THE DIFFERENTIAL GEOMETRY OF PROTEINS AND ITS APPLICATIONS TO STRUCTURE DETERMINATION

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1. Introduction

The accelerating growth of structural biology has created an enormous amount of information which we are only beginning to interpret. Today structural biology is approaching a comprehensive taxonomy of soluble globular proteins, and to make further progress it is imperative to address frontier areas which include membrane proteins, intrinsically disordered proteins, large macromolecular assemblies, and partially organized or heterogeneous structures such as cytoskeletal assemblies and amyloid fibers. These types of problems lie at the experimental frontier, as they severely tax the capabilities of existing physical techniques. Less widely appreciated is that these problems also lie at an intellectual frontier, as we currently lack a language to rigorously describe continuously variable or inexactatively defined structures. This review presents some recent efforts to use a continuous representation of protein structure to model and gain some insight into protein structure. The formalism employed is based on the differential geometry of continuous curves- the fold of a protein is considered as a curve which follows the path of the protein backbone. We have developed methods to interconvert continuous curves and discrete atomic coordinates as a way to create and manipulate protein models. In this review we present new tools based on continuous methods to investigate and utilize...
experimental structures and for examining relationships between the known structures and fold families. The review is organized addresses three scientific objectives:

1. **Structure determination**: How to devise methods for obtaining the geometric parameters needed to specify a curve (and its associated atomic model) from experimental data for the purpose of structure determination by electron microscopy, NMR, and crystallography.

2. **Study of Fold Continua for Structural Prediction and Comparison**: How to explore the possibilities simple protein architectures offer by variation of the underlying geometrical parameters to investigate the relations between known folds and as a novel means of predicting molecular structure.

3. **Continuum Mechanics of Biological Structure**: How to model conformational changes associated with biological function in terms of deformations of elastic bodies.

The mathematical methods developed for creating smooth deformations of protein models can be used to investigate intrinsically disordered regions, evolutionary structural change, distributed conformational changes associated with binding, and accommodation of structures to quaternary structure rearrangements.

The starting point is to represent and model the protein backbone by continuous curves or surfaces. A basic mean of comparison between mathematical structures is to create transformations or mappings between them and then to investigate the properties of the mappings. In particular, properties of continuity and differentiability are crucial for such investigations. The natural representation of molecular structure with atomic coordinates is discrete and so precludes the application of continuous methods. However, continuous supersets which contain the discrete points corresponding to atomic positions can be operated on in this manner. Use of this alternative continuous representation allows the application of powerful analytical and geometric methods to answer questions about protein folds and conformations and their relationships through study of their corresponding space curves and surfaces.

It is critical not to lose sight of the atomic details, which are the foundation for understanding the properties of proteins and provide much of the underpinning for our current mechanistic view of biological processes. The challenge is to integrate both continuous and discrete representations to
best understand the mathematical, physical, chemical, and biological properties of proteins and other macromolecules, using the appropriate viewpoint and theoretical tools for a particular question.

In Section 2 the relevant mathematics for protein curve construction are described, and various avenues for further development and new mathematical challenges will be indicated. In section 3, three distinct mathematical problems which underly extraction of curve parameters from various types of data are posed for study - progress in any of the three will yield new ways of utilizing data for structural analysis. Section 4 describes how the small number of geometric parameters needed to describe simple protein architectures allows the systematic search of an entire fold space. Section 5 applies the capabilities of continuous deformation of folds from section 4 and the optimization of curve parameters from section 3 to the problem of protein design.

2. Methods

The path of a protein backbone is often represented as a series of line segments connecting the alpha-carbon atoms. It could also be represented as a smooth curve passing through the same points. In general, regular three-dimensional curves can be completely specified by their curvatures which describe the local bending and twisting of the curve along its length. The local description in terms of curvatures and the global description in terms of spatial coordinates are entirely equivalent. For the particular case of proteins, we have developed specific methods to construct the curvatures of curves that follow the path of protein backbones, and to construct coordinate models of proteins from such curves. The interplay between the local and global descriptions and in particular the modulation of curvatures to control the three-dimensional shape of protein models plays a central role for the work described here. The rest of this section gives some technical details on how to construct curves from data sets and, conversely, how to build protein models with idealized geometry from curvatures. The next sections explore various applications of this formalism to structure determination, protein fold exploration, and protein design.

Let $r = r(s)$ be a curve in $\mathbb{R}^3$ parameterized by its arc-length $s$. At each point $s$ on the curve, one can define (assuming sufficient regularity of the curve) a local general orthonomal basis $\{d_1(s), d_2(s), d_3(s)\}$ by defining the orientation of the vector $d_3$ with respect to the tangent vector $r' = v_1 d_1 + v_2 d_2 + v_3 d_3$ to the curve. Since the vectors $\{d_1(s), d_2(s), d_3(s)\}$ are
Figure 1. Construction of curves from curvatures. A curve (C.) is constructed from its curvature-torsion profile (A. and B.). Note that the signed curvature (real) is used here rather than the curvature itself (assumed positive). D. The position of the C$_\alpha$ atoms obtained from the curve and the experimentally determined C$_\alpha$ coordinates of bacteriorhodopsin are superimposed. This example utilized 96 curve parameters to specify the curve, where as 3 coordinates are required for each of the 228 atoms in the C$_\alpha$ trace.

orthonormal, their evolution is governed by

$$\frac{\partial d_1}{\partial s} = k_3 d_2 - k_2 d_3, \quad \frac{\partial d_2}{\partial s} = k_1 d_3 - k_3 d_1, \quad \frac{\partial d_3}{\partial s} = k_2 d_1 - k_1 d_2.$$ (1)

That is, $D' = DK$, where $D$ is the matrix whose columns are the basis vectors, $(\ )'$ denotes the derivative with respect to $s$, and $K$ is the skew-symmetric matrix

$$K = \begin{bmatrix} 0 & -k_3 & k_2 \\ k_3 & 0 & -k_1 \\ -k_2 & k_1 & 0 \end{bmatrix}$$ (2)

This general description of local bases for curves allows to define both the shape of the curve but also the evolution of a triad of orthonormal vectors attached to it through the specification of a vector of curvatures $k = (k_1, k_2, k_3)$ and a vector of basis orientations $v = (v_1, v_2, v_3)$. This description becomes more familiar if we specialize the general basis by defining $d_3$ as the tangent vector (that is, $(v_1, v_2, v_3) = (0, 0, 1)$ and $d_1$ as the normal vector). In which case, Eq. (1) becomes the Frenet equations and the curvatures of $r(s)$ are the entries of the Darboux vector $k = (k_1, k_2, k_3)$ and they can only be determined up to a phase factor that describes the phase difference between the normal vector and the first basis vector $d_1(s)$. The general basis has many advantages due to its
compact form, the possibility to define local bases for less regular curves or to assign to the phase factor additional information (such as the twist and shear of a ribbon, or the material property of a tube surrounding the central curve). For clarity sake, in this review, we mostly restrict our analysis to the Frenet frame and define \( D = [n(s), b(s), t(s)] \) as the normal, binormal and tangent vectors and \( k = (0, \kappa, \tau) \) where \( \kappa \) is the signed curvature (defined on the real rather than strictly positive) and \( \tau \) is the usual torsion. The exact description of protein backbones with piecewise helical curves in the next section will require the use of the general basis.

2.1. Curvatures to curve.

If the curvatures \( k \) and basis orientation \( v \) are given, the curve can be readily obtained by integrating Eq. (1) together with \( r' = \sum a_i d_i \). These equations form a system of 12 linear non-autonomous equations for the basis vectors \( D \) and curve \( r \) that can be written in a compact form by introducing the vector \( Z = (d_{1,x}, d_{2,x}, d_{3,x}, d_{1,y}, d_{2,y}, d_{3,y}, d_{1,z}, d_{2,z}, d_{3,z}, x, y, z)^T \) in which case, the linear system reads

\[
Z' = MZ,
\]

with

\[
M = \begin{bmatrix}
K^T & 0 & 0 \\
0 & K^T & 0 \\
0 & 0 & K^T \\
V_1 & V_2 & V_3 & 0
\end{bmatrix},
\]

where \( K \) is defined above and \( V_i \) is the 3-matrix whose only non-zero entry is row \( i \) with value \( v \). The integration of Eq. (3) as a function of \( s \) provides both the curve position but also the evolution of the basis. In the particular case where Frenet frame is used \( v = (0, 0, 1) \) and for given curvature \( \kappa \) and torsion \( \tau \), the curve is reconstructed. An example of such a construction is given in Figure 1.

2.2. Curves from atomic models.

The first problem is to obtain curves describing the protein backbone from a set of \( C_\alpha \) coordinates. One of such construction can be found in Richardson ribbon diagrams, where curves, ribbons and helices are used to build a three-dimensional picture of the protein backbone. However, this remarkable construction, obtained by spline fitting, is mostly used for visualization purposes and its mathematical formulation is not a faithful representation of the protein that can be used for other purposes \cite{footnote1, footnote2}. Therefore, one first needs to obtain an exact curve representation of the protein backbone, that
is a curve that passes exactly through each \( C_\alpha \) atom. Obviously, this can be done in many different ways since there are infinitely many choices of curves passing through a given set of points. A possible choice is through the use of polynomial or rational splines by considering the set of \( N \) first atoms and find the spline of degree \( d \) through this set, then sequentially build another spline for the next \( N \) atoms (with possible overlap). Depending on the degree of smoothness required on the curve the parameters \( N \) and \( d \) can be adjusted within the proper algorithm \(^7\). This representation offers an exact description of the protein backbone but does not carry much information on its global structure. In effect, it replaces one discrete set of points by a discrete set of curves through subsets of points.

A more useful way to represent the \( C_\alpha \) trace is to find a piecewise helical curve (referred to hereafter as a polyhelix) through the points, that is a continuous curve built out of connected helices. Many authors have considered the problem of fitting helices through sets of points in space. This problem arises in protein structure \(^7\), engineering design of cables and springs \(^7\), and nuclear and particle physics for particle tracking \(^7\). To obtain an exact representation by a polyhelix, 4 consecutive \( C_\alpha \) atoms are considered and a unique helix can be constructed (see for instance \(^7\)), the helix is characterized by its first \( C_\alpha \), curvature and torsion, and axis. To pinpoint the position of the other \( C_\alpha \), three arclengths are required. Together, it amounts to 12 data points corresponding to the 12 atomic coordinates, providing a 1-1 map between atomic data and helical data. The construction proceeds by considering the fourth to seventh \( C_\alpha \)'s for the next helical piece and so on (See Figure 2). This construction does not provide a purely local representation of the curve since extrinsic data (position of the axis in space) is required. However, using the general basis described above, a complete local representation of the curve can be obtained by specifying for each local piece the following data for the \( i \)th helix starting at the atom number \( 3i + 1 \): a constant curvature vector \( \mathbf{k}^{(i)} \), the orientation of the basis given by a constant vector \( \mathbf{v}^{(i)} \) and the arclength positions of the atoms on the helix \( S_i = \{s_{3i+2}, s_{3i+3}, s_{3i+4}\} \). This represents 9 data for each successive 3 atoms. The change of orientation of the axis is characterized by the change in the vector \( \mathbf{v} \) between the \( i \)th and \( (i+1) \)th helices. Computationally, the protein backbone and the position of the \( C_\alpha \) is fully characterized by a list of triplets \( H^{(i)} = \{\mathbf{v}^{(i)}, \mathbf{k}^{(i)}, S^{(i)}\} \) and the positions are recovered in extrinsic coordinates by integrating Eq. (3). While this operation seems to be a daunting task, it actually amounts to straightforward matrix algebra due to the exact analytical solution of these differential equations in the cases
where curvatures and orientations are constant (See Section 2.4 on polyhelic). It is important to note that this representation is general and not restricted to the analysis of proteins with alpha-helices. A representation of atomic data in terms of polyhelic as many advantages: it is purely local in nature and so exploits the natural geometry of the protein, the curvatures carry global information on the curve through curvatures and torsion and therefore allows for direct identification of regions of interest (for instance alpha-helices or different types of turns), and modulation of these curvatures over long distances identifies long-range structure (e.g. bending and curving of helices, twist of beta-sheets, etc...). To a certain extent, different authors have explored the local geometry of existing proteins using similar approaches with alternative formulations. This construction is the starting point of our analysis. The relevant aspect of local representation is that it helps connect experimental data to structure determination and modeling performed through the use of curve geometry as presented in the next Sections. There, idealized geometries based on polyhelic are used to explore possible folds in parameter space. Polyhelic provide an exact local representation of the protein backbone. However, in many studies, one may be interested in describing nonlocal properties of these structures. For instance, one may be interested in representing an alpha-helix that may not be strictly helical (due to bending or super-twisting) by one single helix. The problem is then to fit a given structure (helix, plane, ...) through a set of points. This can be done through some averaging process on the exact local representation of curves from coordinates: A continuous polyhelix curve of 4 segments constructed from points. B: Construction of coordinates from curves: The local basis on a curve centered on the 3rd Cα and selected atoms expressed in that local coordinate system. C: Section of an idealized polyserine helix constructed with EDPDB (red) compared with the model constructed in the local coordinates (blue) as in section 2.3.
An example of helix fitting is shown in Figure 1 where the curvature-torsion profile was optimized by fitting helical segments to the Cα coordinates from the protein bacteriorhodopsin. There are many different outstanding mathematical and computational issues associated with the problem of fitting a helix, or a cylinder through which will are addressed in separate papers.

2.3. Atomic models from curves.

Curves can be used to describe proteins and construct models in many different ways. At the basic level, the curve can represent the backbone and Cα can be superimposed by imposing that they are located on the curves at determined position s. In particular, at suitable values of s, the local coordinate system has its origin at the Cα positions, which is a natural choice for the coordinate system in which to express the atomic coordinates for the remaining atoms in the residue. A set of local coordinate system $\mathbf{a} = \{a_1, a_2, a_3\}$ represents the point $\mathbf{p}_a = \mathbf{r}(s) + \sum_{i=1}^{3} a_i \mathbf{d}_i(s)$ in the external coordinates. Conversely, any point $\mathbf{p}_a(s)$ has local coordinates

$$a_i = (\mathbf{p}_a - \mathbf{r}(s)) \cdot \mathbf{d}_i(s), \ i = 1, 2, 3.$$  

We have converted and tabulated the local coordinates for all the rotamers from which allows the construction of the alpha-helical regions of protein models from curves with idealized geometry, and which can also be used, albeit with some distortion of the backbone atomic arrangements, in the rest of the protein. In future work we will improve the local coordinate library by studying the dependence of the local coordinates on curvature and torsion, that is to obtain a local description of the rotamers in terms of the curvatures both for simple geometries and in the general case.

2.4. Polyhelices.

A particularly simple choice of curves is obtained by choosing the curvatures to be piecewise constant so that the curves are piecewise helical. These polyhelices can be used to map precisely atomic coordinates to continuous curves. Conversely, they can also be used to study and classify large families of proteins with idealized geometries. The advantages of this representation are threefold: first, it is consistent with the representation from the atomic coordinates making the comparison with experimental data straightforward; second, large families of proteins can be represented by few parameters and the exploration of fold spaces can be achieved with minimal effort; third, the computation of polyhelices can be reduced to simple linear
algebra, making it computationally exact and reliable. The computation of polyhelices is achieved by integrating Eq. (3). Since $M$ is now a constant matrix, the solution of this system is given by

$$Z(s) = A(\kappa, \tau; s)Z(s_0)$$

(4)

where $A(\kappa, \tau; s) = e^{(s-s_0)M}$ is the matrix exponential. Matrix $A$ can be computed exactly and its entries are linear combinations of trigonometric and polynomial function of $s$ with coefficients depending on $\kappa$ and $\tau$. A polyhelix with $N$ helices is completely characterized by a list of curvatures and length:

$$P = \{ (\kappa(i), \tau(i), L(i)), i = 1..N \}$$

and an initial position and basis orientation $Z(0)$. The $j$th helix on the curve is given by the last three components of the vector $Z_j$:

$$Z^{(j)} = A(\kappa^{(j)}, \tau^{(j)}; s)Z^{(j-1)} = \prod_{k=1}^{j} A(\kappa^{(j)}, \tau^{(j)}; s)Z^{(0)}, \quad L_{j-1} \leq s \leq L_j,$$

(5)

where $L_j = \sum_{k=1}^{j} L^{(k)}$. Examples of such computations are given in Figure 1. This analytic expression for the curves provides an efficient way to explore fold space as shown in the next Sections.

### 2.5. Embedding methods.

While $\alpha$-helices lend themselves to a linear description, beta-sheets are inherently a linear but 2-dimensional structure. It is therefore natural to use a 2-dimensional embedding of a curve to describe them. The construction proceeds in 2 stages as illustrated in Figure 3. The linear character of the chain is accommodated by a description of the backbone as a plane curve specified by its curvature. The second stage is to map the curve into a two-dimensional surface plane in a three-dimensional space. The particular embedding chosen defines the surface characteristics of the beta-sheet model and can be described by the classical Darboux frame field representation. Here again, the atoms can be represented in the local Darboux frame. The choice of mapping is restricted by the constraints on bond angles and distances and possible beta-architecture with such constraint can be systematically explored within this framework (this and related ideas on possible architectures are discussed in Section 4).

### 3. Fold space exploration

The set of protein folds is a subset of the set of all possible space curves that can be constructed by standard differential geometry tools. By in-
vestigating the set of possible curves, we can find within it the possible protein folds. The key problem is to identify among all curves the ones that may constitute the path of a protein backbone. Mathematically, the problem amounts to finding functions to score the potential of a curve to take the shape of a protein. The main idea is then to explore continuous families of curves defined by a set of parameters and isolate good protein candidates that correspond to points within that parameter space. Small parameter spaces which describe simple protein architectures can be exhaustively sampled. The ability to explore the entire realm of possibilities inherent in a particular architecture makes it possible to see relations between folds that may not be apparent. The method has applications to protein structure prediction, genome interpretation, and homology modeling. Sequences can be threaded onto curves obtained by a systematic or guided parameter space search. Postulated folds may also serve as protein design targets. Finally, the ability to “interpolate” or “extrapolate” from existing folds may allow prediction of new folds (and explicit construction of their coordinate models) before they are experimentally observed.

3.1. Protein Quality Functions.

Most curves in space could not be realized as paths of protein backbones, because they have impossibly tight bends, unrealistically straight segments, have regions that approach too close to other regions of the curve, or are too loosely packed to have sufficient interactions to remain folded. However, there are some curves that satisfy all those criteria. A fundamental question
is to identify simple criteria (geometric and physical) to quantify whether a given curve might be realizable as protein backbone conformations if the right amino acid sequence could be found. To address this question, we introduce the idea of protein quality functions, to quantify the potential of a curve to be realized as a protein fold. The curvature space is the space of parameters defining a family of curves (for instance, the family of helices is a three-dimensional space defined by curvature, torsion, and length) and a protein quality function is defined at each point of the curvature space and takes real values. A contour plot of a quality function over the curvature space would have islands in regions that correspond to protein-like curves (for instance, $\alpha$-helices would score very high in the family of helices and a small island would be centered around the ideal value of curvature and torsion for $\alpha$-helices). Once such a function has been identified, it is possible to investigate questions about the density of folds in fold space, or the connectedness of fold space (i.e. are regions of protein folds connected or widely separated?). What are the possible choices for quality functions? Clearly, to conduct a search over large regions of the fold space, the quality functions should be easy to evaluate.

A simple protein quality function can be expressed as a ratio of a term that expresses curve compactness and a term that penalizes a curve which approaches itself too closely. To quantify compactness, the notion of contact order $^7$ serves nicely. Given a set of points $\{P(s_k)\}$ on a curve, two points $P(s_i)$ and $P(s_j)$ form a contact when within a prescribed contact distance in space. The contact order is the sequence distance $|s_j - s_i|$ averaged over all contacts. Contact order is large for curves in which many pairs of points distant on the curve are close in space and so serves as a simple quantitative measure of compactness.

$$C_O = \frac{1}{LN} \sum_{i=1}^{N} |s_j - s_i|$$

(6)

where $N$ is the number of contacts and $L$ the number of points. However, a curve that is too compact will approach too close to itself. Defining a clash as a pair of points which are closer than a prescribed clash distance in space, curves with self-intersections are severely penalized by using the quality function:

$$Q_1 = \frac{C_O}{2M}$$

(7)

where $C_O$ is the contact order of the curve and $M$ the number of clashes. Examination of the distances between $C_\alpha$ coordinates of several repeat
proteins suggested the use of 9 Angstroms as the contact distance and 4 Angstroms as the clash distance. An advantage of this quality function is that it can be used both on $C_\alpha$ coordinates from real proteins and on points obtained from curves.

There many different choices for quality functions emphasizing different features of a candidate curve. Such functions could be evaluated on smooth curves which may have advantages for theoretical investigations or on discrete sets of points obtained from the curve, which have the advantage of ease of comparison with experimentally obtained coordinate sets. Other interesting possibilities include the use of the global radius of curvature $\rho$ providing both a local estimate of curvature and a global estimate of self-contact; simplified versions of energy functionals as used in homology modeling and structure prediction $\rho$; family of Vasiliev knots invariants for protein which have already shown great promise for classification purposes $\rho$; or statistics of distributions of curve parameters in curves fitted to experimentally determined coordinates.

3.2. The fold spaces of polyhelices.

As an example of fold space exploration we use the quality function $Q_1$ defined above to study the fold space of some polyhelical families. As discussed in Section 2, a curve consisting of $N$ helical segments can be specified by a list of $N \{\text{curvature, torsion, length}\}$ triples and the fold space is a $3N$-dimensional which can be systematically explored with the function $Q_1$. This space is very convenient to model proteins with $\alpha$-helices since a few parameters are necessary to describe the main building blocks of the proteins.

First, a simple application of the use of the quality function to rank the turns connecting two $\alpha$ helices is shown in Figure 4. The curvature/torsion profile and its corresponding helical hairpin are shown. The turn is parameterized by the two triples $\{\kappa_1, \tau_1, l_1\}, \{\kappa_2, \tau_2, l_2\}$. This 6-dimensional fold space can be searched exhaustively. Contour plots (in which light colors indicate high (favorable) values of $Q_1$ function) as a function of $\kappa_2$ and $\tau_2$, for different values of $l_2$ are shown for 2 different choices of $\kappa_1, \tau_1$, and $l_1$ in Figure 4. The presence of islands and plateaus indicates that only certain combinations of curvature and torsion gives rise to reasonable turns. By selecting points within the white regions, a list of turns which specify high-scoring protein-like helical hairpin curves can be collected in a library of candidates for connecting turns. Once this library is built, one can pro-
ceed hierarchically with the search for protein candidate by finding helical repeats where the connecting turns are given by helical hairpin from the library.

As a more complex example, consider the various types of helical repeat proteins which share a common architecture of \((\text{helix}_1\text{-turn}_1\text{-helix}_2\text{-turn}_2)_N\). Curves corresponding to this architecture can be specified with 14 parameters. (The curvature and torsion of an alpha helix are fixed, so only 2 parameters for the lengths of the helices are needed. Each turn is described by 2 \(\{\kappa, \tau, l\}\) segments.) A curve is determined by a point the 14 dimensional fold space. A systematic search of this space is still a daunting undertaking and some simplifying assumptions are necessary. We assume that the sections helix\(_1\text{-turn}_1\text{-helix}_2\) and helix\(_2\text{-turn}_2\text{-helix}_1\) as helical hairpins. This reduces the search to a 4-dimensional space, over the two (continuous) helix lengths and two turns from the (discrete) helical hairpin list. For high-scoring curves, we construct polyalanine atomic models and overlay them on experimentally determined repeat protein coordinate sets (PDB entries 1i7x, 1b89, and 1b3u. In future work we will expand the
comparison and use more sensitive methods of structure comparison \(^?\). An example of a “hit” from this search is shown in Figure 5, in which a curve close to the armadillo repeat protein \(\beta\)-catenin (PDB code 1i7x) was obtained. Quite remarkably, this result shows that a construction solely based on simple geometric principles can capture the form of existing proteins accurately, and that simple quality functions can be used to search rapidly through the curve specification parameter space and identify protein-like curves. Suggestively, some regions of the fold space have high quality values and yet describe curves that do not resemble any known proteins. Some appear to have plausible packing arrangements- examples are shown in Figure 5. The left three show examples of a family which can be described as a stack of antiparallel coiled-coils. The right three show examples of a family which most closely resembles leucine-rich repeats but which have a second helix in place of the beta-strand. Once interesting regions of a fold space have been identified with one quality functions, it can be further explored by using other quality functions providing, in effect, a series of filters for plausible proteins.

This methodology has been demonstrated with helical repeat proteins due to the small number of parameters needed to describe curves with such architecture. However, the same idea is applicable both to non-repeat proteins and also to beta- or mixed alpha-beta architectures by using different (or mixed-) representations of curves such as the one given in Section 2.5 for beta sheets.

4. Protein Design

The goal of \textit{de novo} design of proteins is to create sequences that fold into a desired three-dimensional structure \(^?\). The continuous representation of proteins is of great utility for design work as it allows the overall specification of protein architecture without requiring that all the atomic details be considered at the outset. It separates the geometric problem of finding a suitable backbone from the problem of consistent atomic interactions. Therefore, it provides tools to specify a protein scaffold that may not resemble natural protein folds.

The primary determinant of globular protein folds is encoded in the binary pattern of hydrophobic and polar residues which defines the hydrophobic core and hydrophilic surface of the protein \(^?\). Recognizing the fundamental character of the hydrophobic core for structure specification, the computational protein design field initially focused on means to design
cores compatible with existing natural scaffolds. In its simplest form, the problem of protein design is the problem of selecting the best sequence which fills the interior volume of a given scaffold. The need for criteria to distinguish among different sequences that satisfy the crude inside/outside constraint but which led to stable but non-unique structures has led to the development of sophisticated energy functions that take into account the many physical interactions specifying a unique folded protein structure. By incorporating these finely-tuned energy functions with the power and efficiency of new combinatorial algorithms, core-packing algorithms have matured to the point where realizable designs have become routine. However, it has become increasingly clear that protein backbones are not rigid scaffolds but rather can move in unexpected ways in response to changes in core composition. The problem of how to incorporate such backbone freedom into protein design algorithms, and to control the computational cost of exploring these extra degrees of freedom, is the main challenge to the field protein design field.

The greatest appeal of a geometric approach to protein design is that it separates the specification of structure from the validation of structure. The geometric representation can be used to construct new plausible models, and different energy functionals can be used to rank them. By separating the two problems, the constraints of sequence and structure can be looked at as independent, and either can be varied to best satisfy the other. In essence, one can ask the question “What is the best sequence for this particular curve?” as well as “What is the best curve for this particular sequence?” By iterating between the two problems, a solution for which both sequence and structure are mutually optimal can be determined.

The continuous representation provides a theoretical foundation for backbone design as well as natural and inexpensive computational methods especially compared to the computations in dihedral angle space.

Recently hybrid methods which alternate between sequence rearrangement and energy minimization to allow backbone relaxation have shown great promise. We propose to generalize these ideas by adapting the continuous description to allow simultaneous minimization of an externally specified energy function with respect to curve parameters and side-chain rotamers as a way to find mutually optimal sequence and structure solutions.
4.1. *Creation of structure specification and optimization tools.*

An important part of the proposed work in protein design is the development of algorithms for optimization of models optimizing user-supplied design criteria (such as the minimization of an energy function) by adjusting curve parameters, and the identification of an interior volume needed to construct a hydrophobic core. Considered in combination, these two constraints will identify suitable scaffolds for design targets. Section 5.2 will discuss how to create sequences that best conform to these scaffolds.

The optimization of curves and curve-derived coordinate models by variation of curve parameters is based on the same underlying methods as the ones described in Section 3 where agreement with experimental data rather than with design constraints was required. From an atomic perspective, the design task is to create energy functions balancing the relative contributions of the different types of interactions (e.g., electrostatic, Vanderwaal’s, hydrophobic) involved in stabilizing folded proteins. From the coarse-grained perspective given by the continuous description, simple geometric criteria based on protein quality functions have considerable discriminating power. The main idea is therefore to use the optimization algorithms of Section 3 with a protein quality function of Section 4 to identify suitable design targets which can then be used to construct detailed atomic models as in section 5.2.

The starting point for sequence design is the determination of the binary hydrophobic/hydrophilic pattern in the primary sequence. A design target conformation can be characterized by a curve. To determine a binary pattern along the curve, it is necessary to identify the interior volume enclosed by the curve. To do so, a simple method (See Figure 6) consists in locating $C_\alpha$ atoms on the curve, and to construct a grid covering the extent of the $C_\alpha$. For every point on the grid, the number of neighboring $C_\alpha$ atoms is determined. Choosing only the points which have many neighboring $C_\alpha$ atoms as the center of spheres, an approximate "interior" volume is obtained by taking the union of these spheres. Constructing the $C_\beta$ atoms from the curve, the atoms facing the "interior" volume can be assigned a hydrophobic character by standard core-packing algorithm like DEE. Subsequently, the curve can be modified so as to bury more or fewer side-chains according to desired criterion such as buried hydrophobic surface or volume.
Figure 6. Determination of binary patterning. A: A three-helix bundle curve. B: \( \alpha \) atoms on the curve. C: "Inside" volume as the union of spheres centered at points near (within 8 Angstroms) sufficiently many (more than 12) \( \alpha \) atoms. D: \( \beta \) atoms constructed from the curve located within the "inside" volume are colored red.

5. Continuum Mechanics of Biological Structures

The methods described in the previous sections can describe protein models accurately but are of geometric nature and exist without reference to physical assumptions about molecules. In this section we build on this foundation, refining the models by incorporating physical considerations to relate structural and energetic properties of molecules and moreover make experimentally testable predictions. We will use classical elasticity theory for this purpose. Primarily we will seek analytical solutions by employing the semi-inverse method of Saint-Venant, although with modern computational resources, numerical solution of the elastic field equations can be applied when needed as an alternative.

For the purposes of structural modeling, elastic deformations can be used to describe large-scale, distributed conformational changes. But importantly, these responses are described in terms of changes in body coordinate systems, which can be made to coincide with the local coordinate frames we have used for construction of atomic models. Thus elasticity provides a natural formalism for devising physical theories which make structural predictions. For instance the elastic energy of filamentous structures can be described with the equation

\[
E_{\text{elastic}} = \frac{1}{2} \int_0^L \left( B_1(s)\kappa_1^2(s) + B_2(s)\kappa_2^2(s) + B_3(s)\kappa_3^2(s) \right) ds \quad (8)
\]

where \( B_1, B_2, \) and \( B_3 \) are the elastic constants and the \( \kappa_i \) are the deviations from equilibrium values of the curvatures. Additional terms may be included, for instance to include nonlocal effects or to obtain force-extension curves for study of mechanical responses.
5.1. Continuum elastic theory of coiled-coils

Within this formalism, a coiled-coil can be modeled as conjoined elastic filaments with elastic constants $B_1 = B_2 = B$ which describe resistance to bending and $B_3 = C$ which describes resistance to twisting. In a coiled-coil configuration, the center line $r(s)$ of each filament itself is a helix. The axis is along $z$, the radius is written $R$, the pitch is $2\pi R/\tan \theta$ and the super-helical angle $\theta$ is the complement of the pitch angle. We parametrize the (helical) center line as:

$$r(s) = \begin{pmatrix} +R \sin \psi(s) \\ -R \cos \psi(s) \\ s \cos \theta + z_0 \end{pmatrix}, \quad \psi(s) = \frac{\sin \theta}{R} s + \psi_0 \quad (9)$$

where $\psi(s)$ is the equatorial angle in the $(x,y)$ plane. The (constant) curvature and torsion of the super-helical axis are $\kappa = \sin^2 \theta/R$ and $\tau = \sin \theta \cos \theta/R$. The Frenet and Cosserat frame vectors are obtained from this parametrization by standard identities and can so be used to construct atomic models.

Coiled-coils are held together by interactions along one face of their constituent helices, and this interaction face can be parametrized in terms of the Cosserat directors and the parameter $\hat{\tau}$ which describes the twisting of the interaction face along the surface of the alpha helix. Constraining these interaction surfaces to be joined provides a structural constraint, and minimizing Equation 8 subject to this constraint yields the equilibrium conformation of the coiled-coil in terms of a relation between the elastic constants $B$ and $C$, the equilibrium super-helical angle $\theta_0$ and the interface parameter $\hat{\tau}$.

$$-\frac{2B \sin^3 \theta_0 \cos \theta_0}{C \cos 2\theta_0} = \sin \theta_0 \cos \theta_0 - \hat{\tau} R \quad (10)$$

This elastic theory agrees well with structures of leucine zipper coiled-coils (Table 1). Under the present proposal we will employ it to construct atomic models of large macromolecular complexes not easily accessible to experimental structural analysis.

5.2. Modeling the open and closed states of the CusCFBA bacterial efflux complex

An application of the coiled-coil theory is to the bacterial metal efflux complex cusCFBA. This complex allows bacteria to survive high concentrations
of toxic heavy metals such as copper by pumping them out of the cell. Other members of this family of proteins are involved in bacterial drug resistance. The structure of this complex is shown schematically in Figure 7. CusC forms the channel, CusA is thought to function as a pump, and the peripheral subunits CusF and CusB are thought to effect the opening and closing of the channel in response to metal concentration. A crystal structure of TolC provides a structural model for the closed form of CusC in which the channel is blocked, and these authors postulated a distinct open conformation for this molecular complex. The particular bending and twisting which is needed to open and close an iris-like arrangement of helices is also well-described within our coiled-coil theory. We propose to use this theory to devise atomic coordinate models of the states of the complex to investigate the mechanism of action of the cusCFBA complex, and in particular the role of the periplasmic subunits cusF and cusB.

Our theory of coiled-coils relies on a geometric relation between the interface residues by which the individual helices associate. A simple modification allows the constraint to be modified to devise barrel-like structures, which are open in the middle. But it is considerably more complicated to devise a constraint equally compatible with two distinct states which must be accommodated in an iris-like structure. In the simple coiled-coil theory an interaction surface on the individual helices (such as the stripe of leucine residues in a leucine zipper) defines the coiled-coil geometry. Our hypothesis for the general problem of forming an iris is that two distinct interaction surfaces (with different values of $\hat{\tau}$) must necessarily exist, which specify distinct coiled-coil geometries corresponding to the open and closed states. In the context of the cusCFBA system, the periplasmic subunits cusB and cusF could bind to one or the other interaction surface so as to control the state of the channel subunit cusC. This hypothesis is will be tested experimentally in the McEvoy laboratory.

<table>
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<th>GCN4</th>
<th>res./turn</th>
<th>X-ray data</th>
<th>$R$</th>
<th>$2\theta$</th>
<th>$\hat{\tau}$ (rad/Å)</th>
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<td>7.6 Å</td>
<td>$-26.0^\circ$</td>
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</table>
Figure 7. From left: Individual filaments of a coiled-coil in the A) unstressed state B) twisted state C-E) Different coiled-coil configurations constrained by their interaction surfaces. The equilibrium state requires a balance between bending and twisting of the filaments. F) Ribbon diagram representing the bacterial efflux complex. (Coordinates are from the closely related multidrug resistance system components TolC (1EK9.pdb) G) Schematic models of an iris formed from a hexameric coiled-coil. (Note CusC is trimeric but its iris is formed by 12 helices.)

5.3. Modeling oligomerization states of Adiponectin

A second application of the coiled-coil theory is to the anti-diabetic signalling hormone adiponectin, which stimulates insulin sensitivity. This molecule circulates in the bloodstream in three distinct oligomerization states (trimer, hexamer and 18mer) but only the largest appears to be active in signalling. The Acrp30 gene coding for adiponectin is organized into segments which code for a globular headpiece, a collagen-like domain, and a short tail region. The structure of the globular headpiece was determined by crystallography but the detailed structure of the remaining portion of the molecule is unknown. The collagen-like domain is necessary for the assembly of adiponectin monomers into trimers through formation of a triple-helical collagen I-type coiled-coil.

We hypothesize that the organizing principle by which the coiled-coil trimers are assembled into the hexameric and high-molecular weight oligomerization states is a coiled-coiled-coil. Our elastic coiled-coil model is well suited to modeling not only the collagen triple helix, but also the higher-order forms by treating the collagen triple helix itself as a single elastic filament. This work is a collaboration with Tsushuen Tsao.

With the exception of the globular headpiece, high-resolution structural information about this molecule has not been obtainable to date, due to a combination of intrinsic flexibility, glycosylation, and the difficulty of protein expression. We will use our methods to generate atomic coordinate models of the distinct states of this molecule, constrained by diverse types of biochemical data. These models will be used by the Tsao lab to design experiments to understand the determinants of signalling in the general context of diabetes research. In this project, we will combine several aspects from the previous sections of the grant.

The first step in the mathematical approach consists in modeling adiponectin as different collagen strands twined together. This provides a family of possible models depending on the geometric parameters (radii, twist, etc.). Purely geometric requirements on the possible forms already
strongly constrain the possible models. The second step is to identify within this large family of coiled structures, subfamilies for which collagen domains are complementary (see Figure 8). This is done by defining a suitable energy that can be minimized within the family of coils. The third and crucial step is to consider specific adiponectin molecules and use constraints from experimental data to refine the structure. The experimental data comes from various sources. The stoichiometry of the three states has been accurately determined by analytical ultracentrifugation and estimates of the radial and axial dimensions of the different species are available from electron micrographs. This information provides a basis for modeling the oligomers, but more subtle details such as the relative twists of the individual molecules in the oligomers or the juxtaposition of sidechains requires higher-resolution or site-specific information. This information will be obtained from footprinting and crosslinking studies mapped onto the mathematical models, in turn enabling the models to be adjusted to improve their match to the data.

The types of information to be included can be divided into dimensional constraints obtained by electron microscopy, surface exposure constraints obtained from footprinting, and distance constraints obtained from crosslinking. In addition we will use the electron micrographs obtained by the Tsao lab as a test of our methods for calculating radon transforms from curvatures and extracting curvatures from them. Since these diverse observations are subject to error, it is necessary to use statistical methods to reconcile these disparate observations. This requires both a way to quantify error distributions in the various types of individual measurements, and to devise algorithms for adjusting the model parameters to best fit all the data.

Figure 8. Models of the oligomerization states of adiponectin compared to electron micrographs Left: trimer (a single coiled-coil) Center: Hexamer (two trimeric coiled-coils wound around each other to form a coiled-coiled-coil) Right: Octadecamer (a distinct coiled-coiled-coil composed of 6 trimeric coiled-coils)

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