IgBlend: Unifying 3D Structures and Sequences in Antibody Language Models

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Abstract

Large language models (LLMs) trained on antibody sequences have shown significant potential in the rapidly advancing field of machine learning-assisted antibody engineering and drug discovery. However, current state-of-the-art antibody LLMs often overlook structural information, which could enable the model to more effectively learn the functional properties of antibodies by providing richer, more informative data. In response to this limitation, we introduce IgBlend, which integrates both the 3D coordinates of backbone atoms (C-alpha, N, and C) and antibody sequences. Our model is trained on a diverse dataset containing over 4 million unique structures and more than 200 million unique sequences, including heavy and light chains as well as nanobodies. We rigorously evaluate IgBlend using established benchmarks such as sequence recovery, complementarity-determining region (CDR) editing and inverse folding and demonstrate that IgBlend consistently outperforms current state-of-the-art models across all benchmarks. Furthermore, experimental validation shows that the model's log probabilities correlate well with measured binding affinities.

1 Introduction

Antibodies are key components of the adaptive immune system, capable of recognizing and neutralizing a wide range of pathogens, including viruses, bacteria, and other foreign invaders. Their ability to bind specific targets with high affinity makes them essential tools in therapeutic development. Recent advancements in natural language processing (NLP) have led to the creation of foundational language models that can learn from and modify antibody sequences [\[Olsen et al., 2022b,](#page-9-0) [2024,](#page-10-0) [Prihoda et al., 2022\]](#page-10-1). Moreover, the three-dimensional (3D) structure of an antibody is closely linked to its specificity, affinity, and interaction with antigens. Therefore, capturing the relationship between sequence and structure is crucial for tasks such as affinity maturation, de novo antibody design, and optimizing antibody-antigen interactions for therapeutic applications. While current language models excel at either sequence-to-sequence or structure-to-sequence (inverse folding) tasks, relying on only one of these modalities at the input limits their capability and flexibility in more complex antibody engineering tasks [\[Olsen et al., 2022b,](#page-9-0) [2024,](#page-10-0) [Prihoda et al., 2022,](#page-10-1) [Høie et al., 2023\]](#page-9-1). In this paper, we introduce IgBlend, a multi-modal model designed to incorporate both sequence and structural information for antibody engineering. Our approach can utilize either sequence, structure, or both, enabling the model to not only sample sequences that can fold to the same parental backbone but also generate more diverse sequences, providing greater flexibility in designing antibody sequences. Moreover, by utilizing both experimentally resolved structures [\[Dunbar et al., 2014\]](#page-9-2) and synthetic data generated through structure prediction models [\[Abanades et al., 2023b,](#page-9-3) [Ruffolo et al., 2023\]](#page-10-2), we aim to improve model performance on key antibody engineering tasks. Our contributions can be summarized as follows:

- We introduce IgBlend, a model that learns antibody representations from either sequence, structure or sequence-structure pairs when structural data is available.
- We present a pre-training strategy with multiple sub-objectives as well as a procedure for training and dataset processing, all of which can broadly be applied to other multi-modal training settings.
- We empirically demonstrate that integrating structural information, even when synthetically generated, significantly improves the performance of large models across a wide range of benchmarks.

The remainder of the paper is organized as follows. First, we discuss related works and introduce the notations. Section [2](#page-2-0) details the architecture of IgBlend, datasets, and training procedures. Then, we compare the performance of IgBlend against existing models in Section [3.](#page-5-0) Lastly, we point out that a more detailed background on antibodies can be found in Appendix [A.](#page-11-0)

Related work. In recent years, significant progress has been made in developing protein and antibody foundation models, drawing from advances in natural language processing (NLP). These models can be broadly categorized based on their focus—either on general protein design or antibody-specific tasks—and the way they approach the problem, such as sequence-to-structure prediction, structureto-sequence generation, or sequence-structure co-design. For general protein design, sequenceto-structure models, including AlphaFold [\[Jumper et al., 2021\]](#page-9-4) and RoseTTAFold [\[Baek et al.,](#page-9-5) [2021\]](#page-9-5), have significantly improved the prediction of protein structures from amino acid sequences. In the structure-to-sequence domain (inverse folding), ESM-IF [\[Hsu et al., 2022\]](#page-9-6) predicts amino acid sequences that fold into a given structure. Meanwhile, sequence-to-sequence models, such as ESM [\[Rives et al., 2021\]](#page-10-3) and its variants, excel at identifying patterns within sequences for tasks such as sequence recovery and mutation prediction. Moreover, some recent works have focused on sequence-structure co-design, with models such as ESM3 [\[Hayes et al., 2024\]](#page-9-7) incorporating both sequence and structure as well as function to improve protein design. There is also hybrid approaches such as LM-Design [\[Zheng et al., 2023\]](#page-10-4) that leverage both sequence and structural inputs to design new protein sequences, aligning with the approach used in this paper. In the context of antibodies, several models have emerged with a similar framework but tailored for immunoglobulins. For sequence-to-structure prediction, antibody-specific models such as ImmuneBuilder [\[Abanades](#page-9-3) [et al., 2023b\]](#page-9-3) and IgFold [\[Ruffolo et al., 2023\]](#page-10-2) predict 3D antibody structures from sequence data. In the structure-to-sequence domain, AntiFold [\[Høie et al., 2023\]](#page-9-1) addresses the inverse folding problem, predicting antibody sequences that correspond to a given backbone structure. For sequenceto-sequence tasks, models such as AbLang [\[Olsen et al., 2022b\]](#page-9-0), AbLang-2 [\[Olsen et al., 2024\]](#page-10-0), AntiBERTy [\[Ruffolo et al., 2021\]](#page-10-5), and Sapiens [\[Prihoda et al., 2022\]](#page-10-1), which are predominantly based on the BERT architecture [\[Devlin et al., 2018\]](#page-9-8), are specifically designed for antibody sequences, helping to improve performance on tasks such as residue restoration and paratope identification. To the best of our knowledge, although these models focus on either sequence or structure, no existing antibody-specific LLM effectively integrates both modalities. In this paper, we address this gap by introducing a sequence-structure-to-sequence framework, similar to LM-Design [\[Zheng et al.,](#page-10-4) [2023\]](#page-10-4) for proteins, to learn a richer and more informative representation. IgBlend learns joint representations of structure and sequence during pre-training, improving upon models that rely on

Figure 1: (a) Antibody structure with antigen binding (Fab), crystallizable (Fc), and variable (Fv) regions, (b) Zoom over the variable region which contains an heavy and a light chain, CDRs regions are displayed in magenta, (C) Modalities that we exploit in this paper for antibody modeling.

a single modality. Finally, we leave out diffusion, flow matching, and graph-based approaches to antibody design to maintain our focus on language models.

Notations. For any single unpaired chain (heavy, light or nanobody), we denote the backbone structure and sequence of the chain with n residues as follows:

structure:
$$
\mathbf{x} := (\mathbf{x}_1, ..., \mathbf{x}_n) \in \mathbb{R}^{3 \times 3 \times n}
$$
 and sequence: $\mathbf{s} := (\mathbf{s}_1, ..., \mathbf{s}_n) \in \mathbb{A}^n$

where $\mathbf{x}_i \in \mathbb{R}^{3 \times 3}$ represents the 3D coordinates of the C-alpha, N, and C atoms of the i^{th} residue, while $s_i \in A := [A, R, N, D, C, E, Q, G, H, I, L, K, M, F, P, S, T, W, Y, V, *]$ specifies the amino acid type corresponding to the i^{th} residue, where $i \in \{1, \ldots, n\}$. For consistency in notation, we will also use ∗ to denote the unknown token for both structure and sequence tokens, acknowledging a slight abuse of notation. Moreover, we stress that this work solely focuses on unpaired sequences and leaves the fine-tuning on purely paired sequences for future work. In the remainder of this paper, we will use \mathbb{P}, \mathbb{E} , and \mathbb{I} to denote the standard probability, expectation, and indicator functions, with \mathbb{I} specifically taking values in $\{0, 1\}$. To compute the differences between two sequences $(s, \hat{s}) \in \mathbb{A}^{|s|}$ of the same length, we will use the normalized Levenshtein distance: Levenshtein(s, \hat{s}) = (1/|s|) ·
 \hat{s}) To compute differences between two healthene etmotypes (x, \hat{s}) \in $\mathbb{R}^{3 \times 3 \times |\mathbf{x}|}$ $\sum_{i=1}^{|\mathbf{s}|} \mathbb{I}\{\mathbf{s}_i \neq$ \hat{s}_i . To compute differences between two backbone structures $(x, \hat{x}) \in \mathbb{R}^{3 \times 3 \times |\mathbf{x}|}$, we will use the Root Mean Square Deviation (RMSD) defined as RMSD($\mathbf{x} \hat{\mathbf{x}}$) = arg min = $\mathbf{x} = \mathbf{x}(\frac{1}{3}|\mathbf{x}|)$. the Root Mean Square Deviation (RMSD) defined as $RMSD(x, \hat{x}) = \arg \min_{R \in \Omega_3, t \in \mathbb{R}^3} (1/3|x|) \cdot \sum_{i \leq |\mathbf{x}|, j \leq 3} ||\mathbf{x}_i^j - R^* \hat{\mathbf{x}}_i^j - t^*||_2^2)^{1/2}$ where $R^* \in \mathbb{R}^3$ ³ and $t^* \in \mathbb{R}^3$ respectively denote th rotation matrix and translation after finding the optimal rigid alignment with the Kabsch algorithm [\[Kabsch, 1976\]](#page-9-9) between the backbone structures where $\Omega_3 \subset \mathbb{R}^{3 \times 3}$ denotes the set of 3D rotations and ∥·∥² denotes the standard Euclidean distance.

2 Methods

In this section, we present the IgBlend architecture, the dataset, and the pre-training objectives.

2.1 Model architecture

The proposed architecture, IgBlend, is illustrated in Fig [2](#page-2-1) and consists of three primary components: a structure encoder that handles the backbone coordinates of the antibody, a sequence encoder that processes the amino acid sequence, and a multi-modal trunk that processes both structural and sequential representations.

Structure encoder. It generates an abstract representation vector for each set of coordinates $x_i \in$ $\mathbb{R}^{3\times3}$ from the full sequence of coordinates $\mathbf{x} = (\mathbf{x}_1,\ldots,\mathbf{x}_n) \in \mathbb{R}^{3\times3\times n}$. This representation (a 512-dimensional embedding) encapsulates the geometry of the global backbone structure. The architecture comprises four GVP-GNN (Graph Neural Network Geometric Vector Perceptron) layers [\[Jing et al., 2020\]](#page-9-10), followed by two generic Transformer encoder layers [\[Vaswani et al., 2017\]](#page-10-6). This

Figure 2: Architecture of the Ig-Blendmodel. It takes as input both: a series of amino acids (top) and a series of 3D coordinates (bottom). The symbol * denotes either a masked amino acid or a masked set of coordinates. Note that the model can process each modality independently by setting all the tokens of one modality to mask. S_e denotes the sequence embedding (i.e. lookup table), T denotes a transformer block, P_e denotes the sinusoidal position embedding. The sequence encoder is displayed of the bottom left, the structure encoder on the bottom left and on the right the multimodality processor.

design is invariant to rotation and translation of the input coordinates and has been demonstrated to effectively capture protein geometries in various learning tasks [\[Jing et al., 2020\]](#page-9-10), including structure-to-sequence models such as ESM-inverse folding [\[Hsu et al., 2022\]](#page-9-6) and AntiFold [\[Høie](#page-9-1) [et al., 2023\]](#page-9-1). The input to the encoder consists of a series of residue coordinates x, with a local reference frame established for each amino acid, as per the approach used in AlphaFold2 [\[Jumper](#page-9-4) [et al., 2021\]](#page-9-4). A change of basis is then performed according to this local reference frame, rotating the vector features from the GVP-GNN outputs into the local reference frame of each amino acid. Finally, the output of the GVP is processed through two Transformer blocks, producing a 512-dimensional embedding for each residue. Notably, each or all sets of coordinates can be masked using the ∗ token.

Sequence encoder. In parallel to the structure encoder, the sequence encoder generates a vector representation (i.e., embedding of size 512) for each amino acid $s_i \in A$ in the full sequence $s = (s_1, \ldots, s_n)$. The architecture includes a one-hot encoded input followed by two blocks of a standard Transformer model [\[Vaswani et al., 2017\]](#page-10-6). This architecture has already been shown to learn relevant information from antibody sequences [\[Olsen et al., 2022b,](#page-9-0) [2024\]](#page-10-0). Specifically, the module utilizes sinusoidal positional embeddings, a SwiGLU activation function [\[Shazeer, 2020\]](#page-10-7), and has an embedding dimension of 512. Additionally, any amino acid within the sequences can be masked using the masked token ∗.

Multi-modality encoder. The fusion layer processes both modalities in two steps. First, it combines the abstract representations from the sequence and structure encoders by concatenating them along the embedding dimension, forming a single vector of size 1024 for each residue. It then processes the concatenated modalities through a series of four Transformer blocks with SwiGLU activations.

Classification head. Finally, the classification head consists of a multi-layer perceptron (MLP) followed by a softmax function and processes the multi-modal representation to generate a probability distribution over amino acid types at each position. Further details can be found in Appendix [B.](#page-11-1)

2.2 Data preparation

To create a model capable of processing both sequential and structural information, we compiled (1) a structural dataset $\mathcal{D}_{struct} = \{ (\mathbf{s}, \mathbf{x})_1, \dots, (\mathbf{s}, \mathbf{x})_{|\mathcal{D}_{struct}|} \}$, which includes structures paired with their corresponding sequences, and (2) a sequential dataset $\mathcal{D}_{seq} = \{(s, *)_1, \ldots, (s, *)_{|\mathcal{D}_{seq}|}\}$, which consists solely of sequence data. These datasets were derived from four sources: SAbDab [\[Dunbar](#page-9-2) [et al., 2014\]](#page-9-2); PLAbDab [\[Abanades et al., 2023a\]](#page-8-0); OAS datasets [\[Olsen et al., 2022a\]](#page-9-11); and INDI [\[Deszynski et al., 2022\]](#page-9-12) fully described in the Appendix [C](#page-11-2) and summarized in Table [1.](#page-3-0) The datasets ´ are further divided into samples for heavy chains, light chains, and nanobodies, resulting in \mathcal{D}_{struct} = $\mathcal{D}_{struct,H} \cup \mathcal{D}_{struct,L} \cup \mathcal{D}_{struct,N}$ and $\mathcal{D}_{seq} = \mathcal{D}_{seq,H} \cup \mathcal{D}_{seq,L} \cup \mathcal{D}_{seq,N}$. Due to the significant imbalance in the number of samples across modalities, as noted in Table [1,](#page-3-0) we implemented a new sampling scheme to rebalance the data. For each modality $M \in \{seq, struct\}$ and each chain type $C \in \{L, H, N\}$, we clustered the datasets $\mathcal{D}_{M,C}$ using MMseqs2 [\[Steinegger and Söding, 2017\]](#page-10-8), clustering over the full sequences with the parameters "-cov-mode 1", "-c 0.8", and "-min_seq_id 0.8" for the sequential datasets and over the concatenated CDR regions with the parameter "−min_seq_id 0.9" for the structure datasets. This process resulted in a union of n_{cluster} clustered samples $\overline{\mathcal{D}_{M,C}} = \bigcup_{i=1}^{n_{\text{cluster}}} C_{M,C}(i)$ for each modality and chain type. Based on these clusters, we defined the distributions $\mathcal{P}(\bar{\mathcal{D}}_{\text{struct}})$ and $P(D_{\text{seq}})$ over each dataset modality as follows: first, we sample a chain type C with equal probability: $\mathbb{P}(C = H) = \mathbb{P}(C = L) = \mathbb{P}(C = N) = 1/3$, then we select a sample within the corresponding dataset $\mathcal{D}_{\text{C,M}}$ according to the size of its corresponding cluster:

Table 1: Number of unique samples per modalities and chain types after the first pre-processing step.

$$
\mathbb{P}(\mathbf{s}, \mathbf{x})_{|\mathbf{M}, \mathbf{C}} = \begin{cases} 1/|\mathcal{C}_{\mathbf{M}, \mathbf{C}}(i_s)| & \text{if } (\mathbf{s}, \mathbf{x}) \in \mathcal{D}_{\mathbf{M}, \mathbf{C}} \\ 0 & \text{otherwise} \end{cases}
$$
(1)

where i_s denotes the index of the cluster containing s, and $|\mathcal{C}_{M,C}(i_s)|$ indicates the size of its corresponding cluster. This clustering-based distribution approach enables us to preserve the entire dataset while re-weighting each cluster to improve diversity in the training set. After clustering, 10% of the clusters are set aside for validation, and another 10% are reserved for testing. Both sets are completely excluded from the training set and have less than 0.8 sequence identity with the training data, ensuring that the validation and test sets are sufficiently distinct from the training set.

2.3 Pre-training objectives

To train IgBlend, the data distribution defined by Equation [\(1\)](#page-4-0) was used, ensuring a balanced representation of heavy, light and nanobodies chains across the two datasets, \mathcal{D}_{seq} and \mathcal{D}_{struct} . We employ a specialized masked language modeling objective capable of handling both sequential and structural data. The model parameters, θ , are optimized by minimizing the sum of three losses based on cross-entropy:

$$
\mathcal{L}_{\text{multi-modal}} := \mathcal{L}_{\text{seq2seq}} + \mathcal{L}_{\text{seq+struct2seq}} + \mathcal{L}_{\text{struct2seq}} \tag{2}
$$

where:

$$
\begin{cases}\n\mathcal{L}_{seq2seq} = \mathbb{E}_{(\mathbf{s},*)\sim\mathcal{P}(\mathcal{D}_{seq})}\left[\sum_{i \in \mathcal{T}_s} -\log(p_{\theta}(s_i|\mathbf{s}_{/\mathcal{M}_s}, *)\right] \\
\mathcal{L}_{seq+struct2seq} = \mathbb{E}_{(\mathbf{s},\mathbf{x})\sim\mathcal{P}(\mathcal{D}_{struct})}\left[\sum_{i \in \mathcal{T}_s} -\log(p_{\theta}(s_i|\mathbf{s}_{/\mathcal{M}_s}, \mathbf{x}_{/\mathcal{M}_x})\right] \\
\mathcal{L}_{struct2seq} = \mathbb{E}_{(\mathbf{s},\mathbf{x})\sim\mathcal{P}(\mathcal{D}_{struct})}\left[\sum_{i \in \mathcal{T}_s} -\log(p_{\theta}(s_i|*, \mathbf{x}_{/\mathcal{M}_x})\right]\n\end{cases}
$$
\n(3)

with $p_\theta(\mathbf{s}_i | \mathbf{s}, \mathbf{x})$ denoting the output of the softmax layer shown in Figure [2](#page-2-1) at position $i \in \{1, \dots, n\}$, given (s, x) as input. By using this combination, the model dedicates equal time on each task. The masking strategy for each pre-training objective is outlined below, defining the positions of the amino acids to predict \mathcal{T}_s , the masked residues in the sequence \mathcal{M}_s , and the masked structures \mathcal{M}_x :

- seq2seq. This task, used in training sequence-only antibody models [\[Devlin et al., 2018,](#page-9-8) [Olsen et al., 2024\]](#page-10-0), is applied to the sequential dataset \mathcal{D}_{seq} , which lacks structural information (i.e., $x = *$). For each sequence, between 10 and 40 of the amino acids are selected for masking using one of two methods: (i) randomly sampling individual residues throughout the sequence or (ii) masking continuous spans of residues, with the starting position chosen at random. The positions of the residues to be predicted are the same as those masked, $\mathcal{M}_s = \mathcal{T}_s$. The masked residues in \mathcal{M}_s are then processed using one of three strategies: (a) replaced by the unknown token ∗ with 80% probability, (b) substituted with a different amino acid with 10% probability, or (c) left unchanged with 10% probability. The masking distribution is also slightly adjusted to ensure balanced coverage of both CDR and framework regions.
- seq+struct2seq. Both sequential and structural information are used to predict masked amino acids, with masking applied to both the structure and sequence simultaneously. The same residues are used for both prediction and masking, with $T_s = M_x$. Following the seq2seq approach, 10 to 40 of the amino acids are masked, using a mix of continuous spans and random positions. With equal probability, we either (i) mask the corresponding coordinates $\mathcal{M}_x = \mathcal{M}_s$ or (ii) retain the full structural information $\mathcal{M}_x = \emptyset$ to use it as guidance.
- **struct2seq.** Only the structural information from the structural dataset \mathcal{D}_{struct} is used to predict amino acids s_i at specific target positions \mathcal{T}_s . The input sequence data is completely disregarded, replaced by a series of unknown tokens ∗, leaving only the structural information x. The target positions for amino acid prediction and masked structures, $T_s = M_x$, are

chosen using the same distribution as in the seq2seq task, alternating between continuous spans and random positions.

2.4 Training details

The model was trained on 8 A10G GPUs using a distributed DDP strategy and the PyTorch Zero Redundancy Optimizer [\[Rajbhandari et al., 2020\]](#page-10-9). The total number of training steps was predetermined at 125,000. The learning rate was warmed up over the first 200 steps to a peak of 0.001, after which it was gradually reduced to zero using a cosine scheduler. Training was conducted in 16-bit precision. To conserve memory and enable a larger batch size, gradient activation checkpointing was implemented immediately after the structural module. The effective batch size was set to 90 per GPU, resulting in a total batch size of 720 samples per step. The AdamW optimizer was used with a weight decay parameter of 0.1, epsilon of 0.00001, and betas of [0.9, 0.95] for regularization. More details about the hyperparameters can be found in the Appendix [B.](#page-11-1)

3 Empirical results

In this section, we evaluate the impact of incorporating structural information into the pre-training of antibody LLMs. Our evaluation focuses on three tasks: (i) sequence recovery of the variable region, (ii) editing of the CDR regions, and (iii) inverse folding. We compare the performance of IgBlend with five existing open-source antibody and nanobody language models, including AbLang [\[Olsen et al., 2022b\]](#page-9-0), AbLang2 [\[Olsen et al., 2024\]](#page-10-0), AntiBERTy [\[Ruffolo et al., 2021\]](#page-10-5), Sapiens [\[Prihoda et al., 2022\]](#page-10-1) and Nanobert [\[Hadsund et al., 2024\]](#page-9-13) as well as two inverse folding models, including AntiFold [\[Høie et al., 2023\]](#page-9-1) and ESM-IF [\[Hsu et al., 2022\]](#page-9-6).

3.1 Sequence recovery

First, we evaluated the task of recovering missing residues in the variable region of an antibody. This task is particularly relevant for various applications where the goal is either to recover, edit, or mutate specific amino acids within a sequence. Following the benchmark established in [\[Olsen](#page-9-0) [et al., 2022b,](#page-9-0) [2024\]](#page-10-0), we proceeded as follows. First, we sampled 1,000 sequences/structures pairs, (s, x) , per chain type from the test distribution, using Equation [\(1\)](#page-4-0). Then, for each pair (s, x) , we randomly sample a sequential mask \mathcal{M}_s that contains between 10% and 40% of the residue indices from the full sequence $[1, \ldots, |s|]$. To evaluate the benefits of incorporating structural information, we compared the performance of IgBlend using different input types. Specifically, we assessed each model's ability to recover the full sequence under various conditions: for sequence-only models, we used $\hat{s} = \text{Model}(s_{/M_s})$; for structure-guided sequential models, we used $\hat{s} = \text{Model}(s_{/M_s}, \mathbf{x})$;
for sequential models that use mosked structural information, we used $\hat{s} = \text{Model}(s_{/M_s}, \mathbf{x})$; for sequential models that use masked structural information, we used $\hat{s} = \text{Model}(\mathbf{s}/M_s, \mathbf{x}/M_s)$;
and for inverse folding models, we used $\hat{s} = \text{Model}(\mathbf{x})$. For each chain type and each region and for inverse folding models, we used $\hat{s} = \text{Model}(\mathbf{x})$. For each chain type and each region reg∈[FW1, CDR1, FW2, CDR2, FW3, CDR3, FW4], where FW and CDR refer to framework and CDR regions of the chain respectively, we recorded the empirical accuracy of the models by computing the Levenshtein(${s_i, i \in \mathcal{M}_s \cap \text{reg}}$, ${\hat{s_i}, i \in \mathcal{M}_s \cap \text{reg}}$) distance over each region and averaged the results over 1,000 sequences. Results are reported in Table 2 for the CDR3 regions and averaged the results over 1,000 sequences. Results are reported in Table [2](#page-6-0) for the CDR3 regions and the remaining regions can be found in Table [4](#page-13-0) of the Appendix. A few remarks are of order:

- First, sequence-only models (AbLang, AbLang2, AntiBERTy, Sapiens, IgBlend) performed similarly across all regions, with at most a 3% difference in accuracy between the models in most regions. In this sense, it has to be noted that IgBlend, trained using a combination of three different objectives shown in Equation [\(2\)](#page-4-1), performed on par with models trained solely on the seq2seq task, indicating that multi-modal training does not compromise performance on individual tasks.
- Secondly, it is important to note that the performance of IgBlend consistently improves with the addition of more input modalities across all chain types. Specifically, for each chain type, IgBlend shows the same ranking in recovery rate: IgBlend(seq+struct guided) > IgBlend(seq+masked struct) > IgBlend(seq-only) and IgBlend(seq+struct guided) > Ig-Blend(inverse folding). Hence, adding information helps the model to be more precise and we deduce that proposed training procedure allows us to merge both modalities successfully.

Table 2: Sequence recovery results. The task involves masking a proportion of residues in a sequence and having the model predict them. The table shows the average percentage of successfully recovered masked residues by region and chain type, with bold font highlighting the best results in each category. IgBlend in "Seq + Struct Guided" mode demonstrates the best overall performance.

• Most notably, by incorporating structural information alongside the masked sequence (IgBlend(seq+struct guidance)), we achieved consistently better results than sequence-only models across all regions and for all modalities. This improvement was particularly notable in the CDR3 region of the nanobody (N-CDR3), where IgBlend(seq+struct) outperformed the second-best model by 11.8% in accuracy. Similar improvements were observed in the CDR3 regions of the light chain (L-CDR3) with a 6.6% increase, and the heavy chain (H-CDR3) with a 7.7% increase.

3.2 Complementarity-determining region (CDR) editing

Second, we focused on the task of editing/recovering the CDR regions of a single chain, which is of particular importance in the process of optimizing antibodies for affinity. In this task, one of the CDR regions is randomly selected and fully masked, i.e., $M_s \in \{CDR1, CDR2, CDR3\}$, and the models are asked to predict the residues within the selected fully masked CDR. Using the same experimental setup with the masked CDR \mathcal{M}_s , we evaluated the models—seq-only, structure-guided, and inverse folding—on 1,000 sequences sampled from the test set as described in Equation [\(1\)](#page-4-0). These sequences were not seen during the training of IgBlend, and we recorded the percentage of successfully recovered residues for each chain type. The results can be found in the table shown in Figure [3](#page-7-0) with all models being evaluated on the same masked sequences. To further evaluate how well models using structural information adhere to structural instructions, we assessed the structural similarities between the generated sequences and the input structure x. We compared the top models in each category: AbLang2 for heavy/light chains, NanoBert for nanobodies, and AntiFold for inverse folding. We sampled 500 sequences per chain type from the test distribution and tasked each model with recovering a missing CDR. For each recovered sequence \hat{s} , we computed its structural approximation $\hat{\mathbf{x}} = \text{IgFold}(\hat{s})$ using IgFold with PyRosetta refinement. We then measured the Levenshtein($\{s_i, i \in M_s\}$, $\{\hat{s}_i, i \in M_s\}$) distance in the masked region and the RMSD($\{s_i, i \in M_s\}$) between the original and recovered structures to assess RMSD($\{x_i, i \in \mathcal{M}_s\}, \{\hat{x}_i, i \in \mathcal{M}_s\}$) between the original and recovered structures to assess
structural similarity. Results are shown in Figure 3, with extended findings available in Appendix structural similarity. Results are shown in Figure [3,](#page-7-0) with extended findings available in Appendix [D.2.](#page-12-0) From both evaluations, key observations include:

- First, as in previous experiments, the top-performing sequence-only models (AbLang, AbLang2, AntiBERTy, IgBlend) showed similar performance across different CDR regions. However, IgBlend outperformed the best sequence-only model by over 9% in accuracy for nanobodies. Notably, incorporating additional information consistently improved IgBlend's performance across all chain types (i.e., IgBlend (Seq+Struct Guided) > IgBlend (Seq+Masked Struct) > IgBlend (Seq-only)). Specifically, adding structural information to the masked sequence (IgBlend (seq+struct guidance)) significantly enhanced performance compared to the best sequence-only models, with improvements of 11.8% in H-CDR3, 6.74% in L-CDR3, and 15.43% in N-CDR3.
- Second, unlike the previous experiments, IgBlend (seq+struct guidance), which uses both $(s_{\mathcal{M}_s}, x_{\mathcal{M}_x})$, shows performance closer to IgBlend (inverse folding), which relies solely on the structure x, than to IgBlend (seq-only), which depends only on the sequential information $s_{\mathcal{M}_s}$. This suggests that, in the task of re-editing complete CDR regions, IgBlend relies more on structural information than on sequential data.

Figure 3: **CDR in-filling results:** One CDR region (CDR1, CDR2, or CDR3) is fully masked, and the model attempts to recover it. **Top:** The table shows the average percentage of correctly recovered residues for heavy chain (H), light chain (L) and nanobodies (N). **Bottom:** The graphs show Levenshtein distances of generated CDR3 regions from the original sequence on the x-axis, and RMSD of the predicted structures from the original backbone on the y-axis for each chain type.

• In terms of structural similarity, it is noteworthy that IgBlend (structure guided) consistently achieves the highest percentage of structures within the bins corresponding to the smallest RMSD values for each chain type, surpassing even the best inverse folding model. Specifically, IgBlend shows 37%, 46%, and 51% of structures in the smallest RMSD bins (1 Å for heavy chains, 0.2 Å for light chains, and 1 Å for nanobodies), compared to $32\%, 5\%,$ and 26% for AntiFold in the same bins. See Figure [3](#page-7-0) for a detailed comparison. Thus, in addition to outperforming sequential models, IgBlend(seq+struct guided) excels at generating sequences that can more accurately fold to the original backbone structure compared to those produced by AntiFold.

3.3 Inverse folding

Finally, we assessed IgBlend's ability to perform the inverse folding task [\[Hsu et al., 2022,](#page-9-6) [Høie](#page-9-1) [et al., 2023\]](#page-9-1), which involves recovering a sequence s from its structure x alone. As with previous experiments, we sampled 500 structure-sequence pairs (s, x) for each chain type from the test set, which was not seen during IgBlend's training. Each inverse folding model was then asked to predict the sequence $\hat{s} = \text{Model}(\mathbf{x})$ based solely on the structure x. To evaluate model performance, we tested different temperatures: $T = 1e - 4$ for the highest probability sequence, $T = 1$ for unbiased results, and $T = 2$ and $T = 3$ for more diverse sequences. We recorded the normalized Levenshtein(s, \hat{s}) distance between the predicted and original sequences, and the RMSD(\bf{x}, \hat{x}) of the approximated structure $\hat{\mathbf{x}} = \text{IgFold}(\hat{\mathbf{s}})$ as a measure of structural similarity. Results for the lowest temperature are shown in Figure [4,](#page-8-1) with additional details in Table [5](#page-13-1) and Figure [6](#page-15-0) in the Appendix. Key observations include:

• First, we observe a positive correlation between Levenshtein distance and RMSD for every model: as the sequence diverges more from the original (larger Levenshtein distance), the RMSD tends to increase, indicating a trade-off between sequence diversity and structural precision. Consequently, as temperature is increased to generate more diverse sequences,

Figure 4: **Inverse Folding Results:** In this task, the sequence is fully masked, and the model attempts to recover it from the structure. The top graph shows the normalized Levenshtein distance between generated and original sequences, with the y-axis displaying the RMSD of the generated structures relative to the original; greater spread in Levenshtein distance and lower RMSD indicate better performance. The bottom table lists the percentage of times each method produced a sequence with RMSD < 1 Å across 500 samples per modality. More details are available in Table [5](#page-13-1) in the Appendix.

the RMSD increases. However, IgBlend demonstrates greater robustness to temperature changes. Second, all models perform better on light chains compared to heavy chains and nanobodies, suggesting that the inverse folding task is more challenging for heavy chains and nanobodies.

• Second, across all temperatures, chain types, and RMSD thresholds (see Figure [4](#page-8-1) and Table [5](#page-13-1) in the Appendix), the models consistently rank as follows based on the number of samples with RMSD below the threshold: IgBlend> AntiFold > ESM-IF. This shows that IgBlend outperforms current state-of-the-art methods in generating sequences that accurately fold back to the original structure. Notably, IgBlend is the first inverse folding model to achieve results on nanobodies comparable to heavy chains. However, this high accuracy comes with lower Levenshtein distance, a limitation seen in all tested settings.

4 Conclusion and future work

In this study, we investigated how incorporating structural information into antibody LLMs enhances performance. We introduced a model that integrates both structural and sequential data, showing that this combination consistently improves performance across all benchmarks compared to sequencebased and inverse-folding models. However, while effective, our approach sometimes sacrifices sequence diversity for accuracy. Future work will focus on including side-chain information and expanding structural datasets.

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A Background on antibodies

Background. In humans, antibodies are classified into five isotypes: IgA, IgD, IgE, IgG, and IgM. This work primarily focuses on IgG antibodies, which are Y-shaped glycoproteins produced by B-cells (see Figure [1\)](#page-1-0), as well as nanobodies, which are antibody fragments consisting of a single monomeric variable domain. Henceforth, "antibody" will specifically refer to IgG antibodies. Antibodies consist of distinct regions that play specific roles in the immune response. The Fab (fragment antigenbinding) region, composed of both variable (V) and constant (C) domains from the heavy and light chains, is primarily responsible for antigen binding. Within this region, the antigen-binding site is formed by the variable domains — VH for the heavy chain and VL for the light chain — which determine the specificity of the antibody and enable it to recognize and bind to specific antigens. The Fv (fragment variable) region is the smallest functional unit of an antibody that can still bind to an antigen. It consists solely of the variable domains (VH and VL) of the heavy and light chains, without the constant domains. Within the variable domains, there are two key distinct regions: the framework regions and the complementarity-determining regions (CDRs). The framework regions provide structural support, maintaining the overall shape of the variable domains, while the CDRs, comprising three loops on both the VH and VL chains, are directly involved in binding to the antigen. These CDRs are crucial for the precise recognition and interaction with specific antigens. While the Fv region is essential for the initial recognition and binding of antigens, it lacks the effector functions present in the full antibody. The Fab region, being larger and more complex due to the inclusion of both variable and constant domains, is generally more stable and has a higher affinity for antigens. The Fv region, on the other hand, is simpler and more easily engineered for various applications, such as in the development of single-chain variable fragment (scFv) antibodies. The base of the Y-shaped antibody, known as the Fc (fragment crystallizable) region, is involved in regulating immune responses. It interacts with proteins and cell receptors, ensuring that the antibody generates an appropriate immune response. Moreover, nanobodies, which are small, single-domain antibodies derived from heavy-chain-only antibodies found in certain animals such as camels and llamas, are even more compact than traditional Fv regions. They retain full antigen-binding capacity while offering advantages such as increased stability and easier production, making them valuable tools in both therapeutic and diagnostic applications.

B Architectural details

In Table [3,](#page-12-1) we collect the full architectural details of the IgBlendarchitecture used in the paper.

C Data set

Data source. To create a model capable of processing both sequential and structural information, we needed to address the significant asymmetry in the availability of data across these modalities (204M sequences and 3M structures as shown in Table [1\)](#page-3-0). Therefore, we compiled two datasets: (1) a structural dataset \mathcal{D}_{struct} , which includes structures paired with their corresponding sequences, and (2) a sequential dataset \mathcal{D}_{seq} , which consists solely of sequence data. These datasets were derived from four primary sources: SAbDab [\[Dunbar et al., 2014\]](#page-9-2), which contains experimentally determined structures using techniques such as electron crystallography and X-ray diffraction; PLAbDab [\[Abanades et al., 2023a\]](#page-8-0), which provides sequences derived from patents; OAS datasets [\[Olsen et al.,](#page-9-11) [2022a\]](#page-9-11), which compile and annotate immune repertoires; and INDI [\[Deszynski et al., 2022\]](#page-9-12), which ´ contains sequences of nanobodies. Given the relatively small number of experimentally determined structures (e.g., approximately 2,000 samples from SAbDab, as shown in Table [1](#page-3-0) after applying our selection criteria), we expanded our structural dataset by incorporating inferred structures. In addition to the inferred structures already present in the PLAbDab dataset (folded with ImmuneBuilder), we generated additional structures from the OAS paired, unpaired and INDI. The paired sequences from OAS were folded with ImmuneBuilder [\[Abanades et al., 2023b\]](#page-9-3) and a clustered version of the unpaired OAS and INDI dataset were folded using IgFold [\[Ruffolo et al., 2023\]](#page-10-2). This process resulted in approximately 4 million unique structures. For the sequential dataset, we extracted data from four repertoires: OAS paired, OAS unpaired, PLAbDab paired, PLAbDab unpaired and INDI.

For each of the datasets \mathcal{D}_{struct} and \mathcal{D}_{seq} , we begin by removing all duplicates, defined as pairs of data with identical sequences. Next, only the data that meet the following criteria are retained:

(1) no unknown residues, (2) no missing residues, and (3) no shorter than expected IMGT regions [\[Ehrenmann et al., 2010\]](#page-9-14), as determined by running ANARCI [\[Dunbar and Deane, 2016\]](#page-9-15). After these cleaning steps, we are left with two datasets: $\mathcal{D}_{struct} = \{ (\mathbf{s}, \mathbf{x})_1, \dots, (\mathbf{s}, \mathbf{x})_{|\mathcal{D}_{struct}|} \}$, which contains pairs of sequences and structures, and $\mathcal{D}_{seq} = \{(s, *)_1, \ldots, (s, *)_{|\mathcal{D}_{seq}|}\}$, which contains only sequential information.

D Experimental results

D.1 Sequence recovery

Table [4](#page-13-0) records the sequence recovery rate on all regions and for each modality.

D.2 CDR editing

Figure [5](#page-14-0) collects the result of the CDR recovery experiment in all CDR regions.

D.3 Inverse folding

Figure [6](#page-15-0) displays the inverse folding results for the different temperatures. Table [5](#page-13-1) reports the results for different RMSD thresholds.

Table 4: Sequence recovery results. The task consists of masking randomly a proportion of residues within a sequence and asking the model to predict the masked residues. The table display the average percentage of successfully recovered masked residues in each region and for each type of chain. Bold font indicates the best result in the comparison

		Heavy			Light			Nanobodies	
RMSD<0.5	AntiFold	ESM-IF	IgBend	AntiFold	ESM-IF	IgBend	AntiFold	ESM-IF	IgBend
$T=1e-4$	20.40	02.40	27.6	49.20	24.20	72.0	07.20	03.40	32.00
T=1	18.60	00.80	25.00	34.00	09.80	68.00	08.79	02.00	30.00
$T=2$	05.95	00.00	10.60	07.50	00.00	47.40	04.75	00.00	08.80
$T=3$	00.00	00.00	01.00	00.00	00.00	06.60	00.00	00.00	00.80
RMSD<1	AntiFold	ESM-IF	IgBend	AntiFold	ESM-IF	IgBend	AntiFold	ESM-IF	IgBend
$T=1e-4$	63.60	28.20	73.60	94.40	89.20	97.80	43.00	38.00	81.00
$T=1$	60.80	14.40	69.40	85.20	71.00	97.00	45.40	24.80	79.00
$T=2$	35.11	02.01	52.80	61.25	18.84	94.60	37.15	03.01	57.80
$T=3$	06.11	00.00	17.60	10.03	00.43	54.60	05.26	00.00	23.40
RMSD _{1.5}	AntiFold	ESM-IF	IgBend	AntiFold	ESM-IF	IgBend	AntiFold	ESM-IF	IgBend
$T=1e-4$	84.60	57.80	89.60	99.80	99.80	100.0	62.60	67.60	95.60
$T=1$	81.20	42.00	86.60	99.80	95.80	100.0	72.20	56.60	94.00
$T=2$	66.17	11.85	78.40	92.50	46.69	99.40	62.85	17.27	80.60
$T=3$	23.40	00.65	51.00	27.96	04.49	83.40	24.44	00.43	51.20
RMSD < 2	AntiFold	ESM-IF	IgBend	AntiFold	ESM-IF	IgBend	AntiFold	ESM-IF	IgBend
T=1e-4	94.00	77.80	96.40	100.0	99.80	100.0	78.20	84.80	98.20
$T=1$	92.40	66.00	95.40	100.0	99.00	100.0	84.20	75.80	97.80
$T=2$	84.68	27.91	90.80	98.12	68.34	99.60	77.97	34.14	90.00
$T=3$	47.12	02.16	71.60	48.02	11.32	93.00	39.85	02.59	69.80

Table 5: Inverse folding results: The sequence if fully masked, and the model attempts to recover it from the structure. The table displays the percentage of sequences generated by each method with a RMSD below a given threshold and for different temperatures. Higher is better.

Figure 5: CDR recovery results: One series of aminco acid of the sequence is fully masked (one CDR), and the model attemps to recover it. AntiFold only uses the structural information. IgBlend (structure guidance) uses the masked sequence and the structure information. The distances (both Levenshtein and RSME) are only computed in the masked CDR regions. The x-axis displays the Levensthein distance of the generated sequences to the original one and the y-axis reports the RMSE of the generated sequence with regards to the original structure.

Figure 6: **Inverse folding results:** The sequence if fully masked, and the model attempts to recover it from the structure. Top: the graph displays the normalized Levenshtein distance of the generated sequences to the original sequences associated with the input structure and the y-axis reports the RMSD of the folded structure of the generated sequences with regards to the original structure set as input. For both metrics, lower is better.