PhosphoNetworks: a database for human phosphorylation networks

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ABSTRACT

Summary: Phosphorylation plays an important role in cellular signal transduction. Current phosphorylation-related databases often focus on the phosphorylation sites, which are mainly determined by mass spectrometry. Here, we present PhosphoNetworks, a phosphorylation database built on a high-resolution map of phosphorylation networks. This high-resolution map of phosphorylation networks provides not only the kinase–substrate relationships (KSRs), but also the specific phosphorylation sites on which the kinases act on the substrates. The database contains the most comprehensive dataset for KSRs, including the relationships from a recent high-throughput project for identification of KSRs using protein microarrays, as well as known KSRs curated from the literature. In addition, the database also includes several analytical tools for dissecting phosphorylation networks. PhosphoNetworks is expected to play a prominent role in proteomics and phosphorylation-related disease research.

Availability and implementation: http://www.phosphonetworks.org

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1 INTRODUCTION

The reversible phosphorylation of proteins regulates many aspects of cellular physiology. Abnormal phosphorylation is known to be both a cause and a consequence of many diseases, such as cancer, diabetes, heat attack, hypertension and rheumatoid arthritis (Cohen, 2001). The systematic study of phosphorylation can greatly benefit disease-related biomedical research. In a recent project, we developed a combined protein microarray and bioinformatics approach to identify kinase–substrate relationships (KSRs) in a high-throughout manner (Hu et al., 2013; Newman et al., 2013). Based on the results of these studies, we experimentally identified substrates for 289 unique kinases, resulting in 3656 high-quality KSRs. The study generated more human KSRs than all previous studies combined. To facilitate the use of this information, we created the ‘Database for a High-Resolution Map of Human Phosphorylation Networks (PhosphoNetworks)’, an integrated information system for the storage, retrieval, visualization and analysis of human phosphorylation data. To enhance the discovery of phosphorylation relationships between kinases and their downstream substrates, we also connected the kinases with the specific phosphorylation site on substrate protein sequences, which will help users design mutagenesis experiments of phosphorylation sites to block the phosphorylation event. Therefore, the PhosphoNetworks database provides not only a powerful information resource but also an integrated analysis platform.

2 DATABASE CONTENT

PhosphoNetworks currently covers KSRs at different levels. First, it contains 24046 in vitro KSRs (rawKSRs) identified by protein microarray. Such relationships reflect biochemical reactions between kinases and their substrates. On the next level, the database includes 3656 refined, high-confidence, physiologically relevant KSRs (refKSRs), which are filtered by a series of criteria to enrich for KSRs that are likely to occur in cells. The quality of the KSRs was extensively validated by independent transfected cell experiments (Newman et al., 2013). Finally, 744 literature-curated KSRs were integrated with the 3656 refKSRs to generate a combined data set (comKSR). Besides the KSRs, the database also includes 300 predicted consensus phosphorylation motifs for 284 human kinases (~55% of human kinome). For dual-specificity kinases, which phosphorylate both serine/threonine and tyrosine residues, two types of motifs are predicted. The quality of these predicted motifs was supported by the comparisons with motifs from other sources, such as positional scanning peptide libraries or other computational methods (Hutti et al., 2004; Mok et al., 2010; Newman et al., 2013).

To create a high-resolution map of KSRs, we further integrated the 300 consensus phosphorylation motifs and phosphorylation sites determined by mass spectrometry (MS/MS) on substrates of each kinase. Using a computational approach, we connected 230 kinases with 2591 phosphorylation sites on 652 substrate proteins.

A summary of PhosphoNetworks content is shown in Table 1, and all data are freely available for all academic users from our Web site (http://phosphonetworks.org/download.html).

3 DATABASE USAGE AND ACCESS

PhosphoNetworks is composed of four major functional modules, namely ProteinSearch, Site Search, PathSearch and NetworkSearch.
NetworkSearch allows the user to perform network analysis by querying a set of proteins. Given a group of protein names as input, NetworkSearch will find all direct neighbors of these proteins and show the network composed by them in a scalable vector graphics (SVG) figure. The template of NetworkSearch is based on our previous MoReNet database (Hu et al., 2010). It allows users to drag and move nodes in the network to get a better topology. The users are also allowed to download gene and gene pairs in the network in plain text format.

### 4 DISCUSSIONS

PhosphoNetworks database integrates two kinds of high-throughput phosphorylation data, protein microarray-verified KSRs and MS-verified phosphorylation sites. Relative to other human phosphorylation-related databases, such as Phospho.ELM (http://phospho.elm.eu.org/) (Dinkel et al., 2011), PhosphoNetworks has three significant merits. First, it is a high-resolution phosphorylation network database, which connects kinases not only to their downstream substrates but also to specific phosphorylation sites on the substrates. Second, it covers a far greater number of novel identified KSRs (24 046 raw KSRs and 3656 refined KSRs) and novel predicted phosphorylation motifs (300 motifs, over double that of the previous knowledgebase). Finally, it provides some unique tools to make it easier for the user to explore and analyze the data. For example, the PathSearch function allows the user to find the missing link in known incomplete pathways, and the SiteSearch function allows the user to predict possible upstream kinases of their interested proteins or peptides. We expect this database will be valuable for proteomics and phosphorylation-related disease research.

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### REFERENCES


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**Table 1. Summary table of data in PhosphoNetworks**

<table>
<thead>
<tr>
<th>Type</th>
<th>Statistics</th>
</tr>
</thead>
<tbody>
<tr>
<td>rawKSR</td>
<td>24046 (K = 289, S = 1067)</td>
</tr>
<tr>
<td>refKSR</td>
<td>3656 (K = 255, S = 742)</td>
</tr>
<tr>
<td>comKSR</td>
<td>4375 (K = 255, S = 1139)</td>
</tr>
<tr>
<td>Motif</td>
<td>300 (K = 284)</td>
</tr>
<tr>
<td>MS PhosSite</td>
<td>70422 (S_p = 48704, T_p = 15373, Y_p = 6375)</td>
</tr>
<tr>
<td>K-S-PhosSite</td>
<td>4417 (K = 230, S = 652, PhosSite = 2591)</td>
</tr>
</tbody>
</table>

*Note:* K, Kinase; S, Substrate; S_p, phosphorylated serine; T_p, phosphorylated threonine; Y_p, phosphorylated tyrosine.