Database and Prediction of Sequence Motifs on Protein Molecular Interactions

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Abstract
We are developing a new prediction tool for protein-protein interactions based on sequence motifs. For this attempt, we have collected the information of the interaction data from existing databases and literatures, and arranged it into a new interaction sequence motif database. This paper describes the database and the prediction system for protein-protein interaction motifs.

1 Introduction
As technologies in the area of genome research advances, huge amount of genome and proteome data are accumulating. Utilizing the huge data, genome scientists are exploring higher biological systems based on molecular interactions that take parts in metabolic pathway and signaling system. Here, we focus on signalling networks, and describe our attempt to predict protein interactions based on sequence motifs. As for proteins in signaling networks, many signalling domains have been reported [1]. These domains include those for protein-protein interactions, e.g. SH2 and SH3 domains. Motifs recognized by the domains could be different in different groups, even if the recognizing domains belong to the same group. Thus, the precise knowledge of interaction pairs is necessary for developing better prediction methods. And, if we could predict these interaction specificity, the predicted results would be helpful to further analyzing of signal networks.

2 Method

2.1 Database
The protein-protein interaction data is composed of two parts. One is general information of interaction domains listed in literature [1] and the other is cross references to individual sequence entries.

2.2 Classification of specificity
Among proteins in signaling networks, SH2 domain and kinase motif have been well investigated and characterized by experimental work [2, 3]. According to Songyang [3], SH2 domains are classified into four groups. In the recognition of phosphotyrosine, each group of SH2 domain has different specificity to different sequence patterns around the phosphotyrosine. Our classification of SH2 domains are based on Songyang [2], and summarized as follows:

1. SH2 domain was extracted from SWISSPROT.
2. The sequence of SH2 domains from SWISSPROT were aligned against the SH2 profile data in PROSITE.
3. The residues contributing to recognition specificity were detected and SH2 domains were classified into four groups.
3 Results and Discussion

Table 1 shows an example of phosphotyrosine - SH2 domain protein pair data. Table 2 shows the classified SH2 domains. Motifs around phosphotyrosine in Table 2 is described in Songyang [2].

Table 1: Database example

<table>
<thead>
<tr>
<th>phosphotyrosine name</th>
<th>motif</th>
<th>SH2 domain name</th>
<th>domain region</th>
</tr>
</thead>
<tbody>
<tr>
<td>Shc</td>
<td>YVNV</td>
<td>Grb2</td>
<td>60-152</td>
</tr>
<tr>
<td>EGF receptor</td>
<td>YSSD</td>
<td>Grb2</td>
<td>60-152</td>
</tr>
<tr>
<td>PDGF receptor</td>
<td>YVPM</td>
<td>Nck</td>
<td>282-376</td>
</tr>
</tbody>
</table>

These motifs are so short (about 4 residues), that a search a sequence database by the motifs would produce many false positives. For this reason, phosphotyrosine motifs must be more closely examined and refined.

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References