A eukaryotic initiation factor 5C is upregulated during metamorphosis in the cotton bollworm, *Helicoverpa armigera*

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Abstract

**Background:** The orthologs of eukaryotic initiation factor 5C (eIF5C) are essential to the initiation of protein translation, and their regulation during development is not well known.

**Results:** A cDNA encoding a polypeptide of 419 amino acids containing an N-terminal leucine zipper motif and a C-terminal eIF5C domain was cloned from metamorphic larvae of *Helicoverpa armigera*. It was subsequently named *Ha-eIF5C*. Quantitative real-time PCR (QRT-PCR) revealed a high expression of the mRNA of *Ha-eIF5C* in the head-thorax, integument, midgut, and fat body during metamorphosis. Immunohistochemistry suggested that *Ha-eIF5C* was distributed into both the cytoplasm and the nucleus in the midgut, fat body and integument. *Ha-eIF5C* expression was upregulated by 20-hydroxyecdysone (20E). Furthermore, the transcription of *Ha-eIF5C* was downregulated after silencing of ecdysteroid receptor (EcR) or Ultraspiracle protein (USP) by RNAi.

**Conclusion:** These results suggested that during metamorphosis of the cotton bollworm, *Ha-eIF5C* was upregulated by 20E through the EcR and USP transcription factors.

Background

To holometabolic insects, molting is a common physiological process, whose life cycles are characterized by a series of molts. During their larval molts, the larvae progress from one instar to the next. Thereafter, pupation and eclosion ensue during their metamorphic molts. Increasing evidence indicates that some hormones and receptors may contribute to the complex developmental pathways associated with molting and metamorphosis. Many genes have been shown to be involved in molting or metamorphosis, such as the transcription factors ecdysteroid receptor (EcR), Ultraspriacle protein (USP), Hormone receptor 3 (HR3) and Broad complex [1], and the programmed cell death pathway genes [2]. Some key regulatory genes have also been identified, such as *E74B* and *E93* [3]. However, very few genes downstream of Broad complex, *E74B* and *E93* have been identified. Consequently, there is a dearth of available knowledge on the molecular mechanisms that lead to larval molt and metamorphosis. By conducting a research of the molting related genes, we may further understand the molecular mechanism of development and ecdysone regulation, and find the novel molecular targets to effectively control the pest.

Suppression subtractive hybridization (SSH) is a useful method for identifying differentially expressed genes during larval molting. Using the metamorphically committed larvae (6th-72, 96 and 120 h) as the tester and the feeding 5th instar larvae (5th-24 h) as the driver, we obtained an EST, which was similar to basic leucine zipper by BLASTX.
Some eIFs are regulators of signaling pathways, such as translation but also important in other life processes. More and more evidence suggests that translation initiation factors (eIFs) are not only essential in the initiation of protein translation but also important in other life processes. Some eIFs are regulators of signaling pathways, such as eIF4A of Drosophila melanogaster, which functions as a negative regulator of Dpp/BMP (decapentaplagic/bone morphogenetic protein) signaling that mediates SMAD (mother against dpp) degradation [6]. Eukaryotic initiation factor 6 selectively regulates Wnt signaling and β-catenin protein synthesis [7].

eIF5C is a phylogenetically conserved protein, which is said to contain an N-terminal leucine zipper motif and a C-terminal eIF5C domain. Our BLASTX results showed that homologs of eIF5C exist in various organisms, from Cryptococcus neoformans to Homo sapiens. BZAP45, the ortholog of eIF5C in humans, contributes to transcriptional control at the G1/S phase transition [8]. In Rattus norvegicus, brain development-related molecule 2 (Bdm2) is a developmentally regulated gene, which is highly expressed in fetal rat brain [9]. Wang et al. [10] showed that eIF5C was associated with the ribosome through an interaction with D. melanogaster ribosomal protein L5 (dRPL5), suggesting its possible role during protein synthesis in fruit flies. Given that there are no related functional reports to date, the information of eIF5C from other insects have been obtained from gene sequencing.

In this study, we cloned and characterized the eIF5C from the metamorphic larvae of H. armigera and designated it as Ha-eIF5C, which contains an N-terminal leucine zipper motif and a C-terminal eIF5C domain. The expression, distribution and characterization of Ha-eIF5C were studied by employing Quantitative real-time PCR (QRT-PCR), recombinant expression and immunoblotting analysis. Likewise, we also investigated the gene’s hormonal regulation and its position in the 20E signal transduction pathway.

Results
Gene cloning and sequence analysis of Ha-eIF5C
Based on the fragment of Ha-eIF5C obtained from suppression subtractive hybridization (SSH), the 5’ end fragment was obtained using specific reverse primer elf5CR and the T3 primer. The 3’ end fragment was amplified with the specific primer elf5CF and the T7 primer. The full-length elf5C of H. armegera (1675 bp) was obtained through an assemblage of overlapping nucleic acids. This included a 57 bp 5’ untranslated region (UTR), a 1260 bp open reading frame and a 340 bp untranslated region in the 3’ UTR, with a 18 bp poly A tail. The ORF encoded a 419 amino acid protein with a calculated molecular mass of 48 kDa and a predicted isoelectric point of 6.05. Moreover, there were some putative post-translational modification sites including seven protein kinase c phosphorylation sites, two tyrosine kinase phosphorylation sites, three N-myristoylation sites, five casein kinase II phosphorylation sites and one N-glycosylation site (Fig. 1).

Identification of Ha-eIF5C
The result of the BLASTX analysis suggests that Ha-eIF5C has certain similarities to various genes, including elf5C from Bombyx mori (88%), elf5C from Apis mellifera (68%), elf5C from Aedes aegypti (66%) and elf5C from D. melanogaster (63%) (Fig. 2). SMART predicted that Ha-eIF5C contains a C-terminal eIF5C domain (326-411 aa) and an N-terminal leucine zipper motif (39-60 aa).

Recombinant expression and purification of Ha-eIF5C
After IPTG induction, the recombinant GST-eIF5C was expressed in supernatant and purified by Glutathione Sepharose 4B. The deduced molecular weight of the recombinant expressed protein was 48 kDa as shown in Fig. 3. To prepare the antiserum, a gel extraction of recombinant eIF5C after cleavage of GST-eIF5C with thrombin was used.

Tissue distribution and expression patterns of Ha-eIF5C
To study the tissue distribution of Ha-eIF5C, the total RNA of the head-thorax, integument, midgut, fatbody and haemocyte were extracted from 5th 24 h (5th instar larvae 24 h after ec dysis), 5th-HCS (5th instar larvae 36 h after ec dysis, with head capsule slippage, HCS) and 6th 72 h (72 h after ec dysis, wandering 0 d, metamorphically committed larva) stage. As shown in Fig. 4, Ha-eIF5C transcript was detected at a high level in the head-thorax, integument, midgut and fat body but not in haemocytes in metamorphosis stage. QRT-PCR was utilized to analyze the expression of Ha-eIF5C in developmental midgut and fat body. The results showed that there was an obvious increase in the level of Ha-eIF5C transcript during metamorphosis. The immunoblotting revealed that the expression of Ha-eIF5C protein agreed with the mRNA transcription (Fig. 5).

Hormonal regulations on Ha-eIF5C
To examine the effect of ecdysone on Ha-eIF5C expression, 6th instar 0 h larvae (6th-0 h, with white head capsule) were injected with 20-hydroxyecdysone (20E). Compared with the control, a 5-6-fold increase in Ha-eIF5C expression was observed at 1 h and 3 h after the
Figure 1
Complete cDNA sequence and deduced amino acid sequence of Ha-eIF5C. The underlined amino acid sequences indicate predicted phosphorylation sites. Protein kinase C phosphorylation sites (2–4; 17–19; 91–93; 107–109; 193–195; 335–337; 387–389); tyrosine kinase phosphorylation sites (18–26; 345–351); casein kinase II phosphorylation sites (58–61; 100–103; 303–306; 407–410; 414–417). The putative N-glycosylation site is shaded. Predicted N-myristoylation sites are in block.
challenge. However, the expression level of Ha-eIF5C started to decline at 6 h and returned to the basal level at 12 h (Fig. 6).

In order to study whether Ha-eIF5C was upregulated downstream of the 20E-induced transcription cascade, we knocked down EcR and USP in the HaEpi cell line by RNAi. After either EcR or USP was knocked down via RNAi, the transcription of Ha-eIF5C was down regulated compared with the control and it could not be upregulated anymore by treatment with 20E (Fig. 7).

**Immunohistochemistry**

To verify the expression and localization of Ha-eIF5C, we performed an immunohistochemical analysis of the midgut (Fig. 8), fat body and integument (Fig. 9) from feeding 5th larvae (5th-24 h), molting 5th larvae (5th-HCS) and wandering 6th-96 h larvae (6th-96 h). In the 5th-HCS stage, the midgut epithelium consisted of larval polyploid cells (LPC, including columnar and goblet cells) and intestinal stem cells (ISC) (Fig. 8-K). Larval ISCs are the progenitors of the adult midgut epithelium. The larval polyploid cells moving into the lumen from the basement challenge.
membrane were replaced by proliferating and differentiating ISCs at 6th-96 h (Fig. 8-L). At this point, groups of imaginal cells began to form cell layers. Our immunohistochemical analysis shows that Ha-eIF5C was distributed into both the cytoplasm and nucleus in the midgut during the feeding 5th, molting 5th and wandering 6th-96 h stages. Relatively strong fluorescent signals were detected on the outer peripheries of the midgut epithelium from worms during 5th instar feeding and molting stage, as well as the larval polyploid cells and the imaginal cells from larvae at wandering stage. Likewise, ISCs, muscle cells and basement membrane were localized in this area. At the same time, the localization of Ha-eIF5C in the integument and fat body was detected by immunohistochemistry. During the molting 5th and wandering 6th-96 h (Metamorphic molting), a cascade of physiological processes occurred. These included the separation of the old cuticle from the underlying epidermis, followed by the secretion of a new cuticle beneath the old. Finally, a shedding of the old exoskeleton occurred (Fig. 9-E, F). It was obvious that Ha-eIF5C appears localized in both cytoplasm and nucleus in the epidermis and lipocyte.

**Discussion**

In this work, we identify a 1675 bp full-length elf5C from *H. armegera*. This includes a 1260 bp open reading frame encoding a 419 amino-acid protein with a predicted molecular mass of 48 kDa. Protein alignments showed that Ha-eIF5C and elf5C from *A. aegypti*, *A. mellifera*, *D. melanogaster* and *B. mori* are very similar. Ha-eIF5C is a phylogenetically conserved protein predicted to contain an N-terminal leucine zipper motif (39–60 aa) and a C-terminal elf5C domain (326–411 aa). This elf5C domain was first detected at the very C-termini of the yeast protein GCD6, elf-2B epsilon and two other eukaryotic translation initiation factors, elf-4 gamma and elf-5, and may likewise be involved in the interaction of elf-2B, elf-4 gamma and elf-5 with elf-2 [11]. Therefore, this elf5C domain in Ha-elf5C implies that Ha-eIF5C might also function as a novel translation initiation factor.

Leucine zipper motifs are protein-protein dimerization motifs consisting of heptad repeats of leucine residues that form a coiled-coil structure [12]. These motifs have been well described in the context of transcription factors such as c-Fos and c-Jun, where they mediate homo- and hetero-dimerization critical for the DNA binding properties of these transcription factors [13]. Proteins containing
leucine zipper motif have been reported to be related with larval growth, molting and metamorphosis in D. melanogaster [14,8]. Our work provides evidence that the expression of Ha-eIF5C, which contains a leucine zipper motif, is upregulated during metamorphosis. We speculate that through its leucine zipper domain, Ha-eIF5C may be involved in transcriptional regulation during insect development.

The expression profile of Ha-eIF5C was correlated with the metamorphic process of H. armigera. In our study, Ha-eIF5C was upregulated during metamorphosis in the head-thorax, integument, midgut and fat body. Thummel [15] reported that apoptotic and autophagic programmed cell death pathways are involved in tissue histolysis and remodeling during metamorphosis. Gorski et al. [16] confirmed that programmed cell death in the salivary glands of D. melanogaster requires active protein synthesis, even though cell death is a degradative cellular process. Gorski et al. [16] also found significant upregulation of several translation-initiation factors.

Molting and metamorphosis of larvae are very important physiological behaviors in insects, and are governed by two hormones, namely, 20E and juvenile hormone (JH) [17]. Wang et al. [18] showed that H. armigera had a similar developmental schedule as compared with Manduca sexta. 20E levels increase during the late stages of the final (wandering) instar in M. sexta larvae, before pupal ecdysis, and then decrease at the pupal ecdysis [1]. The expression of the Ha-eIF5C transcript went with the titer of 20E and was enhanced after being injected with 20E, which suggested that it was regulated by 20E in vivo. Moreover, the fact that the expression level of Ha-eIF5C in HaEpi cell line decreased after EcR or USP was knocked down demonstrated that Ha-eIF5C was upregulated by 20E via EcR or USP transcription factor.

Ha-eIF5C appears localized in both the cytoplasm and the nucleus in the midgut, integument and fat body. However, it was identified as a cytoplasmic protein in D. melanogaster [10]. In light of its function in the initiation phase of protein synthesis, eIFs were often targeted in the cyto-
plasm. However, some elfs such as elf4E are distributed in the cytoplasm and the nucleus. In the cytoplasm, elf4E acts in the rate-limiting step of translation initiation. In the nucleus, elf4E facilitates the nuclear export of a subset of mRNAs. Both of these functions contribute to elf4E’s ability to oncogenically transform cells [19]. Neither elf5C nor elf4E contain classical nuclear localization signals (NLSs) predicted by the bioinformatics method http://cubic.bioc.columbia.edu/cgi/var/nair/resonline.pl. They might act in consonance with some assistant factors that are imported into the nucleus. Dostie et al. [20] demonstrated that elf4E-Transporter (4E-T) is a nucleocytoplasmic shuttling protein that contains an elf4E-binding site, one bipartite NLS and two leucine-rich nuclear export signals, which mediate the nuclear import of elf4E via the importin αβ pathway by a piggy-back mechanism.

**Conclusion**

Ha-elf5C possibly functions as a novel translation initiation factor in protein synthesis just like elf5C of *D. melanogaster*. However, it was interesting to find that Ha-elf5C was upregulated during metamorphosis. Likewise, it was equally interesting to discover that the expression of Ha-elf5C transcript was enhanced by 20E through EcR and USP. Thus, we hypothesize that Ha-elf5C possibly functions as a regulator of cotton bollworm development, in addition to its role as a translation initiation factor.

**Methods**

**Insects**

The larvae of the cotton bollworm were maintained in this laboratory with an artificial diet described by Zhao et al. at 28 °C under a light:dark ratio of 14:10 h [21]. Moths were fed with 2% sugar water.

**Molecular cloning of Ha-elf5C gene**

A fragment of Ha-elf5C was obtained by suppression subtractive hybridization (SSH) using the metamorphically committed larvae (6th-72, 96 and 120 h) as the tester and the feeding 5th instar larvae (5th-24 h) as the driver [4]. The full-length cDNA was cloned using the cDNA library of *H. armigera* as a template. The 3’ end of the gene was amplified using a gene-specific forward primer, elf5Cf (5’-aactccagcaagggcaagatg-3’) and a T7 primer (5’-taatacgactcactataggg-3’). Similarly, the 5’ end of the cDNA was amplified by a T3 primer (5’-aattaaccctcactaaaggg-3’) and a reverse gene-specific primer elf5Cr (5’-tcttcttcggcgctctgtagc-3’).

**Sequence analysis**

Similarity analysis was performed by BLASTX [http://www.ncbi.nlm.nih.gov/]. Gene translation and prediction of the deduced protein were performed by ExPASy Proteomics Server [http://www.expasy.ch/tools/], including compute pI/Mw, TMpred, NetPhos, NetNGlyc and NetOGlyc. Signal sequence and motif prediction utilized SMART [http://smart.embl-heidelberg.de/]. Alignments were performed with ClustalW [http://www.ebi.ac.uk/clustalw/index.html] and GENDOC computer programs [http://www.psc.edu/biomed/genedoc/].

**Recombinant expression and purification**

A pair of primers (elf5CexpF: tactcagagtagcatcatgagctgtaaaaaac; elf5CexpR: tactcagctgacatctttcttgacgctgtc) were designed to amplify the sequence coding for Ha-elf5C protein (bold indicates BamHI and SalI sites, respectively). The DNA fragment was cut with BamHI and SalI, ligated into expression vector pGEX-4T-1 and transformed into competent *Escherichia coli* BL21 host cells. The recombinant expression of Ha-elf5C was induced by 0.1 mM Isopropyl β-D-1-Thiogalactopyranoside (IPTG). Thereafter, the cells were centrifuged (6000 g, 10 min), resuspended with Phosphate-Buffered Saline (PBS, 140 mM NaCl, 2.7 mM KCl, 10 mM Na2HPO4 and 1.8 mM KH2PO4, pH 7.4) containing 0.1% Triton X-100 and sonicated. The recombinant GST-elf5C was expressed in supernatant and purified by Glutathione Sepharose 4B.

**Antiserum preparation**

Rabbit polyclonal antiserum against Ha-elf5C was prepared using recombinant protein purified from *E. coli* by
SDS-PAGE. About 200 μg protein was diluted with saline and mixed with the same volume of complete Freund’s adjuvant. It was then injected hypodermically into the back of the rabbit. After three weeks, the emulsified mixture of 200 μg purified recombinant protein and incomplete Freund’s adjuvant was then subcutaneously injected into the rabbit. Two weeks later, the rabbit was given booster injections of 500 μg antigen without adjuvant and

Figure 8
Immunocytochemical localization of eIF5C in the midgut. Panels A-C are negative controls with pre-immune rabbit serum; panels D-F are midgut from feeding 5th instar larva (5th-24 h), molting 5th instar larva (5th-HCS) and 6th-96 h (wandering) larva; panels G and J, H and K, I and L are the magnified D, E, F, respectively; nuclear staining was done by DAPI (G, H, I) and the positive signals were detected by ALEXA 488 assay (J, K, L), panels A-F are overlay. LM, lumen of midgut; LPC, larval polyploidy cells; ISC, intestinal stem cell; IMC, imaginal cells; Co, clumnar cells; Go, goblet cells. Scale bar = 100 μm.
the antiserum samples were collected. The specificity of the antiserum was examined by immunoblotting and the antiserum was used in all the immunoassay experiments.

**Immunoblotting**

We followed previously reported procedures [22]. Protein extracts (100 μg) of the *H. armigera* tissues were separated using 12.5% SDS-PAGE and transferred onto a nitrocellulose membrane. Antiserum against Ha-eIF5C was diluted 1:100 in 2% non-fat milk in Tris-buffered saline (TBS, 10 mM Tris-HCl, pH 7.5 and 150 mM NaCl) and the second antibody of Horseradish Peroxidase (HRP) conjugated to goat anti-rabbit IgG was diluted 1:10,000 in the same blocking buffer (2% non-fat milk in TBS).

**Quantitative real-time PCR analysis**

Total RNA was isolated from the head-thorax, integument, midgut, fat body and haemocytes at different developmental stages using Unizol reagent according to the manufacturer's protocol (Biostar, Shanghai, China). A total of 5 μg RNA was used to reverse transcribe the first strand cDNA (First Strand cDNA Synthesis Kit, MBI Fermentas, St. Leon-Rot, Germany). It was subsequently used as a template in the PCR reactions.

SYBR green-based quantitative real-time PCR (Q-PCR) analysis was performed using PTC-200 DNA Engine thermal cycler (MJ Research) and chromo4 four-color real-time detector (Bio-Rad, America). The following primers

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**Figure 9**

**Immunocytochemical localization of eIF5C in the integument and fat body.** Panels A-C are negative controls with pre-immune rabbit serum (merged into DAPI staining outcome); panels D-I are integuments and fat body from feeding 5th instar larvae (5th-24 h, D&G), molting 5th instar larvae (5th-HCS, E&H) and 6th-96 h larvae (wandering, F&I). The nuclei were stained with DAPI (D, E, F) and the positive signals were detected using ALEXA 488 as the secondary antibody (G, H, I). Scale bar = 100 μm.
were used to amplify a specific fragment of 102 bp: elf5CF1 (5'-tattgcaatgtgtgtcctgct-3'); elf5CR1 (5'-cagccaaacgctggttgaatg-3'). A 150 bp fragment of β-actin was also amplified as control, with the primers actinF (5'-ctgttggaaggtgga-3') and actinR (5'-ctgttggaaggtgga-agggga-3'). Amplification conditions were 95°C, 2 min; 40 cycles (95°C, 15 s; 62°C, 50 s; incubated at 72°C for 2 s; plate read; incubated at 82°C for 2 s; plate read); melting curve from 60°C to 95°C, read every 0.5°C, hold 1 s. The data from real-time PCR instrumentation were then prepared for input into Microsoft Excel and analyzed using the 2-ΔΔCT method [23].

**Hormonal regulation of Ha-elf5C**

The 6th instar 0 h larvae (6th-0 h) were injected with steroid 20E (500 ng/larva). 20E was first dissolved to 10 mg/ml in dimethyl sulphoxide (DMSO) and then diluted into 0.1 mg/ml with PBS when injecting worms. Untreated controls were only injected by equivalent amounts of carrier. Total RNA of the midgut was extracted from the injected worms at different developmental periods. A comparison of the differences between the control and the challenged was done by RT-PCR with gene specific primers: elf5CF1 (5'-tatgcaatgtgtgtcctgct-3'); elf5CR (5'-ttcttctggcctgctgta-3'). The following procedure was employed: one cycle (94°C, 2 min); 26 cycles (94°C, 30 s; 53°C, 45 s; 72°C, 45 s), followed by a last cycle (72°C, 10 min). The β-actin gene was used for normalization. Each experiment was repeated three times independently. Ratios of Ha-elf5C to β-actin were calculated with Quantity One (Bio-Rad, Hercules, CA, USA).

**RNAi**

The primers of EcRRNAiF1 (5'-gcctaatgactctatagccacagtggataacagcggga-3') and EcRRNAiR1 (5'-gcctaatgactctatagccacagtggataacagcggga-3'), EcRRNAiF2 (5'-gcctaatgactctatagccacagtggataacagcggga-3') and EcRRNAiR2 (5'-gcctaatgactctatagccacagtggataacagcggga-3'), USPRNAiF1 (5'-gcctaatgactctatagccacagtggataacagcggga-3') and USPRNAiR1 (5'-gcctaatgactctatagccacagtggataacagcggga-3'); USPRNAiF2 (5'-gcctaatgactctatagccacagtggataacagcggga-3') and USPRNAiR2 (5'-gcctaatgactctatagccacagtggataacagcggga-3'); GFPRNAiF1 (5'-gcctaatgactctatagccacagtggataacagcggga-3') and GFPRNAiR1 (5'-gcctaatgactctatagccacagtggataacagcggga-3'); GFPRNAiF2 (5'-gcctaatgactctatagccacagtggataacagcggga-3') and GFPRNAiR2 (5'-gcctaatgactctatagccacagtggataacagcggga-3') were used for PCR to amplify the gene fragments. PCR products were purified using a PCR purification kit. dsRNA was synthesized using the MEGAscript™ RNAi kit (Ambion Inc, Austin, USA). The procedures of culturing HaEpi cell line and RNAi were performed according to Shao et al. [24]. The green fluorescence protein (GFP) was used as control.

**Immunohistochemistry**

The midguts and integuments adhering with fat bodies were dissected in PBS and fixed for 10 h in 4% paraformaldehyde at 4°C. The tissues were dehydrated with a graded series of ethanol. Protein digestion was performed by incubating with proteinase K (20 μg/ml) for 15 min at 37°C. Sections were blocked in 2% bovine serum albumin (BSA), incubated with a primary antibody against Ha-elf5C diluted to 1:100, and then with a goat anti-rabbit-ALEXA 488 antibody (Eugene, United States) diluted to 1:1000 in PBS with 2% BSA at room temperature for 30 min. The nuclei were stained with 4'-6-Diamidino-2-phenylindole dihydrochloride (DAPI, 1 μg/ml in water, San Jose, United States) for 10 min. Negative controls were treated in the same manner, but pre-immune rabbit serum was used in place of the antisemirum against Ha-elf5C. Fluorescence was detected with an Olympus BX51 fluorescence microscope.

**Accession numbers**

The nucleotide sequence reported in this paper has been submitted to GenBank with accession number [GenBank: EU526835].

**Authors’ contributions**

DID performed the study. JXW participated in the design and coordination of the work. XFZ conceived the study and helped to draft the final version of this manuscript. All authors read and approved the final manuscript.

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