Dynamic transcriptome profiling exploring cold tolerance in forensically important blow fly, Aldrichina grahami (Diptera: Calliphoridae)

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Abstract

Background: Aldrichina grahami (Diptera: Calliphoridae) is a forensically important fly, which has been widely applied to practical legal investigations. Unlike other necrophagous flies, A. grahami exhibits cold tolerance which helps to maintain its activity during low-temperature months, when other species are usually not active. Hence, A. grahami is considered an important forensic insect especially in cold seasons. In this study, we aim to explore the molecular mechanisms of cold tolerance of A. grahami through transcriptome.

Results: We collected eggs and larvae (first-instar, second-instar and third-instar) at three different temperatures (4 °C, 12 °C and 20 °C) and performed RNA-seq analyses. The differentially expressed genes (DEGs) associated with the cold-tolerance were screened out. The Venn analysis of DEGs from egg to third-instar larvae at three different temperatures showed there were 9 common genes. Candidate biological processes and genes were identified which refer to growth, and development of different temperatures, especially the chitin and cuticle metabolic process. The series-clusters showed crucial and unique trends when the temperature changed. Moreover, by comparing the results of growth and developmental transcriptomes from different temperatures, we found that DEGs belonging to the family of larval cuticle proteins (LCP), pupal cuticle protein (CUP), and heat shock proteins (HSP) have certain differences.

Conclusions: This study identified functional genes and showed differences in the expression pattern of diverse temperatures. The DEGs series-clusters with increasing or decreasing trends were analyzed which may play an important role in cold-tolerance. Moreover, the findings in LCP, CUP and HSP showed more possible modulations in a cold environment. This work will provide valuable information for the future investigation of the molecular mechanism of cold tolerance in A. grahami.

Keywords: Aldrichina grahami, Cold tolerance, Differentially expressed genes, Transcriptome, Forensic entomology

Background

Forensic entomology, which explores the succession pattern and developmental stages of insects found on the decomposed cadavers, has been increasingly recognized as an important tool in the medico-legal discipline [1]. The knowledge of forensic entomology offers vital clues for the postmortem interval (PMI). The developmental time of the immature stages of necrophagous flies (eggs, larvae, pupae) has been a good indicator in determining the minimum PMI (PMImin) [2, 3]. Comparing with other species, Calliphoridae has been investigated frequently because of predominance in colonizing corpses [4, 5]. Aldrichina grahami, a forensically important blow fly, is mainly distributed in Asia [6, 7]. In 1929, A. grahami was first reported by Aldrich in California [8]. Aldrichina grahami has been widely applied to practical legal investigations [9, 10] and exhibited some unique characteristics like low-temperature tolerance, which...
makes it active in the cold winter when other necrophagous flies almost inactive [11–13]. Several researchers are interested in development of A. grahami on different common temperatures, but only a few Japanese researchers explored its low-tolerance [14, 15]. In 1962, Japanese researchers found that the adults of A. grahami could survive when the temperature was relatively cold (2 to 7 °C) [16]. However, the hatching rates of eggs kept at this environmental condition was zero [16]. In 1985, it also had been reported in Japan that lethal time of 50% (LT50) of egg, larva, and pupal stages of A. grahami exposed to 4 °C was approximately 4, 12, and 6 days, respectively [17]. An insect species’ capacity for cold tolerance is regarded as a dominant factor for its population expansion [18] and geographical distribution [19]. It is well-known that insects always face a great challenge to survive in an extremely cold environment [20]. Recently, there are a few studies about low-temperature adaptability of other flies [21, 22], however, few researches focus on A. grahami and its unique characteristic at a molecular level.

Cold tolerance of insects, an intricate adaptive response which subjects to biochemical and physiological regulation, has become a hot topic in recent years [22–25]. In temperate climates, the survival of insects mainly depends on the adaptations to cold temperature [26]. For instance, the insect cuticle is secreted by epidermal cells and covers the whole surface of the body [27], which provides protection against physical injury and dehydration [28]. Previous studies have shown that the properties of the insect cuticle may be determined by the complex structural interactions between chitin polysaccharides and cuticular proteins (CPs) [29]. Moreover, extreme cold environment can affect cellular membrane integrity by inducing the transition of membrane’s phospholipid bilayer [30]. These changes in membrane fluidity alter function and activity of membrane-bound enzyme [31, 32]. Lots of attempts have been made to elucidate the mechanisms of insect cold tolerance [33, 34]. In cold winter, the metabolic activity rate of insects is generally low. And there has little changed in their state, organ development, and tissue differentiation [35]. However, their physiological metabolic processes are still active, such as energy metabolism, endocrine regulation, lipid metabolism and sugar metabolism [30, 36, 37]. In energy metabolism, it has been proved that low-molecular-weight sugars and polyols are vital intermediate metabolites and energy substances in many insect species in a cold environment [18]. Importantly, polyols and sugars generally accumulate and work as cryoprotectants at low temperature [17]. Although different insects accumulate dissimilar polyols, the increase in glycerol content is associated with the cold environment [33, 34]. The adaptability of these compounds leads to efficient resource utilization and maintain the dynamic balance of nutrition, thus providing a higher level of cold tolerance of overwintering insects [20]. In lipid metabolism, desaturation of fatty acids in membrane’s phospholipid and desaturation of triacylglycerides decrease the melting point, leading to an increasing fluidity and accessibility at low temperatures [38, 39]. Moreover, it has been reported that the inhibition of several metabolic pathways at low temperature may avoid damaging imbalance. For instance, enzyme activities and expressions in insects that regulated the physiological processes were related to survival, growth, and development [40]. Last but not least, the biochemical mechanisms of cold tolerance are reported to involve antioxidant defense [41] and aminoacyl-tRNA biosynthesis [42].

In the present study, we used RNA-seq technique to identify the cold tolerance related genes by building transcriptomes profile of eggs, and larvae (1st instar to 3rd instar) under three different temperatures (4 °C, 12 °C and 20 °C). DEGs among eggs and different instars were identified by comparative transcriptome analysis. Finally, differentially expressed LCP, CUP, and HSP in egg and larvae were examined for transcriptome data validation.

**Results**

**Overview of RNA-Seq data**

A total of 36 libraries (Additional file 1: Table S1) were sequenced from egg and larvae (first-instar, second-instar and third-instar) at three different temperatures (4 °C, 12 °C and 20 °C) of 12 groups (n = 3 for each), representing the egg stage at 4 °C low temperature (L0), 12 °C middle temperature (M0) and 20 °C relatively high temperature (H0); the first-instar larvae stage at low temperature (L1), middle temperature (M1) and relatively high temperature (H1); the second-instar larvae stage at low temperature (L2), middle temperature (M2) and relatively high temperature (H2); and the third-instar larvae stage at low temperature (L3), middle temperature (M3) and relatively high temperature (H3), respectively. Totally, there were 250.58 Gb clean bases was obtained. About 6.26–7.90 Gb clean bases were produced for each library. After discarding low-quality reads, RNA-seq yielded from 62.55 to 79.02 Mb clean reads with average about 90% Q30 bases for each sample, which were used for all further expression analysis. Among the total number of clean reads from 36 samples, 80.49 to 88.12% were successfully mapped against the reference A. grahami genome. The percentage of the unique mapping reads was 48.78 to 67.39% in each sample (Table 1). As the correlation of transcript expression level is a vital indicator for the reliability of the experimental results, we found that in egg stage, the Pearson correlation coefficient between three biological replicates of three groups in this study had high repeatability (i.e., all
Moreover, the results of larvae were showed high repeatability (Additional file 2: Figure S1).

### DEGs involved in the three different temperatures

In a set condition (p-value < 0.05, and fold change ≥2.0), we screened the DEGs to determine the differences under different constant temperatures. In the egg stage, there were 5602, 7592, 9292 DEGs in the comparison of L0 vs M0, M0 vs H0, L0 vs H0, respectively. In first-instar, 3766, 2095, 3075 genes were found to be DEGs in the comparison of L1 vs M1, M1 vs H1; L1 vs H1, respectively. In the second-instar and third-instar larvae stage, thousands of DEGs were also obtained in a similar comparison (Fig. 2). To sum up, the comparison of low temperature and the high temperature had the largest number of DEGs. And the egg stage was the period most affected by environmental
**Fig. 1** The Pearson correlation coefficient between three biological replicates of egg stage at 20 °C (H0), egg stage at 12 °C (M0), and egg stage at 4 °C (L0).

**Fig. 2** The number of differentially expressed genes between the comparison groups. Up-regulated DEGs (red), and down-regulated DEGs (green) were presented by histogram.
temperature. Likewise, as the expression patterns of all the DEGs under different temperature at egg stage were analyzed (Additional file 3: Figure S2), we found out obvious differences between 4 °C and 20 °C. However, other stages didn’t show such distinct differences.

Next, we investigated the DEGs of one development stage under three different temperatures (4 °C, 12 °C and 20 °C) at the same development stages. In egg stage, plenty of DEGs were screened in the two of the three comparisons of L0 vs M0, M0 vs H0, L0 vs H0. The number 3185 was the largest of comparisons among the four groups. In first-instar, 440 DEGs were found to be an intersection in the two of the three comparisons of L1 vs M1, M1 vs H1; L1 vs H1. In the second-instar and third-instar larvae stage, 927 and 530 DEGs were also obtained in a similar comparison (Fig. 3). Subsequently, we found out 9 DEGs (Fig. 3e) were the intersection in the two of these four comparisons, suggesting to be the most important for cold tolerance. However, none has the same trend when the temperature drops gradually in all four stages (Additional file 4: Table S2). To be mentioned, great changes had taken place in flies from eggs to mature larvae, especially in egg stage, both morphologically and genetically. Hence, DEGs of intersection in the two of these three comparisons were also worth studying.

Series-cluster analysis and functional annotation of the clusters

The expression patterns not only indicate the diverse and complex interactions among genes, but also suggest that genes with similar expression patterns may have similar functions when in a cold environment. In the egg libraries (0 libraries), a total of 13,356 genes were found to be DEGs. Then 12 series-clusters (Fig. 4) were obtained based on them. Each gene cluster exhibited a distinctive expression pattern. The largest group of 0 libraries is cluster 12 with 2585 (19.4%). Importantly, it can be found out that cluster 6 and cluster 9 had extremely different trends when the temperature changed. There were 1272 DEGs in the cluster and among them, 122 (9.6%) was novel genes. The cluster 6 showed directly decreasing trend when facing cold. On the contrary, cluster 9 showed a completely opposite trend. In cluster 9, there were 1615 DEGs and 626 (38.8%) was the novel one. Moreover, in the first-instar larvae libraries (1 libraries) (Additional file 5: Figure S3), the second-instar larvae libraries (2 libraries) (Additional file 6: Figure S4), and the third-instar larvae libraries (3 libraries) (Additional file 7: Figure S5), there were 12,563, 12,354, and 11,242 DEGs had been analyzed as series-cluster, respectively. There was also distinct increasing or decreasing trend in their clusters when the environment temperature was cooling down. Series-cluster analysis and functional annotation of the clusters provided

Fig. 3 The Venn diagram showed the shared and unique genes of DEGs were statistically analyzed. a the Venn diagram in egg stage, b the Venn diagram in first-larvae stage, c the Venn diagram in second-larvae stage, d the Venn diagram in third-larvae stage, e the Venn diagram in (a, b, c and d)
crucial clues of the key DEGs that worked in cold temperature. For example, in cluster 7 of 3 libraries, we found out the third instar larval cuticle proteins (LCP) had significant changes.

**GO and KEGG pathway analysis of DEGs in 0 libraries**

A total of 7250 DEGs were annotated into GO terms involved in the egg stage. Undoubtedly, the L0 vs H0 has the largest number of DEGs. Among them, 1992 (31.4%) novel genes were found out. DEGs were classified into 48 subcategories within three standard categories (molecular functions, biological processes and cellular components) (Fig. 5). “Cellular process” and “metabolic process” were the most enriched in the biological process domain. In cellular component category, “membrane” and “membrane part” were the highest enriched, while “binding” and “catalytic activity” were the most enriched in the molecular function category. Meanwhile, the significant enriched GO terms (p-value < 0.05) involved in the egg stage were determined from the DEGs of L0 vs M0, M0 vs H0, L0 vs H0 (See for a full list of GO terms in Table 2). DEGs of chitin metabolic process, glucosamine-containing compound metabolic process, amino sugar metabolic process and structural constituent of cuticle were all significantly enriched in L0 vs M0, M0 vs H0, L0 vs H0.

Meanwhile, KEGG pathway related to temperature changes were also recognized (Fig. 6), including cAMP
signaling pathway, cGMP-PKG signaling pathway, pentose and glucuronate interconversions, oxytocin signaling pathway and circadian entrainment, etc. Importantly, the cAMP and cGMP-PKG signaling pathway, which are involved in energy metabolism and cell growth and differentiation, are suggested to be important during the temperature changes in *A. grahami*. To be mentioned, endocrine system and digestive system might play a vital role in a cold environment.

**GO and KEGG pathway analysis of DEGs in 1 and 2 library**

A total of 5121 DEGs and 6354 DEGs were annotated into GO terms involved in the first-instar and the second-instar, respectively. DEGs of L1 vs H1 (Additional file 8: Figure S6) were classified into subcategories within three standard categories. “Cellular process” and “biological regulation” were the most enriched terms in the biological process domain. Similarly, in cellular component category, “membrane” and “membrane part” were the highest enriched, while “binding” and “catalytic activity” were mostly enriched terms in the molecular function category. Moreover, the significantly enriched GO terms (p-value < 0.05) involved in the first-instar larvae stage were determined from the DEGs of L1 vs M1, M1 vs H1; L1 vs H1 (Additional file 9: Table S3). Response to stimulus, circadian rhythm, regulation of glucose metabolic process, chitin metabolic process, glucosamine-containing compound metabolic
process, structural constituent of cuticle and amino sugar metabolic process were all significantly enriched in the DEGs of L1 vs M1, M1 vs H1; L1 vs H1.

KEGG pathway related to temperature changes was also recognized (Additional file 10: Figure S7), including protein digestion and absorption, endocrine resistance, cGMP-PKG signaling pathway, cAMP signaling pathway, longevity regulating pathway and circadian entrainment, etc. In addition, the results of GO (Additional file 11: Figure S8 and Additional file 12: Table S4) and KEGG pathway (Additional file 13: Figure S9) analysis in 2 library were similar to that in 1 library.

| Tissue comparison | GO Term                                      | Rich Ratio | P-value  |
|-------------------|----------------------------------------------|------------|----------|
| H0 vs L0          | chitin metabolic process                      | 0.78       | 4.14E-09 |
|                   | glucosamine-containing compound metabolic process | 0.78       | 5.92E-09 |
|                   | amino sugar metabolic process                 | 0.77       | 1.20E-08 |
|                   | chitin binding                                | 0.78       | 1.41E-08 |
|                   | structural constituent of cuticle              | 0.76       | 1.39E-07 |
|                   | response to stimulus                          | 0.60       | 2.01E-06 |
|                   | G-protein coupled receptor activity           | 0.73       | 0.0001   |
|                   | DNA-dependent DNA replication                 | 0.80       | 0.0006   |
|                   | circadian rhythm                              | 0.83       | 0.0016   |
|                   | chitin-based cuticle sclerotization           | 0.86       | 0.0030   |
|                   | RNA metabolic process                         | 0.63       | 0.0330   |
| H0 vs M0          | glucosamine-containing compound metabolic process | 0.73       | 8.75E-12 |
|                   | aminoglycan metabolic process                 | 0.70       | 1.71E-11 |
|                   | amino sugar metabolic process                 | 0.72       | 1.76E-11 |
|                   | chitin metabolic process                      | 0.72       | 2.33E-11 |
|                   | chitin binding                                | 0.70       | 1.60E-09 |
|                   | structural constituent of cuticle              | 0.63       | 1.76E-05 |
|                   | anatomical structure development              | 0.50       | 0.0012   |
|                   | muscle cell development                       | 0.80       | 0.0017   |
|                   | DNA binding                                   | 0.51       | 0.0036   |
|                   | chitin-based cuticle sclerotization           | 0.76       | 0.0041   |
|                   | cuticle development                           | 0.62       | 0.0057   |
| M0 vs L0          | structural constituent of cuticle             | 0.76       | 2.13E-27 |
|                   | glucosamine-containing compound metabolic process | 0.68       | 2.68E-21 |
|                   | amino sugar metabolic process                 | 0.68       | 5.85E-21 |
|                   | chitin metabolic process                      | 0.68       | 1.42E-20 |
|                   | aminoglycan metabolic process                 | 0.64       | 4.45E-20 |
|                   | chitin binding                                | 0.64       | 1.09E-15 |
|                   | G-protein coupled receptor activity           | 0.54       | 2.02E-06 |
|                   | muscle cell development                       | 0.70       | 0.0003   |
|                   | cuticle development                           | 0.52       | 0.0003   |
|                   | circadian rhythm                              | 0.62       | 0.0004   |
|                   | chitin-based cuticle sclerotization           | 0.67       | 0.0007   |

Table 2 GO functional enrichment analysis related with temperature of the DEGs of H0 vs L0, H0 vs M0 and M0 vs L0

A total of 5593 DEGs were annotated into GO terms involved in the third-instar. DEGs of L3 vs H3 were classified into subcategories within three standard categories (Additional file 14: Figure S10). “Cellular process” and “biological regulation” were the most enriched in the biological process domain. “Membrane” and “membrane part” were the highest enriched in cellular component category. And “binding” and “catalytic activity” were the most enriched in the molecular function category. Moreover, the significant enriched GO terms (p-value < 0.05) involved in the third-instar larvae stage were determined.

GO and KEGG pathway analysis of DEGs in 3 libraries
from the DEGs of L3 vs M3, M3 vs H3 and L3 vs H3 (Additional file 15: Table S5). Unlike 0, 1 and 2 libraries, DEGs of chitin metabolic process, chitin binding, structural constituent of cuticle were not obviously enriched in 3 libraries. DEGs of response to starvation, energy reserve metabolic process, response to stimulus, glucosamine-containing compound metabolic process and amino sugar metabolic process were all significantly enriched in 3 libraries.

Furthermore, KEGG pathway related to temperature changes were also recognized (Additional file 16: Figure S11), including protein digestion and absorption, glycosphingolipid metabolism, PPAR signaling pathway, PI3K-Akt signaling pathway and RNA polymerase, etc.

**Gene-co-expression**

The gene co-expression net-work of DEGs in the cold temperatures was analyzed (Fig. 7). There are several core genes with the highest degrees connect with most adjacent genes in the network, which are frequently identified as key indicators. Of these, DEGs which were involved in chitin metabolic process (LCP2, LCP8 and CHI10) and structural constituent of cuticle (CU01, CUP7 and CUP9) showed the strongest relationship in net-work. Next, gene co-expression net-work analysis of DEGs in 12 °C (Additional file 17: Figure S12) and 20 °C (Additional file 18: Figure S13) were also constructed. From the results of the high temperature group, there were different core genes compared with low temperature group, but still involved in chitin metabolic process, lipid metabolism, etc. Gene co-expression analysis could provide the vital genes that may regulate the adaptations in response to a cold environment.

**Validation of sequencing data**

A total of 9 DEGs were selected for the purpose of validating whether the sequencing and analysis were reliable. Four DEGs were selected for qRT-PCR quantification from RNA sample in 0 library. Similarly, five DEGs were selected from RNA sample in 3 library. The details of those unigenes and primer pairs used in this study are shown in Additional file 19: Table S6. A comparative analysis of all the selected genes showed a similar expression pattern in the qRT-PCR analysis as observed in RNAseq data (Fig. 8). To be mentioned, the expression trends of several crucial DEGs, such as LCP8, LCP5, HSP68 and OBP99a, were as expected.
Discussion
Developmental transcriptomes comparisons between 4 °C and 20 °C

It is known that 20 °C is the optimum growth temperature for the development of A. grahami, while 4 °C is an extreme low temperature [16, 17]. Hence, the developmental transcriptomes profile of separated instars under two different temperatures were compared to explore genes that related to the cold tolerance. Many biological processes were enriched in both 4 °C and 20 °C libraries. DEGs of Chitin metabolic process, structural constituent of cuticle, glucosamine-containing compound metabolic process, and amino sugar metabolic process were all significantly enriched in H0 vs L0, H1 vs L1, and H2 vs L2. Chitin and sugar metabolic process showed an important role in a cold environment. In the results of gene-co-expression, it also showed the importance of Chitin metabolic process. It was consistent with that chitin is a component of insect cuticle [29], which could protect the body of insect [27], and CPs and sugar were vital intermediate metabolites in many insect species in a cold environment [10]. Moreover, our results also suggested that endocrine system and digestive system played vital roles in cold tolerance. Especially, the cAMP and cGMP-PKG signaling pathway were all significantly enriched in the DEGs of H0 vs L0, H1 vs L1, and H2 vs L2, indicating the involvement of energy metabolism [43, 44] and cell growth and differentiation [45, 46]. However, unlike 0, 1 and 2 libraries, chitin metabolic process, chitin binding, structural constituent of cuticle were not obviously enriched in 3 libraries. In the third-instar larvae, response to starvation, energy reserve metabolic process, glucosamine-containing compound metabolic process and amino sugar metabolic process were all significantly enriched. This may be caused by energy saving for the pupal stage [47].

DEGs may play crucial roles in cold tolerance in A. grahami

Firstly, we found that 9 DEGs were the intersection-based common genes in all libraries, suggesting their involvement in cold tolerance. Unfortunately, none showed consistent trend in the four developmental
stages in our studies when different temperatures. This may be caused by the morphological and genetic changes taken place from eggs to mature larvae in flies [48–50]. In addition, we explored the expression patterns in each library via a cluster way, in which the same cluster not only indicates the diverse and complex interactions among genes, but also suggests that genes may have similar functions when in a cold environment. Twelve series-clusters were obtained in each library. We found that cluster 6, cluster 11, cluster 11, cluster 2 had directly decreasing trend when facing cold in 0 library, 1 library, 2 library, 3 library, respectively. On the contrary,
cluster 9, cluster 8, cluster 4, cluster 11 had a continuous increasing trend in 0 library, 1 library, 2 library, 3 library, respectively. As we aimed to explore the cold tolerance, it was an effective way to search for the key DGEs in these clusters. Moreover, DEGs which were involved in chitin metabolic process (LCP2, LCP8 and CHI10) and structural constituent of cuticle (CU01, CUP7 and CUP9) showed the strongest relationship in net-work, indicating that cuticle protein family may play significant roles in cold tolerance. It is noticeable that lipid metabolism (WAT and FACR1) also played an important role when facing cold. Therefore, we proposed that LCP, CUP and HSP might be responsible for cold-tolerance in *A. grahami*.

Although chitin is one of the vital component of the cuticle, there are various larval cuticle proteins, providing protection in cold [51, 52]. It is reported that when exposed to cold or some other environmental stresses, insects can synthesize heat shock proteins, which function as molecular chaperones in protecting cellular proteins [53]. Moreover, HSPs have also been reported in *Drosophila* [54, 55], *Bemisia tabaci* [53], *Bombyx mori* [56], etc. Surprisingly, we found that odorant-binding proteins (OBPs) had significant increases in H3 vs L3, but no differences in H0 vs L0. OBPs are the key step in the insect olfaction [57], suggesting that *A. grahami* need better olfactory sensation to search for food in cold weather. Finally, we chose 9 unigenes to examine our hypothesis via qPCR method, which were involved in synthesis of cuticle proteins, heat shock proteins, and odorant-binding proteins. These results showed the same trend with transcriptome data (Fig. 8).

The developmental stages of insect include eggs, larvae, pupae and adult. In this study, we focused on eggs and larvae stages depending on the following reasons: Firstly, low-temperature tolerance at egg stage directly affects survival rate. Larvae are also susceptible to cold temperature, while pupae and adult have their outermost shell or fluff to keep warm. Secondly, gene changes in eggs and larvae stages are mostly continuous. Dramatic changes would take place in flies from larvae to adult, both morphologically and genetically. Moreover, as the individual differences of transcriptome in adults are various, it is insensitive to explore its features by adults at first. Hence, we chose eggs and larval stages to explore the cold tolerance in *A. grahami*. Apart from the feature exploration, we also focused on the developmental expression profiles subsequently. Then we selected the key DEGs related to development time, which could be applied in PMI min deduction in the future.

Importantly, when study the low temperature on the development of the blow fly, apart from low temperature, several factors should be taken into consideration. Firstly, the effects of constant temperature and fluctuating temperature on the development of blow fly [15]. Secondly, the insects from different geographic regions might have different cold tolerance [12]. Lastly, as blow fly is a kind of heterothermy animal, its larvae often gathering and feeding together and can produce heat [58, 59]. The larval mass effect can help the larvae resist the cold environment. Hence, it is sensible to consider and avoid these aspects when exploring relative studies. To be mentioned, *A. grahami* has been widely applied to practical forensic investigations [9, 10]. And the low-temperature tolerance of *A. grahami* [16, 17] makes it more important among necrophagous flies, especially in the cold environment. However, it cannot be ignored that 4 °C is an extreme low temperature. Under this condition, the egg of *A. grahami* indeed requires a long period to hatch. Moreover, it is also difficult for the adult to emerge from pupae. Hence, it would be meaningful to explore a low temperature between 4 °C and 12 °C in the future, at which the *A. grahami* could fly, mate, oviposit and develop slowly but more naturally.

**Conclusion**

In this study, we provide a new insight into transcriptional profiles of egg and larvae of *A. grahami* under different temperatures condition. Various pathways and biological processes were annotated that related with dynamic gene expression of *A. grahami* under low temperature treatment, like chitin metabolic process, structural constituent of cuticle, glucosamine-containing compound metabolic process, and amino sugar metabolic process, etc. Differentially expressed gene cluster and molecular network also indicated a complicate mechanism beneath the cold-tolerance of *A. grahami*. Furthermore, the DEGs, such as LCP, CUP and HSP, showed more possible modulations in a cold environment. The accumulated knowledge about *A. grahami* biology makes it to be a promising model for detailed mechanism researches on those forensically important fly species. This work provided valuable information for future research on the mechanism of cold-tolerance of *A. grahami*.

**Methods**

**Specimen collection and fly rearing**

The oriental adult specimens of *A. grahami* were obtained using pork in Changsha city (28°12′ N, 112°58′ E), Hunan province, China, in March 2016. The adult flies were maintained in a rearing box (35 × 35 × 35 cm³) at 25 °C with 70% humidity and 12:12 h light: dark photo-period. One dish contained 1:1 mixture of sugar and milk powder as food for adults, and another dish contained 15 g of fresh pig lung to induce egg laying. After fresh pig lung was set up, the egg laying was checked and eggs were collected every half hour. Each egg was
transferred to a new dish in an artificial climate box until its wandering stage. Three artificial climate boxes were used in our study at constant temperatures of 4 °C, 12 °C, and 20 °C, respectively. Due to the timeframe for development of eggs varies greatly at different temperatures, we have examined the time for egg development under each temperature in our preliminary study. We found that it took approximately 10 days, 52 h, and 28 h to develop into first-instar larvae (from eggs) at 4 °C, 12 °C, and 20 °C, respectively. Therefore, we used the following time points to represent egg stage as 5 days for 4 °C (L0), 26 h for 12 °C (M0), and 14 h for 20 °C (H0). Samples were harvested at designed time points and snap frozen in liquid nitrogen and then stored at −80 °C. Three biological replicates of each group were used for RNA extraction and transcriptome sequencing.

RNA extraction and library preparation for transcriptome sequencing

The Trizol method (Invitrogen, USA) was adopted to extract the total RNA of samples collected. The RNA quality was assessed by formaldehyde gel electrophoresis and quantitated spectrophotometrically (NanoDrop 2000). DNase I (TakaRa, Japan) was used for RNA purification. Double-stranded cDNA was synthesized using random hexamer-primers, reverse transcriptase, DNA polymerase I and RNaseH by taking short fragments as templates. Next, double-stranded cDNA was subsequently subjected to end-repair and ligation with adapters. These modified products were enriched with PCR to construct the final cDNA library. After test the quality of libraries, they were loaded onto the flow cell channels of the BGISEQ500 platform (BGI-Shenzhen, China).

Sequence reads mapping and assembly

Firstly, raw reads of fastq format were filtered by removing reads containing adapters, reads containing poly-N and low quality reads, then high-quality clean reads were obtained (reads contain 20% base quality lower than Q20). At the same time, Q20, Q30, GC-content and sequence duplication level of the clean data were calculated. All the succeeding analyses were carried out using high quality clean reads. Clean reads were mapped to the Aldrichina grahami genome assembly (NCBI: PRJNA513084) by using HISAT2 [60] with following parameter: --dta --phred64 unstranded --new-summary -x index – 1 read_r1–2 read_r2 (PE).

Differential expression analysis

Differential expression analysis of two groups was performed using the parameter FPKM (Fragments per kilo-base of transcript per million mapped reads), which was applied to quantify the gene expression levels. HTseq [61] was used for count calculation and FPKM was calculated with NCBI gtf file through gene length annotation. The DEGseq package was applied to filter the DEGs with a fold change > 2 or fold change < 0.5, and false discovery rate (FDR) < 0.05 [62].

GO enrichment and KEGG pathway enrichment analysis

Gene ontology (GO) analysis was performed to facilitate elucidating the biological implications of unique genes in the significant or representative profiles, which is conducive to find those GOs with more concrete function description in our study [63]. KEGG [64] is a database resource for understanding high-level functions and utilities of the biological system. Pathway analysis was used to find out the significant pathways of the DEGs via the KEGG database. A Fisher exact test was used to find the vital enrichment pathway with the threshold of the significance of p-value < 0.05 and FDR < 0.05, which could find those crucial pathways in our study.

Classification and co-expression of DEGs

Series cluster analysis was performed using STEM [65] to classify the DEGs in twelve clusters based on the FPKM change tendency of genes on three different temperatures of each developmental stage. Fisher’s exact test and the multiple comparison tests were used to calculate the significant levels of profiles [66, 67]. Gene co-expression network analysis was performed to track the interactions among the DEGs, according to the dynamic expression changes on three different temperatures of each developmental stage. Pearson correlation was applied to each pair of genes and the significantly correlated pairs were used to construct the network [68].

Quantitative real-time PCR analysis

To assess the reliability of the sequencing and analysis by quantitative real-time PCR (RT-qPCR), we used the same RNA samples for transcriptome sequencing. The RT-qPCR reactions were carried out on a 7500 Real-Time PCR System (Applied Biosystems) using the 2 × T5 Fast qPCR Mix (SYBR Green I) (Qingke Biotechnology Co., Ltd. Hunan, China) according to the manufacturer’s instructions. Every RT-qPCR amplification mixtures (20 μL) contained 2 × T5 Fast qPCR Mix (10 μL), 50 × ROX Reference Dye II (0.4 μL), each forward and reverse primers (0.8 μL), diluted cDNA (2 μL) and RNase-free water (6 μL). The PCR reaction was set as follows: an initial denaturation at 95 °C for 1 min, followed by 40 cycles of 95 °C for 10 s; 60 °C for 10 s; 72 °C for 15 s. Relative gene expression levels were calculated by 2^\triangle\triangle\Ct, GST1 were selected as internal reference genes [69]. The primers of all selected genes were designed by Primer Premier 5.
expression net-work in first-instar larvae stage at 12°C (M1), (c) the gene co-expression net-work in second-instar larvae stage at 12°C (M2), (d) the gene co-expression net-work in third-instar larvae stage at 12°C (M3).

Additional file 18: Figure S13. The gene co-expression net-work of DEGs in the relatively high temperatures (20°C) was analyzed. (a) the gene co-expression net-work in egg stage at 20°C (H0), (b) the gene co-expression net-work in first-instar larvae stage at 20°C (H1), (c) the gene co-expression net-work in second-instar larvae stage at 20°C (H2), (d) the gene co-expression net-work in third-instar larvae stage at 20°C (H3).

Additional file 19: Table S6. Primers used for qPCR validation.

Abbreviations
CH10: Chitinase 10; CPs: Cuticular proteins; CU01: Cuticle protein 1; CUP: Pupal cuticle protein; CUP7: Pupal cuticle protein 7; DEGs: Differentially expressed genes; FPKM: Fragment per kilobase of exon model per million mapped reads; GO: Gene ontology; H0: Egg stage of Aldrichina grahami at 20°C; H1: First-instar larvae stage of Aldrichina grahami at 20°C; H2: Second-instar larvae stage of Aldrichina grahami at 20°C; H3: Third-instar larvae stage of Aldrichina grahami at 20°C; HSP: Heat shock proteins; HSP68: Heat shock protein 68; KEGG: Kyoto encyclopedia of genes and genomes; L0: Egg stage of Aldrichina grahami at 4°C; L1: First-instar larvae stage of Aldrichina grahami at 4°C; L2: Second-instar larvae stage of Aldrichina grahami at 4°C; L3: Third-instar larvae stage of Aldrichina grahami at 4°C; LCP: Larval cuticle proteins; LCP2: Larval cuticle protein 2; LCP5: Larval cuticle protein 5; LCP8: Larval cuticle protein 8; LT50: Lethal time of 50%; M0: Egg stage of Aldrichina grahami at 12°C; M1: First-instar larvae stage of Aldrichina grahami at 12°C; M2: Second-instar larvae stage of Aldrichina grahami at 12°C; M3: Third-instar larvae stage of Aldrichina grahami at 12°C; PMI: Minimum postmortem interval; RT-qPCR: Quantitative real-time PCR; WAT: Fatty acyl-CoA reductase wAT.

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Authors’ contributions
ZL, HH, FM, and JC conceived the study and participated in its design. ZL and HH performed experiments. ZL, HH, and YJ analyzed the data. ZL and JC wrote the manuscript, which all authors have read and approved the manuscript.

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Availability of data and materials
The raw reads has been submitted to SRA under accession number PRJNA565270 (https://www.ncbi.nlm.nih.gov/bioproject/PRJNA565270/). Aldrichina grahami genome sequence data: NCBI: PRJNA513084.

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

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

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References

1. Kong-Klaev T, et al. Impact of abiotic factor changes in blowfly, Achatocandrus ruficollis (Diptera: Calliphoridae), in northern Thailand. Parasitol Res. 2014;113(4):1353–60.
2. Amendt J, et al. Forensic entomology: applications and limitations. Forensic Sci Med Pathol. 2011;7(4):379–92.
3. Boehme P, et al. The analysis of temporal gene expression to estimate the age of forensically important blow fly pupae: results from three blind studies. Int J Legal Med. 2014;128(3):565–73.
4. Sukontat K, et al. Forensic entomology cases in Thailand: a review of cases from 2000 to 2006. Parasitol Res. 2007;101(5):1417–23.
5. Goff ML. Comparison of insect species associated with decomposing remains recovered inside dwellings and outdoors on the island of Oahu, Hawaii. J Forensic Sci. 1991;36(3):748–53.
6. Chen LS, et al. Estimation time of death by necrophagous flies life cycle. Forensic Sci Int Genet. 2013;7(1):139–43.
7. Xu H, et al. Age-dependent changes in cuticular hydrocarbons of larvae in Aldrichina grahami (Diptera: Calliphoridae). Forensic Sci Int. 2014;242:236–41.
8. Aldrich JM. New two-winged flies of the family Calliphoridae from China. Proc U S Natl Mus. 1931;284:1–5.
9. Guo YD, et al. Application of Aldrichina grahami (Diptera, Calliphoridae) for forensic investigation in central South China. Rom J Leg Med. 2011;19:55–8.
10. Li X, et al. Mitochondrial DNA and STR analyses for human DNA from maggots crop contents: a forensic entomology case from Central-Southern China. Trop Biomed. 2011;28(2):333–8.
11. Gruner SV, et al. Development of the oriental latrine Fly, Chrysomya rufifacies (Macquart) (Diptera: Stratiomyidae) to cold temperatures above and below freezing point. J Econ Entomol. 2017;110(4):1501–7.
12. Cammock JA, Nelder MP. Cool-weather activity of the forensically important blowfly, Achatocandrus ruficollis (Diptera: Calliphoridae), in northern Thailand. Parasitol Res. 2014;113(4):1353–60.
13. Amendt J, et al. Forensic entomology: applications and limitations. Forensic Sci Med Pathol. 2011;7(4):379–92.
14. Boehme P, et al. The analysis of temporal gene expression to estimate the age of forensically important blow fly pupae: results from three blind studies. Int J Legal Med. 2014;128(3):565–73.
15. Sukontat K, et al. Forensic entomology cases in Thailand: a review of cases from 2000 to 2006. Parasitol Res. 2007;101(5):1417–23.
16. Goff ML. Comparison of insect species associated with decomposing remains recovered inside dwellings and outdoors on the island of Oahu, Hawaii. J Forensic Sci. 1991;36(3):748–53.
17. Chen LS, et al. Estimation time of death by necrophagous flies life cycle. Fa Yi Xue Za Zhi. 2010;26(3):332–5.
18. Xu H, et al. Age-dependent changes in cuticular hydrocarbons of larvae in Aldrichina grahami (Diptera: Calliphoridae). Forensic Sci Int. 2014;242:236–41.
19. Aldrich JM. New two-winged flies of the family Calliphoridae from China. Proc U S Natl Mus. 1931;284:1–5.
20. Guo YD, et al. Application of Aldrichina grahami (Diptera, Calliphoridae) for forensic investigation in central South China. Rom J Leg Med. 2011;19:55–8.
21. Li X, et al. Mitochondrial DNA and STR analyses for human DNA from maggots crop contents: a forensic entomology case from Central-Southern China. Trop Biomed. 2011;28(2):333–8.
22. Gruner SV, et al. Development of the oriental latrine Fly, Chrysomya rufifacies (Macquart) (Diptera: Stratiomyidae) to cold temperatures above and below freezing point. J Econ Entomol. 2017;110(4):1501–7.
23. Cammock JA, Nelder MP. Cool-weather activity of the forensically important blowfly, Achatocandrus ruficollis (Diptera: Calliphoridae), in northern Thailand. Parasitol Res. 2014;113(4):1353–60.
24. Amendt J, et al. Forensic entomology: applications and limitations. Forensic Sci Med Pathol. 2011;7(4):379–92.
25. Boehme P, et al. The analysis of temporal gene expression to estimate the age of forensically important blow fly pupae: results from three blind studies. Int J Legal Med. 2014;128(3):565–73.
26. Sukontat K, et al. Forensic entomology cases in Thailand: a review of cases from 2000 to 2006. Parasitol Res. 2007;101(5):1417–23.
27. Goff ML. Comparison of insect species associated with decomposing remains recovered inside dwellings and outdoors on the island of Oahu, Hawaii. J Forensic Sci. 1991;36(3):748–53.
28. Chen LS, et al. Estimation time of death by necrophagous flies life cycle. Fa Yi Xue Za Zhi. 2010;26(3):332–5.
29. Xu H, et al. Age-dependent changes in cuticular hydrocarbons of larvae in Aldrichina grahami (Diptera: Calliphoridae). Forensic Sci Int. 2014;242:236–41.
52. Rebers JE, Riddiford LM. Structure and expression of a Manduca sexta larval cuticle gene homologous to Drosophila cuticle genes. J Mol Biol. 1988; 203(2):411–23.
53. Wang XR, et al. Genome-wide identification and characterization of HSP gene superfamily in whitefly (Bemisia tabaci) and expression profiling analysis under temperature stress. Insect Sci. 2019;26(1):44–57.
54. Ashburner M, Bonner JJ. The induction of gene activity in drosophila by heat shock. Cell. 1979;17(2):241–54.
55. Tower J. Heat shock proteins and Drosophila aging. Exp Gerontol. 2011; 46(5):353–62.
56. Li ZW, et al. The small heat shock protein (sHSP) genes in the silkworm, Bombyx mori, and comparative analysis with other insect sHSP genes. BMC Evol Biol. 2009;9:215.
57. Kim H, et al. A mosquito hemolymph odorant-binding protein family member specifically binds juvenile hormone. J Biol Chem. 2017;292(37):15329–39.
58. Magni PA, Dhalival SS, Dadour IR. Effect of continuous and cyclic exposure to a cold environment on the development of larvae of Lucilia sericata (Diptera: Calliphoridae) in different sized larval masses. J Med Entomol. 2016; 53(4):782–9.
59. Heaton V, Moffatt C, Simmons T. Quantifying the temperature of maggot masses and its relationship to decomposition. J Forensic Sci. 2014;59(3):676–82.
60. Kim D, Langmead B, Salzberg SL. HISAT: a fast spliced aligner with low memory requirements. Nat Methods. 2015;12(4):357–60.
61. Anders S, Pyl PT, Huber W. HTSeq--a Python framework to work with high-throughput sequencing data. Bioinformatics. 2015;31(2):166–9.
62. Anders S, Huber W. Differential expression analysis for sequence count data. Genome Biol. 2010;11(10):R106.
63. Gene Ontology C. The Gene Ontology (GO) project in 2006. Nucleic Acids Res. 2006;34(Database issue):D322–6.
64. Kanehisa M, et al. The KEGG resource for deciphering the genome. Nucleic Acids Res. 2004;32(Database issue):D277–80.
65. Ernst J, Bar-Joseph Z. STEM: a tool for the analysis of short time series gene expression data. BMC Bioinformatics. 2006;7:191.
66. Miller LD, et al. Optimal gene expression analysis by microarrays. Cancer Cell. 2002;2(5):353–61.
67. Ramoni MF, Sebastiani P, Kohane IS. Cluster analysis of gene expression dynamics. Proc Natl Acad Sci U S A. 2002;99(14):9121–6.
68. Prieto C, et al. Human gene coexpression landscape: confident network derived from tissue transcriptomic profiles. PLoS One. 2008;3(12):e3911.
69. Bagnall NH, Kotze AC. Evaluation of reference genes for real-time PCR quantification of gene expression in the Australian sheep blowfly, Lucilia cuprina. Med Vet Entomol. 2010;24(2):176–81.

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