Structural bioinformatics

ProtPOS: a python package for the prediction of protein preferred orientation on a surface

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Abstract

Summary: Atomistic molecular dynamics simulation is a promising technique to investigate the energetics and dynamics in the protein–surface adsorption process which is of high relevance to modern biotechnological applications. To increase the chance of success in simulating the adsorption process, favorable orientations of the protein at the surface must be determined. Here, we present ProtPOS which is a lightweight and easy-to-use python package that can predict low-energy protein orientations on a surface of interest. It combines a fast conformational sampling algorithm with the energy calculation of GROMACS. The advantage of ProtPOS is it allows users to select any force fields suitable for the system at hand and provide structural output readily available for further simulation studies.

Availability and Implementation: ProtPOS is freely available for academic and non-profit uses at http://cbbio.cis.umac.mo/software/protpos

Supplementary information: Supplementary data are available at Bioinformatics online.

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1 Introduction

Understanding the mechanisms of protein adsorption on surfaces is important for many biotechnological applications, such as the design of medical implants, protein biochips, drug delivery systems and point-of-care devices, etc. By virtue of molecular modeling and simulations, the protein adsorption processes, its dynamics and subsequent protein–surface interactions can be studied at the molecular or atomic level of detail. However, a successful simulation of the adsorption process depends largely on the initial protein orientation relative to the surface (Wei et al., 2012). To avoid sampling protein trajectory which will eventually fail to adsorb, a work-around is to firstly determine the preferred orientations of the protein and use them as starting structures in molecular simulations.

Essentially, a search for the low-energy protein orientation on a given surface is an optimization process which includes a conformation generation strategy and a scoring function. For the former, a systematic rotation (and translation) of the protein or a Monte Carlo simulation is commonly used (Hsu et al., 2008; Xie et al., 2010; Zheng et al., 2004). For this structure determination problem, a simplified scoring function accounting for only protein–surface interactions is usually preferred for higher computational efficiency. Interaction parameters can be taken from an empirical force field or modification from the scoring function of a docking method (Sun et al., 2005). Interestingly, unlike other structure prediction problems such as protein–ligand docking or protein–protein docking where numerous tools exist to simplify the prediction task, there are only limited choices available for fast protein-surface structure prediction. One such option is RosettaSurface (Makrodimitris et al., 2007) which is included in the Rosetta molecular modeling package, and hence the use of RosettaSurface entails a deep learning curve. Other prediction methods mentioned above, to our knowledge, are unavailable publicly for general use.

Hence, to fill this gap, we have developed a self-contained, lightweight and easy-to-use software package called prediction of PROTein Preferred Orientation on a Surface (ProtPOS) based on the protein-surface docking method described in (Ngai et al., 2015). It combines particle swarm optimization (PSO) algorithm (Kennedy and Eberhart, 1995) with energy minimization to perform global and local search over six degrees of freedom of the protein translation and rotation on
a given surface. The advantage of ProtPOS is that it allows users to select any force fields suitable for the system at hand, to be provided as inputs in the format of GROMACS topology together with the protein and surface coordinates. In this way, predicted low energy protein conformations are compatible to the selected force field and readily available for further molecular dynamics (MD) simulations.

2 Implementation

ProtPOS is a package of open-source shell scripts and Python programs utilizing the PyMOL library for conformation generation and the popular MD simulation software GROMACS for energy minimization and scoring. The workflow of ProtPOS is explained in Fig. 1. To facilitate ease-of-use, a template wrapper script is provided through which users can customize parameters for the search algorithm, perform prediction and clustering analysis. During the search, all rotational angles around the protein center of mass are allowed. For surfaces with repeating surface patterns, lateral translation (X and Y directions) of the protein can be restrained to defined ranges, typically the dimensions of the surface unit cell; whereas the translation along the surface normal (Z) is set to be [1.0, 5.5] Å from the surface, i.e. the interfacial region where protein-surface interactions are significant. If contacting residues are known a priori, they can be given to ProtPOS to limit further the search on the specified protein surface. It should be mentioned that, for better computational efficiency, the Generalized Born implicit solvent model (GBSA) is used during energy minimization and only nonbonded interactions between the protein and the surface are included in scoring.

3 Validation and performance

ProtPOS was tested by predicting the initial adsorption orientations of lysozyme on a hydrophobic surface of perfluorodecane molecules. The results were compared to experiments and previous computational studies qualitatively and found good agreement (See Supplementary Table S1). Furthermore, the selected low-energy structures were subjected to 50 ns MD simulations in order to assess the stability of the protein orientations. In three out of four cases, the proteins maintained the predicted orientations and stably attached to the surface (Supplementary Table S2) whereas in one case the protein was rotated slightly keeping three contacting residues same as in the initial orientation and remain stably attached throughout the remaining simulation time. MD simulations of randomly selected protein orientations either resulting in the protein converted to predicted orientation or diffused away from the surface, suggesting that ProtPOS predicted structures serve as good starting structures for further simulation studies.

Regarding the time performance, our test case running on a consumer-grade desktop computer (with Intel i7-4790 3.6 GHz CPU and 16 GB memory) took about 1 day to complete one ProtPOS run, with an average of 81.5 iterations to reach convergence in the PSO search. The most expensive calculation is the energy minimization step which is done for all PSO particles in each iteration. Future improvement on ProtPOS should consider pre-filtering of highly unfavorable orientations before performing energy minimization, as well as parallelizing the PSO algorithm by multicore CPU and advanced GPU techniques.

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