**Combining Pairwise Structural Similarity and Deep Learning Interface Contact Prediction to Estimate Protein Complex Model Accuracy in CASP15**

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## **Abstract**

Estimating the accuracy of quaternary structural models of protein complexes and assemblies (EMA) is important for predicting quaternary structures and applying them to studying protein function and interaction. The pairwise similarity between structural models is proven useful for estimating the quality of protein *tertiary* structural models, but it has been rarely applied to predicting the quality of *quaternary* structural models. Moreover, the pairwise similarity approach often fails when many structural models are of low quality and similar to each other. To address the gap, we developed a hybrid method (MULTICOM\_qa) combining a pairwise similarity score (PSS) and an interface contact probability score (ICPS) based on the deep learning inter-chain contact prediction for estimating protein complex model accuracy. It blindly participated in the 15th Critical Assessment of Techniques for Protein Structure Prediction (CASP15) in 2022 and ranked first out of 24 predictors in estimating the global accuracy of assembly models. The average per-target correlation coefficient between the model quality scores predicted by MULTICOM\_qa and the true quality scores of the models of CASP15 assembly targets is 0.66. The average per-target ranking loss in using the predicted quality scores to rank the models is 0.14. It was able to select good models for most targets. Moreover, several key factors (i.e., target difficulty, model sampling difficulty, skewness of model quality, and similarity between good/bad models) for EMA are identified and analayzed. The results demonstrate that combining the multi-model method (PSS) with the complementary single-model method (ICPS) is a promising approach to EMA.

**Keywords**: Estimation of protein model accuracy, protein quaternary structure prediction, deep learning

## **1. Introduction**

Most proteins must fold into some specific 3D shape called tertiary structures to carry out their biological function. Two or more proteins (more precisely protein chains) often interact to form a protein complex or assembly to collaboratively perform a complicated biological function such as transduction of an extracellular signal into a cell to trigger gene expression. Therefore, predicting the tertiary structure of a single protein (monomer) and the quaternary structure of a protein complex/assembly (multimer) from its sequence has been a major pursuit of the scientific community for the last few decades because the experimental methods of determining protein structures are expensive and time-consuming and therefore can only be applied to a small fraction of proteins.

To rigorously and objectively assess the progress in protein structure prediction, the Critical Assessment of Techniques for Protein Structure Prediction (CASP) was launched in 1994 and continued to be held every two years [(Moult et al., 1995; Kryshtafovych et al., 2014; Moult et al., 2016; Kryshtafovych et al., 2019; Kwon et al., 2021)](https://www.zotero.org/google-docs/?0I5FE3), with the latest CASP15 experiment concluded in 2022. Stimulated by the CASP experiments, significant progress has been made in protein tertiary structure prediction. Particularly, with the development of deep learning-based methods over the last decade [(Eickholt & Cheng, 2012; Wang et al., 2017; Kandathil et al., 2019; Hou et al., 2019; Senior et al., 2019; Zheng et al., 2019; Senior et al., 2020; Yang et al., 2020; Baek et al., 2021;Jumper et al., 2021,Liu et al., 2022)](https://www.zotero.org/google-docs/?tHwnPr), the prediction of protein tertiary structures had steadily improved from one CASP experiment to another until a deep learning method - AlphaFold2 (Jumper et al., 2021) drastically advanced the filed with its outstanding capability of predicting tertiary structures with the quality close to that of the experimental techniques such as X-ray crystallography in the 2020 CASP14 experiment. The success of AlphaFold2 and other deep learning methods has revolutionised the prediction and application of protein tertiary structures in various biological research.

As the tertiary structure prediction problem has been largely solved, the field started to focus more on predicting the *quaternary* structures of protein complexes and assemblies, which also had a long history of development but did not progress as fast as the tertiary structure prediction (Lensink et al., 2016, 2018, 2021). However, the situation started to change as more and more deep learning methods were developed to predict inter-protein contacts and quaternary structures[(Quadir et al., 2021; Xie & Xu, 2021; Yan & Huang, 2021; Evans et al., 2022; Guo et al., 2022; Roy et al., 2022)](https://www.zotero.org/google-docs/?whcmv4) . Particularly, adapting AlphaFold2 for protein quaternary structure prediction (i.e., AlphaFold-multimer (Evans et al., 2022)) substantially improved the accuracy of protein complex/assembly structure prediction.

In parallel to the development of the methods for sampling protein quaternary structures, numerous methods have been developed for estimating the accuracy of predicted quaternary structural models (**EMA**) in order to select the best predicted structures and use them effectively in biological research[(Dapkūnas et al., 2021)](https://www.zotero.org/google-docs/?weri3e). The EMA methods can be classified into four categories: (1) single-model energy/statistical potential-based methods; (2) single-model machine learning methods; (3) multi-model consensus methods; and (4) hybrid methods of combining (1), (2) and/or (3). Single-model energy/statistical potential-based methods [(Chen et al., 2003; Lu et al., 2003; Pierce & Weng, 2007, 2008)](https://www.zotero.org/google-docs/?AhFgkr) calculates the physical energy (e.g., electrostatics, hydrophobic interactions, solvation) or statistical potential[(Olechnovič & Venclovas, 2017)](https://www.zotero.org/google-docs/?Q67Jyx) (e.g., residue-residue contact potentials) of a model to estimate its quality. Machine learning methods including deep learning methods [(Wang et al., 2020; Igashov et al., 2021; Wang et al., 2021; Chen et al., 2022; Liu et al., 2023)](https://www.zotero.org/google-docs/?OdP3Ld) use expert-curated or automatically extracted features of a model as input to predict its quality. For instance, DProQA (Chen et al., 2022) uses a gated-graph transformer to take a protein quaternary structure represented as a graph (nodes: residues and edges: residue-residue interactions) as input to predict its quality score. The multi-model consensus methods leverage the similarity between structural models to evaluate their quality, which have been extensively used in assessing the quality of protein tertiary structures [(Zhang & Skolnick, 2004a; Wallner & Elofsson, 2006; Larsson et al., 2009; Cheng et al., 2009; Wang et al., 2011; McGuffin et al., 2021)](https://www.zotero.org/google-docs/?Qv7mk1) and is considered one of the most effective methods for tertiary structure quality assessment[(Cheng et al., 2019)](https://www.zotero.org/google-docs/?bnIwMN). The reason that the model consensus methods work well in many situations is that the good models similar to the native structures must be similar to each other, while bad models can have many ways to be incorrect and are therefore often much less similar to each other, leading to a relatively higher average pairwise similarity score for good models. The approach generally works well when there is a significant or relatively high proportion of good models in a model pool that dominate over the bad models. However, the model consensus approach was rarely used in estimating the accuracy of protein *quaternary* structures until the 2022 CASP15 experiment.

To prepare for the CASP15 experiment, we extended our pairwise model similarity method for *tertiary* structure quality assessment [(Wang et al., 2011)](https://www.zotero.org/google-docs/?o3cBx0) to estimate the quality of protein *quaternary* structural models. However, the pairwise similarity between models cannot work well when many models in a model pool have low quality and are also similar to each other (e.g., structure predictors making the same rather than different folding mistakes), leading to bad models having high pairwise similarity scores (i.e., higher estimated quality scores) to be ranked higher. To address the weakness, we developed a hybrid method (MULTICOM\_qa) combining the pairwise similarity score between a multimer model and other models (PSS) with a single-model-based interface contact probability score (ICPS) that does not depend on the similarity between models to estimate the accuracy of the model. ICPS is calculated from the inter-chain contact map predicted by inter-chain residue-residue contact/distance predictors (Guo et al., 2022; Roy et al., 2022; Xie & Xu, 2022) such as CDPred (Guo et al., 2022). MULTICOM\_qa uses a weighted average of PSS and ICPS to predict the global accuracy (quality) of a quaternary structural model. We first benchmarked MULTICOM\_qa on the CASP14 dataset internally and then blindly tested it in the 2022 CASP15 experiment. It ranked first out of 24 EMA predictors in estimating the global fold accuracy of protein quaternary structural models in CASP15. Our post-CASP15 analysis investigates several important factors influencing MULTICOM\_qa’s performance and identifies its strengths and weaknesses for further improvement.

## **2. Methods**

## To prepare for CASP15, we developed several individual EMA methods to measure the pairwise similarity between multiple quaternary structural models and predict the quality of the contact interfaces of a single structural model, which are described in Section 2.1. We then combined them to design a hybrid method of estimating model accuracy for the CASP15 experiment, which is described in Sections 2.2 and 2.3.

## **2.1 Individual EMA methods**

**Average pairwise similarity score(PSS)**. A complex structure alignment tool - MMAlign [(Mukherjee & Zhang, 2009)](https://www.zotero.org/google-docs/?XzRXc5) - is used to calculate the structural similarity score (i.e., TM-score[(Zhang & Skolnick, 2004b)](https://www.zotero.org/google-docs/?tMAzMD)) between a multimer model and each of the other multimer models in a model pool for a multimer target. The average TM-score between a model and all other models is the average pairwise similarity score (**PSS**) for the model (i.e., *PSSi*, *PSSi* is PSS of model *i*, *j* is the index of any model other than *i*, *TM-scoreij* is the TM-score between model *i* and model *j*, and *n* is the total number of models in the pool). PSS is considered an estimation of the true quality of a model. As a comparison with the PSS, the average similarity between the monomer structural units of one multimer model and their counterparts of each of the other multimer models (called *monomer\_pss*) and the average similarity between the pairs of chains (dimers) of one multimer model and their counterparts of each of the other multimer models (called *dimer\_pss*) are also calculated. monomer\_pss and dimer\_pss are calculated by TM-score [(Zhang & Skolnick, 2004b)](https://www.zotero.org/google-docs/?MNM1uh) and DockQ[(Basu & Wallner, 2016)](https://www.zotero.org/google-docs/?NQjFqT), respectively. Different from monomer\_pss considering only tertiary structure similarity but not inter-chain interactions and dimer\_pss considering both tertiary structure similarity and dimeric interactions, PSS takes into account the overall fold and interface similarity between a multimer model and all the other multimer models and therefore is expected to be more informative about the quality of the model than mnomer\_pss and dimer\_pss. It is worth noting that PSS, monomer\_pss and dimer\_pss are all *multi-model* EMA methods because multiple structural models for the same target are required to compute them.

Diagram

Description automatically generated

**Figure 1: (A)** A simplified illustration of the ICPS calculation. **(i)** A dimer model with chain A in red and chain B in blue. **(ii)** All the inter-chain contacts from a dimeric interface of the model are identified, which are coloured in red and blue. **(iii)** The green lines highlight some inter-chain contacts present in the interface. **(iv)** The predicted probability scores for the inter-chain contacts are extracted from the inter-chain contact map predicted by a deep learning predictor (CDPred). **(v)** The probability scores of the contacts in the interface are averaged as ICPS for the model. **(B)** The pipeline of MULTICOM\_qa for estimating CASP15 assembly model accuracy. The input is a pool of assembly models for a target and other related information such as the stoichiometry of the target, protein sequences, predicted tertiary structures for each unit in the target, and the number of CPU cores. Multiple CPU cores can be used to speed up the calculation of the pairwise similarity scores between assembly models. PSS and ICPS for each model are computed and then averaged as the predicted global fold accuracy score for the model. The ICPS score is also used as the predicted interface score for each model.

**Average interface contact probability score(ICPS)**. The inter-chain residue-residue contacts in a multimer model are first identified. Two residues are considered in contact if their Ca - Ca atom distance is less than 8 Å. The interaction interfaces between two chains that interact in at least 20% of multimers are selected for the ICPS calculation. The inter-chain residue-residue contact probability map for every two chains forming an identified interaction interface are then predicted by an inter-chain residue-residue contact predictor - CDPred([Guo et al., 2022)](https://www.zotero.org/google-docs/?P0fQJ0). CDPred is a deep learning-based tool that can predict the inter-chain distance map for homodimers and heterodimers. It leverages the 2D attention mechanism and deep residual network to capture the interaction between different chains to increase the accuracy of inter-chain distance prediction. The inter-chain distance map predicted by CDPred is converted into an inter-chain contact probability map. The predicted probabilities for the inter-chain residue-residue contacts in each interaction interface in a multimer model are extracted from the inter-chain contact probability maps predicted by CDPred. The average probability of the inter-chain residue-residue contacts in the interaction interfaces is called the average interface contact probability score (**ICPS**), measuring the likelihood of the interaction interfaces in a multimer model. For example, **Figure 1 (A)** illustrates how ICPS is calculated for an interface. ICPS is a *single-model* quality score because it only needs the information of a single model to be computed, which is different from PSS that depends on the comparison between models.

**Number of interface contacts(interface\_size)**. A simple metric - the number of inter-chain contacts in a multimer, which measures the size of the interaction interfaces of a multimer model, was shown to have some discriminative power of ranking multimer models in [(Bryant et al., 2022)](https://www.zotero.org/google-docs/?PJ06TX). In this study, this method serves as a baseline to measure the performance of PSS, monomer\_pss, dimer\_pss and ICPS.

We evaluated the five methods above in terms of average ranking loss on the CASP14 dataset. The CASP14 multimer dataset consists of 21 multimer targets, each of which has about 100 structural models predicted by CASP14 assembly predictors. Each method was used to rank the models for each target. The ranking loss is the true TM-score of the best model for a target minus the true TM-score of the no. 1 model ranked by a method. If a method ranks the truly best model of a target no. 1, the ranking loss for the target is 0. The average per-target ranking loss of each method over all the CASP14 targets was computed (supplementary **Figure S1**).

PSS has the lowest loss of 0.118, which is much lower than the other methods, indicating it is quite effective in ranking CASP14 multimer models. interface\_size has the highest loss of 0.319. The loss of dimer\_pss is 0.147, lower than 0.199 of monomer\_pss and 0.204 of ICPS. dimer\_pss performs better than monomer\_pss because the former considers the interactions between protein chains but the latter does not. It is also interesting to see that the single-model score - ICPS performs similarly to a multi-model score - monomer\_pss and much better than the interface\_size, indicating that ICPS contains valuable information for ranking multimer structural models.

**2.2. Combination of EMA Methods**

After establishing PSS as the best individual model ranking method on the CASP14 dataset, we tried to combine the different methods (multimer\_pss, dimer\_pss, monomer\_pss, and ICPS) to further improve the model ranking performance. In one experiment, we used the average of the four methods ( (PSS + dimer\_pss + monomer\_pss + ICPS) / 4 ) to rank the models of the CASP14 targets. However, the average ranking loss of this combination is 0.156, higher than 0.118 of using PSS alone. The reason for the worse performance is that the dimer\_pss and monomer\_pss are redundant and inferior to PSS and therefore combining them with PSS does not help.

In another experiment, we tested the combination of the multi-model method - PSS and the single-model method - ICPS. We used each of them to rank the models for each CASP14 target. The weighted average of the two ranks for each model (i.e., 0.4 × rank based on ICPS + 0.6 × rank based on PSS) from the two rankings is used as the final rank for the model. The ranking loss of this average ranking approach is 0.105, lower than 0.118 of using only PSS, indicating that ICPS and PSS are complementary and combining them can improve model ranking. Therefore, we chose to combine PSS and ICPS to develop an EMA predictor for the CASP15 experiment. However, the weights of combining them were not fully optimized prior to CASP15 due to the time constraint.

**2.3 MULTICOM\_qa for Estimating Assembly Model Accuracy in CASP15**

Because CASP15 required an estimated global quality score for each model instead of its predicted rank, MULTICOM\_qa used the weighted average of PSS and ICPS (i.e., 0.4 × ICPS + 0.6 × PSS) as the predicted global fold accuracy score for each model. ICPS was also used as the predicted interface score for each model. CDPred was used to predict inter-chain contact maps for the ICPS calculation. The prediction pipeline of MULTICOM\_qa is illustrated in **Figure 1 (B)**.

Given a pool of models for an assembly target, MULTICOM\_qa extracts their backbone structures by filtering out their side chain atoms. The pairwise similarity score (i.e., TM-score) between any two models is calculated by MMalign. To speed up the calculation, multiple CPU cores can be used to compute the scores in parallel. The average pairwise similarity score between one model and all other models (PSS) is computed for each model. The stoichiometry and sequence information are used to find all the acceptable interactions for a target. A dimeric interaction between two chains is considered as an acceptable interaction and is taken into account in the calculation of the ICPS if it is present in at least 20% of the models in the pool. The tertiary structure for each chain of the target predicted by AlphaFold2 is provided for CDPred to predict the inter-chain residue-residue contact map for any two interacting chains, from which the ICPS for the interfaces of each assembly model is computed. The weighted average of ICPS and PSS for each model is used as the predicted global fold accuracy score of the model, while ICPS is used as the predicted interface score.

MULTICOM\_qa was internally benchmarked on the CASP14 dataset with four existing methods: ZRANK[(Pierce & Weng, 2007)](https://www.zotero.org/google-docs/?vQlUj6), ZRANK2[(Pierce & Weng, 2008)](https://www.zotero.org/google-docs/?vQlUj6), GNN\_DOVE[(Wang et al., 2021)](https://www.zotero.org/google-docs/?4EG4eL), and DProQA[(Chen et al., 2022)](https://www.zotero.org/google-docs/?e6ZjXe) (supplementary **Table S1**). MULTICOM\_qa clearly performed better than the other four methods. After being briefly tested, MULTICOM\_qa blindly participated in the CASP15 experiment from May to August 2022.

**3. Results**

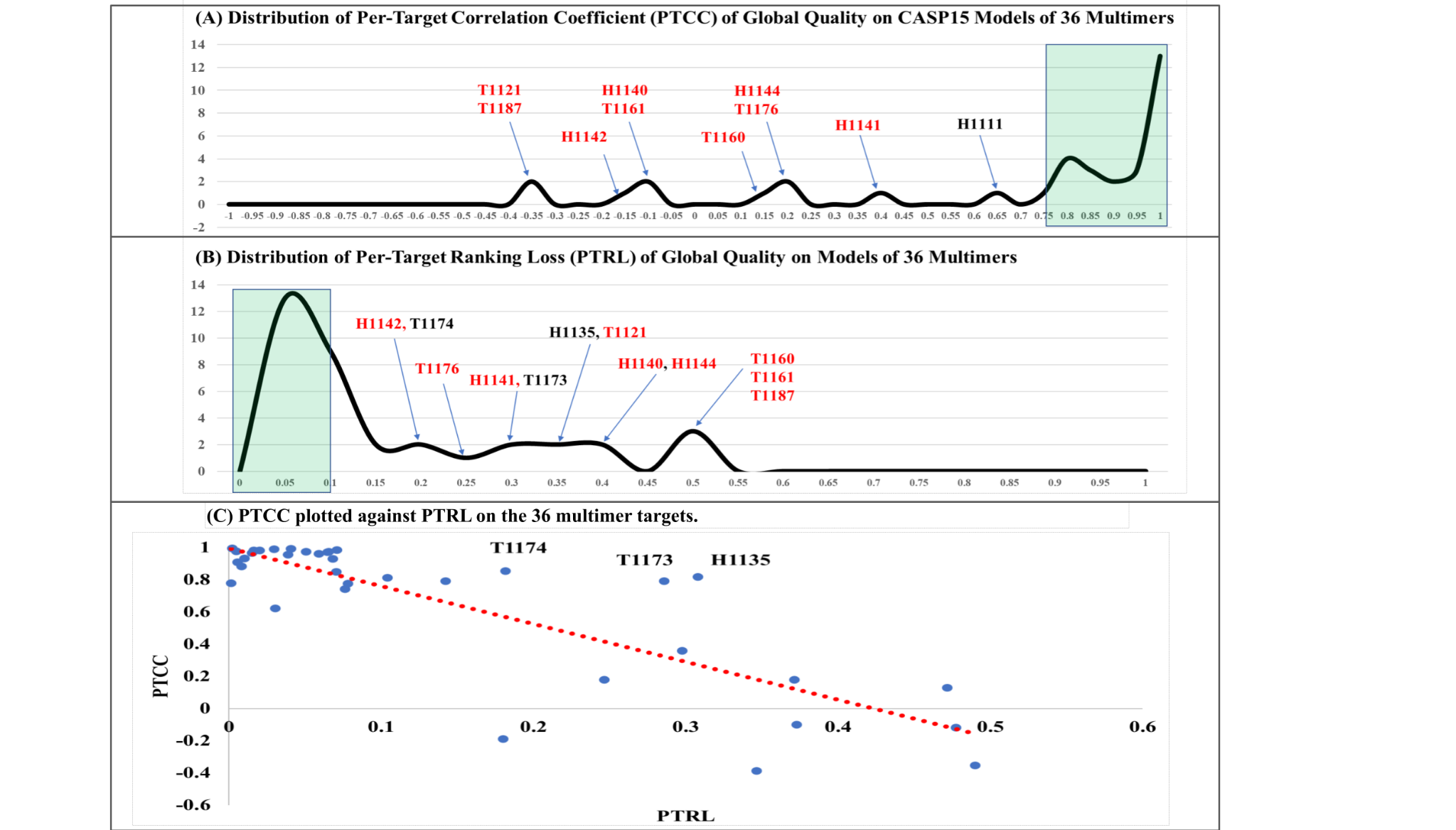
**3.1 Evaluation metrics and CASP15 assembly targets and models**

The CASP15 assessors of the EMA category used a comprehensive evaluation metric (called SCORE) composed of Pearson’s correlation, Spearman’s correlation, ranking loss, and area under ROC curve in terms of both TM-score [(Zhang & Skolnick, 2004b)](https://www.zotero.org/google-docs/?l8pLqz) and GDT-TS score [(Zemla, 2003)](https://www.zotero.org/google-docs/?oR9xtJ) to compare the performance of CASP15 EMA predictors in estimating the global fold accuracy of CASP15 assembly models [(Studer et al., 2022)](https://www.zotero.org/google-docs/?6HuyPs). Based on this metric, MULTICOM\_qa ranks No. 1 in estimating the global accuracy of the CASP15 assembly models (see the CASP15 official ranking in terms of SCORE: <https://predictioncenter.org/casp15/zscores_EMA.cgi> for details).

To complement the CASP15 official assessment, in this study, we mainly use two individual, complementary metrics (i.e., *per-target correlation correlation coefficient* (**PTCC**) and *per-target ranking loss* (**PTRL**)) to evaluate the CASP15 global model accuracy prediction results of MULTICOM\_qa and analyze its strengths and weaknesses. PTCC is the Pearson’s correlation coefficientbetween model quality scores predicted by an EMA method (e.g., MULTICOM\_qa) and true TM-scores (global quality) of the models of a target. Higher PTCC, the better the estimation of the accuracy of the models for the target. PTRL for a target is the difference between TM-score of the truly best model in a model pool and TM-score of the no. 1 model selected by an EMA method for a target. Smaller PTRL, the better the ranking for the target. PTCC and PTRL measures how well an EMA method performs on one target from the two complementary perspectives. PTCC (or PTRL) can be averaged over all the targets to assess the overall performance of an EMA method on a dataset.

We evaluate MULTICOM\_qa’s EMA predictions for the assembly models of all the CASP15 assembly targets except H1171, H1172, and H1185 (i.e., 36 targets in total). H1171 and H1172 with multiple native conformations and H1185 whose native structure was not available to us are excluded. The assembly models for the targets were predicted by CASP15 assembly predictors and were released for CASP15 EMA predictors including MULTICOM\_qa to estimate their accuracy from May to August 2022. Each target has about 275 structural models. The 36 assembly (multimer) targets include 13 homodimers, 10 heterodimers, 6 homomultimers with more than two chains, and 7 heteromultimers with more than two chains. Some targets (e.g., H1111, H1114, T1115, and H1137) are very large. For instance, H1137 has 10 chains with a stoichiometry of A1B1C1D1E1F1G2H1I1. Almost all the models for these targets were generated by the customized AlphaFold-multimer used by the CASP15 assembly predictors, and most targets have some models of good or reasonable quality. However, quite a few targets are very hard and have few good models, posing a significant challenge for EMA.

**3.2 The overall performance of estimating the accuracy of CASP15 assembly models and the factors influencing the performance**

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**Figure 2**. The overall performance of MULTICOM\_qa on the models of 36 CASP15 multimer targets. (**A)** Distribution of the per-target correlation coefficient between MULTICOM\_qa predicted and true global quality scores (PTCC). The green area contains targets with high PTCC > 0.75. Several targets with low/moderate correlation coefficients are labelled. The average PTCC on the 36 targets is 0.6626. (**B)** Distribution of the per-target ranking loss (PTRL). The green area contains targets with low PTRL < 0.1. Several targets with high ranking loss are labelled. The red color highlights nine targets with both high ranking loss and low correlation. The average PTRL on the 36 targets is 0.142. **(C)** PTCC is plotted against PTRL on the 36 multimer targets. The general trend is that a higher PTCC corresponds to a lower PTRL (correlation between them = -0.8358). However, there are three pronounced exceptions (T1174, T1173 and H1135) with both high correlation and high ranking loss.

**Figure 2A** shows the distribution of PTCC between MULTICOM\_qa predicted global quality scores and the true TM-scores of the models for the 36 CASP15 assembly targets. The true TM-score of a model is obtained by using MMalign to compare it with the native structure and is normalized with respect to the length of the native structure. Higher PTCC indicates the more linear consistency between the predicted global accuracy and the true global accuracy of the models of a target. 26 targets have a relatively high PTCC >= 0.7, indicating MULTICOM\_qa can estimate the relative global quality of the models of most targets reasonably well. The average PTCC on the 36 CASP15 targets is 0.6626. Two targets (H1141 and H1111) have a moderate correlation between 0.35 and 0.65, while another three targets (T1160, H1144 and T1176) have a low positive correlation between 0 and 0.2. MULTICOM\_qa completely failed on five targets (T1121, T1187, H1142, H1140 and T1161) as the PTCCs for them are negative.

**Figure 2B** illustrates the distribution of PTRL for the models of 36 CASP15 multimers. The average PTRL is 0.142. Most targets have a low or acceptable ranking loss less than ~0.15, indicating that MULTICOM\_qa can select a reasonable model for most targets as no. 1 model. However, 12 targets (H1142, T1174, T1176, H1141, T1173, H1135, T1121, H1140, H1144, T1160, T1161 and T1187) have a high ranking loss (PTRL > ~0.15), indicating MULTICOM\_qa failed to rank good or acceptable models of these targets at the top. Interestingly, 9 out of the 12 targets (i.e., H1142, T1176, H1141, T1121, H1140, H1144, T1160, T1161 and T1187) also have a low or negative correlation coefficient (PTCC < 0.4), indicating poorly predicted quality scores often lead to a high ranking loss. Indeed, there is a highly negative correlation of -0.84 between PTRL and PTCC (**Figure 2C**). Most of the targets that have high PTCC (>0.7) have a relatively low ranking loss (PTRL < 0.15) as expected. But there are three pronounced exceptions (T1174, T1173 and H1135) that have high PTCC and high PTRL. Supplementary **Figure S2** plots MULTICOM\_qa predicted quality scores against the true quality scores of the models of the three targets, respectively. Interestingly, even though there is an overall strong positive correlation between the predicted quality scores and true quality scores across the entire quality score range, their correlation for good models (true TM-score > 0.6 or 0.7) are negative, indicating that MULTICOM\_qa was not able to select better models among the good models and instead it chose some mediocre ones from them as no. 1. There are no targets that have both low PTCC (< 0.6) and low PTRL (< 0.1).

To analyze what factors influence the PTCC of MULTICOM\_qa, we plot PTCC against the average true TM-score of the models for each target (i.e., a measure of the absolute difficulty of a target; higher, less difficult), the proportion of good models (i.e., a measure of the sampling difficulty of a target; higher, less difficult to sample a good model by a method such as AlphaFold-multimer), and the skewness of the distribution of the quality (true TM-scores) of the models for each target (supplementary **Figure S3A-C**). The skewness is a measure of the asymmetry of model quality distribution, which is equal to , where *n* is the number of the models for a target, is the TM-score of model *i*, is the mean TM-score and is the standard deviation. A large positive skewness indicates that there are more highly concentrated below-average models within a narrow range and there is a wide spread (a long tail) of low-frequency above-average models (supplementary **Figure S4A**). A small negative skewness indicates that there are more highly concentrated above-average models within a narrow range and there is a wide spread (a long tail) of low-frequency below-average models (supplementary **Figure S4B**). A zero skewness means the above-average and below-average models are symmetrically distributed.

The Pearson’s correlation between PTCC and each of the three factors (average TM-score, proportion of good models, skewness) is 0.5852, 0.615, and -0.7314, respectively (supplementary **Figure S3A-C**). A moderate positive correlation between PTCC and the average TM-score indicates that the absolute difficulty of a target plays a moderate role in estimating the accuracy of the models of the target. A slightly stronger positive correlation between PTCC and the proportion of good models suggests that sampling more good models can make the quality estimation easier, which is expected because more good models lead to higher pairwise similarity scores between good models and likely higher estimated quality scores for them than bad models. A very strong negative correlation between PTCC and skewness indicates that the asymmetry distribution of the quality of the models plays an important role in MULTICOM\_qa’s estimation of the model accuracy. A high concentration of below-average-quality models in a small range in conjunction with a low frequency of above-average-quality models spread in a wide range of quality (high positive skewness) makes the estimation of model accuracy harder. On the opposite, a low negative skewness makes the estimation of model accuracy easier. Overall, the skewness influences the model accuracy estimation more than the absolute difficulty of a target (i.e., average TM-score) and the sampling difficulty (i.e., the proportion of good models).

Supplementary **Figure S3D-F** plots PTRL against three factors (average TM-score, proportion of good models, and skewness). Like PTCC,the similar relationship has been observed that PTRL has a moderate (negative) correlation with the average TM-score measuring absolute target difficulty, stronger (negative) correlation with the proportion of good models measuring the sampling difficulty, and very strong positive correlation with the skewness of the distribution of model quality. For instance, three homomultimers (T1160, T1161 and T1187) with the highest ranking loss of 0.47203, 0.47772 and 0.49026 also have the high skewness of 8.114, 2.6317 and 2.3826.

Supplemental **Figure S5** compares MULTICOM\_qa with other 22 CASP15 EMA predictors (excluding APOLLO since it submitted the same global quality scores for all the models for most targets) in terms of the average PTCC and the average PTRL on 36 CASP15 assembly targets. MULTICOM\_qa has the highest average PTCC of 0.6626, which is 17.67% higher than the second highest PTCC of 0.5631 of ModFOLDdock. MULTICOM\_qa has the third lowest average PTRL of 0.142, higher than the lowest average PTRL of 0.1067 of Venclovas and second lowest average PTRL of 0.1383 of VoroIF.

**3.3 Successful and failed CASP15 cases of MULTCOM\_qa**

**Figure 3A-D** illustrates the four typical good cases in which MULTICOM\_qa performed well. **Figure 3A** illustrates a case (a 27-chain heteromultimer H1111 with a stoichiometry of A9B9C9) in which structural models largely fall into two groups: a good group of models with TM-score higher than 0.90 and a bad group of models with TM-score lower than 0.25. Because there is a significant portion of good models and they are highly similar to each other, the average pairwise similarity for good models is relatively high and thus they can be readily picked up by MULTICOM\_qa. Indeed, MULTICOM\_qa selected one model from the bin with TM-scores in (0.95, 1]. The model has a TM-score of 0.95196, slightly lower than 0.9825 of the best model in the pool (the ranking loss - PTRL = 0.03054). The correlation - PTCC between the predicted quality scores and true TM-scores is 0.62, which is only moderate.

**Figure 3B** is a homodimer (T1123) that has a large portion of good models with TM-score between 0.85 and 0.9, several high-quality models with TM-score between 0.9 and 0.95, and many other models with TM-score widely spanning from ~0.15 to 0.85. MULTICOM\_qa was able to select a model from the bin (red) with the highest score range (0.9, 0.95). The top-1 selected model has a TM-score of 0.9063. The PTRL is very low (0.01671). The PTCC is very high (0.98), indicating MULTICOM\_qa, mostly its PSS component, is good at estimating the accuracy of the models with this kind of skewed quality distribution.

**Figure 3C** is H1137, a very large heteromultimer target (stoichiometry: A1B1C1D1E1F1G2H1I1). The true TM-score of its models have a wide range of distribution from ~0.05 to 0.95. However, the mode of the distribution lies in the medium quality score range (0.65, 0.7]. MULTICOM\_qa was able to select a model from the bin with the second highest score range (0.85, 0.9]. The model has a TM-score of 0.87644, which is not the best but a reasonable choice. The PTRL is 0.05916, which is reasonably low considering the size and difficulty of the target. The PTCC is 0.96, indicating that MULTICOM\_qa’s predictions can have high correlation with true quality scores when the true quality scores of the models are somewhat evenly distributed in a large range.

**A picture containing graphical user interface

Description automatically generated**

**Figure 3**. Good and bad examples of applying MULTICOM\_qa to estimate the accuracy of the models of CASP15 multimer targets. In each example, the distribution of the true TM-scores of the models for each target is visualized as histogram. The native structure, true top-1 (best) model, selected top-1 model, and the TM-scores of the latter two are presented. The bin in the histogram from which the top-1 model was selected by MULTICOM\_qa is highlighted in red. (**A)** H1111: stoichiometry = A9B9C9, PTCC = 0.62, PTRL = 0.03054; (**B**) T1123: stoichiometry = A2, PTCC = 0.98, PTRL = 0.01671; (**C**) H1137: stoichiometry = A1B1C1D1E1F1G2H1I1, PTCC = 0.96, PTRL = 0.05916; (**D**) T1179, stoichiometry = A2, PTCC = 0.97, PTRL = 0.06577; (**E**) T1121: stoichiometry = A2, PTCC = -0.39, PTRL = 0.34683; the correct interfaces are circled; and the top-1 model selected by MULTICOM\_qa contains only one of the two correct interfaces, resulting in a high ranking loss of 0.34683.

**Figure 3D** is a case in which combining ICPS with PSS improves the ranking of the models. T1179 is a homodimer whose models have TM-scores largely falling into the range (0.75, 0.9], with some better models on its right side and many worse models on its left side. The pairwise similarity score (PSS) chose a good model with TM-score of 0.85378, but MULTICOM\_qa combining PSS and ICPS selected a better model with TM-score of 0.90331. This case indicates that the single-model quality score - ICPS in MULTICOM\_qa can sometimes improve the selection of the top-1 model by moving it to the higher quality bin.

**Figure 3E** is a typical case where MULTICOM\_qa failed miserably. The quality scores of the models of this homodimer target (T1121) are distributed between ~0.45 and ~0.85. However, the scores of the majority of the models fall into a narrow low score bin (0.50, 0.55], even though there are some better models whose scores are largely evenly distributed from ~0.55 to ~0.85. The distribution of the quality of the models is highly skewed. And because there are too many bad models that are quite similar to each other (average TM-score for the bad models in the (0.5 - 0.55] bin is 0.86), they dominate the calculation of PSS, leading to bad models having higher estimated quality scores. Consequently, MULTICOM\_qa chose a bad model from the bin (0.5, 0.55]. The ranking loss is very high (PTRL = 0.34683). Moreover, the PTCC is negative (-0.39), indicating that PSS of MUTLCOM\_qa completely failed in this situation and ICPS could not rescue it. This is a typical case in which MULTICOM\_qa fails when there is only a small portion of good models and the bad models are similar to each other. In this case, MULTICOM\_qa selected a model with one interaction interface that matches with one of the two interaction interfaces of the native structure, while the best model in the pool has the two correct interfaces.

Supplementary **Figure S6** shows 8 additional good cases where MULTCOM\_qa performs well, each of which has a significant portion of good models making ranking easy. Supplementary **Figure S7** shows 6 additional bad cases where MULTICOM\_qa failed, each of which has a small portion of good models.

Scatter chart

Description automatically generated with medium confidence

**Figure 4. (A)** The model similarity graph for H1111. **(B)** The model similarity graph for T1121. In a model similarity graph, each node denotes a model. An edge is used to connect two models (nodes) if their structural similarity score - TM-score is greater than a threshold. The threshold is determined such that only the top 30% of model pairs are connected by edges. The weight of each edge is TM-score between two nodes (models) calculated by MMalign, which is normalized by the total sequence length of the larger model if the two models have different sizes. The color of the nodes corresponds to the true TM-scores of the models. For H1111, both the best model (true top-1 model) and selected top-1 model come from the largest subgraph with the highest quality. For T1121, the selected top-1 model is from the largest subgraph with mediocre quality, but the best model is in the third largest subgraph.  **(C)** The distribution of TM-scores of the models of H1114 as well as its native structure, top-1 model selected by PSS, top-1 model selected by ICPS, top-1 model selected by MULTICOM\_qa, and the best model in the model pool. Both ICPS and MULTICOM\_qa selected a model that is much better than the model selected by PSS. The four A chains in the good models form a cube in the center (red), which is a key feature of the structure of H1114. **(D)** A homo-dimeric interface between two A chains of the top-1 model selected by ICPS for target H1114. The model is H1114TS119\_2. The red lines represent the true inter-chain contacts in the interface. The numbers are the probabilities for the true contacts predicted by the CDPred. The true contacts have high predicted probabilities, leading to a high ICPS of 0.44 for the interface.

To further analyze how MULTICOM\_qa may succeed or fail, we draw a *model similarity graph* for a good case H1111 (**Figure 4A**) and a bad case T1121 (**Figure 4B**). In each graph, a node denotes a model and an edge connects two models if their structural similarity is higher than a threshold. Here, a threshold is chosen so that 30% of model pairs are connected by edges. The model similarity graph of H1111 has 7 subgraphs (clusters), among which the largest subgraph has higher true quality scores and high within-subgraph similarity. Both the best model and the top-1 model selected by MULTICOM\_qa are from the largest subgraph that contains a lot of good and similar models. In contrast, the model similarity graph of T1121 has five subgraphs (clusters). The best model resides in the third large subgraph, while MULTICOM\_qa selected one model from the largest subgraph containing a lot of bad models that are similar to each other. This example clearly demonstrates that the high similarity between bad / mediocre models in a large cluster makes model ranking difficult for MULTICOM\_qa, mostly its PSS component.

**3.4 Quantification of the difficulty of structural models for model accuracy estimation**

The results in Section 3.2 show that the absolute difficulty of a target (average true TM-score of the models of the target), the sampling difficulty (the proportion of good models), and the skewness of the distribution of quality of the models affect the EMA performance (PTCC and PTRL) of MULTICOM\_qa moderately or strongly. We also calculate the correlation between PTCC and the mean pairwise similarity between all the models in a model pool of a target (i.e., equal to the average of PSS of the models in the pool) for the CASP15 targets (**Table 1**). The correlation between MULTICOM\_qa’s PTCC and the mean pairwise similarity score of all the models of a target is only 0.1220, which is much weaker than the correlation between PTCC and the other three factors (0.5852 for average TM-score; 0.6150 for the proportion of good models, and -0.7314 for the skewness), indicating that the mean pairwise similarity between all the models in a model pool is not a good indicator of the difficulty of estimating the accuracy of the models. The reason is that, even though high similarity between good models makes the model accuracy estimation easier, high similarity between bad models makes it harder. To treat the negative and positive impact of the similarity between good/bad models differently, we design a new metric called *Difficulty Index* to quantify the difficulty of estimating the accuracy of the models for a target.

**Table 1**. Pearson's correlation between PTCC (or PTRL) of MULTICOM\_qa and each of the five factors (average TM-score, proportion of good models, skewness, mean pairwise similarity score between all the models, and difficulty index). Bold numbers denote the strongest correlation.

|  |  |  |  |  |  |
| --- | --- | --- | --- | --- | --- |
| Pearson’s Correlation | Avg true TM-score of models | Proportion of good models (TM-score > 0.8) | Skewness | Mean pairwise similarity score between all models | Difficulty Index |
| PTCC of MULTICOM\_qa | 0.5852 | 0.6150 | -0.7314 | 0.1220 | **0.7723** |
| PTRL of MULTICOM\_qa | -0.6661 | -0.7259 | 0.7784 | -0.1478 | **-0.8360** |

The difficulty index for the models of a target is defined as . PSSi is the average pairwise similarity score of model *i*, *TM-score*i is the true quality score of model *i*, *n* is the number of models. denotes the middle value between the minimum and maximum TM-scores of the models. The difficulty index ranges from -1 to 1. Lower the difficulty index, more difficult the models are for accuracy estimation. In this definition, the PSS of the below-average-quality (bad) models whose TM-score is less than the middle value (*TM-score*i < *TM-scoremiddle*) contributes negatively to the difficulty index, while the above-average-quality (good) models (*TM-score*i > *TM-scoremiddle*) contribute positively. The correlation between the difficulty index and PTCC (or PTRL) of MULTICOM\_qa is 0.7723 (or -0.8360), which is even stronger than the correlation between the skewness and PTCC (or PTRL) and several times stronger than the mean pairwise similarity score between all the models (**Table 1**). The results show that the difficulty index considering the different impacts of the similarity between bad/good models is a very effective measure of the difficulty of estimating the accuracy of the models in a model pool, which is more effective than only considering the skewness of the distribution of the model quality. The correlation for the difficulty index is much higher than for the mean pairwise model similarity confirms the importance of treating the pairwise similarity between good or bad models differently for EMA. The difficulty index can be used to select hard targets for testing EMA methods to improve their performance on them.

Moreover, there is a strong correlation (0.8078) between the difficulty index and the average true TM-score of the models (absolute difficulty) of a target (supplementary **Figure S8**). This indicates that the difficulty index of the models for a target can also quantify the difficulty of predicting structures for the target.

**3.5 Analysis of the properties and contributions of PSS and ICPS**

We compare how the EMA performance (PTCC and PTRL) of the two components of MULTICOM\_qa (i.e., PSS and ICPS) is influenced by the four factors (**Table 2**): (1) the average true TM-score of the models of a target, (2) the proportion of good models, (3) the skewness of the distribution of the quality of models, and (4) the difficulty index of the models. The PTCC of PSS has the strongest correlation with the four factors (**Table 2A**), the PTCC of MULTICOM\_qa has the second strongest correlation with them (see **Table 1**), and the PTCC of ICPS has a much weaker correlation with them than PSS and MULTICOM\_qa (**Table 2A**). This indicates that the performance of the single-model method - ICPS is much less influenced by the four factors than the multi-model method - PSS. The stronger correlation between MULTICOM\_qa with the four factors mostly comes from PSS as they have similar correlation coefficients and trends. The largely similar results are observed in terms of PTRL (see the PTRL’s correlation for PSS and ICPS in **Table 2B** andfor MULTICOM\_qa in **Table 1**). Finally, PTCC/PTRL of PSS has the strongest correlation with the difficulty index, but PTCC/PTRL of ICPS has the strongest correlation with the skewness.

**Table 2.** The correlation between the performance (PTCC or PTRL) of two components (PSS and ICPS) of MULTICOM\_qa and the four factors (average true TM-scores of the models of a target, the proportion of good models, the skewness of the quality of the models, and the difficulty index of the models). **(A)** The correlation between PTCC and the four factors; and **(B)** the correlation between PTRL and the four factors.

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **A. Correlation of PTCC** | Average TM-score of models | Proportion of good models (TM-score > 0.8) | Skewness | Difficulty index |
| PTCC of PSS | 0.6694 | 0.6514 | -0.7467 | 0.7838 |
| PTCC of ICPS | 0.2207 | 0.3163 | -0.3669 | 0.3101 |

|  |  |  |  |  |
| --- | --- | --- | --- | --- |
| **B. Correlation of PTRL** | Average TM-score of models | Proportion of good models (TM-score > 0.8) | Skewness | Difficulty index |
| PTRL of PSS | -0.6900 | -0.7360 | 0.7740 | -0.8351 |
| PTRL of ICPS | -0.2716 | -0.2583 | 0.3190 | -0.2790 |

We also investigate how the two components (PSS and ICPS) of MULTICOM\_qa contribute to its EMA performance on the 36 CASP15 multimer targets. Because MULTICOM\_qa was not able to generate PSS for a very large multimer H1114 ((stoichiometry: A4B8C8)) within the two-day limit, it used ICPS scores alone as the final predicted quality scores for the models of H1114 during the CASP15 experiment. Here, to fairly compare the performance of PSS and ICPS of MULTICOM\_qa, we use MMalign (version 2021/8/16) to regenerate PSS for the models of H1114 and calculate the weighted average of the PSS and ICPS as the final predicted scores.

**Table 3**. The PTCC and PTRL of PSS, ICPS and MULTICOM\_qa (correction) on 36 CASP15 multimer targets. MULITICOM\_qa (correction) differs from the CASP15 MULTICOM\_qa only on one target - H1114. Bold number denotes the best PTCC or PTRL.

|  |  |  |
| --- | --- | --- |
| Method | Average PTCC↑ | Average PTRL↓ |
| PSS | **0.6737** | 0.1454 |
| ICPS | 0.3139 | 0.2938 |
| MULTICOM\_qa (correction) | 0.6645 | **0.1413** |

**Table 3** reports the average PTCC and the average PTRL of the three methods on the 36 CASP15 multimer targets. MULTICOM\_qa has the lowest average PTRL, while PSS has the highest average PTCC, indicating that combining PSS and ICPS in MULTICOM\_qa reduces the average per-target ranking loss, but slightly decreases the average per-target correlation coefficient. Both MULTICOM\_qa and PSS perform much better than ICPS, suggesting that the multi-model method PSS still works much better than the single-model based interface contact score - ICPS on average. However, PSS and ICPS are complementary. H1114 (stoichiometry: A4B8C8) is a notable example showing that combining ICPS and PSS reduces the ranking loss. **Figure 4C** illustrates how ICPS and MULTICOM\_qa selected a model for H1114 that is much better than PSS. The reason is that CDPred predicted high probabilities for some true contacts in an interface of some good models, leading to a higher ICPS for them to be selected (**Figure 4D**).

**3.6 Optimization of the weights of ICPS and PSS in MULTICOM\_qa**

When we developed MULTICOM\_qa for the CASP15 experiment, the weights for its two components (0.4 for ICPS and 0.6 for PSS) were not optimized due to the time constraint. After the CASP15 experiment, we tested different weights for them on the 36 multimer targets to see if better results can be obtained. As in Section 3.5, the incorrect PSS of the models of H1114 was corrected for this analysis. **Figure S9** shows how the average per-target ranking loss (PTRL) of MULTICOM\_qa changes with respect to the weight of ICPS. The lowest average PTRL of 0.1381 is achieved at the ICPS weight of 0.48, which is better than the loss of 0.1413 for the weight 0.4 used in the CASP15 experiment and the loss of 0.1454 of using PSS alone (i.e., ICPS weight = 0). The results show that the performance of combining PSS and ICPS may be further improved if their weights are optimized.

**4**. **Discussion**

Calculating the similarity between two multimers depends on multimer structure alignment tools such as MMalign. The existing multimer alignment tools require a good (or optimal) mapping between the chains of one multimer and those of another multimer to accurately calculate their similarity. However, finding an optimal mapping for two large homomultimers or large heteromultimers containing multiple copies of the same chain is extremely hard and time consuming. Therefore, even though the empirical approach adopted by MMalign can calculate multimer similarity well in most cases, sometimes it severely underestimates the similarity between the models of some large multimers consisting of many chains or yields different similarity scores when the order of the two multimer models is changed, which negatively affects the performance of MULTICOM\_qa. Moreover, the speed of calculating similarity between large multimer models consisting of dozens of chains is slow, which prevents the pairwise similarity calculation from being applied to a large number of models. However, it appears that a new, efficient algorithm of finding the optimal chain mapping between large multimers was designed by CASP15 EMA assessors during the CASP15 meeting[(Studer et al., 2022)](https://www.zotero.org/google-docs/?tRNlpV), which is expected to alleviate the problem. Given an optimal chain mapping as input, some complex alignment tools such as DockQ that requires a chain mapping as input, can be more readily applied to aligning mutimers consisting of more than two chains to calculate their similarity.

As demonstrated in our CASP15 experiment, the multi-model pairwise similarity approach (PSS) works well when the quality scores of the models for a target are either shifted toward the high score range or evenly distributed across the score range. On average PSS still works much better than single-model approach (ICPS) of estimating the quality of quaternary structural model. This is not surprising because the multi-model pairwise similarity measures the frequency of models sampled by the underlying prediction methods. If the underlying prediction methods do a reasonable job, it is expected to produce good models with high probability. Because good models must be similar to the same native/true structure, they must also be similar to each other, leading to a higher average pairwise similarity score for them. Another implicit assumption for the pairwise similarity method to work well is that bad models have different ways to be wrong and therefore are likely different to each other. This is true if the underlying prediction methods do not make the same mistake frequently. However, this assumption can be violated, reflected in that the pairwise similarity approach often fails when there is a large portion of bad models that are similar to each other, which dominates the average pairwise similarity calculation so that bad models are assigned higher scores than good models. This happened for some hard targets in CASP15 when the underlying structure prediction methods generate similar bad models for them (**Figure 4B**). This is a key issue to be addressed to further advance the pairwise similarity approach. The difficulty of structural models for such hard targets can be measured well by the difficulty index introduced in this work.

One way to correct the weakness of the pairwise similarity approach is to combine it with the complementary single-model EMA methods such as the deep learning predicted interface contact probability score (ICPS) that do not depend on the comparison between models. This direction may be important for advancing the state of the art of estimating model accuracy because it is unlikely single-model EMA methods alone can catch up with the performance of the pairwise model similarity approach in a short period of time. However, although this direction is promising, the combination of the two is not trivial. A simple weighted average of the scores of the two can correct some mistakes but may not significantly improve the model accuracy estimation. Therefore, more sophisticated approaches of integrating the strengths of the two complementary approaches need to be developed. For instance, a deep learning method can be trained to combine the pairwise similarity scores with multiple single-model EMA methods to estimate model accuracy. Moreover, for AlphaFold2 predicted multimer models, the AlphaFold2 predicted TM-score or plDDT score can potentially be combined with the pairwise similarity approach if they are available. We have noticed that the model ranking of AlphaFold2 predicted quality score and the pairwise model similarity score are complementary on our in-house models generated for CASP15 targets (data not shown). However, the search for the best approach of combining them is still challenging and ongoing. Moreover, to further advance the combination, more sophisticated and accurate deep learning single-model EMA methods for quaternary structure models need to be developed as what happened in estimating the accuracy of tertiary structure models[(Chen et al., 2023)](https://www.zotero.org/google-docs/?HWpPMG).

## **5. Conclusion**

In this work, we developed a hybrid method of combining the pairwise model similarity score and the interface residue-residue contact prediction to estimate the accuracy of protein assembly models and blindly tested it in the CASP15 experiment. The method is effective in ranking protein assembly models and predicting their global quality scores as demonstrated by its outstanding performance in the CASP15 experiment. Our experiment demonstrates that the average pairwise similarity score (PSS) can estimate the accuracy of models well in most cases, but it fails when a hard target has a large portion of bad models that are similar to each other. The single-model based interface contact probability score (ICPS) derived from the deep learning-based inter-chain contact prediction can provide some valuable information complementary with PSS to evaluate the quality of the inter-chain interaction interface in multimer models. A simple weighted combination of PSS and ICPS can correct some model ranking errors of PSS but does not systematically improve its ranking, suggesting that a more sophisticated (e.g., deep learning-based) integration of the pairwise similarity scores and single-model quality scores is needed to significantly improve the estimation of quaternary model accuracy. More accurate single-model EMA methods are also needed to improve the combination. Finally, it is useful to consider several important factors such as the target difficulty, model sampling difficulty, skewness of model quality, and model difficulty index for designing and evaluating EMA methods.

**Data Availability**

The source code of MULTICOM\_qa and the data are available at <https://github.com/BioinfoMachineLearning/MULTICOM_qa> .

**Conflict of Interest**

### The authors declare that there is no conflict of interest.

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**Authors’ Contributions**

JC conceived and designed MULTICOM\_qa. RR implemented the method and tested it on the CASP14 dataset. ZG adapted CDPred for the ICPS calculation. RR and NG ran MULTICOM\_qa in the CASP15 experiment and collected the CASP15 data. JC, JL and RR designed data analysis methods. JL, JC and RR analyzed the CASP15 results. JC, RR, and JL wrote the manuscript.

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