ASSEMBLEFLOW: RIGID FLOW MATCHING WITH INER TIAL FRAMES FOR MOLECULAR ASSEMBLY

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Paper under double-blind review

ABSTRACT

Molecular assembly, where a cluster of rigid molecules aggregated into strongly correlated forms, is fundamental to determining the properties of materials. However, traditional numerical methods for simulating this process are computationally expensive, and existing generative models on material generation overlook the rigidity inherent in molecular structures, leading to unwanted distortions and invalid internal structures in molecules. To address this, we introduce AssembleFlow. AssembleFlow leverages inertial frames to establish reference coordinate systems at the molecular level for tracking the orientation and motion of molecules within the cluster. It further decomposes molecular SE(3) transformations into translations in \mathbb{R}^3 and rotations in SO(3), enabling explicit enforcement of both translational and rotational rigidity during each generation step within the flow matching framework. This decomposition also empowers distinct probability paths for each transformation group, effectively allowing for the separate learning of their velocity functions: the former, moving in Euclidean space, uses linear interpolation (LERP), while the latter, evolving in spherical space, employs spherical linear interpolation (SLERP) with a closed-form solution. Empirical validation on the benchmarking data COD-Cluster17 shows that AssembleFlow significantly outperforms six competitive deep learning baselines by at least 45% in assembly matching scores while maintaining 100% molecular integrity. Also, it matches the assembly performance of a widely used domain-specific simulation tool while reducing computational cost by 25-fold.

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1 INTRODUCTION

Deep learning methods have been revolutionizing scientific research across various domains, enabling breakthroughs in fields such as drug discovery (Yu et al., 2024), material science (Merchant et al., 2023), and molecular design (Loeffler et al., 2024). For instance, AlphaFold-like systems have demonstrated unprecedented accuracy and creativity in designing protein structures (Jumper et al., 2021), driving innovation in drug discovery. These advancements underscore the transformative potential of machine learning in tackling complex scientific problems.

Molecular assemble or crystallization is one 040 such complex process where rigid molecules transition from a weakly correlated arrangement 041 to a highly ordered, strongly correlated struc-042 ture. During this process, each molecule is ap-043 proximated to maintain its shape and structure 044 unchanged as it moves in the 3D space, as illustrated in Figure 1. Crystallization plays a 046 pivotal role in determining the physical prop-047 erties of materials, including their mechanical 048 strength, electrical conductivity, and thermal stability (Porter et al., 2009; Carter & Norton, 2013), making it a key process in material 051 science (Ashby & Jones, 2012), pharmaceuticals (Hilfiker, 2006), and nanotechnology (Gon-052



Figure 1: Illustration of the assembly of a cluster of three molecules transitioning from a weakly correlated structure (left) to a strongly correlated crystal structure (right). A key challenge for existing generative models in material generation is preserving the **rigidity** of each molecule throughout this transformation in 3D space, and this paper aims to address this.

osalves et al., 2000). For example, the crystalline form of a drug can affect its solubility and bioavailability (Byrn et al., 1999; Healy et al., 2017); Similarly, precise control over molecular arrangements 068

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Figure 2: (a, b): The molecule's center of mass (CoM) is used as the reference point for **translation** and interpolation in Euclidean space. (c, d): An inertial frame serves as the reference coordinate system, tracking rigid molecular **rotation**—where rotation operations are implemented using quaternion representation—and enabling interpolation in spherical space.

is key to optimizing the electronic and catalytic performance of organic semiconductors, polymers, and molecular catalysts (Saparov & Mitzi, 2016).

Traditional numerical methods have long been employed to simulate the crystallization pro-075 cess (Martínez et al., 2009; Van Der Spoel et al., 2005), but they are often computationally expensive 076 and inefficient, limiting their scalability and practical use in large-scale applications. On the other 077 hand, despite the importance of crystallization, existing machine learning methods struggle to capture the physical constraints critical to this process. A major limitation is the failure to account for the 079 inherent rigidity of molecular structures during crystallization (Liu et al., 2024), often leading to unwanted distortions and invalid internal structures, i.e., non-rigid molecules. In an assembly, molecules 081 must retain their rigid atomic structures, as this rigidity is essential for producing meaningful packing arrangements. However, current generative models for molecular crystallization treat molecules as 083 flexible entities (Liu et al., 2024), resulting in physically unrealistic packing structures and atomic arrangements, failing to retain individual molecule's structure intact. 084

- 085 To address these limitations, we introduce AssembleFlow, a novel framework specifically designed to 086 incorporate the rigid body constraints inherent in molecular assembly or crystallization. As illustrated 087 in Figure 2, AssembleFlow leverages inertial frames to establish reference coordinate systems for 880 assembling molecules. Because SE(3) is the semi-direct product of the rotation group SO(3) and the translation group \mathbb{R}^3 , we can further decompose the group SE(3) transformations into translations 089 in \mathbb{R}^3 (Figure 2 (a, b)) and rotations in SO(3) (Figure 2 (c, d)). Such decomposition allows the 090 explicit enforcement of both translational and rotational rigidity at the molecular level effectively, 091 ensuring that each molecule in the cluster moves as a unified, rigid body throughout the crystallization 092 process. During such enforcement, AssembleFlow employs a distinct approach for learning the 093 SE(3)-equivariant velocity functions associated with translations and rotations. For translations, 094 it uses linear interpolation (LERP) in Euclidean space, while for rotations, it leverages spherical 095 linear interpolation (SLERP) in spherical space, with a closed-form solution. This distinction in 096 handling the translation and rotation groups allows AssembleFlow to accurately model the rigid transformations of molecules during each prediction and generation step within the flow matching 098 framework (Lipman et al., 2022; Liu et al., 2022; Albergo & Vanden-Eijnden, 2022).
- 099 We empirically evaluate AssembleFlow using the benchmarking crystallization dataset COD-100 Cluster17. The quantitative results reveal that AssembleFlow significantly outperforms six competi-101 tive deep learning baselines by at least 45% in terms of assembly matching score. Also, AssembleFlow 102 exhibits strong assembly performance compared to a widely used domain-specific simulation tool 103 for molecular assembly, achieving this with a 25-fold reduction in computational cost. Furthermore, 104 we present qualitative results, including DFT energy and atomic collision properties of predicted 105 crystals, which further demonstrate AssembleFlow's effectiveness in preserving and modeling the rigidity of the molecular crystallization and assembly process. Our work is the first to implement 106 rigid generation in SE(3) space for molecular assembly. We also want to mention that in what follows, 107 we use molecular assembly, crystallization, and molecular packing interchangeably.

108 2 PRELIMINARIES

Molecular crystallization. Molecular crystallization is a transition of molecules from weakly
 correlated structures to strongly correlated structures, *e.g.*, from liquid or gas phase to solid phase,
 as illustrated in Figure 1. A common example is liquid water freezing into ice, transitioning from
 a liquid phase to a solid phase. The crystallization from a gas phase directly to a solid phase is called
 deposition.

SE(3)-equivariance. For geometric modeling for crystallization, one critical property of the target function is rotation-equivariant and translation-equivariant (*i.e.*, SE(3)-equivariant). We here provide a brief introduction on the SE(3)-equivariance, and for more detailed discussions of SE(3)-equivariance, we refer the reader to (Smidt et al., 2018; Brandstetter et al., 2021; Liu et al., 2023; Zhang et al., 2023). SE(3)-equivariance is the property for the geometric modeling function $f : X \to Y$ as:

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 $f(\rho_X(\boldsymbol{a})\boldsymbol{x}) = \rho_Y(\boldsymbol{a})f(\boldsymbol{x}), \quad \forall \boldsymbol{a} \in G, \boldsymbol{x} \in X,$ (1)

where $\rho_X(a)$ and $\rho_Y(a)$ are the SE(3) group representations on the input and output space, respectively. SE(3)-equivariant modeling in Equation (1) is essentially saying that the designed deep learning model *f* is modeling the whole SE(3) group transformation trajectory on the molecule conformations, and the output is the transformed \hat{y} accordingly. One concrete example is that when we rotate the input molecular system by a certain angle, the predicted forces by SE(3)-equivariant models will also rotate accordingly.

Conditional flow matching. Conditional flow matching (CFM) (Lipman et al., 2022) and two parallel works (Rectified Flow (Liu et al., 2022) and Stochastic Interpolants (Albergo & Vanden-Eijnden, 2022)) formulate the distribution modeling problem as learning a vector field that can generate a probability path mapping from simple distribution at t = 0 to the target distribution at t = 1. Please refer to the original papers for a more detailed discussion (Lipman et al., 2022).

In the crystallization processes, our geometric data are atomic coordinates in the 3D Euclidean points $r \in \mathbb{R}^3$, and the **atomic type is fixed** during the whole crystallization process, so we may as well ignore that. Then we define time-dependent vector field $v : [0,1] \times \mathbb{R}^3 \to \mathbb{R}^3$. A time-dependent vector field defines a time-dependent diffeomorphic map, called flow, $\phi : [0,1] \times \mathbb{R}^3 \to \mathbb{R}^3$. The vector field defines flow via an ordinary differential equation as

$$l\phi_t(\mathbf{r})/dt = u_t(\phi_t(\mathbf{r})). \tag{2}$$

A probability density path is denoted as $p : [0,1] \times \mathbb{R}^3 \to \mathbb{R}_{>0}$. Existing flow model (Chen et al., 2018) maps a prior distribution p_0 to another distribution p_t with push-forward equation or change of variable rule: $p_t(\mathbf{r}) = [\phi_t]_* p_0(\mathbf{r}) = p_0(\phi_t^{-1}(\mathbf{r})) \det \left| \frac{d\phi_t(\mathbf{r})}{dx} \right|^{-1}$. Thus, modeling the likelihood of data distribution at t = 1 can be transformed into modeling the velocity field matching problem with parameterized velocity field v_{θ} , *i.e.*, flow matching:

$$\mathcal{L}_{\text{FM}} = \mathbb{E}_{t,\boldsymbol{r}} \| u_t(\boldsymbol{r}) - v_\theta(\boldsymbol{r},t) \|^2.$$
(3)

With the continuity equation (Villani et al., 2009), we can further derive an equivalent objective by considering the conditional vector field conditioned on the empirical data r_1 , *i.e.*, $u_t(r|r_1)$, and the resulting objective is the conditional flow matching:

$$\mathcal{L}_{\text{CFM}} = \mathbb{E}_{t,\boldsymbol{r},\boldsymbol{r}_1} \left\| u_t(\boldsymbol{r}|\boldsymbol{r}_1) - v_{\theta}(\boldsymbol{r},\boldsymbol{r}_1,t) \right\|^2.$$
(4)

152 3 METHOD: ASSEMBLEFLOW

154**Problem Formulation.** AssembleFlow is designed to model rigid transformations during crystalliza-155tion, ensuring that each molecule in the cluster remains rigid throughout the transformation process.156Rigorously, we are modeling $P(\{r_f\} | \{r_i\})$, where $\{r_f\}$ and $\{r_i\}$ are the atom conformations in157final and initial positions respectively. During this process, we assume the rigidity of molecules.158Noticeably, we will use a preprocessed dataset where the prior conformations are geometrically159optimized and fixed (Liu et al., 2024).

This section outlines the five key steps in the algorithm's development. Specifically, in Section 3.1, we
 explain how inertial frames can be leveraged to provide a stable reference for tacking the orientation of multiple assembling molecules in the Euclidean space. Such a reference perspective can guarantee

162 rigid structures during molecular rotations throughout the assembly process. Building on these 163 frames, there are multiple ways for rotation representation, so in Section 3.2, we illustrate how to use 164 quaternion representation for capturing the rotation transformation induced from inertial frames. This 165 is followed by a detailed discussion of AssembleFlow, a rigid flow matching method, in Section 3.3. 166 AssembleFlow decomposes the assembly probabilistic paths into SO(3) group path and \mathbb{R}^3 group path to guarantee the rigidity, and learns the time-dependent vector fields through a flow-matching 167 framework on the two path spaces respectively. In Section 3.4, we employ the reparameterization trick 168 to make AssembleFlow more numerically stable. Finally, in Section 3.5, we present the two types of SE(3)-equivariant flow matching velocity functions specifically designed for use in AssembleFlow. 170 Note: the pseudo algorithm of our AssembleFlow is provided in Appendix E.3. 171

- 172 3.1 SO(3) GROUP AND INERTIAL FRAME FOR RIGID PACKING
- The core of AssembleFlow lies in the utilization of the inertial frame as the reference frame. Within this frame, the rotation matrix in the SO(3) group defines how the molecular system rotates rigidly. This serves as the key step in AssembleFlow for modeling rigid transformations in SO(3) group.

SO(3) **group.** The special orthogonal group, denoted as SO(*n*), is a group of rotation matrices that represent rotations in *n*-dimensional Euclidean space. In this paper, we are interested in n = 3dimensional space, and every rotation matrix used to perform a rotation in 3D space can be represented as an element of SO(3) group. The SO(3) group consists of all orthogonal matrices (with determinant 1) $R \in \mathbb{R}^{3 \times 3}$ such that $R^T R = I$, where R^T is the transpose of R and I is the identity matrix.

Inertial frame as the reference frame. An inertial frame is a reference frame such that it can provide a consistent basis for describing a molecule's motion, including rotation. We here utilize the inertial frame to build up a basis to describe how each molecule rotates in the Euclidean space. One example is illustrated in Figure 2. Importantly, an inertial frame provides a coordinate system such that a molecule stays rigid and does not deform over the crystallization or modeling process; we here assume the system is not influenced by external forces. Next, we will detail how inertial frames are used to represent the rotation matrix for rigid molecules.

- First, we employ the following four sequential steps to derive the reference frames that construct the rotation matrix from N atomic positions r:
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- Calculate the mass center: $c = \frac{1}{N} \sum_{i} r_{i}$.
- Adjust position relative to the center $r_i = r_i c$.
- Compute the inertia tensor $\hat{I} = \sum_i ||\mathbf{r}_i||^2 I \mathbf{r}_i \mathbf{r}_i^T$, where I is the unit diagonal matrix.
- Obtain the principal axes of inertia by applying eigen-decomposition on \hat{I} . We have $\hat{I} = Q\Lambda Q^T$, where Q is the orthogonal matrix whose columns are the eigenvectors of \hat{I} , and Λ is the diagonal matrix whose elements are the eigenvalues λ_i of \hat{I} , representing the principal moments of inertial along the principal axes.

The above steps yield the three principal 199 axes in Q. To adopt this for modeling the 200 crystallization process (Figure 1), we build 201 inertial frames for each molecule in the 202 cluster. In our case, for each molecule we 203 need to build two inertial frames, for the 204 weakly correlated and strongly correlated 205 structures, respectively. We call these two 206 frames initial (inertial) frame \mathcal{F}_i and final 207 (inertial) frame \mathcal{F}_f .

208 Second, we apply the eigen-decomposition 209 to obtain the **initial principal axes** Q_i and 210 **final principal axes** Q_f , respectively. As 211 illustrated in Figure 2 (d), we can only per-212 form the rotation based on the aligned prin-



Figure 3: (a, b) show two potential rotational alignments between two coordinate systems (axes). (c, d) show that only one unique rotation is possible for four non-coplanar points.

cipal axes or the aligned coordinate systems. However, as we conduct the eigen-decomposition of frames \mathcal{F}_i and \mathcal{F}_f , it is not guaranteed that the corresponding principal axes Q_i and Q_f are aligned. To align the two coordinate systems, we aim to match both the directions and the orders of the corresponding principal axes in each set. We will detail this alignment process next. 216 Align the directions between initial and final coordinate systems. First, for a given inertial frame 217 \mathcal{F} , we have three axes in Q composing a coordinate system. This can lead to eight possible directions. 218 To align the directions between the initial system Q_i and final system Q_f , we add a first constraint 219 - the three axes must form a right-handed coordinate system. Such a filtering step can be achieved 220 by using cross-product: if the local frame system is right-handed, then the cross-product between any two axes should match the third axis or share the same direction as the third axis. Otherwise, the 221 coordinate system is left-handed, then we randomly revert one basis out of three. This reduces to 222 four potential combinations of directions. To further align the directions, we introduce Lemma 1 and 223 Theorem 1 to help provide us with theoretical guidance. 224

225 **Lemma 1.** For an initial inertial frame \mathcal{F}_i and a final inertial frame \mathcal{F}_f , we build up the correspond-226 ing right-handed principal axes as coordinate systems, Q_i and Q_f , respectively. Suppose we have to 227 change the directions of Q_f to match Q_i , then we should change the directions of two bases in Q_f .

228 *Proof.* There are three bases in Q_f . If we change one or three basis directions in Q_f , then Q_f will 229 change from right-handedness to left-handedness, which violates the assumption. Thus, if we need to 230 change the directions of Q_f to match Q_i , we should change the directions of two bases in Q_f .

231 With Lemma 1, we find that using three base vectors is insufficient to determine the direction 232 alignment of three axes. To define the directions that can match between the initial and final 233 coordinate systems, we need to incorporate an extra node as an auxiliary, as in Theorem 1. 234

Theorem 1. For an initial inertial frame \mathcal{F}_i and a final inertial frame \mathcal{F}_f , we build up the corre-235 sponding right-handed principal axes as coordinate systems, Q_i and Q_f , respectively. Then we need 236 to incorporate a fourth point that is not coplanar with the three basis vectors, to align the directions 237 of two coordinate systems with one unique rotation transformation matrix. 238

239 *Proof Sketch.* We first provide intuitive examples in Figure 3. In Figure 3 (a, b), we can see at least two possible rotation matrices to transform from initial axes to final axes. However, when we add 240 a fourth non-coplanar point in Figure 3 (c), the rotation transformation becomes unique, and the 241 corresponding rotation in Figure 3 (d) is invalid. Then more rigorously, the proof includes the two key 242 steps: (1) Using Lemma 1, we can find multiple rotation matrices for alignment between coordinate 243 systems. (2) After introducing the fourth non-coplanar point, the contradiction proves that there exists 244 only one unique rotation for alignment. For more rigorous proof, please refer to Appendix D. 245

246 Align the ordering between initial and final coordinate systems. We can typically sort the 247 eigenvectors (as for principal axes) through the corresponding eigenvalues. The main challenge comes when there is a tie in eigenvalues. Because this is a rare case, we propose doing a depth-first-248 search to enumerate all the possible combinations of basis orderings of Q_f , to match Q_i . 249

250 Outputs and engineering issue: numerical stability. Without loss of generality, we can assume that 251 we do not change the axis direction or ordering in the initial coordinate system Q_i , and we only change 252 Q_f to \hat{Q}_f , so as to align with Q_i . The ultimate rotation matrix is thus $R = Q_i^T \hat{Q}_f$. Meanwhile, we 253 would like to point out that multiple numerical stability issues exist. This can arise in the following 254 scenarios: (1) when the sampled points are near the origin, (2) when checking if the eigenvalues are 255 tied or not, (3) when extracting a fourth non-planar point for alignment, (4) when verifying whether rotating the initial atoms (points) matches the final atoms. To mitigate these issues, we carefully 256 select a threshold value and clamp the (reconstructed) coordinates to this minimum threshold. 257

258 **Summary.** In this section, we first introduce the basic concepts of SO(3) group and inertial frame. 259 Then we present how we construct the initial and final inertial frames for each molecule, *i.e.*, \mathcal{F}_i 260 and \mathcal{F}_f , in molecular crystallization. Next, by applying the eigen-decomposition on the constructed inertial frames, we obtain the initial and final principal axes (right-handed), Q_i and Q_f , respectively. 261 262 Finally, we align the Q_f to Q_i by checking the directions and ordering of three axes in Q_f . This 263 results in two aligned bases (Q_i and \hat{Q}_f) and a rotation matrix R such that $\hat{Q}_f = R Q_i$. As a result, 264 this enables rigid molecule-level rotations during their transformations in the assembling processes.

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3.2 **OUATERNION REPRESENTATION FOR ROTATION**

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For the SO(3) group in Section 3.1, there are multiple ways to represent a rotation transformation 268 in addition to the rotation matrix R. If we want to model the rotation matrix directly, we must 269 guarantee that the generated matrix variable satisfies the two properties discussed in Section 3.1, namely Orthogonality and Determinant. Such a constrained modeling is challenging. Thus, an alternative way of rotation representation with a more flexible formulation is preferred. To attain this goal, we utilize quaternion representation defined through the inertial frame, as described below.
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Definition. A quaternion q is defined as:

$$\boldsymbol{q} = \boldsymbol{w} + \boldsymbol{x}\boldsymbol{i} + \boldsymbol{y}\boldsymbol{j} + \boldsymbol{z}\boldsymbol{k} = (\boldsymbol{w}, \boldsymbol{v}),\tag{5}$$

where w, x, y, z are real numbers, and i, j, k are the fundamental quaternion units. Equivalently, the w is called the real part, while v = (x, y, z) is a 3D vector representing the imaginary part.

Rotation quaternion. A rotation quaternion is a unitary quaternion, *i.e.*, $w^2 + x^2 + y^2 + z^2 = 1$, and this can be easily achieved by taking the normalization of the quaternion variables. In what follows, we will assume the quaternion is a rotation quaternion unless otherwise specified. Notice that for each rotation matrix, there are two equivalent quaternions, q and -q. Here we manually enforce the real number part of the generated quaternion to be non-negative.

Tranformation from rotation matrix to rotation quaternion. There are multiple ways to extract the quaternion from the rotation matrix, and we provide a more detailed discussion in Appendix B. In this work, we adopt eigendecomposition (Horn, 1987; Bar-Itzhack, 2000) to extract the initial and final quaternion (*i.e.*, q_i and q_f) from the initial and final coordinate systems (*i.e.*, Q_i and Q_f).

287 288 Spherical interpolation (SLERP) for quaternion interpolation. One of the main advantages 289 of using quaternion is that it is friendly to interpolation on the SO(3) space, *i.e.*, the spherical 290 interpolation (SLERP) between two quaternions q_0 and q_1 :

$$SLERP(\boldsymbol{q}_0, \boldsymbol{q}_1, t) = \frac{\sin((1-t)\omega)\boldsymbol{q}_0 + \sin(t\omega)\boldsymbol{q}_1}{\sin(\omega)},$$
(6)

where ω is the angle between q_0 and q_1 , and $t \in [0, 1]$ is interpolation parameter. Thus, we can see that SLERP provides a smooth and uniform rotation between two quaternions. An example is provided in Figure 2.

We provide a comprehensive discussion of various rotation representations in Appendix B, including quaternion multiplication and vector rotation using quaternion. Please consult that section for details.

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3.3 PATH INTERPOLATION IN ASSEMBLEFLOW

To model the crystallization process, our AssembleFlow method integrates the inertial frames and quaternion representation for rotation, as discussed in Sections 3.1 and 3.2, into a conditional flow matching framework. We note that unlike most existing flow matching methods that focus solely on atom-level diffusion paths in the Euclidean space, which suffices for non-rigid transformation, AssembleFlow operates within the full SE(3) group space in the molecule level due to the rigidity requirement.

Recall that the crystallization process involves the movement over a cluster of molecules, and for each molecule in the cluster, AssembleFlow jointly models the rotational transformations in SO(3) space and translational transformations in \mathbb{R}^3 space. Such a decomposition ensures the preservation of rigid molecular structures throughout the crystallization process. We next detail these two transformations.

Modeling translations in \mathbb{R}^3 . The goal is to model the molecule-level translations in each cluster, and AssembleFlow achieves this by modeling the translations on each molecule's mass center, as depicted in Figure 2. For notation simplicity, we will use $x \in \mathbb{R}^3$ to represent the translation vector.

We adopt the flow-matching framework, and the goal here is to learn the probability of final mass center x_f from the initial mass center x_i , *i.e.*, $p(x_f|x_i)$. To this end, we assume that we use linear interpolation (LERP) for path interpolation, by treating $x_0 = x_i$ and $x_1 = x_f$, then for interpolation parameter $t \in [0, 1]$, we have the interpolated translation as:

$$\operatorname{ERP}(\boldsymbol{x}_0, \boldsymbol{x}_1, t) = t\boldsymbol{x}_0 + (1 - t)\boldsymbol{x}_1.$$
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Next, we introduce an SE(3)-equivariant function $v_{\theta,\mathbb{R}^3}(x_t,t)$ as the core module to learn the velocity at time t. Thus the objective function is defined as:

$$\mathcal{L}_{\mathbb{R}^3} = \|\boldsymbol{x}_1 - \boldsymbol{x}_0 - \boldsymbol{v}_{\theta,\mathbb{R}^3}(\boldsymbol{x}_t, \boldsymbol{q}_t, t)\|^2.$$
(8)

Modeling rotations in SO(3). For modeling the rotations in the SO(3) group, recall that we can find the initial bases Q_i and final principal bases Q_f from inertial frames in Section 3.1, and then we

transform them into the rotation quaternions q_i and q_f as introduced in Section 3.2. The task here is to model $p(q_f|q_i)$.

Thus, it is natural to adopt the spherical interpolation (SLERP) as the smooth translation between two quaternions. We treat $q_0 = q_i$ and $q_1 = q_f$ and plug them into Equation (6). This gives us the interpolated rotations at time t. The first-order derivative of SLERP has an analytical formula:

$$\frac{d}{dt} \text{SLERP}(\boldsymbol{q}_0, \boldsymbol{q}_1, t) = \frac{\omega \Big(\cos(t\omega)\boldsymbol{q}_1 - \cos((1-t)\omega)\boldsymbol{q}_0\Big)}{\sin(\omega)}.$$
(9)

Similarly here, we then introduce an SE(3)-equivariant function $v_{\theta,SO(3)}(q_t, t)$ to model the moleculelevel rotation velocity at time t. The objective function becomes:

$$\mathcal{L}_{\text{SO}(3)} = \|\frac{d}{dt} \text{SLERP}(\boldsymbol{q}_0, \boldsymbol{q}_1, t) - \boldsymbol{v}_{\theta, \text{SO}(3)}(\boldsymbol{x}_t, \boldsymbol{q}_t, t)\|^2.$$
(10)

Inference. For inference, AssembleFlow conducts the sampling step in SO(3) and \mathbb{R}^3 alternatively:

$$\boldsymbol{x}_{t+1} = \boldsymbol{x}_t + \delta t \cdot \boldsymbol{v}_{\theta,\mathbb{R}^3}(\boldsymbol{x}_t, \boldsymbol{q}_t, t), \qquad \boldsymbol{q}_{t+1} = \boldsymbol{q}_t + \delta t \cdot \boldsymbol{v}_{\theta,\mathrm{SO}(3)}(\boldsymbol{x}_t, \boldsymbol{q}_t, t).$$
(11)

Thus, both the molecule-level translation x_{t+1} and molecule-level rotation q_{t+1} are applied on each molecule, and we repeat Equation (11) for T steps to obtain the predicted strongly correlated molecule position. However, in Equation (11), it remains an open question on how to obtain the ω for SO(3) generation, since ω is the angle between q_0 and q_1 , and q_1 is unknown during the inference process. We address this by proposing the reparameterization trick as will be discussed next in Section 3.4.

3.4 REPARAMETERIZATION FOR STRONGLY CORRELATED STRUCTURES

We here leverage the reparameterization trick to directly model the SE(3) action at time T instead of the velocity at each time t. Equivalently, the velocity of SE(3) action can be written as:

$$\boldsymbol{v}_{\theta,\mathbb{R}^3}(\boldsymbol{x}_t, \boldsymbol{q}_t, t) = (\hat{\boldsymbol{x}}_{1,\theta}(\boldsymbol{x}_t, \boldsymbol{q}_t, t) - \boldsymbol{x}_t)/(1-t),$$

$$\boldsymbol{v}_{\theta,\mathrm{SO}(3)}(\boldsymbol{x}_t, \boldsymbol{q}_t, t) = \frac{\omega \Big(\cos(t\omega)\hat{\boldsymbol{q}}_{1,\theta}(\boldsymbol{x}_t, \boldsymbol{q}_t, t) - \cos((1-t)\omega)\boldsymbol{q}_0\Big)}{\sin(\omega)}.$$
 (12)

In other words, we directly estimate the translation $\hat{x}_{1,\theta}(x_t, q_t, t)$ and rotation $\hat{q}_{1,\theta}(x_t, q_t, t)$ in the final step or the strongly correlated structure. The objectives over the two spaces thus become:

$$\mathcal{L}_{\mathbb{R}^{3},\text{reparameter}} = \mathbb{E}[\|\boldsymbol{x}_{1} - \hat{\boldsymbol{x}}_{1,\theta}(\boldsymbol{x}_{t},\boldsymbol{q}_{t},t)\|^{2}], \qquad \mathcal{L}_{\text{SO}(3),\text{reparameter}} = \mathbb{E}[\|\boldsymbol{q}_{1} - \hat{\boldsymbol{q}}_{1,\theta}(\boldsymbol{x}_{t},\boldsymbol{q}_{t},t)\|^{2}].$$
(13)

The final objective function is the summation of two terms. Besides, such a reparameterization enables us to conduct inference using the Euler algorithm:

$$\boldsymbol{x}_{t+1} = \boldsymbol{x}_t + \delta_t \cdot (\hat{\boldsymbol{x}}_{1,\theta}(\boldsymbol{x}_t, \boldsymbol{q}_t, t) - \boldsymbol{x}_t)/(1-t),$$

$$\boldsymbol{q}_{t+1} = \boldsymbol{q}_t + \delta_t \cdot \frac{\hat{\omega} \Big(\cos(t\hat{\omega}) \hat{\boldsymbol{q}}_{1,\theta}(\boldsymbol{x}_t, \boldsymbol{q}_t, t) - \cos((1-t)\hat{\omega}) \boldsymbol{q}_0 \Big)}{\sin(\hat{\omega})},$$

(14)

where $\hat{\omega}$ is the angle between q_0 and $\hat{q}_{1,\theta}(x_t, q_t, t)$. The velocity functions of $\hat{x}_{1,\theta}(x_t, q_t, t)$ and $\hat{q}_{1,\theta}(x_t, q_t, t)$ are SE(3)-equivariant and will be discussed next in Section 3.5.

368 3.5 SE(3)-EQUIVARIANT MULTI-GRAINED VELOCITY FUNCTION

Recall that the data structure considered here is the cluster of molecules, thus it is natural to split the modeling into intra-molecule and inter-molecule modeling, as introduced below. For intra-molecule modeling, we adopt the PaiNN (Schütt et al., 2021), which is one of the most widely used SE(3)-equivariant models. It can encode the inherent geometric structural information of individual molecules. Then for inter-molecule modeling, we consider two options of SE(3)-equivariant models: (1) Atomic-level modeling that utilizes all the atoms' positions for learning the molecular-level rotation and translation for the next step. (2) Molecular-level modeling that directly utilizes the molecular-level rotation and translation for next-step prediction. This concludes our discussion on AssembleFlow, and more details are provided in Appendix E. A high-level overview and pseudo algorithm are provided in Algorithms 1 and 2 in Appendix E.3.

	Packing	Matching		Validity	
	PM (atom) \downarrow	PM (center) ↓	Collision ↓	Separation ↑	Compactness ↑
		Dataset: COD-Cl	uster17-5K		
GNN-MD	13.67 ± 0.06	13.80 ± 0.07	27.53 ± 0.49	0.22 ± 0.11	$\textbf{100.00} \pm \textbf{0.00}$
CrystalSDE-VE	15.52 ± 1.48	16.46 ± 0.99	1.20 ± 0.08	27.17 ± 0.86	57.47 ± 7.76
CrystalSDE-VP	18.15 ± 3.02	19.15 ± 4.46	0.84 ± 0.14	53.13 ± 12.89	34.00 ± 30.75
CrystalFlow-VE	14.87 ± 7.07	13.08 ± 4.51	1.37 ± 0.04	35.70 ± 0.73	8.40 ± 4.17
CrystalFlow-VP	15.71 ± 2.69	17.10 ± 1.89	1.38 ± 0.04	35.43 ± 0.88	4.87 ± 1.09
CrystalFlow-LERP	13.59 ± 0.09	13.26 ± 0.09	0.34 ± 0.01	97.38 ± 0.10	$\textbf{100.00} \pm \textbf{0.00}$
AssembleFlow (ours)	$\textbf{7.27} \pm \textbf{0.04}$	$\textbf{6.13} \pm \textbf{0.10}$	$\textbf{0.33} \pm \textbf{0.00}$	$\textbf{97.64} \pm \textbf{0.36}$	$\textbf{100.00} \pm \textbf{0.00}$
		Dataset: COD-Clu	ister17-10K		
GNN-MD	13.83 ± 0.06	13.90 ± 0.05	27.88 ± 0.49	0.23 ± 0.11	$\textbf{100.00} \pm \textbf{0.00}$
CrystalSDE-VE	17.25 ± 2.46	17.86 ± 1.11	0.99 ± 0.27	32.99 ± 10.72	34.93 ± 14.99
CrystalSDE-VP	22.20 ± 3.29	21.39 ± 1.50	0.53 ± 0.35	52.48 ± 15.44	16.83 ± 18.09
CrystalFlow-VE	16.41 ± 2.64	16.71 ± 2.35	1.42 ± 0.03	33.79 ± 0.51	5.47 ± 0.47
CrystalFlow-VP	19.39 ± 4.37	16.01 ± 3.13	1.44 ± 0.03	33.35 ± 0.55	4.23 ± 0.48
CrystalFlow-LERP	13.54 ± 0.03	13.20 ± 0.03	0.32 ± 0.00	97.32 ± 0.05	$\textbf{100.00} \pm \textbf{0.00}$
AssembleFlow (ours)	$\textbf{7.38} \pm \textbf{0.03}$	$\textbf{6.21} \pm \textbf{0.05}$	$\textbf{0.31} \pm \textbf{0.00}$	$\textbf{97.73} \pm \textbf{0.16}$	99.93 ± 0.05
		Dataset: COD-Cl	uster17-All		
GNN-MD	22.30 ± 12.04	14.51 ± 0.82	24.29 ± 4.58	4.13 ± 5.60	98.77 ± 1.73
CrystalSDE-VE	17.28 ± 0.73	18.92 ± 0.03	0.19 ± 0.18	15.47 ± 12.42	2.51 ± 2.37
CrystalSDE-VP	18.03 ± 4.56	20.02 ± 3.70	0.55 ± 0.19	48.78 ± 1.70	6.88 ± 2.82
CrystalFlow-VE	12.80 ± 1.20	15.09 ± 0.34	1.41 ± 0.01	35.34 ± 0.28	2.90 ± 0.02
CrystalFlow-VP	13.50 ± 0.44	13.28 ± 0.48	1.51 ± 0.02	33.06 ± 1.31	6.61 ± 3.17
CrystalFlow-LERP	13.61 ± 0.00	13.28 ± 0.01	0.34 ± 0.00	97.34 ± 0.02	$\textbf{99.99} \pm \textbf{0.01}$
AssembleFlow (ours)	$\textbf{7.37} \pm \textbf{0.01}$	$\textbf{6.21} \pm \textbf{0.01}$	$\textbf{0.31} \pm \textbf{0.00}$	$\textbf{98.15} \pm \textbf{0.22}$	99.98 ± 0.00

Table 1: AssembleFlow against six generative models on COD-Cluster17 with 5K, 10K, and all samples. The
 best results are marked in **bold**.

4 EXPERIMENTS

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4.1 EXPERIMENT SETUP

Datasets. We evaluate our method using the crystallization dataset COD-Cluster17 (Liu et al., 2024).
This COD-Cluster17 contains 133K crystals and is a curated subset derived from the Crystallography
Open Database (COD) database (Grazulis et al., 2009). We consider three versions of COD-Cluster17,
with 5k, 10k, and all data, respectively. Detailed discussion on this dataset is provided in Appendix G.

409 Evaluation metrics. We evaluate the per-410 formance of the compared approaches us-411 ing a comprehensive set of metrics tailored 412 to assess the quality of crystallization pack-413 ing. These metrics include: (1) Packing 414 Matching (PM) (Chisholm & Motherwell, 415 2005): This metric measures how well the generated molecular assemblies match the 416 reference crystal structures in terms of spa-417 tial arrangement and packing density. Fol-418 lowing (Liu et al., 2024), we employ pack-419



Figure 4: AssembleFlow achieved marginally lower results in formation energy.

ing matching on both the atomic level (PM-atom) and the mass-center-level (PM-center) (Chisholm 420 & Motherwell, 2005). (2) Atomic Collision: This follows (Cordero et al., 2008). It measures the 421 percentage of collided atom pairs in the predicted assemblies. Atoms must maintain a minimum 422 covalent distance governed by the balance of attractive and repulsive forces. (3) Separation: We 423 extend the metric from (Xie et al., 2022; Yang et al., 2024) to our setting. A cluster of molecules is 424 valid if the minimum distance between molecules is above 0.5Å (Court et al., 2020). This metric is 425 referred to as *separation* to measure the validity to avoid unphysical interactions at the molecular 426 level. (4) Compactness: We propose this measure by calculating the percentage of simulated clusters 427 where the maximum atomic pairwise distances are below 100Å. A higher compactness value suggests 428 a more efficient arrangement, where the intermolecular spaces are minimized, leading to a denser 429 crystalline structure. (5) DFT Energy: This metric evaluates the stability of crystal structures by calculating their total electronic energy using Density Functional Theory (DFT) (Kohn & Sham, 430 1965; Van Der Spoel et al., 2005). Lower DFT energy generally suggests a more stable molecular 431 configuration. Detailed discussions on these metrics are provided in Appendix G.



Figure 5: Predicted molecular assembly of two randomly picked crystals from the test set (top and bottom subfigures), each consisting of 17 molecules. The leftmost shows the initial structure, the rightmost represents the ground truth, and the intermediate steps display predictions at the 20^{th} , 40^{th} , and 50^{th} iteration, respectively.

Baselines. We compare our method with two categories of baselines: state-of-the-art deep generative
 models and an established domain-specific simulation tool.

452 (1) Deep generative baselines. For generative models, we evaluate our approach against GNN-453 MD (Liu et al., 2024), CrystalSDE (Liu et al., 2024), CrystalFlow (Liu et al., 2024), and different 454 variations of them, including CrystalSDE-VE, CrystalSDE-VP, CrystalFlow-VE, CrystalFlow-VP, 455 and CrystalFlow-LERP. These models employ various mechanisms to handle the challenges of 456 molecular crystallization. CrystalSDE-VE and CrystalSDE-VP use stochastic differential equations 457 to model diffusion processes under different parameterizations. CrystalFlow-VE and CrystalFlow-VP apply flow matching principles for diffusion-based interpolation path, with the latter focusing on 458 variance-preserving methods. CrystalFlow-LERP utilizes linear interpolation to handle molecular 459 transformations, striking a balance between computational complexity and performance. 460

(2) Domain-specific simulation baseline. We also compare our method with PackMol (Martínez et al., 2009), a well-established simulation tool widely used in the field for molecular packing. PackMol has long been a go-to solution for chemistry and material experts due to its ability to generate initial molecular configurations for follow-up simulations, making it an important and relevant baseline for evaluating molecular assembly tasks. More detail on this baseline is in Appendix F.

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The comparison results
with the generative modeling baselines and the simulation model are presented in Tables 1 and 2, respectively.

474 As shown in Table 1, 475 AssembleFlow significantly 476 outperformed all six deep 477 generative models across al-478 most all metrics. For ex-479 ample, AssembleFlow im-480 proved Packing Matching 481 by at least 45% compared 482 to other models. Notably, 483 most baselines struggled with rigid packing, leading 484

Table 2: Ablation studies of PackMol and AssembleFlow variants.							
	PackMol	AssembleFlow-Atom	AssembleFlow-Molecule				
	Data	aset: COD-Cluster17-5K					
PM (atom)↓	7.10 ± 0.05	7.27 ± 0.04	7.67 ± 0.10				
PM (center) \downarrow	6.05 ± 0.04	6.13 ± 0.10	6.77 ± 0.10				
Collision \downarrow	0.32 ± 0.00	0.33 ± 0.00	0.37 ± 0.01				
Separation \uparrow	99.56 ± 0.08	97.64 ± 0.36	92.95 ± 0.16				
Dataset: COD-Cluster17-10K							
PM (atom) \downarrow	7.16 ± 0.01	7.38 ± 0.03	7.65 ± 0.17				
PM (center) \downarrow	6.11 ± 0.01	6.21 ± 0.05	6.69 ± 0.20				
Collision \downarrow	0.30 ± 0.00	0.31 ± 0.00	0.35 ± 0.01				
Separation \uparrow	99.45 ± 0.10	97.73 ± 0.16	92.67 ± 0.32				
Dataset: COD-Cluster17-All							
PM (atom) \downarrow	7.15 ± 0.01	7.37 ± 0.01	7.40 ± 0.11				
PM (center) \downarrow	6.09 ± 0.01	6.21 ± 0.01	6.28 ± 0.13				
Collision \downarrow	0.30 ± 0.00	0.31 ± 0.00	0.33 ± 0.00				
Separation ↑	99.42 ± 0.03	98.15 ± 0.22	95.60 ± 0.11				

to very low Separation scores, except for CrystalFlow-LERP. For example, AssembleFlow achieved a Separation rate of 97.64%, while GNN-MD only reached 0.22%.

4.2 MAIN RESULTS

486 As shown in Table 2, when compared to the domain-specific tool PackMol, our data-driven 487 approach demonstrates strong assembly performance relative to this widely used simulation method. 488 Remarkably, our method achieves 100% validity, matching that of the domain-specific simulation 489 tool. While this outcome highlights the promise of AssembleFlow, it is expected for PackMol, as 490 it leverages well-established domain knowledge and heuristic physical rules to determine molecular orientations. Promisingly, both methods achieved a very low Collision rate. Furthermore, Figure 4 491 shows the total DFT energy distributions of crystals predicted by AssembleFlow and calculated 492 by PackMol. The results indicate that the data-driven AssembleFlow performs comparably to the 493 domain simulation tool PackMol. 494

4.3 Ablation Studies

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497 Velocity function. As discussed in Section 3.5, 498 AssembleFlow can utilize two types of SE(3)-499 equivariant velocity functions. We here evaluate 500 their performance and, as shown in the last two 501 columns of Table 2, AssembleFlow-Atom gen-502 erally outperforms AssembleFlow-Molecule, as 503 atomic-level information provides more detailed 504 geometric insights. Despite this, both variations of AssembleFlow exhibit strong results, signifi-505 cantly surpassing other deep learning baselines. 506

507 Atomic collision. To show atomic collisions of
508 the assemblies, we visualize, in Figure 6, two
509 atomic collisions in assemblies predicted by As510 sembleFlow, where two molecules collide into
511 each other (indicated by the red circle).

512 Computational cost. In Figure 7, we present
513 the computation time needed for Assemble514 Flow and PackMol. Figure 7 reveals that our
515 data-driven method achieves a 25-fold reduc516 tion in computational costs. This suggests that
517 our method can be scaled effectively for larger
518 molecular systems and datasets.

- 519 **Predicted trajectory.** In Figure 5, we visual-
- 520 ize two flow trajectories predicted by Assemble-



Figure 6: Atomic collisions (red circles) in predicted assemblies.



Figure 7: Comparison of computational cost in hours for 10,543 molecule clusters from COD-Cluster17. Pack-Mol requires around 864 hours, while AssembleFlow requires 35 hours.

Flow, showing the processes of how AssembleFlow transformed two clusters of molecules from weakly corrected structures (left) to strongly corrected structures (right); The leftmost structure is the input, while the rightmost represents the ground truth.

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5 CONCLUSION AND OUTLOOK

We introduced AssembleFlow, a generative model that maintains the inherent rigidity of molecular
 structures during assembly. By using inertial frames for positional references at molecular level,
 AssembleFlow accurately tracks molecular orientation and motion. It decomposes transformations
 into translations and rotations, enforcing rigidity throughout the generation process. This innova tive approach enables the model to separately learn velocity functions using linear and spherical
 interpolation for accurate rigid molecular assembly.

Empirical results on COD-Cluster17 show that AssembleFlow outperforms six state-of-the-art
 deep learning models while maintaining molecular integrity and achieves comparable assembly
 performance to an established simulation tool, significantly reducing computational costs.

To the best of our knowledge, AssembleFlow is the first to implement rigid generation in SE(3) space
 for molecular assembly. It has the potential to be generalized to more complex and challenging
 scenarios, such as simulating the crystallization process of polymorphs with diverse configurations
 and structures. We aim to explore these challenges in future work.

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A RELATED WORKS

758 A.1 GEOMETRIC MODELING

Geometric modeling of molecules has predominantly been applied in 3D Euclidean space (Smidt et al., 2018). It requires that the representation and generation function over the molecular system remain equivariant to the rotations and translations, *i.e.*, the SE(3)-equivariance, ensuring that molecular properties are preserved regardless of orientation or position in space (Smidt et al., 2018; Zhang et al., 2023; Liu et al., 2023).

We note that reflection (or chirality) is also an important factor in geometric molecule modeling. For
multi-component molecular systems like protein-ligand binding complexes, each individual molecule
can lead to different energies and corresponding properties with distinct chiralities (Brooks et al.,
2011; McVicker & O'Boyle, 2024). Also, as shown in AlphaFold2, the natural molecules should be
sensitive to the chirality (Jumper et al., 2021). Thus, more physically accurate geometric modeling
should be SE(3)-equivariant and reflection-antisymmetric, and we have shown how our proposed
AssembleFlow satisfies this in Section 3.

772 We also want to highlight another line of research that downplays the importance of symmetry 773 in molecular modeling, as demonstrated by models like AlphaFold3 (Abramson et al., 2024) and 774 XYZTransformer (Flam-Shepherd & Aspuru-Guzik, 2023). These models avoid incorporating geometric symmetries because enforcing group symmetry constraints, such as SE(3)-equivariance, 775 776 can limit a model's expressiveness. In some domain-specific tasks, relaxing these constraints has resulted in strong performance. However, in the case of crystallization, maintaining molecular 777 rigidity—a key symmetric property—is crucial. As demonstrated in previous work (Liu et al., 778 2024), neglecting these equivariance constraints during molecular crystallization leads to unrealistic 779 molecular structures.

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A.2 GENERATIVE MODELS ON GEOMETRIC DATA

The geometric modeling on the continuous 3D Euclidean space can be naturally merged with deep 784 generative frameworks, where the goal is to learn the geometric data distribution and generate new 785 molecules. The deep generative models include but are not limited to denoising score matching (Vin-786 cent, 2011; Song & Ermon, 2019), denoising diffusion probabilistic model (Ho et al., 2020), and flow 787 matching (Lipman et al., 2022; Liu et al., 2022; Albergo & Vanden-Eijnden, 2022). Such geometric 788 generative models have been widely adopted for molecule and material generation (Hoogeboom 789 et al., 2022; Jiao et al., 2024; Luo et al., 2024), protein design and protein folding (Zhang et al., 790 2024; Campbell et al., 2024), structure-based drug design (Guan et al., 2023; 2022), and molecular 791 dynamics simulation (Doerr et al., 2021; Arts et al., 2023; Fu et al., 2022).

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A.3 GENERATIVE MODELS ON RIGID GEOMETRIC DATA

Though our work is the first to apply rigid generation in SE(3) space for molecular pack-796 ing/crystallization, similar ideas have been carried out for protein backbone generation (Jumper 797 et al., 2021; Yim et al., 2023a;b; Bose et al., 2023; Huguet et al., 2024) and relevant works (Köhler 798 et al., 2023; Ganea et al., 2021; Klein et al., 2024)) In the protein backbone generation and folding 799 setting, there exists a well-defined local frame structure for each residue: the backbone atom pairs 800 (C-N) and $(C_{\alpha}-C)$ form two bases, and their cross product leads to the third basis, which is a normal vector perpendicular to the two bases. Thus, the rotation matrix can be easily constructed 801 based on such a local frame structure. Such a modeling paradigm is appealing for macromolecules 802 like proteins to reduce the computational cost. 803

In this field, AlphaFold2 adopts this local-frame idea to model the folding process (Jumper et al., 2021), while FrameDiff applies this idea and denoising diffusion model for protein structure generation (Yim et al., 2023b). Similarly, FrameFlow and FoldFlow integrate local frames with flow matching to learn the protein dynamics (Yim et al., 2023a; Bose et al., 2023; Huguet et al., 2024). However, this approach cannot be easily extended to crystallization tasks, as constructing reliable local frames to establish positional references for assembling molecules is not straightforward. Instead, we innovatively introduce inertial frames to achieve this goal.

We have added related works above: (Köhler et al., 2023; Ganea et al., 2021; Klein et al., 2024; Yim et al., 2023a).

A.4 Comparison Between Spherical Interpolation and Exponential Map Interpolation on SO(3)

In addition to spherical interpolation, we could also consider the exponential map interpolation, as
used in (Riemannian FM, FrameFlow, and FoldFlow). In this section, we would like to compare
the theoretical differences. We further conduct experiments for empirical comparison, and please
check Appendix H.2 for more details.

⁸²⁰ We mark exponential map interpolation as (EMLERP), and it is defined as:

$$\mathrm{EMLERP}(\boldsymbol{r}_0, \boldsymbol{r}_1, t) = \exp_{\boldsymbol{r}_0}(t \log_{\boldsymbol{r}_0}(\boldsymbol{r}_1)). \tag{15}$$

We then discuss how FoldFlow and FoldFlow2 utilized this equation since they are the latest work
 along this line of using EMLERP. By utilizing the tangent space of the manifold and axis-angle
 representation, existing works (FoldFlow, FoldFlow2) have been using an approximated closed-form
 solution for the derivative:

$$\frac{d}{dt} \text{EMLERP}(\boldsymbol{r}_0, \boldsymbol{r}_1, t) = \log_{\boldsymbol{r}_t} \frac{\boldsymbol{r}_0}{t}.$$
(16)

829 Thus, their objective function on SO(3) is:

$$\mathcal{L}_{SO(3)} = \|\frac{d}{dt} \text{EMLERP}(\boldsymbol{r}_0, \boldsymbol{r}_1, t) - \boldsymbol{v}_{\theta, SO(3)}(\boldsymbol{x}_t, \boldsymbol{r}_t, t)\|^2$$

= $\|\log_{\boldsymbol{r}_t} \frac{\boldsymbol{r}_0}{t} - \boldsymbol{v}_{\theta, SO(3)}(\boldsymbol{x}_t, \boldsymbol{r}_t, t)\|^2.$ (17)

To sum up, these two methods do not have a clear methodological advantage over each other; however, the EMLERP considers more approximation tricks in implementation. We summarize the main differences in Table 3.

Table 3: C	Comparison	between t	the inter	polation	paths in	AssembleFlow	and	FrameFlow
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	AssembleFlow (ours)	FoldFlow
Reference Frame	Eigenvectors of inertial frames	Gram-Schmidt on N - C - C_{α}
SO(3) Interpolation	Spherical Interpolation	Exponential Map Interpolation
Equation	$\frac{\sin((1-t)\omega)\boldsymbol{q}_0 + \sin(t\omega)\boldsymbol{q}_1}{\sin(\omega)}$	$\exp_{\boldsymbol{r}_0}(t\log_{\boldsymbol{r}_0}(\boldsymbol{r}_1))$
Derivative	Equation (9)	Equation (16)
Rotation Representation (for Velocity / Objective Function)	Quaternion	Rotation Matrix or Axis-angle
Reparameterization	Yes	No

A.5 COMPARISON BETWEEN OUR WORK AND RIGID STRUCTURE SAMPLING (KÖHLER ET AL., 2023)

We would like to emphasize that our work differs from (Köhler et al., 2023) in the following fundamental aspects. Noticeably, we would like to emphasize that our work can be seen as an extension of (Köhler et al., 2023), yet addressing more practical and challenging problems, including rigid modeling on arbitrary molecules, SE(3)-equivariant modeling, and interpolation modeling.

- 1. **Experiment and Data Difference**: (Köhler et al., 2023) targets at modeling the transition, e.g., water molecules at different temperatures (no code or comments related to methane rigid modeling were found in the GitHub repository). The MD simulation can provide samples at the stable or equilibrium status. Our work models the transition from unstable to stable conformations.
- 2. Dynamic Transition Modeling vs. Stable Structure Sampling: Unlike (Köhler et al., 2023) which focuses on stable structure sampling, our work models the dynamic transition process from weakly correlated (unstable) structures to strongly correlated (stable) structures. Notably, dynamic transition modeling toward stability is identified as a nontrivial next step

in the ICML work (Köhler et al., 2023), which states '... a flow model for the positions that can handle ... phase transitions'. We also want to emphasize that the transition of a molecular cluster from weakly correlated (unstable) structures to strongly correlated (stable) structures is a special case of the general dynamics. The limitation here is the data insufficiency (lack of intermediate snapshots), not modeling.

- 3. Objective Difference: (Köhler et al., 2023) first introduces molecular equilibrium sampling with Boltzmann distribution in Eq 1,2. But this is not the goal, (Köhler et al., 2023) changes to estimating log-ratio: $\nabla F = -\log(Z_{\alpha_1}/Z_{\alpha_0})$ between two configurations in Eq 3. This measure estimates the energy difference and tells which state is more stable. (Köhler et al., 2023) says using trackable priors like Gaussian can be biased, thus it takes the insight from previous work, and targets solving the problems Eq 3 and 7 (learned free energy perturbation). (Notice that other equations Eq 2,4,5 are preliminaries, not directly related to the core method in this work.) This reveals another theoretical difference between this work and our work: (Köhler et al., 2023) is estimating the upper bound of log-ratio between two stable states (with MD simulations), and ours is directly modeling $p(X_1|X_0)$ from unstable to stable transition.
 - 4. Use of Inertial Frames for Rigid Modeling: (Köhler et al., 2023) specifically models the frame for H-O-H (codes here) (no code or comments related to methane rigid modeling were found in the GitHub repository). In contrast, our approach is more generalizable, as the inertial frame can serve as a reference frame for any molecule. Thus, (Köhler et al., 2023) cannot be directly applied to our dataset, as it is limited to few types of constructed frames, and our work can be viewed as an extension of (Köhler et al., 2023) to a more general setting.
 - 5. **SE(3)-equivariant Symmetry Modeling**: (Köhler et al., 2023) states that it has a limitation on not 'exploiting the SE(3) symmetry of jointly moving all rigid bodies'. We solve this issue by introducing the SE(3)-equivariant modeling from two granularities.
 - Limitations of Coupling Layers in Normalizing Flow: (Köhler et al., 2023) does modeling with an extra bijectivity constraint in coupling layers, limiting the model capacity (Ho et al., 2019). Flow matching enables flexible velocity functions under the interpolation framework.
 - We list the comparison in the table below:

Table 4: Comparison between (Köhler et al., 2023) and AssembleFlow.

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	Paper (Köhler et al., 2023)	AssembleFlow (ours)
Experiments	Transition between two stable conformations	Transition from unstable to stable
Data	Water and methane (experiment missing on GitHub repo) molecules	COD organic molecules
Frame construction	H-O-H frame specific for water molecule (No code or comments related to methane rigid modeling were found in the GitHub repository)	Inertial frame for any organic molecule
Data	Water molecules	COD organic molecules
Objective Function	Upper bound of log-ratio between two stable con- formations for energy difference	Direct estimation of conditional density from un- stable to stable
Modeling SE(3) symmetry in moving rigid bodies	No	Yes
Avoid bijectivity constraint in coupling layers	No	Yes

В **ROTATION REPRESENTATION**

In this section, we will be mainly discussing three types of rotation representations. It is important to note that "representation" here refers to the data structure commonly used in the machine learning community, rather than the concept of a representation space.

In Appendix C.1, we will explain how to use inertials to represent the rigid structures of molecules in a cluster. With such a rigid representation, we can then decompose SE(3) transformation into a tuple of SO(3) and \mathbb{R}^3 transformations.

A natural way to represent the SO(3) transformation is by using the rotation matrix, as will be introduced in Appendix B.1. However the rotation matrix is not flexible and it must satisfy specific mathematical properties to make sure it is a valid rotation in space, so we need a more flexible and efficient representation. To this end, we would like to introduce axis-angle representation in Appendix B.2 and quaternion representation in Appendix B.3. The axis-angle representation and quaternion representation are closely related, and their transformation is discussed in Appendix B.4. Last but not least, the transformation between the rotation matrix and quaternion is in Appendix B.5, and the transformation between axis-angle representation and rotation matrix representation is in Appendix B.6. An overview of the rotation representation and the corresponding transformations are listed in Figure 8.



Figure 8: Transformation between rotation matrix, quaternion, and axis-angle representation.

ROTATION REPRESENTATION WITH ROTATION MATRIX B.1

Definition Rotation matrix is $R \in \mathbb{R}^{3 \times 3}$, satisfying two conditions:

The set of all orthogonal matrices of size 3 with determinant 1 is a representation of a group known as the special orthogonal group SO(3). To generate a rotation matrix in SO(3), certain properties for the rotation matrix must be satisfied, and such constrained generation is challenging. Thus, an alternative way of representing the rotation matrix is preferred. To this end, we consider axis-angle representation, as described below.

Rotation with Rotation Matrix To rotation a point, p = (x, y, z), in the 3D Euclidean space, the rotation operation is:

Properties

1. Orthogonality: The rows and columns of R are orthonormal vectors. $R^T R = R R^T = I$. 2. Determinant: The determinant of R must be 1. det(R) = 1.

972 • If we have multiple rotation matrices, and we want to yield a single matrix that combines 973 these rotations into one, then we have two options: 974 - Extrinsic rotations. All rotations refer to a fixed and global coordinate system, and the 975 rotation matrices are ordered from the right to the left. If we apply rotations R_1, R_2 , 976 and R_3 , then it is written as $R = R_3 R_2 R_1$. - Intrinsic rotations. A rotation refers to the last rotated coordinate system, and the 977 rotation matrices are ordered from the left to the right. If we apply rotations R_1, R_2 , 978 and R_3 , then it is written as $R = R_1 R_2 R_3$. 979 980 981 **ROTATION REPRESENTATION WITH AXIS-ANGLE REPRESENTATION B**.2 982 The axis-angle represents the rotation by its angle θ and the rotation axis u. For example, a rotation 983 of 180 degrees around the Y-Axis would be represented as $\theta = 180$, $\boldsymbol{u} = (0, 1, 0)$. The representation 984 is very intuitive, but for actually applying the rotation, another representation is required, such as a 985 quaternion or rotation matrix. 986 987 **Definition** The axis-angle representation of a rotation is defined by two components: 988 989 1. Rotation axis: a unit vector $\boldsymbol{u} = (u_x, u_y, u_z)$ that specifies the direction of rotation, $\|\boldsymbol{u}\| =$ 990 1. 991 2. A scalar θ specifies the angle of rotation around the rotation axis. 992 For annotation, an axis-angle rotation can thus be presented by four numbers as $(\theta, \hat{x}, \hat{y}, \hat{z})$. 993 994 **B.3** ROTATION REPRESENTATION WITH QUATERNION REPRESENTATION 995 996 Quaternion represents a rotation by a 4D vector and it is a more concise representation than a rotation 997 matrix. It requires more math and is less intuitive, but is a much more powerful representation. 998 Ouaternion representation has been widely used in rigid motion modeling, robotics modeling, and 999 quantum mechanics (*e.g.*, the spin of an electron and the polarization of a photon). In this work, we 1000 are focusing on the first case, rigid motion modeling. 1001 1002 **Definition** A quanternion q is: 1003 $\boldsymbol{q} = \boldsymbol{w} + x\boldsymbol{i} + y\boldsymbol{j} + z\boldsymbol{k} = (\boldsymbol{w}, \boldsymbol{v}),$ (19)1004 1005 where w, x, y, z are real numbers, and i, j, k are the fundamental quaternion units. The w is the real part, and v = (x, y, z) is a 3D vector representing the imaginary part. 1007 1008 **Multipliation of basis elements** The multiplication for the basis elements i, j, k is defined below: 1009 1010 $i^2 = i^2 = k^2 = -1$ 1011 ij = -ji = k1012 jk = -kj = i(20)1013 ki = -ik = j1014 ijk = -1.1015 1016 1017 Quanternion multiplication: Hamilton product This can give us the quaternion multiplication, 1018 a.k.a., Hamilton product. For two quaternions $\mathbf{r} = (r_0, r_1, r_2, r_3)$ and $\mathbf{s} = (s_0, s_1, s_2, s_3)$: 1019 t = rs, (21)1020 1021 where 1022 $t_0 = r_0 s_0 - r_1 s_1 - r_2 s_2 - r_3 s_3$ 1023 $t_1 = r_0 s_1 + r_1 s_0 - r_2 s_3 + r_3 s_2$

- 1024 1025 $t_1 = r_0 s_1 + r_1 s_0 - r_2 s_3 + r_3 s_2$ $t_2 = r_0 s_2 + r_1 s_3 + r_2 s_0 - r_3 s_1$
 - $t_3 = r_0 s_3 r_1 s_2 + r_2 s_1 + r_3 s_0.$

(22)

Point rotation with quaternion We rotate a point $v = (v_x, v_y, v_z)$ by the quaternion q = (w, x, y, z) using the following three steps:

- 1. Transform \boldsymbol{v} to quaternion $\boldsymbol{p} = (0, v_x, v_y, v_z)$.
 - 2. Construct the conjugate quaternion $q^* = (w, -x, -y, -z)$.
- 3. There are two types of rotation operations:

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- (a) Active rotation: $p' = q^* pq$, when the point is rotated w.r.t. the coordinate system.
- (b) Passive rotation: $p' = qpq^*$, when the coordinate system is rotated w.r.t. the point.

Notice that the two rotations are opposite from each other. In our case, we use the passive rotation.

4. The resulting vector v' is extracted from the imaginary part of p'.

Spherical Interpolation (SLPER) for Quaternion Interpolation Quaternions are often pre ferred for interpolating between rotations because they offer smoother interpolation than axis-angle
 representation. The spherical interpolation defines the geodesic over the rotation group.

$$\text{LERP}(\boldsymbol{q}_0, \boldsymbol{q}_1, t) = \frac{\sin((1-t)\omega)\boldsymbol{q}_0 + \sin(t\omega)\boldsymbol{q}_1}{\sin(\omega)},$$
(23)

where ω is the angle between q_0 and q_1 .

1044 1045 Properties

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- A quaternion is a unit quaternion if $||q|| = w^2 + x^2 + y^2 + z^2 = 1$.
- All rotation quaternions must be unit quaternions.
 - A rotation of q_a followed by a rotation of q_b can be combined into a single rotation: $q_c = q_b q_a$. Note that order matters.
 - The conjugate of a quaternion is $q^* = (w, -x, -y, -z)$.
- The inverse of a rotation quaternion is $q^{-1} = q^*$. Then we can see that $qq^{-1} = qq^* = (1, 0, 0, 0)$.
- Quaternion multiplication is associative: abc = a(bc).
- Quaternion multiplication is not commutative: $ab \neq ba$.

B.4 TRANSFORMATION BETWEEN AXIS-ANGLE AND QUATERNION

Axis-angle representation to quaternion representation Axis-angle representation is $u = (u_x, u_y, u_z)$ equipped with a rotation angle θ , and the rotation quaternion is unitary, *i.e.*, $w^2 + x^2 + y^2 + z^2 = 1$. The quaternion is thus:

$$q = \cos\left(\frac{\theta}{2}\right) + \sin\left(\frac{\theta}{2}\right)(u_x i + u_y j + u_z k), \tag{24}$$

1063 or equivalently in the vector form:

$$\boldsymbol{q} = \left(\cos\left(\frac{\theta}{2}\right), \sin\left(\frac{\theta}{2}\right)u_x, \sin\left(\frac{\theta}{2}\right)u_y, \sin\left(\frac{\theta}{2}\right)u_z\right).$$
(25)

1067 Quaternion representation to axis-angle representation Quaternion is q = (w, x, y, z), and to 1068 convert it to axis-angle representation:

1. Compute the angle $\theta = 2\cos^{-1}(w)$.

1070 2. Compute the axis u: 1071

 $\boldsymbol{u} = \frac{(x, y, z)}{\sqrt{x^2 + y^2 + z^2}} = \frac{(x, y, z)}{\sin\frac{\theta}{2}}.$ (26)

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1074 1075 B.5 TRANSFORMATION BETWEEN QUATERNION AND ROTATION

1076 Quaternion to rotation matrix Given a quaternion q = (w, x, y, z), the corresponding rotation 1077 matrix is

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$$R = \begin{bmatrix} 1 - 2y^2 - 2z^2 & 2xy - 2wz & 2xz + 2wy \\ 2xy + 2wz & 1 - 2x^2 - 2z^2 & 2yz - 2wx \\ 2xz - 2wy & 2yz + 2wx & 1 - 2x^2 - 2y^2 \end{bmatrix}$$
(27)

1080 **Rotation matrix to rotation quaternion** Given a rotation matrix $R = \begin{bmatrix} R_{11} & R_{12} & R_{13} \\ R_{21} & R_{22} & R_{23} \\ R_{31} & R_{32} & R_{33} \end{bmatrix}$, then 1081 1082 we first calculate the magnitude of four quaternion components as below: 1084 $|q_0| = \sqrt{\frac{1 + R_{11} + R_{22} + R_{33}}{4}}$ $|q_1| = \sqrt{\frac{1 + R_{11} - R_{22} - R_{33}}{4}}$ 1087 1088 (28) $|q_2| = \sqrt{\frac{1 - R_{11} + R_{22} - R_{33}}{4}}$ 1089 1090 1091 $|q_3| = \sqrt{\frac{1 - R_{11} - R_{22} + R_{33}}{4}}$ 1092 1093 1094 To find the signs of the four elements, we can find the largest magnitude: 1095 1096 • If $|q_0|$ is the largest, then $w = q_0,$ $x = \frac{R_{32} - R_{23}}{4w},$ $y = \frac{R_{13} - R_{31}}{4w},$ $z = \frac{R_{21} - R_{12}}{4w}.$ (29)1099 • If $|q_1|$ is the largest, then 1100 1101 $x = q_1,$ $w = \frac{R_{32} - R_{23}}{4x},$ $y = \frac{R_{12} + R_{21}}{4x},$ $z = \frac{R_{13} + R_{31}}{4x}.$ 1102 (30)1103 • If $|q_2|$ is the largest, then 1104 1105 $y = q_2,$ $w = \frac{R_{13} - R_{31}}{4u},$ $x = \frac{R_{12} + R_{21}}{4y},$ $z = \frac{R_{23} + R_{32}}{4y}.$ (31)1106 1107 1108 • If $|q_3|$ is the largest, then 1109

$$z = q_3, \qquad w = \frac{R_{21} - R_{12}}{4z}, \qquad x = \frac{R_{13} + R_{31}}{4z}, \qquad y = \frac{R_{23} + R_{32}}{4z}.$$
 (32)

The sign is ambiguous because any rotation has two possible quaternion representations. If one isknown, the other one can be found by taking the negative of all four terms.

Besides, there exist other solutions, for instance, extracting quaternion from rotation matrix using eigendecomposition (Horn, 1987; Bar-Itzhack, 2000). We first construct a matrix K with:

$$K = \frac{1}{3} \begin{bmatrix} R_{11} + R_{22} + R_{33} & R_{32} - R_{23} & R_{13} - R_{31} & R_{21} - R_{12} \\ R_{32} - R_{23} & R_{11} - R_{22} - R_{33} & R_{12} + R_{21} & R_{13} + R_{31} \\ R_{13} - R_{31} & R_{12} + R_{21} & R_{22} - R_{11} - R_{33} & R_{23} + R_{32} \\ R_{21} - R_{12} & R_{13} + R_{31} & R_{23} + R_{32} & R_{33} - R_{11} - R_{22} \end{bmatrix} .$$
(33)

1121 Then we perform eigendecomposition $K = V\Lambda V^T$, where Λ is a diagonal matrix with eigenvalues 1122 and V is the matrix with eigenvectors as columns. Finally, we pick up the eigenvector w.r.t. the 1123 largest eigenvalue, and this eigenvector is the unit quaternion.

1125 B.6 TRANSFORMATION BETWEEN AXIS-ANGLE REPRESENTATION AND ROTATION MATRIX

1127 **Axis-angle representation to rotation matrix** Construct the skew-symmetric matrix:

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$$K = \begin{bmatrix} 0 & -u_z & u_y \\ u_z & 0 & -u_x \\ -u_y & u_x & 0 \end{bmatrix}$$
(34)

¹¹³² According to the Rodrigue's rotation formula, we have:

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$$R = I + \sin(\theta)K + (1 - \cos(\theta))K^2.$$
(35)

1134 Rotation matrix to axis-angle representation of R, *i.e.*, 1136 The rotation angle can be computed using the trace Tr(R) = 1

 $\theta = \cos^{-1}\left(\frac{\operatorname{Tr}(R) - 1}{2}\right),\tag{36}$

1138 where Tr(R) is the trace, as $Tr(R) = R_{11} + R_{22} + R_{33}$.

Then we calculate the rotation direction or rotation axis $\boldsymbol{u} = (u_x, u_y, u_z)$, as

$$u_x = \frac{R_{32} - R_{23}}{2\sin\theta}, \qquad u_y = \frac{R_{13} - R_{31}}{2\sin\theta}, \qquad u_z = \frac{R_{21} - R_{12}}{2\sin\theta}.$$
 (37)

1144 Notice that when $\theta = 0$, there is no definition of rotation axis and the whole rotation matrix R is 1145 unitary. When $\theta = \pi$, because $\sin \theta = 0$, then we need other methods (*e.g.*, eigendecomposition) to 1146 determine the rotation axis.

¹¹⁸⁸ C INERTIAL FRAME FOR RIGID BODY

1190 C.1 RIGID REPRESENTATION OF MOLECULE

1192 We employ the following four sequential steps to derive the reference frames that construct the 1193 rotation matrix from N atomic positions r:

- Calculate the mass center: $c = \frac{1}{N} \sum_{i} r_i$.
 - Adjust position relative to the center $\vec{r}_i = r_i c$.
- Compute the inertia tensor $\hat{I} = \sum_{i} ||\mathbf{r}_{i}||^{2}I \mathbf{r}_{i}\mathbf{r}_{i}^{T}$, where I is the unit diagonal matrix.
- Obtain the principal axes of inertia by applying eigen-decomposition on \hat{I} . We have $\hat{I} = Q\Lambda Q^T$, where Q is the orthogonal matrix whose columns are the eigenvectors of \hat{I} , and Λ is the diagonal matrix whose elements are the eigenvalues λ_i of \hat{I} , representing the principal moments of inertial along the principal axes.

1202 1203 C.2 Orthogonal Matrix

In linear algebra, an orthogonal matrix or orthonormal matrix is a square matrix whose columns and rows are orthonormal vectors. This can be written as

$$Q^T Q = Q Q^T = I. aga{38}$$

This leads to the equivalent characterization: a matrix Q is orthogonal if its transpose is equal to its inverse:

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$Q^T = Q^{-1}.$ (39)

Notice that when discussing matrices, the two terms (orthogonal and orthonormal) can be used interchangeably.

1214 If Q is a square matrix, then the conditions $RR^T = I$ and $R^TR = I$ are equivalent. Proof sketch: 1215 $R^TR = I$ and $R^TRR^{-1} = R^{-1}$, so $R^T = R^{-1}$. This can give us $RR^T = I$. 1216

1217 C.3 ROTATION MATRIX FROM RIGID BODY

From Appendix C.1, we can construct the inertial tensors. Then we employ eigenvalue decomposition on the inertial tensor. The normalized eigenvectors v_1, v_2, v_3 form an orthonormal basis, which can be used to construct the rotation matrix, *i.e.*,

$$R = \begin{bmatrix} \boldsymbol{v}_1 & \boldsymbol{v}_2 & \boldsymbol{v}_3 \end{bmatrix}. \tag{40}$$

1224 1225 1226 Eigendecomposition of Inertial Tensors For inertial tensor *I*, the decomposition is: with $Iv_i = \lambda_i v_i$, where λ_i are eigenvalues and $v_i \in \mathbb{R}^{3 \times 1}$ are eigenvectors. Thus, we can have

$$I\boldsymbol{v} = \lambda \boldsymbol{v}$$

$$IR = R\Lambda$$

$$I = R\Lambda R^{-1},$$
(41)

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1231 1232 where $\Lambda = \begin{bmatrix} \lambda_0 & 0 & 0 \\ 0 & \lambda_1 & 0 \\ 0 & 0 & \lambda_2 \end{bmatrix}$. Because the inertial tensors are symmetric matrices, we have that

1234 matrices *R* are orthonormal matrices.

1236 C.4 EXPLORATIONS ON OTHER REFERENCE FRAME OPTION

One critical question is in addition to the Inertial frame, do we have other options for modeling the rigidity? One simple solution is to directly apply the eigendecomposition as principal component analysis (PCA) on the point clouds of centered molecules.

1241 First, we would like to clarify that there are two roles and one important property of the inertial frame and its eigenvectors:

1242 1243 1244		Role 1: The three bases in inertial frames act as a reference or a canonical pose.Role 2: The three bases enable modeling the velocity function in SO3 space.What's more important, we expect the three bases to be numerically stable.								
1245 1246 1247	Then pose,	tting back to the question, though both can help build up the reference frame or canonical ere are certain aspects we would like to emphasize when comparing PCA and Inertial frames.								
1247 1248 1249 1250 1251 1252 1253 1254 1255 1256 1257 1258 1259 1260 1261 1261 1262 1263	-	 Canonical pose: The key question is defining a canonical pose. If we just do PCA on the point clouds, this cannot guarantee the group symmetry in SE(3). However, if we remove the center point, then the group symmetry can be guaranteed; then we can use SVD to get the three principal components. Till this step, one can find this is somewhat similar to the inertial frame construction (Sec 3.1). Difference between PCA and Inertial Frame as reference frames: Though both can be used for building up the reference frame or canonical poses, SVD (for PCA) on the set of point clouds N × 3 (N is the number of atoms) can be less numerically unstable, while eigendecomposition Inertial Frame 3 × 3 can be more numerically stable. We conduct an experiment to verify this. Experiment setup: Suppose we have weakly-correlated structures X₀ and strongly-correlated structures X₁, and we find the corresponding bases using either eigendecomposition on centered X or inertial frame I. The two bases are marked as B₀ ∈ ℝ^{3×3} and B₁ ∈ ℝ^{3×3}. Objective: We can obtain the rotation matrix with R^T = B^T₀B₁, and then we can rotate the whole molecular system as X̃₁ = X₀R^T, and we are measuring the reconstruction errors as MSE(X₁ − X̃₁). We mark the MSE using inertial frame and PCA as δ_{Inertial} 								
1264 1265 1266 1267 1268 1269 1270 1271 1272 1273 1274 1275 1276		and δ_{PCA} , respectively. If $\delta_{Inertial} < \delta_{PCA}$, then we can conclude that using the inertial frame is more stable than using PCA, and vice versa. Notice that since the MSE reconstruction is meaningless when it is too small, so we only compare these two frames when at least one of them has reconstruction greater than or equal to a threshold θ . - Results: The comparison results are in Table 5, and we can observe that in general, using the inertial frame is more stable than PCA. We are listing multiple reconstruction threshold θ in Table 5, and we are using $\theta = 1e - 3$ in the main article. Table 5: Comparison of using inertial frame and PCA for reconstruction (%). θ 1e-3 1e-4 1e-5 1e-6 1e-7 1e-8 $P(\delta_{Inertial} < \delta_{PCA})$ 0.434 0.884 1.338 1.794 2.254 2.727 $P(\delta_{Inertial} > \delta_{PCA})$ 0.371 0.756 1.147 1.539 1.934 2.345								
1277 1278 1279	C 5	Kabsch Algorithm								

1279 C.5 KABSCH ALGORITHM 1280

Kabsch algorithm is one way to compute the optimal rotation matrix that minimizes the root-meansquare deviation (RMSD) between two sets of points (atoms in our case). However, it is guaranteed in
the COD-Cluster17 dataset that the molecules in weakly correlated structures can rotate to molecules
in strongly correlated structures; in other words, the RMSD can be approximately 0 if we use the
Kabsch algorithm, which is equivalent to calculating the rotation matrix directly after we fix the
poses. We have shown how to calculate the rotation matrix in the experiment above.

¹²⁹⁶ D Proof of Theorem 1 1297

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Proof. For three vectors, we can easily find a counter-example, as illustrated in Figure 3 (a, b).
Figure 3 (a, b) describes two cases where we have the same initial frame, and we can rotate it to two different final frames with two rotation matrices, yet the righthandness still matches. We can easily see that there are four options of rotation matrices in this case, and we cannot uniquely determine the final inertial frame in this case.

More rigorously, let us first assume that there exists a rotation transformation R that can transform the initial coordinate system Q_i to the final coordinate system Q_f , as:

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 $\begin{bmatrix} Q_{f,0} \\ Q_{f,1} \\ Q_{f,2} \end{bmatrix}^T = R \cdot \begin{bmatrix} Q_{i,0} \\ Q_{i,1} \\ Q_{i,2} \end{bmatrix}^T$ (42)

(43)

(44)

First, as Lemma 1, we should change either zero or two directions for direction alignment.

Then without loss of generality, we can assume the two directions to be the last two axes. Thus, we can obtain a rotation matrix R' such that R' is rotating R along vector $Q_{f,0}$ with 180 degrees. We can represent R' using Rodrigue's rotation formula, as $R' = (2Q_{f,0}Q_{f,0}^T - I)R$. Thus, we can have:

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This is essentially saying starting from one initial frame, we can have multiple matched final frames. Thus, using only three vectors cannot uniquely determine the direction matching. We provide two

 $R' \cdot \begin{bmatrix} Q_{i,0} \\ Q_{i,1} \\ Q_{i,2} \end{bmatrix}^T = (2Q_{f,0}Q_{f,0}^T - I) \begin{bmatrix} Q_{f,0} \\ Q_{f,1} \\ Q_{f,2} \end{bmatrix}^T = \begin{bmatrix} Q_{f,0} \\ -Q_{f,1} \\ -Q_{f,2} \end{bmatrix}^T$

1324 Thus, using only three vecto examples in Figure 3 (a, b).

For the four vectors, we introduce an extra atom into the inertial frame system, and such an extra atom point is nonplanar to the three base axes. Then the problem becomes: starting from an initial frame and an extra point, can we find multiple rotation matrices such that the final frames have reflected directions? To be more rigorous, let us have the following formulation.

First, let us assume we have this rotation matrix:

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1337 1338 As discussed above, we need to guarantee the right-handedness property, thus, without loss generality, 1339 here we also assume the last two axes are reflected. The question turns to: does it exit another rotation 1340 matrix R', such that:

 $\begin{bmatrix} Q_{f,0} \\ Q_{f,1} \\ Q_{f,2} \\ \mathbf{v} \end{bmatrix}^T = R \cdot \begin{bmatrix} Q_{i,0} \\ Q_{i,1} \\ Q_{i,2} \\ \mathbf{v} \end{bmatrix}^T$

 $\begin{bmatrix} Q_{f,0} \\ -Q_{f,1} \\ -Q_{f,2} \\ \boldsymbol{v} \end{bmatrix}^T = R' \cdot \begin{bmatrix} Q_{i,0} \\ Q_{i,1} \\ Q_{i,2} \\ \boldsymbol{v} \end{bmatrix}^T$ (45)

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We now use contradiction. Since we still have the two axes rotated 180 degrees around the first axes, $Q_{f,0}$, so $R' = (2Q_{f,0}Q_{f,0}^T - I)R$. Then given the two conditions $v^T = Rv^T$ and $v^T = R'v^T$, we have $(2Q_{f,0}Q_{f,0}^T - I)v^T = v^T$. 1350 If we let $Q_{f,0} = [k_1, k_2, k_3]$ and $\boldsymbol{v} = [v_1, v_2, v_3]$, then we have

$$\begin{array}{ll} 1352 & (2Q_{f,0}Q_{f,0}^{T}-I)\boldsymbol{v}^{T}=\boldsymbol{v}^{T} \\ 1353 \\ 1354 \\ 1355 \\ 1355 \\ 1355 \\ 1356 \\ 1357 \\ 1358 \\ 1359 \\ 1360 \\ 1361 \end{array} \begin{pmatrix} k_{1}k_{1} & k_{1}k_{2} & k_{1}k_{3} \\ k_{1}k_{2} & k_{2}k_{2} & k_{2}k_{3} \\ k_{2}k_{3} & k_{2}k_{3} & k_{3}k_{3} \\ k_{2}k_{3} & k_{2}k_{3} & k_{3}k_{3} \\ k_{2}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{3}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{3}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{3}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{3}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{3}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{4}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{5}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{6}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{7}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{7}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}v_{3}) \\ k_{7}(k_{1}v_{1}+k_{2}v_{2}+k_{3}v_{3}) \\ k_{7}(k_{1}v_{1}+k_{2}v_{$$

After calculation, we can obtain that $Q_{f,0} = cv$, where c is a coefficient. However, as we claimed in the condition, v does not lie in the same line as $Q_{f,0}$, thus, there does not exist such another rotation matrix $R' \neq R$ satisfying Equation (45). We also provide two examples in Figure 3 (c, d).

1367 By contradiction, we can tell that there is only one unique rotation mapping from the initial inertial frame to the final inertial frame. \Box

To sum up, three points cannot form a rigid structure in Euclidean space, thus there can exist multiple reflection transformations, leading to opposite inertial frames. Four points can form a rigid structure, thus there exists only one reflection transformation.

1404 E PROBLEM FORMULATION AND MORE DETAILS OF ASSEMBLEFLOW

1406 E.1 PROBLEM FORMULATION

We would like to emphasize that previous works aim at atomic level modeling, while our proposed
AssembleFlow focuses on molecular level modeling. Meanwhile, both models need to satisfy the
SE(3)-equivariance, as detailed below.

Atomic Level Modeling Existing deep learning frameworks have been using atomic level modeling.
For each atom r, the inference step is:

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$r_{t+1} = r_t + x_{\theta,t},$ s.t. $x_{\theta,t}$ is SE(3)-equivariant. (47)

Thus, we can observe that such a problem formulation cannot guarantee the rigidity of each molecule during the crystallization process.

¹⁴²⁰ Molecular Level Modeling In our proposed AssembleFlow, it learns the translation and rotation at the molecule level. For each atom r, the inference step is:

$$\boldsymbol{r}_{t+1} = R_{\theta,t}(\boldsymbol{r}_t + \boldsymbol{x}_{\theta,t}),$$

s.t. $\boldsymbol{x}_{\theta,t}$ and $R_{\theta,t}$ are SE(3)-equivariant. (48)

The $x_{\theta,t}$ and $R_{\theta,t}$ are molecular level modeling. Notice that this also holds after we take the reparameterization, as discussed in Section 3.4. In Appendix E.2, we will explore how to define the SE(3)-equivariant models on top of that.

1429 1430 E.2 SE(3)-EQUIVARIANT VELOCITY FUNCTION

We consider two types of SE(3)-equivariant models as the velocity function. As shown in Figure 9, the inputs are the same for learning: the positions at the initial step and the final step, respectively. We take the position of the mass center for each molecule in the cluster to obtain the translation in \mathbb{R}^3 (x_i and x_f), and we take the first principal axes of inertial frames to obtain the reference coordinate system for rotation in SO(3) (q_i and q_f with alignment). Then we adopt Equations (6) and (7) for the interpolation on SO(3) and \mathbb{R}^3 group respectively, which gives us translation x_t and rotation q_t at interpolation time $t \in [0, 1]$.

Recall that the data structure considered here is the cluster of molecules, thus it is natural to split the modeling into intra-molecule and inter-molecule modeling, as introduced below.

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Intra-molecule Modeling. For each molecule in the cluster, we adopt the SE(3)-equivariant
PaiNN (Schütt et al., 2021) to obtain the representation for each atom. Such an atomic representation
can encode the inherent geometric structural information of individual molecules, which can be
passed to inter-molecule modeling in the next step.

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1446 Inter-molecule Modeling. This step aims to model the inter-molecule interactions during the 1447 molecular crystallization process based on the intra-molecule representation. We can have two 1448 options for SE(3)-equivariant inter-molecule modeling: (1) to project x_t and q_t back to obtain the 1449 atom-wise position and do modeling, as in Figure 9(a), or (2) to directly perform molecular level 1450 modeling on molecular-level translation x_t and rotation q_t , as in Figure 9(b).

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• Atomic level modeling. This means we build up the SE(3)-equivariant models on top of atom positions r_t at time t, and the outputs are the rotation and translation for step t + 1 or step T (if we use reparameterization).

- Obtain intra-molecule representation h_a using PaiNN (Schütt et al., 2021).
- Obtain time embedding h_t with positional encoding (Vaswani et al., 2017).
- Build up the vector frame basis (Liu et al., 2023) for each atom F_a , based on its neighborhoods.
 - Then we update the atomic representation h_a as the summation of h_a and h_t .

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rotation matrix) and conduct the alignment between the initial and final frames. (3) Last but not least, we transform the coordinate system from rotation matrix to quaternion as discussed in Appendix B.5.

Algorithm 1 Learning of Assem	ibleFlow.		
 Inputs: For N atoms in N molecular level initial rotation final position r ∈ ℝ^{N×3}, m x₁ ∈ ℝ^{M×3}, timestep T ∈ Ω for epoch e ∈ [1, E] do 	\overline{A} molecules, we have atomic tion quaternion $q_0^{M \times 4}$ and tra- olecular level final rotation qu \mathbb{R} , epoch $E \in \mathbb{R}$, coefficients of	c level initial position anslation $\boldsymbol{x}_0 \in \mathbb{R}^{M\times d}$ naternion $\boldsymbol{q}_1 \in \mathbb{R}^{M\times d}$ $\boldsymbol{x}_0, \boldsymbol{\alpha}_1 \in \mathbb{R}.$	on $\boldsymbol{r}_0 \in \mathbb{R}^{N \times 3}$, $^{\times 3}$, atomic level 4 and translation
3: Sample $t \in [1, T]$.			
4: Conduct LERP to obtain	translation \boldsymbol{x}_t at time t , follow	ing Equation (7).	
5: Perform SLERP to obtain	quaternion q_t at time t, follow	wing Equation (6).	
6: Predict the final quaterni SE(2) against modeli	on $\boldsymbol{q}_1 = \boldsymbol{q}_{1,\theta}(\boldsymbol{x}_t, \boldsymbol{q}_t, t)$ and the provided in A provided to \boldsymbol{x}_t	$x_1 = x_{1,\theta}$	$(\boldsymbol{x}_t, \boldsymbol{q}_t, t)$ using
7. Minimize loss $C = \alpha_0 C_{\pi}$	$\pm \alpha_{\rm s} \int ds ds cuscus d d d d d d d d d d d d d d d d d d d$	as defined in Faux	ation (13)
8: end for	\sim , reparameter + $\alpha_1 \sim SO(3)$, reparameter	er, as defined in Equa	(15).
Algorithm 2 Inference of Assen	nbleFlow.		
molecular level initial rotat \mathbb{R} , learned SE(3)-equivaria $\hat{x}_{1,\theta}(x_t, q_t, t)$. 2: for timestep $t \in [1, T]$ do 3: Predict the final quaterni SE(3)-equivariant modeli 4: Calculate the next-step qu 5: Move the cluster of molec 6: Obtain the corresponding 7: end for 8: The final predicted crystal s E.4 HYPER-PARAMETERS We provide the key hyper-param Table 6: I Model	ion quaternion $q_0^{M \times 4}$ and training in quaternion $q_0^{M \times 4}$ and train the models final rotation quaternion $\hat{q}_{1,\theta}(x_t, q_t, t)$ and translation \hat{q}_{t+1} and translation cules w.r.t. \hat{q}_{t+1} and \hat{x}_{t+1} . atomic positions r_{t+1} at time tructure is \hat{r}_T .	nslation $\hat{x}_0 \in \mathbb{R}^{M \times i}$ ernion $\hat{q}_{1,\theta}(x_t, q_t, t)$ anslation $\hat{x}_1 = \hat{x}_{1,\theta}$ \hat{x}_{t+1} as Equation (14 t + 1, following Equ le 6.	$x_t, q_t, t)$ using (x_t, q_t, t) using (1). (48).
Model	Hyperparamete	er Value	
Intra-modeling	PaiNN ambadding dir	n (128)	
	num of layers	{3}	
	cutoff	{5}	
	read out	{mean}	
Intra-modeling	Atomic Level num of layers	$\{2,5\}$	
	num of head	$\{4, 8\}$	
	num of timeste	eps {50, 200}	
	α_0	$\{1\}$	
T , 1	α ₁	(4.5)	
Intra-modeling	num of head	$\{4,5\}$ $\{4,8\}$	
	num of timeste	eps {50, 200}	
	$lpha_0$	$\{1\}$	
	α_1	{1, 10}	
Optimization	seed	$\{0, 42, 123\}$ $\{1000, 2000\}$	
	cpouns	$\{90, 50\}$	
	cutoff c	{20, 50}	
	cutoff c learning rate	$\{20, 50\}\$ $\{1e-4, 5e-4\}$	

¹⁵⁶⁶ F DETAILS OF BASELINES

1568 F.1 DEEP LEARNING BASELINES AND PARAMETERS 1569

Notice that for all the baselines listed below, we also adopt the PaiNN for atomic level representation (Schütt et al., 2021), and the hyperparameters are the same as Appendix E. We list the remaining hyperparameters of baselines in Table 7.

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Table 7: Hyperparameter specifications for deep learning baselines.

1576	Model	Hyperparameter	Value
1577 1578 1579 1580 1581	GNN-MD	seed epochs num of timesteps cutoff <i>c</i> learning rate optimizer	{0, 42, 123} 3000 {50, 200} {20, 50} {1e-4, 5e-4} {Adam }
1582 1583 1584 1585 1586	CrystalSDE	SDE type seed epochs num of timesteps cutoff <i>c</i> learning rate optimizer	{VE, VP} {0, 42, 123} 3000 {50, 200} {20, 50} {1e-4, 5e-4} {Adam }
1587 1588 1589 1590 1591 1592	CrystalFlow	interpolation type seed epochs num of timesteps cutoff <i>c</i> learning rate optimizer	{VE, VP, LERP} {0, 42, 123} 3000 {50, 200} {20, 50} {1e-4, 5e-4} {Adam }

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1595 F.2 PACKMOL

We want to mention that using PackMol for evaluation with the packing matching metrics is nontrivial. This is because the atom ordering in each molecule and the molecule ordering in each cluster are different between PackMol simulated results and the ground-truth results. We note that deep learning methods do not have this issue because the orderings of initial atoms/molecules match with the final atoms/molecules.

Thus, to address this issue, we first use the Hungarian algorithm to match the mass centers of simulated results to obtain the least matching distance with the ground truth mass centers, *i.e.*, PM (center). This gives us the molecule ordering mapping from simulated clusters to the ground-truth clusters. Then for each molecule simulated-and-ground-truth pair, we apply the Hungarian algorithm again to obtain the minimum distance for alignment.

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1620 G DATASET AND EVALUATION METRICS

1622 G.1 DATASET

We evaluate our method using the crystallization dataset COD-Cluster17 from (Liu et al., 2024). This COD-Cluster17 is a curated subset derived from the COD database (Grazulis et al., 2009). We note that some single molecules/substances can crystallize in different forms, known as polymorphs. This arises due to the change of the configurations when the process happens, such as the environment temperature, pressure, and solvent. COD-Cluster17 simplifies this setup by ignoring the configuration information and treats the crystallization problem as a density estimation problem.

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- 1631 G.2 DETAILS OF EVALUATION METRICS

We illustrate five types of evaluation metrics below. Notice that in the original dataset, the dynamics or trajectories of molecules are missing. Thus, our evaluation is based on the ground truth cluster geometry at the last step.

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Packing Matching (PM) This metric quantifies how well the generated molecular assemblies
match the reference crystal structures in terms of spatial arrangement and packing density (Chisholm
Motherwell, 2005). It is a key indicator of how accurately a model can replicate real-world
crystallization patterns. We provide both the atomic MP, denoted as "PM (atom)" and mass-centerlevel PM, denoted as "PM (center)".

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Collision This follows (Cordero et al., 2008). It measures if there is any atomic collision in the 1642 predicted assemblies. Atoms must maintain a minimum pairwise distance governed by the balance of 1643 attractive and repulsive forces. More concretely, we are using covalent radii as the most strict metric 1644 for atomic collisions in molecular generation. This is because it provides a precise lower bound 1645 for the distances between atoms when they are bonded. In other words, covalent radii represent the 1646 distance at which two atoms form a stable covalent bond, which is a very close and well-defined 1647 interaction compared to non-covalent interactions. However, other types of atomic radii, such as 1648 van der Waals radii or ionic radii, can be used for different purposes, depending on the nature of the 1649 interaction you're modeling.

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Separation We extend the metric from (Xie et al., 2022; Yang et al., 2024) to our setting. A cluster of molecules is valid if the minimum distance between molecules is above 0.5Å (Court et al., 2020).
This metric is referred to as *separation* to measure the validity to avoid unphysical interactions at the molecular level.

1656 Compactness We propose this measure by calculating the percentage of simulated clusters where 1657 the maximum atomic pairwise distances are below 100Å. This assesses the spatial efficiency of 1658 the molecular assemblies, indicating how closely the constituent molecules are packed together. A 1659 higher compactness value suggests a more efficient arrangement, where the intermolecular spaces are 1660 minimized, leading to a denser crystalline structure.

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DFT Energy This metric evaluates the stability of crystal structures by calculating their total
electronic energy using Density Functional Theory (DFT) (Kohn & Sham, 1965). Lower DFT energy
generally suggests a more stable molecular configuration. To be more concrete, we adopt ACPYPE
(AnteChamber PYthon Parser interfacE) (Sousa da Silva & Vranken, 2012) and GROMACS (Van
Der Spoel et al., 2005) for energy calculation.

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ABLATION STUDIES Η

H.1 ABLATION STUDIES ON RANDOM SAMPLING

Here we are adding another baseline by randomly sampling translation and rotation.

- For SO(3), we can do random sampling.

• For \mathbb{R}^3 , we first obtain the range of atom positions in the training data, and then we just do uniform sampling within this range.

The results are in Table 8. As observed in Table 8, the Random baseline performs exceptionally well across all three validity metrics; however, its packing matching is significantly worse by an order of magnitude.

Table 8: AssembleFlow against six generative models on COD-Cluster17 with 5K, 10K, and all samples. The best results are marked in **bold**. Baseline Random has the best validity metrics, but they are meaningless since the packing matching is extremely high, remarking that the results collapse. Thus, we mark them in gray.

	Packing	Matching	Validity		
	PM (atom) ↓	PM (center) \downarrow	Collision \downarrow	Separation ↑	Compactness ↑
		Dataset: COD-Cl	uster17-5K		
Random	54.07 ± 0.42	54.62 ± 0.43	0.31 ± 0.01	99.88 ± 0.01	100.00 ± 0.00
GNN-MD	13.67 ± 0.06	13.80 ± 0.07	27.53 ± 0.49	0.22 ± 0.11	$\textbf{100.00} \pm \textbf{0.00}$
CrystalSDE-VE	15.52 ± 1.48	16.46 ± 0.99	1.20 ± 0.08	27.17 ± 0.86	57.47 ± 7.76
CrystalSDE-VP	18.15 ± 3.02	19.15 ± 4.46	0.84 ± 0.14	53.13 ± 12.89	34.00 ± 30.75
CrystalFlow-VE	14.87 ± 7.07	13.08 ± 4.51	1.37 ± 0.04	35.70 ± 0.73	8.40 ± 4.17
CrystalFlow-VP	15.71 ± 2.69	17.10 ± 1.89	1.38 ± 0.04	35.43 ± 0.88	4.87 ± 1.09
CrystalFlow-LERP	13.59 ± 0.09	13.26 ± 0.09	0.34 ± 0.01	97.38 ± 0.10	$\textbf{100.00} \pm \textbf{0.00}$
AssembleFlow (ours)	$\textbf{7.27} \pm \textbf{0.04}$	$\textbf{6.13} \pm \textbf{0.10}$	$\textbf{0.33} \pm \textbf{0.00}$	$\textbf{97.64} \pm \textbf{0.36}$	$\textbf{100.00} \pm \textbf{0.00}$
		Dataset: COD-Clu	ister17-10K		
Random	54.20 ± 0.90	54.76 ± 0.90	0.30 ± 0.00	99.86 ± 0.01	100.00 ± 0.00
GNN-MD	13.83 ± 0.06	13.90 ± 0.05	27.88 ± 0.49	0.23 ± 0.11	$\textbf{100.00} \pm \textbf{0.00}$
CrystalSDE-VE	17.25 ± 2.46	17.86 ± 1.11	0.99 ± 0.27	32.99 ± 10.72	34.93 ± 14.99
CrystalSDE-VP	22.20 ± 3.29	21.39 ± 1.50	0.53 ± 0.35	52.48 ± 15.44	16.83 ± 18.09
CrystalFlow-VE	16.41 ± 2.64	16.71 ± 2.35	1.42 ± 0.03	33.79 ± 0.51	5.47 ± 0.47
CrystalFlow-VP	19.39 ± 4.37	16.01 ± 3.13	1.44 ± 0.03	33.35 ± 0.55	4.23 ± 0.48
CrystalFlow-LERP	13.54 ± 0.03	13.20 ± 0.03	0.32 ± 0.00	97.32 ± 0.05	$\textbf{100.00} \pm \textbf{0.00}$
AssembleFlow (ours)	$\textbf{7.38} \pm \textbf{0.03}$	$\textbf{6.21} \pm \textbf{0.05}$	$\textbf{0.31} \pm \textbf{0.00}$	$\textbf{97.73} \pm \textbf{0.16}$	99.93 ± 0.05
		Dataset: COD-Cl	uster17-All		
Random	65.94 ± 0.07	66.56 ± 0.07	0.30 ± 0.00	99.91 ± 0.00	100.00 ± 0.00
GNN-MD	22.30 ± 12.04	14.51 ± 0.82	24.29 ± 4.58	4.13 ± 5.60	98.77 ± 1.73
CrystalSDE-VE	17.28 ± 0.73	18.92 ± 0.03	0.19 ± 0.18	15.47 ± 12.42	2.51 ± 2.37
CrystalSDE-VP	18.03 ± 4.56	20.02 ± 3.70	0.55 ± 0.19	48.78 ± 1.70	6.88 ± 2.82
CrystalFlow-VE	12.80 ± 1.20	15.09 ± 0.34	1.41 ± 0.01	35.34 ± 0.28	2.90 ± 0.02
CrystalFlow-VP	13.50 ± 0.44	13.28 ± 0.48	1.51 ± 0.02	33.06 ± 1.31	6.61 ± 3.17
CrystalFlow-LERP	13.61 ± 0.00	13.28 ± 0.01	0.34 ± 0.00	97.34 ± 0.02	$\textbf{99.99} \pm \textbf{0.01}$
	7.27 ± 0.01	(21 ± 0.01)	0.21 0.00	09 15 1 0 22	00.09 \ 0.00

1728 H.2 Ablation Studies on Interpolation on SO(3)

1730 Empirical results. We consider replacing the SLERP with EMLERP in AssembleFlow, and name
1731 it as AssembleFlow-EMLERP. We conduct the experiment on COD-5000, where we are taking the
1732 optimal hyperparameters from AssembleFlow.

The results are in Table 9. As observed, using SLERP is better than EMLERP. We are still running results for COD-10k and COD, and will update the results later.

Table 9: AssembleFlow against six generative models on COD-Cluster17 with 5K, 10K, and all samples (running now). The best results are marked in **bold**.

	Packing	Matching		Validity	
	PM (atom) \downarrow	PM (center) \downarrow	Collision \downarrow	Separation †	Compactness ↑
		Dataset: COD-Clus	ster17-5K		
GNN-MD	13.67 ± 0.06	13.80 ± 0.07	27.53 ± 0.49	0.22 ± 0.11	$\textbf{100.00} \pm \textbf{0.00}$
CrystalSDE-VE	15.52 ± 1.48	16.46 ± 0.99	1.20 ± 0.08	27.17 ± 0.86	57.47 ± 7.76
CrystalSDE-VP	18.15 ± 3.02	19.15 ± 4.46	0.84 ± 0.14	53.13 ± 12.89	34.00 ± 30.75
CrystalFlow-VE	14.87 ± 7.07	13.08 ± 4.51	1.37 ± 0.04	35.70 ± 0.73	8.40 ± 4.17
CrystalFlow-VP	15.71 ± 2.69	17.10 ± 1.89	1.38 ± 0.04	35.43 ± 0.88	4.87 ± 1.09
CrystalFlow-LERP	13.59 ± 0.09	13.26 ± 0.09	0.34 ± 0.01	97.38 ± 0.10	$\textbf{100.00} \pm \textbf{0.00}$
CrystalFlow-LERP	13.59 ± 0.09	13.26 ± 0.09	0.34 ± 0.01	97.38 ± 0.10	$\textbf{100.00} \pm \textbf{0.00}$
AssembleFlow-EMLERP	7.30 ± 0.04	6.32 ± 0.04	0.37 ± 0.01	93.38 ± 0.54	100.00 ± 0.00
AssembleFlow (ours)	$\textbf{7.27} \pm \textbf{0.04}$	$\textbf{6.13} \pm \textbf{0.10}$	$\textbf{0.33} \pm \textbf{0.00}$	$\textbf{97.64} \pm \textbf{0.36}$	$\textbf{100.00} \pm \textbf{0.00}$
		Dataset: COD-Clus	ter17-10K		
GNN-MD	13.83 ± 0.06	13.90 ± 0.05	27.88 ± 0.49	0.23 ± 0.11	$\textbf{100.00} \pm \textbf{0.00}$
CrystalSDE-VE	17.25 ± 2.46	17.86 ± 1.11	0.99 ± 0.27	32.99 ± 10.72	34.93 ± 14.99
CrystalSDE-VP	22.20 ± 3.29	21.39 ± 1.50	0.53 ± 0.35	52.48 ± 15.44	16.83 ± 18.09
CrystalFlow-VE	16.41 ± 2.64	16.71 ± 2.35	1.42 ± 0.03	33.79 ± 0.51	5.47 ± 0.47
CrystalFlow-VP	19.39 ± 4.37	16.01 ± 3.13	1.44 ± 0.03	33.35 ± 0.55	4.23 ± 0.48
CrystalFlow-LERP	13.54 ± 0.03	13.20 ± 0.03	0.32 ± 0.00	97.32 ± 0.05	$\textbf{100.00} \pm \textbf{0.00}$
AssembleFlow-EMLERP	7.51 ± 0.17	6.46 ± 0.22	0.33 ± 0.00	94.68 ± 0.44	99.93 ± 0.05
AssembleFlow (ours)	$\textbf{7.38} \pm \textbf{0.03}$	$\textbf{6.21} \pm \textbf{0.05}$	$\textbf{0.31} \pm \textbf{0.00}$	$\textbf{97.73} \pm \textbf{0.16}$	99.93 ± 0.05
		Dataset: COD-Clus	ster17-All		
GNN-MD	22.30 ± 12.04	14.51 ± 0.82	24.29 ± 4.58	4.13 ± 5.60	98.77 ± 1.73
CrystalSDE-VE	17.28 ± 0.73	18.92 ± 0.03	0.19 ± 0.18	15.47 ± 12.42	2.51 ± 2.37
CrystalSDE-VP	18.03 ± 4.56	20.02 ± 3.70	0.55 ± 0.19	48.78 ± 1.70	6.88 ± 2.82
CrystalFlow-VE	12.80 ± 1.20	15.09 ± 0.34	1.41 ± 0.01	35.34 ± 0.28	2.90 ± 0.02
CrystalFlow-VP	13.50 ± 0.44	13.28 ± 0.48	1.51 ± 0.02	33.06 ± 1.31	6.61 ± 3.17
CrystalFlow-LERP	13.61 ± 0.00	13.28 ± 0.01	0.34 ± 0.00	97.34 ± 0.02	$\textbf{99.99} \pm \textbf{0.01}$
AssembleFlow-EMLERP	$\textbf{7.28} \pm \textbf{0.00}$	6.23 ± 0.01	0.35 ± 0.00	93.17 ± 0.02	99.98 ± 0.00
AssembleFlow (ours)	7.37 ± 0.01	$\textbf{6.21} \pm \textbf{0.01}$	$\textbf{0.31} \pm \textbf{0.00}$	$\textbf{98.15} \pm \textbf{0.22}$	99.98 ± 0.00

1782 H.3 ABLATION STUDIES ON ADDITIONAL ENERGY RESULTS

We computed and compared the formation energy for both systems in Figure 4. Formation energy, defined as the difference between a compound's DFT total energy and the sum of the energies of its constituent elements, is a direct indicator of the relative favorability of a material's energy state. We also provide the pairwise differences between the two systems in terms of formation energy in Figure 10.



Figure 10: Formation energy difference between PackMol-constructed assemblies and AssembleFlow-predicted.