RESEARCH ARTICLE

The Roles of Post-translational Modifications in the Context of Protein Interaction Networks

Guangyou Duan*, Dirk Walther*
Max Planck Institute for Molecular Plant Physiology, Potsdam-Golm, Germany

* walther@mpimp-golm.mpg.de

Abstract

Among other effects, post-translational modifications (PTMs) have been shown to exert their function via the modulation of protein-protein interactions. For twelve different main PTM-types and associated subtypes and across 9 diverse species, we investigated whether particular PTM-types are associated with proteins with specific and possibly “strategic” placements in the network of all protein interactions by determining informative network-theoretic properties. Proteins undergoing a PTM were observed to engage in more interactions and positioned in more central locations than non-PTM proteins. Among the twelve considered PTM-types, phosphorylated proteins were identified most consistently as being situated in central network locations and with the broadest interaction spectrum to proteins carrying other PTM-types, while glycosylated proteins are preferentially located at the network periphery. For the human interactome, proteins undergoing sumoylation or proteolytic cleavage were found with the most characteristic network properties. PTM-type-specific protein interaction network (PIN) properties can be rationalized with regard to the function of the respective PTM-carrying proteins. For example, glycosylation sites were found enriched in proteins with plasma membrane localizations and transporter or receptor activity, which generally have fewer interacting partners. The involvement in disease processes of human proteins undergoing PTMs was also found associated with characteristic PIN properties. By integrating global protein interaction networks and specific PTMs, our study offers a novel approach to unraveling the role of PTMs in cellular processes.

Author Summary

The function of proteins is frequently modulated by chemical modifications introduced after translation from RNA. These post-translational modifications (PTMs) have been shown to also influence the interaction between proteins carrying them. We tested whether specific PTM-types characterized by attaching different chemical groups are associated with proteins with characteristic and possibly strategic positions within the network of all protein interactions in cellular systems. Based on network-theoretic analyses of PTMs in
the context of protein interaction networks of nine selected species, we indeed observed distinctive properties of twelve PTM-types tested. Phosphorylation was found associated with proteins in central locations with the broadest interaction scope, while glycosylation was more prominent in proteins at the periphery of the web of all protein interactions. The involvement in disease processes of human proteins undergoing PTMs was also found associated with characteristic protein interaction network properties. Our study highlights common and specific roles of the various PTM types in the orchestration of molecular interactions in cells.

Introduction

As chief actors within living cells, proteins serve diverse functions such as catalysis, transport, structural building material and many others [1]. While the human gene set was estimated at about 25,000 genes [2], the human proteome size is expected to be far larger and estimated at over 1 million proteins [3]. Beyond alternative splicing of mRNA as a source of protein diversity, post-translational modifications (PTMs) of proteins further modulate and extend the range of possible protein functions by covalently attaching small chemical moieties to selected amino acid residues. More than 200 different types of PTMs have been identified that affect many aspects of cellular functionalities, such as metabolism, signal transduction, and protein stability [4, 5]. These modifications include phosphorylation, glycosylation, methylation, acetylation, amidation and many other types, see http://www.uniprot.org/docs/ptmlist for a more detailed controlled vocabulary of PTMs curated by UniProt [6]. With technological advances, PTMs can be detected at an ever increasing breadth, precision, and quantity e.g. by using mass spectrometry (MS) based methods [7]. Several databases have been established to store the obtained information, such as UniProt [8], dbPTM [9], PTMCuration [10], PTMcode [11] and many others. Among them, a number species-specific databases have been developed [12–14] offering the opportunity to investigate PTMs in an evolutionary context as well.

Many studies on PTMs have focused on specific types and their relevance for protein function with phosphorylation representing the most actively researched PTM-type [15–19]. More recently, the interplay between different PTM-types has moved into the focus of attention [20–23]. For example, evidence of an interdependence of phosphorylation and acetylation was reported for a genome-reduced bacterium Mycoplasma pneumoniae [24]. Furthermore, so-called integrative PTM spots (PTMi) have been identified as site in proteins at which different PTMs operated in a combinatorial manner to modulate protein function [25]. A more global view of the interplay between PTM-types was presented in a study on the co-evolution between 13 frequent PTM types in 8 eukaryotic species [26]. Carboxylation was identified as evolutionarily most conserved, whereas phosphorylation was found among those PTM-types playing a central role in the modulation of the dynamics of protein function. For a recent review on the evolution and functional cross-talk between PTMs, see [27].

In addition to PTMs, protein function is also regulated and mediated by non-covalent protein-protein interactions [28–33]. As many PTMs modulate the binding affinities between proteins by changing the electrostatic or structural properties of the involved interaction sites [29], PTMs and protein-protein interactions are frequently functionally connected. Based on data from the dbPTM database of protein post-translational modifications [9], more than 60% of PTM sites are related to those protein functional domains that were shown to preferentially engage in direct protein-protein interactions suggesting a central regulatory role of PTMs in the modulation of protein interactions, and thus, function.
Therefore, it appears plausible that proteins carrying a particular PTM-type may possess specific interaction characteristics. Indeed, for the important and intensively investigated PTM-type phosphorylation, it was found that in yeast, phospho-proteins engage in many more protein-protein interaction than proteins without phosphorylation sites [34], found similarly in Arabidopsis thaliana [35]. Thus, phosphorylation of a single protein potentially leads to a modulation of many different interactions, and thus, molecular processes, simultaneously.

As the different PTMs modify proteins in specific ways, it appears furthermore likely that their consequences on protein-protein interactions may be different as well. This hypothesis formed the starting point for the present study. Specifically, we asked whether different PTM-types are associated with characteristic protein-protein network properties, such as interaction degree, clustering coefficient, and closeness centrality, for those proteins carrying them. We selected these three properties as they each reflect on potential functional role such as scope of impact (degree), diversity of responses (clustering coefficient), and placement within a possible signaling cascade (closeness centrality). Furthermore, we investigated whether those characteristics are conserved across different species, and if the particular functions the different PTMs fulfil may become apparent when inspected from the viewpoint of protein-protein interactions.

Results

To base our analyses on high-confidence PTM-instances, we restricted our analyses to PTM-sites that have been identified experimentally leaving out all annotated PTMs that are based on computational predictions alone. PTM-site information was collected from 11 different database resources and consolidated into a single set via sequence position information. Currently available datasets proved sufficiently large to conduct statistical analyses on the role of PTMs in the context of protein interaction networks for the following twelve PTM-types: acetylation, amidation, carboxylation, disulfide bond, glycosylation, hydroxylation, methylation, nitrosylation, phosphorylation, proteolytic cleavage, sumoylation, and ubiquitination associated with a set of nine diverse eukaryotic species covering several lineages and kingdoms: mammals (Homo sapiens, Mus musculus, Rattus norvegicus, Bos taurus), an invertebrate (Caenorhabditis elegans), the insect Drosophila melanogaster, fungi (Saccharomyces cerevisiae, Schizosaccharomyces pombe), and the plant Arabidopsis thaliana (Table 1). Acetylation is characterized by the attachment of an acetyl group either to the N-terminus of protein or to lysine residues. Amidation leads to the addition of amide groups to the C-terminus of proteins. During carboxylation, a carboxylic group is added to glutamate residues. Glycosylation includes all O-linked (serine, threonine, tyrosine residues) or N-linked (arginine and asparagine residues) attachments of simple or complex carbohydrates (e.g. monosaccharides, branched polysaccharides) to proteins. Upon Hydroxylation, hydroxyl groups are attached to proline residues. Methylation refers to transfer of methyl groups to arginine or lysine residues. Nitrosylation leads to the incorporation of nitric oxide into the thiol group of cysteine residues. Protein phosphorylation is associated with attaching phosphate groups to serine, threonine, or tyrosine protein residues. Both sumoylation and ubiquitination are characterized by the attachment of small proteins to target proteins modifying their function or stability or, in the case of ubiquitination, tagging them for degradation. While the previous 10 PTM-types are characterized by the attachment of chemical moieties to proteins, disulfide bond and proteolytic cleavage lead to posttranslational modifications via changing the chemical bond structure within a given protein and in the process removing atoms from it, either by forming a covalent bond (disulfide bond between two cysteine residues accompanied by the removal of two hydrogen atoms) or by breaking peptide bonds (proteolytic cleavage removing HOH). As the latter two met our established
Table 1. Frequency table of proteins associated with the selected PTM-types and species.

| NCBI taxonomy ID | Species Name       | acetylation | amidation | carboxylation | disulfide bond | glycosylation | hydroxylation | methylation | nitrosylation | phosphorylation | proteolytic cleavage | sumoylation | ubiquitination |
|------------------|--------------------|-------------|-----------|---------------|----------------|---------------|---------------|-------------|---------------|-----------------|---------------------|--------------|---------------|
| 10090            | Mus musculus       | 1688        | 7         | 11            | 0               | 1808          | 5             | 879         | 996           | 9770             | 0                   | 67           | 5616          |
| 10116            | Rattus norvegicus  | 303         | 25        | 14            | 0               | 164           | 8             | 198         | 85            | 3496             | 0                   | 14           | 523           |
| 9913             | Bos taurus         | 103         | 19        | 34            | 0               | 108           | 11            | 20          | 3             | 175              | 0                   | 3            | 6             |
| 9606             | Homo sapiens       | 4858        | 38        | 88            | 190             | 1823          | 33            | 1118        | 368           | 14425            | 388                 | 327          | 6440          |
| 7227             | Drosophila melanogaster | 10    | 13        | 3             | 0               | 83            | 0             | 7           | 0             | 2415             | 0                   | 0            | 2             |
| 6239             | Caenorhabditis elegans | 16    | 13        | 1             | 0               | 214           | 0             | 2           | 0             | 2138             | 0                   | 0            | 0             |
| 3702             | Arabidopsis thaliana  | 96         | 1         | 6             | 0               | 71            | 6             | 23          | 41            | 9198             | 0                   | 5            | 51            |
| 4932             | Saccharomyces cerevisiae | 494  | 1         | 0             | 0               | 55            | 0             | 27          | 1             | 3195             | 0                   | 17           | 125           |
| 4896             | Schizosaccharomyces pombe  | 2         | 0         | 0             | 0               | 11            | 0             | 8           | 0             | 1072             | 0                   | 0            | 6             |

Listed are the numbers of proteins with the corresponding PTM-type in the respective species.

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count-criteria and do indeed modify the protein, we retained them for the initial analyses. However, as data were available for human only, and to confine the analyses to moiety-addition-type PTMs, both PTM-types were left out in the subsequent analyses of network properties. For several PTM-types, several subtypes exist, e.g. Ser/Thr/Tyr phosphorylation. Generally, we considered PTM-types based on the added moiety, but also repeated selected analyzes with a further subsetting of the datasets based on the receiving group on the protein.

The collected sets of proteins associated with the twelve different posttranslational modifications (PTMs) across the nine selected species were first investigated for PTM-type specific biological process involvement based on GO-term enrichment statistics as well as probed for significant co-occurrence patterns of different PTM-types on the same protein. Subsequently, we investigated whether protein sets associated with particular types of PTMs exhibit characteristic protein interaction network (PIN) properties. For the latter, we computed three basic and commonly used network properties, the degree, the clustering coefficient, and the closeness centrality, associated with proteins belonging to different PTM-specific protein sets when mapped onto the species-specific PIN. Finally, we tested whether pairs of interacting proteins exhibit preferences with regard to PTM-types they carry.

PTM-specific biological process and location enrichment analysis

Frequently, proteins are modified not only by a single PTM event, but by several and of different PTM-types. Thus, if we wish to understand the role of individual PTM-types in the context of protein-protein interactions, we first need to understand their co-occurrence on the same protein as well as their functional profile as it seems plausible that PTM-types associated with similar functional involvement will also exhibit similar characteristics with regard to their protein interactions.

The functional significance of specific PTM-types and the respective proteins carrying them has been amply investigated [26, 32, 36, 37]. To provide a comparative overview of the selected PTM-types studied here, we integrated all species-specific gene ontology (GO) annotations into a merged set and determined preferred biological process involvements, functional roles, and subcellular locations based on GO-enrichment statistics computed for this artificial "super species". Consistently across all GO-term domains (process, function, and location), the 12 PTM-types are grouped into two major groups with sumoylation, nitrosylation, methylation, acetylation, phosphorylation, ubiquitination in one group (group-I), and disulfide bond, carboxylation, hydroxylation, proteolytic cleavage, glycosylation and amidation in the other (group-II) (Fig. 1, S1 Fig.). While group-I PTM-types were found preferentially in proteins located in the cytosol and nucleus and involved in regulatory processes (most noteworthy, transcriptional regulation), group-II PTM-types appear associated with membrane-, subcellular compartment localizations (carboxylation), extracellular locations and secretory processes. In line with several reported observations on their concerted action [38], phosphorylation and ubiquitination were found with similar GO process and location profiles. Acetylation appears to be involved in similar processes as well. Indeed, the combined action of these three PTM-types has been described in selected cases as, for example, for the protein p53 [20]. Furthermore, glycosylation and proteolytic cleavage exhibit similar GO-term characteristics, which may reflect the involvement of and even interplay between both PTMs in the modification of secreted and/or membrane-embedded proteins [39]. Thus, different PTM-types have similar functional involvement and location profiles suggesting that their characteristic protein-interaction network properties may also be similar, which appears implied in particular based on common localizations influencing the scope of potential protein interaction partners.
The co-occurrence patterns of different PTM-types on the same protein are critical confounding factors for the analysis of the individual PTM-types in the context of protein interactions. Evidently, frequent PTM-types will have a greater chance of co-occurring with other PTM-types on the same protein. Indeed, as judged by the Jaccard-distance, protein sets in human associated with phosphorylation, acetylation, and ubiquitination—the three PTM-types with the most observed instances (Table 1)—exhibit large overlaps (Fig. 2). However, as we tested for deviations from the expected chance overlap as well, this co-occurrence on the same protein also seems significant. In general, the overlap amongst all PTM-types studied here is extensive. The reduced overlap for amidation, disulfide bond, hydroxylation carboxylation, and methylation with other PTM-types appears largely caused by their low frequency. Similarly, when expanding the overlap analysis to all species considered here, a large overlap between phosphorylation and acetylation is evident (Fig. 3). Interestingly, when viewed across several species, methylation emerges as a PTM-type with significant co-occurrence with both acetylation and phosphorylation possibly reflecting their joint association with histones [40]. Furthermore, glycosylation and phosphorylation appear to frequently co-exist on the same protein.

In conclusion, the overlap of different PTM-types on the same protein is extensive and greater than expected by chance. Even though suggested by the separate clustering of PTM-types based on their functional and location annotations (Fig. 1), with regard to their co-occurrence pattern, no equivalent segregation is apparent. Therefore, all PTM-types will—when analyzed jointly—likely exhibit similar protein-interaction characteristics. While primarily reporting results on the global protein sets (including overlaps), we also performed analyses on
the one-PTM-type-only protein sets. Evidently, one cannot be certain that those unique sets are truly unique in reality as not all PTM-types and their instances have been identified yet. Furthermore, rendering the data set PTM-type specific, i.e. reducing the protein sets to sets conforming to one PTM-type only must inevitably lead to a massive reduction of statistical power.

PTM-type specific protein interaction network properties

We inspected the protein sets associated with specific PTM-types in the context of known protein-protein interactions. We mapped all proteins with annotated PTMs onto the respective protein interaction networks (PINs) of the nine selected species (Table 2). By computing three network properties, the degree, the clustering coefficient and the closeness centrality, we wished to investigate whether proteins associated with particular PTM-types exhibit distinct interaction characteristics that may be indicative of a PTM-specific function. The degree quantifies the average number of connections a protein engages in. Thus, it reflects on how many interaction partners may be affected by a PTM of a given protein. The clustering coefficient allows estimating whether the proteins connected to a central reference protein are in turn connected amongst themselves. High clustering coefficients would indicate a closely knit network of local interactions, whereas low clustering coefficients would suggest that separate molecular processes with little communication between them are modulated, when a central protein undergoes a PTM. Finally, the closeness centrality allows assessing how centrally a particular protein resides relative to the overall network. Proteins with high closeness centrality are situated in central network positions such that they may serve central information relay functions. By contrast, low closeness centrality corresponds to peripheral locations as typical of initial receptor molecules. (For a formal definition of the three network properties, please see Methods.) Therefore, all three chosen network properties allow a direct interpretation of the specific function of protein modifications in the context of protein interactions.
PTMs with regard to impact (degree) and role as a potential information relay hub (clustering coefficient and closeness centrality) in the network of all interacting proteins in the cell. Fig. 4 shows the frequency distributions of the PTM-type-specific network properties exemplified in *Homo sapiens*. Overall, all PTM-types exhibit a tendency to have higher degrees, lower clustering coefficients, and higher closeness centralities than protein sets not carrying the respective PTM-type, which includes a set of human proteins (1,864 or 20.8% of all human proteins) currently not known to undergo any of the 12 PTM-types considered here. The latter

![Image](Figure 3. Significant co-existence pairs of PTM-types across all selected species. Edge width was set proportionally to the number of species in which a particular PTM pair was found to occur more frequently than expected (see legend to Fig. 2) at significance levels of FDR-corrected $p$-values < 0.01. The values on the edges indicate the number of species with significant co-existence normalized by the number of common species between each pair of PTM-types as not all PTM-types are present in all species based on our filtering criteria (see Methods). Numbers in parentheses are the normalized counts of significant “red” co-existence edges to other PTM-types.

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Table 2. PIN size (number of proteins and interactions) for the nine selected species after excluding the components with the size less than 100.

| NCBI taxonomy ID | Species Name     | STRING     | IntAct      | Common  |
|------------------|------------------|------------|-------------|---------|
|                  |                  | Proteins   | Interactions| Proteins | Interactions| Proteins   | Interactions|
| 10090            | *Mus musculus*   | 6599       | 55812       | 5983    | 14641       | 1068       | 1101       |
| 10116            | *Rattus norvegicus* | 3544     | 19394       | 1307    | 1866        | 180        | 156        |
| 9913             | *Bos taurus*     | 2899       | 16119       | 113     | 183         | 68         | 80         |
| 9606             | *Homo sapiens*   | 8949       | 71153       | 10588   | 57922       | 3592       | 7556       |
| 7227             | *Drosophila melanogaster* | 4319 | 50039       | 7280    | 24368       | 1052       | 1164       |
| 6239             | *Caenorhabditis elegans* | 3796 | 32247       | 3604    | 7785        | 684        | 719        |
| 3702             | *Arabidopsis thaliana* | 5528   | 34129       | 3499    | 8534        | 1186       | 2181       |
| 4932             | *Saccharomyces cerevisiae* | 4743   | 63169       | 5492    | 68891       | 3103       | 11723      |
| 4896             | *Schizosaccharomyces pombe* | 2748   | 22235       | 0       | 0           | 0          | 0          |

High confidence PINs extracted from STRING are listed in the “STRING” column, the “IntAct” column contains the PIN sizes extracted from IntAct. The “Common” column indicates the number of common proteins and interactions in STRING and IntAct. Only those proteins with available UniProt ID were considered.

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set (no PTM), was observed with lower degree, higher clustering coefficient, and lower closure centrality than proteins undergoing a PTM (S2 Fig.). With regard to degree, the largest increases relative to the respective reference sets were observed for sumoylation, proteolytic cleavage, and amidation, albeit for the latter the count of observed instances is low. By contrast, glycosylation shows almost no change of degree relative to its reference set. With regard to the clustering coefficient, sumoylation, proteolytic cleavage, and carboxylation were found with the largest decreases relative to their respective control sets. Finally, sumoylation, proteolytic cleavage, and amidation were the top-three PTM-types associated with the largest relative increase in their median closeness centrality compared to their proteins sets devoid of the respective PTM-type. Again, glycosylation was found with the smallest relative change with regard to closeness centrality. Thus, excluding the PTM-types with very low counts (amidation and carboxylation), sumoylation and proteolytic change were identified as the two PTM-types associated with the largest relative differences across all three network properties examined. In short, both are characterized by high degree, low clustering coefficient, and high closeness centrality. Glycosylation is found at the other end of the spectrum with no change with regard to degree and closeness centrality, but a drop in clustering coefficient. The three most abundant PTM-types in human—based on available data—acetylation, phosphorylation, and ubiquitination, all show significant and comparable degree and closeness centrality increases. With regard to closeness centrality, phosphorylation is signified by the largest drop among the three PTM-types associated with the largest relative differences across all three network properties examined. In short, both are characterized by high degree, low clustering coefficient, and high closeness centrality. Glycosylation is found at the other end of the spectrum with no change with regard to degree and closeness centrality, but a drop in clustering coefficient. The three most abundant PTM-types in human—based on available data—acetylation, phosphorylation, and ubiquitination, all show significant and comparable degree and closeness centrality increases. With regard to closeness centrality, phosphorylation is signified by the largest drop among the three PTM-types associated with the largest relative differences across all three network properties examined. In short, both are characterized by high degree, low clustering coefficient, and high closeness centrality.

Next, we expanded our analyses to the remaining eight species considered here (Fig. 5). To avoid database specific effects, we considered two sources of PIN information, STRING and IntAct [41, 42]. While STRING contains integrated interactions from different sources, IntAct contains experimentally verified interactions extracted from literature and based on direct user submissions. Except for a few cases (13 out of 86 with data available for both PIN-resources), consistent results across the two PIN-data resources were obtained. Furthermore, for some of the IntAct derived PIN-properties, we detected a difference in their mean value compared to their median, reflecting the lower counts of IntAct events resulting in non-Gaussian/asymmetric
distribution. Furthermore, truly comparing the different PTM-types across several species is possible only for the PTM-types acetylation, glycosylation, and phosphorylation, as for the others sufficient PIN-information is lacking. Also, we restricted the analyses to the addition-type PTM-types with sufficient data acetylation, glycosylation, methylation, phosphorylation, nitrosylation, sumoylation, and ubiquitination.

Despite these data limitations, a similar overall picture emerges as observed in human. All examined PTM-types appear to be associated with proteins with increased degree and closeness centrality, but decreased clustering coefficient relative to protein sets not carrying the respective PTM-type. Glycosylation is a notable exception and seems associated with slightly decreased, rather than increased, degree centrality, while exhibiting similarly decreased clustering coefficient.

Figure 5. Degree, clustering coefficient, closeness centrality analysis for proteins with different PTM-types in each considered species and associated high-confidence STRING and IntAct PIN. The species are ordered according to their phylogenetic relationships as shown on the left. For every PTM-type, the log2 of fold difference value for the degree/clustering coefficient/closeness centrality value relative to the respective value associated with proteins not carrying this particular PTM-type are given for PINs based on STRING and IntAct, respectively. Color scale indicates increased (red) or decreased (blue) values in the PTM-set relative to the non-PTM-set with symmetric color intervals (i.e. full color saturation based on the maximal absolute increase or decrease fold difference observed across all values in the table.) Bold-font (underlined) fold-changes indicate significant fold-changes at p < 0.05 (p < 0.01) by Mann-Whitney test with FDR correction, the values in red or blue text represent significantly higher or lower network properties, which are inconsistent with the background color based on mean (not median) values. PTM-types “carboxylation”, “proteolytic cleavage”, “hydroxylation”, and “disulfide bond” are not included in this analysis as associated numbers were available for Homo sapiens only.

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clustering coefficients across all species as the other PTM-types. With regard to closeness centrality, the results for glycosylation are mixed with a few species (the four mammalian species and the plant \textit{Arabidopsis thaliana}) showing slightly increased values for STRING-based PINs and decreased values in the remaining species. However, clearly more table cells are colored blue signifying lowered values than for the other PTM types further supported by negative fold-change values for IntAct PINs. Thus, overall, glycosylation appears to generally be associated with no significantly changed, or slightly lowered closeness centrality relative to control protein sets.

Several of the PTM-types inspected here can be subdivided further into separate sets based on the identity of the targeted group on the protein (Lysine vs. N-terminal acetylation, N- or O-linked glycosylation, arginine-/lysine-methylation, and serine/threonine and tyrosine phosphorylation. Especially in the case of S/T vs. Y-phosphorylation differences in the associated PIN-properties would be of interested given the importance of phosphorylation in general, and in particular, as the two different types are catalyzed by different kinases \cite{43}. However, when subdividing the protein sets into the target-specific PTM types, consistent results were obtained as reported for the merged sets (S3 Fig.). As the only notable exception, the difference in degree observed for O-linked glycosylation (trend for increased degree relative to reference set) compared to N-linked glycosylation (trending towards decreased degree) is worth mentioning.

In an attempt to address a possible confounding influence of co-occurring different PTM-types on the same protein, we repeated the analysis of PTM-type-specific PIN properties shown in Fig. 5 for sets of proteins that are annotated to undergo one PTM-type only (S4 Fig.). Inevitably, this dramatically reduced the number of proteins that can be used (S1 Table). Hence, a meaningful analysis was possible for four PTM-types (acetylation, glycosylation, phosphorylation, and ubiquitination) only. Again, some conflicts between STRING and IntAct derived results render drawing clear conclusions difficult. However, phosphorylation again comes out as being associated with high-degree, low clustering coefficient, and high closeness centrality proteins compared to reference unphosphorylated protein sets. Glycosylation, on the other hand, is found again with low degree, low clustering coefficient, and low closeness centrality compared to control sets of proteins that are not glycosylated. Acetylation and ubiquitination both appear less consistent with the results reported for the whole protein set (Fig. 5). Ubiquitination was found with low degree, high clustering coefficient, and low closeness centrality; i.e. opposite the trend reported in Fig. 5. For acetylation, no clear trends are evident also because of many conflicts between STRING and IntAct based results. Thus, PIN-properties for phosphorylation and glycosylation are confirmed in the unique protein sets, whereas acetylation and ubiquitination either behave differently when protein sets are properly reduced to unique sets, or clear conclusions cannot be drawn as of yet because of data limitations.

Cross-protein PTM interaction patterns

Above, we examined the co-occurrence of different PTM-types detected on the same protein (Fig. 3). We extended the co-occurrence analysis to pairwise physically interacting proteins carrying different PTMs based on protein sets characterized by one PTM-type only (Fig. 6). For all PTM-types but disulfide-bond proteins, there is a tendency to self-interact, i.e. two separate proteins carrying the same PTM-type interact more often than randomly expected. Phosphorylated proteins display the broadest interaction range with significantly more interactions than expected to 8 other PTM-type proteins, followed by glycosylated proteins (6 distinct PTM-type partners), and acetylation (5 distinct PTM-type partners). By contrast, proteins associated with methylation, disulfide-bond formation or amidation exhibit a reduced interaction spectrum with likely interactions to only three of fewer other PTM-type proteins including interactions between two proteins carrying the same PTM-type. Acetylation, glycosylation, and
phosphorylation form a clique of more than expected interactions among these three PTM-types. Especially phosphorylated proteins were found to interact more often with acetylated proteins than expected by chance. Please note that the analysis displayed in Fig. 6 is controlled for abundance; i.e., the reported interactions are above the expected chance-encounters. Interaction statistics including proteins including those with multiple different PTM-types is provided as S5 Fig. As frequently, proteins carry multiple different PTM-types, conclusions with regard to preferred cross-protein PTM-type interactions is less meaningful. However, the trends described above are apparent as well.

Disease association of PTM-type-specific network properties in human
We tested whether proteins are more likely to be implicated in human disease when their associated PIN property values were at the high or low end of the spectrum. Most significantly for phosphorylation, but also evident for ubiquitination, acetylation, and glycosylation, we detected a larger than expected overlaps with known human disease proteins for high-degree proteins only, but not for proteins undergoing the same PTM-type but with low interaction degree proteins. Similarly, glycosylated and phosphorylated proteins are more likely disease associated when they have high closeness centrality. By contrast, low clustering coefficient appears correlated with disease association for proteins undergoing phosphorylation, acetylation, and ubiquitination (Table 3).

Discussion
The modulation of protein function via different types of post-translational modifications (PTMs) and their combinatorial interplay has attracted considerable attention in recent years [15–27]. In this study, we added the interaction layer to the study of PTMs by performing a systematic investigation of the network properties of the different PTM-types in the context of the physical interactions of PTM-carrying proteins. For twelve different PTM-types and across nine diverse species, we determined characteristic and informative network parameters with the goal to investigate whether particular PTM-types are associated with specific and possibly
strategic placements in the context of all protein interactions such that their individual role in the orchestration of the combined action of all proteins becomes apparent. Generalized across all PTM-types and species investigated here, PTM-carrying proteins appear engage in more physical contacts, with a reduced clustering coefficient among those proteins they are interacting with, and elevated closeness centrality than their respective protein sets devoid of the particular PTM-type (Fig. 5) or that, as far as we currently know, do not harbor any PTM of any type (S2 Fig.). Differences between the twelve studied PTM-types proved less pronounced with essentially all—except for glycosylation (see below)—following the same trend of high degree, low clustering coefficient, and high closeness centrality with only subtle differences in magnitude between them. However, given the present data coverage, it is not yet possible to conclusively decide whether these differences are statistically significant and biologically relevant. When further subsetted into special types of PTMs (e.g. S/T/Y phosphorylation), no significant sub-type differences were evident (S3 Fig.). As motivated above, the three selected network properties were selected specifically to allow conclusions as for the “strategic” roles of PTM in the context of interactions. According to this logic, proteins with PTMs engage in more and different process than non-PTM proteins and play central information relay functions.

Focusing on human PIN and PTM data, sumoylation and proteolytic cleavage stand out as being associated with the largest relative increase of degree and closeness centrality relative to reference sets. Proteolytic cleavage has been associated with activation processes and protein targeting events (cleavage of targeting N-terminal peptide) and constitute a “dramatic” modification as the relative change of molecular composition of a protein can be significant. Furthermore,

| PTM-type           | degree | clustering coefficient | closeness centrality |
|--------------------|--------|------------------------|----------------------|
| acetylation        | 4.94E-07 | 2.52E-03               | 9.94E-01             |
| amidation          | 1.00E+00 | 1.00E+00               | 1.00E+00             |
| carboxylation      | 1.57E-01 | 9.36E-01               | 6.37E-02             |
| disulfide bond     | 9.49E-02 | 1.00E+00               | 5.05E-01             |
| glycosylation      | 1.04E-05 | 2.42E-01               | 8.15E-05             |
| hydroxylation      | 1.00E+00 | 3.46E-01               | 1.00E+00             |
| methylation        | 4.35E-02 | 1.00E+00               | 1.00E+00             |
| nitrosylation      | 1.00E+00 | 1.00E+00               | 7.10E-01             |
| phosphorylation    | 9.41E-17 | 4.94E-07               | 2.89E-03             |
| proteolytic cleavage| 4.71E-02 | 7.16E-01               | 6.62E-02             |
| sumoylation        | 2.42E-01 | 5.05E-01               | 4.71E-02             |
| ubiquitination     | 5.06E-09 | 3.70E-03               | 2.10E-01             |

Protein sets were sorted in descending order of the respective PIN-property and the top/bottom 25% tested for overlap with human disease genes reported in OMIM (http://www.ncbi.nlm.nih.gov/omim) based on a Fisher’s exact test with FDR-correction with respective counts of proteins binned according to binary decisions top/bottom 25% property and disease association yes/no forming the 2x2 contingency table. p-values highlighted red indicate significant overlap of the top-25% set of proteins for a particular network property with human disease proteins, while blue values signify larger than expected overlaps of the bottom-25% protein sets with proteins annotated to be involved in disease processes at p-value thresholds of p-value 0.05, 0.01 (if underlined), respectively. Black font color indicates no significant overlap of neither the top nor the bottom 25% set.

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transporting proteins to different compartments will inevitably influence the possible interaction scope. The significance of sumoylation in a range of regulatory processes has been increasingly recognized [44]. Our results underscore the importance of this PTM-type.

Phosphorylation, the PTM-type with the largest data support, was identified as the PTM-type with the consistently central and with the largest potential influence scope (Fig. 5). Phosphorylated proteins reside in central network positions (high closeness centrality) and interact with many other proteins (high degree) including specifically pairwise interactions with proteins carrying any of the other four PTM-types as well as other phosphorylated proteins (Fig. 6 — pairwise interaction figure). Examples from human of phospho-proteins interacting with proteins carrying other PTM-types include the kinases: mitogen-activated protein kinase 1 (MAPK1), interleukin-1 receptor-associated kinase 2 (IRAK2), and spleen tyrosine kinase (SYK). Those proteins each interact with other proteins representing four different PTM-types. These findings underscore once again the central importance of phosphorylation as perhaps the most important and central PTM-type identified so far. Similar characteristics were found for acetylation, albeit the detected magnitude and statistical support is lower.

By contrast, glycosylation was found associated with proteins of low degree, low clustering coefficient, and low closeness centrality (Fig. 5). In particular the low degree and low closeness centrality of glycosylated proteins may be interpreted as consistent with their preferred location in cytosolic membranes and to act as receptors and cell-cell communication mediators (Fig. 1, GO-term clustering) [45]. Unlike the other four PTM-types, the transferred glycosyl-groups can be large leading to impeded protein-protein interactions of glycosylated proteins. In addition, because of their frequent embedding in membranes, they operate in two dimensions, not three as for soluble cytosolic proteins, effectively cutting down the interaction potential.

As shown in Fig. 6, all PTM-types are found on proteins that exhibit a tendency to interact with other proteins carrying the same PTM-type. In the case of phosphorylation, such interactions are interpretable as the well known as phosphorylation/kinase cascades [46, 47]. It is also possible that the detected tendency of PTM-types to self-interact originates from protein complexes, in which all partners undergo the PTMs of the same type. For example, in histone complexes, lysine residues on different proteins are acetylated modifying the binding affinity of histones to DNA [48, 49]. Similar considerations apply to methylation events in histone [50] and other protein complexes [51].

By including nine species from different kingdoms and lineages, we aimed to extract both general and species/lineage-specific trends. However, currently available datasets proved comprehensive enough for a few species only (human, mouse, rat). In the case of phosphorylation, sufficient data were available across all nine species and provided a consistent result of increased degree and closeness centrality and a decreased clustering coefficient (Fig. 5). The increased likelihood of a functional association of proteins with high interaction degree and their involvement in human disease has been reported before [52, 53]. In selected cases, proteins carrying PTMs have also been reported to be more likely related to disease processes than non-PTM proteins [54–56]. Our dataset allowed us to expand this analysis to testing specific PTM-types combined with their PIN-characteristics. Our results suggest that not only does a PTM render proteins more likely disease associated, but that this association may depend on what PIN context it is embedded in. High degree, low clustering coefficient, and high closeness centrality proteins are more likely to be disease associated (Table 3) than their respective counterpart sets at the respective other end of the property PIN-property spectrum, especially for the PTM-types phosphorylation and glycosylation, albeit it for the latter, no significant clustering coefficient trend was detected. Examples of disease-associated phosphorylated or glycosylated proteins detected with high degree and closeness centrality or low clustering coefficient are provided in Table 4. It may be speculated that proteins with the
properties identified as more likely disease associated based on their PIN properties may constitute promising candidates for intensified research. Evidently, the relevance of the protein p53 in human cancer development has long been recognized [57]. In our study, it was identified as one with characteristic network properties typical of disease associated proteins in general.

Evidently, this study hinges on the completeness and accuracy of the available PTM and PIN data as well. Any bias towards a specific detection of particular protein classes and their associated PTM may further skew our results. By imposing a high significance cutoff for the PIN-data (confidence score > 0.9), and furthermore exploiting two data sources (STRING and IntAct), we believe to have taken proper precautionary steps even though some discrepancies were detected (Fig. 5). However, at this point it cannot be decided whether the size of the

| PTM-type | PIN-characteristics | Protein | Ensembl ID | Disease |
|----------|---------------------|---------|------------|---------|
| Glycosylation | high degree, high closeness centrality | Pro-epidermal growth factor | ENSP00000265171 | Hypomagnesemia 4 (HOMG4) [MIM:611718][67] |
| | | Transforming growth factor beta-1 | ENSP00000221930 | Camurati-Engelmann disease (CE) [MIM:131300][68] |
| | | Interleukin-6 | ENSP00000258743 | Rheumatoid arthritis systemic juvenile (RASJ) [MIM:604302][69] |
| Phosphorylation | high degree, high closeness centrality | Cellular tumor antigen p53 | ENSP00000269305 | Esophageal cancer (ESCR) [MIM:133239][70]; Li-Fraumeni syndrome (LFS) [MIM:151623][71] |
| | | RAC-alpha serine/threonine-protein kinase | ENSP00000270202 | Breast cancer (BC) [MIM:114480]; Colorectal cancer (CRC) [MIM:114500][72]; Proteus syndrome (PROTEUS) [MIM:176920] [73] |
| Phosphorylation | low clustering coefficient | Histone acetyltransferase p300 | ENSP00000263253 | Rubinstein-Taybi syndrome 2 (RSTS2) [MIM:613684][74] |
| | | Autoimmune regulator | ENSP00000291582 | Autoimmune polyendocrine syndrome 1, with or without reversible metaphyseal dysplasia (APS1) [MIM:240300] [75] |
| | | ALK tyrosine kinase receptor | ENSP00000373700 | Neuroblastoma 3 (NBLST3) [MIM:613014][76] |
| | | Ataxin-2 | ENSP00000366843 | Spinocerebellar ataxia 2 (SCA2) [MIM:183090][77]; Amyotrophic lateral sclerosis 13 (ALS13) [MIM:183090] [78] |

Table 4. Examples of human PTM-carrying proteins with characteristic PIN-properties and their known disease involvement.

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dataset (relatively small IntAct data set) or the type of PINs that are recorded cause these differences. With regard to PTMs, we used experimentally verified PTMs only. Future investigations of the PIN characteristics of PTMs will benefit from the expected significant increase of experimentally verified sites. In addition, a larger set of different PTMs with sufficient numbers will likely become available, allowing also to further specify the PTM-types used in this study.

A possible selection bias may also come from preferentially profiling those proteins for PTMs that possess “interesting” properties such as high degree. However, as PTMs are increasingly identified in massive, “shotgun” style omics studies, such selection bias may not be that critical. Rather, abundance may be a concern then. However, for phosphorylation it was reported that protein abundance is not correlated with network properties [34, 35]. Furthermore, we also found that network properties are largely independent of the number of PTMs on a given protein (S6 Fig.). While significant due to the large number of observations, no relevant correlation was found neither for degree nor clustering coefficient with the number of phosphorylation sites taken as the PTM-type with the largest available dataset. However, for closeness centrality, a more sizable positive correlation ($r = 0.164$) was detected suggesting that more heavily phosphorylated proteins occupy more central positions in the network of protein-protein interactions.

In conclusion, proteins carrying different types of PTMs differ from average non-PTM-proteins and differ between each other with regard to their protein interaction characteristics. Thus, their location within the web of physical protein-protein interactions is not only non-random, but very likely indicates their specific functional roles in the orchestration of molecular processes mediated by the physical interactions between proteins.

Materials and Methods

Datasets

Post-translational modifications. Post-translational modifications annotated as “experimentally verified” and the associated proteins were extracted from UniProt [58], PhosphoSitePlus [59], dbPTM [9], Phospho ELM [12], PhosphoGRID [60], PHOSIDA [13], HPRD [61], OglycBase [62], PhosPhAt [14], P3DB [63], PTMcode [11] as of 2014 April. Subsequently, the sets obtained from the different data sources were consolidated to create a single set based on protein sequence position information. Initially, only those PTM-types were considered further, for which more than 1000 sites were reported across the various data resources, regardless of species. Subsequently, only those species were retained for which the count of PTM-sites across all PTM-types was 1000 or more. The intersection of both sets yielded the primary PTM-type-species dataset for analysis. To remove outliers, extremely long or short proteins as determined by falling below the 1-percentile or above the 99-percentile of observed protein sequence length distribution were removed. Furthermore, proteins with an extremely high number of PTM sites greater than or equal to. $average(number of PTM sites)+3+sd(number of PTM sites)$ were discarded as well. Twelve PTM types met those criteria in at least one species including 10 PTM-types in which a chemical moiety is attached: acetylation, amidation, carboxylation, glycosylation, hydroxylation, methylation, nitrosylation, phosphorylation, sumoylation, ubiquitination, and two PTM-types that modify the protein via forming or breaking bonds within the protein: disulfide bond and proteolytic cleavage. Nine species met those criteria for at least one PTM-type including representatives from the animal (mammals: Mus musculus, Rattus norvegicus, Bos taurus, Homo sapiens; insects: Drosophila melanogaster, invertebrates: Caenorhabditis elegans), plant (Arabidopsis thaliana), and fungal (Saccharomyces cerevisiae, Schizosaccharomyces pombe) kingdom, respectively (Table 1).
Many proteins carry more than one PTM-type. Some analyses were meaningful only if the considered proteins carry one PTM-type only (e.g. interaction between proteins carrying different PTM-types). Then, protein sets were filtered further and referred to as “one-PTM-type-only” (S1 Table). For statistics of the associated sub-sets of the dataset, see S2 Table.

**Protein-protein interaction networks.** For the nine species selected based on available PTM information, high confidence (confidence score $\geq 0.9$) protein-protein interactions (PINs) were extracted from STRING (version 9.05) [41]. In order to avoid database biases, each species’ protein-protein interactions were also extracted from the IntAct database (downloaded on 9.26.2013) [42], which stores interactions derived from literature curation or direct user submissions. To remove isolated interactions significantly affecting some of the network properties (e.g. closeness centrality), the network components with the size less than 100 were excluded (S3 Table). Only one component (the “giant”) component was left for each selected species. The sizes of the different PINs (species and data source) are summarized in Table 2. As the overlap between the STRING and IntAct interaction set is relatively small, the two datasets can be seen as largely disjoint and independent.

**Network properties.** The PINs associated with the selected species correspond to undirected networks. The degree of a node $n$ is the number of edges linked to $n$ [64]. The clustering coefficient [64] of a node $n$ is defined as: $C_n = 2e_n/(k_n(k_n - 1))$, where $k_n$ is the number of neighbors of $n$ and $e_n$ is the number of connected pairs between all neighbors of $n$. It reflects the connectivity of adjacent nodes. The closeness centrality $C_c(n)$ of a node $n$ is defined as the reciprocal of the sum of shortest path length originating from $n$ to all other nodes $m$: $C_c(n) = \frac{1}{\sum_{m \neq n} L(n,m)}$, where $L(n,m)$ is the length of the shortest path between two nodes $n$ and $m$. It ranges between 0 and 1. It corresponds to the inverse of the number of steps needed to traverse from all other nodes in the network to a selected node. The R package “igraph” (http://cran.r-project.org/web/packages/igraph/index.html) was used to compute the above-mentioned network properties.

**Protein interaction network characteristics tests**

For each PTM-type and across the selected species, the three selected network properties (degree, clustering coefficient, and closeness centrality) were computed for all proteins carrying the respective PTM-type and compared to those proteins not carrying this particular PTM-type. Significant differences between the respective two distributions were detected based on a Mann–Whitney test. The $p$-values were corrected for multiple testing considering as the total number of tests all tests across species, all PTM-types for each network property. In all cases of multiple testing correction, the FDR method was used [65]. Significance testing was applied to only those PTM-types and species with 30 proteins instances or more.

**Biological function enrichment analysis**

For all species selected in this study, the available genome gene ontology (GO) process, function and cellular compartment annotations were extracted from UniProtKB-GOA [66] as the reference set. All the selected species were combined into one ‘species’ in the enrichment analysis. The method “elim” and the Fisher’s exact test with FDR were used for enrichment analysis using the “topGO” R package. The cutoff $P$-value was set to 0.01.

**Jaccard index for PTMs co-existence**

For each pair of PTMs (A and B) in one species, The Jaccard index for the co-existence of A and B is defined as $\frac{|\text{Intersection}(S_A,S_B)|}{|\text{Union}(S_A,S_B)|}$, where the set of proteins associated
with A and B are denoted as \( S_A \) and \( S_B \). Fisher’s exact test was used to test the significance of co-existence.

**Fisher’s exact test for cross-protein PTM interaction**

For each pair of PTM-types and across the selected species, Fisher’s exact test was designed to test the over and under protein interaction frequency. The \( p \)-values were corrected for multiple testing considering as the total number of tests across all pair of PTM-types for each species. The cutoff \( p \)-value was set to 0.01. Significance testing was applied to only those pairs of PTM-types in each species with at least 10 proteins instances or more separately.

**Overlap between human disease proteins and PTMs associated proteins**

Human disease proteins were downloaded from OMIM (http://www.ncbi.nlm.nih.gov/omim) as of May, 2014. In order to test the overlap between human diseases proteins with proteins associated with high (top 25%) or low (bottom 25%) network property values, Fisher’s exact test was used for all PTM-types. The \( p \)-values were corrected for multiple testing (FDR) across all PTM-types and network properties.

**Supporting Information**

**S1 Fig.** Heatmap for significant molecular function terms across all studied PTM-types. The top five GO-terms were included that were found significantly enriched for each PTM-type. Each element in the heat map (Euclidean distance hierarchical clustering, average linkage) represents the grey-scale-encoded \( p \)-value, in which a particular combination of PTM-type and GO-term was found significantly enriched. The combined whole UniProtKB-GOA for all the selected species was used as the background set, Fisher’s exact test with FDR correction was used for the enrichment analysis, and the \( p \)-value (FDR) threshold indicating significance was set to 0.01.

**S2 Fig.** Protein interaction network properties associated with proteins without any and with a PTM in *Homo sapiens*. The red (blue) asterisks on the top of violin plot represents the corresponding non-PTM group has a significantly higher (lower) median value compared to the non-PTM group (*: \( p \)-value 0.05, **: \( p \)-value 0.01) according to a Mann-Whitney test.

**S3 Fig.** Degree, clustering coefficient, closeness centrality analysis for proteins with different PTM-subtypes in each species and associated high-confidence STRING and IntAct PIN (refined dataset). The species are ordered according to their phylogenetic relationships as shown on the left. For every PTM-type, the log-2 of fold difference value for the degree/clustering coefficient/closeness centrality value relative to the respective value associated with proteins not carrying this particular PTM-type are given for PINs based on STRING and IntAct, respectively. Color scale indicates increased (red) or decreased (blue) values in the PTM-set relative to the non-PTM-set with symmetric color intervals (i.e. full color saturation based on the maximal absolute increase or decrease fold difference observed across all values in the table.) Bold-font (underlined) fold-changes indicate significant fold-changes at \( p<0.05 \) (\( p<0.01 \)) by Mann-Whitney test with FDR correction, the values in red or blue text represent significantly higher or lower network properties which are inconsistent with the background color.
S4 Fig. Degree, clustering coefficient, closeness centrality analysis for proteins with different PTM types in each species and associated high-confidence STRING and IntAct PIN. Protein sets were selected to contain one PTM-type only (one-PTM-type-only dataset). The species are ordered according to their phylogenetic relationships as shown on the left. For every PTM-type, the log-2 of fold difference value for the degree/clustering coefficient/closeness centrality value relative to the respective value associated with proteins not carrying this particular PTM-type are given for PINs based on STRING and IntAct, respectively. Color scale indicates increased (red) or decreased (blue) values in the PTM-set relative to the non-PTM-set with symmetric color intervals (i.e., full color saturation based on the maximal absolute increase or decrease fold difference observed across all values in the table.) Bold-font (underlined) fold-changes indicate significant fold-changes at $p < 0.05$ ($p < 0.01$) by Mann-Whitney test with FDR correction, the values in red or blue text represent significantly higher or lower network properties, which are inconsistent with the background color based on mean (not median) values.

(TIF)

S5 Fig. Pairwise interactions of proteins carrying different PTM-types (full dataset). Increased frequencies of protein-protein interactions (designated as protein A and B, respectively) carrying the respective PTM-types relative to expectation. Line width is proportional to the number of species, which exhibit significant interactions of PTM-types carried by interacting proteins. The contingency table for the Fisher exact test contained the respective counts for number of proteins associated with a particular PTM-pair versus all alternative pairings and whether they have been reported to interact or not with applied FDR-corrected p-value threshold of $<0.01$.

(TIF)

S6 Fig. Scatterplot between the number of PTMs, here phosphorylation sites, and corresponding network properties (degree, clustering coefficient, and closeness centrality) for phosphorylated proteins based on the STRING network (the giant component was considered only) of Homo sapiens. The red line connects the mean network property values of proteins associated with different numbers of PTMs. Associated Pearson linear correlation coefficients, $r$, (and p-values) were: degree $r = 0.056$ ($1.46\times10^{-6}$), clustering coefficient: $r = -0.068$ ($6.27\times10^{-8}$), closeness centrality: $r = 0.164$ ($0.00$).

(TIF)

S1 Table. Frequency table of proteins associated with the selected PTM-types and species after excluding proteins with more than one PTM-type (one-PTM-type-only dataset); i.e. every protein was annotated to harbor one PTM-type only.

(DOCX)

S2 Table. Frequency table of proteins associated with the selected PTM-subtypes and species (PTM-subtypes dataset).

(DOCX)

S3 Table. PIN size (number of proteins and interactions) for the nine selected species before excluding the components of size less than 100.

(DOCX)

Author Contributions

Conceived and designed the experiments: GD DW. Performed the experiments: GD. Analyzed the data: GD DW. Wrote the paper: GD DW.
References

1. Lodish H, Berk A, Zipursky SL, Matsudaira P, Baltimore D, et al. (2000) Molecular cell biology.
2. Human Genome Sequencing Consortium (2004) Finishing the euchromatic sequence of the human genome. Nature 431: 931–945. doi: 10.1038/nature03001
3. Jensen ON (2004) Modification-specific proteomics: characterization of post-translational modifications by mass spectrometry. Curr Opin Chem Biol 8: 33–41. doi: 10.1016/j.cbpa.2003.12.009 PMID: 15036154
4. Deribe YL, Pawson T, Dikic I (2010) Post-translational modifications in signal integration. Nat Struct Mol Biol 17: 666–672. doi: 10.1038/nsmb.1842 PMID: 20495563
5. Zhao S, Xu W, Jiang W, Yu W, Lin Y, et al. (2010) Regulation of cellular metabolism by protein lysine acetylation. Science 327: 1000–1004. doi: 10.1126/science.1179689 PMID: 20167786
6. The UniProt Consortium (2010) The Universal Protein Resource (UniProt) in 2010. Nucleic Acids Res 38: D142–8. doi: 10.1093/nar/gkp846 PMID: 19843607
7. Kim SC, Sprung R, Chen Y, Xu Y, Ball H, et al. (2006) Substrate and functional diversity of lysine acetylation revealed by a proteomics survey. Mol Cell 23: 607–618. doi: 10.1016/j.molcel.2006.06.026 PMID: 16916647
8. The UniProt Consortium (2013) Activities at the Universal Protein Resource (UniProt). Nucleic Acids Res 42: D191–D198. doi: 10.1093/nar/gkt1140 PMID: 24253303
9. Lu C-T, Huang K-Y, Su M-G, Lee T-Y, Bretaña NA, et al. (2013) dbPTM 3.0: an informative resource for investigating substrate site specificity and functional association of protein post-translational modifications. Nucleic Acids Res 41: D295–D305. doi: 10.1093/nar/gks1229 PMID: 23193290
10. Khoury GA, Baliban RC, Floudas CA (2011) Proteome-wide post-translational modification statistics: frequency analysis and curation of the swiss-prot database. Sci Rep 1: 90. doi: 10.1038/srep00090
11. Minguez P, Letunic I, Parca L, Bork P (2013) PTMcode: a database of known and predicted functional associations between post-translational modifications in proteins. Nucleic Acids Res 41: D306–D311. doi: 10.1093/nar/gks1230 PMID: 23193284
12. Dinkel H, Chica C, Via A, Gould CM, Jensen LJ, et al. (2011) Phospho.ELM: a database of phosphorylation sites–update 2011. Nucleic Acids Res 39: D261–7. doi: 10.1093/nar/gkq1104 PMID: 21062810
13. Gnaf D, Gunawardena J, Mann M (2011) PHOSIDA 2011: the posttranslational modification database. Nucleic Acids Res 39: D253–60. doi: 10.1093/nar/gkq1159 PMID: 21081558
14. Zulawski M, Braginets R, Schulze WX (2013) PhosPhAt goes kinases–searchable protein kinase target information in the plant phosphorylation site database PhosPhAt. Nucleic Acids Res 41: D1176–1184. doi: 10.1093/nar/gks1081 PMID: 23172287
15. Choudhary C, Kumar C, Gnaf D, Nielsen ML, Rehman M, et al. (2009) Lysine acetylation targets protein complexes and co-regulates major cellular functions. Science 325: 834–840. doi: 10.1126/science.1175371 PMID: 19608861
16. Zielinska DF, Gnaf F, Wiciejski JR, Mann M (2010) Precision mapping of an in vivo N-glycoproteome reveals rigid topological and sequence constraints. Cell 141: 897–907. doi: 10.1016/j.cell.2010.04.012 PMID: 20510933
17. Oliveira AP, Ludwig C, Picotti P, Kogadeeva M, Aebersold R, et al. (2012) Regulation of yeast central metabolism by enzyme phosphorylation. Mol Syst Biol 8: 623. doi: 10.1038/msb.2012.55 PMID: 23149688
18. Ubersax JA, Ferrell JE (2007) Mechanisms of specificity in protein phosphorylation. Nat Rev Mol Cell Biol 8: 530–541. doi: 10.1038/nrm2235 PMID: 17958314
19. Roux PP, Thibault P (2013) The coming of age of phosphoproteomics: from large data sets to inference of protein functions. Mol Cell Proteomics 12: 3453–3464. doi: 10.1074/mcp.R113.032862 PMID: 24037665
20. Brooks CL, Gu W (2003) Ubiquitination, phosphorylation and acetylation: the molecular basis for p53 regulation. Curr Opin Cell Biol 15: 164–171. doi: 10.1016/S0955-0674(03)00003-6 PMID: 12648672
21. Latham JA, Dent SYR (2007) Cross-regulation of histone modifications. Nat Struct Mol Biol 14: 1017–1024. doi: 10.1038/nsmb.1307 PMID: 17984964
22. Danielsen JMR, Sylvestersen KB, Bekker-Jensen S, Szklarczyk D, Poulson JW, et al. (2011) Mass spectrometric analysis of lysine ubiquitylation reveals promiscuity at site level. Mol Cell Proteomics 10: M110.003590. doi: 10.1074/mcp.M110.003590 PMID: 21139048
23. Hunter T (2007) The age of crosstalk: phosphorylation, ubiquitination, and beyond. Mol Cell 28: 730–738. doi: 10.1016/j.molcel.2007.11.019 PMID: 18082598
24. Van Noort V, Seebacher J, Bader S, Mohammed S, Vonkova I, et al. (2012) Cross-talk between phosphorylation and lysine acetylation in a genome-reduced bacterium. Mol Syst Biol 8: 571. doi: 10.1038/msb.2012.4 PMID: 22373819

25. Woodsmith J, Kamburov A, Stelzl U (2013) Dual Coordination of Post Translational Modifications in Human Protein Networks. PLoS Comput Biol 9: e1002933. doi: 10.1371/journal.pcbi.1002933 PMID: 23505349

26. Minguez P, Parca L, Diella F, Mende DR, Kumar R, et al. (2012) Deciphering a global network of functionally associated post-translational modifications. Mol Syst Biol 8: 599. doi: 10.1038/msb.2012.31 PMID: 22806145

27. Beltrao P, Bork P, Krogan NJ, van Noort V (2013) Evolution and functional cross-talk of protein post-translational modifications. Mol Syst Biol 9: 714. doi: 10.1002/msb.201304521 PMID: 24366814

28. Gavin A-C, Bösche M, Krause R, Grandi P, Marzioch M, et al. (2002) Functional organization of the yeast proteome by systematic analysis of protein complexes. Nature 415: 141–147. doi: 10.1038/nature00380 PMID: 11805826

29. Seet BT, Dikic I, Zhou M-M, Pawson T (2006) Reading protein modifications with interaction domains. Nat Rev Mol Cell Biol 7: 473–483. doi: 10.1038/nrm1960 PMID: 16829979

30. Arabidopsis Interactome Mapping Consortium (2011) Evidence for Network Evolution in an Arabidopsis Interactome Map. Science (80-) 333: 601–607. doi: 10.1126/science.1203877

31. De Las Rivas J, Fontanillo C (2012) Protein-protein interaction networks: unraveling the wiring of molecular machines within the cell. Brief Funct Genomics 11: 489–496. doi: 10.1093/bfgp/els036 PMID: 22908212

32. Nishi H, Hashimoto K, Panchenko AR (2011) Phosphorylation in protein-protein binding: effect on stability and function. Structure 19: 1807–1815. doi: 10.1016/j.str.2011.09.021 PMID: 22153503

33. Vinayagam a., Stelzl U, Fouill e R, Plassmann S, Z enkner M, et al. (2011) A Directed Protein Interaction Network for Investigating Intracellular Signal Transduction. Sci Signal 4: rs8–rs8. doi: 10.1126/scisignal.2001699 PMID: 21900206

34. Yachie N, Saito R, Sugiyama N, Tomita M, Ishihama Y (2011) Integrative Features of the Yeast Phosphoproteome and Protein–Protein Interaction Map. PLoS Comput Biol 7: e1001064. doi: 10.1371/journal.pcbi.1001064 PMID: 21298081

35. Duan G, Walther D, Schulze W (2013) Reconstruction and analysis of nutrient-induced phosphorylation networks in Arabidopsis thaliana. Front Plant Sci 4: 540. doi: 10.3389/fpls.2013.00540 PMID: 24400017

36. Walsh CT, Gameau-Tsodivoka S, Gatto GJ (2005) Protein posttranslational modifications: the chemistry of proteome diversifications. Angew Chem Int Ed Engl 44: 7342–7372. doi: 10.1002/anie.200501023 PMID: 16267872

37. Creixell P, Linding R (2012) Cells, shared memory and breaking the PTM code. Mol Syst Biol 8: 598. doi: 10.1038/msb.2012.33 PMID: 22866336

38. Nguyen LK, Kolch W, Kholodenko BN (2013) When ubiquitination meets phosphorylation: a systems biology perspective of EGFR/MAPK signalling. Cell Commun Signal 11: 52. doi: 10.1186/1478-811X-11-52 PMID: 23902637

39. Chan C-P, Mak T-Y, Chin K-T, Ng IO-L, Jin D-Y (2010) N-linked glycosylation is required for optimal proteolytic activation of membrane-bound transcription factor CREB-H. J Cell Sci 123: 1438–1448. doi: 10.1242/jcs.067819 PMID: 20356926

40. Tweedie-Cullen RY, Reck JM, Mansuy IM (2009) Comprehensive mapping of post-translational modifications on synaptic, nuclear, and histone proteins in the adult mouse brain. J Proteome Res 8: 4966–4982. doi: 10.1021/pr9003739 PMID: 19737024

41. Franceschini A, Szklarczyk D, Frankild S, Kuhn M, Simonovic M, et al. (2013) STRING v9.1: protein-protein interaction networks, with increased coverage and integration. Nucleic Acids Res 41: D808–D815. doi: 10.1093/nar/gks1094 PMID: 23203871

42. Kerrien S, Aranda B, Breuza L, Bridge A, Broackes-Carter F, et al. (2012) The IntAct molecular interaction database in 2012. Nucleic Acids Res 40: D841–D846. doi: 10.1093/nar/gkr1088 PMID: 22121220

43. Manning G, Whyte DB, Martinez R, Hunter T, Sudarsanam S (2002) The protein kinase complement of the human genome. Science (80-) 298: 1912–1934. doi: 10.1126/science.1075762

44. Flotho A, Melchior F (2011) Sumoylation: A Regulatory Protein Modification in Health and Disease. Annu Rev Biochem 82: 357–385. doi: 10.1146/annurev-biochem-061909-093311 PMID: 23746258

45. Moremen KW, Tiemeyer M, Nairn A V (2012) Vertebrate protein glycosylation: diversity, synthesis and function. Nat Rev Mol Cell Biol 13: 448–462. doi: 10.1038/nrm3383 PMID: 22722607
46. Popescu SC, Popescu G V, Bachan S, Zhang Z, Gerstein M, et al. (2009) MAPK target networks in Arabidopsis thaliana revealed using functional protein microarrays. Genes Dev 23: 80–92. doi: 10.1101/gad.174009 PMID: 19095804

47. Keshet Y, Seger R (2010) The MAP kinase signaling cascades: a system of hundreds of components regulates a diverse array of physiological functions. Methods Mol Biol 661: 3–38. doi: 10.1007/978-1-60761-795-2_1 PMID: 20811974

48. Grunstein M (1997) Histone acetylation in chromatin structure and transcription. Nature 389: 349–352. doi: 10.1038/38664 PMID: 9311776

49. Struhl K (1998) Histone acetylation and transcriptional regulatory mechanisms. Genes Dev 12: 599–606. doi: 10.1101/gad.12.5.599 PMID: 9499396

50. Zhang Y, Reinberg D (2001) Transcription regulation by histone methylation: interplay between different covalent modifications of the core histone tails. Genes Dev 15: 2343–2360. doi: 10.1101/gad.927301 PMID: 11562345

51. Boisvert F-M, Côté J, Boulanger M-C, Richard S (2003) A proteomic analysis of arginine-methylated protein complexes. Mol Cell Proteomics 2: 1319–1330. doi: 10.1074/mcp.M300088-MCP200 PMID: 14534352

52. Wachi S, Yoneda K, Wu R (2005) Interactome-transcriptome analysis reveals the high centrality of genes differentially expressed in lung cancer tissues. Bioinformatics 21: 4205–4208. doi: 10.1093/bioinformatics/bti688 PMID: 16188928

53. Jonsson PF, Bates P a (2006) Global topological features of cancer proteins in the human interactome. Bioinformatics 22: 2291–2297. doi: 10.1093/bioinformatics/btl390 PMID: 16844706

54. Anbalagan M, Huderson B, Murphy L, Rowan BG (2012) Post-translational modifications of nuclear receptors and human disease. Nucl Recept Signal 10: e001. doi: 10.1621/nrs.10001 PMID: 22438791

55. Ito K (2007) Impact of post-translational modifications of proteins on the inflammatory process. Biochem Soc Trans 35: 281–283. doi: 10.1042/BST0350281 PMID: 17371260

56. Vidal CJ (2011) Post-Translational Modifications in Health and Disease. doi: 10.1007/978-1-4419-6382-6

57. Levine AJ, Momand J, Finlay CA (1991) The p53 tumour suppressor gene. Nature 351: 453–456. doi: 10.1038/351453a0 PMID: 2046748

58. The UniProt Consortium (2014) Activities at the Universal Protein Resource (UniProt). Nucleic Acids Res 42: 7486. doi: 10.1093/nar/gku469

59. Hornbeck P V, Komhauser JM, Tkachev S, Zhang B, Skrzypek E, et al. (2012) PhosphoSitePlus: a comprehensive resource for investigating the structure and function of experimentally determined post-translational modifications in man and mouse. Nucleic Acids Res 40: D261–D70. doi: 10.1093/nar/gkr1122 PMID: 22135298

60. Sadowski I, Breitkreutz B-J, Stark C, Su T-C, Dahabieh M, et al. (2013) The PhosphoGRID Saccharomyces cerevisiae protein phosphorylation site database: version 2.0 update. Database 2013: bat026. doi: 10.1093/database/bat026

61. Goel R, Harsha HC, Pandey A, Prasad TSK (2012) Human Protein Reference Database and Human Proteinpedia as resources for phosphoproteome analysis. Mol Biosyst 8: 453–463. doi: 10.1039/c1mb05340j PMID: 22159192

62. Gupta R, Birch H, Rapacki K, Bruna S, Hansen JE (1999) O-GLYCBASE version 4.0: a revised database of O-glycosylated proteins. Nucleic Acids Res 27: 370–372. doi: 10.1093/nar/27.1.370 PMID: 9847232

63. Yao Q, Ge H, Wu S, Zhang N, Chen W, et al. (2014) P3DB 3.0: From plant phosphorylation sites to protein networks. Nucleic Acids Res 42: D1206–D1213. doi: 10.1093/nar/gkt1135 PMID: 24243849

64. Assenov Y, Ramirez F, Schelhom S-E, Lengauer T, Albrecht M (2008) Computing topological parameters of biological networks. Bioinformatics 24: 282–284. doi: 10.1093/bioinformatics/btm554 PMID: 18006545

65. Benjamini Y, Hochberg Y (1995) Controlling the false discovery rate: a practical and powerful approach to multiple testing. JR Stat Soc: 289–300.

66. Dimmer EC, Huntley RP, Alam-Faruque Y, Sawford T, O’Donovan C, et al. (2012) The UniProt-GO Annotation database in 2011. Nucleic Acids Res 40: D565–D570. doi: 10.1093/nar/gkr1048 PMID: 22123736

67. Groenestege WMT, Thébault S, van der Wijst J, van den Berg D, Janssen R, et al. (2007) Impaired basolateral sorting of pro-EGF causes isolated recessive renal hypomagnesemia. J Clin Invest 117: 2260–2267. doi: 10.1172/JCI31680 PMID: 17671655
68. Kinoshita A, Saito T, Tomita H, Makita Y, Yoshida K, et al. (2000) Domain-specific mutations in TGFB1 result in Camurati-Engelmann disease. Nat Genet 26: 19–20. doi: 10.1038/79128 PMID: 10973241

69. Fishman D, Faulds G, Jeffery R, Mohamed-Ali V, Yudkin JS, et al. (1998) The effect of novel polymorphisms in the interleukin-6 (IL-6) gene on IL-6 transcription and plasma IL-6 levels, and an association with systemic-onset juvenile chronic arthritis. J Clin Invest 102: 1369–1376. doi: 10.1172/JCI2629 PMID: 9769329

70. Hollstein MC, Metcalf RA, Welsh JA, Montesano R, Harris CC (1990) Frequent mutation of the p53 gene in human esophageal cancer. Proc Natl Acad Sci U S A 87: 9958–9961. doi: 10.1073/pnas.87.24.9958 PMID: 2263646

71. Frebourg T, Barbier N, Yan YX, Garber JE, Dreyfus M, et al. (1995) Germ-line p53 mutations in 15 families with Li-Fraumeni syndrome. Am J Hum Genet 56: 608–615. PMID: 7887414

72. Carpten JD, Faber AL, Horn C, Donoho GP, Briggs SL, et al. (2007) A transforming mutation in the pleckstrin homology domain of AKT1 in cancer. Nature 448: 439–444. doi: 10.1038/nature05933 PMID: 17611497

73. Lindhurst MJ, Sapp JC, Teer JK, Johnston JJ, Finn EM, et al. (2011) A mosaic activating mutation in AKT1 associated with the Proteus syndrome. N Engl J Med 365: 611–619. doi: 10.1056/NEJMoa1104017 PMID: 21793738

74. Roelfsema JH, White SJ, Aniyürek Y, Bartholdi D, Niedrist D, et al. (2005) Genetic heterogeneity in Rubinstein-Taybi syndrome: mutations in both the CBP and EP300 genes cause disease. Am J Hum Genet 76: 572–580. doi: 10.1086/429130 PMID: 15706485

75. Heino M, Scott HS, Chen Q, Peterson P, Mäebpää U, et al. (1999) Mutation analyses of North American APS-1 patients. Hum Mutat 13: 69–74. doi: 10.1002/(SICI)1098-1004(1999)13:1%3C69::AID-HUMU8%3E3.0.CO;2-6 PMID: 9888391

76. Janoueix-Lerosey I, Lequin D, Brugières L, Ribeiro A, de Pontual L, et al. (2008) Somatic and germline activating mutations of the ALK kinase receptor in neuroblastoma. Nature 455: 967–970. doi: 10.1038/ nature07398 PMID: 18923523

77. Pulst SM, Nechiporuk A, Nechiporuk T, Gispert S, Chen XN, et al. (1996) Moderate expansion of a normally biallelic trinucleotide repeat in spinocerebellar ataxia type 2. Nat Genet 14: 269–276. doi: 10.1038/ng1196-269 PMID: 8896555

78. Elden AC, Kim H-J, Hart MP, Chen-Piotkin AS, Johnson BS, et al. (2010) Ataxin-2 intermediate-length polyglutamine expansions are associated with increased risk for ALS. Nature 466: 1069–1075. doi: 10.1038/nature09320 PMID: 20740007