PROTEINS POCKETS ANALYSIS AND DESCRIPTION

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Keywords: Protein-ligand docking, Curvature analysis, Concavity tree, Travel depth, Pocket mouth area and perimeter.

Abstract: The development of computational techniques to guide the experimental processes is an important step for the determination of the protein functions. The purpose of the activity here described is the characterization of the active sites in protein surfaces and their quantitative representation. A few pocket parameters like volume, travel depth, mouth area and perimeter, amplitude parameters, interfacial area ratio, summit density and mean summit curvature are hierarchically accessible through a concavity tree that topologically represents the entire protein molecule. This structural representation is particularly useful for the evaluation of binding pockets, the comparison of the morphological similarity and the identification of potential ligand docking.

1 INTRODUCTION

When a novel experimentally determined protein is discovered, bioinformatics tools are used to screen the datasets of proteins with known functionalities, searching for candidate binding sites for the 3D structure of the new protein. Specifically, if a region in the surface of the novel protein has similarities to the binding site of a known protein the interactions of the latter are expected candidates in order to discover the unknown functionalities of the former protein.

The analysis of the binding sites of proteins and their detection, have been developed on the basis of different protein representations and matching strategies.

CAST (Liang et al., 1998) and CASTp (Binkowski et al., 2003) automatically locate and compute pockets volume and mouth opening area and circumference of pockets. The algorithm applies 3D computational geometry: a triangulation of the protein’s surface atoms (which covers a conglomerate of 3D tetrahedra) is analyzed through convex hull, alpha shapes and discrete flow theory. A tetrahedron having at least a triangle that cross the void region is designated as ‘empty tetrahedron’. Empty tetrahedra sharing a common triangle are grouped so ‘flowing’ towards neighboring larger tetrahedra which act as sink. A pocket, which is a potential binding site, is then obtained as collection of empty tetrahedra.

PASS (Brady and Stouten, 2000) is a purely geometrical method composed of several steps. First, the protein surface is completely covered by probe spheres. Second, each probe is associated with a “burial” value, which corresponds to the number of atoms contained within a sphere of radius 8 Å. Third, the probes with a “burial” value lower than a predefined threshold are eliminated. Fourth, these three steps are iterated (with step one applied only to the parts of the surface covered by the probes) until no new buried probe are added. Fifth, a probe weight, which is dependent to the number of their neighboring spheres and the extent to which they are buried, is assigned. Finally, a shortlist of active site points (ASPs), ranked by the probe weight, is selected by identifying the central probes in regions that contain many spheres with high burial count. These ASPs represent the loci of potential binding sites.

In POCKET (Levitt and Banaszak, 1992), the protein is mapped onto a 3D grid in which a grid point belongs to the protein molecule if it is within 3 Å from an atom coordinate. The pockets are characterized as a set of grid points, not belonging to the protein for which a scanning along the x, y, or z-axes presents a protein-solvent area-protein sequence (called protein-solvent-protein event).

LIGSITE (Huang and Schroeder, 2006) extends POCKET by scanning also along the four cubic diagonals in addition to the coordinates axis (in fact, the previous pocket’s definition produces classifications that change with the angle between the reference system and the protein). The grid points that present a
The number of protein-solvent-protein events greater than a threshold are candidate active sites.

SURFNET-ConSurf (Glaser et al., 2006), represents the pocket surface by combining geometrical features together with an evolutionary characteristics based on the degree of conservation of the amino acids involved. SURFNET-ConSurf is based on a two-stage process. In the former stage, through SURFNET (Laskowski, 1995) the potential binding sites in the protein surface are identified. The clefts are detected by placing a sphere between all pairs of atoms such that the sphere just touches each atom, then this sphere is progressively reduced in size up to no further intersection with other atoms are present. The resulting sphere is retained only if its radius is greater than a minimum size predefined. Once the clefts have been filled by spheres they can be individually described by geometric features (e.g., the volume). In the latter stage, the regions of the resulting clefts that are distant from highly conserved residues, as defined by the ConSurf-HSSP database (Glaser et al., 2005), are removed, thus reshaping the cleft volumes. The second stage is based on two parameters: the maximum allowed distance and the minimum conservation score cutoff. The remaining clefts are candidate active sites (in particular the largest ones).

In this paper, we just introduce a feature vector and describe in details a data structure in order to represent the pocket extracted by the various methods to improve the analysis and to reach the needed performance quality. The paper is organized as follows: in section two a short survey of features for pockets characterization is given; in section three a data structure to describe the protein’s pockets, based on a concavity tree representation, which allows the complete description, at different scales and abstraction levels, supporting at each level the feature vector description previously introduced, is presented. In section four a practical case is described. The last section five contains the conclusion and the near future subsequent activity.

2 FEATURES FOR POCKET DESCRIPTION

The goal of this paper is the characterization of each pocket, which is extracted by segmenting the protein surface, through morphological and topological quantitative descriptors. These features are enriched with local biochemical features (types of residues and their characteristics) to detect and specialize the active sites of a protein.

We are considering now a second step of the process of active sites identification, that is, we start knowing the segmentation of the protein surface in a number of pockets and tunnels. In particular, we will refer to the method given in (Cantoni et al., 2009c) but the technique described in the sequel is not limited to this particular solution.

2.1 Preliminary Statements

In the discrete space the protein is defined in a 3D grid (CG) of dimension L x M x N voxels. Note that the grid is extended one voxel beyond the minimum and maximum coordinate of the SES (Solvent-Expanded Surface, also known as the molecular surface or Connolly surface), generated by the envelope of a rolling sphere over the van der Waals atoms surface\(^1\) in each orthogonal direction, in this way both SES and Convex Hull (CH) borders are inside the CG border. The voxel resolution adopted is 0.25 Å, so as to be small enough to ensure that, with the used radii in biomolecules atoms\(^2\), any concave depression or convex protrusion is represented by at least one voxel.

The CH of a molecule is the smallest convex polyhedron that contains the molecule points. In \(\mathbb{R}^3\) the CH is constituted by a set of facets, that are triangles, and a set of ridges (boundary elements) that are edges. A practical O(n log n) algorithm for general dimensions CH computing, is Quickhull (Barber and Dobkin, 1996), that uses less space and executes faster than most of the others algorithms. Let us call \(R\) the region between the CH and the SES (the concavity volume (Borgefors and Sanniti Di Baja, 1996)), that is:

\[
R = CH \cap \overline{SES}
\]

and \(Re\) the connected component of \(R\) adjacent to the CH border; that is \(Re\) and \(R\) differ for the cavities \(C\): the volumes completely enclosed in the macromolecule \(M\):

\[
C = CH - Re - M
\]

The region \(Re\) has been partitioned into a set of disjoint segments \(P_{SES} = \{P_1, \ldots, P_j, \ldots, P_N\}\), where \(N\) is the number of pockets and tunnels. The partition must satisfy the following condition:

\[
P_i \cap P_j = \emptyset, i \neq j
\]

\[
P_1 \cup \cdots \cup P_j \cup \cdots \cup P_n = Re
\]

\(\text{\textsuperscript{1}}\)The radius of the solvent sphere is usually set to the approximate radius of a water molecule.

\(\text{\textsuperscript{2}}\)The smallest used atom is oxygen having a Van der Walls radius of 1.4 Å.
Our goal is to define a feature vector to represent, in a discriminant way to facilitate processing and statistical analysis, each pocket of $P_{SEs}$. Many vector’s parameters need a reference plane that we have identified with the one to which it belongs the largest CH’s triangle involved in the pocket.

2.2 Basic Features

The problem of defining an optimal set for feature selection is complicated because, besides building robust models, it is also important simplifying the amount of resources required to describe the data accurately, without ambiguity, in a very large set of redundant and relevant information. The expert can help, but can usually construct only a set of application-dependent features.

An extended set of general features that can be, in peculiar cases, partitioned in well-organized proficient subsets, is the following: (i) Pocket Volume (Laskowski et al., 1996), (ii) Surface to Volume Ratio, (iii) Skewness and Kurtosis of Height Distribution (Blunt and Jiang, 2003), (iv) Mouth Aperture (in details we consider area, perimeter and the perimeter to area ratio), (v) Travel Depth (Coleman and Sharp, 2006) (Giard et al., 2008), (vi) Top Peaks and Valleys, (vii) Summit Density, Mean Summit Curvatures (both the average of the principal curvatures of peaks and valleys (Coleman et al., 2005), (Cantoni et al., 2009a)), (viii) Interfacial Area Ratio, and (ix) Residue Conservation (Glaser et al., 2006) (the conservation score for each residue in a given protein can be obtained from the ConSurf-HSSP database (Glaser et al., 2005)).

3 THE DATA STRUCTURE

One of the most successful approaches for shape analysis and description is the structural one. We think that this is particularly fruitful in proteomics in which the morphology plays a fundamental role. A complex shape, like the Re, is segmented into its component (the pockets set), and each pocket can be subsequently decomposed into simpler region, and the complete description is given in terms of the region’s features and their spatial relationship. Nevertheless, pocket shapes can be rather complex and not directly decomposable into simpler regions. However we can re-apply the segmentation process of the Re into the pockets. This process can be executed recursively. In this way a sequence of approximations is built, and, at each stage, exact measures of the remaining concavities, based on the parameters described above, are given. This structural hierarchical description and analysis, guided by the concavities (Borgefors and Sanniti Di Baja, 1992), seems to us a very promising effective description.

The basic structure of the approach was firstly introduced in (Arcelli and Sanniti Di Baja, 1978) and in (Borgefors and Sanniti Di Baja, 1996) has been finally called concavity tree: “components of C (Re in our 3D case) for which the internal section of the perimeter (surface in 3D) exceeds the external section are structured concavities. A more sophisticated analysis of these regions is performed to extract further features. The envelopes CH of the concavity regions are computed using the same process as that applied to the original pattern. Merging can occur while filling meta-concavities, so the concavity regions must be labeled and processed individually. For each concavity region, its meta-concavities are identified. The process continues until all regions are convex”.

The final result is a hierarchical structure, the (meta) concavity tree. At each level the concavities can be analyzed and described on the basis of the above feature vector - computed at each node: obviously the features defined for concavities can also be computed for the meta-concavities.

The “concavities” (three “pockets” and one “tunnel”) and four second level meta-concavities of a 2D example are shown in Figure 1. Figure 2 shows concavities and meta-concavities of level two, three and fourth for the tunnel of level one. The corresponding concavity tree is shown in Figure 3. Note that terminating nodes, i.e. regions without significant concavities, are highlight with a bordeaux contour.

Figure 1: A 2D representation example with a tunnel and three pockets in a section composed of three connected components (in brown). The closed curve in black corresponds to the first level convex hull, and the border in brown dotted-dashed line embodies the area under analysis. Part of the second level with three meta-concavities (A, B, C) is shown; in evidence also five third level termination-node components (A1, A2, C1, D1, and D2).
Figure 2: Continuing the 2D representation example of Figure 1, the details of the tunnel B component (light green) are shown. The second level is composed of the three meta-concavities (B1, B2, and B3). Each one has concavities at the third level: B1 has a termination-node concavity (B12) and a second component B11 which maintains a meta concavity at the fourth level B111; B2 has a termination-node concavity (B22) and a second component B21 which maintains a meta concavity at the fourth level B211; finally, B3 has three node concavities (B31, B32, B33) each one maintaining a meta-concavity node-component B311, B321 and B331 respectively, at the fourth level.

Figure 3: Representation of the concavity tree for the 2D section of Figures 1 and 2. Note that each node contains the information of the feature vector previously presented.

Figure 4: On the top the 'space filling' representation of 1MK5. The colours follow the standard CPK scheme. On the bottom the correspondent secondary structure representation.

4 EXAMPLE OF A PRACTICAL REPRESENTATION

A practical example of our hierarchical representation is given in figure 4 referring to the Apostreptavidin Wildtype Core-Streptavidin with Biotin structure (1MK5 in PDB).

As mentioned above, the analysis is accomplished with a resolution of 0.25 Å, which entails a van der Waals radius of more than five voxels to the smallest represented atoms. The representation is related to the SES obtained from the quoted surface, after the execution of a closure operator, using a sphere with radius of 1.4 Å, approximately 6 voxels, (corresponding to the conventional size of a water molecule) as structural element (see figure 5).

The segmentation of concavities and meta-concavities is computed with the technique given in (Cantoni et al., 2009b), where the two parameters that characterize the execution, the minimum passage section $\theta_1$ and the maximum mouth aperture $\theta_2$ have been fixed to 200 voxels (which for a circle corresponds to a radius of about 8 voxels) and 2000 voxels (which for a circle corresponds to a radius of about 25 voxels).

Figure 6 represents the complete concavity tree for the 4 main pockets and tunnels. The molecule is represented inside its own convex hull which is drawn completely on the backside and only through its edges in the front side.

This process is conducted on the basis of other two thresholds: $\theta_3$ represents the minimum concav-
5 CONCLUSIONS

As morphology is important for protein recognition and function, the development of shape representa-
Figure 8: Complete concavity tree of the five main pockets of the 1MK5.

calculation of a subset of relevant features for building robust models. The exclusion of redundant features, by reducing the dimension of the parameter space, improves the time performance and facilitates fast searching, processing and statistical analysis.

REFERENCES


