iTRAQ-Based Proteomics Reveals Novel Members Involved in Pathogen Challenge in Sea Cucumber Apostichopus japonicus

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Abstract
Skin ulceration syndrome (SUS) is considered to be a major constraint for the stable development of Apostichopus japonicus culture industries. In this study, we investigated protein changes in the coelomocytes of A. japonicus challenged by Vibrio splendidus using isobaric tags for relative and absolute quantification (iTRAQ) over a 96 h time course. Consequently, 228 differentially expressed proteins were identified in two iTRAQs. A comparison of the protein expression profiles among different time points detected 125 proteins primarily involved in response to endogenous stimuli at 24 h. At 48 h, the number of differentially expressed proteins decreased to 67, with their primary function being oxidation reduction. At the end of pathogen infection, proteins responsive to amino acid stimuli and some metabolic processes were classified as the predominant group. Fifteen proteins were differentially expressed at all time points, among which eight proteins related to pathologies in higher animals were shown to be down-regulated after V. splendidus infection: paxillin, fascin-2, aggrecan, olofactomedin-1, nesprin-3, a disintegrin-like and metallopeptidase with thrombospondin type 1 motif (Adams7), C-type lectin domain family 4 (Clec4g) and n-myc downstream regulated gene 1 (Ndrg1). To gain more insight into two SUS-related miRNA (miR-31 and miR-2008) targets at the protein level, all 129 down-regulated proteins were further analyzed in combination with RNA-seq. Twelve and eight proteins were identified as putative targets for miR-31 and miR-2008, respectively, in which six proteins (5 for miR-31 and 1 for miR-2008) displayed higher possibilities to be regulated at the level of translation. Overall, the present work enhances our understanding of the process of V. splendidus-challenged sea cucumber and provides a new method for screening miRNAs targets at the translation level.

MicroRNAs (miRNAs) are a type of approximately 22 nt-sized small noncoding RNAs that exist in diverse organisms [6]. It has been confirmed that in plants, miRNA-target interactions are often within the coding region and nearly perfectly complementary, which triggers miRNA cleavage. By contrast, animal miRNA/target duplexes generally are interrupted by gaps and mismatches and occur in the 3’UTR of mRNAs. Pioneering genetic studies in Caenorhabditis elegans have identified that lin-4 can inhibit the translation of lin-14 and lin-28 miRNA without affecting the cellular level of lin-14 and lin-28 [7][8]. Recently, miRNAs have emerged as key regulators of a broad spectrum of cellular activities, including immune response [9], insulin secretion [10], and viral replication [11]. In miRNA research, the identification of the targets of individual miRNAs is of utmost importance. Our understanding of the molecular mechanisms by which individual miRNAs modulate cellular functions remains incomplete until a full set of miRNA targets is identified and validated. Since miRNAs can function through partial or full complementary base pairing to their target messenger RNAs (mRNAs), resulting in
translational inhibition or mRNA degradation [12], it seems to be feasible to identify the putative targets of miRNAs by detecting the cellular level of mRNAs or proteins from the same biological sample. In our previous work, miR-31 and in vitro miRNA were demonstrated to be involved in SUs outbreak [13], and their candidate targets were also predicted by RNA-seq analysis by miRanda toolbox based on the reverse expression pattern between miRNA and mRNA [14]. However, negative expression correlations were not detected by qRT-PCR for most of their putative targets at the mRNA level in bacteria challenged samples, indicating that the targets might be regulated at the protein level without affecting mRNA abundance. Therefore, proteomics is a suitable method to reveal the full spectrum of miRNA targets and quantify the contribution of translational repression by miRNAs because it provides a rapid and comprehensive evaluation of protein profiles in complex protein samples. Over the past few decades, many proteomic platforms have been developed for the qualitative and quantitative characterization of protein mixtures and post-translational modifications, such as 2D gel-MS [15], LC-MS/MS [16]. Currently, isobaric tags for relative and absolute quantification (iTRAQ) has unique advantages over other conventional proteomics techniques because iTRAQ identifies and quantifies many proteins from specific biological environments using labeled peptides identifiable by sensitive mass spectrometers [17]. iTRAQ analysis is further strengthened by using robust bioinformatic tools and statistical analyses to support observations [18]. Using the infection model and iTRAQ approach, many researchers have made great advances in identifying proteins involved in the pathogenic process [19] [20]. Moreover, the iTRAQ approach has also been successfully employed for the identification of miRNA targets in other species [21] [22]. However, the identification of proteins by tandem mass spectrometry requires reference protein databases that are only available for model species. Because of the recent contributions to the transcriptomic characterization of the A. japonicus immune system, including transcriptome analysis of diseased [14] and LPS-challenged A. japonicus [23], a protein database for mass spectrometry-based identification of non-model organisms has been generated. Meanwhile, NMR-based metabolomics has been performed to explore the metabolic changes in the muscle tissues of pathogen-challenged and diseased A. japonicus, providing a comparative understanding of the metabolic profiles under different conditions [24]. To understand the intrinsic pathogenic mechanism, we analyzed protein expression patterns over a 96 h time course pathogen infection to identify proteins and peptides changes in response to V. splendidus challenge and reveal miRNA targets at the translational level, thus increasing our knowledge of cellular pathways important for infection and pathogenesis.

Materials and Methods

Ethics statement

The sea cucumbers (A. japonicus) here are commercially cultured animals, and all the experiments were conducted in accordance with the recommendations in the Guide for the Care and Use of Laboratory Animals of the National Institutes of Health. The study protocol was approved by the Experimental Animal Ethics Committee of Ningbo University, China.

Experimental animals and conditions

One hundred healthy adult sea cucumbers A. japonicus (165±25 g) were obtained from Bowang Aquaculture Company (Ningbo, China) and evenly assigned to four tanks randomly. The animals were then acclimatized in aerated natural seawater (salinity 25 psu, temperature 16°C) for three days prior to be treated. V. splendidus were then re-suspended in filtered seawater (FSW). For the challenge experiments, one tank used as a control, and the other three tanks were immersed with high density of V. splendidus with a final concentration of 10^7 CFU mL^-1. After being challenged for 24 h, the coelomocytes were collected from control and challenged group to confirm the existence of V. splendidus by 16S rRNA PCR. The sea cucumbers were then dissected and coelomic fluids were collected from five individuals in each tank, and approximately 50 ml of coelomic fluids were gathered at 0, 24, 48 and 96 h, respectively. Of which, 45 mL was separated as sample for iTRAQ analysis, and 5 mL for RNA quantity. The coelomic fluids were then centrifuged at 1,000 g for 5 min to harvest the coelomocytes and the supernatant were then stored at −80°C before protein and RNA extraction.

Protein extraction, quantization, digestion and iTRAQ labeling

Two biological replicates of each group were prepared for the proteomics experiments. Briefly, the total protein of each sample was grinded to powder with liquid nitrogen and dissolved in lysis solution [9 M Urea, 4% CHAPS, 1% DTT, 1% IPG buffer (GE Healthcare)]. The mix was incubated at 30°C for 1 hour and centrifuged at 15,000 g for 15 min at room temperature. The supernatant was collected and quantified by the Bradford method [25]. For each sample, 100 µg of protein was dissolved in a dissociation buffer (AB Sciex, Foster City, CA, USA). After being reduced, alkylated and trypsin-digested, the samples were labeled following the manufacturer’s instructions for the iTRAQ Reagents-plex kit (AB Sciex). Samples taken at 0, 24, 48 and 96 h were then labeled with iTRAQ reagents with molecular masses of 113, 114, 115, and 116 Da, respectively. Additional independent biological replicates were labeled with other reagents with molecular masses of 117, 118, 119, and 121 Da. After labeling, all samples were pooled and purified using a strong cation exchange chromatography (SCX) column by Agilent 1200 HPLC (Agilent) and separated by liquid chromatography (LC) using a Eksigent nanoLC-Ultra 2D system (AB SCIEX). The LC fractions were analyzed using a Triple TOF 5600 mass spectrometer (AB SCIEX). Mass spectrometer data acquisition was performed with a Triple TOF 5600 System (AB SCIEX, USA) fitted with a Nanospray III source (AB SCIEX, USA) and a pulled quartz tip as the emitter (New Objectives, USA). Data were acquired using an ion spray voltage of 2.5 kV, curtain gas of 30 PSI, nebulizer gas of 5 PSI, and an interface heater temperature of 150°C. For information dependent acquisition (IDA), survey scans were acquired in 250 ms and as many as 35 product ion scans were collected if they exceeded a threshold of 150 counts per second (counts/s) with a 2° to 5° charge-state. The total cycle time was fixed to 2.5 s. A rolling collision energy setting was applied to all precursor ions for collision-induced dissociation (CID). Dynamic exclusion was set for 75 of peak width (18 s), and the precursor was then refreshed off the exclusion list.

Protein Identification and Quantification

The iTRAQ data were processed with Protein Pilot Software v4.0 against the A. japonicus database using the Paragon algorithm [26]. Protein identification was performed with the search option
of emphasis on biological modifications. The database search parameters were the following: the instrument was TripleTOF 5600, iTRAQ quantification, cysteine modified with iodoacetamide; biological modifications were selected as ID focus, trypsin digestion. For false discovery rate (FDR) calculation, an automatic decoy database search strategy was employed to estimate FDR using the PSPEP (Proteomics System Performance Evaluation Pipeline Software, integrated in the ProteinPilot Software). The FDR was calculated as the number of false positive matches divided by the number of total matches. Then, the iTRAQ was chosen for protein quantification with unique peptides during the search, and peptides with global FDR values from fit less than 1% were considered for further analysis. Within each iTRAQ run, differentially expressed proteins were determined based on the ratios of differently labeled proteins and p-values provided by Protein Pilot; the p-values were generated by Protein Pilot using the peptides used to quantitate the respective protein. Finally, for differential expression analysis, fold change was calculated as the average ratio of 114/113 and 118/117 at 24 h, 115/113 and 119/117 at 48 h, 116/113 and 121/117 at 96 h, respectively, and proteins with a fold change of >1.5 or <0.67 and p value less than 0.05 were considered to be significantly differentially expressed.

**GO and KEGG pathway enrichment analysis**

For GO and KEGG pathway enrichment analysis, the homology search was first performed for all query protein matches with blastp against the *Mus musculus* protein database. The E-value was set to < 1e-10, and the top 10 best hits for each query sequence were taken. Among the 10 best hits, the hit with the best identity to the query was picked as homologous. The GO analysis was performed with different mapping steps to link all blast hits to the functional information stored in the Gene Ontology database using the DAVID toolkit. Public resources such as NCBI, PIR and GO are used to create links with protein IDs and corresponding gene ontology information. All annotations are associated to an evidence code that provides information regarding the quality of this functional assignment. For KEGG Pathway enrichment analysis, the pathway enrichment was performed using annotated proteins in the query dataset against the KEGG database, and Cytoscape was used for protein-protein interaction network construction.

**miRNA target prediction**

miRNA target prediction was performed only for down-regulated proteins. The corresponding miRNA sequences were acquired from miRNA-seq analysis [14]. Computational identification of miR-31 and miR-2008 targets was performed using the miRanda toolbox to search complementary regions between miRNA and the 3’UTR of mRNA with default parameters of S>90 (single-residue pair scores) and ΔG<-17 kal/mol.

**miRNA and RNA quantification analysis**

Total RNA was extracted with the RNAise plus reagent (Takara, Japan) following the manufacturer’s instructions. The SYBR green qRT-PCR assay was used for miRNA and mRNA quantification in identical samples. In brief, 500 ng RNA containing miRNAs was polyadenylated by poly(A) polymerase and converted to cDNA by reverse transcriptase using the miScript Reverse Transcription Kit (Qiagen, Germany). For miRNA analysis, the qRT-PCR was performed using the miScript SYBR Green PCR kit (Qiagen, Germany) with the manufacturer-provided miScript Universal primer and the miRNA-specific forward primers (for mRNA, the primers were gene-specific forward and reverse primers) in a Rotor-Gene Q 6000 Real-time PCR detection system (Qiagen, Germany).

The miRNA-specific primers were designed based on the miRNA sequences obtained from [13] and the gene-specific primers were designed based on the mRNA sequences obtained from [14]. All the primers used in qRT-PCR were shown in table 1. Each reaction was performed in a final volume of 20 μl containing 2 μl of the cDNA, 10 μM of each primer, 6 μl RNA-free water and 10 μl SYBR Green PCR Master mix (Qiagen). The amplification profile was: denaturation at 94°C for 15 min, followed by 40 cycles of 94°C for 15 s, 60°C for 30 s and 70°C for 30 s, in which fluorescence was acquired. At the end of the PCR cycles, melting curve analyses were performed. Each sample was run in triplicates for analysis. The expression levels of miRNAs were normalized to RNU6B[27], and the expression levels of mRNAs were normalized to 18S rRNA[28].

The $2^{-\Delta\Delta CT}$ method was used to analyze the expression levels of both miRNA and mRNA, and the obtained values represented the n-fold difference relative to the control (untreated samples). The data are presented as relative expression levels (mean±S.D., n = 3), and all experimental data were subjected to one-way Analysis of Variance (ANOVA) followed by multiple Duncan tests to determine differences in the mean values among the controls. Significant differences between the treated and corresponding control groups at each time point are indicated with one asterisk for P<0.05 and two asterisks for P<0.01. The error bars in the graphs represent standard deviations.

**Results**

**Overview of changes in host proteome induced by *V. splendidus* infection**

Compared with the control group, the *V. splendidus* were detected in the challenged groups. And the samples were then used for iTRAQ analysis. In conclusion, we describe global proteome changes in the *A. japonicus* coelomocytes during the short-term course of *V. splendidus* infection. We identified 2793 distinct proteins, of which 2049 were identified and quantified reliably at a global false discovery rate (FDR) of 1%. Compared with the control group, 228 identified proteins had significant changes in expression at different time points, among which 15 were observed to be differentially expressed at all examined time points (Fig. 1). A total of 99 proteins displayed increased expression trends (fold change >1.5, p≤0.05), and 129 proteins displayed decreased expression levels (fold change <0.67, p≤0.05) compared with the control group (Table S1). Notably, eight proteins related to pathology in higher animals exhibited down-regulated expression after *V. splendidus* infection: paxillin, fascin-2, nesprin-3, aggrecan, olofactomedin-1, a disintegrin-like and metallopeptidase with thrombospondin type 1 motif (Adamts7), c-type lectin domain family 4 (Clec4g) and N-myc downstream regulated gene 1 (Ndrg1).

**Proteomic analysis at each time point after pathogen infection**

Among these differentially expressed proteins, there were 125, 67 and 114 proteins identified in pathogen-challenged *A. japonicus* at 24, 48 and 96 h, respectively, compared with the control group. Of these, 67, 30, and 67 proteins were uniquely identified at 24, 48, 96 h post infection, respectively (Table S2). Some well-documented immune-related proteins were also included into this group. Glutathione S-transferase were uniquely expressed at 24 h and sharply increased to 3.587-fold compared to control group. However, folacin B and guanine nucleotide binding protein were
down-regulated their expression to 0.026-fold and 0.171-fold at 48 h and 96 h, respectively. More importantly, some apoptosis-related proteins, such as sarcoma oncogene (Src), vitronectin and vinculin, displayed time-dependent depressed expression (Table S2). Protein-protein interaction (PPI) analysis further indicated a diversified functional network of these novel proteins at different time points (Fig. 2).

GO and KEGG analysis of the differential expression proteins

GO enrichment demonstrated that 80, 44, and 26 protein categories were enriched in the Biological process (BP), Cellular component (CC) and Molecular function (MF) categories, respectively (Table S3). Among these, actin filament depolymerization, intracellular non-membrane-bounded organelle, and actin binding were the most abundant categories in BP, CC and MF, respectively. The differentially expressed proteins were predominately binding proteins involving in cytoskeleton organization processes at 24 h post infection. At 48 h, differentially expressed proteins were also primarily binding proteins involving in protein complex assembly for extracellular structure organization, and at 96 h, differentially expressed proteins were predominately proteins related to ribosomal assembly and regulating the actin cytoskeleton (Fig. 3). Further KEGG pathway enrichment revealed that these proteins were mainly involved in Focal adhesion and cytoskeleton regulation pathways (24 h), various sugar metabolisms and related to protein complex assembly (48 h); and ribosome, amino sugar and nucleotide sugar metabolism (96 h) (Fig. 4).

Target prediction of miR-31 and miR-2008

By integrating these results with our RNA-seq and miRanda toolbox analyses, 12 and 8 proteins were identified as putative targets for miR-31 and miR-2008 from the 129 down-regulated proteins, respectively (Table S4). Heat shock protein 90 (Hsp90b1) was the only common target for both of these miRNAs.

Table 1. Primers used in qRT-PCR.

| Gene name | Forward primer (5’-3’) | Reverse primer (5’-3’) |
|-----------|------------------------|------------------------|
| Spu-miR-31 | AGGCAAGATGGTGCCATAGCT | Qiagen miScript universal primer |
| Spu-miR-2008 | ATCCACCTCGTGTCATACATC | |
| Actr3 | CTACCATGTTCAGGGATTTCGG | GGTGAGAGATGACCTCCTTGCG |
| Clec4g | CCAACGGGAGCAAAACAAATG | TGGCAACGGCCAACATACAC |
| Cndp2 | GTCGGAGATGGTGGGATACGT | CATCAGGAGCAAGAACGAG |
| Coro6 | GCCATACGGGATTGCACTGG | GATTTGTTCCTTCCTCCTC |
| Gnb2 | GCAAAATTAACCCGAGAGCAG | CAAGTAGCAGAACGGGACATC |
| H10 | CAATCCGGGACCTGTAATG | AGCGCTGAAGATGACCTC |
| Hmmpa1 | GAGGTGATTATGCCATGGTCTGC | CACCACTGCTCTTCTTCT |
| Hmml | CAGGGAAGCTACACCAAAATG | AACAAATCCCTCCATTCAG |
| HSP90b1 | TACCTCGGAGATGGTTCACTCC | TCTTCCCCTTCGTCCTCCT |
| Hspa8 | CTGGCAACACATCATCACCA | ACTGACCTGAAACCCCTC |
| Klia0196 | GGTGATTGCGGATCAACGAC | GACCTGTCGCCGCAGTGAC |
| Myo10 | GCCCTGCTCTCTAGCAGTATG | GAGTTCGCTTCTGTCAGAAAT |
| Naca | AACAGTTCCGACCCAGGAAAC | CTCCTCACAACCCACCTCATC |
| Nnt | GAGTACGCTTGGTCAAGATG | CATCTCTGCAAGTGATAGTGG |
| Plekhd1 | AGGCGGAGAGTTCCTCGTG | CTGTGCGGAGATGGAGGATG |
| Rpl12 | CTTGGAGAGAAGTGGGAGACC | CTTGCAGGGGATGGATGCA |
| Slc25a5 | GCAGTTACCTCGGGTCTCTATG | GGTCTTCCCAGAGTGGCAG |
| Tgd5 | CTCGGAAGCTATCCACACAGG | CGACAAAGGAGGATGTTT |
| Yes1 | CGAAGATCCGAAATCACCGC | AGGACACATAGGACACAGT |

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Quantitative analysis of miRNAs and mRNAs

The expression patterns of these two miRNAs (miR-31 and miR-2008) and their 20 putative targets were examined by quantitative PCR and the results are shown in Fig. 5. miR-31 and miR-2008 were both significantly up-regulated after pathogen infection. For the targets of miR-31, five genes—WASH complex subunit strumpellin (Kiaa0196), myosin X (Myo10), TDP-glucose 4,6-dehydratase (Tgds), 60S ribosomal protein L12 (Rp112) and H1 histone family, member 0 (H1f0)—displayed unchanged expression at the mRNA level. Guanine nucleotide binding protein beta 2 (Gnb2), nascent polypeptide-associated complex alpha (Naca), nicotinamide nucleotide transhydrogenase (Nnt) and heterogeneous nuclear ribonucleoprotein A1 (Hnrnpl) maintained constant expression of the miR-2008 targets.
Figure 4. KEGG pathway enrichment analysis of differentially expressed proteins at each time point. A: 24 h; B: 48 h; C: 96 h. doi:10.1371/journal.pone.0100492.g004

Figure 5. Time-course expression patterns of miRNAs and its putative targets of *A. japonicus* upon *V. splendidus* treatment. Hnmpa1: Heterogeneous nuclear ribonucleoprotein L; Yes1: Yamaguchi sarcoma viral oncogene homolog 1; Cndp2: CNDP dipeptidase 2; Coro6: Coronin 6; Hspa8: Heat shock 70 kDa protein 8; Plekhd1: PH domain-containing family D member 1; Slc25a5: solute carrier family 25. doi:10.1371/journal.pone.0100492.g005
Unique proteins revealed the temporal and spatial host-pathogen interaction progress

The biological roles of unique proteins identified from each comparison (Fig. 1) were further examined to understand the changes occurring at each time point. From the perspective of temporal progress of host-pathogen interaction, there was a function-biased conversion from 24 h to 96 h post-infection. In the early phase after *V. splendidus* infection (24 h), the unique proteins predominantly had actin or cytoskeletal protein binding activities and were involved in actin polymerization or depolymerization and cytoskeleton organization processes. At 48 h, the function of the uniquely differentially expressed proteins included calcium ion, vitamin or metal ion binding activities involved in oxidation-reduction and some metabolic processes. Finally, in the late phase of infection (96 h), the function of its unique proteins were transferase activities, exopeptidase activities and cofactor or coenzyme binding activities involved in translation, protein folding and some catabolic processes. The different distributions of functions among these unique proteins at each time point demonstrated a temporal process initiating from environmental information management to the final stress respondent.

On the other hand, diversified immune- and apoptosis-related proteins were also identified at each time points (Table S2), indicating that modulating immune system and apoptosis pathway were the utmost event in this host-pathogen interaction process. A number of effectors like lectin, lysozyme and Toll had been addressed to be involved into *V. splendidus* challenged sea cucumber. In the present study, diverse expression profiles of immune-related genes were detected between 24 h and other two time points. The up-regulation of Gstl at 24 h might result from the host immune response upon pathogen attack. As the GST family had been known as phase II detoxification enzymes which involved in response to pathogen attack, oxidative stress, and heavy-metal toxicity [29]. And the gene expression pattern of theta-class of GST in *Vibrio tabetus* and LPS challenged molluscan *Ruditapes philippinarum* also showed the similar tendency marked as the sharply raised expression level at 6 h and 24 h post infection [30]. The Ficolins were serum complement lectins which were able to recognize pathogen-associated molecular patterns (PAMPs) and trigger the activation of immune system by initiating activation of complement pathway or stimulating secretion of the inflammatory cytokines [31]. Another down-regulated immune related protein, Gnb2 belong to the G protein family, which function as G protein signaling to modulate cell motility and killing of invading pathogen. The significantly down-regulation of these two factors might represent negative modulation of the innate immune response as the insufficient of FcRb and Gnb2 might result in higher susceptibility to infection in individuals.

However, apoptosis related molecules like Src, vitronectin and vinculin were scarcely investigated in sea cucumber. Src is the canonical member of the non-receptor family of tyrosine kinases, which could inhibit apoptosis through the Erk1/2-dependent degradation of the death accelerator Bik [32]. Previous study also demonstrated that in the human primary macrophages, the Src tyrosine kinase activity were essential for the inflammammasome activation during influenza A virus infection [33]. Vitronectin was considered to modulate neutrophil adhesion, chemotaxis, to contribute to neutrophil-associated proinflammatory processes [34], and to inhibit neutrophil apoptosis through activation of integrin-associated signaling pathways [35]. Vinculin was known to regulate extracellular signal-regulated kinase (ERK) by modulating the accessibility of paxillin for FAK interaction, thus controlling the survival and motility of cells which were critical to metastasis [36]. But recent study also shown that vinculin was emerging as a regulator of apoptosis and Shigella entry into host cells [37][38]. What is more important is that Pxn served as a “hub” protein for Src, vitronectin and vinculin, indicating a potential apoptosis related signal transportation role of paxillin under bacterial infectious conditions.

Metabolism and fuel usage during pathogen infection period

Recently, an NMR-based metabolomics study was conducted to explore the metabolic changes in muscle tissues of pathogen-challenged *A. japonicus*. This study observed that the pathogen did not induce obvious biological effects in *A. japonicus* samples 24 h after infection. Enhanced energy storage and immune responses were observed in *V. splendidus*-challenged *A. japonicus* samples at 48 h, marked by increased glucose and branched chain amino acids, respectively. Finally, infection of *V. splendidus* induced significant increases in the energy demand of *A. japonicus* samples at both 72 and 96 h, confirmed by decreased glucose and glycogen and increased ATP levels [24]. Because of technology restrictions, it is difficult to examine metabolic changes in pathogen-challenged coelomocytes. However, by reference to the metabolomics in the muscle tissues of *V. splendidus*-challenged samples and combined with our proteome analysis, we can infer metabolic changes in coelomocytes.

At 24 h post-infection, down-regulation of glucose phosphate isomerase-1 and transaldolase-1 in the pentose phosphate pathway will result in the accumulation of α-D-Glucose-6P and β-D-Fructose-6P and deficiency of D-Glycer-aldehyde-3P, which may increase the concentration of pyruvate and inhibit the glycolysis pathway. Excessive pyruvate will promote the biosynthesis of leucine, isoleucine and valine, resulting in increased levels of glucose and branched amino acids at 48 h post-infection, the up-regulation of triosephosphate isomerase-1 and transaldolase-1 in the fructose and mannose metabolism pathway will increase the level of intracellular Glycer-aldehyde-3P, which will accelerate the glycolysis pathway. Moreover, the up-regulation of ornithine aminotransferase and argininosuccinate synthetase-1 in arginine and proline metabolism will facilitate the urea cycle, which will further promote alanine metabolism. In addition, the typical decrease of
Coupling transcriptional and post-transcriptional miRNA and L8, that was observed at 96 h post infection may have resulted in decreased cellular amino acid levels and increased energy demand.

Supporting Information

Table S1 Summary of the total differential expressed proteins by iTRAQ.

Table S2 Common and unique expressed proteins at each times.

Table S3 GO enrichment of the total differential expressed proteins.

Table S4 Putative targets for miR-31 and miR-2008 from the down-regulated expressed proteins.

Author Contributions

Conceived and designed the experiments: CL CJ. Performed the experiments: PJZ PZ. Analyzed the data: PJZ CL. Contributed reagents/materials/analysis tools: CL DP YB. Contributed to the writing of the manuscript: PJZ CL.
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