

GeoAB: Towards Realistic Antibody Design and Reliable Affinity Maturation

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Abstract

Increasing works for antibody design are emerging to generate sequences and structures in Complementarity Determining Regions (CDRs), but problems still exist. We focus on two of them: *(i) authenticity of the generated structure* and *(ii) rationality of the affinity maturation*, and propose GEOAB as a solution. In specific, GeoAB-Designer generates CDR structures with realistic internal geometries, composed of a generative geometry initializer (Geo-Initializer) and a position refiner (Geo-Refiner); GeoAB-Optimizer achieves affinity maturation by accurately predicting both the mutation effects and structures of mutant antibodies with the same network architecture as Geo-Refiner. Experiments show that GEOAB achieves state-of-the-art performance in CDR co-design and mutation effect predictions, and fulfills the discussed tasks effectively.

1. Introduction

Antibodies are immune proteins that can bind to a kind of antigen protein so as to recognize and neutralize the pathogen (Basu et al., 2019). There are two heavy chains and two light chains in an antibody, leading the antibodies to Y-shaped proteins. In each chain, three variable regions determine the binding property of the antibody towards the antigen, which are called Complementarity Determining Regions (CDRs). Therefore, to design antibodies that bind to specific antigens with desirable properties, the key problem is to establish computational approaches to accurately identify the 3D structures and 1D sequence of the amino acids in the CDRs because the large combinatorial space of amino acid types results in infeasible and unaffordable validation of wet-lab experiments.

Recently, deep learning has revolutionized fields like drug

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Proceedings of the 41st International Conference on Machine Learning, Vienna, Austria. PMLR 235, 2024. Copyright 2024 by the author(s).

discovery and protein design (Jumper et al., 2021; Dauparas et al., 2022; Lin et al., 2022; 2023). To be specific, increasing works are emerging for antibody design, which considers antigens and conserved regions (non-CDRs) as context information, to achieve the co-design of the sequence and structure of the target CDRs (Kong et al., 2023a; Jin et al., 2021). Besides, based on the co-design models, the optimization of the antibody by mutating amino acids in the CDRs to enhance the binding affinity can be realized, which is called affinity maturation (Cai et al., 2023). Iterative Target Augmentation (ITA) is proposed to fulfill the affinity maturation tasks (Yang et al., 2020), by iteratively adding co-design models' prediction of the mutant antibodies structures and sequences with higher affinity predicted by another pretrained model to the training set and thus to guide and retrain the co-design models to generate antibodies of high affinity. However, several problems still exist, and we focus on two of them: *(i) authenticity of the generated structure* and *(ii) rationality of the affinity maturation*.

To illustrate problem *(i)*, we claim that in the structure design of the CDRs, these methods usually ignore geometry constraints of protein structures, such as peptide planar and inflexible geometries like bond lengths (Robinson et al., 2014), causing the unrealistic modeling of the CDR structures. Figure 1 gives an illustration of the distribution gap between partial inflexible internal geometries (bond length, angles, and inflexible torsions) and redundant ones (flexible torsions) of the structures generated by state-of-the-art methods and real-world proteins (Sec. 5.1 gives details).

For problem *(ii)*, these methods first employ a trained model with sequences and structures as input to predict the change in binding free energy ($\Delta\Delta G$) and exploit the ITA algorithm to tackle the optimization. This procedure is irrational due to two problems. First, the used $\Delta\Delta G$ -predictor requires mutant types' structures as input, while they are usually unknown, so it uses predicted structures by methods like Rosetta programs (Park et al., 2016) as training instances. However, the generated mutant antibodies by the co-design models are of different domains from the training ones, resulting in the unreliability of the predicted $\Delta\Delta G$ (See Fig. 5 as an example). Secondly, the inaccurate prediction of $\Delta\Delta G$ cannot guide the ITA reliably to generate antibodies toward affinity maturation targets. The other ap-

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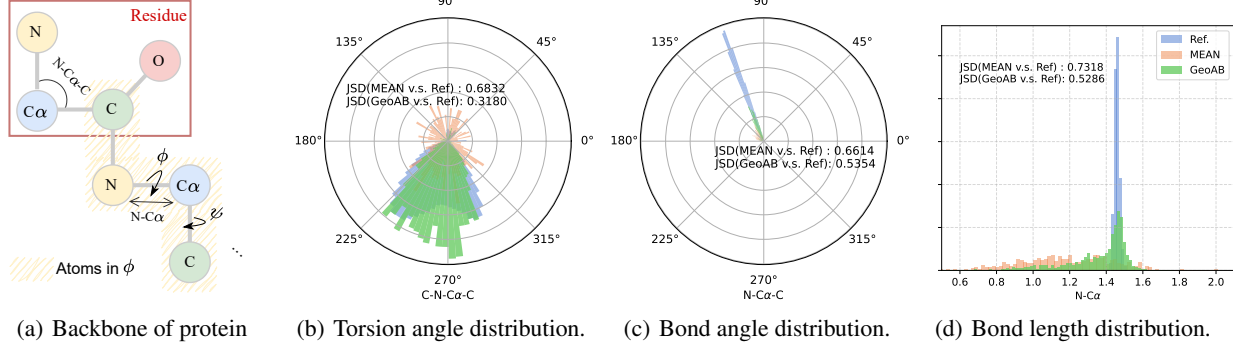


Figure 1: Distribution gap between several internal geometries of true protein structures as a reference and of generated ones. Jensen–Shannon divergence (JSD) is used as a measurement. (a) gives a sketch of the demonstrated geometries. (b,c,d) The torsion angle ‘C-N-Cα-C’ is a redundant (flexible) geometry, while the bond angle ‘N-Cα-C’ and length ‘N-Cα’ are inflexible geometries. Internal geometries obtained from CDR structures generated by MEAN usually deviate from the true distribution greatly. In comparison, GeoAB narrows the gap and generates more realistic and reasonable CDR structures.

proach for affinity maturation is to enumerate amino acid types on the points of interest. Previous works show that multiple-point mutations usually achieve successful affinity maturation (Sulea et al., 2018), but the enumeration of mutations on multiple points is unaffordable in wet-lab validation. Therefore, it is urgent to establish an effective computational method for narrowing down the search space for the task.

To address them, we proposed GEOAB. *On antibody CDR co-design*, it can jointly sample CDR sequences and structures with realistic protein geometry, by using heterogeneous residue-level encoder and equivariant atom-level interaction layers and employing energy-informed geometric constraints. In detail, GeoAB-Designer consists of a geometry initializer (Geo-Initializer) and a position refiner (Geo-Refiner). Geo-Initializer is a generative model with informative prior, which samples the redundant internal geometries and reconstructs the structures with NeRF (Parsons et al., 2005). Further, Geo-Refiner as GNNs represents the binding site as hierarchical graphs, in which a high-level graph models heterogeneous residue-level relations and a low-level one captures atoms’ interactions considering mechanics for bond lengths, bond angles, and torsion angles. Equivariance is ensured in updating the atom positions. *On mutation-based affinity maturation*, we propose a novel structure-aware GeoAB-Optimizer, based on the proposed network architecture and geometric constraints. It uses amino acid sequences and the context structures as input, and generates both structures of the CDRs and $\Delta\Delta G$, to avoid the problem of domain differences in input CDR structures. Based on the fact that the representations obtained by structure-related tasks usually assist energy-related prediction (Jin et al., 2023), we propose a structure-aware joint training strategy on paired labeled data (with $\Delta\Delta G$ labels) and unpaired ones. Extensive experiments demonstrate the superiority of GEOAB in both realistic generation and reli-

able affinity maturation for antibody CDRs.

2. Background

Notations. For a binding complex composed of an antigen-antibody pair as \mathcal{C} , which contains N_{aa} amino acids, there are N_{ag} amino acids in the antigen and N_{cdr} and N_{ncdr} amino acids of a particular CDR and other non-CDRs in the antibody. We represent their index set by \mathcal{I}_{ag} , \mathcal{I}_{cdr} and $\mathcal{I}_{\text{ncdr}}$ according to the sequential orders, where $|\mathcal{I}_{\text{ab}}| = N_{\text{ab}}$, $|\mathcal{I}_{\text{cdr}}| = N_{\text{cdr}}$ and $|\mathcal{I}_{\text{ncdr}}| = N_{\text{ncdr}}$. For a protein, we consider four backbone atoms for each amino acid, so one amino acid can be represented by its type $a_i \in \{1, \dots, 20\}$ and atom coordinates of $\{\text{N}, \text{C}\alpha, \text{C}, \text{O}\}$ as $(\mathbf{x}_{i,1}, \dots, \mathbf{x}_{i,4})$, where $\mathbf{x}_{i,j} \in \mathbb{R}^3$. Therefore, $\mathcal{C} = \{(a_i, \mathbf{x}_{i,1}, \dots, \mathbf{x}_{i,4})\}_{i=1}^{N_{\text{ag}}+N_{\text{cdr}}+N_{\text{ncdr}}}$ can be split into three sets as $\mathcal{C} = \mathcal{G} \cup \mathcal{R} \cup \mathcal{N}$, according to the index sets of antigen, CDR and non-CDR. For each \mathcal{C} , Gibbs free energy of association is the physical quantity used to measure the binding affinity between antigens and antibodies, denoted by $\Delta G = E(\mathcal{C})$. When mutations occur, $\mathcal{C} = \mathcal{C}^{(\text{wt})}$ as wild types will be changed into $\mathcal{C}^{(\text{mt})}$ as mutant types, with both structures and sequences changed. The mutation effect is usually measured by the change of Gibbs free energy as $\Delta\Delta G = E(\mathcal{C}^{(\text{mt})}) - E(\mathcal{C}^{(\text{wt})})$.

CDR co-design. For CDR co-design, our goal is to establish a model to learn the distribution of CDRs conditioned on the antigens and non-CDRs of the antibodies, *i.e.* $p(\mathcal{R}|\mathcal{G} \cup \mathcal{N})$. For realistic structure design, the internal geometries of the protein structures generated by models should be close to the true ones that are governed by physicochemical rules. For example, the planarity of peptide bonds usually constrains the torsion angle of ‘O=C-N-Cα’ to be 0, and steric collisions between atoms lead torsion angles of ϕ and ψ to fall into defined regions in a graph called the Ramachandran plot (Agnihotry et al., 2022) (Details in Appendix. A.1).

Mutation-based CDR affinity maturation. For affinity maturation, we aim to develop a model that can generate $\mathcal{C}^{(\text{mt})}$ that satisfies $E(\mathcal{C}^{(\text{mt})}) - E(\mathcal{C}^{(\text{wt})}) < 0$, based on $\mathcal{C}^{(\text{wt})}$. Since $E(\cdot)$ is intractable and the mutation data is limited, one key issue is the reliability of direct prediction of $\Delta\Delta G(\mathcal{C}^{(\text{mt})}, \mathcal{C}^{(\text{wt})})$. Otherwise, the maturation is unreasonable given an inaccurate evaluation of the mutation effects. In addition, as a single mutation of amino acid types may affect the structures on a larger scale, it is also important to consider the structure flexibility of mutant types, since their structures are usually unknown. Note that CDRs make a disproportionately large contribution to the binding interactions in the antigen-antibody complexes, so we focus on the mutation effects of points located in the CDRs. Therefore, the problem can be formulated as to develop a model $q(\mathcal{R}^{(\text{mt})} | \mathcal{C}^{(\text{wt})})$, s.t. $\Delta\Delta G(\mathcal{C}^{(\text{mt})}, \mathcal{C}^{(\text{wt})}) < 0$ for $\mathcal{R}^{(\text{mt})} \sim q$, based on our assumption that the structures of non-CDRs and antigens are inflexible.

3. Method

3.1. Graph Construction

We denote the antibody-antigen residue graph as heterogeneous graph $(\mathcal{V}_{\text{aa}}; \mathcal{E}_1, \mathcal{E}_2, \mathcal{E}_3)$, where the nodes are denoted by the residues with C α coordinates $\mathcal{V}_{\text{aa}} = \{(a_i, \mathbf{x}_{i,2})\}_{i=1}^{N_{\text{aa}}}$, and the edge sets are composed of three kinds: sequence edge set \mathcal{E}_1 obtained by whether the two amino acids are consecutive in the protein sequence, i.e. $(i, j) \in \mathcal{E}_1$ if they are consecutive, intra- and inter-structure edge sets \mathcal{E}_2 and \mathcal{E}_3 obtained by whether the two amino acids are close in distance and belong to the same/different proteins. For atom graph $(\mathcal{V}_{\text{at}}; \mathcal{E}_{\text{at}})$, the nodes are represented by its atom types $e_j \in \{1, 2, 3, 4\}$ and positions \mathbf{x}_j as $\mathcal{V}_{\text{at}} = \{(e_j, \mathbf{x}_j)\}_{j=1}^{4N_{\text{aa}}}$, and the edge set \mathcal{E}_{at} is constructed according to the chemical bonds as Figure 1.(a) shows.

3.2. Geo-Refiner

Given the initial states of the backbone atom positions and amino acid types which are often inaccurate, the Geo-Refiner both refines the structures of the backbone atoms and predicts the amino acid types in CDRs. The refinement process is deterministic and includes two steps: residue-level encoding and atom-level updating.

Heterogeneous residue-level encoding. In the residue-level graph $(\mathcal{V}_{\text{aa}}; \mathcal{E}_1, \mathcal{E}_2, \mathcal{E}_3)$, we use a simple heterogeneous GNN to encode the residues. Considering a L -layer heterogeneous residue encoder, the formulation of the l -th equivariant layer can be defined as follows:

$$\begin{aligned} \mathbf{h}_i^{(l+1)} &= \sum_{k=1}^3 \sum_{(i,j) \in \mathcal{E}_k} \text{MLP}_k^{(l)} \left([\mathbf{h}_i^{(l)}, \mathbf{h}_j^{(l)}, d_{i,j}^{(l)}] \right); \\ \mathbf{x}_i^{(l+1)} &= \mathbf{x}_i^{(l)} + \sum_{k=1}^3 \sum_{(i,j) \in \mathcal{E}_k} (\mathbf{x}_i^{(l)} - \mathbf{x}_j^{(l)}) (\mathbf{h}_i^{(l)} \mathbf{W}_k), \end{aligned} \quad (1)$$

for $1 \leq l \leq L$. $\mathbf{x}_i^{(l)}$ is i -th node's updated position of C α atom after $(l-1)$ layers, and $\mathbf{x}_i^{(1)}$ is the initialized C α positions which can be obtained by an initializer (see Sec. 3.3). $\text{MLP}_k^{(l)}$ as a shallow multi-layer perceptron (MLP), encodes the i and j nodes' embeddings and inter distance $d_{i,j}^{(l)} = \|\mathbf{x}_i^{(l)} - \mathbf{x}_j^{(l)}\|_2$ as messages, and summation is used to aggregate messages. \mathbf{W}_k projects embeddings of high dimension into one. $\mathbf{h}_i^{(1)}$ as the input of the GNN, is the initial embedding of residues, which considers amino acid's sequence position i , types a_i , intra-residues' bond lengths, angles and torsion angles, and inter-residues' distances, angles, and dihedrals. $\mathbf{h}_i = \mathbf{h}_i^{(L+1)}$ is the output E(3)-invariant representation, which will be used for updating the atom representations and other downstream tasks.

Mechanics-informed atom-level updating. In the atom-level graph $(\mathcal{V}_{\text{at}}; \mathcal{E}_{\text{at}})$, atom representations are first encoded by $\mathbf{y}_i = \text{Emb}(e_i) + \mathbf{h}_i$, for $1 \leq i \leq 4N$, where $\text{Emb}(\cdot)$ is a simple lookup table, and \mathbf{h}_i is the representation of the residue which the atom belongs to, in Eq. 1. A three-layer GAT (Veličković et al., 2018) is stacked for atom-level message passing, as $\mathbf{z}_i = \text{GAT}(\mathbf{y}_i, \{\mathbf{y}_j\}_{(i,j) \in \mathcal{E}_{\text{at}}})$.

In position updating, we generalize the graph mechanics networks (Huang et al., 2022) to binary, ternary, and quaternary interactions, and harness generalized coordinates to describe the kinematics of the atoms. In specific, for a group of connected atoms as $\mathcal{S} = \{(\mathbf{z}_i, \mathbf{x}_i)\}_{1 \leq i \leq M}$, position and velocity in the generalized coordinates as \mathbf{q}_i and $\dot{\mathbf{q}}_i$, and velocity in global coordinates as $\dot{\mathbf{x}}_i$, the updating process is

$$\begin{aligned} \dot{\mathbf{q}}_i &= f \left(\sum_{j=1}^M \mathbf{z}_j, g(\{\mathbf{x}_j\}_{1 \leq j \leq M}) \right) \dot{\mathbf{q}}_i^{(0)}; \\ \dot{\mathbf{x}}_i &= \text{FK} \left(\{(\mathbf{q}_j^{(1)}, \dot{\mathbf{q}}_j)\}_{1 \leq j \leq M} \right) \end{aligned} \quad (2)$$

where $\mathbf{q}_j^{(1)}$ is the initial atom position in generalized coordinates, $\dot{\mathbf{q}}_i^{(0)}$ is velocity direction, $g(\cdot)$ obtains the representation of invariant intra-geometries in the group, $f : \mathbb{R}^3 \rightarrow \mathbb{R}$ is an arbitrary MLP, and forward kinematics $\text{FK}(\cdot)$ projects the positions in updated generalized coordinates back to the global states. We detail how to update the position states in atom-level interaction layers for the three kinds.

To model the **binary interaction** with $M = 2$, we consider the two connected atoms indexed by (i, j) . The two atoms connected by a chemical bond cannot form a generalized coordinate, and thus we use the global coordinate for updating the positions. The velocity direction $\dot{\mathbf{q}}_i^{(0)}$ is parallel to the bond for scaling the bond length, as $\dot{\mathbf{q}}_i^{(0)} = \frac{\mathbf{x}_i - \mathbf{x}_j}{\|\mathbf{x}_i - \mathbf{x}_j\|_2}$, and $\dot{\mathbf{x}}_i = \dot{\mathbf{q}}_i \cdot g(\mathbf{x}_i, \mathbf{x}_j) = \|\mathbf{x}_i - \mathbf{x}_j\|_2$ is the bond length. Note that (i, j) can be commutative, and the binary interaction is equivalent to vanilla EGNN (Satorras et al., 2021).

To model the **ternary interaction** with $M = 3$, we consider

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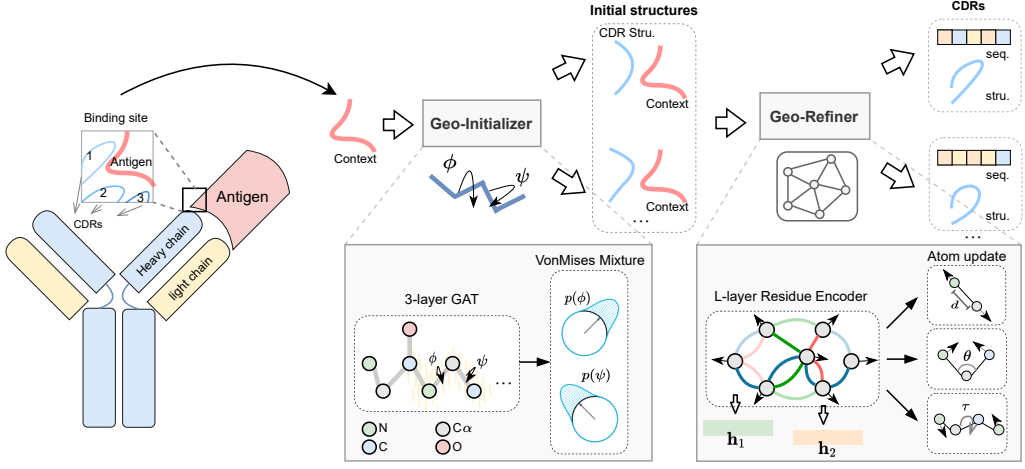


Figure 2: Workflows of **GeoAB-Designer** in antibody design task.

the three consecutively connected atoms indexed by (i, j, k) . To update the bond angle $\theta = \theta_{i,j,k}$, the center atom position \mathbf{x}_j is fixed, and $\dot{\mathbf{q}}_i^{(0)} = \frac{(\mathbf{x}_i - \mathbf{x}_j) \times (\mathbf{x}_k - \mathbf{x}_j)}{\|(\mathbf{x}_i - \mathbf{x}_j) \times (\mathbf{x}_k - \mathbf{x}_j)\|_2}$ represents unit angular velocity. $g(\mathbf{x}_i, \mathbf{x}_j, \mathbf{x}_k) = [\sin(\theta); \cos(\theta)]$ is the angle encoding. The forward kinematics written as $\dot{\mathbf{x}}_i = \mathbf{x}_j + \text{rot}(\dot{\mathbf{q}}_i)(\mathbf{x}_i - \mathbf{x}_j) - \mathbf{x}_i$ model the velocity in the global coordinates, where $\text{rot}(\dot{\mathbf{q}}_i)$ indicates the rotation matrix around the direction of $\dot{\mathbf{q}}_i$ by absolute angle $\|\dot{\mathbf{q}}_i\|$. The dynamic updates for atom k are similar.

To model the **quaternary interaction** with $M = 4$, the four consecutively connected atoms are indexed by (i, j, k, l) . To update the torsion angle $\tau = \tau_{i,j,k,l}$, the atom positions \mathbf{x}_j and \mathbf{x}_k are fixed, and the atoms i and l turn round on the axis $(\mathbf{x}_j - \mathbf{x}_k)$, so that $\dot{\mathbf{q}}_i^{(0)} = \frac{\mathbf{x}_j - \mathbf{x}_k}{\|\mathbf{x}_j - \mathbf{x}_k\|_2}$ and $\dot{\mathbf{q}}_k^{(0)} = -\dot{\mathbf{q}}_i^{(0)}$ are unit torsion angular velocities. $g(\cdot)$ encodes torsion angles by $[\sin|\tau|; \cos|\tau|]$, in which the absolute values of τ are taken to avoid parity (Jing et al., 2023). The FK(\cdot) is written as $\dot{\mathbf{x}}_i = \mathbf{x}_j + \text{rot}(\dot{\mathbf{q}}_i)(\mathbf{x}_i - \mathbf{x}_j) - \mathbf{x}_i$ for atom i and $\dot{\mathbf{x}}_l = \mathbf{x}_k + \text{rot}(\dot{\mathbf{q}}_l)(\mathbf{x}_l - \mathbf{x}_k) - \mathbf{x}_l$ for atom l .

As each atom may belong to several groups, the final updates of the positions in the global coordinates are the summation of all the updates in different groups. Because the binary and quaternary interaction layers are both linear combinations of $\{\mathbf{x}_i\}_{1 \leq i \leq M}$, they are $E(3)$ -equivariant (Villar et al., 2023); The cross product operation in ternary interaction layer leads it to be roto-translational but not reflectional equivariant (Geiger & Smidt, 2022), *i.e.* $SE(3)$ -equivariant. In practice, we find Geo-Refiner does not need multi-round iterations, *i.e.* One-shot prediction achieves comparable performance.

3.3. Geo-Initializer

For antigens and non-CDRs, the structures of ground truth are used as initial states. For CDRs, linear interpolation

is usually used to initialize atom positions (Kong et al., 2023a), where the equispaced atoms are distributed according to the $(\min \mathcal{I}_{\text{cdr}} - 1)$ -th and $(\max \mathcal{I}_{\text{cdr}} + 1)$ -th amino acids' positions. However, the initialization cannot give any informative prior knowledge for realistic structure generation.

To initialize realistic structures of CDRs, we propose Geo-Initializer to directly generate internal geometries. We observed that the redundant geometries in modeling the backbone atoms are three torsion angles of 'N-C α -C-N', 'C α -C-N-C α ' and 'C-N-C α -C' (Appendix A.2). Therefore, we use Von Mises distributions to generate the three torsion angles (Swanson et al., 2023), since it is a continuous probability distribution on the circle with support in $[-\pi, \pi]$. As the multi-modal nature of rotatable bond torsion angles in 'N-C α -C-N' and 'C-N-C α -C' is observed (Appendix B.1), a mixture of K von Mises distributions is employed to capture the modality, as $p(\tau) = \sum_{k=1}^K w_k \frac{e^{\kappa_k \cos(\tau - \mu_k)}}{2\pi I_0(\kappa_k)}$, in which τ is the torsion angle, $I_0(\cdot)$ is the modified Bessel function of order 0, w_k is the weight, μ_k is the mean, and κ_k is the concentration of the i -th distribution. To obtain the parameters of $\{w_k, \mu_k, \kappa_k\}_{k=1}^K$, a shallow GAT is used to encode the atom-level graph to obtain representation for each atom. After summing the representations of the four atoms that make up the torsion angle τ , three shallow MLPs project the summation to K -dimensions, leading to $3 \times K$ predicted parameters in $p(\tau)$. To train the Geo-Initializer, we minimize the negative log-likelihood of the ground truth torsion angle samples for a given rotatable bond under a mixture of K Von Mises distributions defined by the predicted parameters.

Geo-Initializer thus is a generative model from which the redundant torsions of the backbones are sampled. For the rest of the inflexible geometries, the ideal values are used. In this way, NeRF is employed to reconstruct the backbone atom

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positions as the initial structures for further refinements.

3.4. Geometric Constraints

For structure refinement, the outputs of atom-level interaction layers are the backbone atom positions in CDRs in global coordinates, denoted by \mathcal{X} , and the superscripts 0 denote the true values to differentiate the predictions. Inspired by AMBER (Maier et al., 2015), we propose to incorporate the following energy terms as geometry constraints, to govern the generated structure with chemical rules, including constraints on bond length, angles, and torsion angles.

$$\begin{aligned} Loss_{\text{geo}} &= E_{\text{bon}}(\mathcal{X}) + E_{\text{ang}}(\mathcal{X}) + E_{\text{tor}}(\mathcal{X}); \\ E_{\text{bon}}(\mathcal{X}) &= \sum_{(i,j) \in \mathcal{E}_{\text{at}}} (d_{i,j} - d_{i,j}^0)^2; \\ E_{\text{ang}}(\mathcal{X}) &= \sum_{\substack{(i,j) \in \mathcal{E}_{\text{at}}, \\ (j,k) \in \mathcal{E}_{\text{at}}}} (\theta_{i,j,k} - \theta_{i,j,k}^0)^2; \\ E_{\text{tor}}(\mathcal{X}) &= \sum_{\substack{(i,j), (j,k), \\ (k,l) \in \mathcal{E}_{\text{at}}}} (1 - \cos(\tau_{i,j,k,l} - \tau_{i,j,k,l}^0)); \end{aligned} \quad (3)$$

where $d_{i,j}$ is the bond length between i, j , $d_{i,j}^0$ denotes the expected bond length between atom i, j , which can be obtained by the true structures. For $\theta_{i,j,k}$ and $\theta_{i,j,k}^0$, $\tau_{i,j,k,l}$ and $\tau_{i,j,k,l}^0$, the definitions are similar.

Besides, two loss functions are employed: (i) residual-level error on $\text{C}\alpha$ positions and (ii) atom-level error on all backbone atoms. Since the objective region for designing is CDRs, for $i \in \mathcal{I}_{\text{cdr}}$, the loss for position can be written as

$$\begin{aligned} Loss_{\text{pos}} &= \text{Hub} \left(\{\mathbf{x}_i^{(L+1)}\}, \{\mathbf{x}_{i,2}^0\} \right) \\ &\quad + \text{Hub} \left(\{(\mathbf{x}_{i,1}, \dots, \mathbf{x}_{i,4})\}, \{(\mathbf{x}_{i,1}^0, \dots, \mathbf{x}_{i,4}^0)\} \right), \end{aligned} \quad (4)$$

where $\mathbf{x}_i^{(L+1)}$ is the predicted position of residue i , output by L -layer residue-level encoders and defined as $\text{C}\alpha$'s positions in Eq.1, and $\mathbf{x}_{i,2}^0$ is the true $\text{C}\alpha$'s position of residue i ; $(\mathbf{x}_{i,1}, \dots, \mathbf{x}_{i,4})$ is the four backbone atoms of residue i , obtained by atom-level interaction layers, and $(\mathbf{x}_{i,1}^0, \dots, \mathbf{x}_{i,4}^0)$ is that of ground truth. $\text{Hub}(\cdot)$ is the Huber loss used for training stability. The designated position loss aims first to determine the coarse positions of residues and then refine the grained positions of the backbone atoms.

3.5. Antibody CDR Co-Design

In the task of antibody CDR design, we aim to co-design both the amino acid sequence and the backbone atom's structures in CDRs. For sequence design, a masked state as an absorbing type is used to initialize the amino acid types. For structure design, we propose two schemes for the task: (i) **GeoAB-Designer** composed of '(pre-trained) Geo-Initializer + Geo-Refiner' as a generative model due to the stochasticity of Geo-Initializer and (ii) **GeoAB-Refiner**

composed of 'Linear Initialization + Geo-Refiner' as a refinement model. By this means, the loss function is

$$Loss = \alpha_1 Loss_{\text{type}} + \alpha_2 Loss_{\text{pos}} + \alpha_3 Loss_{\text{geo}}. \quad (5)$$

$\{\alpha_j\}_{j=1,2,3}$ are loss weights. $Loss_{\text{type}} = \text{CE}(\{\hat{\mathbf{h}}_i\}, \{a_i^0\})$, in which $\{\hat{\mathbf{h}}_i\}$ are the logits, obtained by a linear classifier stacked after the L -layer residue encoder, projecting $\mathbf{h}_i^{(L)}$ from D -dimension to $\hat{\mathbf{h}}_i \in \mathbb{R}^{20}$, $\{a_i^0\}$ are the true amino acid types in CDRs, and $\text{CE}(\cdot)$ is the cross entropy loss.

Dynamic weights. In **GeoAB-Designer**, the inflexible internal geometries in the initialized backbone structures are almost identical to the real ones (see Appendix. C.3). To preserve the initialized realism of the geometries, we hope that in the training stage, the constraints on internal geometries as $Loss_{\text{geo}}$ can be more emphasized at the beginning, and gradually decreasing losses in positions $Loss_{\text{pos}}$ in the training process can also lead to a further reduction in $Loss_{\text{geo}}$, given that when the model can make perfect predictions of position such that $Loss_{\text{pos}} \rightarrow 0$, $Loss_{\text{geo}}$ is also minimized. Motivated by this, we use a training trick of dynamic loss weights, by setting α_3 large at the beginning and gradually making it smaller with exponential moving average (See Appendix. B.2), to keep the geometry from deviating too much from the initial state during training.

3.6. Mutation-Based Affinity Maturation

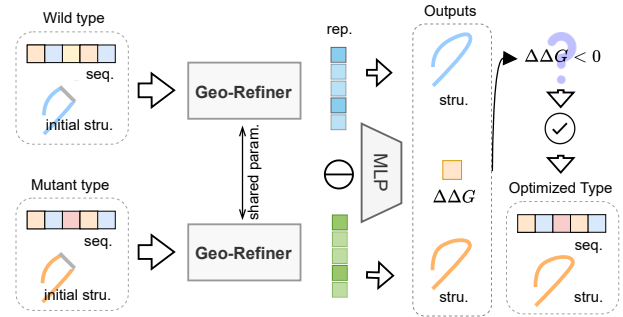


Figure 3: Workflows of **GeoAB-Optimizer** in antibody affinity maturation task.

For affinity maturation, two sequences of amino acids are given, *the wild type* and *the mutant type*. The reliable maturation requires our model to accurately predict both the structures and the mutation effect $\Delta\Delta G$. To tackle the problem, we propose an architecture composed of twin Geo-Refiners with shared parameters and a joint training strategy to make full use of the paired labeled and unpaired data.

In detail, we first assume that the mutant amino acid will affect the surrounding structures around it, but keep the rest unchanged, so we aim to predict the structure of the surrounding N_{flx} residues. A simple 'Linear Initialization'

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initializes the backbone atoms in the affected regions. For an instance of *paired data* with two types of sequence, the initial amino acid sequences as inputs are the true lists of types, and we write the index set of mutant points as \mathcal{I}_{mt} . Two Geo-Refiners with shared parameters will generate the backbone structure of two types in the affected regions by the atom-level interaction layers. Besides, following (Shan et al., 2022), $\Delta\Delta G$ is predicted by

$$\Delta\Delta G = \text{Proj} \left(\frac{1}{|\mathcal{I}_{\text{mt}}|} \sum_{i \in \mathcal{I}_{\text{mt}}} (\mathbf{h}_i^{(\text{mt})} - \mathbf{h}_i^{(\text{wt})}) \right), \quad (6)$$

in which $\text{Proj}(\cdot)$ is an MLP, projecting D -dimensional representation to a scalar. In this way, the loss function reads

$$\text{Loss} = \alpha_1 \text{Loss}_{\text{ddg}} + \alpha_2 \text{Loss}_{\text{pos}} + \alpha_3 \text{Loss}_{\text{geo}}, \quad (7)$$

where $\text{Loss}_{\text{ddg}} = \text{MSE}(\Delta\Delta G, \Delta\Delta G^{(0)})$, and $\Delta\Delta G^{(0)}$ is the labeled change in Gibbs free energy. Loss_{pos} and Loss_{geo} are usually calculated with the structures of wild types since the structures of the mutant ones are usually unavailable. For an instance of *unpaired data* with a single type of sequence that is usually defined as wild type, we select a point of interest and mask its surrounding N_{flx} residuals structure by the same initialization method. The mutant type is defined as itself, and the loss on mutation effect will change into $\text{Loss}_{\text{ddg}} = \text{MSE}(\Delta\Delta G, 0)$.

Once the model is trained, we can obtain the mutant sequences and structures such that $\Delta\Delta G < 0$ by enumerating the other 19 amino acids and replacing the point of interest.

4. Related Work

Antibody Design. Classical works are usually based on hand-crafted potential functions, including physical force-field terms (Li et al., 2014; Lapidoth et al., 2015; Adolf-Bryfogle et al., 2018) and statistical potential (Min-yi & Andrej, 2006). This results in intensive computation and unavoidable inaccuracy, as complex mechanisms of protein structure cannot be described by simple potentials. Deep-learning-based methods concentrated on 1D sequence design (Alley et al., 2019; Liu et al., 2020; Saka et al., 2021; Akbar et al., 2022), and then, thanks to great progress in graph neural networks (Wu et al., 2021; Liu et al., 2021; Wu et al., 2023) and protein modeling (Huang et al., 2023; Wu et al., 2024; Tan et al., 2024; 2023), more works have been focused on structure-sequence co-design. For example, RefineGNN (Jin et al., 2021) auto-regressively generate the amino-acid types and structures; HERN (Jin et al., 2022) achieves co-design and docking tasks by using a hierarchical graph; MEAN (Kong et al., 2023a) employ multi-channel equivariant attention networks to generate CDRs with multi-round refinements; DyMEAN (Kong et al., 2023b) similarly can fulfill the co-design tasks, and simultaneously dock to

the epitope at a full-atom level. DiffAB as diffusion models generates translation, orientation, and type variables (Luo et al., 2022). HTP (Wu & Li, 2023) and ABGNN (Gao et al., 2023) focus on antibody pretraining. These methods hardly generate antibodies that conform to physicochemical constraints, while GeoAB focuses on realistic generation.

Mutation Effect Prediction. Similarly, traditional approaches utilize energy functions to model the interactions and compute the $\Delta\Delta G$ (Steinegger & Söding, 2017). Statistical methods rely on feature engineering and use the invariant features and statistical learning methods to predict mutation effects (Lei et al., 2023). In deep learning, the effectiveness in predicting $\Delta\Delta G$ is validated. For example, ESM-1v (Meier et al., 2021) proposes sequence-based pretraining tasks and predicts mutation effects on protein functions. DDG-Predictor (Shan et al., 2022) takes structures and sequences of both types as input, and predicts the $\Delta\Delta G$ effectively. However, the structures of mutant types are usually unknown, RDE-PPI (Luo et al., 2023) avoids the problem and uses a structure-aware pretrained network on side-chain rotamers to assist the prediction. Besides, there are also works on designing pretraining tasks for $\Delta\Delta G$ predictions (Hsu et al., 2022; Yang et al., 2023; Cai et al., 2023). In comparison, GEOAB also considers mutation effects attributing to structural flexibility but uses joint training strategies instead of pretraining paradigms.

5. Experiment

5.1. Antibody CDR Co-Design

Baselines. We select 6 methods as benchmarks. For refinement methods, **RefineGNN**, **MEAN** and **DyMEAN** are used. Their variants of **C-RefineGNN** and **C-DyMEAN** are included, meaning that the two methods use the same contextual condition. RefineGNN only models the heavy chains with less context and DyMEAN is designed for both docking and co-designing with the unknown paratope. For generative methods, we choose **DiffAB** as a baseline, which generates translation, orientation, and type variables, with the intra-residue geometries fixed with ideal values. As discussed in Sec. 3.5, we propose two variants of GeoAB. The first is **GeoAB-Refiner** in comparison with the refinement methods; the second is **GeoAB-Designer**, as a generative model compared with DiffAB. For **GeoAB**, our model is open to the public through <https://github.com/Edapinenut/GeoAB>.

Setup. Following (Kong et al., 2023a), we use the SAbDab (Dunbar et al., 2014) with complete antibody-antigen structures for training. IMGT scheme is used to renumber the complexes (Lefranc et al., 2003). The splits of training, validation and test sets are according to the clustering of CDRs via MMSeqs2 (Steinegger & Söding, 2017). After the 10 cross-validations on SAbDab, we benchmark all methods

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Table 1: Metrics on generated CDR-H3 for RAbD compared with the reference. ‘GeoAB-R’ is the refinement variant and ‘GeoAB-D’ is the generative one. Values in **bold** are the best metrics.

Method	AAR(\uparrow)	A-RMSD(\downarrow)	UA-RMSD(\downarrow)	Iddt(\uparrow)	TM(\uparrow)
RefineGNN	0.2965	3.6622	7.5417	0.8415	0.8312
C-RefineGNN	0.2990	3.3124	7.2014	0.8387	0.8320
MEAN	0.3706	1.5764	1.7866	0.9814	0.9819
DyMEAN	0.3926	1.6970	10.9927	0.9540	0.9617
C-DyMEAN	0.3928	1.5656	1.7641	0.9821	0.9824
GeoAB-R	0.4053	<u>1.3978</u>	<u>1.5852</u>	0.9860	0.9858
	20%	0.2771	5.9337	9.4393	0.7029
DiffAB	50%	0.2678	5.9528	9.6680	0.7081
	100%	0.2693	5.9372	9.8248	0.6953
	20%	0.3939	1.4027	1.5038	0.9848
GeoAB-D	50%	0.3940	1.4119	1.5488	0.9844
	100%	0.3869	1.4217	1.6143	0.9857

Table 2: Metrics on flexible and inflexible geometries obtained by generated CDR-H3 v.s. reference. Values in **bold** are the best metrics, and in underline are the second. The full comparison is given in Appendix C.3

Geometry	JSD(\downarrow)					MAE/MCE(\downarrow)				
	MEAN	C-DyMEAN	DiffAB	GeoAB-R	GeoAB-D	MEAN	C-DyMEAN	DiffAB	GeoAB-R	GeoAB-D
C α -C	0.7318	0.7673	0.1582	0.5286	<u>0.5084</u>	0.3209	0.5310	0.0298	0.2182	<u>0.1883</u>
C-N	0.6720	0.7350	0.6537	<u>0.5916</u>	0.5477	0.3824	0.4784	0.6202	<u>0.3201</u>	0.2537
N-C α -C	0.6614	0.5646	0.3207	0.5354	<u>0.5309</u>	0.2512	<u>0.1646</u>	0.1239	0.1853	0.2207
C α -C-N	0.6819	0.7583	0.6108	<u>0.6031</u>	0.5628	0.2368	0.5441	0.2181	0.1479	<u>0.1908</u>
O=C-N-C α	0.6015	0.6670	0.4354	<u>0.4193</u>	0.3657	0.7799	0.9813	0.8524	<u>0.3158</u>	0.2992
C-N-C α -C	0.6832	0.3506	0.4992	0.3180	<u>0.3250</u>	0.5611	0.9952	0.7277	0.3313	0.3634

with the 60 diverse complexes as RAbD (Adolf-Bryfogle et al., 2018), and give comparisons. The training is still conducted on the SAbDab dataset, but we eliminate all antibodies from SAbDab whose CDR-H3s share the same cluster as those in RAbD to avoid potential data leakage. We use AAR metrics to evaluate amino-acid recovery ratio; A-RMSD and UA-RMSD as the RMSD between C α with/without Kabsch alignment; Iddt and TM-score measuring global structural similarities. For the generative methods, group comparisons are made by selecting percentile of 20%, 50%, and 100% according to UA-RMSD. For details see Appendix C.1.

Results. Table 1 gives results on RAbD CDR-H3 designing, and for 10-fold cross-validation on SabDab, it is given in Appendix C.2. It shows that in refinement models, GeoAB-R achieves the overall best performance, especially in structure evaluation, with 10.71% and 10.02% improvements in A-RMSD and UA-RMSD respectively over state-of-the-art C-DyMEAN. In generative models, GeoAB-D also outperforms DiffAB in all aspects by a large margin.

Geometry analysis. Since GeoAB aims to generate realistic structures, we analyze internal geometries. We use Jensen–Shannon divergence (JSD) to evaluate the distribution differences and use mean absolute error (MAE) and mean cosine error (MCE) as two metrics to measure the instance differences in lengths and angles, respectively. In specific, MCE is written as $MCE(\theta^0, \theta) = 1 - \cos(\theta^0 - \theta)$, reaching 0 when $\theta = \theta^0 \pmod{2\pi}$. Table 2 gives the comparison, including inflexible geometries like bond lengths ‘C α -C’ and ‘C-N’, bond angles ‘N-C α -C’ and ‘C α -C-N’,

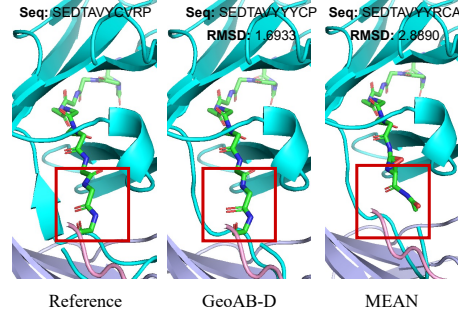


Figure 4: Backbone atoms in CDR-H3 (shown in sticks) on PDB-3nid, of the reference and designed by GeoAB and MEAN. RMSDs are calculated with all backbone atoms.

and torsion angles ‘O=C-N-C α ’ and flexible torsions ‘C-N-C α -C’. GeoAB-D generates the most realistic internal geometries in overall evaluations. For inflexible geometries, because DiffAB regards the residues as rigid bodies, the intra-residue geometries are set as ideal values, leading to the least deviation from references in ‘C α -C’ and ‘N-C α -C’. However, in inter-residue geometries such as ‘C-N’ and ‘C-N-C α -C’, GeoAB shows superiority over DiffAB. Further, GeoAB-D as a generative model, initialized by the prior torsion angle initializer, outperforms GeoAB-R in JSD, but is less competitive in MAE/MCE due to its stochasticity. Figure 1 shows the distribution gap between GeoAB-R and MEAN; Figure 4 gives an example to show that GeoAB-D generates more realistic CDR-H3 structures than MEAN.

5.2. Mutation-Based Remodeling

Baselines. We compare our method with 4 baseline $\Delta\Delta G$ -predictors, including **Rosetta** ddG (Alford et al., 2017; Leman et al., 2020), **FoldX** (Delgado et al., 2019), **DDG-Pred** and **RDE-PPI**. The first two are classical energy-based, and the last two are structure-based deep models. Note that all these methods are not pretrained, and **RDE-PPI** is tested as an unpretrained version for fair comparisons.

Performance Evaluation. Following (Luo et al., 2023), we use **SKEMPI2** (Jankauskaitė et al., 2019) as the evaluation datasets. The datasets are split into three folds by structure, in which two of them are used for training and validation, and the rest are used for testing. Cross-validation is evaluated for $\Delta\Delta G$ -predictors. Four metrics including Pearson

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Table 3: Metrics for cross-validation evaluation of $\Delta\Delta G$ prediction on SKEMEPI2 dataset. Results of the methods with ‘*’ are from RDE-PPI (Luo et al., 2023). Values in **bold** are the best metrics, and in underline are the second.

	Per-Structure		Overall			
	PCC(↑)	SRCC(↑)	PCC(↑)	SRCC(↑)	RMSE (↓)	MAE (↓)
Rosetta*	0.3284	0.2988	0.3113	0.3468	1.6173	1.1311
FoldX*	0.3789	0.3693	0.3120	0.4071	1.9080	1.3089
RDE-PPI	<u>0.3924</u>	0.3529	<u>0.6368</u>	0.4761	1.6071	1.1492
DDG-Pred*	0.3750	0.3407	0.6580	0.4687	<u>1.4998</u>	<u>1.0821</u>
GeoAB	0.3936	<u>0.3610</u>	0.6324	0.4972	1.4758	1.0611

correlation coefficient (PCC), Spearman’s rank correlation coefficient (SRCC), RMSE, and MAE are used to measure the accuracy of predictions. Besides, the per-structure PCC and SRCC are calculated in each structure of the protein complex. Table. 3 shows that GeoAB achieves competitive performance in the $\Delta\Delta G$ prediction tasks. In addition, GeoAB does not require the structures surrounding the mutant points as input, and predicts both the structures and $\Delta\Delta G$, so the domain difference problems resulting from the unavailability of mutant types’ structures are avoided.

Affinity Maturation. After ensuring our models’ ability of reliably predicting $\Delta\Delta G$, we retrained it with the joint training strategy. In detail, we divide the SKEMEPI2 datasets into two sets, where the test instances are the antibody-antigen complexes that appear in the RAbD, and the rest of SKEMEPI2 as *paired data* together with the training instances in SAbDab as *unpaired data* make up of the training set. PDB-REDO (Joosten et al., 2014) are also used as *unpaired data* included in the training set. For training instance in SKEMEPI2, we set $N_{\text{flx}} \sim U(11, 15)$ which mimics the length of CDR-H3 regions and masks the structures of the N_{flx} residues surrounding the mutant points by linear interpolation initialization, and Geo-Refiner is used to predict the masked structures and $\Delta\Delta G$ as discussed in Sec. 3.6. For the training instance in SAbDab, the masked region is CDR-H3, and the mutant types are regarded the same as the wild types, leading to $\Delta\Delta G = 0$. For PDB-REDO, we randomly select one residue on a chain and mask the surrounding N_{flx} residues’ structures for prediction, utilizing more data to enable the model to perceive the local structures.

Maturation Results. Table. 4 gives performance on the accuracy of predicting $\Delta\Delta G$ and structures on the test set on SKEMEPI2. For affinity maturation tasks, we conduct the optimizing processes as shown in Figure. 3, by randomly choosing 1 or 2 points in the CDR-H3 regions and enumerating the sequences as the model inputs 400 times for each antibody-antigen complex. The sequences and the predicted structures with the lowest $\Delta\Delta G$ predictions are selected as the optimized samples with affinity maturation. We also give the average $\Delta\Delta G$ of the maturation results on the 60 complexes in Table. 4, with results of ITA with GeoAB-Refiner also provided. Figure. 5 gives a sample optimized by **GeoAB-Optimizer**, and an unrealistic sample optimized

Table 4: The results for structure and $\Delta\Delta G$ prediction on SKEMEPI2 test set and affinity maturation on RAbD CDR-H3.

	SKEMEPI2	RAbD
A-RMSD	3.4733	1.5982
UA-RMSD	2.6785	1.4006
Per-Stru	PCC SRCC	0.3977 /
Overall	PCC SRCC	0.6657 /
Optimized $\Delta\Delta G$	/	-3.1598
Optimized $\Delta\Delta G$ (ITA)	/	-7.2816

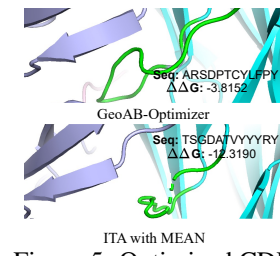


Figure 5: Optimized CDR-H3s for PDB-4u6h.

by MEAN with ITA, where $\Delta\Delta G$ is predicted by DDG-Pred in ITA. It shows that the training set of $\Delta\Delta G$ -predictor (generated by Rosetta) is of great difference from the testing instance (generated by MEAN), so it gives a low $\Delta\Delta G$ even if the generated structure is very unrealistic (broken chains in CDRs). In comparison, our **GeoAB-Optimizer** avoids the problem because it does not need the mutant type’s structures as input. Besides, the detailed results of ITA are given in Appendix. C.4 for fair comparisons.

6. Analysis

We test if each proposed technique is necessary in GeoAB. We conduct ablation studies on SAbDab training and RAbD testing following the *setup in Table. 1*. In Table. 5, ‘w/o Bin/Ter/Quail’ means GeoAB-R without Binary or Ternary or Quaternary interaction layers for position update; ‘w/o Bon/Ang/TorGC’ means GeoAB-R without Bond length or Angle or Torsion geometric constraints; ‘w/o DyLW’ means training GeoAB with or without dynamic loss weight tricks; Besides, we also conduct experiments on the effects of ESM pretraining embedding, as ‘w/o ESM’. We choose representative metrics including AAR, UA-RMSD, and JSD in ‘C α -C’, ‘C α -C-N’, and ‘C-N-C α -C’ to illustrate how these techniques affect the designation. It can be concluded from Table. 5 that (1) The interaction layers help to improve UA-RMSE through a better prediction on flexible torsion angle of ‘C-N-C α -C’ except that ‘TerIL’ contributes little; (2) Realistic inflexible geometries are usually generated by models with corresponding geometric constraints, according to ‘w/o BonGC’ and ‘w/o AngGC’. ‘TorGC’ also benefits the torsion angle prediction; (3) ESM embedding brings little improvements to the co-design tasks; (4) ‘DyLW’ helps GeoAB-G to generate more realistic inflexible geometries, since the JSDs are close to GeoAB-R when it is removed.

7. Conclusion and Limitation

A method called GeoAB is proposed, which focuses on realistic CDR structure design and reliable affinity maturation. Two models are established as solutions. GeoAB-Designer is able to generate realistic CDR structures; GeoAB-Optimizer used for affinity maturation is able to predict $\Delta\Delta G$ accurately and generate CDR structures, avoiding

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Table 5: Ablation on proposed techniques. Geometries like ‘C α -C’ are JSDs of them. Values in **bold** are the best, and values in underline deviate from GeoAB-R obviously.

Ablation	AAR	UA-RMSD	C α -C	C α -C-N	C-N-C α -C
GeoAB-R	0.4053	1.5852	0.5286	0.6031	0.3180
w/o BinIL	0.3994	<u>1.6695</u>	0.5414	0.5986	<u>0.3786</u>
w/o TerIL	0.3893	1.6101	0.5370	0.6133	0.3382
w/o QualIL	0.3849	<u>1.6779</u>	0.5251	0.6022	<u>0.4274</u>
w/o BonGC	0.3990	1.6134	0.6136	0.6357	0.4412
w/o AngGC	0.4016	1.6073	0.5427	<u>0.6941</u>	0.3437
w/o TorGC	0.4019	<u>1.6477</u>	0.5376	0.5801	<u>0.5696</u>
w/o ESM	0.4096	1.5824	0.5297	0.6118	<u>0.3177</u>
w/o DyLW	0.3814	1.6231	0.5428	0.6152	0.3218

the problem of domain differences in input structures.

Still, limitations exist. First, the proposed interaction heads require more computational complexity than the vanilla EGNN, for each requires extra $\mathcal{O}(m)$ updates on atoms’ positions, for a CDR composed of m corresponding groups. Secondly, the accuracy of prediction $\Delta\Delta G$ is not as good as the models with structure-aware pretraining paradigms like RED-PPI, which will be our future concentration.

Acknowledgements

This work was supported by the Science & Technology Innovation 2030 Major Program Project No. 2021ZD0150100, National Natural Science Foundation of China Project No. U21A20427, Project No. WU2022A009 from the Center of Synthetic Biology and Integrated Bioengineering of Westlake University, and Project No. WU2023C019 from the Westlake University Industries of the Future Research. Finally, we thank the Westlake University HPC Center for providing part of the computational resources. Besides, we thank the help of Dr. Tailin Wu, and his great efforts in his deep insights into cutting-edge issues, the guidance provided in the rebuttal, and the funding that supplied us with the necessary equipment for our research.

Impact Statement

The world is revolutionizing itself in one epidemic after another. With the outbreak and long-term continuation of COVID-19, more and more AI scientists are beginning to turn their research interests to drug design. The focus of this paper is on one of them – antibody drugs. There is some work aimed at using AI algorithms to design and optimize antibodies. However, antibodies are a type of protein and few work focuses on the quality of their generation. Here, the present work presents a considerable contribution to improving both the quality of antibody generation and the reliability of optimization. We emphasize the social significance of the work here and hope to get the attention of the reviewers and the review committee.

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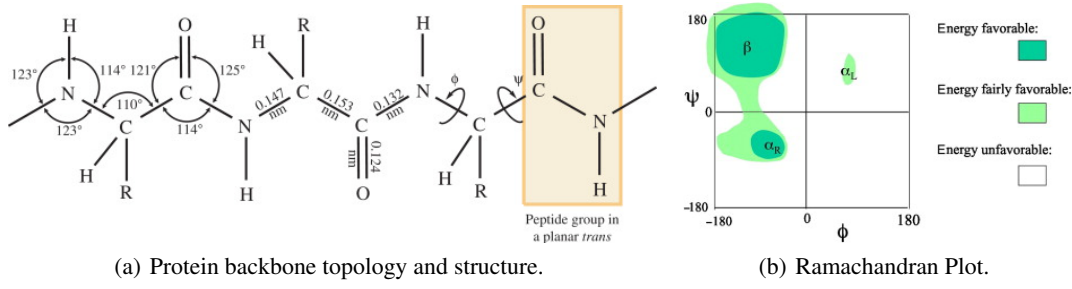
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(a) Protein backbone topology and structure. (b) Ramachandran Plot.

Figure 6: Illustration of peptide planar and the redundant geometry.

A. Protein Backbone Geometry Analysis

A.1. Flexible and Inflexible Geometries

The topology of a part of proteins consists of residues, which are shown in Figure 6(a) (Lieberman, 2014). The bond lengths and bond angles are ideally inflexible and governed by physicochemical laws, with the ideal values shown in Figure 6(a). However, in molecules, the torsion angles are usually flexible, leading to the conformation flexibility of biomolecules. In the protein backbones, there are four main torsion angles, including ϕ as 'C-N-C α -C', ψ as 'N-C α -C-N', ω as 'C α -C-N-C α ', and 'C-N-C α =O'. Besides, 'C-N' as the peptide bond, usually has two states: trans, $\omega \approx 180^\circ$, and cis, $\omega \approx 0^\circ$. In the trans configuration, the two alpha carbon atoms of the connected amino acids are on the opposite sides of the peptide bond, whereas in cis configuration they are on the same side of the peptide bond. In most cases, the peptide bonds in proteins are trans. This preference can be explained by the steric clashes that occur between groups attached to the alpha carbon atoms in cis form, which hinder the formation of this configuration. G.N. Ramachandran recognized that steric collisions between atoms prevent some combination of ϕ and ψ angles and, for the trans configuration, ranges of ϕ and ψ angles fall into defined regions in a graph called the Ramachandran plot as Figure 6(b) shows. According to permitted ϕ ψ and ω angles, preferred conformations of the main chain lead to recurrent structures in proteins namely alpha helix, beta sheets, and turns (Cacabelos et al., 2014). Besides, due to the peptide planar, the 'C α -C-N-C α ' torsion angles are also restricted as 0° .

In this way, we can conclude that the flexibility of the backbone is mainly determined by the fluctuations in torsion angles, specifically ϕ and ψ , and the four flexible torsion angles degenerate into two. However, in real-world observation, ω are not strictly 0° or 180° , generating a deviation of $\pm 7^\circ$ from ideal values, which is also redundant geometry in accurate backbone generation tasks (MacArthur & Thornton, 1996). By this means, to generate a backbone, one can establish a generative model on the three torsion angles, such as FoldingDiff (Wu et al., 2022). Together, with the ideal values of inflexible geometries and generated redundant torsions, the protein backbone structures can be reconstructed by NeRF (Parsons et al., 2005), by sequentially placing the atom in the local coordinate systems, where the positions are determined by the length as distance, angles and dihedrals in the spherical coordinate systems.

A.2. Analysis on SAbDab CDRs

Beyond the general proteins, we now focus on the antibodies' CDRs structures, to figure out if the highly flexible regions follow the same rules as the former conclusions.

We first intercept the CDRs of all antibodies in the training set, and obtain empirical and kernel density estimated distributions of the chemical bond lengths, bond angles, and torsion angles of individual fragments, in Figure 7.

It shows that the internal geometries basically follow the conclusions obtained in the last section. Distributions of ϕ , ψ and ω show the highest stds among all the 12 geometries, and all the ω torsion is trans. Besides, we give the statistical values on MAE between the ideal values and the observed ones and stds of each distribution, to further prove the conclusions in Table 6. The same conclusion can be reached.

B. Method

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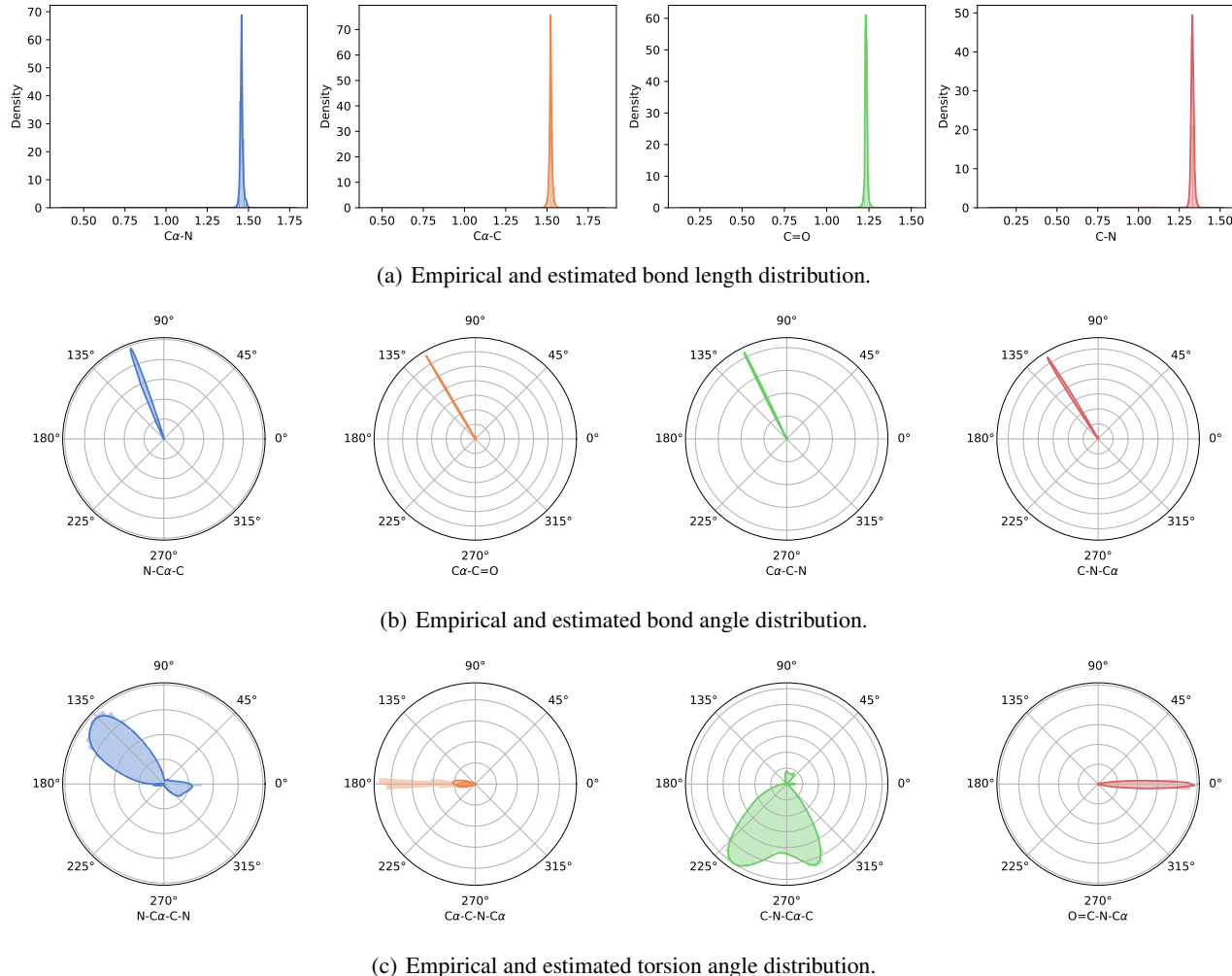


Figure 7: Distributions of flexible and inflexible geometries obtained by CDRs in SabDab datasets.

Table 6: Statistics of flexible and inflexible geometries.

Geometry	Cα-N(Å)	Cα-C(Å)	C=O(Å)	C-N(Å)	N-Cα-C(°)	Cα-C=O(°)	Cα-C-N(°)	C-N-Cα(°)	Cα-C-N-Cα(°)	O=C-N-Cα(°)
Ideal Value	1.32	1.53	1.24	1.47	114	121	110	123	180	0
MAE	0.0158	0.0128	0.0111	0.0851	2.7286	2.9057	5.0941	4.5620	7.6987	3.6468
Std	0.0108	0.0107	0.0086	0.0852	2.6354	2.8098	4.8009	6.4211	5.8353	2.9836

B.1. Geo-Initializer

The multi-modality of the redundant torsions is shown in Figure. 7(c), with ‘N-C α -C-N’ and ‘C-N-C α -C’ demonstrating 3 to 4 peaks in the estimated distributions. In this mean, we select $K = 4$ in the von Mises mixtures. Besides, the encoders mapping the atom-level graph ($\mathcal{V}_{at}; \mathcal{E}_{at}$) is a three-layer GAT in implementation. Since the edge sets are composed of the bonds, the inter-chain messages cannot be passed. Therefore, the Geo-Initializer focuses more on the single-chain structure, leading the RMSDs to high values because of insufficient contextual conditions.

B.2. Dynamic Weights

The weight of $Loss_{geo}$ as α_3 is changed gradually with exponential moving average updating (EMA). For the α_3 in t -th iteration, it reads

$$\alpha_3^{(t)} = \max\{\exp(-\beta t)\alpha_3^{(1)} + \gamma\alpha_3^{(t-1)}, \alpha_{\min}\}, \quad (8)$$

In which α_{\min} is set as the minimum bond of the weight, t is the training iteration, γ and β are two decayed coefficients, which are hyper-parameters, set as 0.999 and 0.9999 in practice.

C. Experiments

C.1. Hyper-parameters

Heterogeneous residue-level encoder is parameterized as 9 layers of heterogeneous GNNs. In each layer, the MLP is constructed by ‘Linear + SiLu + Linear’, with Dropout probability equaling 0.1 to avoid over-fitness. The embedding dim is set as 128.

Equivariant atom-level interaction layers are composed of a 3-layers GAT. For each, the function f is parameterized by a three-layer MLP.

Training weights. α_1, α_2 are set as 1.0 and 0.8 respectively for co-design models. α_3 is set as 0.4 in GeoAB-R and dynamic as Appendix. B.2 discussed. For $\Delta\Delta G$ prediction, α_1 is set as 1.0, and α_2 and α_3 are set as 0.4, because we hope the model can focus more on predicting accuracy and perceiving the structures gradually.

Training parameters. The learning rate lr is $5e - 4$. In all training, the max training epoch is 20. LambdaLR schedule is used, with lr_lambda is set as $0.95 \times lr$.

Epitope selection. In our main experiments on antibody CDR design, we select the 48 residues of the antigen closest to the antibody in terms of the C α distance as the epitope like (Jin et al., 2021).

C.2. 10-Cross Validation Results on SAbDab

Here we give the 10-CV evaluation results in SAbDab to show that GeoAB achieves the state-of-the-art performance in Table 8.

Table 7: Results of $\Delta\Delta G$ optimized by ITA with different methods.

Region	$\Delta\Delta G$ - H3
Random	1.25
RefineGNN	-3.41
C-RefineGNN	-3.66
Mean	-5.79
C-Dymean	-6.56
GeoAB-R	-7.28

Table 8: Results of 10-cross validation on different baselines. GeoAB-R and GeoAB-D reach overall the best performance.

Methods	CDR-H1		CDR-H2		CDR-H3	
	AAR	RMSD	AAR	RMSD	AAR	RMSD
RefineGNN	39.40	3.22	37.06	3.64	21.13	6.00
C-RefineGNN	33.19	3.25	33.53	3.69	18.88	6.22
C-HERN	47.36	1.45	41.45	1.20	28.47	2.64
MEAN	58.29	0.98	47.15	0.95	36.38	2.21
C-DyMEAN	61.13	0.89	53.14	0.87	37.09	2.09
GeoAB-R	64.87	0.82	56.09	0.74	37.58	1.94
GeoAB-D	64.03	0.84	54.98	0.75	37.09	1.97

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Table 9: JSDs of the compared methods for different geometries *v.s.* the reference. GeoAB-I is the Geo-initializer.

JSD	C-RefineGNN	Mean	C-Dymean	GeoAB-R	Diffab	GeoAB-G	GeoAB-I
N-CA	1.6648	0.7318	0.7673	0.5286	0.1582	0.5084	0.1314
CA-C	1.5251	0.7579	0.7635	0.5359	0.2252	0.4949	0.1487
C=O	1.6801	0.7130	0.7509	0.5744	0.1289	0.6834	0.1106
C-N	1.5546	0.6720	0.7325	0.5916	0.6537	0.5477	0.1465
N-CA-C	0.8325	0.6614	0.5646	0.5354	0.3207	0.5309	0.0907
CA-C=O	0.8280	0.7209	0.7019	0.6767	0.2693	0.6456	0.1114
C-N-CA	0.8325	0.6468	0.7496	0.5574	0.5624	0.6123	0.0876
CA-C-N	0.8326	0.6819	0.7583	0.6031	0.6108	0.5628	0.0904
O=C-N	0.6721	0.7075	0.7382	0.6128	0.6384	0.5749	0.1451
N-CA-C-N	0.3892	0.3121	0.4230	0.2717	0.4249	0.2470	0.2289
CA-C-N-CA	0.4566	0.5231	0.6704	0.2994	0.4278	0.3011	0.2061
C-N-CA-C	0.5468	0.6832	0.3506	0.3180	0.4992	0.3250	0.1980
O=C-N-CA	0.7352	0.6015	0.6670	0.4193	0.4354	0.3657	0.2194

C.3. Complete Analysis on Internal Geometries

We give the complete analysis of the generated internal geometries in Table. 9.

C.4. ITA Results of GeoAB-R

Here we also conduct experiments on ITA with GeoAB-R, following the protocol of MEAN (Kong et al., 2023a). To ensure the expected generalizability, we select a total of 53 antibodies from SKEMPI V2.0 training set for affinity optimization and split SAbDab into training and validation sets in a ratio of 9:1 for pretraining the model. The results of ITA for antibody optimization are given in Table. 7, following the protocol of MEAN (Kong et al., 2023a).

C.5. Computational Comparison

We have conducted a comparison of our models with others on the CDR3 co-design task to address this concern. ‘Batch_size’ is set to 8, layer number is 6 and the embed size is all set to 128 for a fair comparison. Table. 10. gives the complexity analysis. We can conclude that: In comparison to MEAN, which costs the least computational resources, our model has comparable GPU memory cost and training time, especially in GeoAB-R, because MEAN takes multi-head attention as its graph updating modules, while GeoAB-R uses a faster and simpler MLP-based heterogenous GNN. However, in the atom-updating layer, it is the key to increasing the training time because of a larger parameter number. For GeoAB-D, the initialization takes more time than Linear Initialization in GeoAB-R, which leads to more time in training and inference. However, since the initializer is pertained, so the trainable parameter number and GPU memory cost are of tiny significance compared with GeoAB-R. Note that DiffAB as a diffusion-based method, usually requires more training epochs and the iterative denoising process leads the inference time much longer than others.

Methods	GPU Memory (MiB)	Inference Time (s/100samples)	Training Time (s/epoch)	Paramater Number
MEAN	5594	14	81	1349515
DyMEAN	16122	48	201	2250270
DiffAB	19041	212	124	3995270
GeoAB-R	5934	18	96	1984663
GeoAB-D	6033	25	154	1984663

Table 10: Complexity comparison of the included models.