qPhos: a database of protein phosphorylation dynamics in humans

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ABSTRACT

Temporal and spatial protein phosphorylation dynamically orchestrates a broad spectrum of biological processes and plays various physiological and pathological roles in diseases and cancers. Recent advancements in high-throughput proteomics techniques greatly promoted the profiling and quantification of phosphoproteome. However, although several comprehensive databases have reserved the phosphorylated proteins and sites, a resource for phosphorylation quantification still remains to be constructed. In this study, we developed the qPhos (http://qphos.cancerbio.info) database to integrate and host the data on phosphorylation dynamics. A total of 3,537,533 quantification events for 199,071 non-redundant phosphorylation sites on 18,402 proteins under 484 conditions were collected through exhaustive curation of published literature. The experimental details, including sample materials, conditions and methods, were recorded. Various annotations, such as protein sequence and structure properties, potential upstream kinases and their inhibitors, were systematically integrated and carefully organized to present details about the quantified phosphorylation sites. Various browse and search functions were implemented for the user-defined filtering of samples, conditions and proteins. Furthermore, the qKinAct service was developed to dissect the kinase activity profile from user-submitted quantitative phosphoproteome data through annotating the kinase activity-related phosphorylation sites. Taken together, the qPhos database provides a comprehensive resource for protein phosphorylation dynamics to facilitate related investigations.

INTRODUCTION

As one of the most important post-translational modifications (PTMs), protein phosphorylation is involved in almost all biological processes and plays physiological and pathological roles in diseases and cancers (1–3). Phosphorylation is reversibly catalysed by kinases and phosphatases, which dynamically control the phosphorylation dynamics based on a temporal and spatial context (4,5). In 1992, Edmond H. Fischer and Edwin G. Krebs shared the Nobel Prize in Physiology or Medicine for their discovery of reversible protein phosphorylation as a biological regulatory mechanism (6), while the prize in 2001 was awarded to Leland H. Hartwell, Tim Hunt and Paul M. Nurse for identifying key regulators, including cyclin-dependent kinases (CDKs) and cyclins, which accurately orchestrate the cell cycle through phosphorylation (7). In recent decades, many studies have dissected the molecular mechanisms and biological functions of phosphorylation dynamics, and kinases and phosphatases are popular research areas for the development of target therapies (5,8).

Recently, the advancement of high-throughput proteomics techniques greatly promoted the profiling and quantification of the phosphoproteome in various cells and tissues under different conditions (9,10). For example, based on phosphoproteome quantification, Wojcechowski et al. dissected the molecular mechanism of cellular reprogramming during HIV-1 entry (11), and van den Biggelaar et al. unveiled the dynamic phosphorylation of thrombin signalling in human primary endothelial cells (12). Since dynamic phosphorylation events could provide helpful clues for kinase-mediated signalling, Ho et al. identified the activation of PI3K/AKT/mTORC1 signalling for cell survival by the ELABELA peptide in human embryonic stem cells during heart development (13), Bai et al. profiled the driver tyrosine kinases in sarcoma (14) and Casado et al. inferred the aberrant kinase activation in leukaemia cells through the quantitative phosphoproteome-based compu-
tional analyses (15). Thus, the quantitative phosphoproteomics data could provide great help in understanding phosphorylation-controlled biological processes.

Previously, several pioneering studies have contributed to constructing resources to host phosphorylation-related data. The currently available databases contain many human protein phosphorylation data. UniProt (16) is the most important infrastructure for protein annotations, which contain massive experimentally identified phosphorylation sites and annotations in the Human Protein Reference Database (HPRD) (17) also contain protein phosphorylation information. Databases including dbPTM (18), PhosphoSitePlus (19), SysPTM (20), PHOSIDA (21), dbPAF (22), Phospho.ELM (23) and PhosphoPep (24) curate and host experimentally identified phosphorylation sites from the published literature, and iPTMnet (25) contains information on the phosphorylation regulatory network and conservation. Furthermore, the PhosSNP (27) and ActiveDriverDB (28) databases analyse the genetic variations that influence phosphorylation, and the PTMcode (26) database provides the functional associations of phosphorylation sites. With the continuous improvement of high-throughput phosphoproteome techniques and the rapid increase of phosphoproteome datasets, ProteomeScout (29) and ProteomicsDB (30) were developed to store proteome and PTM proteome datasets and provide online analysis tools. However, although the ProteomeScout database contains protein quantification information, the quantification of phosphorylation events is still missing. Taken together, although databases for various aspects of protein phosphorylation are available, the database for the quantification of phosphorylation is still absent. Since the quantification/stoichiometry/dynamics of phosphorylation are critical for the molecular mechanisms under different temporal and spatial contexts, a comprehensive resource for protein phosphorylation quantification could facilitate the reuse of published quantitative phosphoproteome datasets and provide great help for phosphorylation-related studies.

In this study, we developed the qPhos database to host the quantitative phosphoproteome data generated in Homo sapiens. In total, 3 537 533 quantification events for 199 071 non-redundant phosphorylation sites on 18 402 proteins under 484 conditions were collected from 190 published studies. The primary references for the data were provided to ensure their quality and repeatability, and various annotations were integrated into the database. The sequence and structure preferences of the quantified phosphorylation sites were presented in Figure 2A–D. It seemed that the phosphorylation sites were generally equally distributed among the protein sequences but slightly enriched in C-terminal areas (E-ratio = 1.06, P-value < 10\(^{-17}\)) (Figure 2A). Through computational annotation of secondary structures, surface accessibility and disorder region by NetSurfP (37) and IUPred (38), it was observed that the phosphorylation sites were enriched in the coil region (E-ratio = 1.21, P-value < 10\(^{-18}\)) (Figure 2B), exposed region (E-ratio = 1.21, P-value < 10\(^{-18}\)) (Figure 2C) and disordered region (E-ratio = 2.09, P-value < 10\(^{-18}\)) (Figure 2D). As shown in Figure 2E, the distribution of phosphorylated serine, threonine and tyrosine residues was consistent with previous studies (19,22). Furthermore, the potential upstream kinases were annotated for these phosphorylation sites by various resources. The experimentally identified kinase-site regulatory relations were retrieved from previous studies, including Phospho.ELM (23), PhosphoSitePlus (19) and MusiteDeep (33). Additionally, the potential site-specific kinase-substrate relationships were predicted by sequence-based

**CONSTRUCTION AND CONTENT**

To establish the resource for protein phosphorylation dynamics, the published quantitative phosphoproteomics datasets were collected from the literature and annotations from various resource were integrated (Figure 1). Keywords including ‘phosphoproteomic’, ‘phosphoproteomics’ and ‘phosphoproteome’ were used to search PubMed to retrieve the phosphoproteome-related literature. Based on the criteria of high-throughput mass spectrometry-based site-level protein phosphorylation quantification, the published datasets of quantitative phosphoproteome datasets in *H. sapiens* were manually curated. Besides the quantified phosphorylation sites, details about the quantification including the experimental condition, phosphopeptide enrichment method, mass spectrometry and raw peptide were collected. All quantified phosphopeptides and modified residues were mapped to the *H. sapiens* reference proteome sequences downloaded from UniProt database (Release 2018.04) (Figure 1) (16). Due to the differences among the miscellaneous reference proteome datasets used in these literature, about 4.31% of raw phosphopeptides could not be mapped to the UniProt reference human proteome, and these limited unmapped data was dropped. In addition, the annotations from databases such as UniProt (16), ExpASy (31) and DrugBank (Release 2018.07) (32) were integrated into the qPhos database to provide comprehensive information for phosphorylation events (Figure 1). The experimental verified kinase–substrate relationships were integrated from Phospho.ELM (23), PhosphoSitePlus (19) and MusiteDeep (33), and potential relationship were predicted by sequence-based predictor GPS (34) and network-based predictor iGPS (35) with a high threshold (Figure 1). Furthermore, the activity-related phosphorylation sites in kinases were also curated from the literature. The human kinases and their names were retrieved from the EKPD database (36), and coupled with keywords including ‘phosphorylation’ and ‘activity or activate or activation’ to search PubMed for experimental evidence of activity-related phosphorylation (Figure 1).

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predictor GPS (34) and network-based predictor iGPS (35) with a high threshold. The annotations of kinases were summarized at the family level in Figure 2F.

**USAGE**

The qPhos database was developed for scientists to quickly access the quantitative phosphoproteome data in a user-friendly manner. For convenient usage, qPhos provided browse and search functions to query the database. Three browse options, including condition, sample and gene, were provided to browse the database by selecting the item from the list (Figure 3A). The experimental conditions, samples including cell lines and tissues, and gene symbols were sorted and organized in alphabetical order, which enabled the users to quickly choose the interesting data (Figure 3A). Simple and complex search functions were implemented at the home (Figure 3B) and search (Figure 3C) pages, respectively, which provided keyword-based queries in protein and gene names, protein functions and descriptions of conditions and samples. Furthermore, the retrieved phosphorylation sites from the browse or search functions could be further filtered by conditions, samples and methods in the results page (Figure 3D).

In the results page, the information was organized by the quantification events in a tabular format with UniProt accession, gene name, position, sequence window, sample information, sample type, experimental condition, quantification method, log2-transformed ratio and P-value (Figure 3D). Users could click the plus button to view the detailed information about the protein and phosphorylation sites. The detailed information had four sections, including ‘About experiment’, ‘Potential kinases and their inhibitors’, ‘Sequence and structure’ and ‘About protein’ (Figure 3E–I). The detailed description of the condition, raw quantifications, source reference, experimental method and instrument and raw peptide were shown in the ‘About Experiment’ section (Figure 3E). ‘Potential kinases and their inhibitors’ showed the experimental identified and predicted upstream kinases for the phosphorylation site (Figure 3F). Furthermore, the inhibitors annotated by DrugBank for the kinases were shown (Figure 3F). The sequence and structure properties of the protein were visualized in the ‘Sequence and Structure’ section, which included the quantified phosphorylation sites, activity-related phosphorylation sites if the protein was a kinase, disorder region, secondary structure and surface accessibility (Figure 3G). A magnifier was implemented to show the details by enlarging the selected region (Figure 3G). Furthermore, users could access descriptions about visualization by hovering over the content (Figure 3I). The ‘About Protein’ section...
Figure 2 Summary of the sequence and structure preferences and kinase families of the quantified phosphorylation sites, including the summary of the position along the protein sequence (A), secondary structure (B), surface accessibility (C), disorder region (D), serine/threonine/tyrosine (E) and regulator kinase family (F) for the quantified phosphorylation sites.

presented the protein information including database accessions, protein/gene name/alias, functional descriptions, PTMs and sequences adopted from the UniProt database (Figure 3H).

Since autophosphorylation or phosphorylation of the specified segment could activate or inactivate the kinases (39), the identification and quantification of these activity-related phosphorylation sites could indicate the activation status of these kinases. Through exhaustive curation of literature, we collected the experimentally identified activity-related phosphorylation sites for the human kinome. In total, 829 activity-related phosphorylation sites were curated in 272 kinases (Figure 4A), which covered over half of the kinome. Among the activity-related phosphorylation sites, nearly half were autophosphorylation sites. Most autophosphorylationsites could activate the kinases, while seven autophosphorylationsites could inactivate the kinases. Furthermore, most activity-related phosphorylation sites were positively related to the activities of the kinase, while only a small fraction were negatively related (Figure 4A). The distributions of activity-related phosphorylation sites among kinases and kinase families were summarized in Figure 4B and C, respectively. The qKinAct service was developed to query the quantification events of activity-related phosphorylationsites to annotate the kinase activity profile from quantitative phosphoproteome data (Figure 4D). The activity-related phosphorylation sites could be annotated through straightforward submission of the identified phosphorylated peptide and their quantifications (Figure 4E).

Here, we provide two examples to show the applicability of the qKinAct service for kinase activity analysis. Paulo et al. globally analysed the phosphorylation dynam-
Figure 3. The detailed information in qPhos. (A) Browse function. (B) Simple search function. (C) Advanced search function. (D) The returned search results. (E) The information about the protein. (F) The information about the quantification of the phosphorylation site. (G) The information on potential kinases and their inhibitors for the quantified phosphorylation site. (H) The sequence and structure properties of the phosphorylation site. (I) The enlarged view of the sequence and structure properties.
The qKinAct service for the analysis of kinase activities. (A) The distribution of different types of activity-related phosphorylation sites in kinases. ‘+’, ‘−’ and ‘auto’ represent positively related, negatively related and autophosphorylation sites. (B) The distribution of kinases with activity-related phosphorylation sites in kinases. (C) The distribution of kinase families with activity-related phosphorylation sites in kinases. (D) The example for submission of quantitative phosphoproteome data. (E) The returned results for the query of kinase activity-related phosphorylation sites. The kinase activity profile for phosphorylation dynamics in nicotine-treated pancreatic stellate cells (F) and TNFα-stimulated phosphorylation dynamics (G).
inAct service was developed to dissect the kinase activity profile from user-submitted quantitative phosphoproteome data through direct annotations of activity-related phosphorylation sites for kinases. Meanwhile, there were also various limitations in the database. The database focused on human health-related phosphorylation dynamics, and currently only data from human tissues and cell lines were collected. However, several quantitative phosphoproteomics studies were based on a mouse model or other model organisms (45), which also provided helpful clues for medical investigations. Furthermore, previous studies showed that single nucleotide polymorphisms (SNPs) and somatic mutations at or around the phosphorylation sites could affect the phosphorylation dynamics (27), and currently qPhos has not linked the phosphorylation sites to the large quantity of SNP and mutation data. Taken together, although improvements remain to be achieved, qPhos could serve as a comprehensive resource to enable researchers to systematically and conveniently access the phosphorylation dynamics data under different experimental conditions, and its qKinAct service could help users to easily analyse the kinase activity profile from their own quantitative phosphoproteome data. The qPhos database will be regularly updated to keep pace with the progress of the quantitative dynamics of phosphorylation.

**FUNDING**

National Natural Science Foundation of China [31501069, 81802438]; National Key R&D Program of China [2018YFC1313300]; Program for Guangdong Introducing Innovative and Entrepreneurial Teams [2017ZT07S096]; Natural Science Foundation of Guangdong Province [2014A030312015]; Science and Technology Program of Guangdong [2015B020230088]; Science and Technology Program of Guangzhou [15570006, 201508020250, 201604020003]; Fundamental Research Funds for the Central Universities [SYSU: 16ykzd06]; Pearl River S&T Nova Program of Guangzhou [Ze-Xian Liu]; Special Program for Applied Research on Super Computation of the NSFC-Guangdong Joint Fund (the second phase). Funding for open access charge: Program for Guangdong Introducing Innovative and Entrepreneurial Teams [2017ZT07S096].

**Conflict of interest statement.** None declared.

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