
Accurate Protein Structure Prediction by Embeddings and Deep Learning Representations

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Abstract

Proteins are the major building blocks of life, and actuators of almost all chemical and biophysical events in living organisms. Their native structures in turn enable their biological functions which have a fundamental role in drug design. This motivates predicting the structure of a protein from its sequence of amino acids, a fundamental problem in computational biology. In this work, we demonstrate state-of-the-art protein structure prediction (PSP) results using embeddings and deep learning models for prediction of backbone atom distance matrices and torsion angles. We recover 3D coordinates of backbone atoms and reconstruct full atom protein by optimization. We create a new gold standard dataset of proteins which is comprehensive and easy to use. Our dataset consists of amino acid sequences, Q8 secondary structures, position specific scoring matrices, multiple sequence alignment co-evolutionary features, backbone atom distance matrices, torsion angles, and 3D coordinates. We evaluate the quality of our structure prediction by RMSD on the latest Critical Assessment of Techniques for Protein Structure Prediction (CASP) test data and demonstrate competitive results with the winning teams and AlphaFold in CASP13 and supersede the results of the winning teams in CASP12. We make our data, models, and code publicly available.

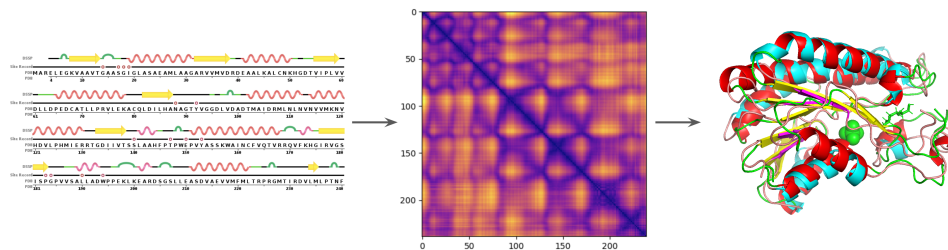


Figure 1: Our method operates by i) predicting backbone atom distance matrices and torsion angles ii) recovering backbone atom 3D coordinates, and iii) reconstructing the full atom protein by optimization

1 Introduction

Proteins are necessary for a variety of functions within cells including transport, antibody, enzyme and catalysis. They are polymer chains of amino acid residues, whose sequences dictate stable spatial conformations, with particular torsion angles between successive monomers. To perform their functions, the amino acid residues must fold into proper configurations. The sequence space of proteins is vast, 20 possible residues per position, and evolution has been sampling it over billions of years. Thus, current proteins are highly diverse in sequences, structures, and functions. The high throughput acquisition of DNA sequences, and therefore ubiquity of known protein sequences stands in contrast to the limited availability of 3D structures, which are more functionally relevant. From a physics standpoint, the process of protein folding is a search for the minimum energy conformation which happens in nature paradoxically fast [10]. Unfortunately, explicitly computing the energy of a protein conformation, along with its surrounding water molecules, is extremely complex. Inferring local *secondary structure* [7] consists of linear annotation of structural elements along the sequence [9]. Inferring *tertiary structure* consists of resolving the 3D atom coordinates of proteins. When highly similar sequences are available with known structures, this homology can be used for modeling. Recently, PSP had been tackled by first predicting contact points between amino acids, and then leveraging the contact map to infer structure. A primary indicator of contact between a pair of amino acids is their tendency to have correlated and compensatory mutations during evolution. The availability of large scale data on DNA, and therefore protein sequences allows detection of such co-evolutionary constraints from sets of sequences that are homologous to a protein of interest. Registering such contacting pairs in a matrix facilitates a framework for their probabilistic prediction. This contact map matrix can be generalized to register distances between amino acids [13].

Machine learning approaches garnered recent success in PSP [3, 2] and its sub-problems [12]. These leverage available repositories of tertiary structure [8] and its curated compilations [11] as training data for models that predict structure from sequence. Specifically, the recent biannual critical assessment of PSP methods [4] featured multiple such methods. Most prominently, a closed-source, ResNet-based architecture [5] has achieved impressive results in the CASP evaluation settings, based on representing protein structures both by their distance matrices as well as their torsion angles. In this work, we design a novel representation of biologically relevant input data and construct a processing flow for PSP as shown in Figure 1. Our method leverages advances in deep sequence models and proposes a method to learn transformations of amino acids and their auxiliary information. The method operates in three stages by i) predicting backbone atom distance matrices and torsion angles ii) recovering backbone atom 3D coordinates, and iii) reconstructing the full atom protein by optimization.

We demonstrate state-of-art protein structure prediction results using deep learning models to predict backbone atom distance matrices and torsion angles, recover backbone atom 3D coordinates and reconstruct the full atom protein by optimization. Our three key contributions are:

- A gold standard dataset of around 75k proteins described in Table 1 which we call the CUProtein dataset which is easy to use in developing deep learning models for PSP.
- Competitive results with the winning teams on CASP13 and a comparison with AlphaFold (A7D) [5] and results mostly superseding the winning teams on CASP12.
- Publicly available source code for both protein structure prediction using deep learning models and protein reconstruction.

Feature	Notation	Source	Dims	IO	Type
AA Sequence	x_1	PDB	$n \times 1$	Input	21 chars
PSSM	x_2	AA/HHBlits	$n \times 21$	Input	Real [0,1]
MSA covariance	x_3	AA/jackHMMER	$n \times n$	Input	Real [0,1]
Secondary Structure (SS)	x_4	DSSP	$n \times 8$	Input	8 chars
C_α, C_β Distance Matrices	$g_1(\cdot)$	PDB	$n \times n$	Output	Ångstrom
Torsion Angles (ϕ, ψ)	$g_2(\cdot)$	DSSP	$n \times 2$	Output	Radians

Table 1: Our CUProtein dataset consists of amino acid sequences, secondary structure, PSSM’s, MSA covariance matrices, backbone distance matrices, and torsion angles.

This work explores encoded representation for sequences of amino acids alongside their auxiliary information. We offer full access to data, models, and code, which remove significant barriers to entry for investigators and make publicly available methods for this important application domain.

2 Methods

We address two problems: (i) predicting backbone distance matrices and torsion angles of backbone atoms from amino acid sequences, Q8 secondary structure, PSSMs, and co-evolutionary multiple sequence alignment; and (ii) reconstruction of all atom coordinates from the predicted distance matrices and torsion angles.

We begin with a one-hot representation of each amino acid and secondary structure sequence, and real valued PSSM’s and MSA covariance matrices. These are passed through *embedding layers* and then onto an encoder-decoder architecture. To leverage sequence homology, we compute the covariance matrix of the MSA features by embedding the homology information along a k dimensional vector to form a 3-tensor and contract the tensor A_{ijk} along the k dimensional embedding which is then passed as input to the encoder: $\Sigma = A_{ji}^k A_{ijk}$.

We use encoder-decoder models with a bottleneck to train prediction models. The encoder f receives as input the aggregation A (by concatenation) of the embeddings e_i of each input x_i , and two separate decoders g_1 and g_2 that output distance matrices and torsion angles for $i \in \{1, \dots, 4\}$: $g_j \left(f \left(\text{Agg}_{x_i \in \mathcal{X}} (e_i(x_i)) \right) \right)$. In addition, we also use a model which consists of separate encoders f_i for each embedded input $e_i(x_i)$, that are aggregated by concatenation after encoding, and separate decoders g_1 and g_2 for torsion angles and backbone distance matrices: $g_j \left(\text{Agg}_{x_i \in \mathcal{X}} (f_i(e_i(x_i))) \right)$.

Using a separate encoder model involves a larger number of trainable parameters. Our models differ in the use of embeddings for the input, their models, and loss functions.

Building on techniques commonly used in natural language processing (NLP) our models use embeddings and a sequence of bi-directional GRU’s and LSTM’s with skip connections, and include batch normalization, dropout, and dense layers. We experimented with various loss functions including MAE, MSE, Frobenius norm, and distance logarithm to handle the dynamic range of distances. To learn the loss function we have also implemented distance matrix prediction using conditional GAN’s and VAE’s for protein sub-sequences.

Once backbone distance matrices and torsion angle are predicted we address two reconstruction sub-problems: (i) reconstructing the protein backbone coordinates from their distance matrices, and (ii) reconstructing the full atom protein coordinates from the C_α or C_β coordinates and torsion angles.

We employ three different techniques for reconstructing the 3D coordinates X between backbone atoms of a protein from the predicted matrix of their pairwise distances [6]. Given a predicted distance matrix \hat{D} our goal is to recover 3D coordinates \hat{X} of n points. We notice that $D(X)$ depends only on the Gram matrix $X^T X$: $D(X) = \mathbf{1} \text{diag}(X^T X)^T - 2X^T X + \text{diag}(X^T X) \mathbf{1}^T$. In multi-dimensional scaling we optimize: minimize $\left\| D(\hat{X}) - \hat{D} \right\|_F^2$. In semi-definite programming and relaxation we optimize: minimize $\left\| K(G) - \hat{D} \right\|_F^2$ s.t. $G \in C$, where $K(G) = \mathbf{1} \text{diag}(G)^T - 2G + \text{diag}(G) \mathbf{1}^T$ operates on the Gram matrix G . In alternating direction method of multipliers [3] we optimize:

PDB	CASP	Tar-id	Best RMSD	A7D	Ours
5Z82	13	T0951	1.01 (Seok)	NA	1.79
6D2V	13	T0965	1.72 (A7D)	1.72	1.60
6QFJ	13	T0967	1.13 (BAKER)	NA	1.18
6CCI	13	T0969	1.96 (Zhang)	2.27	2.53
6HRH	13	T1003	0.88 (MULTICOM)	2.12	2.95
6QEK	13	T1006	0.58 (YASARA)	0.78	1.02
6N91	13	T1018	1.24 (Wallner)	1.77	3.89
6M9T	13	T1011-D1	1.58 (A7D)	1.58	1.64
5J5V	12	T0861	0.49 (MULTICOM)	NA	1.00
2N64	12	T0865	1.87 (HHPred)	NA	1.58
5JMU	12	T0879	1.35 (MULTICOM)	NA	1.31
5JO9	12	T0889	1.31 (Seok)	NA	1.79
4YMP	12	T0891	1.10 (GOAL)	NA	1.36
5XI8	12	T0942-D2	1.73 (EdaRose)	NA	1.60

Table 2: Comparison between the winning CASP12/13 models for each protein, best AlphaFold (A7D) model for CASP13, and our model for CASP12/13.

$$\underset{G, Z, \eta}{\text{minimize}} \lambda \|\eta\|_1 + \frac{1}{2} \left(\sum_{i,j=1}^n (G_{ii} + G_{jj} - 2G_{ij} + \eta_{ij} - \hat{D}_{ij}^2) \right)^2 + \mathbb{1}\{Z \in S_+^n\} \text{ s.t. } G - Z = 0.$$

We have found multi-dimensional scaling to be the fastest and most robust method of the three, without depending on algorithm hyper-parameters, which is most suitable for our purposes.

Given backbone coordinates we assign plausible coordinates to the rest of the protein’s atoms. We begin with an initial guess or prediction for ϕ and ψ torsion angles. We maintain a look-up table of mean ϕ , ψ values for each combination of two consecutive α torsions and three α -angles (the angles defined by three consecutive atoms). Using these values and the C_α positions we generate an initial model. This model is then relaxed by a series of energy minimization simulations under an energy function that includes: standard bonded terms (bond, angle, plane and out-of-plane), knowledge based Ramachandran and pairwise terms, torsion constraints on the ϕ and ψ angles, and tether constraints on the C_α position. The latter term reduces the perturbations of the initial high forces. Finally, we add side-chains using a rotamer library, and remove clashes by a series of energy minimization simulations. We develop a very similar method for reconstructing the full atom protein from C_β atom distance matrices.

3 Results

We have compared our predictions on CASP12 and CASP13 [1] test targets. Deep learning methods were widely used only starting from CASP13. AlphaFold was introduced starting from CASP13. The use of deep learning methods in CASP13, due to availability of DL programming frameworks, significantly improved performance over CASP12. Table 2 provides a representative comparison between the winning CASP12 and CASP13 competition models, AlphaFold models for CASP13 for which A7D submitted predictions to CASP, and our models. These proteins were selected such that they do not contain multiple chains and are sufficiently complex. The average length of the CASP13 target proteins selected is 325 residues, and the average for CASP12 is 247 residues. Results of RMSD around 2 Angstrom on test targets are considered accurate in CASP. Our results supersede the winning teams of CASP12 compared with each best team for each protein which highlights the improvement using deep learning methods. Our approach is on par with the winning teams in CASP13, compared with the winning team for each protein, which highlights that our methods is state-of-the-art. We measure the sequence independent RMSD which is consistent with the CASP evaluation reports and matches the deposited structures and our predictions. CASP competitors such as AlphaFold provide predictions for selected proteins. Overall, our performance on CASP is highly competitive. Training our models on a cloud instance takes two days using GPUs. Prediction of backbone distance matrices and torsion angles takes a few seconds per protein, and reconstruction of full atom proteins from distance matrices and torsion angles takes a few minutes per protein, depending on protein length. Limitations of this work are: (i) we only handle single domain proteins and not complexes with multiple chains, (ii) PSSM and MSA data for several of the CASP targets are limited to a subsequence of the full length protein, and (iii) we do not use available methods for detecting and reconstructing beta-sheets.

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