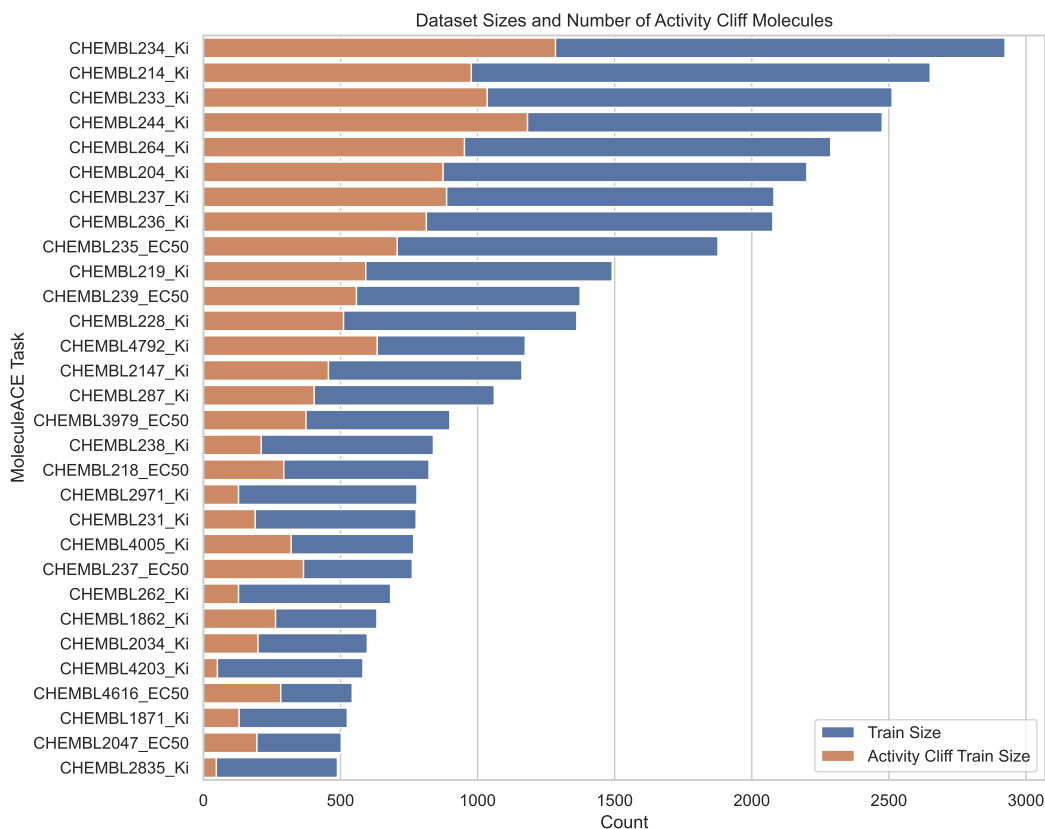


Figure 2: Training data sizes for each task in MoleculeACE.

A.4 Examples of molecular pairs

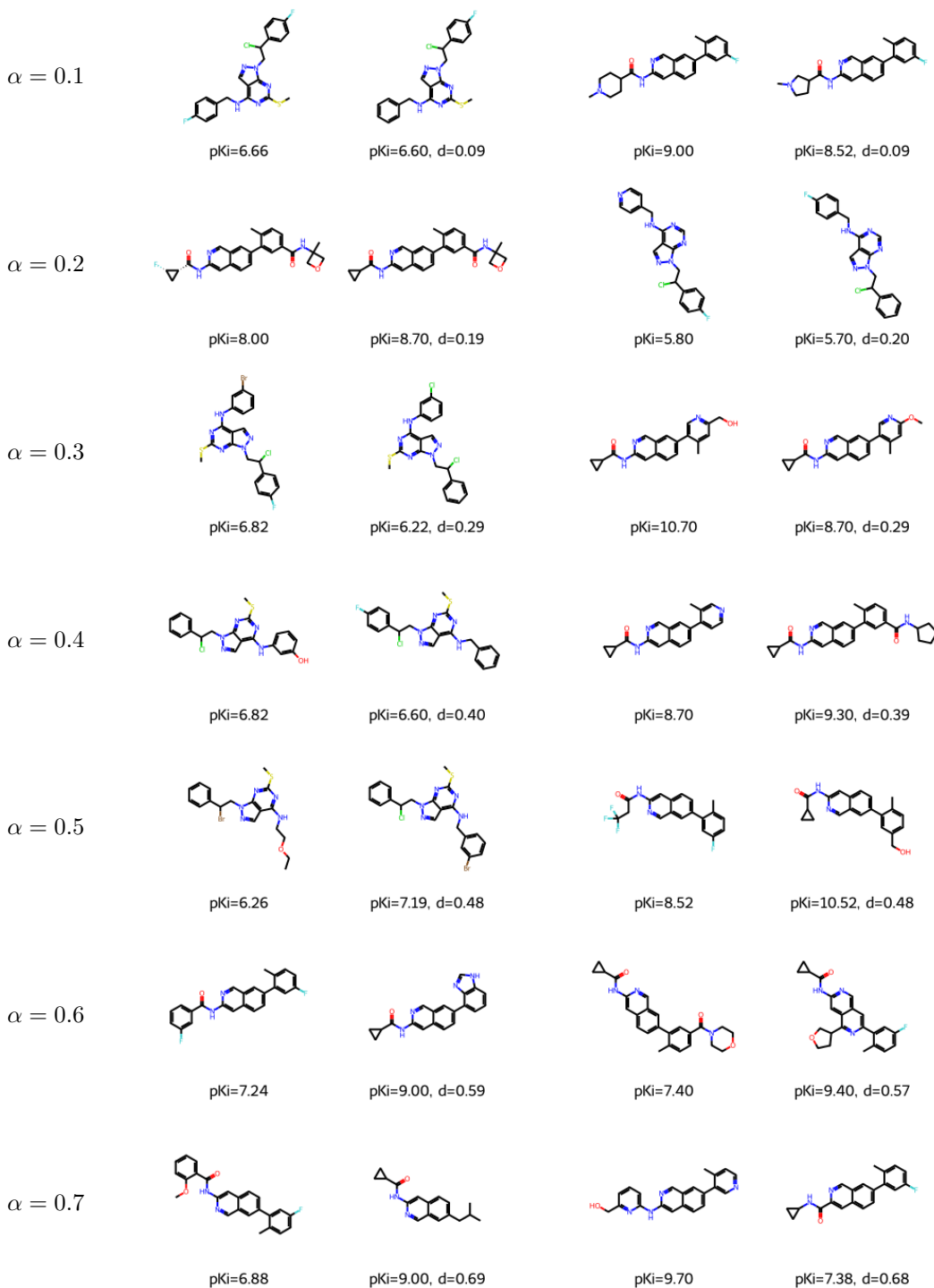


Figure 3: Molecular pairs obtained by the data matching procedure described in Section 3 at different Tanimoto distance thresholds α for MoleculeACE task CHEMBL1862_Ki.

Figure 4: Molecular pairs of *activity cliff molecules* obtained by the data matching procedure described in Section 3 at different Tanimoto distance thresholds α for MoleculeACE task CHEMBL1862_Ki.

A.5 Additional results

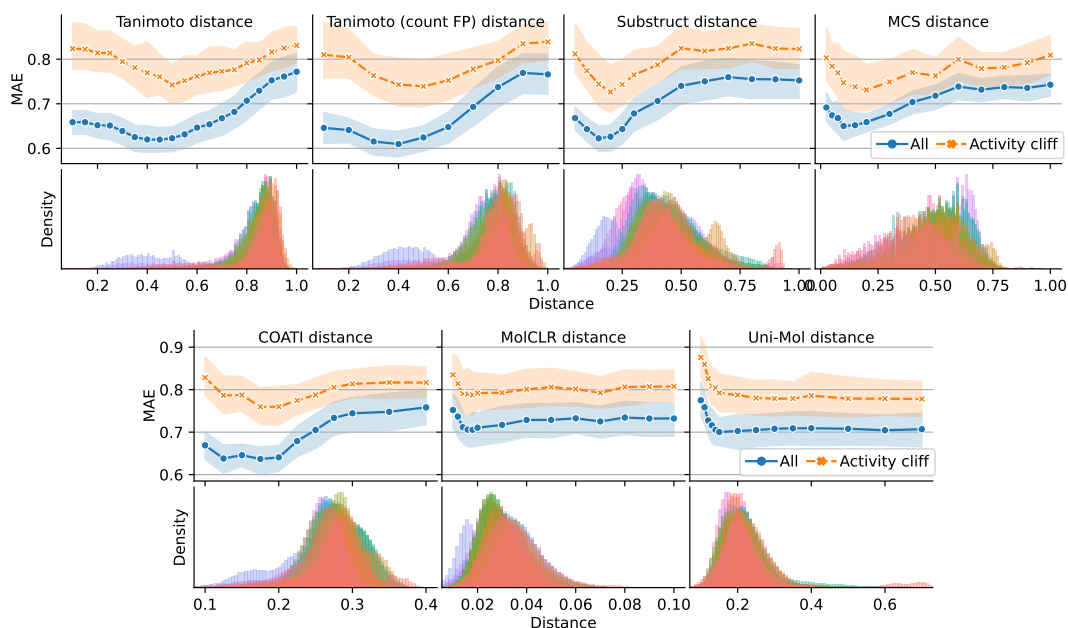


Figure 5: **Leveraging local structural information enhances predictive performance.** MAE (\downarrow) as a function of distance threshold α for several additional distance metrics compared to Figure 1, as well as pairwise distance distributions for each metric. *Tanimoto*: Tanimoto (Jaccard) distance between binary Morgan fingerprints. *Tanimoto (count FP)*: Tanimoto (Jaccard) distance between count-based Morgan fingerprints. *Substruct*: Tanimoto (Jaccard) distance between substructure count vectors using a list of 1242 predefined substructures from Ehrlich and Rarey [46]. *MCS*: Distance metric based on maximum common substructure (MCS) defined as $1 - 2N_{\text{MCS}}/(N_i + N_j)$ where N_{MCS} is the number of atoms in the MCS, N_i is the number of atoms in molecule i , and N_j is the number of atoms in molecule j . *COATI* [43], *MolCLR* [42], *Uni-Mol* [45]: Euclidean distances between neural network embeddings obtained with these pre-trained models.

Table 3: **Mean Absolute Error (MAE) (\downarrow) comparison of predictive performance.** MAE with standard deviation across tasks for Standard and SQRL methods using Tanimoto distance with $\alpha = 0.7$. **Bold values** indicate the better performing method (lower MAE) for each model and dataset. Uni-Mol was evaluated on a smaller subset of tasks due to computational constraints (see Appendix A.1.)

Model	MoleculeACE		MoleculeACE-Cliff		Internal Targets	
	Standard	SQRL	Standard	SQRL	Standard	SQRL
<i>Baselines</i>						
XGBoost	0.54 \pm 0.07	0.59 \pm 0.08	0.64 \pm 0.12	0.71 \pm 0.13	0.41 \pm 0.18	0.55 \pm 0.19
RF	0.54 \pm 0.07	0.57 \pm 0.08	0.63 \pm 0.10	0.68 \pm 0.12	0.41 \pm 0.18	0.50 \pm 0.21
KNN	0.69 \pm 0.12	0.77 \pm 0.13	0.83 \pm 0.15	0.88 \pm 0.18	0.71 \pm 0.37	0.95 \pm 0.35
MLP	0.94 \pm 0.22	0.67 \pm 0.10	0.96 \pm 0.20	0.79 \pm 0.12	0.67 \pm 0.41	0.65 \pm 0.20
<i>GNNs</i>						
AttentiveFP	0.80 \pm 0.12	0.58 \pm 0.11	0.86 \pm 0.12	0.65 \pm 0.16	0.56 \pm 0.24	0.50 \pm 0.15
GINE	1.14 \pm 0.24	0.64 \pm 0.09	1.23 \pm 0.32	0.72 \pm 0.14	0.75 \pm 0.13	0.47 \pm 0.19
PNA	0.82 \pm 0.21	0.65 \pm 0.09	0.85 \pm 0.18	0.75 \pm 0.14	0.63 \pm 0.27	0.58 \pm 0.26
MolCLR	0.91 \pm 0.12	0.62 \pm 0.09	0.95 \pm 0.13	0.72 \pm 0.14	0.62 \pm 0.39	0.50 \pm 0.21
<i>Transformers</i>						
COATI	0.71 \pm 0.10	0.65 \pm 0.09	0.74 \pm 0.20	0.75 \pm 0.14	0.59 \pm 0.30	0.61 \pm 0.20
Uni-Mol	0.94 \pm 0.26	0.69 \pm 0.09	0.92 \pm 0.22	0.77 \pm 0.14	0.62 \pm 0.41	0.49 \pm 0.48
SAFE-GPT	0.76 \pm 0.12	0.68 \pm 0.11	0.78 \pm 0.10	0.80 \pm 0.13	0.55 \pm 0.22	0.60 \pm 0.26



Figure 6: **SQRL enhances local molecular consistency.** Spearman's rank correlation coefficient (\uparrow) plotted as a function of the distance between test points and their nearest neighbors in the training set. SQRL-trained models that benefit from this training strategy demonstrate the most significant performance gains for test points with close neighbors, while generally maintaining comparable performance to standard-trained models for more distant points. Models in this plot were trained with the distance threshold $\alpha = 0.7$ using Tanimoto distance.