Van der Waals packing facilitates membrane protein association

Abstract

Visual Abstract

Design algorithm, experiment, fluorescence correlation

Introduction

Membrane proteins comprise 25-30% of the proteins found within protein-coding genes of various organisms (Fagerberg et al., 2010). Proper membrane protein folding is critical for essential biological functions, including cell signaling, ion balance, and gene regulation. Misfolding of membrane proteins has been found to be involved in several human diseases such as Parkinson’s, cystic fibrosis, and cancer (Sanders and Myers, 2004; Gregersen et al., 2006).

To understand how protein misfolding plays a role in disease states and progression, it is necessary to investigate how these proteins fold. However, studying membrane protein folding is inherently a difficult challenge because of their hydrophobic nature. Membrane proteins are difficult to express in yields high enough for biophysical experiments, and purification and solubilization of these proteins often lead to aggregation or unfolding (Carpenter et al., 2008). To combat these challenges, much of the research studying membrane protein folding is focused on understanding the biophysical forces that govern the folding process. Investigation of the biophysical forces that govern protein folding will allow us to determine why specific mutations result in diseases caused by misfolding. In addition, this knowledge can be applied to design new therapeutics that specifically target proteins in these misfolded states. Understanding the forces that enable membrane proteins to fold will contribute to our knowledge of how these fundamental forces are involved in human health and disease.

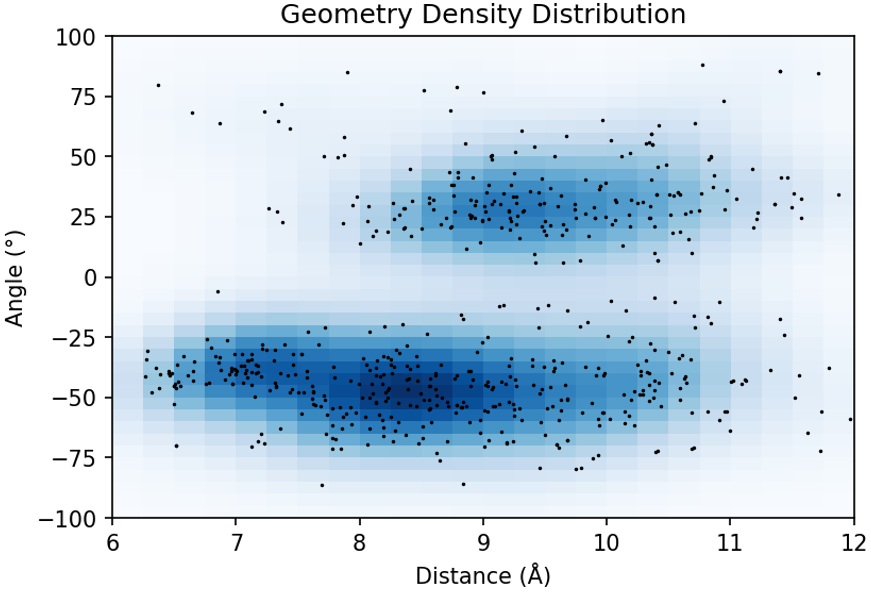
Proper membrane protein folding is regulated by a distribution of stabilizing hydrogen bonds, weak polar interactions, and van der Waals forces between the unfolded and folded states. Previous research has measured the contributions of both hydrogen bonding and weak polar interactions in the membrane and determined that these forces can drive membrane protein folding (Zhou et al., 2001; Yano et al., 2002; Johnson et al., 2007), but research is lacking on the contribution of van der Waals packing. This force is particularly important due to the nature of van der Waals interactions: Even if hydrogen bonding or polar interactions play a significant stabilizing role, because van der Waals occurs between any nonbonded atoms in close contact, it is a necessary force that is always present within the folded state. This means that van der Waals packing is essential for folding, but the extent at which packing can be a driving force for membrane protein folding is unclear. Without understanding the extent at which van der Waals packing contributes to folding, we cannot complete our understanding of how these other forces contribute to membrane protein association and folding.

The contribution of van der Waals packing to membrane protein folding can be broken down into three distinct interactions: lipid-lipid packing, lipid-protein packing, and protein-protein packing. Protein-protein (or sidechain) packing, is a technically feasible starting point because of the ability to manipulate sequences and determine changes in stability due to mutation. Previous research has demonstrated that disruption of packing within the core of bacteriorhodopsin destabilizes protein structure (Faham et al., 2004; Joh et al., 2009). In addition, a recent study using membrane protein design has shown that optimized sidechain packing can stabilize the folded state of phospholamban (Mravic et al., 2019). Although it is known that sidechain packing plays a role in stabilizing membrane protein structure in these individual systems, the energetic contribution of sidechain packing to the folded state of membrane proteins more generally has not yet been determined. My research aims to characterize and quantify the extent at which sidechain packing is a driving force for membrane protein association. To do so, I am investigating the role that sidechain packing plays in the association of homodimers, a simple and tractable model for studying membrane protein folding (Popot and Engelman, 1990). Using large scale computational design on common dimeric backbone geometries found within the PDB and a complementary high throughput assay, we have demonstrated that sidechain packing can facilitate association for a variety of membrane protein structures. This research gives insight into the extent at which fundamental sidechain packing impacts membrane protein association and folding, something that has not yet been done despite the importance of van der Waals packing to membrane protein folding.

The contribution of van der Waals packing to stabilizing membrane protein structure remains a significant gap in our understanding of membrane protein folding (Hong et al., 2014). Determining how sidechain packing impacts the association of membrane protein subunits will give us a better understanding of how membrane proteins assemble to fold stabilized structures. By understanding how these forces contribute to stability, we will be closer to obtaining a holistic view of how all forces involved combine to stabilize membrane protein structure. My research will increase our knowledge of the fundamental rules of membrane protein folding, and add to our understanding of complex membrane protein mechanisms, such as oligomerization and conformational change necessary for essential biological processes including signal transduction and ion transport (Sanders and Myers, 2004; Gregersen et al., 2006). Eventually, this knowledge can be used to design new functional membrane protein structures, advancing the field of synthetic biology.

Results

**Design of membrane protein homodimers**



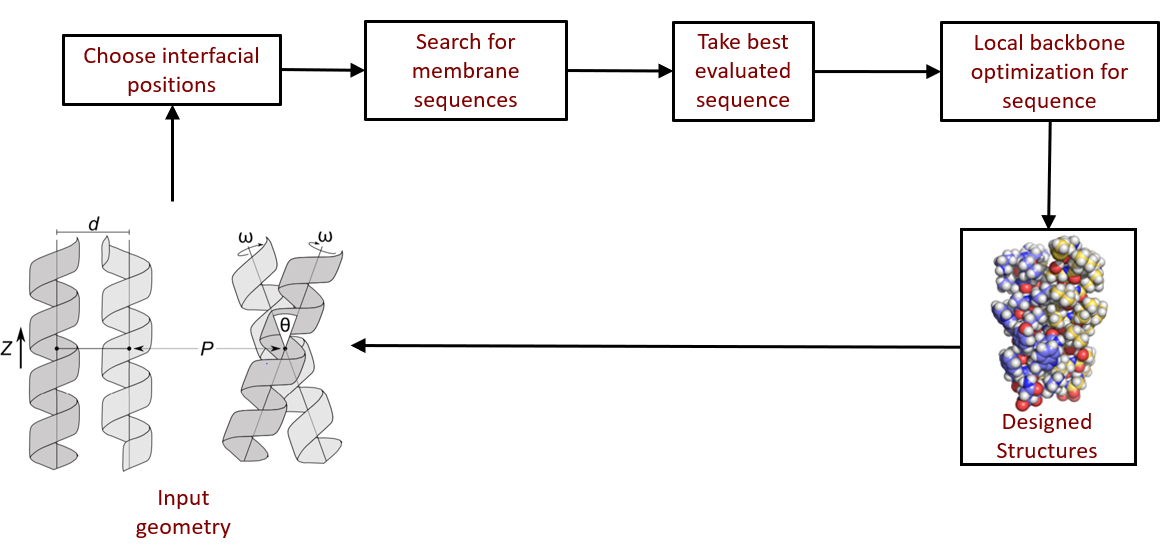


Figure 1. Membrane protein dimer design. (a) Helices within close contact (define how) were extracted from all membrane protein dimers submitted to the PDB in (date; 2019). Orientations of Proteins in membranes (OPM, cite) was used to only choose the proteins found in membranes. These geometric terms are plotted and overlaid over the density of these in space. (b) Design algorithm… (c)

[Insert design algorithm name here] designs homodimer membrane protein sequences by using an iteration of fixed backbone design and local backbone optimization. The sequence is initially optimized by evaluating van der Waals, hydrogen bonding, and an implicit membrane solvation model (cites). Simultaneously, we have developed a entropic term that determines the likelihood of a sequence to be found in membrane proteins by the comparison of amino acids found in membrane proteins (methods, cite). [design name] allows for the efficient design of hundreds to thousands of membrane proteins from a large variety of membrane protein geometries.

To design membrane protein dimers using [design name], we extracted the density of membrane protein helices within close contact found within the PDB (Figure 1A). This resulted in three distinct regions of high density: left handed, right handed, and GASright. ...

**Sort-Seq: Assessing membrane protein association in high-throughput**

To experimentally assess the dimerization of our 1000 designed proteins and their corresponding mutants, we used sort-seq, a high-throughput method that utilizes TOXGREEN which has been used previously to assess TM homo-oligomerization *in vivo* (cite TOXCAT and green). This method utilizes the expression of a chimera of the designed proteins fused to the dimeric ToxR transcriptional activator. Dimerization of the TM helices results in the dimerization of ToxR, allowing it to bind to a promoter that activates the expression of the reporter gene green fluorescent protein (GFP). This fluorescence can be used as an output for the association of the designed proteins in a biological membrane. An oligo pool gene library of the designed proteins is cloned into *Escherichia coli* bacterial cells that are run (better word) through a fluorescence activated cell sorter (FACs), allowing us to measure the association strength of our designed proteins in high throughput (cite sort-seq/SMA thesis; methods for the code?).

Fluorescence and association of designed structures and mutants

Figure 2A: Fluorescence Reconstruction general data (distribution of all proteins fluorescence, maybe as the pi charts or bar graphs; or both)

2B: more general data about fluorescence; maybe separate into different graphs (what other types of graphs can I use to represent fluorescence? Bar graphs with number of designs from each region that made it through; could also have bar graphs for their expected geometries here

2C: Another schematic about their geometries here? Maybe a quick graph to compare fluorescence between each region (with no expectation/expected result, just the data and explanation of it)

Figure 3A: Energy score vs fluorescence

3B: separate energy scores vs fluorescence here

3C: if we do regression analysis, could add that here

3D: also can add in geometry data for this set of structures; just the parsed data from clashing mutants

3E: void mutant data, also potentially from void mutants

Figure 4A: Conversion to deltaG

4B: snippets of westerns? Or at least quantification from westerns? Maltose test in supplement?

Discussion

Van der waals facilitates…x amount…approximately x% of GASright, known for Hbonding. Can we conclude anything about the contribution of van der Waals in GASrights and other regions of proteins? Are there any other studies that have left/right handed structures that do experiments that demonstrate packing/dimerization ability? Or portions of proteins found in some of these structures (maybe coiled-coil FtsLB data? What conclusions if any can be drawn from these? If they exist, could I take a look at the distribution of known proteins and their sequences/interfacial residues (any way to do this computationally?) and compare to them? I could at the very least compare to the residues found in the structures that I extracted from and see if this gives me any insight? Would it be possible to look at them positionally? Or at least look at the most similar structures by RMSD or something?

Could have a figure here for the confirmation of designed structures using a different method? TMDOCK could really be useful here because of the compared deltaG. Honestly, I likely could run TMDOCK in low throughput on just the proteins that match up with our clashing data, and maybe compare the deltaG values just to see how an experimental deltaG compares to an alternate energetic prediction than mine? Assuming all of the proteins work, I think it could be worth it

Acknowledgements

Twist, Flow core, Raman lab, CBI, SciMed, TMDOCK maybe, should send an email

Methods

* Many supplementary figures
  + More details in the starter outline

**References**

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