pTARGET: a web server for predicting protein subcellular localization

Chittibabu Guda

Gen*NY*Nsis Center for Excellence in Cancer Genomics and Department of Epidemiology and Biostatistics, University at Albany, State University of New York, 1 Discovery drive, Rensselaer, NY 12144-3456, USA

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ABSTRACT

The pTARGET web server enables prediction of nine distinct protein subcellular localizations in eukaryotic non-plant species. Predictions are made using a new algorithm [C. Guda and S. Subramaniam (2005)] pTARGET [corrected] a new method for predicting protein subcellular localization in eukaryotes. Bioinformatics, 21, 3963–3969], which is primarily based on the occurrence patterns of location-specific protein functional domains in different subcellular locations. We have implemented a relational database, PreCalcDB, to store pre-computed prediction results for all eukaryotic non-plant protein sequences in the public domain that includes about 770 000 entries. Queries can be made by entering protein sequences or by uploading a file containing up to 5000 protein sequences in FASTA format. Prediction results for queries with matching entries in the PreCalcDB will be retrieved instantly; while for the missing ones new predictions will be computed and sent by email. Pre-computed predictions can also be downloaded for complete proteomes of Saccharomyces cerevisiae, Caenorhabditis elegans, Drosophila, Mus musculus and Homo sapiens. The server, its documentation and the data are accessible from http://bioinformatics.albany.edu/~ptarget.

INTRODUCTION

Protein subcellular localization is a key functional characteristic of proteins. Subcellular localization of proteins in appropriate compartments is vital for the internal structure of the cell itself and for its functional integrity. Determination of the subcellular localization by experimental means is not practical for all proteins owing to time and cost constraints. Alternatively, several computational methods have been developed for the prediction of subcellular localization in eukaryotic proteins. These methods are broadly categorized into four classes. (i) Methods based on sorting signals relying on the presence of sorting signals that are recognized by location-specific transport machinery to enable their entry (1–3). (ii) Methods based on the differences in amino acid composition (AAC), pseudo-AAC (includes AAC plus sequence order) or amino acid properties of proteins from different subcellular locations (4–6). (iii) Methods based on lexical analysis of keywords (LOCkey) in the functional annotation of proteins (7). (iv) Methods using phylogenetic profiles (8) or domain projection method (9). Nevertheless, only a handful of methods are accessible online to the research community. We compiled a list of currently available web servers that offer access to web-based prediction of subcellular localizations for eukaryotic proteins (Table 1). Some of these methods can predict only a few types of locations owing to inherent limitations while some lack the robustness to handle the heterogeneity expected in eukaryotic proteomes. Moreover, some web servers are designed to process only one sequence or a limited number of sequences in batch queries, thus, limiting their use for proteome-wide predictions (Table 1).

Recently, we developed two prediction methods: MITOPRED, for predicting nucleus-encoded mitochondrial proteins (10,11) and pTARGET, for predicting nine distinct subcellular locations in eukaryotic proteomes (12) based on location-specific functional domains and AAC. pTARGET method is relatively robust for proteome-wide predictions since it does not rely on the presence of a signal or target peptides. Based on this method, here we present the pTARGET web server that can process proteome-scale queries, backed by a relational database, PreCalcDB, containing pre-computed predictions.

DESIGN AND IMPLEMENTATION

The pTARGET server has been designed using PERL-CGI interface to process user queries and display or email the...
prediction results. A relational database, PreCalcDB containing pre-computed predictions has been developed to back the web server and to provide instant access to predictions for most of the eukaryotic protein sequences in the public domain. Perl DBD module was used to interface with the MySQL database. Query sequences are first searched against this database and predictions will be retrieved for matching entries; while for others, a new prediction process will be launched. The new prediction process includes searching the Protein family database (Pfam database, http://pfam.wustl.edu), which is the most time-consuming step in the prediction process. Pre-calculated prediction results are instantly displayed on the screen while those from new predictions are emailed to the user upon completion of the computation steps.

Algorithm
The pTARGET method (12) predicts proteins targeted to nine distinct subcellular locations in eukaryotic non-plant species. This prediction algorithm calculates two distinct scores, i.e. first, a score based on the presence or absence of location-specific Pfam domains (Pfam score) and second, a score based on the relative amino acid weights calculated from AAC (AAC score). The nine subcellular locations predicted by pTARGET include cytoplasm, endoplasmic reticulum, extracellular/secretory, golgi, lysosome, mitochondria, nucleus, plasma membrane and peroxisome.

Pre-computed prediction database
To expedite the response time, pre-computed predictions have been provided for all non-redundant eukaryotic protein sequences (excluding plant sequences) in the public domain (~770 000 sequences). We have created a relational database, PreCalcDB, using the open source database MySQL 4.0 (downloaded from http://dev.mysql.com). PreCalcDB contains several relational tables to store protein sequences, headers and pTARGET prediction results. Protein sequence strings are treated as primary keys that are indexed to sequence accession IDs and to the prediction results. Programs for database development were written in SQL and supporting programs for accessing and manipulating the database were implemented in JAVA using the JDBC (Java Database Connectivity) API. Query sequences are searched against those in the PreCalcDB and for matching entries, predictions are retrieved and displayed instantly in the browser window. For the missing entries, the response time depends on the number of sequences requiring new predictions, which is ~1 h for 60 sequences on the current server. If approved by the user, new query sequences and the prediction results will be automatically loaded back to the PreCalcDB to make them available to the next user. Since PreCalcDB stores comprehensive sets of protein sequences from major databases including Swiss-Prot and TrEMBL from EBI and the ‘nr’ database from GenBank, pre-computed predictions are available for the majority of user queries. Predictions for complete proteomes of important eukaryotic species including yeast (Saccharomyces cerevisiae), nematode (Caenorhabditis elegans), fruit fly (Drosophila melanogaster), mouse (Mus musculus) and human (Homo sapiens) can also be downloaded from the web server.

Input and output formats
Users can enter protein sequences in the text box or upload a file containing up to 5000 protein sequences in FASTA format.
Results will be displayed on the screen and emailed in plain text format. Users should be aware that some spam filtering programs installed on the user’s mail client could sort the emails from pTARGET server into a SPAM folder. As shown in the screen shot (Figure 1), each prediction is followed by a prediction confidence value. Prediction confidence value is calculated as the ratio of calculated score to the total score required to make a true prediction, and it is expressed in percentage. For example, a score of 50 is required to make a prediction with 100% confidence. If the calculated score for a query sequence is 45, the prediction confidence is 90%. All query sequences with calculated scores equal to or exceeding 50 are predicted with 100% confidence.

**DISCUSSION**

pTARGET web server is intended for performing genome-scale prediction of protein subcellular localizations in eukaryotic organisms excluding plant species. Since several metabolic pathways and organelles in plants are not the same as in animals, the distribution of protein functional domains in these two systems is different. Similarly, bacterial species do not possess all the subcellular locations as eukaryotic cells do. Hence, predictions with this web server should be used only in the context of eukaryotic non-plant proteomes. Prediction capabilities of other similar web servers are limited to a few types of locations in some cases while some can process only one query sequence at a time (Table 1), making them unsuitable for genome-scale predictions. Moreover, the underlying prediction algorithm (12) for this web server is primarily based on the Pfam domain occurrence patterns and hence is more robust than some other methods that require the presence of a signal or target peptide to make an accurate prediction. Since pTARGET predictions are compute-intensive, we implemented the PreCalcDB as a back-end server to this web resource to significantly improve the response time. One limitation of pTARGET method is that proteins containing Pfam domains that exist in multiple subcellular locations or those without Pfam annotations are predicted solely based on their amino acid composition resulting in reduced prediction accuracy. However, over 70% of the protein sequences in the public domain are currently covered with Pfam annotations and this coverage is rapidly expanding. Hence, the prediction accuracy of pTARGET is expected to improve as more Pfam domains and more information on subcellular localization become available. We have implemented a self-enriching feature for the PreCalcDB which, with user’s approval, can store the newly calculated predictions in the relational database to make them instantly accessible to another user. We will update the PreCalcDB every 6 months to provide the current and most accurate information to the research community.

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