Sequence analysis

CSS-Palm: palmitoylation site prediction with a clustering and scoring strategy (CSS)

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ABSTRACT

Summary: Palmitoylation is an important post-translational lipid modification of proteins. Unlike prenylation and myristoylation, palmitoylation is a reversible covalent modification, allowing for dynamic regulation of multiple complex cellular systems. However, in vivo or in vitro identification of palmitoylation sites is usually time-consuming and labor-intensive. So in silico predictions could help to narrow down the possible palmitoylation sites, which can be used to guide further experimental design. Previous studies suggested that there is no unique canonical motif for palmitoylation sites, so we hypothesize that the bona fide pattern might be compromised by heterogeneity of multiple structural determinants with different features. Based on this hypothesis, we partition the known palmitoylation sites into three clusters and score the similarity between the query peptide and the training ones based on BLOSUM62 matrix. We have implemented a computer program for palmitoylation site prediction, Clustering and Scoring Strategy for Palmitoylation Sites Prediction (CSS-Palm) system, and found that the program’s prediction performance is encouraging with highly positive Jack-Knife validation results (sensitivity 82.16% and specificity 83.17% for cut-off score 2.6). Our analyses indicate that CSS-Palm could provide a powerful and effective tool to studies of palmitoylation sites.

Availability: CSS-Palm is implemented in PHP/PERL+MySQL and can be freely accessed at http://bioinformatics.lcd-ustc.org/css_palm/

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Supplementary information: Supplementary data are available at Bioinformatics online.

INTRODUCTION

Many proteins are post-translationally modified by the addition of a palmitate molecule to a cysteine by thioesterification. Biochemically, palmitoylation enhances the surface hydrophobicity of protein substrates and promotes their interactions with membranes (Kleuss and Krause, 2003). Thus, some hydrophilic proteins like Ras and G proteins make use of this modification to attach themselves to membranes, often in combination with prenyl groups. Palmitoylation can also regulate intracellular trafficking (Kang et al., 2004), sorting (Schneider et al., 2005), subcellular localization (Van Itallie et al., 2005) and functional activities of the proteins (Sudo et al., 1992). Although palmitoylation is increasingly recognized as a frequent and important modification of eukaryotic signaling proteins, the molecular mechanism underlying protein palmitoylation remains elusive.

Identification of palmitoylation sites could provide an effective approach to understand the molecular mechanism of palmitoylation. To date, only a few palmitoylation sites have been identified experimentally, mainly through mutagenesis studies of candidate cysteine residues using conventional biochemical methods. The distinguishing features of palmitoylation sites had not been well-characterized, and most previous studies suggested that there is no canonical consensus sequence/motif for the palmitoylation sites (Bijlmakers and Marsh, 2003; el-Husseini Ael and Bredt, 2002; Linder and Deschenes, 2003; Smotrys and Linder, 2004; ten Brinke et al., 2002). Just as in the phosphorylation site prediction (Zhou et al., 2004), we propose that such a pattern may consist of multiple consensus motifs.

METHODS

Based on the above hypothesis, we present a novel computational method/software CSS-Palm—Palmitoylation Site Prediction using a Clustering and Scoring Strategy. We have collected 210 experimentally verified palmitoylation sites from 83 distinct proteins as training set, and grouped them into several subsets based on their sequence similarity, and the common characteristics of each subset are described by the high sequence similarity between the palmitoylation sites. For each given cysteine residue, its final score as a potential palmitoylation site is defined as the highest similarity score among all similarity scores against the above partitioned subsets. A detailed description of the algorithm can be found in the Supplementary Material. The sequence logo (WebLogo, Crooks et al., 2004)
of each of the subsets (Supplementary Fig. S1, S2 and S3) indicates that such strategy achieves better specificity over the whole set of collected palmitoylation sites (Supplementary Fig. S4). Its prediction performance on the curated dataset is highly encouraging with Jack-Knife sensitivity of 82.16% and specificity of 83.17%, respectively. To facilitate its application, we have developed an easy-to-use web server that is freely accessible from http://bioinformatics.lcd-ustc.org/css_palm/

APPLICATION

To explain how to use our CSS-Palm web server, we choose human Centromeric protein E (CENP-E, Q02224), a mitotic kinesin, as an example. Human CENP-E protein, as an important component of Kinetochore protein complexes, is localized in outerplate kinetochores during mitosis, attaches spindle microtubules to centromeres and orchestrates the fidelity of chromosomal segregation (Yao et al., 1997, 2000).

We first retrieved the primary sequences of both human and mouse CENP-E from ExPASy and pasted both the sequences in FASTA format into the textbox of CSS-Palm (http://bioinformatics.lcd-ustc.org/css_palm/prediction.php) and pressed the ‘Submit’ button to get the predicted palmitoylation sites by CSS-Palm (Fig. 1a). As in Figure 1b, the prediction result suggests that both of the CENP-E proteins may be palmitoylated at a conserved pair of peptides in human (1477-EELKVAHCCNLKEQET-1492) and mouse (1378-EELNLARCCLEQENK-1393), which needs further experimental verification.

Fig. 1. (a) Prediction page of CSS-Palm web server. (b) Prediction results of human (Q02224) and mouse CENP-E (Q6RT24). The default cut-off score is 2.6 (Sn 82.16% and Sp 83.17%).
By pressing on ‘here’ in the button ‘Download the TAB-delimited data file from here’ (as shown in Fig.1b), a user can get the TAB-delimited result file that is convenient for further automatic processing.

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