

Supplementary Materials

Table S1. 19 kinds of combinations of MSA_{paired} and structural templates for homo-multimer targets.

MSA and Template Combinations	MSA_{paired}		Structural templates
	Sequence database	Interaction source	Template database
<i>default_multimer</i>	UniRef30, BFD, MGnify clusters	-	pdb70
<i>default_pdb</i>	UniRef30, BFD, MGnify clusters	-	pdb_sort90
<i>default_pdb70</i>	UniRef30, BFD, MGnify clusters	-	pdb70
<i>default_comp</i>	UniRef30, BFD, MGnify clusters	-	pdb_complex
<i>default_struct</i>	UniRef30, BFD, MGnify clusters	-	pdb_complex
<i>default_af</i>	UniRef30, BFD, MGnify clusters	-	Predicted chain structures

<i>default_img</i>	UniRef90, Integrated Microbial Genomes (IMG), metagenome sequence databases	-	pdb70
<i>uniclust_oxmatch_a3m</i>	UniClust30	Species annotation	pdb70
<i>spec_iter_uniref_a3m</i>	UniRef30	Species annotation	pdb70
<i>spec_iter_uniref_sto</i>	UniRef90	Species annotation	pdb70
<i>spec_iter_uniprot_sto</i>	UniProt	Species annotation	pdb70
<i>spec_pdb</i>	UniRef30	Species annotation	pdb_sort90
<i>spec_pdb70</i>	UniRef30	Species annotation	pdb70
<i>spec_comp</i>	UniRef30	Species annotation	pdb_complex
<i>spec_struct</i>	UniRef30	Species annotation	pdb_complex
<i>spec_af</i>	UniRef30	Species annotation	Predicted chain structures
<i>pdb_iter_uniref_a3m</i>	UniRef30	PDB	pdb70
<i>pdb_iter_uniref_sto</i>	UniRef90	PDB	pdb70
<i>pdb_iter_uniprot_sto</i>	UniProt	PDB	pdb70

Table S2. 29 kinds of combinations of MSA_{paired} , MSA_{unpaired} and structural templates for hetero-multimer targets.

MSA & Template Combination	MSA_{paired}		MSA_{unpaired}	Structural template	
	Sequence database	Interaction source	Sequence database	Template database	Concatenation
<i>default_multimer</i>	UniProt	Species annotation	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>default_pdb</i>	UniProt	Species annotation	UniRef30, BFD, MGnify clusters	pdb_sort90	PDB code
<i>default_pdb70</i>	UniProt	Species annotation	UniRef30, BFD, MGnify clusters	pdb70	PDB code
<i>default_comp</i>	UniProt	Species annotation	UniRef30, BFD, MGnify clusters	pdb_complex	PDB code
<i>default_struct</i>	UniProt	Species annotation	UniRef30, BFD, MGnify clusters	pdb_complex	PDB code

<i>default_af</i>	UniProt	Species annotation	UniRef30, BFD, MGnify clusters	Predicted chain structures	-
<i>uniclust_oxmatch_a3m</i>	UniClust30	Species annotation	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>spec_iter_uniref_a3m</i>	UniRef30	Species annotation	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>spec_iter_uniref_sto</i>	UniRef90	Species annotation	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>spec_iter_uniprot_sto</i>	UniProt	Species annotation	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>spec_pdb</i>	UniRef30	Species annotation	UniRef30, BFD, MGnify clusters	pdb_sort90	PDB code
<i>spec_pdb70</i>	UniRef30	Species annotation	UniRef30, BFD, MGnify clusters	pdb70	PDB code
<i>spec_comp</i>	UniRef30	Species annotation	UniRef30, BFD, MGnify clusters	pdb_complex	PDB code
<i>spec_struct</i>	UniRef30	Species annotation	UniRef30, BFD, MGnify clusters	pdb_complex	PDB code
<i>spec_af</i>	UniRef30	Species annotation	UniRef30, BFD, MGnify clusters	Predicted chain structures	-
<i>unidist_uniref_a3m</i>	UniRef30	UniProt accession ID	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>unidist_uniref_sto</i>	UniRef90	UniProt accession ID	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>unidist_uniprot_sto</i>	UniProt	UniProt accession ID	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>str_iter_uniref_a3m</i>	UniRef30	STRING database	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>str_iter_uniref_sto</i>	UniRef90	STRING database	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>str_iter_uniprot_sto</i>	UniProt	STRING database	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>str_pdb</i>	UniRef90	STRING database	UniRef30, BFD, MGnify clusters	pdb_sort90	PDB code
<i>str_pdb70</i>	UniRef90	STRING database	UniRef30, BFD, MGnify clusters	pdb70	PDB code

<i>str_comp</i>	UniRef90	STRING database	UniRef30, BFD, MGnify clusters	pdb_complex	PDB code
<i>str_struct</i>	UniRef90	STRING database	UniRef30, BFD, MGnify clusters	pdb_complex	PDB code
<i>str_af</i>	UniRef90	STRING database	UniRef30, BFD, MGnify clusters	Predicted chain structures	PDB code
<i>pdb_iter_uniref_a3m</i>	UniRef30	PDB	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>pdb_iter_uniref_sto</i>	UniRef90	PDB	UniRef30, BFD, MGnify clusters	pdb_seqres	-
<i>pdb_iter_uniprot_sto</i>	UniProt	PDB	UniRef30, BFD, MGnify clusters	pdb_seqres	-

Table S3. The average TM-score of the best of five models of the top 15 out of 26 server predictors including NBIS-AF2-multimer (the standard AlphaFold-Multimer predictor) on the 41 multimers, 14 TBM multimers, 27 TBM/FM and FM multimers. When calculating the average TM-score, if a predictor did not submit a prediction for a target, the TM-score is set to 0. The bold font highlights the best result. The underline denotes the second best result.

Server predictor	Sum of Z-scores (> 0.0)	Avg TM-score on 41 multimers	Avg TM-score on 14 TBM multimers	Avg TM-score on 27 FM and FM/TBM multimers	Target count
Yang-Multimer	28.4019	0.7542	0.8619	0.6984	39
MULTICOM_deep	<u>24.6154</u>	0.7781	0.8536	0.7389	41
MULTICOM_qa	23.1823	<u>0.7963</u>	<u>0.8541</u>	<u>0.7663</u>	41
Manifold-E	22.8345	0.8161	0.8504	0.7984	41
DFolding-server	20.2187	0.6816	0.7835	0.6287	36
ColabFold	18.3899	0.6715	0.7584	0.6265	39
UltraFold_Server	18.2015	0.7641	0.7999	0.7456	41
MultiFOLD	17.4288	0.6900	0.7671	0.6500	41
Kiharalab_Server	16.561	0.6995	0.7839	0.6557	40
MUFold	16.4772	0.7478	0.8481	0.6958	41
NBIS-AF2-multimer	14.8905	0.7375	0.8438	0.6824	41
RaptorX-Multimer	14.7853	0.7061	0.8168	0.6487	40
DFolding-refine	12.8844	0.6973	0.8362	0.6253	36
GuijunLab-DeepDA	12.6262	0.7361	0.8309	0.6870	41
Yang-Server	12.2828	0.3839	0.6557	0.2430	19

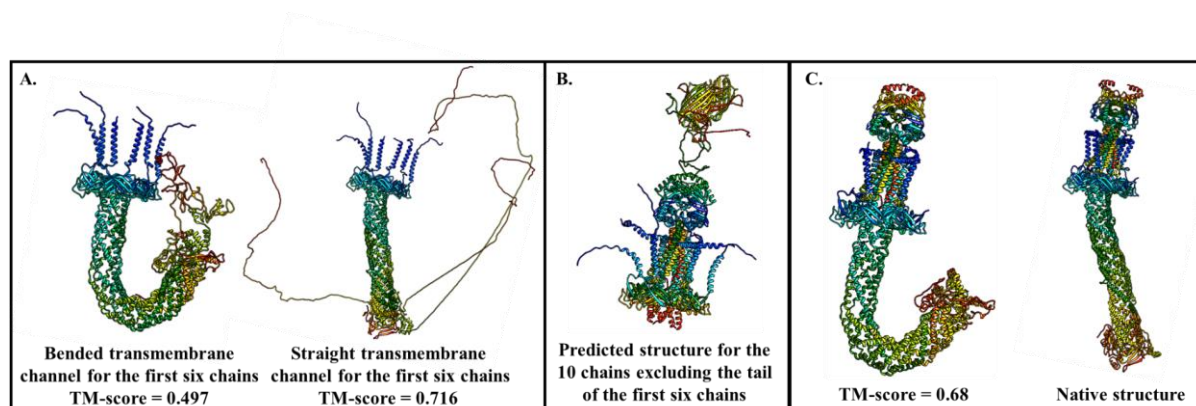


Figure S1. A. The two conformations (with TM-score = 0.497 and 0.716) of predicted complex structure for the first six chains of H1137 (A1B1C1D1E1F1); **B.** The predicted structure for the 10 chains excluding the tail of the first six chains; **C.** The native structure and the full-length complex structure (TM-score = 0.68) for H1137 built from the bended transmembrane channel for the first six chains (**left structure in plot A**) and the predicted structure for the 10 chains excluding the tail of the first six chains (**structure in plot B**).