In 1973, Christian Anfinsen and colleagues postulated that the information required for a protein to fold into its native structure is encoded in its amino acid sequence. Almost 40 years have passed, but determining the three-dimensional structure of a large protein solely from its amino acid sequence remains a formidable challenge. In the past decade and a half, remarkable progress in the development of cost-effective high-throughput sequencing technologies has led to an explosion of genomic sequence information from evolutionarily diverse organisms. Exploiting this information with statistical tools and constraints derived from evolutionary principles has begun to show promise toward determining protein structure from sequence.

Recently, there has been increasing excitement in predicting protein structure by using co-evolving sites within protein sequences as a structural constraint [1-6]. These studies have demonstrated how improved statistical tools and the growing amount of sequence information hold enormous potential to predict the three-dimensional structure of proteins (Box 1). Along these lines, Hopf et al. [7] have developed a computational approach that exploits protein sequences to predict membrane protein structures. They demonstrate that the three-dimensional structure of α-helical membrane proteins can be determined with very good accuracy using their algorithm EVfold_membrane [7]. Their approach identifies positions in protein sequences that show correlated patterns of mutation and uses this information as a structural constraint to model membrane protein structures [7].

**Ab initio** structure prediction of membrane proteins

Reliable prediction of structures could have a major impact on our understanding of membrane protein function. This is underscored by the fact that less than 1% of the structures in the Protein Data Bank are of integral membrane proteins despite these comprising over 20% of all genes in mammalian genomes. Membrane proteins are physiologically crucial given their function as a vital communication interface between the intracellular and extracellular environments, and between the cytosol and diverse membrane-bound organelles. Hence, many membrane proteins are pharmacologically important and are potential drug targets. While efforts in structural genomics have led to the elucidation of structures of numerous soluble proteins, determining the structure of membrane proteins remains challenging due to difficulties involved in their expression, purification and crystallization. Thus, any approach that provides insights into structures of membrane proteins will be very useful in explaining their function.

Unlike soluble proteins, membrane proteins predominantly adopt α-helical (as seen in G-protein-coupled receptors) or β-barrel (as seen in porins) structures due to the constraints posed by the lipid environment. Membrane proteins with α-helical structure dominate most cell membranes, while those with β-barrel structure are confined to the outer membrane of bacteria and eukaryotic organelles. Due to the limited structural diversity seen in the membrane protein fold compared to soluble proteins, the former lends itself better to **ab initio** structure prediction.

In previous approaches, the topology of membrane proteins has been predicted using machine-learning approaches that are trained on lipid exposure and residue contact information [8] and energy-based folding methods that incorporate the knowledge of helix packing...
Box 1. Identification and interpretation of co-evolving positions

Metrics to quantify pairwise co-evolution of positions from multiple sequence alignments

State of the art approaches to obtain pairwise co-evolutionary correlations between positions in a multiple sequence alignment mainly rely on multivariate statistics. Such approaches take into account the frequencies at which certain amino acid residues appear at aligned sites. Among the most popular are Pearson correlation, maximum likelihood approaches, Bayesian statistics, chi-square-like methods and statistical coupling analysis [10]. In particular, approaches from information theory that use mutual information and Shannon entropy have recently been applied to three-dimensional structure prediction with promising results [1-8].

Filtering relevant and significant correlations from high background noise

One main obstacle in quantifying co-evolution is to filter the relevant information from noise that typically appears in the same order of magnitude as the signal seen for significant correlations [10]. This noise can be due to two main sources: (1) statistical noise due to incomplete or unrepresentative sequence sets and random co-variance, and (2) phylogenetic noise arising from the tree-like phylogenetic relationship of homologous sequences. Sophisticated normalization methods such as interdependency mutual information or comparison to test sets with random distributions can significantly enhance the signal to noise ratio [10].

Disentangling direct and indirect correlations

Maximum-entropy-based algorithms have recently been successfully applied to co-evolutionary sequence correlation in order to find residue contacts in proteins and have been implemented using a numerical method from mean field theory [7]. Another method is based on calculating the inverse covariance matrix to obtain partial correlations. In this approach, the accuracy of the strongest direct interactions can significantly be improved by sparse inverse covariance estimation, in which the inverse covariance matrix is numerically estimated while iteratively removing weaker interactions. An efficient algorithm known as graphical LASSO has been used to transfer this principle to co-evolutionary correlation problems in order to predict residue contacts [1,4].

From sequences to membrane protein structure prediction

According to Anfinsen’s hypothesis, the native protein structure is intrinsically encoded in its sequence. But how is it possible to infer this information solely from sequence data? The EVfold_membrane algorithm developed by Hopf et al. [7] uses a global statistical approach (entropy maximization) and identifies evolutionary coupling (co-evolution) between positions on a sequence from a multiple sequence alignment (Figure 1). The basic principle of the method is that spatially interacting residues tend to co-evolve; thus, co-evolving sites on a sequence can be used as a spatial constraint to predict the three-dimensional structure of a protein. Using the evolutionary coupling between positions as constraints, distance geometry and simulated annealing methods are then used to generate three-dimensional models of proteins, akin to the approach employed in deciphering solution structures of proteins using NMR spectroscopy. The predicted models are assessed and ranked based on various criteria, such as the quality of secondary structure, lipid accessibility of the residues and satisfaction of evolutionary constraints.

The performance of this algorithm was benchmarked against the experimentally determined structures of 25 membrane proteins from diverse protein families. The predictions showed general agreement with the overall fold and expected deviations in loop regions, unstructured segments in the transmembrane region, and in the orientation of side chains of amino acids. The power of this algorithm was further highlighted by applying the method to predict the structures of 11 membrane proteins with unknown structure. Some of the predicted structures show similarity to folds of known families, thereby making it possible to detect evolutionary relationships between protein families that have significantly diverged.

Beyond membrane protein structure prediction: function and dynamics

Unlike RNA structures, in which pairwise interactions between bases are highly specific, a single residue in a protein typically contacts multiple residues in its proximity to maintain the three-dimensional fold of a protein. This results in complex co-evolution profiles between different positions that may become hard to interpret. In other words, this aspect of proteins leads to
the appearance of background noise while computing pairwise co-evolution. This phenomenon makes reliable identification of significant co-evolutionary correlations difficult (Box 1). While there has been significant progress in improving the signal-to-noise ratio over the last years, the methods still have a high false-positive rate, making it difficult to differentiate between direct and indirect interactions between positions in the protein [8]. However, if one can disentangle this information, which is still an area of intense research, it can reveal key molecular aspects of the structure, dynamics and function of the protein.

In addition to inferring the structural proximity of specific residues, co-evolution between positions in a sequence can provide information about the conformation dynamics (including allosteric communication and alternative conformations) and molecular function (such as which residues are involved in substrate recognition, regulation and interaction) of proteins (Figure 1). By resolving contradictory distance constraints and investigating residues that co-evolve with a large number of positions, Hopf et al. [7] harnessed this information to identify different conformational states (as shown for membrane proteins glpT and OCTN, which transport glycerol 3-phosphate and organic cations, respectively). This has also provided hints about contacts in homooligomerization interfaces (as shown for the ABC transporter MsbA), and has facilitated prediction of functionally important sites (as shown for adiponectin receptor protein 1, AdipoR1). Thus, information on co-evolving positions can be exploited to gain important insights into the dynamics and function of proteins.

**Future directions and novel applications**

The computational study undertaken by Hopf et al. [7] demonstrates the potential of what one can learn about the structure of membrane proteins from residue co-evolution. While this is an impressive and important landmark in membrane protein structure prediction, there is still scope for improvement. For instance, constraints derived from biochemical experiments on protein function (such as restraining the distances between side chains of substrate binding residues in transporters) could be incorporated. Moreover, the method could be extended further by creating hybrid computational approaches that exploit Rosetta or other classical folding methods to leverage the strengths of constraints both in terms of co-evolution and energy.

In terms of applications, there are a number of problems where one can exploit this method. For instance, a variety of meta-genomics studies from diverse niches are providing us with a deluge of sequence information of several proteins that are completely uncharacterized. In this context, sequence co-evolution-based approaches can provide insights into structures of novel families of predicted proteins, which could be important for biotechnological and protein engineering applications. The algorithm can also be useful in medicine and drug development as a number of membrane proteins are implicated in several human disease conditions and many of them are difficult targets for structural analysis (for example, the cystic fibrosis transmembrane conductance regulator, whose structure remains to be determined). Thus, ab initio structure determination of such medically important membrane proteins could provide structural models to better interpret disease mutations and serve as potential starting points for structure-based drug design. Such
models can help identify structurally proximal residues that could be engineered to increase the stability of membrane proteins for expression, making them suitable for crystallization and for structure determination. With major advancements in the field of electron cryomicroscopy, it is now routinely possible to obtain low-resolution maps (8 to 12 Å) of proteins either by single particle reconstruction or electron crystallography. Inferring topology, subunit orientation and the oligomeric arrangement of membrane proteins using this method might provide models to interpret and complement the electron microscopy maps of macromolecular complexes involving membrane proteins.

Conclusions
Presently, structural biology struggles to hold pace with the rapid growth in genome sequencing. Only a fraction of all known sequence families is represented in the Protein Data Bank with at least one structure of its members. Despite the great progress in structure prediction using protein fragments, improved force fields, molecular dynamics simulations and homology modeling, structures of membrane proteins remain widely inaccessible. Computational methods such as EVfold-membrane thus hold the key to accelerating the structure determination of membrane proteins, thereby bridging the widening sequence-structure gap.

Abbreviations
NMR, nuclear magnetic resonance.

Competing interests
The authors declare that they have no competing interests.

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