

# LEARNING THE NEIGHBORHOOD: CONTRAST-FREE MULTIMODAL SELF-SUPERVISED MOLECULAR GRAPH PRETRAINING

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## ABSTRACT

High-quality molecular representations are essential for property prediction and molecular design, yet large labeled datasets remain scarce. While self-supervised pretraining on molecular graphs has shown promise, many existing approaches either depend on hand-crafted augmentations or complex generative objectives, and often rely solely on 2D topology, leaving valuable 3D structural information underutilized. To address this gap, we introduce C-FREE (*Contrast-Free Representation learning on Ego-nets*), a simple framework that integrates 2D graphs with ensembles of 3D conformers. C-FREE learns molecular representations by predicting subgraph embeddings from their complementary neighborhoods in the latent space, using fixed-radius ego-nets as modeling units across different conformers. This design allows us to integrate both geometric and topological information within a hybrid Graph Neural Network (GNN)-Transformer backbone, without negatives, positional encodings, or expensive pre-processing. Pretraining on the GEOM dataset, which provides rich 3D conformational diversity, C-FREE achieves state-of-the-art results on MoleculeNet, surpassing contrastive, generative, and other multimodal self-supervised methods. Fine-tuning across datasets with diverse sizes and molecule types further demonstrates that pretraining transfers effectively to new chemical domains, highlighting the importance of 3D-informed molecular representations. Our code and checkpoints are publicly available at <https://github.com/ariguiba/C-FREE>.

## 1 INTRODUCTION

High-quality molecular representations are critical for predicting properties, interpreting chemical behavior, and accelerating compound discovery (Wigh et al., 2022; Elton et al., 2019). Many existing approaches, however, rely on a single modality, such as SMILES strings (Hirohara et al., 2018; Wang et al., 2019), 2D graph structures (Gilmer et al., 2017; Kipf & Welling, 2017; Xu et al., 2019), or 3D conformations (Schütt et al., 2017; Gasteiger et al., 2021). While effective, each of these methods captures only part of the molecular information and overlooks complementary aspects available in other modalities (Liu et al., 2023b). Beyond modality limitations, these models often require large, curated datasets, which restricts their use in low-data settings.

Because such curated datasets are often unavailable, especially in low-data regimes, self-supervised learning (SSL) provides a promising alternative to fully supervised training. Recent advances in vision and language modeling (Devlin et al., 2019; Chen et al., 2020; Grill et al., 2020a; Caron et al., 2021; He et al., 2022; LeCun & Courant, 2022) have motivated similar methods for molecular graphs, especially approaches that aim to combine structural and geometric information. While these approaches have advanced molecular representation learning, each comes with trade-offs: contrastive methods hinge on carefully chosen negative samples (You et al., 2021b;a), generative methods often require discrete reconstruction in the input graph space with graph tokenization (Hu

et al., 2020b), and latent-predictive methods can depend on augmentations or expensive procedures such as clustering (Skenderi et al., 2025). These challenges motivate simpler predictive frameworks that combine 2D topology and 3D conformations without relying on negatives or input-space reconstruction, and that work reliably across both high- and low-data settings.

**Current work.** Our self-supervised framework C-FREE (Contrast-Free Representation learning on Ego-nets) adopts a predictive learning strategy with subgraphs as the basic modeling unit. It is motivated by three goals: (i) *avoiding computationally intensive or ambiguous design choices*, including expensive augmentations, heavy subgraph-construction procedures, and complex negative-sampling schemes. For example, clustering-based subgraph algorithms such as METIS (Skenderi et al., 2025) can be costly, and defining suitable augmentations or negatives is often non-trivial, since molecules with nearly identical structures (e.g., chiral isomers) may still have very different properties; (ii) *leveraging the success of subgraph-based methods in 2D graph supervised learning* (Bevilacqua et al., 2022; Wollschläger et al., 2024), which suggest that aggregating information from substructures can yield richer graph-level representations, and (iii) *harnessing the benefits of multi-modal architectures in the supervised setting* (Zhu et al., 2024; Nguyen et al., 2024; Manolache et al., 2024). Since many molecular properties depend on multiple conformations and their probabilities (Cao et al., 2022), using multiple high-probability conformers alongside 2D topology helps capture this variability and improves predictive performance. Building on JEPA (Assran et al., 2023) and Equivariant Subgraph Aggregation Networks (ESAN) (Bevilacqua et al., 2022), our method segments graphs into disjoint subgraphs, similar to image patches or language tokens, and learns to align each subgraph with its context in latent space. Unlike GraphJEPA (Skenderi et al., 2025) and I-JEPA (Assran et al., 2023), it avoids positional encodings, hierarchical objectives, and costly clustering, and instead leverages the inductive bias of 2D and 3D encoders together with subgraph-based pre-training to learn rich embeddings. Our contributions are as follows:

1. **A new multi-modal pretraining task for molecular graphs.** We introduce a broadly applicable predictive objective based on  $k$ -EgoNet subgraphs, avoiding costly hand-crafted augmentations and utilizing both 2D and 3D views of the molecule.
2. **Robust performance in both multimodal and 2D-only settings.** Our framework leverages 2D topology together with multiple 3D conformations when available, but also performs strongly in purely 2D settings where conformers are absent.
3. **A simple and effective training scheme.** We adopt non-contrastive predictive learning, avoiding the pre-train/fine-tune mismatch and removing the need for negative samples or augmentations. Moreover, when fine-tuning, our framework simulates ESAN (Bevilacqua et al., 2022) and is provably more expressive than 1-WL (Weisfeiler & Leman, 1968)
4. **State-of-the-art results.** Our approach matches or surpasses other self-supervised models under both linear-probe evaluation, where the backbone is frozen, and full fine-tuning. It achieves the best average performance on MoleculeNet (Wu et al., 2018) and shows strong transfer to novel multimodal molecular benchmarks such as MARCEL (Zhu et al., 2024).

## 2 RELATED WORK

Existing approaches to graph self-supervised learning can be grouped into three main categories: *contrastive learning*, *generative pre-training*, and *latent representation learning*. Each of these has also been extended to molecular graphs, with varying degrees of multimodal integration.

*Contrastive learning* aligns representations of similar instances while pushing apart dissimilar ones and has become central to graph representation learning. GraphCL (You et al., 2021b) and JOAO (You et al., 2021a) pioneered this idea through graph augmentations, while InfoGraph (Sun et al., 2020) maximized mutual information across views. Extensions to molecules incorporate 3D information: GraphMVP (Liu et al., 2022a) aligns 2D topology and 3D conformations with generative objectives, MoleculeSDE (Liu et al., 2023a) introduces symmetry-aware stochastic differential equations, and 3D InfoMax (Stärk et al., 2022) encodes 3D from 2D via mutual information. While effective (Wang et al., 2023a), these methods depend on negative samples and large batches (You et al., 2021b), a limitation exacerbated by irregular graph structures.

*Generative pre-training* forms the second category of self-supervised learning, where models reconstructs masked or missing components of a graph from its surrounding context. Early works such as

AttrMask (Hu et al., 2020a), ContextPred (Hu et al., 2020a), and EdgePred (Hamilton et al., 2017) predict attributes or edges, while GPT-GNN (Hu et al., 2020b) and GROVER (Rong et al., 2020) extend to autoregressive reconstruction and chemically informed motif prediction. More recent methods incorporate multimodal and geometric signals: unified cross-modal generation of 2D/3D (Zhu et al., 2022), modality integration via MoleBlend (Yu et al., 2023), and geometry-aware prediction in 3D PGT (Wang et al., 2023b). Despite their promise, generative methods must reconstruct both discrete graph structure and continuous features, and autoregressive variants are further complicated by the lack of a natural ordering over graph nodes.

*Latent representation learning* forms the third category of self-supervised methods. Instead of reconstructing raw graph structures or features, these approaches predict target embeddings directly in latent space, yielding compact, denoised, and often across multimodal representations. BGRL (Thakoor et al., 2023) employs a bootstrapped online–target encoder scheme under augmentations, while LaGraph (Xie et al., 2022) frames the task as latent graph prediction, optimizing an upper bound with context-aware regularization on masked nodes. While latent prediction methods avoid the costly generation of negatives, they remain sensitive to augmentation quality and model update stability, and are prone to representation collapse (Assran et al., 2023; Grill et al., 2020b).

Within latent representation learning, GraphJEPa (Skenderi et al., 2025) extends the Joint Embedding Predictive Architecture (JEPa) (Assran et al., 2023) to graphs by masking METIS-generated clusters and predicting them as patch-like substructures, while also encoding hierarchical information via hyperbolic subgraph coordinates. While effective, this approach incurs significant computational cost and depends on auxiliary components—such as clustering, hierarchical encodings, positional embeddings—that add complexity without being clearly essential for representation learning.

Another direction explores large-scale supervised pretraining on massive labeled molecular datasets, aiming to transfer knowledge to downstream tasks (Beaini et al., 2024). While effective in some cases, this approach still depends on labeled data and may be domain-specific, since source and target distributions can differ. In contrast, self-supervised methods avoid this reliance on labels and can transfer more flexibly. Importantly, the two strategies are complementary: self-supervised pretraining can provide strong initializations that are later fine-tuned on labeled data.

To the best of our knowledge, our work is the first non-contrastive, non-generative predictive framework for multimodal molecular representation learning. We now turn to its design.

### 3 CONTRAST-FREE MULTIMODAL SELF-SUPERVISED PRETRAINING

In the following, we outline our proposed training pipeline, illustrated in Fig. 1. Unlike most generative methods (Hu et al., 2020a; Hamilton et al., 2017), we apply our training objective fully in the latent space, without reconstructing the original features of the masked components.

The core principle of our approach is to learn representations by aligning the embedding of a target view with that of a related context view. Specifically, we represent each molecule as a 2D graph  $G = (V, E)$ , where  $V$  is the set of nodes (atoms) and  $E$  the set of edges (covalent bonds). For each atom  $v \in V$ , we include 3D coordinates  $r_v \in \mathbb{R}^3$ , taken from multiple conformers of the molecule. Using these graph and geometric features, we construct complementary context–target views that serve as inputs for our contrast-free pretraining scheme. Each 2D and 3D view is encoded independently, and the resulting embeddings are concatenated into a single multimodal sequence processed by a transformer (Vaswani et al., 2017). We then align the embedding of the target subgraph with that of its associated context subgraph. This design is loosely inspired by ESAN (Bevilacqua et al., 2022), but adopts a simplified variant: we use fixed-radius ego-nets to obtain complementary views during pretraining, and at fine-tuning we evaluate both linear probing on whole-graph embeddings and an aggregation of subgraph embeddings using DeepSets (Zaheer et al., 2017).

**Context–Target View Generation.** We generate complementary 2D views by sampling  $k$ -EgoNets, where the  $k$ -hop neighborhood of a node defines one subgraph and the remaining nodes and edges define its complement (see Fig. 2). The 3D coordinates are added to generate the corresponding 3D views for all the conformers. Either view can serve as the target while the other acts as context, and their roles are alternated during training to avoid prediction bias. We adopt fixed-radius neighborhoods with  $k \in \{3, 4\}$ , analogous to fixed-size patches in vision-based methods (Assran et al., 2023). Although graphs vary in size and structure, this ensures that each subgraph captures a comparable amount of local information.

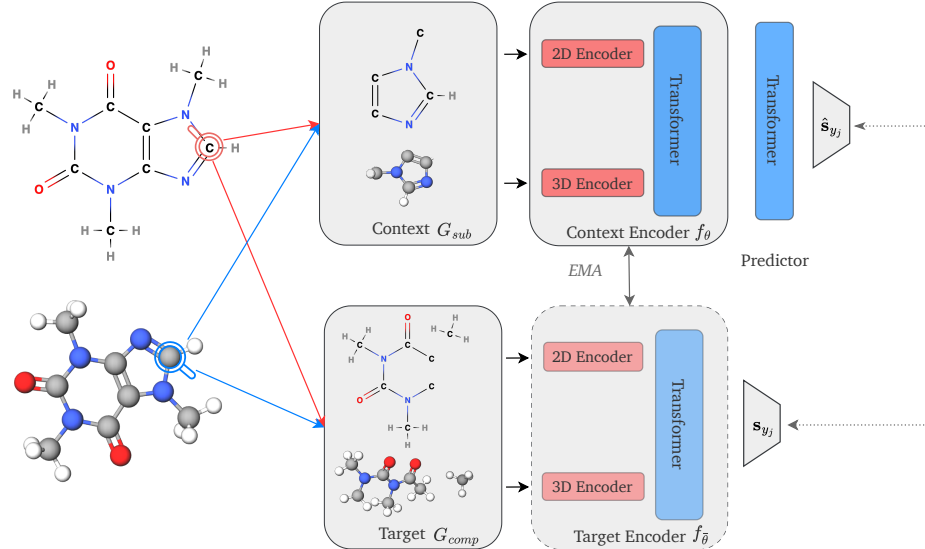


Figure 1: From each molecular graph, we sample a random node and extract its  $k$ -EgoNet (Bevilacqua et al., 2022) with  $k \in \{3, 4\}$  to form complementary context and target subgraphs. Both 2D and 3D views are encoded with a GINE and a SchNet, concatenated, and passed through a transformer; the context embedding is further processed by a predictor to estimate the target. Training minimizes the mean squared  $\mathcal{L}_2$  loss between predicted and encoded targets, with the target encoder updated as an exponential moving average (EMA) of the context encoder (Grill et al., 2020b; Assran et al., 2023). For clarity, only one 3D conformation is shown, though in practice we use three.

To further diversify training, we sample multiple nodes  $v_1, v_2, \dots, v_n$  per molecule and construct their corresponding  $k$ -EgoNets  $E(v_1), E(v_2), \dots, E(v_n)$ , yielding multiple complementary context–target pairs without increasing dataset size.

**Context Encoder.** We aim to learn subgraph representations that generalize to whole-molecule embeddings. Following the architecture proposed in Manolache et al. (2024), we use a message-passing neural network (MPNN) with GINE (Hu et al., 2020a; Xu et al., 2019) as the 2D encoder, and SchNet (Schütt et al., 2017) as the 3D encoder used to process multiple conformers. From GINE, we obtain node-level embeddings  $\{\mathbf{h}_v^{2D}\}$  for all atoms in the subgraph by averaging their intermediate representations across layers. From SchNet, we extract node-level embeddings  $\{\mathbf{h}_{v,c}^{3D}\}$  for each conformer  $c$ , preserving per-atom detail across conformations. To build the multimodal sequence, we prepend a learnable classification token  $\mathbf{h}_{CLS}$  and insert a learnable separation token  $\mathbf{h}_{SEP}$  between the 2D and 3D components, resulting in the following multi-modal sequence:

$$\mathbf{H} = [\mathbf{h}_{CLS}, \mathbf{h}_{SEP}, \{\mathbf{h}_v^{2D}\}, \mathbf{h}_{SEP}, \{\mathbf{h}_{v,c}^{3D}\}, \mathbf{h}_{SEP}]$$

To distinguish between modalities, we add learnable modality embeddings that mark whether a token comes from the 2D or 3D graph. The full sequence is then passed through a Transformer with multiple self-attention layers to capture global dependencies both within and across modalities.

**Predictor Network.** The predictor takes the multimodal embedding of the context subgraph, given by the  $\mathbf{h}_{CLS}^{out}$  embedding from the context encoder, which fuses outputs from the 2D GINE and 3D SchNet. The predictor is a lightweight transformer followed by an MLP, which maps the context embedding to the representation of the complementary subgraph. Since the upstream modality

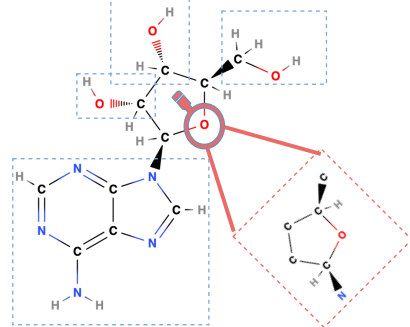


Figure 2: To generate subgraphs, we sample a random node from the original graph (here, the oxygen atom) and extract its 2D ego-net as the context subgraph (outlined by red square). The remaining components (outlined by blue squares) constitute the target subgraph.

encoders already capture spatial and relational information, we do not add explicit positional encodings, unlike image-based JEPA (Assran et al., 2023) and 2D GraphJEPA (Skenderi et al., 2025).

**Target Encoder.** The target subgraph  $f_{\bar{\theta}}$  is encoded by a separate instance of the context encoder. Maintaining two distinct networks stabilizes training and mitigates representation collapse, a strategy widely adopted in self-predictive frameworks such as BYOL (Grill et al., 2020b), I-JEPA (Assran et al., 2023), and BGRL (Thakoor et al., 2023). The target encoder’s weights are updated via an exponential moving average (EMA) of the context encoder’s parameters:

$$\bar{\theta}^{(t)} = \tau \bar{\theta}^{(t-1)} + (1 - \tau) \theta^{(t)},$$

where  $\bar{\theta}^{(t)}$  are the exponentially moving averaged parameters at step  $t$ ,  $\theta^{(t)}$  are the current context encoder parameters, and  $\tau \in [0, 1]$  is the decay rate controlling the contribution of past parameters.

**Pretraining task.** Each subgraph is represented by a single multimodal embedding, taken from the final classification token embedding  $\mathbf{h}_{CLS}^{out}$ . For the *context* subgraph, we feed the entire multimodal token sequence from the context encoder into the predictor transformer and take its output CLS token as the predicted embedding. For the *target* subgraph, we use the CLS token directly from the target encoder. The self-supervised objective minimizes the mean squared  $L_2$  distance between the predicted context embedding and the target embedding:

$$\frac{1}{M} \sum_{i=1}^M \sum_{j=1}^k \|\hat{\mathbf{s}}_{y_j} - \mathbf{s}_{y_j}\|^2,$$

where  $\hat{\mathbf{s}}_{y_j}$  and  $\mathbf{s}_{y_j}$  denote the predicted and target subgraph embeddings,  $M$  is the batch size, and  $k$  the number of sampled views (ego-nets and their complements). All views are treated as separate instances when computing the loss.

**Fine-tuning.** When fine-tuning, we use the target encoder as our pretrained backbone to generate graph embeddings, and add lightweight task-specific heads. We consider two types of task heads: (i) linear probing on whole-graph embeddings with a single linear layer ( $C\text{-FREE}_{LIN}$ ) to evaluate representation quality, and (ii) aggregating  $k$ -EgoNet subgraph embeddings with DeepSets (Zaheer et al., 2017) ( $C\text{-FREE}_{DS}$ ), showing that subgraph pretraining transfers both to whole-molecule prediction and to ESAN-style fine-tuning schemes. We find that  $C\text{-FREE}_{DS}$  is especially beneficial in the 2D-only setting, while for multimodal inputs both heads perform similarly, likely because 3D information compensates for the lower expressiveness. Nevertheless, downstream convergence is faster with DeepSets, suggesting advantages in aligning pretraining and fine-tuning.

### 3.1 INVARIANCE AND EXPRESSIVENESS

We make two theoretical observations. First, with the DeepSets head,  $C\text{-FREE}_{DS}$  simulates ESAN (Bevilacqua et al., 2022) and is more expressive than 1-WL (Weisfeiler & Leman, 1968). An informal statement is given below; the full theorem and proof are in Appendix Section A.1.

**(Informal) Lemma 1.** *Under the assumptions from Theorem 2 of (Bevilacqua et al., 2022),  $C\text{-FREE}$  with a DeepSets task head is as expressive as ESAN, hence it is strictly more expressive than the 1-WL algorithm (Weisfeiler & Leman, 1968).*

Second,  $C\text{-FREE}$  preserves the invariances of its modality encoders. Prior work has shown that architectures of this form inherit invariance from their encoders (Manolache et al., 2024), and the same holds for our framework. For completeness, we include the lemma in Appendix Section A.1.

## 4 EMPIRICAL EVALUATION

We evaluate our framework through four complementary sets of experiments:

- (i) **Frozen backbone evaluation.** We assess representation quality by freezing the backbone and training linear probes (Section 4.1). On MoleculeNet (Wu et al., 2018),  $C\text{-FREE}$  outperforms contrastive and non-contrastive baselines and remains effective even with 2D-only inputs (Table 1). On Kraken (Gensch et al., 2022), pretrained backbones not only converge faster but also achieve lower error than random initialization, with even the 2D-only pretrained model surpassing a randomly initialized multimodal one (Fig. 3).

Table 1: Performance on MoleculeNet (Wu et al., 2018) with frozen backbones. **Non-CL** denotes non-contrastive and **CL** contrastive methods. We report **C-FREE<sub>2D</sub>** (2D-only) and **C-FREE<sub>MM</sub>** (multi-modal), each with linear probing on whole-molecule embeddings (**LIN**) or subgraph aggregation with DeepSets (Zaheer et al., 2017) (**DS**). Metric: ROC-AUC ( $\uparrow$ ). **Red** marks the best model and **Blue** the second best. **C-FREE** ranks first or second on 6 of 8 datasets, with **MM-LIN** best overall, while even the 2D-only variants of C-FREE outperform all baselines on average.

MOLECULENET DATASETS (LINEAR PROBE)									
	BBBP ( $\uparrow$ )	Tox21 ( $\uparrow$ )	TOXCAST ( $\uparrow$ )	SIDER ( $\uparrow$ )	ClINTOX ( $\uparrow$ )	MUV ( $\uparrow$ )	HIV ( $\uparrow$ )	BACE ( $\uparrow$ )	Avg ( $\uparrow$ )
RANDOM INIT.	50.7 $\pm$ 2.5	64.9 $\pm$ 0.5	53.2 $\pm$ 0.3	53.2 $\pm$ 1.1	63.1 $\pm$ 2.3	62.1 $\pm$ 1.3	66.1 $\pm$ 0.7	63.4 $\pm$ 1.8	59.60
CL	INFOGRAPH	65.9 $\pm$ 0.6	65.8 $\pm$ 0.7	54.6 $\pm$ 0.1	57.2 $\pm$ 1.0	61.4 $\pm$ 4.8	63.9 $\pm$ 1.9	71.4 $\pm$ 0.6	67.4 $\pm$ 4.9
	GROVER	67.0 $\pm$ 0.3	63.9 $\pm$ 0.3	53.6 $\pm$ 0.4	59.9 $\pm$ 1.7	65.0 $\pm$ 6.4	62.7 $\pm$ 1.4	67.8 $\pm$ 1.0	69.0 $\pm$ 4.7
	GRAPHCL	64.7 $\pm$ 1.7	69.1 $\pm$ 0.5	56.2 $\pm$ 0.2	59.5 $\pm$ 0.9	60.8 $\pm$ 3.0	60.6 $\pm$ 1.8	72.5 $\pm$ 1.4	77.0 $\pm$ 1.7
	JOAO	66.1 $\pm$ 0.8	68.1 $\pm$ 0.2	55.1 $\pm$ 0.4	58.3 $\pm$ 0.3	65.3 $\pm$ 6.1	62.4 $\pm$ 1.2	73.8 $\pm$ 1.2	71.1 $\pm$ 0.8
NON-CL	EDGE PRED	54.2 $\pm$ 1.0	66.2 $\pm$ 0.2	54.4 $\pm$ 0.1	56.1 $\pm$ 0.1	65.4 $\pm$ 5.0	59.5 $\pm$ 0.9	73.6 $\pm$ 0.4	71.4 $\pm$ 1.2
	ATTRMASK	62.7 $\pm$ 2.7	65.7 $\pm$ 0.8	56.1 $\pm$ 0.2	58.3 $\pm$ 1.5	61.9 $\pm$ 6.4	60.9 $\pm$ 1.8	65.5 $\pm$ 1.4	64.8 $\pm$ 2.6
	GPT-GNN	62.0 $\pm$ 0.9	64.9 $\pm$ 0.7	55.4 $\pm$ 0.2	55.3 $\pm$ 0.8	55.0 $\pm$ 5.1	61.2 $\pm$ 1.5	71.2 $\pm$ 1.5	61.0 $\pm$ 1.2
	CONT. PRED	55.5 $\pm$ 2.0	67.9 $\pm$ 0.7	54.0 $\pm$ 0.3	57.1 $\pm$ 0.5	67.4 $\pm$ 4.3	60.5 $\pm$ 0.9	66.2 $\pm$ 1.5	54.4 $\pm$ 3.2
	<b>C-FREE<sub>2D-LIN</sub></b>	60.5 $\pm$ 1.7	76.1 $\pm$ 0.2	62.7 $\pm$ 0.4	59.0 $\pm$ 0.6	62.7 $\pm$ 1.0	67.6 $\pm$ 0.5	68.7 $\pm$ 0.4	<b>75.8<math>\pm</math>0.9</b>
	<b>C-FREE<sub>2D-DS</sub></b>	64.2 $\pm$ 3.8	<b>76.7<math>\pm</math>0.6</b>	63.9 $\pm$ 0.3	58.0 $\pm$ 0.7	71.4 $\pm$ 3.7	64.6 $\pm$ 3.1	65.5 $\pm$ 0.6	73.9 $\pm$ 0.7
	<b>C-FREE<sub>MM-LIN</sub></b>	<b>69.8<math>\pm</math>2.6</b>	<b>79.9<math>\pm</math>1.1</b>	<b>65.8<math>\pm</math>0.7</b>	58.5 $\pm$ 2.5	<b>69.9<math>\pm</math>1.9</b>	<b>76.6<math>\pm</math>2.8</b>	72.8 $\pm$ 0.7	75.3 $\pm$ 1.1
	<b>C-FREE<sub>MM-DS</sub></b>	<b>73.8<math>\pm</math>2.1</b>	<b>76.7<math>\pm</math>0.7</b>	<b>66.8<math>\pm</math>0.2</b>	56.4 $\pm$ 1.5	<b>75.7<math>\pm</math>2.2</b>	<b>70.6<math>\pm</math>1.0</b>	71.9 $\pm$ 1.5	<b>75.5<math>\pm</math>1.9</b>
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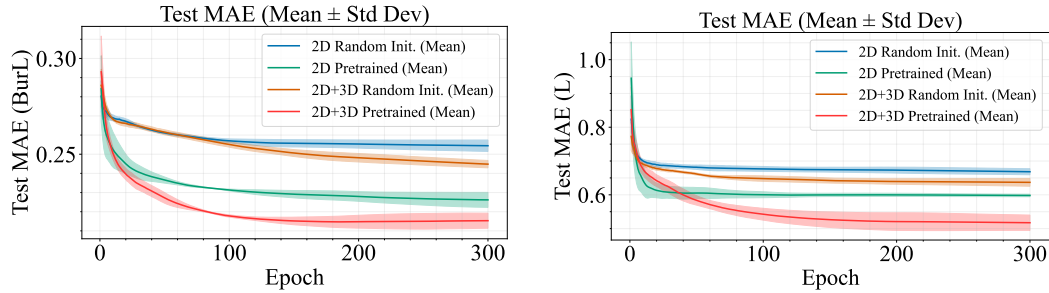


Figure 3: Test MAE on the Kraken regression tasks (Sterimol BurL and Sterimol L) with frozen backbones. GEOM-pretrained models consistently outperform random initialization for both 2D-only and multimodal variants. Pretrained models begin with lower error and converge faster, while randomly initialized models fail to match performance even after 300 epochs. Incorporating the 3D modality yields further gains over 2D-only backbones, with pretraining amplifying this advantage. Curves show the mean over 3 runs, with shaded regions indicating the standard deviation.

- (ii) **Full fine-tuning.** We further evaluate end-to-end adaptability by updating the full backbone with task-specific heads. On MoleculeNet, this setup tests how well pretraining transfers to dataset-specific classification tasks (Section 4.2.1). On Kraken, we find that pretraining improves downstream regression performance over random initialization (Table 3). Finally, on the larger Drugs-75K dataset, we study label efficiency by fine-tuning on progressively larger subsets of labeled data (Section 4.2.2).
- (iii) **Ablations.** We conduct two ablations: (i) removing each modality to assess its contribution (Section 4.2.3), and (ii) removing the predictor network to evaluate its impact on representation quality and training stability (Section 4.3).
- (iv) **Theory alignment.** We verify whether the empirical expressiveness aligns with the theoretical result from Lemma 1 (Section 4.4).

Implementation details for pretraining and evaluation are provided in Section A.4 in the Appendix.

#### 4.1 COMPARISON WITH FROZEN BACKBONES

For the first experiments, we compare our framework against state-of-the-art contrastive and non-contrastive self-supervised methods on molecular property classification tasks. Following Wang et al. (2023a), we pre-train two backbones on 0.33M molecules from GEOM (Axelrod & Gomez-Bombarelli, 2022) and evaluate them on MoleculeNet (Wu et al., 2018). One backbone has 4M parameters and uses 2D inputs, while the other has 9.1M parameters and incorporates both 2D graphs and 3D conformer ensembles available in GEOM, with three additional conformers generated using

Table 2: Performance on MoleculeNet (Wu et al., 2018) with full end-to-end fine-tuning. **Non-CL** denotes non-contrastive, **CL** contrastive, and **Multi** multi-modal methods. Our model is reported as **C-FREE<sub>MM-Full</sub>**, with the multi-modal variant using both modalities. The evaluation metric is ROC-AUC ( $\uparrow$ ). **Red** highlights the best results and **Blue** the second best. **C-FREE** achieves the best results on 4 out of 8 datasets and ranks first overall, outperforming both multi-modal baselines.

		MOLECULENET DATASETS (FULL FINE-TUNING)								
		BBBP ( $\uparrow$ )	Tox21 ( $\uparrow$ )	ToxCast ( $\uparrow$ )	SIDER ( $\uparrow$ )	CLINTOX ( $\uparrow$ )	MUV ( $\uparrow$ )	HIV ( $\uparrow$ )	BACE ( $\uparrow$ )	Avg ( $\uparrow$ )
CL	INFOGRAPH	67.5 $\pm$ 0.1	73.2 $\pm$ 0.4	63.7 $\pm$ 0.5	59.9 $\pm$ 0.3	76.5 $\pm$ 1.0	74.1 $\pm$ 0.7	75.1 $\pm$ 0.9	77.8 $\pm$ 0.8	70.98
	GROVER	70.0 $\pm$ 0.1	74.3 $\pm$ 0.1	65.4 $\pm$ 0.4	<b>64.8<math>\pm</math>0.6</b>	81.2 $\pm$ 3.0	67.3 $\pm$ 1.8	62.5 $\pm$ 0.9	82.6 $\pm$ 0.7	71.01
	GRAPHCL	69.7 $\pm$ 0.6	73.9 $\pm$ 0.6	62.4 $\pm$ 0.5	60.5 $\pm$ 0.8	76.0 $\pm$ 2.6	69.8 $\pm$ 2.6	78.5 $\pm$ 1.2	75.4 $\pm$ 1.4	70.78
	MOLCLR	66.6 $\pm$ 1.8	73.0 $\pm$ 0.1	62.9 $\pm$ 0.3	57.5 $\pm$ 1.7	86.1 $\pm$ 0.9	72.5 $\pm$ 2.3	76.2 $\pm$ 1.5	71.5 $\pm$ 3.1	70.79
	GRAPHLOG	72.5 $\pm$ 0.8	75.7 $\pm$ 0.5	63.5 $\pm$ 0.7	61.2 $\pm$ 1.1	76.7 $\pm$ 3.3	76.0 $\pm$ 1.1	77.8 $\pm$ 0.8	<b>83.5<math>\pm</math>1.2</b>	73.40
NON-CL	ATTRMASK	65.0 $\pm$ 2.3	74.8 $\pm$ 0.2	62.9 $\pm$ 0.1	61.2 $\pm$ 0.1	<b>87.7<math>\pm</math>1.1</b>	73.4 $\pm$ 2.0	76.8 $\pm$ 0.5	79.7 $\pm$ 0.3	72.68
	CONTEXTPRED	65.7 $\pm$ 0.6	74.2 $\pm$ 0.0	62.5 $\pm$ 0.3	62.2 $\pm$ 0.5	77.2 $\pm$ 0.8	75.3 $\pm$ 1.5	77.1 $\pm$ 0.8	76.0 $\pm$ 2.0	71.28
	GRAPHMAE	72.0 $\pm$ 0.6	75.5 $\pm$ 0.6	64.1 $\pm$ 0.3	60.3 $\pm$ 1.1	82.3 $\pm$ 1.2	76.3 $\pm$ 2.4	77.2 $\pm$ 1.0	83.1 $\pm$ 0.9	73.85
	MGSSL	69.7 $\pm$ 0.9	76.5 $\pm$ 0.3	64.1 $\pm$ 0.7	61.8 $\pm$ 0.8	80.7 $\pm$ 2.1	78.7 $\pm$ 1.5	<b>78.8<math>\pm</math>1.2</b>	79.1 $\pm$ 0.9	73.70
	MOLE-BERT	71.9 $\pm$ 1.6	76.8 $\pm$ 0.5	64.3 $\pm$ 0.2	62.8 $\pm$ 1.1	78.9 $\pm$ 3.0	78.6 $\pm$ 1.8	78.2 $\pm$ 0.8	80.8 $\pm$ 1.4	74.04
MULTI	GRAPHMVP	68.5 $\pm$ 0.2	74.5 $\pm$ 0.4	62.7 $\pm$ 0.1	62.3 $\pm$ 1.6	79.0 $\pm$ 2.5	75.0 $\pm$ 1.4	74.8 $\pm$ 1.4	76.8 $\pm$ 1.1	71.69
	3D INFOMAX	69.1 $\pm$ 1.0	74.5 $\pm$ 0.7	64.4 $\pm$ 0.8	60.6 $\pm$ 0.7	79.9 $\pm$ 3.4	74.4 $\pm$ 2.4	76.1 $\pm$ 1.3	79.7 $\pm$ 1.5	72.34
	MOLECULESDE	71.8 $\pm$ 0.7	76.8 $\pm$ 0.3	65.0 $\pm$ 0.2	60.8 $\pm$ 0.3	87.0 $\pm$ 0.5	<b>80.9<math>\pm</math>0.3</b>	<b>78.8<math>\pm</math>0.9</b>	79.5 $\pm$ 2.1	75.07
	MOLEBLEND	<b>73.0<math>\pm</math>0.8</b>	<b>77.8<math>\pm</math>0.8</b>	<b>66.1<math>\pm</math>0.0</b>	<b>64.9<math>\pm</math>0.3</b>	<b>87.6<math>\pm</math>0.7</b>	77.2 $\pm$ 2.3	<b>79.0<math>\pm</math>0.8</b>	<b>83.7<math>\pm</math>1.4</b>	<b>76.16</b>
<b>C-FREE<sub>MM-Full</sub></b>		<b>78.9<math>\pm</math>1.1</b>	<b>84.2<math>\pm</math>0.4</b>	<b>71.7<math>\pm</math>0.9</b>	62.5 $\pm$ 1.9	83.7 $\pm$ 2.9	<b>82.5<math>\pm</math>0.1</b>	77.9 $\pm$ 1.2	78.6 $\pm$ 1.1	<b>77.50</b>

RDKit (Landrum, 2016) at fine-tuning. This setup enables fair comparison to 2D-only baselines while also testing the benefit of 3D information. Performance is reported as mean ROC-AUC over three scaffold splits, with frozen checkpoints chosen by best self-supervised loss and linear probes selected by downstream validation loss. We evaluate two strategies: linear probing on whole-graph embeddings and on subgraph embeddings aggregated with DeepSets (Zaheer et al., 2017).

As shown in Table 1, our framework achieves the best average performance, outperforming baselines on 6 of 8 tasks. In the 2D-only setting, DeepSets aggregation provides clear gains and yields the best average result, with further improvements from adding 3D information. We also see strong gains on multi-task datasets such as Tox21, ToxCast, and MUV, suggesting that our model captures more generalizable features. While DeepSets helps in 2D-only, its effect is minimal in the multimodal case, likely because 3D inputs already encode rich structural detail.

We evaluate transfer to molecular property regression on Kraken (Gensch et al., 2022), which contains 1,552 ligands labeled with four 3D descriptors (Sterimol B5, Sterimol L, buried Sterimol B5, buried Sterimol L). Since Kraken molecules are disjoint from GEOM, it provides a strong test of generalization. As shown in Fig. 3, GEOM-pretrained backbones start with lower error, converge faster, and consistently outperform random initialization, with even the 2D-only pretrained model surpassing a randomly initialized multimodal one. Adding 3D information yields further gains, amplified by pretraining. Results for BurB5 and B5 are included in Appendix Section A.5.

## 4.2 FULL FINE-TUNING

Beyond evaluating frozen representations, we next assess the adaptability of our pretrained models through full end-to-end fine-tuning. In this setting, the entire backbone is updated jointly with a task-specific head, enabling us to test how pretraining improves convergence and downstream performance. We consider two scenarios: (i) property classification on MoleculeNet, where we compare against both contrastive, non-contrastive and multimodal self-supervised baselines, and (ii) property regression on the MARCEL benchmark, where we evaluate transferability to the Kraken dataset and study label efficiency on the larger Drugs-75K dataset.

### 4.2.1 FULL FINE-TUNING FOR PROPERTY PREDICTION

Building on the frozen backbone results, we next evaluate the adaptability of our models through end-to-end fine-tuning. Using the same linear classifier head as in Yu et al. (2023), we update the entire backbone on each MoleculeNet classification dataset. This setting tests how well the pretrained representations adjust to dataset-specific distributions and whether the multimodal backbone provides additional benefits. Performance is reported as mean ROC-AUC over three scaffold splits.

As shown in Table 2, our method achieves notable gains on half of the datasets and the best overall average results. The strongest competitor, MoleBlend (Yu et al., 2023), also uses multimodal inputs but relies on contrastive training. In contrast, our approach matches or surpasses its performance without requiring negative samples.

Table 3: Modality ablation on the Kraken dataset (MAE  $\downarrow$ ). Using the same pretrained backbone, we feed only the 2D sequence for the 2D variant and only the 3D sequence for the 3D variant. Pretraining consistently outperforms training from scratch. The multimodal model delivers the strongest performance, with 3D-only close behind, indicating that 3D features have greater impact than 2D.

	METHOD	B5 $\downarrow$	L $\downarrow$	BURB5 $\downarrow$	BURL $\downarrow$
2D	RANDOM INIT.	0.297 $\pm$ 0.006	0.396 $\pm$ 0.026	0.205 $\pm$ 0.006	0.152 $\pm$ 0.006
	FINE-TUNED	0.276 $\pm$ 0.012	0.340 $\pm$ 0.028	0.176 $\pm$ 0.002	0.146 $\pm$ 0.005
3D	RANDOM INIT.	0.197 $\pm$ 0.006	0.345 $\pm$ 0.011	0.162 $\pm$ 0.006	0.135 $\pm$ 0.009
	FINE-TUNED	0.194 $\pm$ 0.003	0.329 $\pm$ 0.002	0.134 $\pm$ 0.005	0.131 $\pm$ 0.004
MM	RANDOM INIT.	0.203 $\pm$ 0.008	0.378 $\pm$ 0.003	0.161 $\pm$ 0.002	0.142 $\pm$ 0.001
	FINE-TUNED	0.193 $\pm$ 0.017	0.306 $\pm$ 0.011	0.134 $\pm$ 0.009	0.126 $\pm$ 0.004

Table 4: Fine-tuning on the Drugs-75K dataset (Zhu et al., 2024) with limited labeled data. We compare our pretrained backbone (**FFT**) against a model trained from scratch (**RND**) using 1%, 10%, 50%, and 100% of the labels. Pretraining offers clear gains in low-label regimes, while performance is comparable when using the full dataset.

		IP $\downarrow$	EA $\downarrow$	$\chi$ $\downarrow$
1%	RND	0.638 $\pm$ 0.001	0.613 $\pm$ 0.002	0.334 $\pm$ 0.001
	FFT	<b>0.608</b> $\pm$ 0.001	<b>0.583</b> $\pm$ 0.001	<b>0.317</b> $\pm$ 0.001
10%	RND	0.561 $\pm$ 0.002	0.526 $\pm$ 0.001	0.277 $\pm$ 0.001
	FFT	<b>0.520</b> $\pm$ 0.005	<b>0.494</b> $\pm$ 0.002	<b>0.267</b> $\pm$ 0.001
50%	RND	0.457 $\pm$ 0.002	0.433 $\pm$ 0.002	0.233 $\pm$ 0.001
	FFT	<b>0.454</b> $\pm$ 0.001	<b>0.421</b> $\pm$ 0.002	<b>0.230</b> $\pm$ 0.001
100%	RND	<b>0.419</b> $\pm$ 0.005	0.403 $\pm$ 0.002	<b>0.211</b> $\pm$ 0.003
	FFT	<b>0.419</b> $\pm$ 0.002	<b>0.395</b> $\pm$ 0.003	0.213 $\pm$ 0.001

Table 5: Expressiveness results on EXP (Aboud et al., 2021), where 1-WL GNNs cannot surpass random guessing. Even the 1-EgoNet variant of C-FREE approaches the theoretical upper bound, while 2- and 3-EgoNet variants achieve the highest accuracy, outperforming GINE (Hu et al., 2020a) and GraphJEPA (Assran et al., 2023). Unlike GraphJEPA, our method avoids costly METIS clustering (Karypis & Kumar, 1998).

Method	Accuracy ( $\uparrow$ )
GINE	50.69 $\pm$ 1.39
GraphJEPA	98.77 $\pm$ 0.99
C-FREE (1-Ego)	96.03 $\pm$ 1.22
C-FREE (2-Ego)	<b>99.33</b> $\pm$ 0.18
C-FREE (3-Ego)	99.08 $\pm$ 0.20

#### 4.2.2 LABEL-EFFICIENT FINE-TUNING

Building on the observation that fine-tuning outperforms supervised training, we next study label efficiency on the Drugs-75K dataset (Zhu et al., 2024), a GEOM-Drugs subset with 75,099 molecules and at least 5 rotatable bonds. For each molecule, Auto3D (Liu et al., 2022b) generates conformer ensembles, and three DFT-based reactivity descriptors serve as targets: ionization potential (IP), electron affinity (EA), and electronegativity ( $\chi$ ).

We fine-tune our pretrained backbone using 1%, 10%, 50%, and 100% of the available data and compare against a model trained fully supervised from random initialization. With the full dataset, performance is broadly comparable, which is reasonable given the scale of the dataset. More importantly, in low-data regimes, initializing from self-supervised pretraining provides clear gains, consistently outperforming training from scratch (Table 10).

#### 4.2.3 MODALITY ABLATION

After establishing the benefits of our multimodal backbone, we next analyze the contribution of each modality through targeted ablations. Starting from the same GEOM-pretrained backbone, we fine-tune on Kraken while feeding either only the 2D encoder sequence, only the 3D sequence, or both. This setup keeps the architecture and pretraining signal fixed, isolating the effect of each modality and mimicking transfer scenarios where only 2D or 3D data is available. Table 3 shows that pretraining consistently improves over random initialization across all settings, confirming that useful information is transferred even when restricted to a single modality. Among unimodal variants, the 3D-only backbone performs best, suggesting that geometric information has greater impact than 2D topology alone. Combining both modalities achieves the strongest results overall, reinforcing the view that 2D and 3D provide complementary signals.

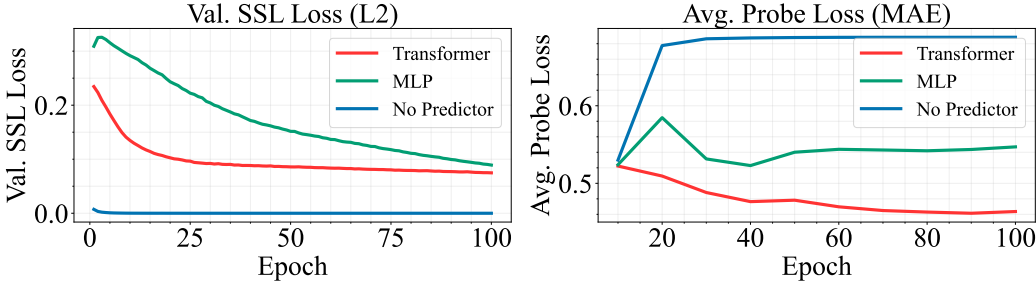


Figure 4: Predictor ablation. **Left:** SSL validation loss on GEOM during pretraining. **Right:** Average linear-probe MAE ( $\downarrow$ ) on Kraken with frozen backbones. Without a predictor, training collapses (loss  $\sim 0$ ) and probes perform worst; an MLP predictor helps but underperforms, while a Transformer predictor achieves the best results.

#### 4.3 ABLATION ON PREDICTOR TYPES

We hypothesize that our model’s strong performance stems from the predictor network, which serves as a guiding signal to refine the representations produced by the encoder. To test this, we pretrain on GEOM following the same setup previously described and perform an ablation with three predictor variants: none, a linear predictor, and a transformer. We then evaluate downstream performance using either a linear probe or full fine-tuning. To keep the study computationally lightweight, we restrict these experiments to the smaller-sized 2D-only pretrained backbone. Since the predictor design is identical in both the 2D-only and multimodal variants, this enables a focused analysis of the predictor’s contribution without incurring excessive computational cost.

As shown in Appendix Table 10, removing the predictor leads to poor downstream performance across regression tasks, with the self-supervised loss collapsing to zero (Fig. 4). While the EMA target encoder stabilizes training (Tervainen & Valpola, 2017; Grill et al., 2020b; Assran et al., 2023), it is insufficient on its own: without an asymmetric architecture, the model collapses to trivial solutions. Adding a predictor breaks this symmetry and prevents collapse (Richemond et al., 2020), consistent with theory showing that a trainable prediction head enables richer representations (Wen & Li, 2022). Even a simple MLP improves results, but the transformer predictor performs best, likely because it operates at the node level before pooling, yielding more informative graph-level embeddings. We therefore adopt it as the default.

#### 4.4 THEORY ALIGNMENT

Finally, we validate our theoretical findings with an experiment on the EXP dataset, designed by Abboud et al. (2021) such that any 1-WL GNN cannot exceed random guessing. We train a smaller version of our encoder and compare it against GINE (Hu et al., 2020a) and GraphJEPAP (Assran et al., 2023). Results are averaged over three runs with resampled EgoNets. As shown in Table 5, even the 1-EgoNet variant of C-FREE approaches the theoretical upper bound, while 2- and 3-EgoNet variants achieve the highest accuracy, outperforming both GINE and GraphJEPAP, the latter relying on the costly METIS (Karypis & Kumar, 1998) algorithm. Nevertheless, the practical significance of expressiveness for molecular learning is limited, as recent work (Pellizzoni et al., 2025) shows that 1-WL already distinguishes most samples in molecular datasets nearly perfectly. Although less decisive for molecules, expressiveness may be more important in other domains, where the flexibility of our framework could be advantageous.

## 5 CONCLUSIONS AND FUTURE WORK

We introduced C-FREE, a contrast-free multimodal self-supervised framework for molecular representation learning. Its core idea is to align embeddings of complementary subgraphs, enabling predictive pretraining without negatives, positional encodings, or costly clustering. By combining 2D molecular graphs and 3D conformer ensembles, C-FREE achieves state-of-the-art results on classification (MoleculeNet) and regression (Kraken, Drugs-75K), with clear gains in low-label regimes. Ablations show that the predictor is critical to avoid collapse and that 2D and 3D provide complementary signals, with 3D offering stronger performance but 2D remaining competitive. Finally, experiments on the EXP benchmark confirm that C-FREE is more expressive than 1-WL.

Looking ahead, several extensions are promising. Our largest model has only 9.1M parameters and was pretrained on 0.33M molecules; scaling to larger architectures and datasets, as the 100M-molecule collections in (Beaini et al., 2024), could unlock further gains. Another extension is combining self-supervised pretraining with an additional supervised stage on such large-scale data before transferring to downstream tasks. Extending to new modalities, such as SMILES strings, is also appealing, though it may require modality-specific masking. Finally, more chemically informed subgraph selection or alternative alignment objectives could further enrich representations.

## 6 REPRODUCIBILITY STATEMENT

We have taken several steps to ensure reproducibility of our work. A full description of the C-FREE framework, including architecture choices, training setup, and pretraining objectives, is provided in Section 3. Implementation details for all experiments, including hyperparameters, optimizers, and fine-tuning protocols, are included in Section A.4. Theoretical claims, such as expressiveness and invariance, are clearly stated in Section 3.1 with proofs in Appendix Section A.1. Datasets used in this work (GEOM, MoleculeNet, Kraken, Drugs-75K, and EXP) are publicly available, and all preprocessing steps are described in Section 4 and Section A.4. Our code and checkpoints are publicly available at <https://github.com/ariguiba/C-FREE>.

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## LLM USAGE STATEMENT

During the course of this work, AI-assisted tools were employed to support specific aspects of the research and writing process:

- **Literature Search:** LLMs were used to identify potentially relevant papers. All references included in this work were thoroughly read, analyzed, and vetted by the authors to ensure their accuracy and relevance.
- **Summarization and Writing Support:** LLMs aided in editing, reformulating, and refining text for clarity, conciseness, and academic style. All AI-generated content was reviewed, adapted, and integrated by the authors to maintain coherence, accuracy, and contextual relevance.

All ideas, critical analyses, interpretations, and conclusions presented in this work are solely those of the authors.

## A SUPPLEMENTARY MATERIALS

### A.1 EXPRESSIVENESS AND INVARIANCE

We preserve the invariance properties of the encoders used in our framework. In particular, we restate the result of Manolache et al. (2024) for our setting, restricted to 2D graphs and 3D conformers:

**Lemma 1.** *Let  $G$  be a 2D molecular graph and  $\{c_1, \dots, c_k\}$  a set of  $k$  3D conformers for the same molecule. Let  $\hat{y} = f_\theta(G, \{c_1, \dots, c_k\})$  be the encoder output as defined in Section 3. Suppose the 3D encoder is invariant to the actions of a group  $\mathcal{G}$ . Then  $f_\theta$  is also invariant to any  $T_1, \dots, T_k \in \mathcal{G}$ , i.e.,*

$$f_\theta(G, \{T_1 c_1, \dots, T_k c_k\}) = f_\theta(G, \{c_1, \dots, c_k\}).$$

The proof of this result is given in Manolache et al. (2024); it applies directly in our case after removing the SMILES modality.

Next, we provide the proof for Lemma 1 in the main paper.

**Lemma 2.** *Let  $C\text{-FREE}_{DS}$  be a model as defined in Section 3 with a subgraph encoder  $f_\theta$  consisting of a 1-WL MPNN (e.g., GIN/GINE) followed by a Transformer without positional encodings, and a DeepSets task head  $DS$ . For any  $k$ -EgoNet policy with  $k \geq 1$  under the assumptions of Theorem 2 from (Bevilacqua et al., 2022),  $C\text{-FREE}_{DS}$  is as expressive as ESAN (Bevilacqua et al., 2022) with an EGO policy, therefore it is as most as expressive as DS-WL and strictly more expressive than 1-WL.*

*Proof.* Fix a  $k$ -EGO policy  $\pi = \text{EGO}_k$  with  $k \geq 1$  and let  $S_\pi(G)$  be the multiset of  $k$ -ego-nets with their complements (edge-covering in the ESAN sense). Define:

$$f_{C\text{-FREE}}(G) = DS(\{f_\theta(S) : S \in S_\pi(G)\}),$$

Due to Lemma 1, we have that  $f_\theta$  maintains the permutation equivariance of the MPNN; moreover, since there exists a parametrization of the Transformer that can approximate the identity map arbitrarily well, the Transformer does not lower the expressive power of the MPNN. We therefore have that  $f_\theta$  is as powerful as 1-WL.

Since we have a DeepSets encoder  $DS$  and an edge-covering  $k$ -EGO policy, we can use the same proof argument as in Theorem 2 from (Bevilacqua et al., 2022), i.e. we apply  $f_\theta$  to each  $S \in S_\pi(G)$  and then aggregate the multisets with  $DS$ , therefore  $f_{C\text{-FREE}}$  simulates ESAN, and is at most as expressive as DS-WL and strictly more expressive than 1-WL.

□

### A.2 BACKBONE PARAMETER EFFICIENCY

Table 6 compares the number of trainable parameters across different SSL backbones. For our method, we report both the total parameters and the encoder-only count used during downstream evaluation. This distinction arises because only the target encoder is retained as the backbone, during downstream tasks, while the predictor is discarded. As a result, nearly half of the parameters are removed at this stage, allowing our method to remain competitive without increasing the parameter load for downstream evaluation, further underscoring its efficiency.

### A.3 COMPUTATIONAL COMPLEXITY ANALYSIS

For generating the subgraphs used as input units in our pre-training scheme, we employ  $k$ -EgoNets with fixed radii  $k \in \{3, 4\}$ . We extract  $k$ -hop neighborhoods using PyTorch Geometric’s (Fey & Lenssen, 2019) `k_hop_subgraph` function, which performs a breadth-first search (BFS) from each node and collects all nodes reachable within  $k$  hops. For constant  $k$ , the BFS cost is bounded by the number of explored edges, yielding a worst-case complexity of  $O(|E|)$ . When repeated for all  $k$  radii, this results in  $O(k \cdot |E|)$ , where  $|E|$  denotes the total number of edges. In practice, the number of explored edges is proportional to the average degree  $d$ , giving a total cost of  $O(k \cdot d \cdot |V|)$ , where  $|V|$  is the number of nodes. Since molecular graphs are sparse, neighborhood growth is modest: an analysis of the GEOM dataset used for pre-training shows that the average degree is only  $d = 2.1$ . As a result,  $k$ -hop neighborhoods remain small, and `k_hop_subgraph` is computationally efficient, scaling linearly with the number of nodes  $O(|V|)$  in practice.

Table 6: Computational efficiency of different SSL methods from Wang et al. (2023a), showing the number of trainable parameters for each backbone. We report both the total parameters of our backbone and those of the encoder alone, since only the latter is used for downstream evaluation. By discarding nearly half of the backbone parameters in this stage, our approach remains competitive without increasing the parameter count for downstream tasks, further highlighting its efficiency.

METHOD	#PARAMETERS (MILLION)
EDGE PRED	7.46
ATTRMASK	7.61
GPT-GNN	7.61
INFOGRAPH	7.82
GROVER	7.57
CONT.PRED	12.00
GRAPHCL	8.19
JOAO	8.19
GRAPHMVP	15.84
<b>C-FREE<sub>2D</sub> (FULL)</b>	8.09
<b>C-FREE<sub>2D</sub> (ENCODER)</b>	4.67
<b>C-FREE<sub>MM</sub> (FULL)</b>	14.65
<b>C-FREE<sub>MM</sub> (ENCODER)</b>	9.12

Table 7: Average runtime (in milliseconds) for generating a single subgraph on the GEOM dataset, comparing METIS partitions with  $n \in \{16, 32\}$  patches and  $k$ -EgoNets with  $k \in \{3, 4\}$ .

	METHOD	AVG. RUNTIME (MS)
METIS	$N = 32$	1.123
	$N = 16$	1.031
K-EGONETS	$K = 3$	0.171
	$K = 4$	0.185

In comparison, the METIS algorithm (Karypis & Kumar, 1998) used in GraphJEPa (Skenderi et al., 2025) employs a multilevel graph partitioning approach. While the algorithm is often cited with an overall complexity of  $O(|E| \cdot \log |V|)$  in practice, it consists of three main phases: (1) a coarsening phase that uses heavy-edge matching to successively reduce the graph size, (2) an initial partitioning phase that partitions the smallest coarsened graph (with negligible complexity due to its small size), and (3) an uncoarsening/refinement phase that projects the partition back to the original graph while refining it at each level. The coarsening and refinement phases dominate the computational cost, each contributing  $O(|E| \cdot \log |V|)$  complexity. However, since molecular graphs are sparse, similar to above, this results in a total complexity of  $O(d \cdot |V| \cdot \log |V|)$ , which is theoretically higher than that of fixed-radius EgoNets.

To further validate this, we ran timed experiments comparing the generation of  $k$ -EgoNet subgraphs with the generation of METIS partitions on the GEOM dataset. Section A.3 reports the average runtime of each method, computed over all graphs in the dataset.

#### A.4 ADDITIONAL DETAILS ON EMPIRICAL EVALUATION

For the multi-modal variant, the context encoder consists of three components: a 6-layer GINE (Xu et al., 2019) with hidden dimension 128, a SchNet with hidden dimension 128, 6 interaction steps, and a cutoff of 10, and 6 Transformer layers with 8 heads and hidden dimension 512. For the pre-trained 2D variant, we use the same GINE configuration and the same number of Transformer layers and heads, but reduce the hidden dimension to 387. In both variants, the predictor is implemented as 4 Transformer layers with 4 heads each. The parameters are updated via backpropagation using the Adam optimizer, while the target encoder is updated through an exponential moving average (EMA) schedule, with the decay rate  $\tau_t$  gradually increasing from 0.995 to 1.0 over the course of training.

Table 8: Overview of tasks and sizes for the MoleculeNet datasets.

	BBBP	Tox21	TOXCAST	SIDER	CLINTOX	MUV	HIV	BACE
# MOLECULES	2,039	7,831	8,575	1,427	1,478	93,087	41,127	1,513
# TASKS	1	12	617	27	2	17	1	1

Table 9: Overview of tasks and sizes for the GEOM, Drugs-75K and Kraken datasets.

	GEOM	DRUGS-75K	KRAKEN
# MOLECULES	304,466	75,099	1,552
# CONFORMERS	25M	558,002	21,287
# TASKS	-	3	4

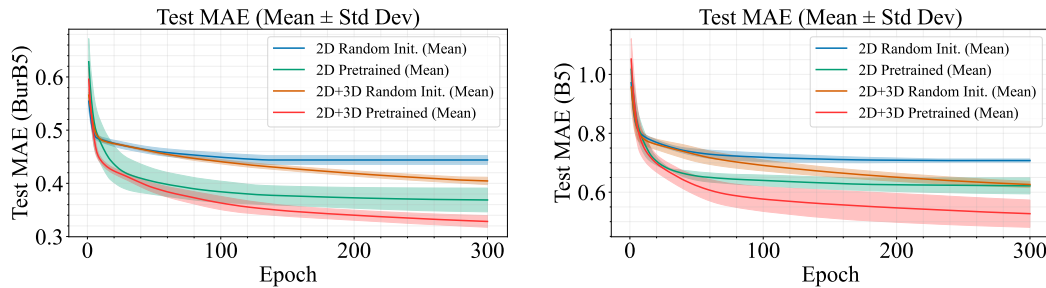


Figure 5: Test MAE on the Kraken regression tasks (Sterimol BurL and Sterimol L) with frozen backbones. GEOM-pretrained models consistently outperform random initialization for both 2D-only and multimodal variants. Pretrained models begin with lower error and converge faster, while randomly initialized models fail to match performance even after 300 epochs. Incorporating the 3D modality yields further gains over 2D-only backbones, with pretraining amplifying this advantage. Curves show the mean over 3 runs, with shaded regions indicating the standard deviation.

Since the EMA decay reaches  $\tau_t = 1$  in the final epoch, the context and target encoders converge to identical parameters. Nevertheless, we follow Assran et al. (2023) and report results using the target encoder.

For the choice of the scheduler we opt for a cosine scheduler without warmup. We notice that using a very small learning rate prevented convergence, while a moderate learning rate caused an early loss drop followed by stagnating representations. Adding a warmup phase allows the model to adapt gradually before the cosine decay, improving stability and representation learning. Thus we begin with a learning rate of  $2 \times 10^{-6}$  over 30 epochs warmup up to  $5 \times 10^{-5}$  and a patience of 50 epochs. For the batch size we use 256 and a weight decay of 0.04 and train for 200 epochs.

All experiments were performed on a mix of Nvidia A100/RTX 4090 GPUs and AMD EPYC 7713/Intel Xeon W-2225 CPUs for both the pre-training and downstream experiments. All experiments consumed a total of approximately 500 GPU hours, with the longest compute being consumed on the pre-training backbone run on the GEOM dataset with a total of 25 hours.

### A.5 ADDITIONAL DETAILS AND RESULTS

In Table 8, we summarize the tasks and dataset sizes of the MoleculeNet benchmarks. Likewise, in Table 9, we present an overview of the GEOM, Drugs-75K, and Kraken datasets, including the corresponding number of conformers.

In Fig. 5 we report the results on the Sterimol B5 and Sterimol BurB5 targets from the Kraken dataset following the experiment in Section 4.1.

In addition to the linear probe, we also report results for full fine-tuning on the Kraken dataset. For this, we use the same setup as in Section 4.1, with an MLP head, and fine-tune the model end-to-end. In Fig. 6, we show results for the 2D-only variant to highlight the initial performance gap, where the pre-trained backbone converges faster than the randomly initialized one.

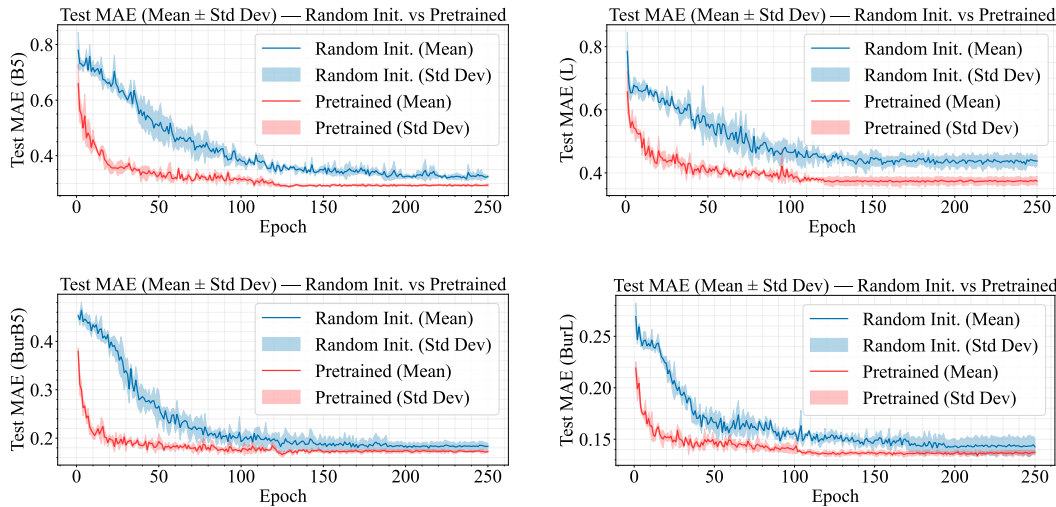


Figure 6: Test MAE on Kraken regression tasks (Sterimol L,B5, BurB5 and BurL) comparing random initialization and 2D-only GEOM-pretrained models. Pretrained models start with lower error and converge faster, while randomly initialized models fail to match their performance even after 250 epochs. Curves show the mean over 3 runs, with shaded regions indicating the standard deviation.

Table 10: Ablation study on the Kraken dataset (MAE  $\downarrow$ ). We keep the encoder fixed and compare three predictors: (1) none, (2) a 2-layer MLP, and (3) a transformer. The transformer consistently achieves the best performance. The gap is especially pronounced in the linear probe (LIN. P.) setting, where the quality of the learned representations matters most. Even with full fine-tuning (FT), the no-predictor and MLP variants fail to match the transformer predictor.

	METHOD	B5 $\downarrow$	L $\downarrow$	BURB5 $\downarrow$	BURL $\downarrow$
FT	NONE	$0.381 \pm 0.023$	$0.494 \pm 0.020$	$0.202 \pm 0.009$	$0.157 \pm 0.004$
	2-LAYERS MLP	$0.315 \pm 0.017$	$0.396 \pm 0.018$	$0.185 \pm 0.009$	$0.144 \pm 0.004$
	TRANSFORMER	$0.292 \pm 0.006$	$0.380 \pm 0.023$	$0.180 \pm 0.014$	$0.146 \pm 0.004$
LIN. P.	NONE	$1.065 \pm 0.001$	$0.814 \pm 0.001$	$0.624 \pm 0.001$	$0.296 \pm 0.001$
	2-LAYERS MLP	$0.817 \pm 0.002$	$0.687 \pm 0.008$	$0.514 \pm 0.001$	$0.266 \pm 0.001$
	TRANSFORMER	$0.588 \pm 0.004$	$0.554 \pm 0.007$	$0.347 \pm 0.003$	$0.202 \pm 0.008$

Finally, in Table 10, we report the explicit numerical results of the probe presented in Section 4.3, evaluated after the full convergence of the model. Additionally, we include the results from end-to-end fine-tuning using the same experimental setup, providing a complete view of how the model performs when the entire backbone is updated. These fine-tuning results further confirm our observation that the predictor transformer consistently outperforms the MLP predictor, while omitting the predictor altogether leads to substantially worse performance, highlighting the importance of the predictor design in our framework.

#### A.6 ADDITIONAL RELATED WORK

**Contrastive Learning** UniCorn (Feng et al., 2024) presents a unified contrastive learning framework that integrates multiple molecular views and existing methods into a single pre-training approach. 3D-Mol (Kuang et al., 2024) leverages 3D conformational information by constructing hierarchical graphs and applying contrastive learning to differentiate molecular conformations. GraphFP (Luong & Singh, 2023) captures higher-level connectivity through fragments—representations of molecular substructures—aligning fragment embeddings with their corresponding graph regions to enable multi-resolution structural learning. MolCLR (Wang et al., 2022) employs three graph augmentations—atom masking, bond deletion, and subgraph removal—and uses contrastive learning to bring augmented views of the same molecule closer, and Galformer (Bai et al., 2023) applies dual-view contrastive learning. Finally, GraphLoG (Xu et al., 2021) captures

both local similarities and global semantic clusters in whole-graph representations using hierarchical prototypes trained via an online EM algorithm.

**Generative Learning** GraphMAE (Hou et al., 2022) adapts the Masked Autoencoder (MAE) from the vision domain to graphs, focusing on reconstructing node attribute features using a scaled cosine error. Similarly, MoleBERT (Xia et al., 2023) extends this idea with a VQ-VAE-based context-aware tokenizer that encodes atom attributes into a larger, chemically meaningful discrete vocabulary, enabling a Masked Atoms Modeling (MAM) task where GNNs predict masked atom codes rather than raw features. Finally, MGSSL (ZHANG et al., 2021) leverages a BRICS-based fragmentation method to extract molecular motifs and pre-trains GNNs to predict motif topology and labels, incorporating multi-level pre-training to capture both local and global graph information.

**Latent Representation Learning** CCA-SSG (Zhang et al., 2021) introduces an alignment objective based on Canonical Correlation Analysis, encouraging the latent features of two augmented views to be maximally correlated while remaining de-correlated across dimensions. Complementing these augmentation-based methods, AFGRL (Lee et al., 2022) proposes a more principled strategy for view generation by identifying structurally and semantically similar anchor nodes, mitigating the reliance on handcrafted augmentations.