GPCR-I-TASSER: A Hybrid Approach to G Protein-Coupled Receptor Structure Modeling and the Application to the Human Genome

**Highlights**
- New approach to ab initio GPCR structure assembly
- Use of mutagenesis data to assist 3D structure construction
- High-resolution structure models for 923 human GPCRs
- Provide reliably model for GPCR families that have no experimental structure

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**In Brief**
Zhang et al. developed a hybrid approach, GPCR-I-TASSER, for GPCR structure predictions, which combines experimental mutagenesis data with ab initio transmembrane helix assembly simulations. The method was applied to 1,026 GPCRs in the human genome, with successfully modeled targets containing many pharmaceutically important families with no previously solved structures.
GPCR-I-TASSER: A Hybrid Approach to G Protein-Coupled Receptor Structure Modeling and the Application to the Human Genome

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SUMMARY

Experimental structure determination remains difficult for G protein-coupled receptors (GPCRs). We propose a new hybrid protocol to construct GPCR structure models that integrates experimental mutagenesis data with ab initio transmembrane (TM) helix assembly simulations. The method was tested on 24 known GPCRs where the ab initio TM-helix assembly procedure constructed the correct fold for 20 cases. When combined with weak homology and sparse mutagenesis restraints, the method generated correct folds for all the tested cases with an average Cα root-mean-square deviation 2.4 Å in the TM regions. The new hybrid protocol was applied to model all 1,026 GPCRs in the human genome, where 923 have a high confidence score and are expected to have correct folds; these contain many pharmaceutically important families with no previously solved structures, including Trace amine, Prostanoids, Releasing hormones, Melanocortins, Vasopressin, and Neuropeptide Y receptors. The results demonstrate new progress on genome-wide structure modeling of TM proteins.

INTRODUCTION

G protein-coupled receptors (GPCRs) are integral membrane proteins that transmit chemical signals into a wide array of different cell types. Many diseases, including those associated with differentiation, proliferation, angiogenesis, cancer, development, and cell survival, involve malfunctions of the receptors, which make GPCRs one of the most widely used drug targets, accounting for over 40% of all pharmaceuticals approved by the US Food and Drug Administration (Eglen et al., 2007). While knowledge of GPCR structures provides important information for function elucidation and drug design, experimental determination of 3D structures of GPCR proteins has proved to be difficult. Significant efforts have been made on the technical improvement of GPCR expression and crystallization, which resulted in successful solution of 15 human GPCRs in the last 8 years since 2007 (Jaakola et al., 2008; Rasmussen et al., 2007). Although remarkable, these account for only a small portion of all GPCRs in the human genome, which is estimated to be approximately 1,000 (Takeda et al., 2002). The lack of atomic-level protein structure information for GPCRs has considerably hindered function annotation and structure-based drug discovery.

Significant efforts have also been made recently in the computational structure modeling of GPCR proteins, with progress witnessed on both new method development and modeling accuracy (Fanelli and De Benedetti, 2011). For instance, Barth et al. developed a structure modeling method to assemble helix-helix packing of membrane proteins with limited constraints. In 4 of 12 proteins, the method produced models of root-mean-square deviation (RMSD) < 4 Å to the X-ray structure (Barth et al., 2009). Chen et al. presented an interesting attempt to assemble protein transmembrane (TM) helices using distance restraints from sparse nuclear magnetic resonance (NMR) paramagnetic relaxation enhancement data. Constrained with a simple geometry pattern, TM-helix bundles up to seven helices can be correctly constructed using one to three restraints (Chen et al., 2011). Yang et al. combined multiple machine learning classifiers for generating inter-TM-helix contact predictions, which have an average accuracy of 62% in the top L/5 predictions. When incorporated in fragment assembly simulations, the predicted inter-helix contact restraints increased the TM-score of the final GPCR models by 37% (Yang et al., 2013). The contact-assisted structure assembly approach has also been exploited by several recent modeling studies for GPCR and other TM proteins (Hopf et al., 2012; Nugent and Jones, 2012).

Despite these advances, the majority of computational approaches to GPCR modeling rely on the detection of homologous templates (Fanelli and De Benedetti, 2011; Zhang et al., 2006). It is well known that pairwise sequence identity between GPCR families is low, and close homologous templates are not available for most of the unknown GPCR families (Archer et al., 2003). Despite the limited availability of global X-ray structures, numerous mutagenesis experiments have been performed on GPCRs to identify the critical residues and motifs, which contain spatial information for improving the modeling accuracy of GPCR structures. For example, the coupled activation and deactivation of residues in mutagenesis experiments usually indicate...
that the residues are spatially adjacent because they are binding to common ligands (Shi and Javitch, 2002). Furthermore, the orientation of mutated functional residues is usually toward the inside core of the seven-helix bundle due to the conservation of inter-helix contacts (Schushan et al., 2010). Thus, specific contacts and distance maps and residue orientations can be derived from the mutagenesis experimental data and converted into 3D restraints to guide the GPCR structure modeling simulations. This is particularly helpful for the modeling of structurally variable regions that cannot be directly transferred by homology inference.

In this work, we aim to develop a new hybrid structure assembly algorithm, GPCR-I-TASSER, by extending the iterative threading assembly method (I-TASSER). The major advantages of GPCR-I-TASSER over existing homology-based methods are:

1. A new GPCR-specific database, GPCR-RD (Zhang and Zhang, 2010a) containing experimental contact and helix orientation data from the literature and database mining, is exploited to improve the structural assembly accuracy;
2. When homology templates are unavailable, a new ab initio folding method is introduced for assembling the TM-helix bundle topology from scratch;
3. A set of new GPCR- and TM-specific energy terms is developed and incorporated into the I-TASSER force field to improve the structure assembly and refinement of both ab initio and threading template models. The major focus of this work is to construct reliable models for the GPCRs that lack close homologous templates.

To examine the efficiency, we first test GPCR-I-TASSER on all known GPCRs in the PDB and report the blind test results from the community-wide GPCR Dock experiments. It was found that the new pipeline can significantly improve the modeling accuracy of template structure identified from threading. For GPCRs without homologous templates, the ab initio folding process can construct an approximately correct fold for all receptors with assistance from sparse mutagenesis data. The algorithm was finally applied to the modeling of all putative GPCRs in the human genome. The comparison with new mutagenesis data and confidence scoring system showed that nearly 90% of targets are expected to have correct folds, including many GPCRs from the families that have no previously solved experimental structures.

RESULTS

GPCR-I-TASSER, as depicted in Figure 1, has three steps consisting of template identification (or ab initio TM-helix construction) and experimental restraint collection, Monte Carlo fragment assembly simulation, and atomic-level structural refinement (see Experimental Procedures and Supplemental Experimental Procedure, for details).

Benchmark Test on 24 Solved GPCRs

To benchmark GPCR-I-TASSER, we collected a set of test structures containing all 24 GPCRs solved so far in the PDB. Since there are multiple entries solved for single GPCRs, we used CD-HIT (Fu et al., 2012) to reduce the redundancy of these entries, which retains the entries having the longest structural coverage for each GPCR family. Table S1 lists the name and organism of the test GPCRs. Since many GPCRs were solved with fused external domains for facilitating crystal nucleation and structure determination, these domains have been excluded in our structure modeling. Table S2 lists the GPCR domains after manual trimming and the TM-helix annotations taken either from the original literature source or from manual inspection of the PDB structure. An updated list of all GPCRs solved in the PDB can be found at http://zhanglab.ccmb.med.umich.edu/GPCR-EXP.

Distant Homology Modeling

We first tested GPCR-I-TASSER by excluding all homology templates that have a sequence identity to target > 30% or are detectable by PSI-BLAST with an E value < 0.05. Despite the relatively stringent filters, many GPCR targets still have some analogous templates, which can be detected by LOMETS (Wu and Zhang, 2007). The threading search generated templates with an average RMSD = 5.74 (or 3.7) Å to the entire chain (or the TM-helix domains) of the native. The average TM-score of the templates is 0.675 (or 0.755). Here and afterward, the RMSD is calculated on Cα atoms only. TM-score is a sequence length-independent metric for measuring structure similarity with a range (0, 1). A TM-score > 0.5 generally corresponds to similar structures in the same SCOP/CATH fold family (Xu and Zhang, 2010). Such a high TM-score of the template detection by LOMETS probably reflects the focus of the experimental efforts that have been made on a set of similar GPCRs; therefore, templates can be inferred easily for the benchmark targets from other solved homologous GPCR structures. We have conducted a simple exercise by counting the homologous templates defined by the LOMETS alignments. The average number of homologous templates with a LOMETS Z-score above the confidence Z-score cutoff is 3.9 in this benchmark set, which is 2.4 times higher than the average for all other human GPCRs (1.6).
Despite the good quality of the threading alignments, GPCR-I-TASSER repacked the structure of the TM helices and drew the threading templates considerably closer to the native. Compared with the experimental structure, the first GPCR-I-TASSER models have the average RMSD reduced from 5.74 Å to 4.22 Å by 1.52 Å in the same threading alignment regions. The TM score increased from 0.675 to 0.806 by 19.4%. A detailed list of the threading templates and GPCR-I-TASSER models is given in Table S3, where values in parentheses are RMSD and TM-score data in the TM regions, and values after ‘/’ are RMSD of the GPCR-I-TASSER models in the threading-aligned region.

In Table S3, we also present the results by the widely used comparative modeling tool, MODELLER (Sali and Blundell, 1993), based on the best LOMETS templates. Since MODELLER is designed to construct models by optimally satisfying spatial restraints from templates, there is not much improvement of the final models over templates. Compared with LOMETS templates, the average RMSD of the MODELLER models increases from 5.74 to 8.07 Å and TM-score increases from 0.675 to 0.694 in the TM region; these moderate (RMSD/TM-score) increases are probably mainly a result of the length increase in the MODELLER modeling.

Goddard and colleagues developed a program, MembStruk, for GPCR structure prediction (Vaidehi et al., 2002). At the time of the MembStruk modeling, only one GPCR with experimental structure was available (i.e., bovine rhodopsin). MembStruk built a model with an RMSD = 3.1 Å in the TM-helix region and an RMSD = 8.3 Å in full-length regions of bovine rhodopsin. As the models generated by MembStruk are not available publicly, we compare GPCR-I-TASSER with MembStruk on this GPCR only. As shown in Table S3, the RMSD of the GPCR-I-TASSER model for bovine rhodopsin (2hpB) is 1.35/5.25 Å in the TM-helix/all regions, which is 1.75/3.05 Å lower than the MembStruk model. However, we note that this comparison might not be entirely fair because there are now more GPCR structures that can serve as templates. We have re-run GPCR-I-TASSER by excluding all GPCR templates (but keeping other membrane structures) in the template library, which resulted in the first predicted model of bovine rhodopsin with 1.82/6.31 Å in the TM-helix/all regions; these RMSD values are slightly higher than the data in Table S3 but still considerably lower than that of the MembStruk results.

In Figure 2A, we present two examples from human opioid receptor (PDB: 4e4A1) (Granier et al., 2012) and human serotonin receptor (PDB: 4iarA1) (Wang et al., 2013), which represent two targets with the most significant structure refinements, where the threading templates have a TM-score = 0.644 and 0.645 but GPCR-I-TASSER refined the models to TM-score = 0.894 and 0.884, respectively. The major improvement occurs at the TM-helix regions, where the RMSD was reduced from 4.66 and 4.67 Å to 1.44 and 1.7 Å, respectively. This improvement is mainly attributed to the new GPCR-specific helical packing potential and the atomic-level fragment-guided molecule dynamic (FG-MD) refinements.

Compared with the TM-helix regions, the modeling of loop structure is more challenging since these regions are less conserved and the threading programs often have alignment gaps. In the 24 proteins, there are on average 7.9% of residues without threading alignments, which are mainly located on the loops/tails. The GPCR-I-TASSER pipeline constructs models for these regions by a lattice-based, ab initio structure assembly procedure extended from the I-TASSER protocol, which resulted in models with an average RMSD = 5.37 Å for the six intra- and extracellular loops. For the functionally important second extracellular loop (EL2) that is often involved in ligand recognition and receptor activation, the average RMSD is 3.85 Å, with an average length of 20.4 amino acids in this test.

It should be mentioned that the quality of template-based structure modeling is sensitive to the level of homologous template filtering. For instance, if we only filtered out the templates of sequence identity > 30% (i.e., dropping off the PSI-BLAST E-value filter) as done in many previous benchmark experiments of structure prediction (Simons et al., 1999; Zhang and Skolnick, 2004a), the TM-score and RMSD of the threading templates will improve to 0.756 and 4.65 Å, respectively, while the quality of the GPCR-I-TASSER models will be improved accordingly with an average TM-score = 0.912 and RMSD = 3.21 Å (or 1.57 Å in the TM-helix and 3.35 Å in the loop regions).

### Ab Initio GPCR Folding

Most GPCRs in the human genome are not closely homologous to the solved GPCRs in the PDB. To examine the ability of GPCR-I-TASSER in ab initio structure assembly, we exploited a second level of template filtering, i.e., to regenerate the models by excluding all GPCR and membrane proteins from our template library.

Since all correct templates have been excluded, it is expected that the templates detected by threading will now have a completely different topology from the native structures. The average TM-score of the templates with the highest Z-score is 0.096, which is well below the average of random structure pairs (0.17) (Xu and Zhang, 2010; Zhang and Skolnick, 2004b). When we applied MODELLER (Sali and Blundell, 1993) to these
templates for full-length model construction using the default setting, a similar set of random models were obtained with an average TM-score = 0.142 and RMSD = 21.74 Å (Table 1). This is expected again because MODELLER was designed to construct structure models by satisfying spatial restraints from templates, an approach best suitable to the targets with close homologous templates.

To build a de novo TM-helix bundle topology, GPCR-I-TASSER first performs a rapid ab initio Monte Carlo assembly simulation, which starts from ideal helix bundles (Figure 3), with the conformational search guided mainly by the generic atomic contact and membrane transfer potentials (Equations S2 and S3 in Supplemental Experimental Procedure). The structural decoys were clustered by SPICKER (Zhang and Skolnick, 2004c), which resulted in the first ab initio models with an average TM-score = 0.389 and RMSD = 11.39 Å (Table 1). In nine cases, the models have a TM-score > 0.4, which indicates an approximately correct topology of the TM-helix assembly (Xu and Zhang, 2010). If we consider the best in the top five models, this number increases to 17 (see Table S4).

Starting from the ab initio TM-helix models and the low-resolution threading template alignments, GPCR-I-TASSER Monte Carlo simulations were conducted to reassemble the TM helices that have the relative orientations restricted by the loop structures. Meanwhile, 294 spatial restraints were extracted from the GPCR-RD database for the 24 test GPCRs. On average, seven residue-residue contact restraints and five helix orientation restraints per target were used to constrain the simulations. This procedure generated full-length models with an average TM-score = 0.517, which is 32% higher than that of the models created by ab initio folding. All the targets have a TM-score > 0.4, and 20 of 24 targets have a TM-score > 0.5 (Table S4).

To test the effect of the mutagenesis restraints, we also ran a version of GPCR-I-TASSER without restraints from GPCR-RD. The average TM-score of the final model decreased by 3.9%. The TM-score reduction in this set of models was found considerably larger than that of the template-based models from the last section (1.4%); this is understandable because the mutagenesis restraints are implemented using a relatively large distance cutoff (i.e. \( d_{ij} < 10 \) Å in Equation S9) or with helix orientation adjustment (Equation S10), which should have a stronger effect on refining models with low resolution.

To illustrate the procedure of ab initio folding, in Figure 2B we show the structural superposition of the predicted models for the adenosine \( A_{2a} \) receptor over the experimental structure (PDB: 3emlA1) (Jaakola et al., 2008) from the three modeling steps. The LOMETS programs hit incorrect templates, which resulted in the MODELLER model with a different topology (TM-score = 0.188). The ab initio folding procedure rearranged artificial helices and constructed a TM-helix bundle with approximately correct topology (TM-score = 0.496). The GPCR-I-TASSER refinement simulations improved the structural model to a TM-score = 0.581.

Figure 2. Illustrative Examples of GPCR Structural Modeling, with Blue and Red Representing Model and Experimental Structure, Respectively

(A) Template-based modeling on human opioid receptor (PDB: 4ej4A1) (Granier et al., 2012) and serotonin receptor (PDB: 4iarA1) (Wang et al., 2013) receptors with close homologous templates detectable by PSI-BLAST or sequence identity > 30% excluded. Left to right columns are the LOMETS templates and GPCR-I-TASSER models overlaid on the native in the whole chain and TM-helix regions, respectively.

(B) Ab initio modeling on human adenosine \( A_{2a} \) receptor (PDB: 3emlA1) (Jaakola et al., 2008) with all homologous and membrane templates excluded. Left to right columns are the models built by MODELLER, ab initio assembly, and GPCR-I-TASSER models overlaid with TM regions of the native structure, respectively.
The data for the 24 benchmark proteins, including template alignments, ab initio folding, and GPCR-I-TASSER models, are downloadable at http://zhanglab.ccmb.med.umich.edu/GPCR-I-TASSER/benchmark.

Blind Test in the GPCR Dock Experiment
As a blind test of GPCR-I-TASSER, we participated (as UMich/0460) in the community-wide GPCR Structure-based Homology Modeling and Docking Assessment 2010 (or GPCR Dock 2010), organized by Kufareva et al. (2011). In the experiment, the organizers requested structure predictions for three GPCR-ligand complexes that were solved by Stevens and coworkers (Chien et al., 2010; Wu et al., 2010): the human CXCR4 chemokine receptor bound either to the small molecule antagonist IT1t or to the peptide antagonist CVX15, and the human dopamine D3 receptor with eticlopride. The predictions were blind, as the target structures were not released until the predictions were completed.

In Figure 4, we show the GPCR models built by the GPCR-I-TASSER pipeline in GPCR Dock 2010, where the GPCR-RD restraint data were not exploited. First, LOMETS threading identified B1AR and B2AR as the templates for the CXCR4 and D3 receptors, respectively, which have a TM-score of 0.695 and 0.627, respectively. The RMSDs of the templates in the threading-aligned region of the TM helices are 3.06 Å and 1.61 Å, respectively. After GPCR-I-TASSER reassembly, the final models have a TM-score = 0.771, 0.768, 0.917 for CXCR4/IT1t, CXCR4/CVX15, and D3/eticlopride, respectively, which are 11%, 11%, and 46% higher than the initial templates. In the same threading-aligned TM region, RMSDs of the final models are 2.08 Å, 2.58 Å, and 1.26 Å, respectively, which are 0.98 Å, 0.48 Å, and 0.35 Å lower than the initial templates. These results confirm that GPCR-I-TASSER has the ability to draw threading templates considerably closer to the native structure.

The ligand-bound GPCR models were generated by BSP-SLIM (Lee and Zhang, 2012), which first identified the ligand-binding pocket positions on the receptor protein by structurally aligning the receptor models to known complex structures in the PDB using TM-align (Zhang and Skolnick, 2005). The ligand-docking models were then generated from a conformational search with ligands constrained in the predicted binding pocket. The ligand-GPCR binding energy in BSP-SLIM consists of hydrogen-bonding, statistical contact potential, solvation, and van der Waals interactions. The RMSDs of the final ligand models by BSP-SLIM are 9.61, 7.35, and 3.51 Å, for CXCR4/IT1t, CXCR4/CVX15, and D3/eticlopride, respectively (Figure 4).

Table S5 lists the top ten groups in GPCR Dock 2010 based on the cumulative Z-scores of the receptor and ligand models for all three targets. Among the 35 participant groups, the UMich-Zhang/0460 groups using GPCR-I-TASSER had the highest Z-score in the receptor models and the second highest in the ligand-docking positions, which resulted in the highest total Z-score of receptor and ligand models, according to the analysis by Kufareva et al. (2011). The most noticeable success is on the distant homologous target CXCR4/CVX15, whereby the assessors commented in the assessment article that “Modeling the CXCR4/CVX15 peptide complex represented the biggest challenge of GPCR Dock 2010. The top model of this complex (#5 by UMich-Zhang) has the Z-score of 2.4, thus far exceeding other models in accuracy” (Kufareva et al., 2011). For the two other less challenging targets (CXCR4/IT1t and D3/eticlopride), however, although the TM-backbone RMSD of the receptor models is ranked at the top for both targets, the accuracy of the functionally important EL2 and the ligand-docking score are considerably worse than the top performing groups (http://abl.uccs.edu/GPCRDock2010), highlighting the need to improve EL2 modeling and BSP-SLIM docking.

We note that the GPCR Dock experiment aims to benchmark the modeling of GPCR-ligand complexes with an emphasis on the ligand-docking technique. The receptor structure submitted by the other groups may not reflect the best receptor models due to the consideration of ligand-docking interactions. Nevertheless, the data provide a partial but independent assessment of GPCR-I-TASSER on the GPCR structure modeling compared with other state-of-the-art approaches.

Structure Modeling of 1,026 GPCRs in the Human Genome

GPCR-I-TASSER Modeling
A total of 1,063 distinct GPCR sequences in the human genome were collected by scanning the databases GPCR-DB (http://www.gpcr.org/7tm/data/) and UniProt (http://www.uniprot.org/docs/7tmlist). Since errors often exist in automated data
collection, we used a semi-manual procedure to examine these GPCR sequences: First, we generated TM-helix prediction by three TM prediction programs from HMMTOP (Tusnady and Simon, 1998), MEMSAT (Jones et al., 1994), and TMHMM (Krogh et al., 2001). If the number of TM helices predicted by any of the programs is less than seven or the number of overlapped residues between the TM regions by the three programs is less than five, we manually examined these sequences (about 400) by checking the UniProt annotation on the TM helices. In case there is no UniProt annotation, we used the GPCR-I-TASSER structure models to extract the TM helices. With this manual verification, we identified 37 non-GPCR sequences where most of them are extracellular domains attached to the receptor but mis-classified as GPCRs. 1,026 validated GPCR sequences were retained for GPCR-I-TASSER modeling.

The GPCR sequences were first threaded through the PDB library using LOMETS (Wu and Zhang, 2007). In 862 cases, at least one of the programs used by LOMETS identified template structures with a significant Z-score above the confidence cutoff of the corresponding program. For the rest of proteins, we constructed the initial TM-helix bundle conformations using the ab initio folding procedure.

In the next step, we collected the sparse experimental data from GPCR-RD (Zhang and Zhang, 2010a), a manually curated database containing multiple GPCR data on site-directed mutagenesis, electron microscopy, neutron diffraction, Fourier transform infrared spectroscopy, and disulfide bridge. The experimentally identified disulfide bridges and functionally important residues (binding to a particular ligand) indicate that these residues should be close to each other to perform their functions. So we applied contact restraints to these residue pairs as described in Equation S9. Besides, the majority of the functionally related point mutations should face to the inside core of the TM-helix bundle (Schushan et al., 2010), which are used to guide the packing of helix orientations as described in Equation S10.

These resulted in 3,425 contacts and 1,401 orientation restraints for the 1,026 human GPCRs. These restraints, together with the threading alignments and ab initio TM-helix models, were used to guide the GPCR-I-TASSER assembly simulations. The atomic details were finally refined by the FG-MD simulation program (Zhang et al., 2011).

For the sequences containing extra domains, which are detected by TheaDom (Xue et al., 2013), models are created for each domain individually using GPCR-I-TASSER (for TM domain) or I-TASSER (for globular domain). The full-length GPCR models are then constructed by assembling the domain structures as described in the Supplemental Experimental Procedure. This domain parsing and assembly procedure can improve the confidence score and modeling accuracy of the individual domains as demonstrated in previous benchmark tests (Zhang, 2014). A multiple-domain example from Q6ZM19, which contains a TM and a globular domain, is presented in Figure 5A, where the domain parsing and assembly procedure increased the C-score (defined below) from −1.79 of the full-chain GPCR-I-TASSER model to 1.11 for the globular domain and 1.32 for the GPCR domain, respectively.

All the models for the 1,026 human GPCRs by GPCR-I-TASSER, together with the template alignment, local and global confidence scoring annotations, and the secondary structure and solvation predictions, are deposited in the GPCR-HGmod database (http://zhanglab.ccmb.med.umich.edu/GPCR-HGmod/). Due to the sensitivity of the model quality to the templates in the PDB, the model prediction for all human GPCRs will be updated every 12 months (the old models will be archived in the online database for tracking progress).

**Global Confidence Score Analyses**

In Figure 6, we present a histogram distribution of the confidence scores (C-score) of the GPCR-I-TASSER models. Here, C-score is defined as the product of the normalized Z-score from LOMETS threading and the cluster density from SPICKER, i.e.

\[
C\text{-score} = \ln \left( \frac{M / M_{\text{tot}}}{\text{RMSD}} \right)^{1/g} \left[ \sum_{i=1}^{9} \left( \frac{Z_i}{Z_{i,\text{cut}}} \right) \right] \quad (\text{Equation 1})
\]

where \(M / M_{\text{tot}}\) is the normalized multiplicity of the structure decoys in the cluster, \(\text{RMSD}\) is the average RMSD of the decoys to the cluster centroid, \(Z_i\) is the highest Z-score of the template detected by the \(i\)th threading program in LOMETS, and \(Z_{i,\text{cut}}\) is the corresponding Z-score cutoff for distinguishing between good and bad template alignments (see Supplemental Experimental Procedure). The C-score has a strong correlation coefficient (0.91) to the actual TM-score of the predicted models based on large-scale benchmark tests (Zhang, 2008).

From the histogram of TM-score data obtained from the benchmark study, we roughly estimated the number of GPCRs expected to have a TM score > 0.5, which indicates a similar fold to the target, i.e., \(\sum_{m=15}^{N_{\text{bin}}} N_{\text{Mtot}}\) where \(M_{\text{bin}} = 15\) is the number of bins split in the C-score space, \(N_{\text{Mtot}}\) is the number of GPCR-I-TASSER models in the \(m\)th C-score bin, and \(r_{\text{Mtot}}\) is the folding rate for the GPCR-I-TASSER/I-TASSER models in the \(m\)th bin based on large-scale benchmark tests on 1,107 known proteins, including the 24 GPCR proteins from the PDB library. We found that there are 923 cases out of the 1,026 GPCRs that should have the highest ranked model with a correct topology.
(TM-score > 0.5). This number is similar to the direct counting of GPCRs with a C-score > -1.5, which is a cutoff that approximately corresponds to the correct models in the benchmark data (Zhang, 2008). In addition, all the models predicted by GPCR-I-TASSER have the typical seven TM-helix bundle topology because of the ab initio folding algorithm and the GPCR-RD experimental restraints, although a number of GPCRs (~200) did not have any TM templates detected by the threading search. Here, we note that the C-score histogram in Figure 6 was calculated based on the whole-chain GPCR sequences, which may contain multiple domains. If we count only for the TM domains, the number of folded cases should be slightly higher since the domain parsing and assembly procedure can increase the C-score and modeling accuracy of individual domains, as illustrated in Figure 5.

The 923 high C-score GPCRs cover 53 of the 54 families in the human genome; the only missed family is Family 3 metabotropic glutamate and calcium receptors, none of the four members of which (Q8NFJ5, Q9NQ84, Q9NZD1, Q9NZH0) has a confident prediction from GPCR-I-TASSER. Since the experimentally solved GPCRs cover only 16 families (Table S1), such a high family coverage partly demonstrates the ability of GPCR-I-TASSER to model distant homology proteins across different families.

In Table S6, we list the top 20 families that have the highest number of GPCRs with a C-score > -1.5. As expected, for the families that have some members with experimentally solved structures, all the GPCRs have high C-score models generated due to the easily detected homologous templates. While most of the high C-score GPCRs are from the Odorant/olfactory and gustatory family, GPCR-I-TASSER also generated models of high C-scores for many families that have no experimentally solved members but are pharmaceutically important drug targets, including Trace amine-associated (brain monoamine regulation; Panas et al., 2012), Prostanoids (initiating cancer and inflammation pathways; Breyer et al., 2001), Releasing hor-

Figure 5. Illustration Examples of the Multi-Domain Assembly and TM Structure Packing

(A) Multi-domain modeling for UniProt: Q6ZM19. The N-terminal domain was excised from the UniProt sequence following the ThreaDom linker prediction. This globular domain and the core TM domain were modeled separately by I-TASSER and GPCR-I-TASSER; the final model was assembled from the structure of two domains with the domain orientation decided by the full-chain GPCR-I-TASSER model. An FG-MD refinement was conducted on the full-chain models to eliminate possible steric clashes.

(B) GPCR-I-TASSER models for UniProt: Q8NGQ3. The left panel shows the model generated when the membrane repulsive energy was turned off, where the N-terminal tail entered into the transmembrane region. The right panel shows that the tails were moved out of the membrane when the membrane repulsive energy in Equation S4 was included.
In Figure 7, we present an example of the estimated local structure accuracy and B-factor profiles, in control with the X-ray crystallography data from the histamine H1 receptor (PDB: 3rzeA1) (Shimamura et al., 2011). The distance errors are mainly located in the unaligned loop and tail regions, which are highly consistent with the ResQ estimation. These profiles are provided for each of the GPCR-I-TASSER models in the GPCR-HGmod database.

**Sequence and Structure Networks of Human GPCRs**

Given the sequence and structural models generated, we present in Figure 8 a 2D view of the sequence and structure distributions of all 1,026 GPCRs in human prepared using Cytoscape 2.8 (Cline et al., 2007). The sequence similarity matrix is measured by pairwise sequence identity calculated by NW-align (http://zhanglab.ccmb.med.umich.edu/NW-align/), where a cutoff of 50% is used to ensure that the connected nodes have conserved functionality. There are 151 GPCR clusters or orphans with an average number of neighbors = 8.9, and the average number of neighbors of non-orphan clusters is 10.9.

In the structure space, the distance matrix is measured by the pairwise TM-score of the GPCR-I-TASSER models, where a cut-off of TM-score > 0.95 is used for node connections to distinguish subtle structural similarity. The total number of the clusters or orphans is 171 in structure space, similar to the number in sequence space. However, the average number of neighbors for the non-orphan clusters is 41.2, which is much higher than that in the sequence space, despite the stringent TM-score cutoff. These data suggest that human GPCRs are much more converged in structure space than in sequence space. We have re-examined the data using more permissive sequence identity cutoffs in the 30%-50% range or TM-score cutoffs in 0.6-0.95 but the clustering data did not qualitatively change.

The high degree of conservation in structure space is partly because GPCR structures are largely constrained by the seven TM-helix bundle topology, despite considerable variations existing in the relative location and orientation of helices and arrangement of loops. There are, however, a few big families, such as olfactory receptors, which have a highly similar structure but with very diverse pairwise sequence identity. The biggest cluster in structure space includes 711 members, which all belong to the class A rhodopsin-like receptors and have a sequence identity as low as 17%. Thus, high-resolution structure modeling should serve as a useful complement to sequence-based analysis for GPCR function annotation.

**Cross-Validation of GPCR-I-TASSER Models with Experimental Mutagenesis Data**

Although the number of experimental 3D structures for GPCRs is low, numerous experiments have been performed on GPCRs to identify the critical residues and motifs from site-directed mutagenesis, solid-state NMR, and neutron diffraction data. Many of these data have been collected in the GPCR-RD database (Zhang and Zhang, 2010a) and converted into the 3D spatial restraints to guide the GPCR-I-TASSER structure modeling. To validate the GPCR-I-TASSER structure models, we compared the predictions with recently collected mutagenesis data that had not yet been incorporated into the GPCR-RD at the time of modeling.

To test the high-confidence models, we collected 58 GPCR-I-TASSER models that have a C-score > 1.0 and at least one contact residue pair from the new mutagenesis experiments. Excluding the N- and C-terminal tails, we found that all the first models by GPCR-I-TASSER have their residue contacts consistent with the mutagenesis data, i.e., with Cα distance < 10 Å for

**Figure 6. C-score Distribution of GPCR-I-TASSER Models for 1,026 GPCRs in the Human Genome**

(Inset) Percentage of the cumulative counts along with the C-score distribution.

**Figure 7. An Illustration of ResQ-Based Local Structure Quality Estimation on the Human Histamine H1 Receptor**

(A) Overlay of the GPCR-I-TASSER model (blue) and the X-ray structure; (B) secondary structure assignment by STRIDE; (C) predicted and actual distance deviations of the model from the X-ray structure; (D) predicted and actual B-factor profiles of the target.

PDB: 3rzeA1 (Shimamura et al., 2011).
contact restraints or $E_{\text{orientation}} < 0$ from Equation S10 for the orientation restraints. In Figure 9, we present a set of randomly selected examples from the high-confidence GPCR-I-TASSER models where the key functional residues are highlighted.

Figure 9A shows the GPCR-I-TASSER model for the formyl peptide receptors, which respond to chemokines and chemottractants found on the surface of phagocytes. There are three residue pairs (D106-R205, A68-N44, and N44-N66) and two functionally related residues (D106 and R205, which are supposed to be in contact with each other based on the mutation and ligand-binding analysis experiments (Lala et al., 1993; Mills et al., 2000; Prossnitz et al., 1999). These residue pairs are all in contact in our formyl peptide receptor model with distances $< 10 \text{ Å}$ (Figure 9A).

Figure 9B shows a second example from the C5a anaphylatoxin chemotactic receptors that mediate cell activation and receptor desensitization. One disulfide bond (C293-C86), two residue pairs (P257-C285 and G210-M120), and two functionally important residues (P170 and Q259) should be in contact according to the experimental data (Baranski et al., 1999; Giannini et al., 1995; Kolakowski et al., 1995; Raffetseder et al., 1996), which is also consistent with the GPCR-I-TASSER models.

Figures 9C and 9D are two other examples from the galanin receptor and the type 1 angiotensin II receptor, respectively. In Figure 9C, one contact pair (H263-R285) and four functional residues (H263, H267, H285, and H289) from the mutagenesis experiments (Berthold et al., 1997; Kask et al., 1996) are all consistent with the GPCR-I-TASSER model. In Figure 9D, function-related residues (N111, A104, S115, W153, T260, and N295) form a well-shaped binding pocket in the GPCR-I-TASSER model, which were identified in the mutagenesis experiments as critical binding residues with the non-peptide ligands (Perlman et al., 1995, 1997; Schambye et al., 1994).

Conclusions

Progress in experimental GPCR structure determination has been slow due to difficulties in acquiring high-resolution experimental data. Computational approaches can also produce high-resolution models, but so far they have been limited to cases where a homologous template is available. To address these limitations, we have developed a new hybrid method, GPCR-I-TASSER, which can exploit distant homology templates and including membrane repulsion, hydrophobic moment, and enhanced aromatic and cation-$\pi$ interactions, were introduced to guide the GPCR-I-TASSER structure assembly simulations. Our unpublished data showed that the inclusion of these TM-specific potentials resulted in a TM-score increase of the GPCR structure models by 3.5% on the test proteins with a $p$ value $< 10^{-5}$. For the targets that do not have close homologies, a new ab initio folding procedure was developed to construct the TM-helix bundles from scratch, which are further refined by the fragment assembly simulations. This hybrid pipeline enables the structure construction of different families of GPCRs, which is essential for genome-wide GPCR modeling and GPCR-ligand screening. Although progress was made to advance computational methods for GPCR modeling, accuracy can still be limited, especially in the de novo cases and in the loop and tail regions. We provide local confidence scores to help identify these uncertain regions.

The GPCR-I-TASSER method was tested on two benchmarks. First, it was tested on 24 GPCR proteins that have an experimentally solved structure. After excluding all homologous proteins with a sequence identity $> 30\%$ and templates detectable by PSI-BLAST, the threading programs successfully identified templates of correct topology with an average TM-score $= 0.675$ and RMSD $= 5.74 \text{ Å}$. After the GPCR-I-TASSER structural reassembly refinement, the TM-score of final models increased to $0.806$ by $19.4\%$ and RMSD reduced to $4.22 \text{ Å}$. To address these limitations, we have developed a new hybrid method, GPCR-I-TASSER, which can exploit distant homology templates and spatial restraints from low-resolution but more easy to acquire experimental data to assist high-resolution GPCR structure modeling.

In addition to the generic knowledge-based force field, a set of new GPCR- and TM-protein-specific energy terms, computed from the high-confidence GPCR-I-TASSER models, were introduced. First, it was tested on two GPCR proteins that have an experimentally solved structure. After excluding all homologous proteins with a sequence identity $> 30\%$ and templates detectable by PSI-BLAST, the threading programs successfully identified templates of correct topology with an average TM-score $= 0.675$ and RMSD $= 5.74 \text{ Å}$. After the GPCR-I-TASSER structural reassembly refinement, the TM-score of final models increased to $0.806$ by $19.4\%$ and RMSD reduced to $4.22 \text{ Å}$. To address these limitations, we have developed a new hybrid method, GPCR-I-TASSER, which can exploit distant homology templates and spatial restraints from low-resolution but more easy to acquire experimental data to assist high-resolution GPCR structure modeling.

In addition to the generic knowledge-based force field, a set of new GPCR- and TM-protein-specific energy terms, including membrane repulsion, hydrophobic moment, and enhanced aromatic and cation-$\pi$ interactions, were introduced to guide the GPCR-I-TASSER structure assembly simulations. Our unpublished data showed that the inclusion of these TM-specific potentials resulted in a TM-score increase of the GPCR structure models by 3.5% on the test proteins with a $p$ value $< 10^{-5}$. For the targets that do not have close homologies, a new ab initio folding procedure was developed to construct the TM-helix bundles from scratch, which are further refined by the fragment assembly simulations. This hybrid pipeline enables the structure construction of different families of GPCRs, which is essential for genome-wide GPCR modeling and GPCR-ligand screening. Although progress was made to advance computational methods for GPCR modeling, accuracy can still be limited, especially in the de novo cases and in the loop and tail regions. We provide local confidence scores to help identify these uncertain regions.

Second, we tested GPCR-I-TASSER in the community-wide blind GPCR Dock experiment. The final models of the CXCR4 and D3 receptors have a TM-score $11\%$ and $46\%$ higher than the threading templates. The RMSD of the TM regions was $2.08$, $2.58$, and $1.26 \text{ Å}$, which are $0.98$, $0.48$, and $0.35 \text{ Å}$ lower than the corresponding initial templates, respectively. These
predictions have a higher average significance score (Z-score) than the other 34 predictor groups.

We applied the GPCR-I-TASSER pipeline to the modeling of all 1,026 putative GPCR proteins collected from the UniProt and GPCR-DB databases. There are 923 cases that are expected to have a correct global fold with a predicted TM-score > 0.5, based on the correlation between C score and TM score. The targets with high-confidence models include many unsolved but pharmaceutically important GPCR families including Trace amine, Prostanoids, Releasing hormones, Melanocortins, Vasopressin, and Neuropeptide Y receptors. The sequence and structure-based clustering studies showed that the structures of GPCRs are more conserved than the sequences during evolution. As part of cross-validations, we compared the GPCR-I-TASSER models with experimental mutagenesis data, which were not used in our structure modeling. Consistency with the experimental data was demonstrated in all GPCR-I-TASSER models that have a confidence score above 1.0. These results demonstrated new progress on genome-wide structure modeling of GPCRs.

EXPERIMENTAL PROCEDURES

GPCR-I-TASSER is designed to construct 3D models of GPCRs and consists of three steps of TM-helix assembly, full-length structure reassembly simulations, and model selection and atomic-level structure refinement (Figure 1). The processes are outlined below, with detailed procedures described in Supplemental Experimental Procedure.

Generation of Transmembrane Helix Framework

The query GPCR sequence is threaded through the PDB by LOMETS (Wu and Zhang, 2007), a meta-threading approach containing nine cutting-edge threading programs, to identify appropriate structure templates. The regions of extra-/intra-cellular loops and TM helices are predicted separately and introduced as additional alignment constraints to enhance the accuracy of the threading alignments for GPCRs (see Equation S1 in Supplemental Experimental Procedure). If no significant template is identified, a new ab initio folding approach is developed to construct the TM framework by replica-exchange Monte Carlo (MC) simulation, starting from seven ideal helices located sequentially along a perimeter of 8 Å. The MC movements involve translation, rotation, and tilting of the helices, and sequence shifts along the helix, addition/deletion of residues, and helix kinking (Figure 3). The simulations are guided by a simple force field consisting of a knowledge-based, distance-specific contact potential, RW (Random Walk; Zhang and Zhang, 2010b), and the free energy change of GPCR and water/lipid interactions (Lomize et al., 2006) (see Equations S2 and S3 in Supplemental Experimental Procedure).

Template-Based Fragment Assembly Simulations

Full-length GPCR models were constructed by reassembling the continuous fragments (mainly TM helices) excised from LOMETS threading alignments or ab initio TM-helix models, following the I-TASSER protocol (Roy et al., 2010; Yang et al., 2015). The force field of the GPCR-I-TASSER simulation consists of three components. The first component is a generic knowledge-based potential extended from I-TASSER that includes statistical Cα and side-chain contact potentials, backbone-orientation specific hydrogen-bond, solvation from neural network prediction, and predicted secondary structure propensities; the second is spatial restraints derived from LOMETS templates and/or ab initio TM-helix models, which consists of Cα distance maps and Cα and side-chain contacts; and the third component consists of six GPCR- and/or TM-specific energy terms as described in Equations S4–S10 in Supplemental Experimental Procedure.

Two types of spatial restraints are derived from the site-directed mutagenesis and affinity labeling experiments collected from the GPCR-RD database (Zhang and Zhang, 2010b). These include contact restraints accounting for the experimentally identified disulfide bridges and the functionally important interactions. (A) Formyl peptide receptor; (B) C5a anaphylatoxin chemotactic receptor; (C) type 1 galanin receptor; (D) type 1 angiotensin II receptor. Dashed lines connect the residue pairs supposed to be in contact in the mutagenesis data, which are all within 10 Å in the GPCR-I-TASSER models.
residues (Equation S9), and an orientation restraint of TM-helix to count for the functionally related point mutations (Equation S10). A general membrane repulsive potential is introduced in Equation S4 to enhance the GPCR-specific topology, i.e., all non-TM-helix residues should be excluded from the TM regions (see Figure 5B).

**Model Selection and Fragment-Guided Structure Refinement**

Structure decoys generated in GPCR-I-TASSER are submitted to SPICKER (Zhang and Skolnick, 2004c) for structure clustering. The decoys with the highest number of structural neighbors are selected, with full-atomic models refined by the FG-MD simulations (Zhang et al., 2011). Furthermore, the SPICKER centroid model is used as a probe to identify analog fragments from the PDB by TM-align (Zhang and Skolnick, 2005), which provides additional spatial restraints to improve the energy landscape funnel in atomic-level structure refinements in FG-MD.

**Multiple-Domain Assembly**

For the GPCRs of multi-domains, we first use ThreDom (Xue et al., 2013) to identify the domain boundary and then use GPCR-I-TASSER and I-TASSER to fold the receptor and globular domains separately. The full-length models are finally built by docking the domain models using the whole-chain model as a reference template.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedure and seven tables and can be found with this article online at http://dx.doi.org/10.1016/j.str.2015.06.007.

**AUTHOR CONTRIBUTIONS**

Y.Z. conceived the project; J.Z., J.Y., and R.J. conduct the calculation and data analysis; J.Z., J.Y., and Y.Z. wrote the article.

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