

Phosphoproteomic analysis of the posterior silk gland of *Bombyx mori* provides novel insight into phosphorylation regulating the silk production



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Cellular and Tissue Research, 366(1), 866–876, 2016. © Springer Science+Business Media Dordrecht. DOI: 10.1007/s00441-016-2400-0

ARTICLE INFO

Accepted: 14 May 2016
Received in revised form: 3 August 2016
Accepted: 9 August 2016
Available online: 13 August 2016

ABSTRACT

To understand phosphorylation event regulating silk synthesis in the posterior silk gland of *Bombyx mori*, phosphoproteome was profiled in a pair of near-isogenic lines, a normal cocooning strain (IC) and a naked pupated strain (IN) that the silk production is much lower than IC. In the posterior silk gland of the IC and IN, 714 and 658 phosphosites resided on 554 and 507 phosphopeptides from 431 and 383 phosphoproteins, were identified, respectively. Of all the phosphosites, the single phosphosite was the dominant phosphorylation form, comprising > 60% of all the phosphosites in two phenotypic of silk production. All these phosphosites were classified as acidophilic and proline-directed kinase classes, and three motifs were uniquely identified in the IC. The motif S-P-P might be important for regulating phosphorylation network of silk protein synthesis. The dynamically phosphorylated proteins participated in ribosome, protein transport and energy metabolism suggest that phosphorylation may play key roles in regulating silk protein synthesis and secretion. Furthermore, fibroin heavy chain, an important component of silk protein, was specifically phosphorylated in the IC strain, suggesting its role to ensure the normal formation of silk structure and silk secretion. The data gain new understanding of the regulatory processes of silk protein synthesis and offer as starting point for further research on the silk production at phosphoproteome level.

Significance statement: Despite the knowledge on regulation of silk protein synthesis in the posterior silk gland has gained at the gene or protein levels, how phosphorylation event influences the silk yield is largely unknown.

In recent years, with the increasing development of proteomic technologies, the biological processes regulating silk yield in silkworms have been investigated. It is reported that the up-regulation of ribosomal proteins, protein degradation and energy metabolism is important for fibroin synthesis and highly efficient protein synthesis system [9,10]. Several synthesis-related proteins from silkworms with adequate nutrient absorption are up-regulated, thus guaranteeing normal function of silk gland cells and enhancing protein synthesis [11]. However, knowledge regarding how phosphorylation events regulating the silk production in PSG is still lacking. Hence, the aim of this work is investigating the regulation of silk production by comparing the phosphoproteome of a pair of isolines. These results are expected to offer better comprehension and utilization of the silk gland in future applications.

2. Materials and methods

2.1. Cross-facilitation experiment

A pair of near-isogenic lines, bred by our lab, named IC with normal pupated and IN with naked pupated was utilized as the experimental materials. To breed them, two silkworm strains, the naked pupa (Nd) with normal markings and the naturally pupated strain whose larvae have striped markings (p^s), were selected and multiplied hybridized (Fig. S1). The Nd and p^s gene were dominant on pupation form and body color, respectively. First, the Nd strain and the p^s strain were hybridized to produce F_1 generation, characterized by naked pupation at the pupal stage. Then, the F_1 generation was backcrossed with the p^s strain, and their offspring, named BC₁ generation, represented two phenotypes, including pupation without cocoons and with normal cocoons. Among them, the normally pupated individuals were discarded and the naked pupated individuals were kept. This backcross method was repeated until the BC₁₇ generation to ensure that only the Nd gene was transferred into the p^s strain. Then, the naked pupated larvae of BC₁₈ generation were inbred with compatriots from the same brood. The offspring presented clear segregation of character, including naked pupated individuals and normal cocooned individuals (the phenotypic ratio was 3:1). To obtain homozygotes, the naked pupated individuals were crossed with their siblings with identical phenotype from the same brood. After the BC₂₀ generation was generated, we merely reserved the broods with single phenotype of naked pupa. Then some of the naked pupated individuals from one brood were chosen to cross with the p^s strain, and the remaining individuals were crossed with their compatriots from the same brood to produce the BC₂₁ generation. When the offspring from more than eight broods obtained by backcrossing with the p^s strain didn't represent character segregation, the Nd gene of this brood was homozygous ($P > 0.996$). Here, we named this family as IN, and the backcross parent was named IC. The key difference between IC and IN was whether the larvae spun and cocooned at the pupal stage.

2.2. Overall experimental design

The experiment setups were basically followed previous description with some modifications [3]. The PSGs from 10 silkworms of each strain were sampled on day 3 of fifth instar, a key time point for massive silk protein synthesis. Samples were dounce-homogenized in lysis buffer (8 M urea, 40 mM Tris-base, 65 mM dithiothreitol (DTT), 4% 3-[(3-cholamidopropyl)dimethylammonio]-1-propanesulfonate (CHAPS), each tablet complete mini protease inhibitor mixture (Roche, Basel, Switzerland) per 10 mL and each tablet phosphatase inhibitor cocktail (Roche, Basel, Switzerland) per 10 mL). The mixture was sonicated for further lysis, and centrifuged at 15,000 g for 45 min. DTT (1 M, 2 μ L) was added into the quantified supernatant (500 μ g), and the sample was incubated at 37 °C for 2.5 h. Then iodoacetamide (1 M, 10 μ L) was added to incubate for 40 min at room temperature in the dark. The

lysate was precipitated overnight by 100% acetone and then diluted by 100 mM ammonium bicarbonate (pH 8.5). Sequencing grade trypsin (Promega, Madison, MI) was used at a ratio of 1/50 (trypsin/protein) and enzymolysed overnight. TFA was added to terminate the digestion. The insoluble materials were removed by centrifuging and then desalted by C18 SepPak cartridges (Waters, Milford, MA). The obtained elutes were lyophilized and stored for the following experiment. During the entire process, protein concentration was measured by using a Bio-Rad Protein Assay Kit (Bio-Rad, Hercules, CA).

2.3. Phosphopeptide enrichment procedure

Phosphopeptide enrichment procedures were performed on a previous method [12]. Briefly, TiO₂ beads were equilibrated by addition of 100 μ L loading buffer (saturated glutamic acid, 65% ACN, 2% TFA), and the prepared peptide mixture was loaded onto the TiO₂ column. The mixture was incubated with end-over-end rotation for 60 min at room temperature. After incubation, the slurry was centrifuged and washed with 100 μ L washing buffer 1 (65% ACN, 0.5% TFA), and 100 μ L washing buffer 2 (65% ACN, 0.1% TFA) and centrifuged again. Then, the phosphopeptides were eluted with 100 μ L elution buffer (300 mM ammonium hydroxide, 50% ACN). The flow through was then collected, dried and prepared for further analysis.

2.4. Nano LC-MS/MS analysis

The nano LC-MS/MS was carried out as previously described [13]. All of the digested phosphopeptides were subjected to LC-LTQ-Orbitrap-MS (Thermo Fisher Scientific, San Jose, CA) in two replicates in each sample. The mixture was analyzed by nano-flow C18 reverse-phase liquid chromatography using a 15 cm fused silica monolithic column (0.075 mm inner diameter) (Column Technology, Fremont, CA). The peptide mixtures were separated using a separating gradient of 2%–35% solvent B (0.1% formic acid in ACN solution) over 120 min at a flow rate of 200 nL/min. The peptide mass spectrum was obtained in positive ion mode using data-dependent automatic switching between MS and MS/MS acquisition modes. The ten most intense ions were selected for MS/MS analysis. Fourier transformed full scan mass spectra were acquired in the range of m/z of 400–2000. The enabling dynamic exclusion and collision energy were set to 90 s (repeat count 2) and 35%, respectively.

2.5. Peptide identification and data analysis

MS/MS spectra were collected for searching by using the target-decoy database searching strategy of the Sequest algorithm [14] against the Silkworm Knowledgebase (SilkDB) (<http://www.silkdb.org/silkdb/doc/download.html>) and the National Center for Biotechnology Information (NCBI) (<http://www.ncbi.nlm.nih.gov/>). The parameters included tryptic specificity, a mass tolerance of 10 ppm, up to 2 miscleavage sites, and a unique static modification of 57.0215 Da (carboxamidomethylation) on cysteine. The results were analyzed as described [15,16], which used decoy matches as a guide. The filtering criteria, like Delta Cn, were applied to achieve a 1% false discovery rate. To localize phosphorylation sites, manual interpretation of the MS/MS ion spectra was utilized. The primary criteria of manual interpretation on MS/MS spectra was that the matched number of the b or y ions of continuous amino acids was designed as over four [12]. And most high abundant peaks in MS/MS spectra could match to b or y ions and the neutral loss ions. To count and classify the significantly altered phosphoproteins, the fold change of the identified phosphopeptide quantities in the IN strain against the IC strain was calculated. Phosphoproteins with fold changes ≥ 1.5 or ≤ 0.67 were considered to be increased or decreased.

2.6. Motif analysis

Phosphopeptide sequences were submitted to the Motif-X algorithm (<http://motif-x.med.harvard.edu/>) for discovering phosphorylation motifs [17]. The background was the uploaded *B. subtilis* proteome (a < 10 m database that was randomly generated from *B. subtilis* proteome). The sequence was centered on each phosphorylation site and extended to 13 amino acids (± 6 residues). Because the Motif-X algorithm excluded the N or C termini, these sites could not be extended. The number of motif occurrences was 20, and the significance threshold was set to $P < 10^{-6}$. The extracted motifs were used to determine the kinase classes, including acidic, proline-directed, basic, tyrosine and others, based on substrate sequence specificity [18]. The simplified categorization is useful because kinase specificity is often defined by amino acid motifs that surrounded serine (Ser), threonine (Thr) and tyrosine (Tyr) residues of the substrate proteins.

2.7. Bioinformatics

Identified differentially changed phosphoproteins were classified according to cellular component, molecular function, and biological process by retrieving the Gene Ontology (GO) annotation database, an integrated web-based GO analysis toolkit for the <http://www.geneontology.org/>

IC and IN strain were 143 and 95, respectively (Supplemental Table 2 and Fig. 1B). For both the IC and IN, the most phosphorylated protein

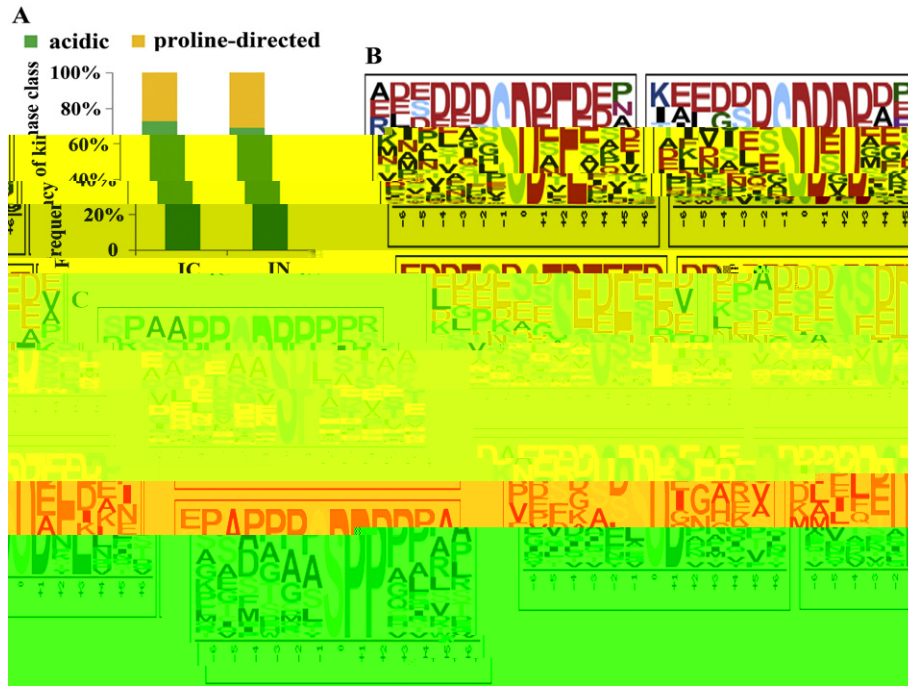


Fig. 3. Characteristics of the motifs identified in *B. b.* The frequencies of each identified kinase class are counted in the IC and IN (A). Logo-like representations of the phosphorylation motifs were identified by the Motif-X algorithm. Two kinase classes, acidic (B) and proline-directed (C), are enumerated individually. The last two on right line (B) and the lower motif (C) are the IC-specific motifs. The remaining five motifs are shared by the IC and IN strains.

synthesis processes, such as vincristine binding protein 2. Hence, the kinase recognized motif S-P-P is important to normal synthesis and secretion of silk protein.

Compared our data with those of *D. p. a. e. a. e. embr. os* [3] and *A. g. e. f. a. c. h. p. o. p. h. a. r. i. n. g. e. a. l. g. l. a. n. d.* [23], the only distinction was multiple phosphopeptides possessed higher proportion on fly, however, most proteins preferred single phosphorylation in both silkworm and honeybee. Ser phosphorylation occupied the largest proportion, and acidic and proline-directed motifs were the first two most abundant classes on all the three species. In all, for the silkworm, most proteins were phosphorylated on a single site, suggesting that a single site phosphorylation regulates the majority of the functions of a protein in most cases. That Ser phosphorylation occupied the largest proportion was also in accordance with the existing theory that protein phosphorylation mainly occurs at Ser in eukaryotic cells [24]. And phosphorylation events are mainly accomplished by acidic and proline-directed kinase classes in the PSG of *B. b.*

3.3. D. f. f. e. a. e. e. d. e. a. a.

Comparing IN (total 383 phosphoproteins) with IC (total 431 phosphoproteins), 237 phosphoproteins, 143 specific phosphoproteins and 94 common phosphoproteins, showed decreased their expression. Whereas 154 phosphoproteins (95 to specific phosphoproteins and 59 to common phosphoproteins) showed increased expression (Supplemental Table 4 and Fig. 1B). Among the down-regulated phosphoproteins, several proteins involved in genetic material synthesis and secretion were found, such as regulator factors and secretion-related proteins. Protein translation proceeds in four phases: activation, initiation, elongation, and termination. During the entire process, regulator factors play important roles. Eukaryotic translation initiation factor 5 (eIF5) is involved in forming a multifactor translation-initiation complex [25]. Once phosphorylation is reduced, the complex formation might be disrupted, thereby influencing translation initiation. As a member of the heteromeric $\beta\gamma\delta$ complex, elongation factor 1 delta (EF-1 δ) is phosphorylated and subsequently recycles inactive EF-1 α GDP to the active GTP-bound form and transfers aminoacyl-tRNAs to

the ribosome [26]. In our study, undetected phosphorylation suggests that the protein elongation process is blocked. Calnexin, normally phosphorylated on the cytosolic domain, participates in ER quality control and ensures the secretion of correctly folded proteins and combines with the ribosome and providing favorable conditions for the translation process [27,28]. Here, unphosphorylation suggests that proteins are accumulating, and de novo protein synthesis is impaired in the cells. In cells, it is only when proteins are trafficked to their final destination and assembled into structural and functional complexes that they can play their roles [29]. Therefore, correct protein trafficking and secretion is important. Consequently, the translation process was hindered and generated low protein yields, and thus blocking protein secretion in the IN strain. The down-regulated ribosomal proteins are likely to contribute to decreased protein synthesis at the translation level has been verified [10]. In the meantime, previous research on in vitro phosphorylation has shown that changes of phosphorylated ribosomal proteins alter the substrate binding sites or the conformation in the ribosome, which leads to a 50% loss of activity in protein synthesis [30, 31]. Therefore, the most phosphorylated ribosomal proteins decreased their abundance level might also be responsible for the reduced protein synthesis observed in the IN strain. Acidic ribosomal protein P1 and P2, members of the key protein synthesis organelle that make up the eukaryotic ribosomal stalk. Their phosphorylated forms are implicated in modulating ribosomal function [32]. In the IN strain, the reduced level of these two proteins may lead to decreased partial activity and further decreased ribosomal productivity. The phosphorylated ribosomal protein S2 (Rps2) participates in aminoacyl-transfer RNA binding to the ribosome [33]. Low Rps2 phosphorylation may suppress elongation process, thus preventing normal protein synthesis in the IN strain. Some phosphoproteins, such as programmed cell death protein 4-like (PDCD4), involved in the regulation of cell development and apoptosis, showed down-regulated phosphorylation in the IN strain. PDCD4 is associated with transduction of anti-apoptotic signaling by phosphorylated by Akt kinase [34,35]. In the naked pupated strain, low level of phosphorylated PDCD4 may accelerate the apoptosis of PSG. In addition, phosphoproteins related to ATP metabolism were specifically expressed in the IC, namely H⁺ transporting ATP synthase beta subunit isoforms 1

and 2. Phosphorylation of ATP synthase-β lowers ATP synthase activity and depresses energy synthesis [36]. Because of the negative phosphorylation, energy production was enhanced in the naked strain, which was consistent with previous results indicating that the strain with low silk yield demand intensive energy metabolism [10]. Notably, higher energy consumption did not cause a positive productivity change. Comparing with our results and previous proteome data [9, 10], the enhanced energy consumption might be used for ontogenetic processes rather than the synthesis of silk protein. Additionally, of the three main silk components, only phosphorylated fibroin heavy chain (Fib-H) was found to specifically exist in the IC strain. Previous research has shown that the phosphorylation of Fib-H drives its function. Phosphorylation of Fib-H causes a conformational change with functional consequences as well as possible binding between silk proteins [37]. The reason for the dephosphorylation of Fib-H might be the distinct *Fb-H* sequence of the Nd strain. Compared with the normal strain, partial bases change generates a distinguishing protein structure and thus preventing phosphate group combination [7,38]. Therefore, in the IN strain, unphosphorylated Fib-H might inhibit the formation of the silk structure and affect silk protein secretion to consequently cause the naked pupated phenotype. At the translational level, the expression of Fib-H was not altered, and it has been confirmed that its quantity does not contribute to the difference in cocoon weight [10]. Hence, phosphorylation may be an important factor that influences silk synthesis by varying the protein structure.

Among the up-regulated phosphoproteins, some molecular chaperones were identified, such as 90-kDa heat shock protein (HSP90) and 97 kDa heat shock protein. HSPs can assist in the formation of mature

proteins and participate in degrading misfolded proteins [39,40]. As mentioned above, large quantities of non-phosphorylated Fib-H with an abnormal conformation must be eliminated with the aid of abundant HSPs. The phosphorylated HSP90 can participate in the regulation of the G₂/M checkpoint, thus mediating the cell cycle [41]. In the IN strain, over-phosphorylated HSP90 might generate a late nuclear division, supporting the existing conclusion that the aberrant nuclear division of the PSG cell generates a shorter PSG [42]. Several phosphoproteins associated with transport showed increased expression level. Among these proteins, transport protein SEC31 (Sec31) is a typical coat protein II, and its phosphorylation appears to reduce the association with Sec23, thereby limiting the budding of vesicles from the ER membranes and regulating ER-to-Golgi trafficking [43]. Hence, in the IN strain, the phosphorylation of Sec31 may depress vesicle transport and impair proper protein function.

3.4. B . f a c a a . f e d f f e e . a e e e d e

For up-regulated phosphoproteins, there were 13 items were annotated, including 6 to cell component, 2 to molecular function, 5 to biological process; for down-regulated phosphoproteins, total 31 items were annotated (Fig. 4). Specifically, in the cell component, proteins with decreased phosphorylation were involved in 12 items, and the number of specific items was 8 terms. Thereinto, macromolecular complexes are an essential aspect in cellular processes, including gene expression, cell cycle regulation, cell signaling and metabolism. Thus, the number of bioactive complexes is higher in the IC to ensure normal physiological function, such as sufficient active ribosomes ensured

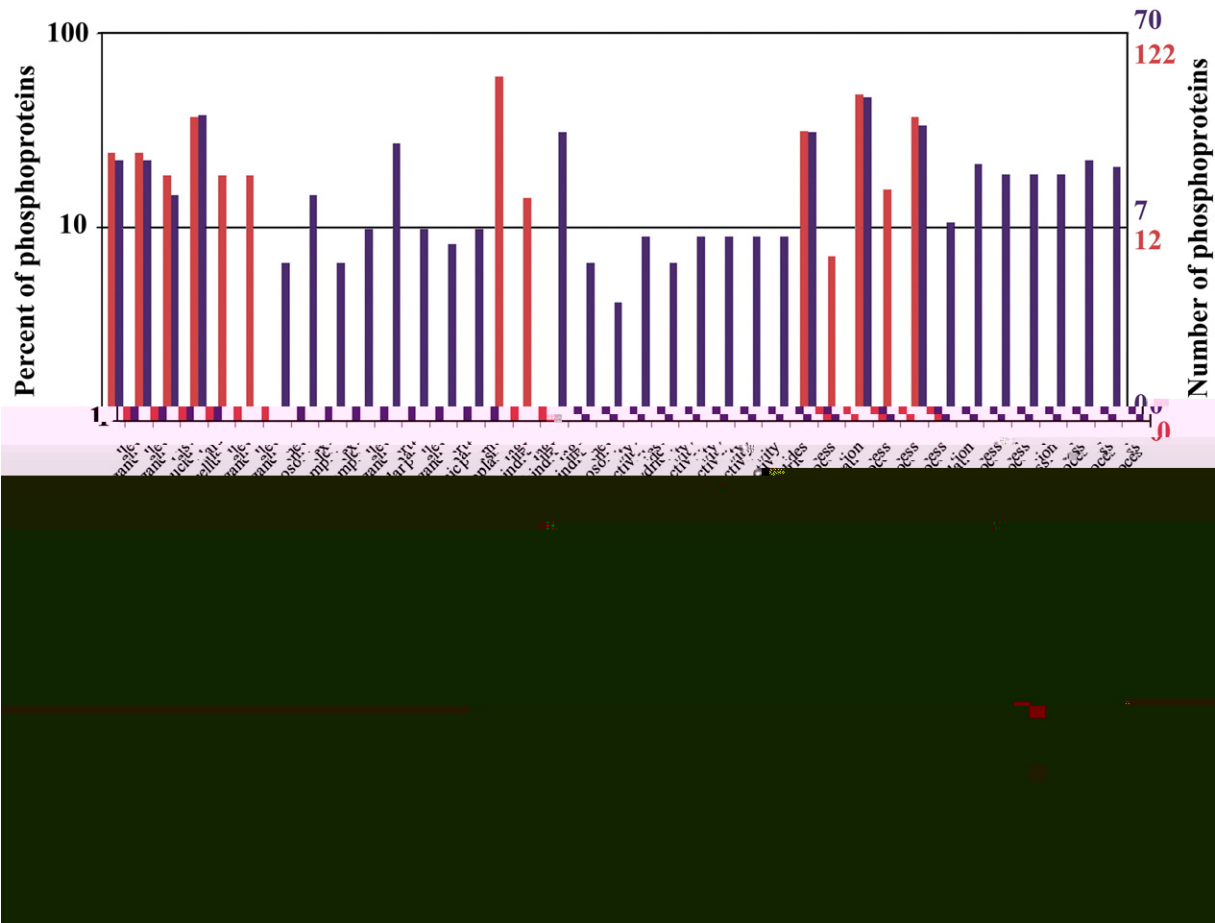


Fig. 4. GO categories of the differentially phosphorylated proteins. The number of phosphoproteins mapping to the GO terms is shown in the right panel. The left panel represents the proportion of phosphoproteins mapping to the GO terms. The light red bar represents the up-regulated phosphoproteins, and the purple bar represents the down-regulated phosphoproteins.

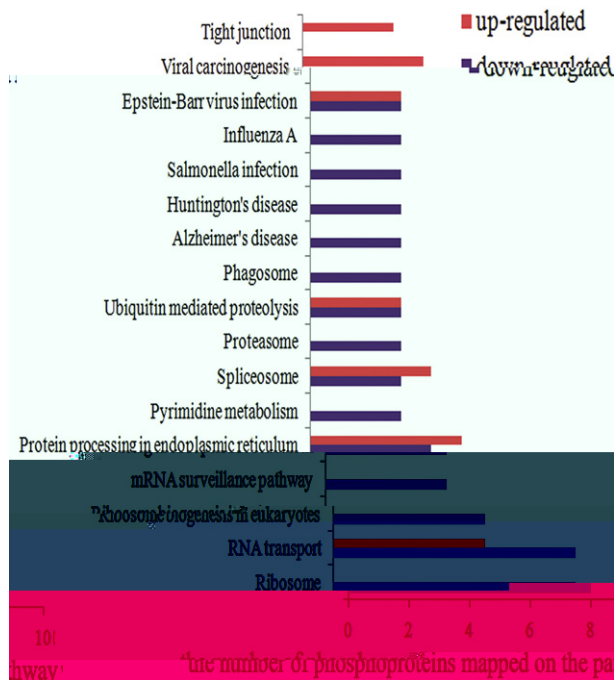


Fig. 5. KEGG pathways mapped by the differentially phosphorylated proteins. The lower panel shows the number of proteins mapping to the pathways. The light red pillar represents the increased phosphoproteins, and the purple pillar represents decreased phosphoproteins.

protein synthesis. In the biological process, a total of 10 GO terms were enriched by down-regulated phosphoproteins. The enriched translation, gene expression process and several biosynthetic processes indicate that protein synthesis and material accumulation are more active in the IC strain.

There were 7 pathways were enriched by up-regulated phosphoproteins, however 15 pathways were enriched by the down-regulated phosphoproteins (Fig. 5). Of these 15 pathways, most of them were related to protein synthesis. For instance, the RNA transport pathway enriched by most phosphoproteins in low abundance level suggest the function of this pathway may be hindered. Synthesizing large quantities of fibroin proteins requires active tRNA transport, thereby ensuring the efficiency of fibroin synthesis of the IC strain. The numerous down-regulated phosphoproteins mapped to ribosome pathway suggest that the translational machinery might be functionally reduced in the IN strain. Consequently, the synthetic quantity of fibroin and other proteins may be reduced, thus resulting in disintegration of PSG with no proteins filled. In addition, ribosome biogenesis in eukaryotes and mRNA surveillance pathway were specifically annotated by the down-regulated phosphoproteins. All these observations support a fact that the declined protein synthesis might be accounting for significantly reduced capacity in producing fibroin in the IN strain.

4. Conclusions

We reported the first phosphoproteome data of PSG of *B. mori*. Ser is preferentially phosphorylated, and acidic and proline-directed kinase classes may carry out the main phosphorylation events. The motif S-P-P is uniquely identified in the IC strain, and its kinase might be important for phosphorylation regulation of producing a large number of silk proteins. Phosphorylation may drive low silk production in the following four aspects. First, the decreased phosphorylation of several ribosomal proteins and regulator factors may associate with the reduced protein synthesis. Second, the decreased phosphoproteins related to transport and secretion imply their roles in negative regulation of de novo protein synthesis. Third, the up-regulation of energy

metabolism in IN indicates that energy turnover is likely related to the metabolism of substances other than silk proteins. Last but not least, the dephosphorylation of Fib-H may inhibit the formation of the silk structure by blocking silk protein secretion into the silk gland cavity. This reported data here gain a new understanding that the phosphorylation events implicate in regulating metabolism of silk synthesis.

Supplementar data to this article can be found online at <http://dx.doi.org/10.1016/j.jprot.2016.08.007>.

Conflict of interest

The authors declare that no conflict of interest exists in the submission of this manuscript, and manuscript is approved by all authors for publication.

Acknowledgment

This work was supported by the National Basic Research Program of China (No. 2012CB114601), China Postdoctoral Science Foundation (No. 518000-X91601) and the Projects of Zhejiang Provincial Science and Technology Plans (No. 2013C32048).

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