First comprehensive analysis of lysine acetylation in *Alvinocaris longirostris* from the deep-sea hydrothermal vents

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**Abstract**

**Background:** Deep-sea hydrothermal vents are unique chemoautotrophic ecosystems with harsh conditions. *Alvinocaris longirostris* is one of the dominant crustacean species inhabiting in these extreme environments. It is significant to clarify mechanisms in their adaptation to the vents. Lysine acetylation has been known to play critical roles in the regulation of many cellular processes. However, its function in *A. longirostris* and even marine invertebrates remains elusive. Our study is the first, to our knowledge, to comprehensively investigate lysine acetylome in *A. longirostris*.

**Results:** In total, 501 unique acetylation sites from 206 proteins were identified by combination of affinity enrichment and high-sensitive-massspectrometer. It was revealed that Arg, His and Lys occurred most frequently at the +1 position downstream of the acetylation sites, which were all alkaline amino acids and positively charged. Functional analysis revealed that the protein acetylation was involved in diverse cellular processes, such as biosynthesis of amino acids, citrate cycle, fatty acid degradation and oxidative phosphorylation. Acetylated proteins were found enriched in mitochondrion and peroxisome, and many stress response related proteins were also discovered to be acetylated, like arginine kinases, heat shock protein 70, and hemocyanins. In the two hemocyanins, nine acetylation sites were identified, among which one acetylation site was unique in *A. longirostris* when compared with other shallow water shrimps. Further studies are warranted to verify its function.

**Conclusion:** The lysine acetylome of *A. longirostris* is investigated for the first time and brings new insights into the regulation function of the lysine acetylation. The results supply abundant resources for exploring the functions of acetylation in *A. longirostris* and other shrimps.

**Keywords:** Alvinocarididae, Hemocyanins, Hydrothermal vent, Lysine acetylation, Post-translational modification

**Background**

Deep-sea hydrothermal vent is one of the chemosynthetically-driven ecosystems and characterized with high pressure, continuous darkness, and enriched hydrogen sulfide (H₂S), methane (CH₄), heavy metals in the ejected fluid [1]. Surprisingly, particular animals are thriving under these harsh conditions, which must develop specific mechanisms to tolerate the extremes and yet benefit from the chemoautotrophic production. As the development of sequencing technology and bioinformatics, large genomic resources have been discovered in hydrothermal vent faunas, such as mussels [2–4], worms [5], crabs [6] and shrimps [7], which have greatly expanded our understanding for the molecular mechanisms of their adaptation to these extreme environments. However, proteomics researches in hydrothermal vent animals have received relatively little attention.

Post-translational modifications (PTMs) modulate the activity of proteins in most eukaryotes [8, 9]. Lysine acetylation is a dynamic and reversible PTM involved in a broad array of biological functions, e.g. gene regulation, DNA-protein interactions, subcellular localization, and protein stability [10]. Changes in cellular lysine acetylation status can also alter metabolic enzyme activity and provide a metabolic adaptive mechanism for the cells [11–14]. Besides, more studies indicate that lysine acetylation plays pivotal roles in mitochondrial function...
and stress response [15–17]. Despite intensive research on lysine acetylation over the past decade, most studies for faunas have been performed in mammals. Recently, with the combination of antibody-based affinity enrichment and high sensitive mass spectrum (MS), high-throughput lysine acetylome analyses are available and becoming popular [18–21]. However, there is no comprehensive study for any marine animal species. Shrimps of the family Alvinocarididae are specifically inhabiting in deep-sea hydrothermal vent and cold seep chemosynthetic areas [22]. Among them, A. longirostris is the only species found co-distributed in both environments thus far [23, 24], and is mostly found on the mussel beds surrounding the base of active hydrothermal chimneys, suggesting they can survive in a fairly reducing environment [25]. These characters make them an excellent model for studying extreme environmental adaptation mechanisms. In this study, we globally identified lysine acetylation sites in proteins of A. longirostris from the Iheya North hydrothermal vent in Okinawa Trough by taking advantage of affinity enrichment and LC (liquid chromatography)-MS/MS, and characterized the protein functions systematically. Further study on the role of lysine acetylation in the shrimp can enhance our understanding of how macroorganisms have adapted to deep-sea extreme environments.

**Methods**

**Sample collection and protein extraction**

Samples of A. longirostris were collected in August, 2016 during the cruise by the scientific research vessel (RV) KEXUE (Institute of Oceanology, Chinese academy of Sciences, China). The sampling sites were near the Iheya North hydrothermal vent in Okinawa Trough (27°16.2' N, 127°04.6' E, depth 1329 m). Materials were taken by the remotely operated vehicle (ROV) Quasar MkII of SMD in the United Kingdom, which was deployed using the RV KEXUE. The in situ environmental conditions were detected by using the sensors of the ROV. The temperature was ~ 3.82 °C, the conductivity was ~ 5.85 s/m, the dissolved carbon dioxide was ~ 304 ppm and the methane concentration was ~ 3.92 mmol/l, while many other data was missing. The samples were immediately frozen in liquid nitrogen and stored at −80 °C. After the cruise, the abdomens of shrimps were dissected for protein extraction in the lab. The overall technological process was outlined in Fig. 1a.

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**Fig. 1** Overview of the global identification of lysine acetylation sites in *Alvinocaris longirostris*. **a** General workflow for the acetylome analysis. **b** Mass error distribution of the identified peptides. **c** Length distribution of the peptides.
Protein of two shrimps was extracted separately as two biological replicates. The tissue was grinded into powder with liquid nitrogen. Four volumes of lysis buffer (8 M urea, 1% Triton-100, 10 mM dithiothreitol, and 1% Protease Inhibitor Cocktail) with inhibitors (3 μM TSA and 50 mM NAM) was added to the powder, followed by sonication three times on ice. The residues were removed by centrifugation at 20,000 g, 4 °C for 10 min. The protein was then precipitated with cold 20% TCA for 2 h at –20 °C. After centrifugation at 12,000 g, 4 °C for 10 min, the supernatant was discarded. The remaining precipitate was washed with cold acetone for three times, and the protein was finally re-dissolved in 8 M urea. The protein concentration was measured with BCA kit accordingly.

Trypsin digestion and HPLC fractionation
Before digestion, the protein solution containing approximate 10 mg protein was reduced with 5 mM dithiothreitol for 30 min at 56 °C and then alkylated with 11 mM iodoacetamide for 15 min at room temperature in darkness. The urea concentration of the protein solution was then diluted to less than 1 M by adding 100 mM NH₄HCO₃. Trypsin (V5111, Promega, Madison, USA) was first added at 1:50 trypsin-to-protein mass ratio and the sample was digested overnight. For the second time, 1:100 trypsin-to-protein mass ratio was selected and the digestion lasted for 4 h. The trypptic peptides were then fractionated into fractions by high pH reverse-phase HPLC using Thermo Betasil C18 column (5 μm particles, 10 mm internal diameter, 250 mm length). Peptides were first separated into 60 fractions with a gradient of 8 to 32% acetonitrile (pH 9.0) for 60 min. After that, the peptides were combined into four fractions and dried by vacuum centrifuging.

Affinity enrichment for acetylated peptides
The peptides were dissolved in NETN buffer (100 mM NaCl, 1 mM EDTA, 50 mM Tris-HCl, 0.5% NP-40, pH 8.0) and incubated with pre-washed antibody beads (Lot number PTM104, PTM Bio, Hangzhou, China) at 4 °C overnight. During the incubation, it was shaken gently. The beads were washed four times with NETN buffer and twice with ddH₂O. The bound peptides were eluted with 0.1% trifluoroacetic acid. Then they were combined and vacuum-dried. The enriched peptides were desalted with C18 ZipTips (Millipore) accordingly and submitted to LC-MS/MS analysis.

LC-MS/MS analysis
The peptides were dissolved in 0.1% formic acid (solvent A) and separated on an EASY-nLC 1000 UPLC system. The gradient of solvent B (0.1% formic acid in 98% acetonitrile) included an increase from 6 to 23% over 26 min, 23 to 35% in 8 min, climbing to 80% in 3 min, holding at 80% for the last 3 min, and finally at a constant flow rate of 400 nl/min.

The peptides were subjected to nanospray ionization (NSI) source followed by tandem mass spectrometry (MS/MS) in Orbitrap Fusion™ Tribrid™ (Thermo). The electrospray voltage was set as 2.0 kv. For MS scans, the scan range was 350 to 1550 m/z. Intact peptides were detected with a resolution of 60,000 in the Orbitrap. Peptides were selected for MS/MS with normalized collisional energy (NCE) being set as 35 and ion fragments were detected with a resolution of 15,000. A data-dependent procedure that alternated between one MS scan and the following 20 MS/MS scans was processed for the top 20 precursor ions above a threshold intensity of 5E3 in the MS scan with 15.0 s dynamic exclusion. Automatic gain control (AGC) was performed to prevent overfilling of the Orbitrap and 5E4 ions were accumulated for generation of MS/MS spectra.

Database search and motif analysis
The identified MS/MS data were analyzed by Maxquant search engine (v.1.5.2.8) [26, 27]. Tandem mass spectra were searched against a protein database of A. longirostris concatenated with reverse decoy database. The database was constructed with the deduced peptide sequences (45,313) from the transcriptome of A. longirostris created by our lab (accession number SRX3177689, unpublished). Trypsin/P was specified as cleavage enzyme allowing up to four missing cleavages. The mass tolerance was set as 20 ppm in first search and 5 ppm in main search for precursor ions, and 0.02 Da for fragment ions. Carbamidomethyl on Cys was specified as fixed modification, while oxidation on Met, acetylation on Lys and acetylation on protein N-terminal were specified as variable modifications. False discovery rate (FDR) was adjusted to <1% and minimum score for modified peptides was set >40. The models of protein sequences constituted with amino acids in specific positions of modify-21-mers (10 amino acids upstream and downstream of the site) were analyzed by using motif-x (http://motif-x.med.harvard.edu/motif-x.html). In order to further investigate the lysine acetylation profile of the microbial community of A. longirostris, tandem mass spectra were similarly searched against the protein database of the family Methylcoccaceae downloaded from Uniprot, considering the high concentration of methane (~3.92 mmol/l) in the sampling sites.

Protein annotation and functional enrichment
Different databases were selected for protein functional annotation. Gene Ontology (GO) annotation proteome was derived from the UniProt-GOA database (http://www.ebi.ac.uk/GOA/). If the proteins were not annotated by UniProt-GOA database, the InterProScan
the two proteins were predicted by NetSurfP [30]. The hemocyanin subunit 1 (AlHc1). Secondary structures of which were described as hemocyanin (AlHc) and According to annotation, deuced amino acid sequences of Structure and evolution analyses of hemocyanin was also used for protein domain functional description. Then proteins were then classified by GO annotation (http://www.geneontology.org/). For each GO and protein domain category, a two-tailed Fisher’s exact test was employed to test the enrichment of the acetylated proteins against all predicted proteins from the transcriptome of A. longirostris. The item with a corrected p-value < 0.05 was considered significant. Further, KEGG (Kyoto Encyclopedia of Genes and Genomes; http://www.genome.jp/kegg/) online service tools KAAS was utilized to predict the pathways in which the acetylated proteins were involved. Then the annotation results were mapped on the KEGG pathway database using KEGG mapper. These pathways were also enriched as performed in GO enrichment and further classified into hierarchical categories according to the KEGG website. Subcellular localizations of the protein were predicted by Wolf PSORT (http://wolfpsort.seq.cbrc.jp/) [28].

Protein-protein interaction (PPI) network of acetylated proteins
Due to absence of protein information of A. longirostris in the STRING database, all identified acetylated protein sequences in A. longirostris were aligned with protein data of its relatively closely related species, Daphnia pulex. The matched homologous proteins in D. pulex were then searched against the STRING database version 10.5 for protein-protein interaction analysis. Only interactions between the proteins belonging to the searched data set were selected, and thereby external candidates were excluded. Interactions that had a confidence score ≥ 0.4 (high confidence) were extracted. Interaction network form STRING was visualized in Cytoscape [29]. A graph theoretical clustering algorithm, molecular complex detection (MCODE) in Cytoscape was applied to analyze densely connected regions.

Structure and evolution analyses of hemocyanin
According to annotation, deuced amino acid sequences of two hemocyanins with acetylation sites were identified, which were described as hemocyanin (AlHc) and hemocyanin subunit 1 (AlHc1). Secondary structures of the two proteins were predicted by NetSurfP [30]. The presumed tertiary structure was established using SWISS-MODEL prediction algorithm (http://swissmodel.expasy.org/) and displayed by DeepView/Swiss-Pdb Viewer version 4.0.2. The two hemocyanin sequences were aligned with those downloaded from GenBank of other caridean shrimps using MEGA version 6.0 [31]. Phylogenetic trees were constructed using Maximum Likelihood (ML) method with RAxML Black-Box webserver (http://phylobench.vital-it.ch/raxml-bb/) [32]. For the ML analysis, we selected the Gamma model of rate heterogeneity and assessed support for the tree nodes using bootstrap procedure with 1000 heuristic replicates.

Results and discussion
The first lysine acetylome map of the deep-sea hydrothermal vent shrimp
Deep-sea hydrothermal vents are extreme chemosynthetic ecosystems where abundant endemic animals inhabit, including many crustaceans, like shrimps. However, lack of genomic information impedes the clarification of molecular mechanisms in the adaptation to these harsh environments. As one way of the PTMs, protein acetylation has been revealed to play critical roles in various physiological processes related to adaptive reaction [33–35]. Recently, we have sequenced the transcriptome of A. longirostris and identified the protein coding domains, which supply baseline data for PTM studies. Therefore, we characterized the lysine acetylated proteins in A. longirostris systematically by utilizing highly sensitive immuno-affinity purification in combination with high-resolution LC-MS/MS in order to facilitate further research for their biological functions in the adaptation of the hydrothermal shrimps.

In the identification, the mass errors were lower than 5 ppm and the majorities were near zero (Fig. 1b), showing the mass accuracy of the MS data. The length of the peptides ranged from 7 to 34 amino acids and most were between 7 and 20 amino acids (Fig. 1c), which confirmed with the standard of sample preparation. Altogether 501 unique acetylation sites from 206 proteins were identified (Additional file 1: Table S1) with 217 sites and 102 proteins co-detected in the two experimental replicates (Fig. 2). All 501 acetylation sites were included in the following analyses. The numbers of modification sites per protein were from 1 to 28, in which more than 60% of proteins contained only one acetylation site (Fig. 3a), which was in common with many other studies [33, 36]. However, probably due to the limited genome information in A. longirostris, the numbers of detected acetylation sites and proteins are much less than those in Drosophila with well assembled whole genome sequences [33]. Additionally, only the abdomen has been selected for analysis in this study, and more tissues and organs should be included considering the tissue specific acetylation [11], which may contribute to the integrity of the acetylome. The acetylation level of proteins varying significantly between different species maybe another reasonable interpretation as suggested in previous studies [36, 37]. However, as the first lysine acetylome map of marine invertebrate and hydrothermal vent fauna, it is expected to supply valuable resources for PTM study in the future.
Motif characters of the acetylated peptides in *A. longirostris*

To further explore the acetylation pattern in *A. longirostris*, flanking amino acid residues from position −10 to +10 around the acetylated lysine were analyzed. Strong bias in amino acids of specific acetylation site motifs was observed. Arg (KacR), His (KacH) and Lys (KacK) occurred most frequently at the +1 position (Fig. 3b, c), which were all alkaline amino acids and positively charged. In the previous study for plants [20, 21] and bacteria [37, 38], KacH has been also identified in the consensus motif. However, these biases are different from those of *Drosophila* and human, in which no significant bias of specific motif is detected and the Tyr, Phe and Pro occur relatively more frequent at the +1 position [33, 39]. Therefore, the motif analysis results suggest that acetylation preferentially occurs at alkaline
and positively amino acid nearby regions in *A. longirostris*, which may be functionally important for acetylation in the shrimps.

**Functional annotation and enrichment of acetylated proteins in A. longirostris**

GO analysis was performed to gain insight into the potential functional implications of acetylation in the shrimp. In total, 170 (82.52%) of the acetylated proteins were annotated and classified into three categories: 'Biological process', 'Cellular component' and 'Molecular function' (Fig. 4; Additional file 1: Table S1). In the 'Biological process' category, the most prevalent GO terms were metabolic process (35%), cellular process (30%) and single-organism process (22%), while localization, biological regulation, cellular component organization or biogenesis, and the other terms accounted for 6, 3, 3 and 1%, respectively (Fig. 4a). Cell (37%), organelle (26%) and macromolecular complex (25%) were the main 'Cellular component' (Fig. 4b). In the 'Molecular function' classification, we found that most acetylated proteins were related to catalytic activity (43%) and binding (43%) (Fig. 4c). According to the functional annotation, many metabolism related enzymes were identified to be acetylated (Additional file 1: Table S1), such as acyl-coenzyme A oxidase 3 (c94891_g1_orf1), citrate synthase (c77391_g1_orf1), glycogen phosphorylase (c75875_g1_orf1), fructose-bisphosphate aldolase (c73820_g1_orf1), malate dehydrogenase (c72173_g1_orf1, c90371_g1_orf1), phosphoglycerate kinase (c89458_g1_orf1), which possibly function in the cell metabolic regulation as previously revealed in various species [38, 40, 41]. In these enzymes, some of the homologous proteins in *Drosophila* and human have also been characterized to be lysine acetylated, indicating their evolutionary and functional conservation [33, 39].

To further determine the proteins more prone to be acetylated, we enriched the acetylated proteins against all predicted proteins from the transcriptome. In the GO enrichment, it was noticed that several protein complexes were preferred to be acetylated, including the proton-transporting ATP synthase complexes (≥5.6%), macromolecular complex (7.72%), ribosome (6.21%), nucleosome (5.38), myosin complex (5.35%), and DNA packaging complex (5.19%) (Fig. 5a). Meanwhile, diverse molecular functions were enriched, such as catalytic activity (14.02%), oxidoreductase activity (11.64%), cofactor binding (9.04%) and fructose-bisphosphate aldolase activity (6.61%), which were involved in various metabolic processes (≥11.7%) and oxidation-reduction process (13.68%) (Fig. 5a). In the KEGG enrichment, 23 pathways were detected to be significantly enriched, e.g. carbon metabolism, biosynthesis of amino acids, citrate cycle, fatty acid degradation, oxidative phosphorylation and peroxisome (Fig. 5b). In the protein domains, thioloase, hemocyanin and TCP-1 like chaperonin intermediate domains were revealed to be significantly enriched, which might be involved in the adaptation to the high H₂S and hypoxia, as well as toxin transport in the hydrothermal vents of *A. longirostris*. Therefore, lysine

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Fig. 4 Functional distribution and subcellular localization of the lysine acetylated proteins in *Alvinocaris longirostris*. **a** Classification based on biological process. **b** Classification based on cellular component. **c** Classification based on molecular function. **d** Subcellular location prediction.
Acetylation trends to target large macromolecular complexes as reported in human [39] and are associated with various processes, like substance transport and metabolism, oxidation-reduction, protein synthesis, chromatin remodeling, which may function in the cell metabolism regulation and stress response to the hydrothermal vents of the shrimp. However, further experimental verification is indispensable. Since optimal conditions to assay unique sites deacetylation are formidable, pivotal acetylated proteins involved in certain biological processes may be purified and selected for functional study by in vitro site-directed mutagenesis of lysine to arginine or glutamine initially [42–44].

Abundant lysine acetylation in mitochondrion and peroxisome of *A. longirostris*
Analysis of acetylated proteins subcellular localization showed that most of them were predicted to localize in cytoplasm (34%), mitochondria (26%) and nucleus (21%), while the others took a proportion of 19%. Previously, large-scale proteomics studies have demonstrated that lysine acetylation is widespread in mitochondria [10, 39, 45, 46], which is the key regulator of cellular energy production and pivotal in the regulation of metabolism and maintaining of cellular homeostasis. In this study, 14 acetylated proteins were identified to be involved in the oxidative phosphorylation which occurred in mitochondria (Fig. 6a), such as ATPase, Cytochrome c oxidase and reductase, and NADH dehydrogenase. Oxidative phosphorylation has been reported to play central roles in various stress responses of many crustacean species by altering gene expression or protein activity [47–50]. Peroxisome is another ubiquitous subcellular organelle participating in metabolic and pathological/stress response processes [51, 52]. In the KEGG pathway enrichment, six acetylated proteins were discovered to be involved in peroxisome, which were acyl-CoA oxidase, carnitine O-acetyltransferase, carnitine O-octanoyltransferase, isocitrate dehydrogenase, superoxide dismutase, Fe-Mn family and peroxisomal 3,2-trans-enoyl-CoA isomerase (Fig. 6b). Limited PTMs have been identified in peroxisome thus far. Acetylation of a peroxisomal enzyme of peroxisomes has been explored in rat liver and suggests that acetylation may play a functional role in the organelle proliferative process [51]. We speculate that acetylation or deacetylation of these mitochondrial and peroxisome proteins may probably regulate the protein function in order to adapt to the hydrothermal vent environments in *A. longirostris*.
Interaction network of acetylated proteins in *A. longirostris*

Protein interactions play an important part in various biological pathways and orchestrate virtually all cellular processes [53]. We further built the PPI network for all acetylated proteins in order to deeply investigate the processes regulated by acetylation and how these modified proteins mutually intertwined. A total of 155 acetylated proteins were identified as nodes and connected with each other (Fig. 7, Additional file 2: Table S2), which represented a global PPI network of *A. longirostris*. The degree of node is the key parameter to evaluate the correlation of protein in network. Among them, six proteins displayed the highest degree (≥40, Additional file 2: Table S2), which were actin 6 (Act6), actin alpha cardiac muscle 1 (Actc1), DNA topoisomerase 2-beta (Top2b), delta-1-pyrroline-5-carboxylate dehydrogenase mitochondrial (Aldh4a1), 10-formyltetrahydrofolate dehydrogenase (Aldh1l1) and trifunctional enzyme subunit alpha mitochondrial (Hadha). By extraction of the highly enriched interaction clusters, we found that the top enriched Cluster 1 consisted proteins involved in binding function, while Cluster 2 and Cluster 3 were metabolism and ribosome associated proteins, respectively (Fig. 7, Additional file 2: Table S2). It was noted that Act6, Actc1 and Top2b were located in Cluster 1, whereas Aldh4a1, Aldh1l1 and Hadha were all included in metabolic pathways, Cluster 2. Massive lysine acetylation in different metabolic processes and ribosome also have been revealed from eukaryotes and prokaryotes [20, 54]. The potential acetylated protein interactions in this study provide valuable candidates in the following studies for their coordination and regulation in the hydrothermal vent shrimps.

Acetylation associated with stress adaptation and hemocyanins in *A. longirostris*

Considering the challenging environment in hydrothermal vents where *A. longirostris* inhabit, such as low oxygen and high levels of toxic substances, we particularly analyzed the acetylated proteins which might be directly response to the stress. According to the annotation, two arginine kinases, Aks, (c18620_g2_orf1, c85882_g1_orf1), lethal (2) essential for life, L(2)efl (c40235_g1_orf1), heat shock protein 70, Hsp70 (c57316_g1_orf1) and hemocyanins (c93924_g1_orf1, c95510_g4_orf1) were identified (Additional file 1: Table S1). As phosphagen kinase, Aks mainly distribute in invertebrates, and play critical roles in the energy mechanism, as well as various stress responses [55]. In *Macrobrachium rosenbergii*, Ak has been revealed involved in the immune response of this shrimp [56]. In *A. longirostris*, Ak1 and Ak2 displayed hyper-acetylation with 11 and 5 acetylation sites, respectively, which may contribute to the physiological regulation in the adaptation to the extreme environments. Moreover, L(2)efl and Hsp70 with acetylation sites both belong to HSP family, which were well-known for their function in cell response to stress, such as thermal stimulus and osmotic pressure [48, 57, 58].

Oxygen (O₂) transport is an essential biological process in animals. In the deep-sea hydrothermal vent environments, there are lack of O₂ and therefore specific
regulation mechanism in organisms is compulsory in order to adapt to the hypoxia. Hemocyanin (Hc) is one of the key respiratory proteins in the blood of arthropods [59], with a principal structure of a hexamer of six similar subunits [60]. Recent studies show that Hc potentially also involved in other stress response of decapods [6, 61–63]. In another alvinocaridid shrimp, Rimicaris exoculata, Hc encoding gene shows a strong up-regulation when the shrimp is exposed to heat stress [64]. Evidence of acetylation in Hcs has been suggested previously [65, 66], but the information is quite limited. In this study, two Hcs were revealed to be acetylated, named AlHc and AlHc1 with two and seven acetylation sites, respectively. The deduced amino acids (aa) of AlHc and AlHc1 contained 681 and 674 aa (Fig. 8a). Two Hc sequences in A. longirostris were then aligned with 10 Hc sequences of other Caridea species. They all contained six conserved copper-binding histidines and a signal peptide (Fig. 8a). Hc sequences of A. longirostris showed the highest similarity with those of M. nipponense, which were also supported by the phylogenetic tree (Fig. 8b).

Previously, three distinct types of hemocyanin subunits have been defined and referred to α, β and γ, in which α and γ are more related with each other than with β [67, 68]. It was obvious that AlHc and AlHc1 were classified into α and γ types based on the tree (Fig. 8b). By second structure analysis, we found site 199 with acetylation of AlHc was located in coli, while site 673 with acetylation was in β-strand (Fig. 8c). In AlHc1, two acetylation sites (sites 47, 555) were contained in α-helix, one in β-strand (site 185), and four (sites 87, 234, 628, 634) in coli (Fig. 8c). It indicates that acetylation in Hcs of A. longirostris prefers to occur at regions with ordered secondary structure, which may be involved in the stability of the Hc complex. On the other hand, more acetylation sites were discovered in AlHc1 than those in AlHc of A. longirostris. Moreover, the acetylation site 628 in AlHc1 was unique in A. longirostris when compared with other shallow water shrimps (Fig. 8a), K (acetylated) in A.

![Fig. 7 Protein-protein interaction network of all acetylated proteins in Alvinocaris longirostris](image-url)
whereas H, N, or R in other species. We infer that this lysine acetylation probably participate in the specific adaptation to the hydrothermal vent environments in *A. longirostris*, but further verification is essential.

**Acetylation of microbes associated with abdomen of A. longirostris**

At the base of the food web, microbes are thriving in the extreme hydrothermal vent and cold seep environments, which are dominated by chemoautotrophic bacteria and archaea fuelled by simple reduced molecules [69]. The symbiont between invertebrates and microbes is also critical for the survival of macroorganisms. Therefore, it is interesting to investigate PTMs in these microbes and their potential roles in the adaptation to the extreme conditions and in the symbiont with the host. However, we have no protein database of microbes from the *A. longirostris* in Iheya North hydrothermal vent area. Methylococcaceae has been discovered rich in a deep-sea mussel *Bathymodiolus platifrons*, another species co-distributed in hydrothermal vent and methane seep [4]. Considering the methane concentration is also high in our sampling site, the mass spectra have been only searched against the protein database of the family Methylococcaceae in order to narrow the range.

Totally, 32 unique acetylation sites from 23 proteins were identified (Additional file 3: Table S3), including methanol dehydrogenase associated peptide, ABC transporter permease, Trk system potassium uptake protein, peptidase M16, ATP-dependent RNA helicase, translocase subunit, 3′,5′-cyclic AMP phosphodiesterase, DNA polymerase I, IS256 family transposase, ATPase of the AAA+ class, tyrosine recombinase, chromosome partition protein and other uncharacterized proteins. According to the GO annotation, these PTMs are probably participate in the adaptation to the extreme environment of microbes mainly by regulating methane oxidation, cell redox homeostasis, substance transmembrane transport and DNA repair. It is essential to obtain the exact protein and
Various acetylation data of microbes from different tissues of the host, such as gill, seta and gut, where more microbes are available. However, we have firstly identified candidate acetylation sites in the microbes of deep-sea chemosynthetic ecosystems preliminarily. Intensive study on the PTMs in the microbes is also expected to facilitate clarifying the symbiont mechanism.

Conclusion
In conclusion, this work provides the first global lysine acetylation data in the deep-sea hydrothermal vent shrimp, *A. longirostris* and even in the marine invertebrates. A total of 501 unique acetylation sites from 206 proteins were identified. Three consensus sequence motifs were extracted. Functional analysis revealed that the protein acetylation was involved in diverse metabolic pathways, oxidative phosphorylation, and many other cellular processes. Abundant lysine acetylation was discovered in mitochondrion and peroxisome. Acetylation associated with stress response was further detected and the acetylation of key respiratory protein Hcs was specifically characterized. Therefore, our study provides new insights into the regulation function of the reversible lysine acetylation, and supplies rich candidates for exploring the functions of acetylation in the adaptation to hydrothermal vents in *A. longirostris*, and in other shrimps.

Additional files

Additional file 1: Table S1. Information for the acetylated proteins in *Alvinocaris longirostris* (XLSX 143 kb)
Additional file 2: Table S2. Information for the protein-protein interaction network in *Alvinocaris longirostris* (XLSX 20 kb)
Additional file 3: Table S3. Information for the acetylated proteins of microbes associated with abdomen of *Alvinocaris longirostris* (XLSX 15 kb)

Abbreviations
Act6: Actin 6; Actc1: Actin alpha cardiac muscle 1; AGC: Automatic gain control; Aks: Arginine kinases; Aldh1l1: 10-formyltetrahydrofolate dehydrogenase; Alh4a1: Delta-1-pyruvoly-2-carboxylic dehydrogenase mitochondrial; AlHc1: Alvinocaris longirostris hemocyanin; AlHc1: Alvinocaris longirostris hemocyanin subunit 1; CH4: Methane; FDR: False discovery rate; GO: Gene Ontology; H3: Hydrogen sulfide; Hadha: Trifunctional enzyme subunit alpha mitochondrial; Hc: Hemocyanin; Hsp70: Heat shock protein 70; IACUC: Institutional Animal Care and Use Committee; KEGG: Kyoto Encyclopedia of Genes and Genomes; LC: Liquid chromatography; MCODE: Molecular complex detection; MS: Mass spectrum; NCE: Normalized collisional energy; NSI: Nanospray ionization; O2: Oxygen; PPI: Protein-protein interaction; PTMs: Post-translational modifications; ROV: Remotely operated vehicle; RV: Research vessel; Top2b: DNA topoisomerase 2-beta

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Availability of data and materials
All data in this study are included in the article and its additional files.

Authors’ contributions
ZY and MH conceived and designed the project. ZS performed lab work, analyzed data and wrote the manuscript. JC and ZS revised the manuscript. All authors read and approved the final version of the manuscript.

Ethics approval
The cruise and field study were approved by Chinese Academy of Sciences. The experiments proceeded according to the guidelines of the Institutional Animal Care and Use Committee (IACUC) of the Chinese Academy of Sciences (No. 2011–2). The research was approved by the Committee on the Ethics of Animal Experiments of the Institute of Oceanology in the Chinese Academy of Sciences.

Competing interests
The authors declare that they have no competing interests.

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