eProS—a database and toolbox for investigating protein sequence–structure–function relationships through energy profiles

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ABSTRACT

Gaining information about structural and functional features of newly identified proteins is often a difficult task. This information is crucial for understanding sequence–structure–function relationships of target proteins and, thus, essential in comprehending the mechanisms and dynamics of the molecular systems of interest. Using protein energy profiles is a novel approach that can contribute in addressing such problems. An energy profile corresponds to the sequence of energy values that are derived from a coarse-grained energy model. Energy profiles can be computed from protein structures or predicted from sequences. As shown, correspondences and dissimilarities in energy profiles can be applied for investigations of protein mechanics and dynamics. We developed eProS (energy profile suite, freely available at http://bioservices.hs-mittweida.de/Epros/), a database that provides ~76000 pre-calculated energy profiles as well as a toolbox for addressing numerous problems of structure biology. Energy profiles can be browsed, visualized, calculated from an uploaded structure or predicted from sequence. Furthermore, it is possible to align energy profiles of interest or compare them with all entries in the eProS database to identify significantly similar energy profiles and, thus, possibly relevant structural and functional relationships. Additionally, annotations and cross-links from numerous sources provide a broad view of potential biological correspondences.

INTRODUCTION

The amino acid sequence-based predictions of protein structure features, stability analyses of known protein structures as well as secondary structure predictions are important tasks in protein modelling (1). Several energy functions and force fields that model the protein free-energy landscape have been developed to address these protein modelling problems. On the one hand, they contribute to protein modelling (i.e. comparative modelling, threading or ab initio folding) and protein model assessment. On the other hand, force fields are essential in molecular simulations and can account for the understanding of dynamics in molecular systems (2). They can also help to comprehend the relations between protein structure and function.

Energy models can be based on first principles approaches using physics laws. In addition, statistical analyses of experimentally derived structures form the basis for the development of so-called knowledge-based energy potentials (2–4). Although the approaches for computing knowledge-based energy potentials are simplified, they can reproduce experimental data with a high level of accuracy if adapted to a specific problem.

For example, elastic network models use simplified coarse-grained interaction models and have proven themselves to accurately determine protein dynamics (5,6). In general, the continuous application of coarse-grained interaction models is because of the reduction of system complexity and, thus, computational demands.

In 2006, Kozielski and colleagues (7) proposed that the sequences of energy values, so-called energy profiles, derived from protein structures by using potential functions can be compared using modified Needleman and Wunsch (8) and Smith and Waterman (9) alignment procedures. They have shown that pairwise comparisons and detected energy profile similarities can lead to the identification of proteins assigned to the same protein families. Additionally, conformational modifications as a result of enzymatic reactions or, in general, protein–environment interactions can be inspected (7,10,11). These studies substantiate the possible fields of application of energy profile-based methods. However, to allow large-scale or
even databank-wide investigations, the generation of large data sets is required. Because of the semi-automatic computation and error-prone nature often implicated by all-atom-based models, generating data sets on a large scale, for example, comparable with the Protein Data Bank (PDB) (12), becomes difficult. This holds especially if physics-based approaches are used as proposed by Kozielski and colleagues (7,10,11).

To allow large-scale energy profile-based analyses, we have developed eProS (energy profile suite), a database and toolbox for energy profile-based studying and comparing sequence–structure–function relationships and protein stability. Energy profiles are derived by using a straightforward coarse-grained energy model, which is suitable for globular and α-helical membrane protein structures. In the process of energy profile calculation, spatial and physicochemical information is integrated. As a result, energy profiles can be interpreted as protein-specific representations (see the Supplementary Data for further details). For example, the eProS output of the human angiogenin variant H13A (PDB ID 1b1j) is illustrated in Figure 1. The sequential visualization of an energy profile (Figure 1B and D) is the most intuitive. However, energy-to-structure mappings, as shown in Figure 1C, can contribute to identify low-energetic and, thus, stabilizing regions and properties in protein structures.

Currently, eProS stores 74 900 pre-calculated energy profiles derived from experimental globular protein structures and ~1300 pre-calculated profiles of α-helical membrane protein structures.

The eProS toolbox and the underlying eProS database provide various ways of visualizing, downloading and accessing energy profile data. The toolbox also includes database-wide searching for similar energy profiles. Here, the query energy profile can be defined by specifying a structure by PDB ID, by uploading a structure in PDB format from which the query energy profile is generated or by uploading an energy profile file that, for example, has been retrieved from the eProS database. Additionally, an amino acid sequence can be used as input. Starting from this, an energy profile prediction algorithm is used, leading to an energy profile that can be used for database-wide searching. The best matching hits are visualized by the eProS toolbox. Various sources of annotation [e.g. Gene Ontology (GO) (13), PDB, CATH (14), SCOP (15) and Pfam (16)] provide a wide view on structural and functional features of the best hits, which can be further broadened through the reverse annotation lookup provided by eProS. The reverse annotation lookup lists all energy profiles that match with the annotation specified by the user. For example, energy profiles of all proteins sharing the same structural topology or molecular function can be investigated concerning common energetic features that point to their similar structure or function. Thus, starting from a protein structure or sequence, estimations about correspondences of protein function and structural features can be drawn from these results and annotations.

DATABASE AND SEARCH TOOL DESCRIPTION

Content and data organization

At present, eProS supplies energy profiles for ~76 200 PDB entries that are internally separated into 74 900

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**Figure 1.** In this figure, the eProS output of the human angiogenin variant H13A (PDB ID 1b1j) is depicted. In (B) and (D), the energy profile is shown as a sequence of energy values. Colouring schemes for energy-to-structure mappings (C and D) and structure-to-sequence mappings (A and B) are provided. The energy colouring is discretized by assigning each energy value to one of the 4-quantiles in the energetic spectrum derived from the eProS database. This measure is because of visualization and performance purposes. The colour mappings provide insights that can contribute to identify low-energetic and, thus, stabilizing regions and properties in protein structures.
globular protein and 1300 α-helical transmembrane protein energy profiles. The corresponding PDB flat files, containing protein structure information, are stored in a local directory on the Web server hosting eProS. Based on these files, energy profiles for each available PDB entry have been pre-computed. Energy profiles are stored in files of a specifically tailored format. The following file formats have been defined and are available for download and analyses:

- *.ep: Tab character separated files with each row containing five columns (chain identifier, PDB residue index, amino acid one-letter code, secondary structure assignment and energy value). The first two lines are reserved for listing the PDB ID and the header row.
- *.ep2: Extended *.ep file. Each line represents a record, whereat the first four characters specify the record type. Record fields are tab character-separated. The following record types are currently defined: ‘NAME’ (PDB ID), ‘TYPE’ (‘TM’ for α-helical transmembrane protein structures, ‘nTM’ for globular protein structures), ‘HEAD’ (header row), ‘ENGY’ (energy value for a single residue, the five record fields correspond to the five columns of a row in a *.ep file) and ‘REMK’ (indicating a comment line).
- *.eps: Binary files. This file format is going to be used in upcoming standalone software applications. It is only in use in server-internal routines at present.

For each protein structure in the local PDB file repository, an energy profile has been saved in each of these three file formats.

The annotation entries displayed on the detailed view of a protein are retrieved from internal relational databases at runtime. All information provided by the database has been obtained from external sources and related databases. For an overview of the integrated data and their sources see Figure 2. The data integration has been achieved as follows:

**TMDET prediction**

TMDET is a service that implements a neural network-based method for the prediction of membrane spanning regions in 3D structures (17). For each α-helical membrane protein structure present at eProS database, a prediction has been computed. These predictions are essential for deriving an energy profile from α-helical membrane protein structures by using the coarse-grained model (for details see the elucidations given in the Supplementary Data). Additionally, if the user has uploaded an α-helical membrane protein structure for analyses, TMDET is applied for predicting the location of the membrane bilayer, and according to these predictions, the energy profile is generated.

**Pfam classification**

The current release of the Pfam database is available in terms of an SQL dump (16). As for the annotation retrieval, only two tables are required, ‘pfama’ (~2200 rows), containing the Pfam classifications, and ‘pdb_pfama_req’ (~110000 rows), mapping PDB IDs to their corresponding classifications.

**SCOP classification**

The SCOP classification releases are not provided in terms of an SQL database dump but as character-separated value files instead (15). According to this data, the following tables have resulted: ‘des’ (~144000 rows) contains descriptions for all SCOP classifications, ‘hie’ (~144000 rows) represents the SCOP classification hierarchy and ‘cla’ (~111000 rows) assigns PDB IDs to SCOP classifications.

**CATH classification**

Two files of the current CATH database release (14) served as source of annotation data, ‘CathDomainList’ and ‘CathNames’. The resulting tables ‘domains’ (~153000 rows) and ‘names’ (~3900 rows) provide information about assigned CATH domains and names of CATH classification nodes, respectively.

**GO term annotation**

Of the 44 tables found in the images of the GO database (13), the following are required to gain the GO terms associated with a protein structure: ‘term’ (~37000 rows) containing the GO terms, ‘gene_product’ (~1300000 rows) containing gene products and ‘species’ (~89000 rows) containing species the gene products originate from. The ‘association’ table lists (~77000000 rows) assignments of GO terms to gene...
products. To find GO terms matching to a PDB ID, the name of each macromolecule of the entry and their sources (NCBI Tax ID) are required. Therefore, two auxiliary tables have been created (~122,000 rows and 108,000 rows, respectively). However, searching for GO terms and especially performing reverse annotation lookups had caused unacceptable response times because of complex query statements that had been originated in the database development. For performance improvements, an additional table (~610,000 rows) has been created, which assigns the GO terms to each protein directly. Thereby a speedup (>5 min to 100 ms) has been achieved for reverse annotation lookup.

Working with the eProS database

The eProS and the collection of energy profiles are freely available to the scientific community in a separate download section (accessible through the ‘Dataset’ link at the eProS homepage). In the download section, it is possible to browse the database and inspect energy profiles of interest. For this purpose, eProS provides the access of energy profile data through flat HTML page tables or by automated download programs, such as wget (http://www.gnu.org/software/wget/) or similar software. This ensures large-scale downloading of energy profile files and high-throughput analyses. The eProS toolbox permits accessing the data by more sophisticated energy profile visualizations, for example, plotting of energy profiles and viewing the protein structure of interest highlighted with energy value-based colouring schemes (Figures 1C and 3). Cross-links and annotations retrieved from various foreign sources (Figures 2 and 3B) are available. From these annotations, the reverse annotation lookup can be accessed, and, subsequently, energy profiles of proteins matching the user-specified annotation are listed (e.g. Figure 3C). As an example, after querying the N-terminal domain of the riboflavin synthase by specifying its PDB ID (1pkv) at the eProS home page (Figure 3A), the corresponding energy data and structure as well as the related annotations are listed (Figure 3B). Reverse annotation lookup is accessed by clicking the annotation of choice, which leads to the list of energy profile data available at eProS that share the specified annotation, in this case riboflavin synthase activity (Figure 3C).

Further methods have been implemented and integrated into the toolbox that allow energy profile analysis, calculation, sequence-based prediction and a database-wide searching for identical or similar energy profiles. An overview of these tools and the implemented data flow is
given in Figure 5. The following elucidations explain these tools briefly:

**eAlign**
This tool provides modified Needleman Wunsch (8) and Smith-Waterman-like (9) alignment procedures for computing pairwise energy profile alignments. Generated alignments are presented as graphs, ASCII-formatted texts as well as dotplots in which energetic identities and similarities are highlighted. In addition, eAlign computes a so-called distance Score (dScore). The dScore gives a hint about the energy profile similarity observed in the alignment (see the Supplementary Data for details).

**eCalc**
eCalc provides the energy profile computation. On the one hand, a PDB ID can be specified, and the corresponding energy profile data are displayed if they are present in the database. If this is not the case, the corresponding coordinates are retrieved from the PDB, and the energy profile is computed. On the other hand, the user can upload a protein structure from which the energy profile is generated, subsequently. The output can be investigated, downloaded for further analyses or reused as input for the eProS toolbox.

**eGOR**
Adopted to the concepts and implementations of protein secondary structure prediction (e.g. GOR I–V) discussed by Garnier and colleagues (18–20), eGOR applies information theory-based methods and knowledge derived from known energy profiles to predict a residues energy value according to its sequence neighborhood composition. This algorithm is the basis for the eGOR tool, and it allows the prediction of an energy profile starting from a user-specified sequence.

**eMut**
This tool visualizes the energetic similarities and dissimilarities of proteins of the same length. Thus, analysis of, for example, (point-)mutated proteins, structure trajectories obtained from molecular dynamics or coarse-grained dynamics simulations or influences of temperature variations on protein stability is supported by eMut.
structures of unknown function can be facilitated.

In the process, pairwise alignments of the query energy profile to all entries of the specified entry set (e.g. globular proteins or \( \alpha \)-helical membrane proteins) are generated. From each alignment, the corresponding dScore is heuristically computed and recorded. This process requires \(~3\) h of computation for querying an average-sized protein structure (\( \approx 120 \) amino acids) to the set of globular protein energy profiles. Because of the time demands, the user has to specify a valid e-mail address to run an eSearch query. After the computation has finished, an e-mail is sent to the user, which provides a link to the result session as well as a session id. The results are presented as an interactive list ranked according to the derived dScores. An example of using eSearch is illustrated in Figure 4. In this case, the energy profile of Trypanosoma brucei brucei thioredoxin (PDB ID 1r26) has been queried, and numerous similar energy profiles have been identified (Figure 4A and B). As a representative example for the general observations that can be made from this query, the energy profile alignment to the ninth match (PDB ID 3h/hv) indicates numerous global energetic correspondences (Figure 4C). As shown, the best matching energy values are located in the first helix and second strand in both structures. By using the reverse annotation lookup of functional annotations (e.g. ‘cell redox homeostasis’ and ‘glycerol ether metabolic process’) and structural annotations (e.g. ‘glutaredoxin’) of 1r26, corresponding energy profile entries are listed. Note that most proteins present in the reverse annotation lookup list are reported as best-matching energy profiles. As 3h/hv has not been annotated by GO-terms; yet, it can be proposed that the GO-terms associated to the best matches can be applied for annotating 3h/hv. Furthermore, integrating the profiles reported by eSearch and reverse annotation lookup to the analyses, the energetic properties can be identified that are responsible for stabilizing the fold. For example, mainly low-energetic residues can be found in the second strand (Figure 4C). In contrast, the third strand is consisting of residues with alternating energy values. Both observations are in agreement in all proteins sharing this topology. On the other hand, functional energetic features might be basically corresponding to residues located in the first helix and second strand, as these residues are found to be energetically conserved in all energy profiles listed by the functional reverse annotation lookup.

In a similar way, the functional clarification of protein structures of unknown function can be facilitated.
Future developments of eProS are going to include the improvement of time-performance of eSearch. Additionally, implementing cross-links between energy profile data will provide a more user-friendly data access. Furthermore, an automated energy profile and annotation retrieval system is currently work in progress. This system is going to be capable of updating the eProS database automatically on a weekly or monthly basis. At the moment, an approach for predicting the topology of an α-helical membrane protein based on its predicted energy profile is under evaluation and will be integrated to the toolbox.

**SUPPLEMENTARY DATA**

Supplementary Data are available at NAR Online: Supplementary Information, Supplementary Figure 1 and Supplementary References [22–27].

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