RESEARCH ARTICLE

Revised: 24 July 2021

PROTEINS WILEY

Protein inter-residue contact and distance prediction by coupling complementary coevolution features with deep residual networks in CASP14

Yang Li^{1,2} | Chengxin Zhang² | Wei Zheng² | Xiaogen Zhou² | Eric W. Bell² | Dong-Jun Yu¹ | Yang Zhang²

¹School of Computer Science and Engineering, Nanjing University of Science and Technology, Nanjing, China

²Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, Michigan, USA

Correspondence

Dong-Jun Yu, School of Computer Science and Engineering, Nanjing University of Science and Technology, Nanjing, China. Email: njyudj@njust.edu.cn

Yang Zhang, Department of Computational Medicine and Bioinformatics, University of Michigan, Ann Arbor, MI, USA. Email: zhng@umich.edu

Funding information

National Institute of Allergy and Infectious Diseases, Grant/Award Number: Al134678; National Institute of General Medical Sciences, Grant/Award Numbers: GM136422, S10OD026825; National Natural Science Foundation of China, Grant/Award Numbers: 61772273, 62072243; National Science Foundation, Grant/Award Numbers: DBI2030790, IIS1901191, MTM2025426; Natural Science Foundation of Jiangsu Province, Grant/Award Number: BK20201304

Abstract

This article reports and analyzes the results of protein contact and distance prediction by our methods in the 14th Critical Assessment of techniques for protein Structure Prediction (CASP14). A new deep learning-based contact/distance predictor was employed based on the ensemble of two complementary coevolution features coupling with deep residual networks. We also improved our multiple sequence alignment (MSA) generation protocol with wholesale meta-genome sequence databases. On 22 CASP14 free modeling (FM) targets, the proposed model achieved a top-*L*/5 long-range precision of 63.8% and a mean distance bin error of 1.494. Based on the predicted distance potentials, 11 out of 22 FM targets and all of the 14 FM/template-based modeling (TBM) targets have correctly predicted folds (TM-score >0.5), suggesting that our approach can provide reliable distance potentials for ab initio protein folding.

KEYWORDS

CASP, coevolution, contact-map prediction, deep learning, protein structure prediction

1 | INTRODUCTION

Ab initio protein structure prediction has been a longstanding challenge in the field of computational biology.¹ We have witnessed evident signs of progress^{2,3} in the recent Critical Assessment of techniques for protein Structure Prediction (CASP) experiments with respect to protein structure prediction based on long-range predicted geometric potentials, usually from supervised machine learning methods. To accurately predict the long-range geometric restraints, for example, inter-residue contacts, early methods derive correlations between positions of multiple sequence alignments (MSAs) inspired by the coevolution phenomenon.⁴ Those coevolution analysis methods can be classified into local⁵⁻⁷ and global,⁸⁻¹¹ that is, direct coupling analysis (DCA), approaches, according to whether all other positions are considered when computing the coupling between a residue pair. The residue-wise correlation maps are later used by supervised machine learning methods as features,¹²⁻¹⁵ and have been incorporated into multiple successful methods¹⁶⁻¹⁸ when coupled with deep convolutional networks. An improved feature extraction strategy¹⁹⁻²³ is employed by feeding raw coevolution to a deep ResNet²⁴ with features to avoid possible information leakage during post-processing. Recent approaches further extend the pipeline for contact map prediction to distance²⁵⁻²⁷ and orientations²⁸ and provide more precise restraints to assist protein folding.

In this article, we introduce DeepPotential, which participated in CASP14 for protein contact/distance prediction. DeepPotential collects both local and global raw coevolution matrices with postprocessing and derives sequence-specific descriptors using a set of candidates from a progressive MSA construction pipeline. By coupling features with deep dilated residual convolutional networks,²⁹ DeepPotential is capable of predicting high precision contact maps superior to our previous approach in CASP13.²¹ We also have found significant improvements by searching against multiple gigantic databases. High-quality protein 3D structure models can be obtained based on the low-error distance prediction by DeepPotential, without the need for template information.

2 | MATERIALS AND METHODS

In CASP14, we tested DeepPotential, and the pipeline can be broken down into a procedure of steps as shown in Figure 1. The prediction contains steps of multi-MSA construction and selection, complementary coevolution feature extraction, and neural network prediction. We also build tertiary protein structures from the predictions of DeepPotential.

2.1 | Datasets

DeepPotential was trained on 26 151 experimentally solved structures collected from PDB,³⁰ with the latest structure being timestamped as 2019.11.12. We first set the maximum sequence length to 1000 and kept the representative sequences after a 35% sequence identity clustering procedure with CD-HIT.³¹

2.2 | Collection of multiple sequence alignments

In CASP14, the six candidate MSAs for each target sequence were generated by searching against two whole-genome databases and four metagenome sequence databases (Metaclust,³² BFD,³³ Mgnify,³⁴ and IMG/M³⁵). Following our previous work,³⁶ a progressive strategy, that is, stopping the search if the target Neff (Equation (1)) threshold is satisfied, was implemented. This early stopping criterion is found to reduce database search time without sacrificing MSA quality for targets with a great deal of homologs.³⁶ The Neff value can be calculated by

$$Neff = \frac{1}{\sqrt{L}} \sum_{n=1}^{N} \frac{1}{1 + \sum_{m=1}^{N} \mathbb{I}[S_{m,n} \ge 0.8]},$$
 (1)

where N is the total number of sequences in the MSA; I is the indicator function; $S_{m,n}$ is the sequence identity between sequence n and sequence m. Figure 2 summarizes the entire MSA generation pipeline. We first search against Uniclust30 (version 2020 01) and UniRef90 using HHblits³⁷ and Jackhmmer³⁸ and obtain MSAs for Stages 1 and 2, respectively. The resulting Stage 2 MSA will be used as the initial profile for HMMsearch and HHblits to search against the Metaclust (Stage 3 MSA) and BFD (Stage 4 MSA) databases, respectively. When searching against the Mgnify and IMG/M databases in Stage 5 and Stage 6, the MSA from Stage 4 is considered as the initial MSA profile. Note that custom HHblits databases will be constructed from raw hits to ensure that the output MSAs come from the same algorithm (Hhblits2) from Stage 2 to Stage 6. For training sequences, the ordinarily HHblits MSA (Stage 1 MSA) was employed with Uniclust30 (version 2017 04). Once the desired MSAs are obtained, the distance map with the highest confidence score will be considered as the final prediction. In CASP14, we explored two confidence score

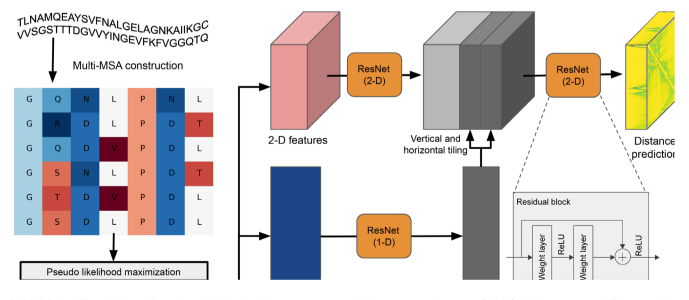


FIGURE 1 The pipeline of DeepPotential. Starting from a sequence, multiple sequence alignments (MSAs) will be constructed. Three models, that is, the pseudolikelihood maximization of the Potts model, mutual information, and a hidden Markov model will be used to extract 2D and 1D features. The two features will go through two sets of residual convolutional layers and tiled together. The combined hidden features will go through another set of residual blocks for the final prediction

PROTEINS WILEY 1913

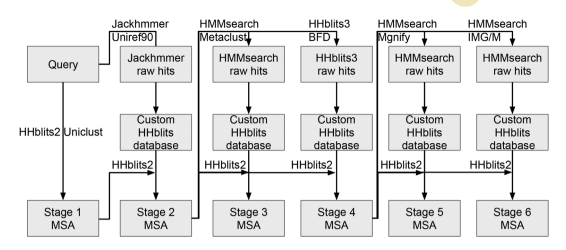


FIGURE 2 The multi-stage progressive multiple sequence alignment (MSA) construction pipeline in DeepPotential

configurations. For Group 010, the confidence score is defined as the mean of the cumulative probability under 12 Å of the top 10^*L predicted C_{β} - C_{β} distance distributions for all residue pairs, while for Group 024, the corresponding threshold is 8 Å.

2.3 | Feature extraction

DeepPotential extracts a set of complementary features as the input of the deep learning model. Coevolution analysis represents conditional or marginal correlations between residues and thus can be critical discriminative features. In DeepPotential, two types of coevolutionary features, the pseudolikelihood maximized Potts model (PLM)¹¹ and mutual information (MI), are extracted. Given the input MSA (*X*) with *N* alignments and *L* sequence length, the PLM feature can be obtained by minimizing the following loss function:

$$\mathcal{L}_{\mathsf{PLM}} = -\sum_{l=1}^{L} \sum_{n=1}^{N} \log \frac{\exp\left(e_{i}(X_{n,i}) + \sum_{j=1, j \neq i}^{L} P_{i,j}(X_{n,i}, X_{n,j})\right)}{\sum_{q=1}^{Q} \exp\left(e_{i}(q) + \sum_{j=1, j \neq i}^{L} P_{i,j}(q, X_{n,j})\right)} + \lambda_{\mathsf{single}} \sum_{i=1}^{L} \|e_{i}\|_{2}^{2} + \lambda_{\mathsf{pair}} \sum_{i, j=1}^{L} \|P_{i,j}\|_{2}^{2},$$

$$i \neq j$$
(2)

where Q = 22 represents 20 regular amino acid types, an undetermined amino acid type state and a gap state; $e \in R^{L \times Q}$ and $P \in R^{L \times L \times Q \times Q}$ are field and coupling parameters of Potts model, respectively; $\lambda_{\text{single}} = 1$ and $\lambda_{\text{pair}} = 0.2 \times (L-1)$ are the regularization coefficients for e and P. The parameter $P_{ij}(q_1, q_2)$ measures the linear coefficient of the q_1 state of residue i and the q_2 state of residue j conditioned on other residues and states, which can eliminate transitive interactions in the observed interactions.

The PLM feature eliminates transitional noise in marginal correlations, which should be more relevant to the structural interaction terms between residue pairs. However, the optimization of PLM could be ill-posed when there are no sufficient aligned sequences in the MSA. In this regard, we utilize a raw marginal correlation measurement, that is, MI, as another pairwise feature. The MI feature of residue i and j is defined as:

$$M_{ij}(q_1,q_2) = f_{ij}(q_1,q_2) \ln \frac{f_{ij}(q_1,q_2)}{f_i(q_1)f_j(q_2)},$$
(3)

here, $f_i(q_1)$ is the frequency of a residue type q_1 at position *i* of the MSA, $f_{ii}(q_1, q_2)$ is the co-occurrence of two residue types q_1 and q_2 at positions *i* and *j*. Note that the raw MI matrix $M_{i,j} \in R^{Q \times Q}$ for residue pair i and j will be kept as features without summation over residue types. Such a formula can capture residue type pair-specific information and provide the fine-grained feature to deep learning model. Compared to the regular Pearson correlation. MI is capable of measuring the non-linear relationships between variables. For each residue pair, another three post-processed residue-wise features will be extracted from the coevolution residue pair matrix ($P_{i,j}$ and $M_{i,j}$), that is, Frobenius norms of (1) the whole residue pair matrix, (2) residue pair matrix excluding the gap state, and (3) residue pair matrix excluding both the gap and undetermined amino acid state. The above coevolutionary feature set records the features for the whole MSA, while $P_{ij}(X_{1,i},X_{1,j})$ and $M_{ij}(X_{1,i},X_{1,j})$ record the target sequencespecific parameters in the PLM and MI matrices. Thus, they are considered as extra 2D features.

In addition to the 2D features, there are four 1D features collected. The first two are the field parameter *e* of the Potts model and the MI matrix where j=i. HMM profile features (30 descriptors for each position) are also considered as the third component by building a profile hidden Markov models from the input alignment using the hhmake program in the hhsuite package.³⁹ The last feature encodes the categorical target sequence with one-hot-encoding.

2.4 | Deep ResNet for multiple geometric map prediction

As shown in Figure 1, the above 1D and 2D input features are fed into 10 1D residual blocks²⁴ and 10 2D residual blocks, respectively. A

general structure of a residual block is shown in Figure 1, where a shortcut link is added from previous layers to the output, compared to the traditional convolutional neural networks. Here one weight layer contains two layers, that is, one convolutional layer and an instance normalization layer,⁴⁰ collected sequentially. The transformed 1D features with 32 channels will be tiled vertically and horizontally and concatenated with transformed 2D features (64 channels). The composited 2D features ($32^*2 + 64 = 128$ channels) will further go through 40 2D residual blocks. The prediction layer for each potential term performs a simple pixel-wise linear transformation to the desired channel size, prior to a softmax layer. Dilation is applied for both 1D and 2D convolutional layers with cycling of 1, 2, 4, 8, and 16. The padding size is then set accordingly to ensure the consistency of feature signal spatial shapes. Dropout is used in all residual blocks and the dropout rate was set to 0.2 globally.

The model was trained by the supervision of distance between inter-residue C_{β} atoms (C_{α} for glycine). The distance value is discretized into 36 equal-width bins from 2 Å to 20 Å, with additional two bins representing distance less than 2 Å and over 20 Å. Thus, the model can be trained with cross-entropy loss over all residue pairs. In addition, some auxiliary tasks, that is, inter- C_{α} atom distance, interresidue orientation angles,²⁸ and H-bond geometry terms defined on long-range neighboring C_{α} atoms⁴¹ are also considered in a multi-task learning strategy.⁴²

2.5 | Template free structure modeling built on DeepPotential restraints

To explore the effectiveness of predicted inter-residue distance for protein structure prediction, we also utilized the predicted C_{β} distance distribution, along with the outputs of auxiliary tasks (predicted C_{α} distance and inter-residue orientation angles) as restraints for ab initio protein structure prediction in the form of potentials. The negative log of geometric distributions was smoothed by a cubic spline into smooth potentials so that they could be optimized by gradient-descent-based methods, for example, L-BFGS implemented by the PyRosetta package.⁴³ Starting from a random initial conformation, the structure was iteratively optimized using L-BFGS. To find the global lowest energy conformation, we used an iterative strategy, and at each iteration, random noise ($\pm 20^{\circ}$) in the backbone torsion angle space was added to the previous confirmation for further energy minimization. The conformation with the lowest energy value was kept for final submission.

3 | RESULTS

We analyzed the performance of DeepPotential on 22 free modeling (FM) (T1070-D1 was excluded since there were not sufficient longrange contacts in this target) and 14 template-based modeling (TBM)/ FM targets from CASP14 (T1085-D2 was excluded because of the unavailability of its experimental structure). The performance was evaluated mainly in two tasks, that is, inter-residue contact and distance prediction. For the evaluation of predicted contacts, the conventional long-range Top-N (N = L/10, L/5, L/2, and L) precision is reported. For predicted discrete distance distribution, we report the results with two different evaluation indexes. The first index is the long-range Top-N bin classification ACCuracy (BACC), considering that the distance prediction is formulated as a multi-class classification problem:

$$BACC = \frac{1}{N} \sum_{i=1}^{N} \mathbb{I}[\operatorname{argmax}(P_i) = k(D_i)].$$
(4)

Here, P_i is the distance distribution of *i*th residue pair ranked by the probability of the distance less than 20 Å. argmax(P_i) finds the predicted distance bin of residue pair *i*; $k(D_i)$ returns the bin of experimental distance D_i . In CASP14, there are 10 bins for distance prediction, representing <4 Å, 4–6 Å, 6–8 Å, ..., 16–18 Å, 18–20 Å, and > 20 Å. It should be noted that there is a natural order among distance bins in such a formulation. Such order sorts the distance bins according to the definition of bins in distance. Thus, we also utilize a second index, long-range Top-*N* mean bin error (MBE), to provide a closer evaluation of distance prediction by considering the closeness in distance bins:

$$MBE = \frac{1}{N} \sum_{i=1}^{N} |\operatorname{argmax}(P_i) - k(D_i)|.$$
 (5)

A lower MBE index means better prediction, and the ideal case of the MBE value should be 0. For distance prediction, a larger threshold (20 Å) was considered in CASP14, compared to the threshold of 8 Å for contacts. In this regard, we set N = L/2, L, 2L, and 5L when evaluating distance prediction.

3.1 | Contact and distance prediction performance in CASP14

The accuracy of long-range contact prediction on CASP14 FM and FM/TBM targets for Groups 010 and 024 are listed in Table 1. On average, the Top-L/5 precisions were 0.638 and 0.615, and 16 and 15 out of all 22 FM targets have a precision ≥0.5 for Group 010 and Group 024, respectively. On 14 FM/TBM targets, the average Top-L/5 precisions increased to 0.868 and 0.862, and 13 out of 14 have a precision ≥0.5 for both of the two Groups. Such a difference is likely because FM/TBM targets have more related samples in the training set of DeepPotential than FM targets, because by definition they are evolutionarily closer than the FM targets to the PDB structure by which DeepPotential was trained. To verify, we searched each of CASP14 domains against the training structures of DeepPotential by TM-align. The average TM-score between FM/TBM targets and their best template is 0.587, significantly higher than 0.497 for FM targets, with an unpaired t-test p-value of 8e-04. FM/TBM targets also, on average, have 258 related structures (TM-score >0.5) in the training

TABLE 1 Mean accuracy and standard error (in brackets) of long-range predicted contacts for DeepPotential's two Groups in CASP14

	FM				TBM/FM			
Group	L/10	L/5	L/2	L	L/10	L/2	L/5	L
010	0.686 (0.086)	0.638 (0.081)	0.534 (0.068)	0.396 (0.051)	0.898 (0.057)	0.868 (0.061)	0.771 (0.062)	0.620 (0.057)
024	0.653 (0.090)	0.615 (0.084)	0.502 (0.068)	0.386 (0.051)	0.872 (0.064)	0.862 (0.059)	0.784 (0.061)	0.629 (0.059)

Abbreviation: FM, free modeling; TBM, template-based modelling.

set. Meanwhile, for FM targets, the corresponding number is only 28. Such data suggest very different relationships to training samples of DeepPotential for FM and FM/TBM targets in CASP14. In addition, FM/TBM targets also have more sequence homologs than FM targets (167.8 vs. 79.7 of average Neff), which provide more reliable features, especially coevolutionary features. In Table 1, we also compare the performance of two Groups based on the DeepPotential pipeline. Although Group 010 had slightly higher precisions for all evaluation indexes, the differences are not statistically significant, with 2-tailed test p-values of .196, .410, .350, and .787, respectively, for both FM and FM/TBM targets. Figure S1 shows the correlation between the confidence score of selected MSAs and contact precision for Group 010 and Group 024 respectively. Both two configurations of confidence score have high correlations with their Top-L long-range precisions, with Pearson correlation coefficient (PCC) of 0.743 and 0.752 for Group 010 and Group 024 respectively. Such data suggest that MSA selection based on the mean Top-N probability scores has robust performance regardless of the specific choice of the threshold for confidence score definition.

In Table 2, we report the two metrics for distance prediction, that is. BACC and MBE. A similar pattern can be observed that FM/TBM targets achieved higher accuracy or lower bin error relative to FM targets. Interestingly, unlike contact precisions that drop sharply when evaluating Top-L/10 to Top-L contacts, we observe a relatively gentle slope even evaluating from Top-L/2 to Top-5L ranked residue pairs. Taking the performance of Group 010 on TBM/FM targets as an example, the accuracy of predicted contacts, with a threshold of 8 Å, drops from 0.898 to 0.620 (31.0% of decrease) when evaluating from Top-L/10 to Top-L contacts. Meanwhile, for distance, the Bin accuracy only drops from 0.562 to 0.456 (18.9% of reduction) when selecting from Top-L/2 to Top- 5^*L distance predictions, sorted by the cumulative probability less than 20 Å. Such an observation suggests a stable accuracy for distance prediction when sorting according to $P(d < 20\text{\AA})$. In addition to the bin accuracy, Table S1 also shows other multi-class classification evaluation indexes, that is, precision, recall and F1-score, from Top-L/2 to Top-5*L distance predictions. The evaluation indexes for each target are the average over all classes (bins). Similar to the precision of contact maps, Group 024 tends to have more accurate distance prediction for FM while Group 010 favors FM/TBM targets, respectively. For the average Top-N bin error presented in Table 2, it can be observed that both of the two Groups maintain relatively low values. For example, the long-range Top-L average bin errors on FM and FM/TBM targets are 1.336 (1.415) and 0.593 (0.532) for Group 010 (024), respectively. It is notable that both

Groups successfully provide 11 FM and 10 FM/TBM targets that have an MBE lower than 1.0, even up to Top-5*L* selected residue pairs. Precise continuous distance prediction, utilized as potentials, will be critical to guide high-accuracy protein structure prediction.

We address the improvements of DeepPotential compared to our previous model, TripletRes, in CASP13 (denoted as TripletRes_CASP13) by head-to-head comparisons in Figure 3. We compare the long-range Top-L/5 TripletRes from their own default MSAs (Figure 3A) and from the same MSAs by DeepPotential (Figure 3B), with the results of Group 010 in CASP14 as an illustration. With the original pipeline, including the previous MSA construction pipeline, TripletRes CASP13 achieved a Top-L/5 precision of 0.310 on 36 FM and FM/TBM targets; when the newly constructed MSAs were applied, the precision rose to 0.398, an improvement of 28.7%, suggesting the new pipeline generates higherquality MSAs over multiple sequence databases. Meanwhile, the precision for Group 010 in CASP14 on the same dataset was 0.483. 56.1%. and 21.3% higher than the control TripletRes CASP13 from the default and the same MSAs, respectively. The corresponding p-values are 3.58e-08 and 2.35e-07 respectively. Besides, a head-to-head comparison of Top-L long-range contact precision between TripletRes (Post-CASP13), introduced in our previous research.²² and Group 010 is presented in Figure S2. The TripletRes (Post-CASP13) was trained using the same dataset with DeepPotential. The Top-L precision increases from 0.398 to 0.418 after the re-training, but still lower than the precision of 0.483 (p-value = 1.0e-05) obtained by DeepPotential. These significant improvements could be attributed to the complete set of features coupled with multi-task learning framework and the larger amount of training data. However, the results show that for those targets that DeepPotential and TripletRes models had similar performance, where limited superiority could be found for DeepPotential. Our next work should be focusing on proper training strategies for those extremely hard targets.

3.2 | Constructing MSAs from gigantic databases improves performance

In CASP14, we utilized a progressive pipeline for MSA construction from whole-genome and meta-genome databases with an enormous number of sequences. The employment of those sequence databases becomes a critical element to achieve higher quality contact/distance prediction, especially for FM targets. In Figure 4A, we show the distributions of long-range Top-*L*/5 precisions of 22 FM targets by gradually adding the considered sequence databases. Other factors, for

example, MSA selection score configuration, are fixed. Comparable performance, that is, precisions of 0.402 versus 0.416 can be observed with two whole-genome databases. In contrast, significant improvements can be observed when meta-genome databases are utilized by our pipeline. When Metaclust and BFD databases were added, the precision increased to 0.495 and 0.610, 19.0% and 46.6% higher than the best results of whole-genome databases, respectively. Additionally, when two extra databases, that is, the Mgnify and IMG/M databases, are considered, mean precisions of 0.647 and 0.660 can be observed respectively. There is also a consistent improvement in median precision from 29.2% to 77.2% with the consideration of more databases. Such data suggest the fundamental role of the MSA construction pipeline and the great usefulness of the utilization of meta-genome databases. It can be observed in Figure 4B that the Neff of selected MSAs is continuously increasing along with the addition of the corresponding databases. The detailed mean (median) logarithm Neff values are 0.21 (-0.11), 0.54 (0.06), 1.21 (0.52), 1.9 (2.07), 2.19 (2.41), and 2.69 (3.35) when adding the corresponding databases. The rapid improvements of contact-map precision are likely highly driven by the MSAs with continuously increasing Neff values at each MSA stage.

A more quantitative analysis about the impact of MSA quality on the performance of contact/distance prediction of DeepPotential is shown in Figure 5A–C. In Figure 5A, the long-range Top-L/5 precision of Group 024 in CASP14 versus the Neff values of the MSAs is presented. The PCC between precision and the common logarithm of Neff is 0.364, indicating a modest correlation. The precision of DeepPotential is less dependent on Neff than TripletRes_CASP with the same input MSAs, whose corresponding PCC is 0.419. As shown in the left-upper block in Figure 5A, 10 out of 14 targets with a Neff value lower than 10 have a precision \ge 0.5, which is 3 more than that of TripletRes_CASP; this may help explain the lower correlation coefficient seen for DeepPotential than TripletRess_CASP.

In addition, we also found that the PCCs are very different for FM and FM/TBM targets. When only FM targets are considered, the correlation is 0.382, slightly higher than the combination of FM and FM/TBM targets. The reason for higher correlation coefficient for FM targets should be the extreme cases (red dots in the lower-left block in Figure 5A) where their Neff values are all less than 1, and the corresponding precisions are <0.3. In contrast, for FM/TBM targets, the correlation became 0.181 with a *p*-value of .535 where a statistically significant correlation could not be detected. Still, we can find one FM/TBM target that has a high Neff value and precision <0.5, T1080-D1. Nevertheless, the 3D model built based on the geometric restraints predicted by DeepPotential eventually had the correct fold, with a TM-score⁴⁴ of 0.503, possibly because of the modest Top-*L* bin error of 1.37.

For FM targets with Neff \geq 10, 10 out of 13 had precisions over 0.5, and the 3 exceptions are T1029-D1, T1093-D1, and T1093-D3. Interestingly, for T1093-D1 and T1093-D3, if we re-run DeepPotential based on MSAs built with domain sequences, the corresponding precisions will be boosted to 0.821 and 1.0, even with lower Neff values of 10.91 and 17.13. We simply visualize the MSA

Performance of long-range Top-N distance prediction for DeepPotential's two Groups in CASP14 with standard error in brackets 2 TABLE

Σ

TBM/FM

Index	Group	L/2	7	21	5L	L/2	r	21	5L
Bin acc	010	0.387 (0.048)	0.369 (0.044)	0.370 (0.041)	0.338 (0.037)	0.562 (0.039)	0.535 (0.037)	0.502 (0.038)	0.456 (0.039)
	024	0.374 (0.047)	0.373 (0.045)	0.362 (0.043)	0.336 (0.039)	0.594 (0.034)	0.581 (0.040)	0.526 (0.039)	0.477 (0.044)
Bin error	010	1.341 (0.258)	1.336 (0.248)	1.343 (0.248)	1.495 (0.229)	0.505 (0.057)	0.593 (0.083)	0.717 (0.155)	0.835 (0.162)
	024	1.384 (0.268)	1.415 (0.266)	1.441 (0262)	1.559 (0.241)	0.453 (0.052)	0.532 (0.084)	0.699 (0.156)	0.826 (0.174)

Abbreviation: FM, free modeling; TBM, template-based modelling

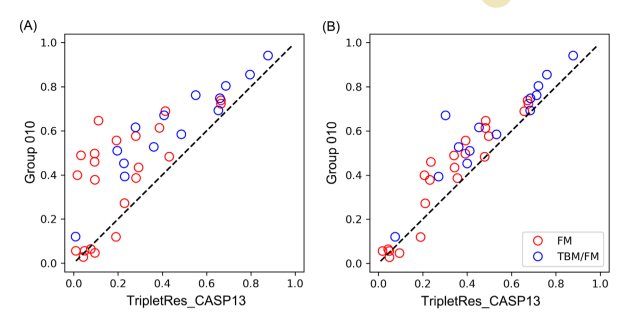


FIGURE 3 Head-to-head comparison of long-range Top-*L* precision between our pipeline in CASP13 and CASP14. (A) Contacts predicted using their own multiple sequence alignments (MSAs). (B) Contacts predicted using same MSAs generated by the newer pipeline

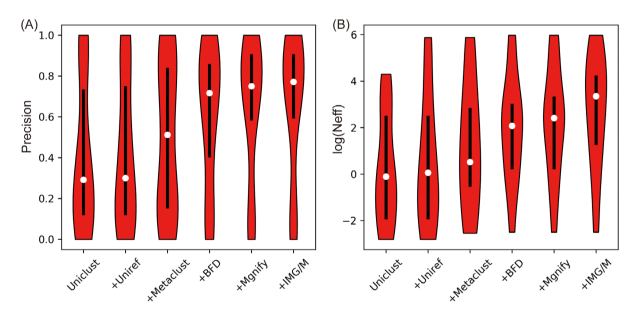


FIGURE 4 Increases of contact long-range Top-*L*/5 precision and average log(Neff) with the consideration of sequence databases illustrated in violin plots. White dots indicate the median value. Vertical black lines indicate 25% to 75% percentile. (A) Precision changes as more databases are used. (B) Log(Neff) changes when more databases are employed

consensus using Sequence logos⁴⁵ with MSAs generated with fulllength sequence and only domain sequence in Figures S3 and S4 for T1093-D1 and T1093-D3, respectively. The graphical characters displayed in Figures S3 and S4 represent significant residues and subtle sequence patterns in the corresponding MSAs. The height of each letter is made proportional to its frequency, and the letters are sorted so the most common one is on top. We found that the patterns of consensus sequences are visibly different for domain and full-length MSAs. The average Kullback–Leibler divergencies over positional amino acid compositions between domain and full-length MSAs are

1.30 and 1.19 for T1093-D1 and T1093-D3 respectively. For MSAs from the full-length sequence, the average sequence identity between aligned sequences and query sequence are 0.098 and 0.118 for T1093-D1 and T1093-D3, respectively, which are much lower than those of the domain sequence MSAs (0.202 and 0.232). The differences in distribution and sequence identity suggest that there could be noisy alignments in our selected MSAs, despite their high Neff value.

Figure 5B,C shows the long-range Top-*L* bin classification accuracy and bin error versus Neff respectively. We found a relatively

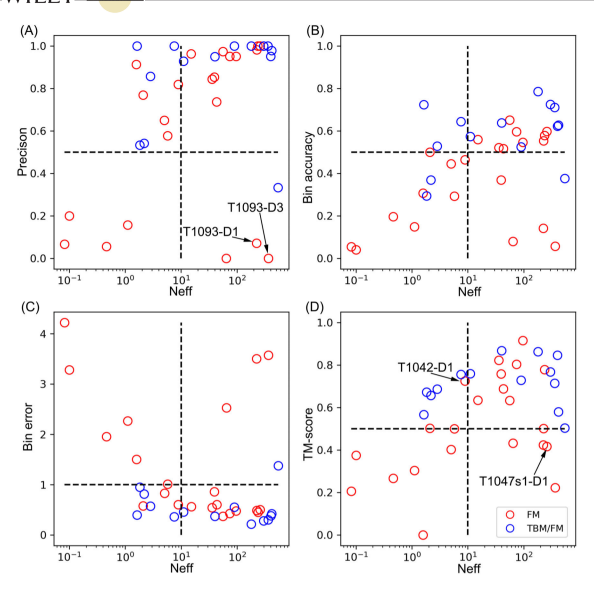


FIGURE 5 Illustration of the effect of multiple sequence alignments (MSAs) on the performance DeepPotential. (A) The precision of longrange Top-*L*/5 contact prediction versus Neff of MSAs. (B, C) Mean bin accuracy and mean bin error of long rang Top-*L* distance prediction versus Neff of MSAs. (D) TM-score of 3D models based on the prediction of DeepPotential versus Neff of MSAs

higher PCC of 0.465 between classification accuracy and the common logarithm of Neff, compared to that between contact precision and the latter. The correlation indicates that there is much room for improvement of the current model for further higher resolution distance prediction, especially for FM targets (PCC = 0.474) with limited sequence homologs. For Top-*L* bin error, we can observe a modest correlation; the corresponding PCCs are -0.363 and -0.386 for all targets and FM targets, respectively.

3.3 | Structure modeling based on DeepPotential in CASP14

The final goal of contact/distance prediction is to assist protein structure prediction, and the best way to evaluate the quality of distance prediction is to analyze the quality of predicted protein structure purely based on predicted distance. On 36 FM and FM/TBM target domains, the 3D models based on predicted geometric descriptors by DeepPotential achieved a mean TM-score of 0.591. When FM domains are evaluated, the mean TM-score becomes 0.514, where 12 have correctly predicted global folds (TM-score ≥0.5). Surprisingly, our submissions had all 14 FM/TBM folds correctly predicted, with a mean TM-score of 0.712, ranging from 0.503 to 0.868, even though structural templates were not used. Figure 5D shows the TM-score and the Neff of 36 FM and FM/TBM targets. Similar to the previous observations, there is a modest correlation between the TM-score and the common logarithm of Neff (PCC = 0.482), but for FM/TBM targets, the correlation is weak, even not statistically significant (PCC = 0.158, p-value = .61). Figure S5 plots the correlation between the TM-score of predicted structures and the precision of long-range Top-L/2 predicted contacts. There is a significant correlation between TM-score and precision (PCC = 0.788), indicating that the accuracy of

predicted structures is highly dependent on the quality of deep learning predictions. There are hardly any cases with low contact precision but high TM-score, which is, however, reasonable since we used a basic protein folding procedure without other information sources, for example, template information.

In Figure 6, we present an example FM domain, T1042-D1, to show that DeepPotential can predict reliable distance potentials with limited sequence alignments (Neff = 8.85) for accurate protein structure prediction. T1042-D1 is the 8th domain of a viral protein⁴⁶ (PDB ID: 6VR4) officially partitioned by CASP. In this case, the long-range Top-*L*/5 precision was 0.818 and for contact map prediction, the Top-*L* bin accuracy and bin error were 0.464 and 0.601, respectively. As shown in Figure 6A, the predicted distance bin map in general successfully recovered the distance patterns of the experimental structure except one region with some geometric interactions between N and C-terminals. Nevertheless, the predicted structure based on the distance map has a TM-score of 0.725.

Despite this successful prediction, we found that domain T1047s1-D1 has an unexpectedly low TM-score. The contact Top-L/5 precision, Top-L bin accuracy, and Top-L bin error are 1.0, 0.597, and 0.502 respectively. However, we observed some noisy distance predictions between a beta-sheet region (residue 50–125) and the C-terminal structure region (residue 125–211) in Figure S6B. Those noisy predictions pulled the two regions together and destroyed the structure of the beta-sheet region (Figure S6D). Thus, our 3D model had only a TM-score of 0.416, although the TM-score of the region near the C-terminal achieved 0.632 (Figure S6E). In Figure S6A, we plot the residue-wise prediction based on Potts model coupling parameters and found observable noisy signals. Thus, the noisy prediction partly came from the MSA and its features.

This example also exposed one weakness of our protein contact/ distance and structure prediction pipeline. The current distance prediction is only formulated to predict distance under a fixed threshold, that is, 20 Å. Thus, the subsequent restraint-based protein folding strategy will be influenced by possibly noisy distance (or other geometric descriptors) potentials. If a region has sparse connections with other parts, the noise could completely mislead the folding. Seeking better formulations, for example, real-distance, to obtain reliable geometric restraints without the limitation of a threshold should help build better 3D models, not only for targets similar to T1047s1, but also for the modeling of interdomain or inter-chain structures. One feasible way could be predicting the parameters of distribution (e.g., the expected value [or mean] and standard deviation of the variable's natural logarithm for log-normal distribution) for the inter-atom distance modeling. The predicted distribution map could be easily converted to smooth potentials for protein folding. Whether such prediction would result in more accurate protein folding still requires further examination.

4 | DISCUSSION

We have introduced DeepPotential, which participated in CASP14 for contact and distance prediction. Our model takes an ensemble of complementary features directly extracted from selected MSAs constructed by progressive searching against multiple sequence databases. Detailed analysis showed that the proposed method can produce relatively high precision contact maps that significantly outperform our previous method in CASP13. We also showed that the distance predictions can be used as reliable restraints for protein structure prediction.

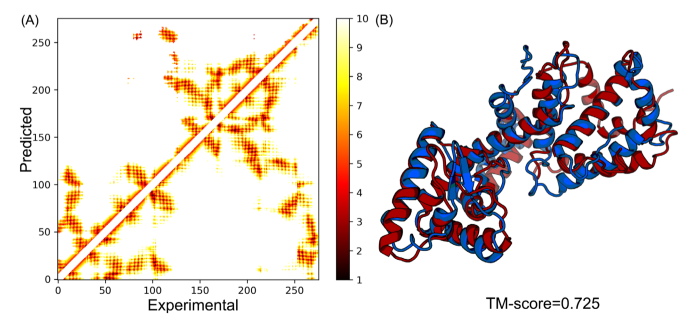


FIGURE 6 An illustrative example of CASP14 domain T1042-D1. (A) Comparison of predicted discrete distance map by DeepPotential and the distance map of the experimental structure. The distance bins are defined according to CASP format. (B) Superposition of submitted first model (blue) and experimental structure (red)

4.1 | What went right?

The results in CASP14 confirm the conclusion of our previous strategy for contact/distance prediction in CASP13, that is, constructing and selecting MSAs from multiple protein sequence databases can significantly improve performance, especially for FM targets. In CASP14, we further extended our MSA construction pipeline by the utilization of large-scale meta-genome databases, which brought a further boost in contact/distance and structure prediction. In addition, the ensemble of multiple raw coevolution features which extract complementary information from MSAs, together with a multi-task learning scheme, contributed to the advantage of DeepPotential over previous approaches.

4.2 | What went wrong?

The DeepPotential model predicts distance and other geometric terms marginally, ignoring the inherent relationships among geometric terms and residue pairs in the loss function. Designing a formulation to effectively model the joint distributions between residue pairs or geometric terms should be a necessity in our future work. Also, the convolutional neural networks used in DeepPotential have a relatively long max path between features of residue pairs. Considering the revolutionary results obtained by AlphaFold2⁴⁷ in CASP14 and the recent success of Transformer⁴⁸ applied in protein sequence modeling,⁴⁹⁻⁵¹ the Transformer framework with a direct max path should be considered in our future work to model direct long-range interactions in space.

ACKNOWLEDGMENTS

We thank Dr. Jianyi Yang and Dr. Ivan Anishchenko for their insightful discussions. This work is supported in part by the National Institute of General Medical Sciences (GM136422, S10OD026825 to Yang Zhang), the National Institute of Allergy and Infectious Diseases (Al134678 to Yang Zhang), the National Science Foundation (IIS1901191, DBI2030790, MTM2025426 to Yang Zhang), the National Natural Science Foundation of China (62072243, 61772273 to Dong-Jun Yu), and the Natural Science Foundation of Jiangsu (BK20201304 to Dong-Jun Yu). DeepPotential was trained using the Extreme Science and Engineering Discovery Environment (XSEDE), which is supported by National Science Foundation (ACI-1548562). The work was done when Yang Li visited the University of Michigan.

DATA AVAILABILITY STATEMENT

All relevant data are within the manuscript and its Supporting Information files.

ORCID

Yang Li D https://orcid.org/0000-0003-2480-1972 Chengxin Zhang D https://orcid.org/0000-0001-7290-1324 Wei Zheng D https://orcid.org/0000-0002-2984-9003 Xiaogen Zhou D https://orcid.org/0000-0001-6839-1923

Eric W. Bell https://orcid.org/0000-0002-3419-4398 Yang Zhang https://orcid.org/0000-0002-2739-1916

REFERENCES

- Zhang Y. Progress and challenges in protein structure prediction. Curr Opin Struct Biol. 2008;18(3):342-348.
- Abriata LA, Tamò GE, Dal PM. A further leap of improvement in tertiary structure prediction in CASP13 prompts new routes for future assessments. *Proteins*. 2019;87(12):1100-1112.
- Abriata LA, Tamò GE, Monastyrskyy B, Kryshtafovych A, Dal Peraro M. Assessment of hard target modeling in CASP12 reveals an emerging role of alignment-based contact prediction methods. *Proteins*. 2018;86(S1):97-112.
- Fitch WM, Markowitz E. An improved method for determining codon variability in a gene and its application to the rate of fixation of mutations in evolution. *Biochem Genet*. 1970;4(5):579-593.
- Korber BT, Farber RM, Wolpert DH, Lapedes AS. Covariation of mutations in the V3 loop of human immunodeficiency virus type 1 envelope protein: an information theoretic analysis. *Proc Natl Acad Sci.* 1993;90(15):7176.
- Dunn SD, Wahl LM, Gloor GB. Mutual information without the influence of phylogeny or entropy dramatically improves residue contact prediction. *Bioinformatics*. 2008;24(3):333-340.
- 7. Göbel U, Sander C, Schneider R, Valencia A. Correlated mutations and residue contacts in proteins. *Proteins*. 1994;18(4):309-317.
- Morcos F, Pagnani A, Lunt B, et al. Direct-coupling analysis of residue coevolution captures native contacts across many protein families. *Proc Natl Acad Sci.* 2011;108(49):E1293.
- Jones DT, Buchan DWA, Cozzetto D, Pontil M. PSICOV: precise structural contact prediction using sparse inverse covariance estimation on large multiple sequence alignments. *Bioinformatics*. 2012; 28(2):184-190.
- Ma J, Wang S, Wang Z, Xu J. Protein contact prediction by integrating joint evolutionary coupling analysis and supervised learning. *Bioinformatics*. 2015;31(21):3506-3513.
- 11. Ekeberg M, Hartonen T, Aurell E. Fast pseudolikelihood maximization for direct-coupling analysis of protein structure from many homologous amino-acid sequences. *J Comput Phys.* 2014;276:341-356.
- Jones DT, Singh T, Kosciolek T, Tetchner S. MetaPSICOV: combining coevolution methods for accurate prediction of contacts and long range hydrogen bonding in proteins. *Bioinformatics*. 2015;31(7):999-1006.
- He B, Mortuza SM, Wang Y, Shen H-B, Zhang Y. NeBcon: protein contact map prediction using neural network training coupled with naïve Bayes classifiers. *Bioinformatics*. 2017;33(15):2296-2306.
- 14. Buchan DWA, Jones DT. Improved protein contact predictions with the MetaPSICOV2 server in CASP12. *Proteins*. 2018;86(S1):78-83.
- Wang Z, Xu J. Predicting protein contact map using evolutionary and physical constraints by integer programming. *Bioinformatics*. 2013; 29(13):i266-i273.
- Wang S, Sun S, Li Z, Zhang R, Xu J. Accurate de novo prediction of protein contact map by ultra-deep learning model. *PLoS Comput Biol.* 2017;13(1):e1005324.
- Adhikari B, Hou J, Cheng J. DNCON2: improved protein contact prediction using two-level deep convolutional neural networks. *Bioinformatics*. 2017;34(9):1466-1472.
- Liu Y, Palmedo P, Ye Q, Berger B, Peng J. Enhancing evolutionary couplings with deep convolutional neural networks. *Cell Syst.* 2018; 6(1):65-74.e63.
- Golkov V, Skwark MJ, Golkov A, et al. Protein contact prediction from amino acid co-evolution using convolutional networks for graphvalued images. Paper presented at: NIPS; 2016.
- Jones DT, Kandathil SM. High precision in protein contact prediction using fully convolutional neural networks and minimal sequence features. *Bioinformatics*. 2018;34(19):3308-3315.

- Li Y, Zhang C, Bell EW, Yu D-J, Zhang Y. Ensembling multiple raw coevolutionary features with deep residual neural networks for contact-map prediction in CASP13. *Proteins*. 2019;87(12):1082-1091.
- Li Y, Zhang C, Bell EW, et al. Deducing high-accuracy protein contactmaps from a triplet of coevolutionary matrices through deep residual convolutional networks. *PLoS Comput Biol.* 2021;17(3):e1008865.
- Li Y, Hu J, Zhang C, Yu D-J, Zhang Y. ResPRE: high-accuracy protein contact prediction by coupling precision matrix with deep residual neural networks. *Bioinformatics*. 2019;35(22):4647-4655.
- He K, Zhang X, Ren S, Sun J. Deep residual learning for image recognition. Paper presented at: Proceedings of the IEEE Conference on Computer Vision and Pattern Recognition; 2016.
- Senior AW, Evans R, Jumper J, et al. Improved protein structure prediction using potentials from deep learning. *Nature*. 2020;577(7792): 706-710.
- Xu J. Distance-based protein folding powered by deep learning. Proc Natl Acad Sci. 2019;116(34):16856-16865.
- Greener JG, Kandathil SM, Jones DT. Deep learning extends de novo protein modelling coverage of genomes using iteratively predicted structural constraints. *Nat Commun.* 2019;10(1):3977.
- Yang J, Anishchenko I, Park H, Peng Z, Ovchinnikov S, Baker D. Improved protein structure prediction using predicted interresidue orientations. *Proc Natl Acad Sci.* 2020;117(3):1496.
- Yu F, Koltun V, Funkhouser T. Dilated residual networks. Paper presented at: Proceedings of the IEEE Conference on Computer Vision and Pattern Recognition; 2017.
- Berman HM, Westbrook J, Feng Z, et al. The Protein Data Bank. Nucleic Acids Res. 2000;28(1):235-242.
- Fu L, Niu B, Zhu Z, Wu S, Li W. CD-HIT: accelerated for clustering the next-generation sequencing data. *Bioinformatics*. 2012;28(23): 3150-3152.
- Steinegger M, Söding J. Clustering huge protein sequence sets in linear time. Nat Commun. 2018;9(1):2542.
- Steinegger M, Mirdita M, Söding J. Protein-level assembly increases protein sequence recovery from metagenomic samples manyfold. *Nat Methods*. 2019;16(7):603-606.
- Mitchell AL, Almeida A, Beracochea M, et al. MGnify: the microbiome analysis resource in 2020. *Nucleic Acids Res.* 2020;48(D1):D570-D578.
- Chen IMA, Chu K, Palaniappan K, et al. The IMG/M data management and analysis system v.6.0: new tools and advanced capabilities. *Nucleic Acids Res.* 2021;49(D1):D751-D763.
- Zhang C, Zheng W, Mortuza SM, Li Y, Zhang Y. DeepMSA: constructing deep multiple sequence alignment to improve contact prediction and fold-recognition for distant-homology proteins. *Bioinformatics*. 2020;36(7):2105-2112.
- Remmert M, Biegert A, Hauser A, Söding J. HHblits: lightning-fast iterative protein sequence searching by HMM-HMM alignment. *Nat Methods*. 2012;9(2):173.

- Johnson LS, Eddy SR, Portugaly E. Hidden Markov model speed heuristic and iterative HMM search procedure. *BMC Bioinformatics*. 2010;11(1):431.
- Steinegger M, Meier M, Mirdita M, Vöhringer H, Haunsberger SJ, Söding J. HH-suite3 for fast remote homology detection and deep protein annotation. *BMC Bioinformatics*. 2019;20(1):473.
- Ulyanov D, Vedaldi A, Lempitsky V. Instance normalization: the missing ingredient for fast stylization. arXiv preprint arXiv:160708022. 2016.
- 41. Yang J, Yan R, Roy A, Xu D, Poisson J, Zhang Y. The I-TASSER suite: protein structure and function prediction. *Nat Methods*. 2015;12(1):7-8.
- Thrun S. Is learning the n-th thing any easier than learning the first? Paper presented at: Advances in Neural Information Processing Systems; 1996.
- Chaudhury S, Lyskov S, Gray JJ. PyRosetta: a script-based interface for implementing molecular modeling algorithms using Rosetta. *Bioinformatics*. 2010;26(5):689-691.
- 44. Xu J, Zhang Y. How significant is a protein structure similarity with TM-score = 0.5? *Bioinformatics*. 2010;26(7):889-895.
- 45. Schneider TD, Stephens RM. Sequence logos: a new way to display consensus sequences. *Nucleic Acids Res.* 1990;18(20):6097-6100.
- Drobysheva AV, Panafidina SA, Kolesnik MV, et al. Structure and function of virion RNA polymerase of a crAss-like phage. *Nature*. 2021;589(7841):306-309.
- Jumper J, Evans R, Pritzel A, et al. Highly accurate protein structure prediction with AlphaFold. *Nature*. 2021. https://doi.org/10.1038/ s41586-021-03819-2
- Vaswani A, Shazeer N, Parmar N, et al. Attention is all you need. arXiv preprint arXiv:170603762. 2017.
- Elnaggar A, Heinzinger M, Dallago C, et al. ProtTrans: towards cracking the language of Life's code through self-supervised deep learning and high performance computing. *bioRxiv*. 2020.
- 50. Rao R, Liu J, Verkuil R, et al. MSA transformer. bioRxiv. 2021.
- 51. Rao R, Meier J, Sercu T, Ovchinnikov S, Rives A. Transformer protein language models are unsupervised structure learners. *bioRxiv*. 2020.

SUPPORTING INFORMATION

Additional supporting information may be found in the online version of the article at the publisher's website.

How to cite this article: Li Y, Zhang C, Zheng W, et al. Protein inter-residue contact and distance prediction by coupling complementary coevolution features with deep residual networks in CASP14. *Proteins*. 2021;89(12):1911-1921. doi: 10.1002/prot.26211