



# High-resolution structure prediction of $\beta$ -barrel membrane proteins

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**$\beta$ -Barrel membrane proteins ( $\beta$ MPs) play important roles, but knowledge of their structures is limited. We have developed a method to predict their 3D structures. We predict strand registers and construct transmembrane (TM) domains of  $\beta$ MPs accurately, including proteins for which no prediction has been attempted before. Our method also accurately predicts structures from protein families with a limited number of sequences and proteins with novel folds. An average main-chain rmsd of 3.48 Å is achieved between predicted and experimentally resolved structures of TM domains, which is a significant improvement ( $>3$  Å) over a recent study. For  $\beta$ MPs with NMR structures, the deviation between predictions and experimentally solved structures is similar to the difference among the NMR structures, indicating excellent prediction accuracy. Moreover, we can now accurately model the extended  $\beta$ -barrels and loops in non-TM domains, increasing the overall coverage of structure prediction by  $>30\%$ . Our method is general and can be applied to genome-wide structural prediction of  $\beta$ MPs.**

structure prediction |  $\beta$ -barrel membrane proteins | strand register | Covariation | loop prediction

The outer membrane proteins are found in the gram-negative bacteria, mitochondria, and chloroplast (1). They form  $\beta$ -barrels, so are also known as  $\beta$ -barrel membrane proteins ( $\beta$ MPs).  $\beta$ MPs are involved in outer membrane biogenesis, membrane anchoring, pore formation, translocation of virulence factors, and enzyme activities (2–5). Recent progress in engineering protein nanopores using  $\beta$ MPs for protein profiling (6–8), DNA sequencing (9, 10), small molecule detection (11), and targeted drug delivery for cancer therapy (12) has increased the significance of understanding the organizing principles of  $\beta$ MPs.

A major obstacle in studies of  $\beta$ MPs is the limited availability of structural data. Only  $\sim 320$   $\beta$ MP structures, of which  $\sim 59$  are nonhomologous, have been deposited in the Protein Data Bank (PDB) that contains  $>135,000$  protein structures (13). Computational studies have contributed to expand our knowledge of  $\beta$ MPs by successfully predicting  $\beta$ MP sequences at a genome-wide scale (14, 15), identifying transmembrane (TM) segments (16, 17) and uncovering sequence and spatial motifs (18, 19). The stability, oligomerization state, protein–protein interaction interfaces, and the transfer free energy of residues in the TM regions of  $\beta$ MPs can also be accurately computed (20–26).

Template-based methods for structure prediction have been successfully applied in studies of globular proteins (27). They have also been used to predict 3D structures of  $\beta$ MPs but have achieved limited success with novel folds like the ones found in VDAC, FimD, PapC, and LptD proteins (28) due to the limited availability of templates for  $\beta$ MPs. General purpose template-free structure prediction methods do not generate accurate structures of  $\beta$ MPs, as these proteins can be large, with the number of residues reaching 800.

A recently published  $\beta$ MP-specific method that combines sequence covariation for contact prediction with a machine-learning-based method achieved limited progress, with a main-chain rmsd of 6.66 Å for predicted structures of TM regions,

before it was adjusted to a better published value of 4.45 Å when only a subset of residues were aligned instead of all TM residues (29). Another template-free  $\beta$ MP-specific method, 3D-SPoT (3D structure predictor of transmembrane  $\beta$ -barrels), can predict the TM regions of  $\beta$ MPs with an average main-chain rmsd of 4.14 Å (30). Despite such progress, further improvement in prediction methods to generate accurate structural models is required to bridge the gap between identified  $\beta$ MP sequences and resolved  $\beta$ MP structures, so that modeled structures can be used directly for applications such as nanopore engineering and drug design/delivery.

In this study, we describe a template-free method for predicting 3D structures of  $\beta$ MPs, which provides significant improvement over previous methods. Our approach, named 3D  $\beta$ -barrel membrane protein predictor (3D-BMPP), is based on a statistical mechanical model (31) that incorporates sequence covariation information and is built upon a parametric structural model of intertwined zigzag coils. In a blind test of 51 nonhomologous  $\beta$ MPs, our prediction generates accurate 3D structures of TM regions with an average main-chain rmsd of 3.48 Å. This represents a significant improvement of  $\sim 3.1$  Å compared with a recent study (29) over a much bigger dataset (51 proteins vs. 17 proteins). In addition, predictions are expanded to include non-TM regions, including both extended  $\beta$ -sheets and loops, resulting in significant increase in the coverage of residues compared with previous methods. Furthermore, our method can be applied to model structures of  $\beta$ MPs with novel folds, including those from mitochondria of eukaryotes, as evidenced by the accurately modeled structures of VDAC and FimD. Our method is general and can be applied to genome-wide structural prediction of  $\beta$ MPs.

## Significance

**$\beta$ -Barrel membrane proteins ( $\beta$ MPs) are drawing increasing attention because of their promising potential in bionanotechnology. However, their structures are notoriously hard to determine experimentally. Here we develop a method to achieve accurate prediction of  $\beta$ MP structures, including those for which no prediction has been attempted before. The method is general and can be applied to genome-wide structural prediction of  $\beta$ MPs, which will enable research into bionanotechnology and drugability of  $\beta$ MPs.**

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The authors declare no conflict of interest.

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**Table 1. Comparison of different methods for strand register and 3D structure prediction for TM regions of  $\beta$ MPPs**

Method	No. $\beta$ MPPs	No. strands	Strand register accuracy, %	Average main-chain TM-rmsd, Å	Average all-atom TM-rmsd, Å
Jackups and Liang (31)	19	256	46	—	—
TMBpro-server (28)	14	214	48	—	7.3
3D-SPoT (30)	23	324	73	4.12	5.6
EVfold_bb (29)	17	265	44	6.66	—
3D-BMPP (this study)	51	771	85	3.48	4.26

3D-BMPP can predict strand registers with an accuracy of  $\sim 85\%$  and 3D structures of TM regions with an average main-chain rmsd of 3.48 Å and average all-atom rmsd of 4.26 Å for a much bigger dataset (51  $\beta$ MPPs vs. 14–23  $\beta$ MPPs).

interactions. Incorrect strand register can lead to erroneous side-chain orientation prediction. The correct prediction of strand register is therefore an important requirement in structure prediction of  $\beta$ MPPs and is well recognized in the literature (28). Our method can predict strand register at 85% accuracy. In contrast, the criteria were relaxed to allow +1 or  $-1$  difference in strand register in a previous study (29). While this relaxation made the register prediction results more presentable (65% after relaxation vs. 44% before relaxation), it is problematic, as it would lead to prediction of TM residues to adopt erroneous orientation opposite to that of the native structures. Such incorrect TM residue orientations would imply completely different properties of the barrel interior and exterior. Here we report correct prediction only when we are able to exactly match the register with the experimentally resolved structure.

**Predicting side-chain orientations.** We use the reduced state space and a single body potential (20) calculated from the updated dataset to predict the side-chain orientation of each strand. Since the side-chain orientations of a strand follow an alternative lipid-facing–pore-facing pattern, only the orientation of the first residue of each strand needs to be predicted. The accuracy of our prediction is 98% (see *SI Appendix, section 3.4* for details).

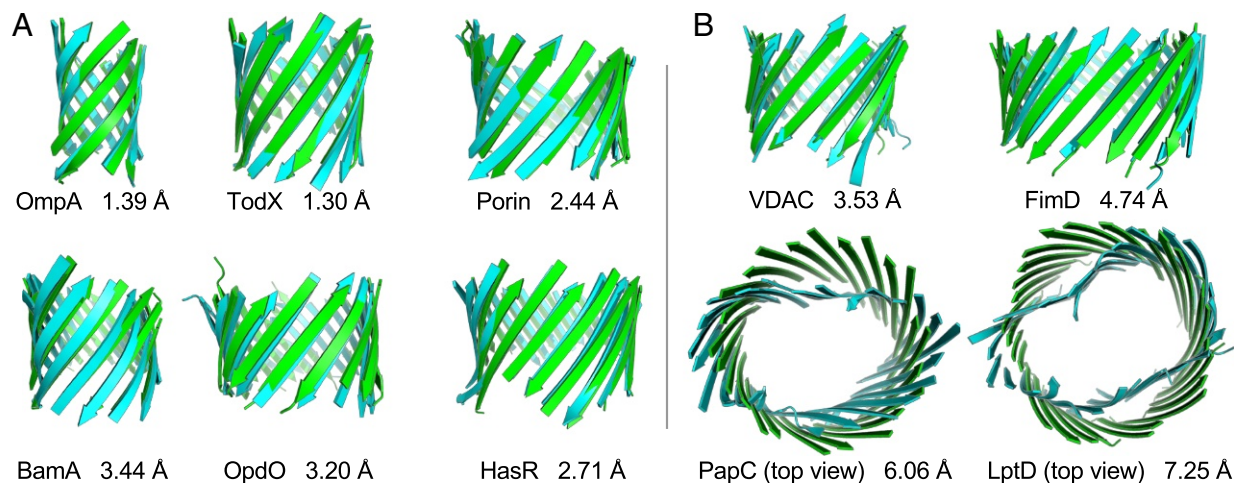
**Optimizing protein shear.** We next optimize the shear number which characterizes the global hydrogen bond pattern of a  $\beta$ MPP. The shear number is the displacement of the relative positions in

the TM strands if one starts to follow the strong hydrogen bond or side-chain interaction between strands, beginning from one strand and returning after a full circle to the same strand (*SI Appendix, Fig. S5*). The predicted shear number of a  $\beta$ MPP can be calculated as the sum of the predicted strand registers.

In the step of register prediction, we keep the register with the lowest score and the one with the second lowest score as candidates for each strand pair. They are then evaluated against the predicted side-chain orientations of the strand, based on the fact that residues sharing a strong hydrogen bond or side-chain interaction have the same side-chain orientation. One of the two registers is then selected so that the predicted shear number is as close as possible to the most common shear number of the  $\beta$ MPPs of the same strand number (*SI Appendix, Table S3*), while keeping the sum of the strand register scores as small as possible (see *SI Appendix, section 3.5* for technical details). After optimization, the error in predicted shear numbers is decreased from  $-0.69 \pm 3.63$  to  $0.12 \pm 1.34$ . The improved global shear accuracy will lead to overall more accurate 3D structure prediction of  $\beta$ MPPs.

### Predicting 3D Structures of TM Regions of $\beta$ MPPs

**Parametric model for 3D structures of the TM regions.** Parametric models have had recent successes in modeling and designing structures of  $\alpha$ -helical proteins (36, 37). We have developed a parametric structural model, named the intertwined zigzag coil model, to generate 3D structures of  $\beta$ MPPs from predicted strand registers (*SI Appendix, Fig. S4*). Following previous studies (30,



**Fig. 2.** Structure prediction of TM regions. (A) Predicted structures of the TM regions (green) superimposed on experimentally determined structures (cyan): OmpA (1bxw), TodX (3bs0), Porin (1prn), BamA (4n75), OpdO (3szv), and HasR (3csl). (B) Predicted structures of the TM regions of proteins with novel folds (green) superimposed on experimentally determined structures (cyan): VDAC (3emn), FimD (3rfz), PapC (2vqi), and LptD (4q35). PapC and LptD are shown in top view.

**Table 2. Flexibility of TM regions of  $\beta$ MPs and the accuracy of the prediction of 3D-BMPP**

PDB ID	$D_{\text{nmr,nmr}}^{\text{TM}}$	$D_{\text{nmr,X-ray}}^{\text{TM}}$	$D_{\text{pred,nmr}}^{\text{TM}}$	$D_{\text{pred,X-ray}}^{\text{TM}}$
1bxw	1.41 ± 0.42	1.99 ± 0.31	1.83 ± 0.15	1.36
1qj8	2.50 ± 0.74	2.48 ± 0.80	3.11 ± 0.46	2.65
1thq	1.99 ± 0.58	4.53 ± 0.38	5.30 ± 0.42	3.32
2f1c	2.42 ± 0.37	2.80 ± 0.21	3.93 ± 0.21	3.06
2f1t	2.13 ± 0.35	4.30 ± 0.11	4.08 ± 0.14	3.12
2lhf	0.82 ± 0.22	No X-ray	1.60 ± 0.08	1.48*
2mlh	1.48 ± 0.28	No X-ray	1.49 ± 0.14	1.44*
Mean	2.11 ± 0.79	3.18 ± 1.16	3.09 ± 1.39	2.35 ± 0.82

$D_{s_1,s_2}^{\text{TM}}$  is the average of the mutual  $C_{\alpha}$ -rmsd between structures  $s_1$  and  $s_2$ .

\*As no X-ray structures for these proteins are available, we used the first model of the NMR data.

38), we model the overall shape of the  $\beta$ -barrel as an ideal cylinder. The  $C_{\alpha}$  trace of each strand is described as a coiled zigzag wrapping around the hypothetical cylinder (see *SI Appendix, section 4.1* for details). This model captures the zigzag nature of a polypeptide in the  $\beta$ MP and the varied distance between  $C_{\alpha}$  atoms on adjacent strands (*SI Appendix, Fig. S3*), which improves positioning of  $C_{\alpha}$  atoms.

**Predicting 3D atomic structures.** We then use these  $C_{\alpha}$  atoms to construct the main-chain atoms using Gront et al.'s (39) algorithm. The side-chain atoms are then added using side-chains with a rotamer library 4 (Scwrl4) (40).

Fig. 2A depicts the predicted structures (green) of the TM regions of proteins OmpA, TodX, Porin, BamA, OpdO, and HasR, which are shown superimposed on experimentally determined structures (cyan). The rmsds of the main-chain atoms between the computed and experimentally resolved structures are 1.39 Å, 1.30 Å, 2.44 Å, 3.44 Å, 3.20 Å, and 2.71 Å for OmpA, TodX, Porin, BamA, OpdO, and HasR, respectively. The structures of the TM regions of 51  $\beta$ MPs are predicted with an average rmsd of 3.48 Å for main-chain atoms and 4.26 Å for all atoms (see *SI Appendix, Table S4 and Fig. S7* for details). The accuracy of predicted structures is maintained for large proteins such as Iron(III) dicitrate transport protein FecA protein (237 TM residues). This is in contrast to other prediction methods, where there is considerable deterioration in the quality of predicted structures (*SI Appendix, Table S5 and Fig. S6*). The average TM scores of our predicted structures also compare favorably with those of a recent study (0.73 vs. 0.54) (29). Furthermore, our results are over a much bigger dataset (51 proteins vs. 17 proteins). Thus, these results represent a very significant improvement. Moreover, the parametric structural model of intertwined zigzag coils improves accuracy of side chains, as the all-atom rmsd has improved by more than 1.30 Å (4.26 Å vs. 5.60 Å) compared with a previous study (30).

TM regions of  $\beta$ MPs have considerable intrinsic flexibility: The NMR structures have an average mutual  $C_{\alpha}$ -rmsd of  $2.11 \pm 0.79$  Å for the seven  $\beta$ MPs with known NMR data (Table 2, column 2). The difference between the NMR and X-ray structures is more pronounced, with an average  $C_{\alpha}$ -rmsd of  $3.18 \pm 1.16$  Å (Table 2, column 3). In contrast, the average  $C_{\alpha}$ -rmsds of our predicted structures against NMR and X-ray structures are  $3.09 \pm 1.39$  and  $2.35 \pm 0.82$ , respectively (Table 2, columns 4 and 5). These differences are similar to the structural differences originating from the intrinsic flexibility of the proteins, suggesting that our prediction of TM regions of  $\beta$ MPs has excellent accuracy comparable to NMR structures.

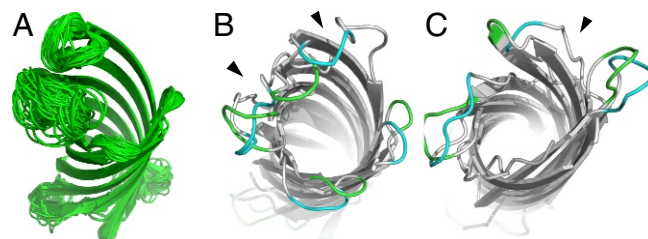
**Predicting structures of  $\beta$ MPs with novel folds.** It is challenging to predict the structures of  $\beta$ MPs with novel folds.  $\beta$ MPs were

considered to have even numbers of strands from 8 to 22 (41). A  $\beta$ MP is considered to have a novel fold when its number of strands has not been observed in other experimentally determined structures. For example, VDAC in mitochondria has an odd number (19) of strands (42); PapC, FimD, and LptD all have more than 22 strands (24, 24, and 26, respectively). Predicting structures of a number of  $\beta$ MPs including VDAC, FimD, and LptD with reasonable accuracy was not possible in a recent study (29), likely due to inaccurate residue contact predictions and limitations in the machine-learning-based procedure. Template-based prediction methods either fail to build any model or generate very poor structures. With the improved modeling procedure of 3D-BMPP, we are able to model the TM regions of the VDAC, FimD, PapC, and LptD proteins with a main-chain rmsd of 3.53 Å, 4.74 Å, 6.06 Å, and 7.25 Å, respectively (Fig. 2B). While the structure of VDAC was previously predicted with an accuracy of 3.9 Å (30) and 7.41 Å (29), to the best of our knowledge the structures of FimD, PapC, and LptD have not been successfully predicted before this study. The large rmsds of predicted structures of PapC and LptD show that our current idealized cylindrical structural model cannot yet model deformed barrels effectively.

### Predicting Structure of non-TM Regions of $\beta$ MPs

**Predicting structures of extended  $\beta$ -sheets.** We also model the structures of the non-TM regions of  $\beta$ MPs, including the extended  $\beta$ -sheets (extended barrels) and loops connecting adjacent strands. The extended barrels have overall similar structures to those of the TM barrels. Including the extended barrel in our prediction increases the coverage of the modeled structures by 20% when measured by the average number of residues modeled in the 51 structures (159 in TM regions vs. 191 in whole-barrel regions, with the largest modeled barrel structure containing 350 residues), with little deterioration in the average main-chain rmsd (3.48 Å vs. 3.80 Å).

**Predicting structures of loops.** Loops are the most flexible regions of  $\beta$ MPs and are important for their functions (43). NMR structures of  $\beta$ MPs show that these loops adopt multiple conformations (44, 45), which likely contribute to the challenges in predicting binding affinity of  $\beta$ MP–ligand interactions (46). We model loops by investigating a large ensemble of loop conformations generated using an improved version of the multi-loop distance-guided sequential chain-growth Monte Carlo (m-DiSGro) algorithm (47) that guarantees clash-free conformations of the sampled loops. For each of the seven  $\beta$ MPs with available NMR structures, once the structure of the barrel domain is predicted, we sample  $3 \times 10^4$ – $3 \times 10^5$  multiloop conformations, with the specific number of conformations dictated by the number and the lengths of loops. We then perform clustering to generate an ensemble of  $\sim 400$  multiloop conformations as a prediction



**Fig. 3.** Structure prediction of loop regions. (A) Ensemble of predicted loop structures of OmpX (1qj8). (B and C) Examples of predicted loops on the extracellular side (B, green) and on the periplasmic side (C, green) superimposed on the corresponding NMR structure (cyan) (49). The black arrowheads indicate the big fluctuations in the barrel region.

**Table 3. Comparison of the accuracy of loop prediction for  $\beta$ MPs**

PDB id	$D_{\text{nmr,nmr}}^{\text{barrel}}$	$\Delta D_{\text{nmr,nmr}}^{\text{loop}}$	$D_{\text{nmr,pred}}^{\text{barrel}}$	$\Delta D_{\text{nmr,pred}}^{\text{loop}}$
1bxw	2.78 ± 0.72	3.83 ± 1.25	3.35 ± 0.51	3.00 ± 0.55
1qj8	3.31 ± 0.80	0.61 ± 0.26	4.14 ± 0.57	0.67 ± 0.27
1thq	1.99 ± 0.58	0.79 ± 0.35	5.30 ± 0.42	0.52 ± 0.21
2f1c	3.33 ± 0.61	3.76 ± 0.94	5.29 ± 0.50	2.78 ± 0.48
2f1t	2.58 ± 0.54	1.01 ± 0.55	4.35 ± 0.15	0.45 ± 0.20
2lhf	0.85 ± 0.24	1.94 ± 0.60	1.63 ± 0.09	2.05 ± 0.27
2mlh	1.48 ± 0.28	1.51 ± 0.64	1.49 ± 0.14	0.99 ± 0.26
Mean	3.65 ± 1.21	1.03 ± 0.89	3.64 ± 1.46	1.12 ± 0.89

We are able to sample most of the loop conformations seen in the NMR structures with <3 Å deterioration in  $C_{\alpha}$ -rmsd.

for each protein. The predicted loop conformations are diverse (Fig. 3A) and represent the broad conformational space that is accessible to loops (48). Examples of predicted loops are shown in Fig. 3.

To assess the quality of the predicted loop conformations, we define a metric  $\Delta D_{s_1,s_2}^{\text{loop}}$  that measures how  $C_{\alpha}$ -rmsd between structures  $s_1$  and  $s_2$  is changed upon incorporation of the loop regions:  $\Delta D_{s_1,s_2}^{\text{loop}} = D_{s_1,s_2}^{\text{whole}} - D_{s_1,s_2}^{\text{barrel}}$ , where  $D_{s_1,s_2}^{\text{whole}}$  is the  $C_{\alpha}$ -rmsd between the structures  $s_1$  and  $s_2$  including both the barrel and loop regions, and  $D_{s_1,s_2}^{\text{barrel}}$  is the  $C_{\alpha}$ -rmsd between the barrel domains only. Since the number  $M$  of available NMR structures for each protein is limited compared with our predictions (~10–20 vs. ~400), we selected  $M$  predicted conformations closest to the NMR structures by  $\Delta D_{\text{nmr,pred}}^{\text{loop}}$  from the modeled ensemble for each protein. The resulting  $\Delta D_{\text{nmr,pred}}^{\text{loop}}$ s calculated using these structures are <3 Å, with an average of  $1.12 \pm 0.89$  (Table 3, column 5), which is on par with the values of  $\Delta D_{\text{nmr,nmr}}^{\text{loop}}$  (Table 3, column 3), suggesting that we are able to sample the loop conformations observed in the NMR structures accurately.

## Discussion

Due to the difficulties in experimental determination of membrane protein structures, there are a limited number of structures of nonhomologous  $\beta$ MPs. However, it is estimated that there are 15,000  $\beta$ MPs across 600 different gram-negative chromosomes (50). Computational modeling has the promise to provide working 3D models for these sequences, enabling novel applications in nanopore engineering and drug design/delivery, as well as further understanding of the structural basis of the function and mechanism of these  $\beta$ MPs. We have developed a method for predicting structures of  $\beta$ MPs, which combines a statistical mechanical model, sequence covariation information, and global register optimization with a parametric structural model of intertwined zigzag coils. The results show that we can accurately predict structures of  $\beta$ MPs with a significantly expanded coverage of extended  $\beta$ -sheets and loops.

The incorporation of global register optimization increases the accuracy of the predicted structures by 0.24 Å on average, suggesting that the global hydrogen bond network cannot be approximated accurately using local strand register alone. As an example, for the  $\beta$ MPs OmpA (PDB ID: 1bxw), hypothetical protein HB27 (PDB ID: 3dzm), and PagL (PDB ID: 2erv), the strand registers were predicted correctly for six of eight strands before global register optimization, with an error in shear number of –4, –6, and –6, respectively. After global register optimization, the strand register was predicted correctly for eight, six, four strands, respectively, and the error in shear number becomes 0 in all three cases. Moreover, the main-chain rmsd of these predicted structures is improved by 2.7 Å, 2.5 Å, and 1.5 Å, respectively.

Our parametric model of intertwined zigzag coils captures the zigzag nature of a polypeptide and the varied distance between  $C_{\alpha}$  atoms of two adjacent strands, which depends on whether the respective residues share a main-chain hydrogen bond. This results in significant improvement in rmsd for all atoms in general and side-chain atoms in particular. When we constructed structures of all 51  $\beta$ MPs using our parametric model with true registers, the average main-chain rmsd of these structures was 2.5 Å. Given our prediction accuracy of 3.48 Å in this study, only ~1 Å error on average is due to incorrect register prediction, while the 2.5-Å error is due to the structural deviation of  $\beta$ MPs from the ideal cylindrical shape.

Currently this ideal cylindrical model cannot capture ellipticity, twist, and curvature of local surface of the deformed barrel domains such as those observed in PapC and LptD (Fig. 2B), and alternative hyperboloid models have been discussed in the literature (51, 52). However, as current understanding of the physical factors determining these geometric properties is incomplete, further investigation of the heterogeneity of interactions in the TM region is required to develop a more accurate geometric model that can account for the deformed barrel domain.

In a recent study, structures for only 17 proteins (compared with 51 proteins in this study) were predicted (29), as the number of sequences available for the remaining proteins was insufficient to analyze sequence covariation. Here, we show that this limitation can be removed by combining patterns of hydrogen bond and side-chain interactions derived from experimentally determined 3D structures with the sequence covariation information (SI Appendix, Fig. S8). Our method predicts the 3D structures of 51  $\beta$ MPs with an average rmsd of 3.48 Å, which compares favorably with the recent study that has an average rmsd of 6.66 Å (29). Detailed technical issues comparing the two methods are discussed in SI Appendix, section 6.

Our method revealed basic organizational principles of  $\beta$ MPs and requires no template structures. In addition, TM regions of  $\beta$ MPs with a novel fold can also be modeled effectively, as evidenced by the predicted structures of VDAC and FimD. Furthermore, non-TM regions including both extended  $\beta$ -sheets and loops can be predicted accurately. Overall, our method opens the possibility of structural studies of many  $\beta$ MPs, including those in eukaryotic mitochondria and chloroplasts.

## Materials and Methods

We use 59  $\beta$ MPs with known structures as our dataset. The mutual sequence similarity is below 30%. Predictions are made only for 51  $\beta$ MPs, after excluding multichain  $\beta$ -barrels to avoid overestimation of repeated interaction types. Leave-one-out cross-validation is performed to assess the accuracy of the predictions.

Here, we describe our methods briefly. More details of the methods can be found in SI Appendix, sections 2–5. We take the canonical model of TM strands based on the physical interactions between strands described in refs. 31 and 33. The energetic contributions incorporate interactions with adjacent strands, interstrand loop entropy, a penalty for left-handedness, and sequence covariation. For each pair of adjacent strands, we enumerate all possible registers in a reduced conformational space and predict the registers. This is followed by the global shear optimization. We use a parametric structural model of intertwined zigzag coils to calculate the positions of  $C_{\alpha}$  atoms. Main-chain atoms and side chains are added using Gront et al.'s (39) algorithm and Scwrl4 (40). We then use an improved version of the m-DiSGro algorithm (47) to sample loop ensembles.

The 3D-BMPP code and the corresponding data are available at [sts.bioe.uic.edu/3dbmpps/](https://sts.bioe.uic.edu/3dbmpps/).

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