

Simultaneous prediction of protein folding and docking at high resolution

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Interleaved dimers and higher order symmetric oligomers are ubiquitous in biology but present a challenge to de novo structure prediction methodology: The structure adopted by a monomer can be stabilized largely by interactions with other monomers and hence not the lowest energy state of a single chain. Building on the Rosetta framework, we present a general method to simultaneously model the folding and docking of multiple-chain interleaved homo-oligomers. For more than a third of the cases in a benchmark set of interleaved homo-oligomers, the method generates near-native models of large α -helical bundles, interlocking β sandwiches, and interleaved $\alpha\beta$ motifs with an accuracy high enough for molecular replacement based phasing. With the incorporation of NMR chemical shift information, accurate models can be obtained consistently for symmetric complexes with as many as 192 total amino acids; a blind prediction was within 1 Å rmsd of the traditionally determined NMR structure, and fit independently collected RDC data equally well. Together, these results show that the Rosetta “fold-and-dock” protocol can produce models of homo-oligomeric complexes with near-atomic-level accuracy and should be useful for crystallographic phasing and the rapid determination of the structures of multimers with limited NMR information.

homo-oligomers | molecular replacement | NMR structure inference | protein structure prediction | symmetry

The majority of expressed proteins function within symmetrical homomeric complexes (1–3). Although a boon for evolving functional diversity (4), this ubiquity of oligomeric structures poses numerous challenges for modern structural biology. The phasing of crystallographic data by molecular replacement and NMR structural inference are complicated by the increasing number of degrees of freedom and spectral degeneracy, respectively, in multimeric systems. The ability to predict de novo the structures of multiple interacting molecular chains would potentially alleviate these problems, allowing unphased diffraction measurements or ambiguously assigned NMR spectra to be resurrected as constraints or as independent validation for *in silico* structural inference.

Accurate modeling of previously unseen homomeric structures has not been demonstrated at high resolution. Significant progress has occurred in modeling the folds of individual monomeric soluble proteins (5–7) and the docking arrangements of predefined monomers (8, 9). However, as proteins interact with other proteins, nucleic acids, or smaller molecules, their lowest free-energy backbone conformations typically shift in response to their partners. These often dramatic structural changes have been a persistent and unsolved issue in blind prediction trials of recent years (8).

In this report, we show how two different methods developed for de novo conformational sampling (for folding and for docking) can be melded into a more general procedure that permits the blind prediction of intertwined complexes of proteins at near-atomic resolution. Some of the test systems involve up to several hundred residues, much larger than previous targets of high-resolution de novo modeling; symmetry plays a crucial role in reducing the

number of degrees of freedom that need to be sampled. The new approach is based on the Rosetta framework for molecular modeling (10).

Results and Discussion

Overview of Rosetta “Fold-and-Dock” Protocol. For each molecular complex, we start from fully extended monomer chains in randomly generated symmetric rigid body arrangements (Fig. 1). The first of two stages of automated de novo modeling alternates sets of fragment insertion moves that perturb the backbone conformation of protein monomers (11) with moves that perturb the symmetric docking arrangement of the monomers (12). The conformational energy is determined by the low-resolution energy function previously developed for monomer protein structure prediction (6, 11) with coarse-grained terms for backbone hydrogen bonds and hydrophobic interactions. Cyclic or dihedral symmetry is maintained by cloning the moves to separate monomers, and each move is accepted or rejected based on the standard Metropolis criterion (12). In the second, high-resolution refinement stage, side chains are built into the backbone conformations in all-atom detail, and small, symmetric perturbations are tested in the context of a reasonably accurate, high-resolution energy function that includes van der Waals interactions, the costs of desolvating atoms, and hydrogen bonds (Fig. 1) (10, 13). The lowest energy models are then clustered, and the five most populous clusters are compared with the experimentally determined structures.

We first tested the fold-and-dock protocol on a benchmark set of 27 protein complexes with structures previously determined by high-resolution crystallography. Most of these molecules form intertwined structures in which the number of intermolecular contacts approaches one third of the number of intramolecular contacts (Table 1). We have assumed that the stoichiometry of the crystallized complex is known (as can be rapidly ascertained experimentally by, for example, analytical ultracentrifugation), but tested all possible symmetries available to a fixed number of monomers. For example, tetramer complexes were modeled with both D2 and C4 symmetries.

Benchmark of Homo-Oligomeric Proteins Solved by X-Ray Crystallography. Initial tests were carried out on coiled coil structures with simple cyclic geometries as well as more complex dihedral symme-

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Table 1. Result of the fold-and-dock protocol on a benchmark of protein complexes solved by crystallography and NMR

PDB ID	Modeled residues [†]	Symmetry	Intermolecular contacts, %	rmsd oligomer prediction, Å ^{*§}	Cluster size [¶]	Lowest rmsd top 400, Å ^{*§}	rmsd monomer extracted from oligomer, Å	Traditional de novo single subunit rmsd, Å [§]
Crystal structure benchmark								
2mlt	26x4	D2	23	7.9	20	1.5	3.1 (1.2)	5.9
2zta	31x2	C2	17	1.6	251	0.6	1.4 (1.0)	10.8
1bdt	44x2	C2	24	12.9	20	0.8	7.0 (5.9)	6.2
2bti	44x2	C2	28	0.9	27	0.8	0.9 (1.5)	4.9
1wvp	47x2	C2	12	11.3	20	2.7	7.2 (7.6)	8.3
1g6u	48x2	C2	12	1.7	355	0.9	1.5 (1.9)	10.1
1irq	48x2	C2	27	11.9	10	3.3	5.6 (5.2)	10.6
2akf	32x3	C3	20	2.0	343	1.0	1.5 (1.6)	3.6
2odk	50x2	C2	15	12.5	10	3.6	11.8 (6.1)	5.9
2p64	51x2	C2	21	4.8	40	2.6	3.4 (3.6)	6.0
2h3r	54x2	C2	18	5.3	26	2.1	2.9 (3.3)	10.2
1rop	56x2	C2	14	1.4	40	0.8	1.1 (1.2)	0.8
1igu	59x2	C2	14	14.6	10	5.3	3.7 (6.5)	6.3
1zv1	59x2	C2	11	1.3	29	1.3	0.9 (1.8)	1.2
1g2z	31x4	D2	30	8.7	70	2.4	3.0 (2.8)	4.9
2fqm	63x2	C2	29	15.6	10	2.4	3.7 (6.3)	5.6
2or1	63x2	C1**	3	8.9	10	2.6	2.6 (1.7)	1.6
1c26	32x4	D2	29	2.7	6	1.8	1.5 (2.8)	5.6
1utx	66x2	C2	8	8.8	10	2.6	2.7 (3.2)	3.4
2dlb	69x2	C2	38	16.2	10	9.1	3.7 (14.1)	10.1
1mby	75x2	C2	15	15.5	10	7.0	3.7 (12.8)	12.6
2nzc	79x2	C2	8	13.5	10	4.5	3.9 (3.8)	2.9
2o1j	43x4	C4	24	1.2	343	1.0	1.8 (0.7)	16.4
1qx8	47x4	D2	30	1.7	56	1.0	1.4 (0.8)	17.8
1fe6	52x4	C4	23	2.4	212	1.9	1.1 (1.9)	20.4
1mz9	45x5	C5	28	1.3	132	1.1	1.7 (1.8)	7.7
1uis	64x6	C6	9	11.6	10	9.8	3.8 (9.9)	2.4
1i8f	71x7	C7	16	11.4	10	10.0	10.1 (7.5)	10.0
Crystal structure blind prediction ^{††}								
3bbp	29x2	C2	15	0.9 (13.7)	—	0.9	1.0 (3.6)	7.9
PDB ID ^{††}	Modeled residues [*]	Symmetry	Intermolecular contacts, %	rmsd, Å ^{*§§}		Lowest rmsd top 400, Å ^{*§§}	rmsd monomer, Å ^{*§§}	rmsd monomer prediction, Å
NMR benchmark (chemical-shift-assisted)								
2nwt (gr83)	38x2	C2	26.5	1.9 (9.3)	—	0.9	1.6 (6.0)	6.5
2bzb	48x2	C2	14.4	2.6 (1.7)	—	1.4	2.3 (1.3)	2.1
2js5 (mcr1)	63x2	C2	5.9	7.6 (18.6)	—	3.6	2.6 (2.9)	3.6
2rmm	56x2	C2	7.5	3.2 (1.7)	—	0.8	0.6 (1.3)	1.2
1ns1	73x2	C2	9.5	1.0 (1.4)	—	0.9	0.9 (2.3)	2.5
2b95 (hr2106)	96x2	C2	9.5	1.7 (2.0)	—	1.7	1.7 (1.7)	1.7
NMR blind predictions								
2k5j (sft1)	44x2	C2	24.3	1.3 (1.1)	—	0.9	1.0 (1.0)	3.9
2k7i (AtT3)	62x2	C2	24.6	5.3 ^{¶¶} (8.0)	—	2.7	1.7 (6.5)	6.6

^{*}Only residues in the regular secondary structures are used for rmsd calculation. For definition, *SI Materials and Methods*.

[†]The C and N termini were trimmed to remove flexible tails. The length refer to the number residues modeled in the simulation.

[‡]Values for the most accurate of the five models selected through clustering.

[§]Calculated over C α atoms over all residues.

[¶]Number of models in largest cluster, after clustering of 400 lowest energy complexes. This provides a measure of the extent of simulation convergence.

^{||}Values in parenthesis come from clustering the 400 lowest energy complexes with a single subunit extracted from the complex. The clustering method was identical to the one used for the full oligomer.

^{**}Forms dimer only in the presence of DNA (38). This protein was used as a reference in the benchmark and the results indicate that, in the absence of strong intermolecular interactions, the correct monomeric fold can be found in the folding-and-docking simulation.

^{††}Two competing orientations are observed: parallel and antiparallel. The largest clusters involve antiparallel coiled-coil orientations while biological data dictates the presence of a parallel orientation. The predicted model was the lowest energy parallel coiled-coil.

^{†††}NESG target ID in parentheses.

^{§§}Values for the model with lowest energy + chemical shift score. The values in parenthesis are from a clustering analysis using the same method as for the crystal benchmark.

^{¶¶}Convergence was not observed for this blind prediction. Predictions were selected based on clustering of decoys with lowest energy + chemical shift score. The third of the five submitted models was the most accurate and has a rmsd of 3.0 Å (Fig. 3E).

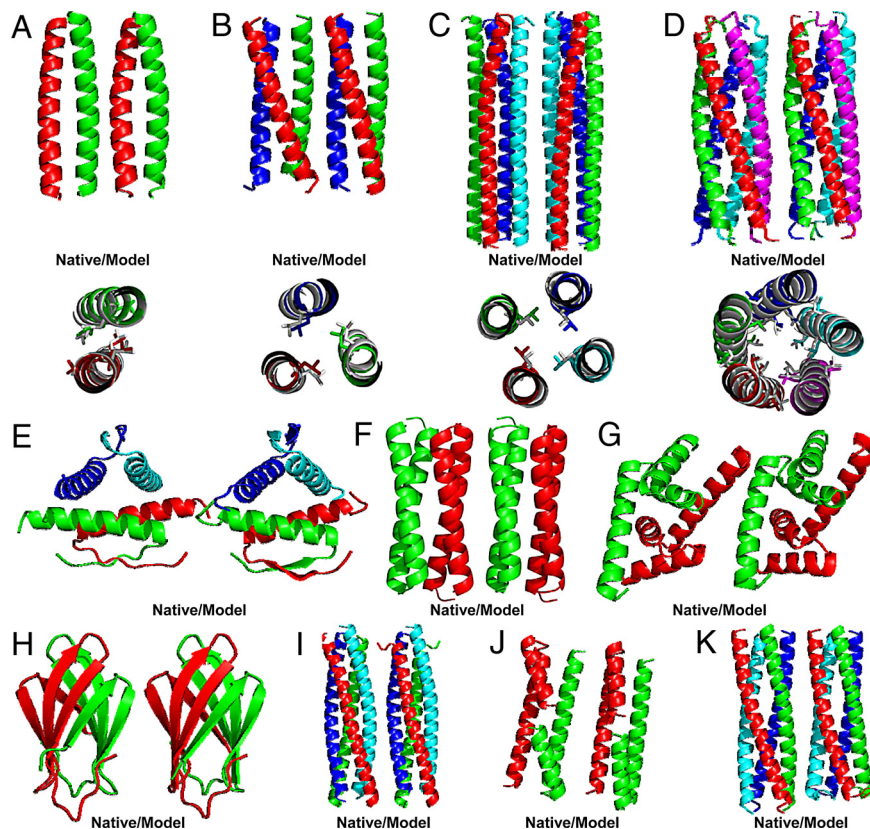


Fig. 2. High-resolution structure prediction using the fold-and-dock protocol. (A–I) Comparison between native crystal structure (*Left*) and most accurate of the five models selected by clustering of the low energy models (*Right*); PDB ID, 2zta (A); 2akf (B); 1fe6 (C); 1mz9 (D); 1c26 (E); 1rop (F); 1zv1 (G); 2bti (H); 2o1j (I); 1g6u (J); 1qx8 (K). For the cyclic coiled-coil structures (A–D), the packing of core leucine side chains is illustrated.

the two types of data, but these techniques require the preparation of mixed isotopically labeled samples, and typically yield only a few intermolecular constraints. To address this problem, we built on recent findings that NMR structure determination of monomers can be greatly accelerated by coupling *de novo* modeling with backbone chemical shift data without further NOE, scalar coupling, residual dipolar coupling (RDC), or isotope-edited measurements (24, 25). The structure of a dimeric complex was recently predicted from models of monomeric components generated using chemical shift fragments (26). We brought together the chemical-shift-derived fragment method with our symmetric folding-and-docking method to model eight complexes under investigation by the Northeast Structural Genomics consortium (NESG; Table 1). The set contains proteins with all- α , α/β , as well as all- β secondary structure in the range of 44 to 96 residues per monomer. Models were ranked based on a sum of the all-atom energy and a chemical shift score, which measures how well chemical shifts estimated from the models agree with the experimental values (25). The single model with lowest value of this combined score is taken as the prediction (Table 1 and Fig. 3; Fig. S2).

In six of the eight cases, the lowest energy Rosetta model had near-atomic accuracy ($<3\text{-}\text{\AA}$ C^α rmsd) (Fig. 3B–D; Fig. S2). The overall rate of success (75%) is higher than the crystal structure benchmark described above (41%), due to enhanced conformational sampling with higher quality chemical-shift-derived fragments. The most remarkable result is seen for the protein HR2106, a mixed α/β protein with a total of 192 residues, resulting in a 1.7- \AA model (Fig. 3D). This topologically complex protein is significantly larger than previous reported successes of chemical-shift-aided protein modeling (24, 25).

Blind Prediction of NMR Structures. For two of the NMR targets, blind predictions were made before NMR structures were released. First, significant convergence was not observed in the modeling of 2k7i, a complex of two 62-residue chains, suggesting that blind

prediction would be inaccurate (Fig. S2). Five models were instead selected based on a combination of the all-atom energy and the chemical shift score. The third of these models is quite accurate, giving an rmsd of 3.0 \AA (2.7–3.3 \AA over the 20 conformers) over secondary structure elements compared with the model obtained with the complete NMR data (Fig. 3E). We investigated whether this quite good model could be distinguished using experimentally measured residual dipolar coupling (RDC) data, which provide orientational information useful for validating NMR structures (27). The correlation between measured and back-calculated residual dipolar couplings (the quality(Q)-factor (28)) was determined for each of the five models, and the third model indeed fit the data the best (Q-factor of 1.06, 1.41, 0.55, 0.73, and 1.03, respectively, for these five models). Thus, additional data can make possible accurate structure determinations when convergence is too poor for the correct structure to be unambiguously identified (Fig. S3A).

The second blind NMR test case, for a slightly smaller complex, produced even more striking results. Modeling runs for the protein 2k5j, a complex of two 44-residue (number of structured residues) chains, gave outstanding convergence, with 52 of the 200 lowest energy models within 1.0- \AA C^α rmsd of each other, suggesting that the modeling had reached near-atomic accuracy. Indeed, when compared with 20 NMR conformers determined subsequently and independently using a full suite of 2136 NOE constraints (Fig. 3F), the lowest energy Rosetta model achieved an accuracy of 0.79–1.20 \AA over all modeled residues, and 0.79–1.09 \AA over the secondary structure elements. Despite making use of far less data, the Rosetta model achieves equally good agreement with RDC data compared with the model obtained with the traditional NMR method (Q-factors of 0.36 for the lowest 10 energy models versus 0.38 for 10 first conformers, respectively) (Fig. 3G; Fig. S3B). The RDC comparison suggests that even higher accuracy might be achievable if RDC data are used for fold-and-dock refinement rather than just validation (29); Rosetta models with quality factors as low as 0.24 are

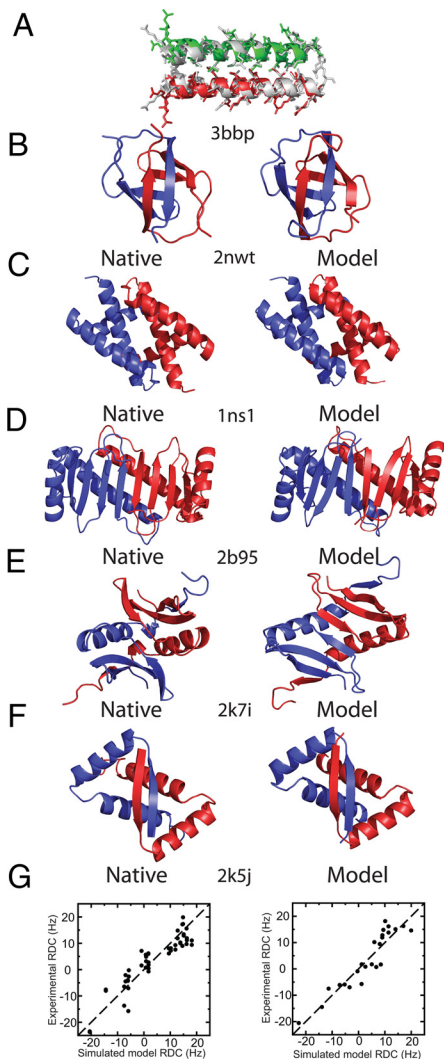


Fig. 3. High-resolution structure prediction using the fold-and-dock protocol. (A) Blind prediction of the structure of a Golgi-associated coiled coil region (PDB ID 3bbp). Red and green crystal structure, white blind prediction. (B–E) High-resolution structure prediction using the fold-and-dock protocol and chemical shift data. (B–D) Comparison between experimentally determined NMR structure (Left) and lowest energy Rosetta model (Right) for 2nwt (B), 1ns1 (C), and 2b95 (D). (E) Blind prediction of 2k7i. Experimentally determined structure (Left) and best blind prediction (Right). (F) Blind prediction of 2k5j. Experimentally determined structure (Left) and best blind prediction (Right). (G) Comparison between experimentally determined RDCs and simulated values for the first conformer in the NMR-ensemble of 2k5j refined without using NOE-data (Left) and best Rosetta model (Right).

present in the generated pool of models (Fig. S3B). Based on these and previous results, chemical-shift-based Rosetta has become an integral part of the NESG high-throughput structure determination pipeline for multimers (30). A third blind prediction of a dimeric structure was validated as this paper went to press. The predicted (lowest energy) structure, a non-intertwined monomer, had a backbone rmsd of 0.8 Å to the subsequently determined crystal structure. This illustrates that the method can produce excellent models for non-intertwined as well as intertwined oligomers—whether the lowest energy structure is intertwined or not is a function of the amino acid sequence, and need not be specified in advance.

Conclusion

We have demonstrated that simultaneous prediction of the fold and docking arrangement for protein complexes with many

different symmetries and modest monomer sizes is feasible at high resolution. Further applications of the presented method can be envisioned in conjunction with experimental techniques beyond crystallography and NMR spectroscopy. First, large symmetric complexes are common targets of cryoelectron microscopy. Attaining high-resolution models from these low-resolution data may be possible by incorporating density maps into the conformational search. Second, oligomeric membrane complexes, which have been difficult to characterize experimentally, are attractive targets for the method. Many physiologically important channels and transporters are in a suitable size range for the protocol described here, and chemical cross-linking and accessibility data may provide rapid validation of high-resolution models. Last, the techniques described here, when coupled to existing design algorithms, provide a computational foundation for rationally engineering proteins that self-assemble into symmetric containers, fibrils, and channels of new and useful function.

Materials and Methods

The computational protocol developed in this work builds on previously described methods for de novo (5) and symmetrical protein assembly (12) structure prediction in Rosetta.

Description of Symmetry. The computational setup used previously in Rosetta to dock already folded monomeric structures into symmetric homo-oligomers has been described previously (12); in this previous work the internal backbone coordinates of each monomer were held fixed. This approach works for non-interleaved homo-oligomers but in general will fail for interleaved structures as the structure of the monomer depends strongly on interactions with other subunits. Here, we have retained the basic framework for modeling symmetric structures while adding in full backbone flexibility to allow monomers to fold up in the context of symmetrically related copies of themselves. During the simulations, backbone, side chain, and rigid body degrees of freedom are all varied. Backbone symmetry is enforced by maintaining identical bond angles, bond lengths, and torsion angles in symmetry-related partners in the system. Side chains are placed using two different methods; combinatorial and one-at-a-time noncombinatorial optimization (31, 32) using a backbone-dependent rotamer library (33). Rotamers are simultaneously inserted into all symmetry related partners of the system, and the energy of the full symmetrical oligomer is evaluated (12, 32). The most straightforward manner to describe rigid body symmetry is to maintain a global reference coordinate system and use symmetry transforms to find the coordinates of symmetry-related partners in the system. This system leads to difficulties in describing complex symmetries and to perform gradient based minimization. Instead, in Rosetta, each symmetry-related partner maintains its own reference coordinate systems that are in themselves related by symmetry. In this setup, the symmetry-related partners have identical coordinates, but in their own reference frames. During the simulation protocol, only rigid body degrees of freedom that maintain symmetry are allowed to vary. The nature of these perturbations depends on the type of symmetry that is simulated. In this work, two types of symmetries are used, cyclic and dihedral symmetry; complete descriptions of the samples degrees of freedom are given in Fig. S4.

Subsystems. For larger oligomers, it is unnecessary to explicitly simulate the whole systems, because a smaller subsystem can describe all of the interactions in the system if interactions at very long distances are ignored. The minimal number of monomers that explicitly needs to be simulated is equal to the number of different interaction surfaces between subunits in the system. To avoid edge effects, the smallest subsystem is a single monomer with all directly neighboring subunits present, and the total energy is calculated as a multiple of the energy of the central monomer.

Energy Function. The energy function in the low-resolution search is a linear combination of mostly knowledge-based terms modeling residue-environment and residue-residue interactions, secondary structure packing, chain density, and excluded volume (34). The high-resolution energy function is composed of a Lennard-Jones potential to model side-chain packing, an orientation-dependent hydrogen bond term parameterized from quantum mechanics (35) and analysis of high-resolution structures (36), the Lazaridis-Karplus implicit solvation model (37), pair-interaction terms modeling long range electrostatics, a side-chain torsional potential derived from the Dunbrack backbone-dependent rotamer library (33), and backbone torsional potential dependent on secondary structure

and amino acid type. Detailed description of the energy terms can be found in Rohl et al. (13) and in supporting information in Kuhlman et al. (38).

Conformational Search Protocol. The initial state of the simulated system is generated with random rigid body degrees of freedom and the subunit backbones in extended conformations. The protocol starts with a low-resolution symmetrical fragment insertion where torsion angles of randomly selected 3 or 9 residue fragments in the protein chains are replaced with torsion angles from nonhomologous proteins with known structure (6); 40,000 fragment insertions attempts are made. At every 1 in 10 fragment insertions, the rigid body of the protein chains are adjusted using two type of moves: random perturbations (translation/rotation) and sliding of subunits into atomic contact (translation). In the random perturbation step, the subunits are randomly and symmetrically rotated or translated up to 5° or 0.5 Å, respectively. In the sliding step, the subunits are translated into atomic contact along the symmetry axis of the system. In the case of dihedral symmetry, the subunits are consecutively translated into atomic contact along the different symmetry axis in a randomly selected order. Fragment and rigid body moves are accepted based on the Metropolis criteria.

The models produced by the low-resolution protocol are energy minimized using an all-atom Monte Carlo Minimization protocol as previously described (5). A number of different symmetrical moves are used in the refinement: small random perturbation of backbone torsion angles, one-at-a-time (noncombinatorial) rotamer optimization using a backbone-dependent rotamer library (33), gradient-based minimization of the backbone, and side chain degrees of freedom and insertion of nonperturbing three or nine residue fragments. In addition to these moves, random rigid body perturbations were also incorporated into the protocol.

The 2×10^4 to 5×10^6 models were generated using Rosetta@home, and the 400 lowest energy models were clustered based on rms similarity to select the best models. As in previous work (6), modeling runs that did not pass 50th percentile energy cuts half-way and three-quarters of the way through the simulation were terminated. A clustering threshold of 2 Å was used, and five models representing the center of the five largest clusters were selected as predictions. NMR models, which can make use of additional limited experimental information, were selected as described below.

To compare the energies of models generated by the de novo protocol with

native structures, the native structures were subjected to all-atom refinement. The native structures were first idealized by replacing the bond angles and bond lengths with the “ideal” values used in the de novo protocol (using a quasi-Newton optimization to adjust the backbone torsion angles to minimize the overall perturbation to the 3D structure) to facilitate comparison.

NMR Structure Inference. For NMR targets, the standard protein fragment selection procedure in Rosetta was replaced with the chemical shift filtered fragment selection used in CS-Rosetta (25). In the CS-Rosetta method, the software SPARTA (39) is used to predict the backbone and C^β chemical shifts of a database protein structures for which high-resolution crystal structures are available. Three and nine residue fragments are selected from this structural database based on similarity between predicted chemical shifts and experimentally determined chemical shifts of the target protein (for further details, see ref. 25). The standard fold-and-dock protocol is then used to generate high-resolution models of homo-oligomers.

A single model is predicted for each target protein by selecting the model with lowest Rosetta energy adjusted with a term measuring the similarity between the predicted chemical shift of the model and the experimentally determined shifts (25).

Availability. The code is freely available to academic users at www.rosetta-commons.org in release Rosetta 2.3.0. For command line, see *SI Materials and Methods*.

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