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Molecular Cloning and Expression Profiling of CncC in Bactrocera dorsalis Hendel

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Simple Summary: We cloned the CncC gene of Bactrocera dorsalis and performed a domain analysis of the protein to clarify the expression levels of the gene in different developmental stages and tissue sites of B. dorsalis and to provide a theoretical basis for further investigating the function of CncC in regulating pesticide resistance in B. dorsalis.

Abstract: The cap ‘n’ collar isoform C (CncC) transcription factor is thought to be a regulator associated with antioxidant and detoxification genes that can enhance pest resistance by regulating the expression of detoxification enzyme genes. However, this transcription factor has not been well studied in the important agricultural pest Bactrocera dorsalis. In this study, the cDNA sequence of CncC in B. dorsalis was cloned, and the complete ORF sequence was obtained; it had a sequence length of 3378 bp, encoding a total of 1125 amino acids. Phylogenetic tree analysis showed that B. dorsalis CncC belonged to the CNC family and that its amino acid sequence showed the closest relationship with B. tryoni. The conserved structural region of BdCncC was analyzed and was found to include a conserved bZIP superfamily structural domain. Spatiotemporal expression analysis revealed that BdCncC was most highly expressed in the adult Malpighian tubules, followed by the antennae, foregut, and midgut, and then the brain, hemolymph, hindgut, and fat body. BdCncC was expressed at every developmental stage, and the highest expression was found in mature males. This study provides a theoretical basis for an in-depth investigation of the function of BdCncC in regulating pesticide resistance in B. dorsalis.

Keywords: Bactrocera dorsalis; BdCncC; gene cloning; expression profiling

1. Introduction

Transcription factors (TFs) are members of the family of regulatory proteins. The sequences of TFs first contain DNA structural domains that recognize the corresponding cis-acting elements in the promoter regions of bound genes, followed by transcriptional regulatory structural domains, nuclear localization signal sequences, and oligomerization sites [1]. Due to the highly conserved structural domains of TF DNA sequences, they are classified into families such as bZIP, NAC, MYB, MYC, and WARK TFs [2,3]. The cap ‘n’ collar isoform C (CncC) TF is a member of the bZIP TF family and was first identified in Drosophila. It was then found to be directly homologous to the vertebrate protein nuclear factor erythroid 2-like Nfe2l2 (Nrf2) and the nematode protein skinhead family member 1 (SKN-1), which plays a key role in regulating cellular defense against oxidative stressors or electrophilic xenobiotics [4–6]. When oxidative stress is generated by exogenous stimuli, the conformation of kelch-like ECH-associated protein 1 (keap1) changes, releasing CncC into the nucleus and promoting the functional expression of CncC [7]. CncC has seven functional structural domains, where the main domain is Neh1, which contains a leucine bZIP structural domain for the recognition and binding of antioxidant response elements (AREs) [8]. Pests regulate the expression of downstream detoxification enzyme genes.
through the CncC/keap1-ARE signaling pathway, thereby increasing their resistance to pesticides. A study on the potato beetle (*Leptinotarsa decemlineata*) showed that CncC was required for the expression of 79% of P450 genes induced by imidacloprid, including genes encoding detoxification enzymes (P450, glutathione S-transferases) [9]. In *Tetranychus cinnabarinus*, CncC was found to affect the sensitivity of cinnabar mites to benzothiurian by regulating the expression of P450 genes [10]. Misra (2011) used resistant Drosophila species with significantly upregulated expression levels of CncC and P450 detoxification enzyme family genes after pesticide treatment. All genes related to detoxification showed a significant decrease after the RNAi knockdown of CncC, suggesting that CncC induces the expression of detoxification enzyme genes in Drosophila, thereby regulating pesticide resistance in Drosophila [6].

The oriental fruit fly, *Bactrocera dorsalis* (Hendel) (Diptera: Tephritidae), is a polyphagous pest that is considered highly invasive in horticulture. The oriental fruit fly infests hundreds of fruits and vegetables, including apricot, avocado, banana, citrus, coffee, etc. For the past 20 years, this fly has been introduced to and spread throughout sub-Saharan Africa [11,12]. Its existence has caused significant financial losses in orchard crops [13]. Furthermore, the overuse of insecticides to control *B. dorsalis* has led to the development of insecticide resistance [13]. Thus, the oriental fruit fly is an important quarantine pest. Chemical-based control strategies are still the main management method for oriental fruit flies, and resistance to these chemicals has become increasingly prominent in recent years, representing a major bottleneck in the sustainable control of infestation [14]. In this study, we obtained the ORF sequence of the *B. dorsalis* *BdCncC* gene by cloning, translated it into the corresponding amino acid sequence, and performed domain analysis to clarify its expression levels in different developmental stages and tissue sites of *B. dorsalis*, with the aim of providing a theoretical basis for an in-depth investigation of the function of *BdCncC* in the resistance of *B. dorsalis*.

2. Materials and Methods

2.1. Insects

*B. dorsalis* was reared at 27 °C with 75% relative humidity under a 14:10 h light:dark photoperiod. Adults were fed an artificial diet consisting of yeast extract and dry sugar mixed at a 1:1 ratio (w/w) and housed in transparent plastic cages.

2.2. RNA Isolation, Reverse Transcription, and *BdCncC* Cloning

Total RNA was isolated from adults using TRizol® reagent (Invitrogen, Carlsbad, CA, USA). RNA quality was checked by 1% agarose gel electrophoresis. Reverse transcription was performed using the PrimeScript™ II 1st strand cDNA Synthesis Kit (Takara, Dalian, China). After reverse transcription, the synthesized cDNAs were stored at −20 °C for future use. The reference sequence of the *BdCncC* cDNA was acquired from NCBI (GenBank number, KJ957012). Primers were designed (Table 1) using Primer Premier 5.0 (Premier Biosoft International, Palo Alto, CA, USA). PCR was performed using Q5 High-Fidelity DNA Polymerase (New England Biolabs, Ipswich, MA, USA) according to the manufacturer’s instructions. The purified PCR product was cloned into the pEASY-Blunt Zero Cloning Vector (TransGen, Beijing, China) following the manufacturer’s instructions before being sequenced. Positive clones were sent for sequencing by Sangon Biotech (Shanghai, China).

2.3. Analysis of the Protein Sequence and Biological Information of *BdCncC*

The obtained full length of *CncC* was compared with the Blast (https://blast.ncbi.nlm.nih.gov/Blast.cgi, accessed on 6 June 2022) in NCBI. The cloned *BdCncC* sequences were analyzed using DNAMAN 9.0. ORF prediction was performed using NCBI ORF Finder (http://www.ncbi.nlm.nih.gov/orffinder, accessed on 7 June 2022). The analysis of conserved structural regions was performed using NCBI Conserved Domains (https://www.ncbi.nlm.nih.gov/cdd, accessed on 7 June 2022). The analysis of protein functional structural domains was performed using SMART (http://smart.embl-heidelberg.de/,
accessed on 7 June 2022). The molecular weight, isoelectric point, instability index (II), and aliphatic index were predicted with ProtParam (https://web.expasy.org/protparam/, accessed on 10 June 2022). Hydrophilicity prediction was conducted using ProtScale (https://web.expasy.org/protscal/, accessed on 10 June 2022). Signal peptide prediction was carried out using SignalP (http://www.cbs.dtu.dk/services/SignalP/, accessed on 10 June 2022). Transmembrane prediction was performed using TMHMM (http://www.cbs.dtu.dk/services/TMHMM-2.0/, accessed on 10 June 2022). Subcellular localization analysis was performed using PSORT II (https://www.genscript.com/psort.html, accessed on 11 June 2022). The prediction of 3D structures was conducted using SWISS-MODEL (https://swissmodel.expasy.org, accessed on 11 June 2022).

Table 1. Primers used for BdCncC in the study.

| Experiment          | Primer Names | Sequences (5′-3′)          |
|---------------------|--------------|-----------------------------|
| Full-length         | CncC-F       | ATGGGTTTGCCCGCTTCGGA         |
|                     | CncC-R       | TCAATCTTTCTGATGCGTTT         |
| RT–qPCR             | CncC-q-F     | CGTGGCAATAGTGGCTT           |
|                     | CncC-q-R     | TATGGAATCTCGGATAGTGAG        |
| Reference genes-1   | a-Tubulin-F  | GGCAATTGATGGTATAACG         |
|                     | a-Tubulin-R  | GGGCAACCAAGTTACGCGAG        |
| Reference genes-2   | RPL-F        | CCCGCTCATATGCTGCCAACT       |
|                     | RPL-R        | GCGCGCTCAACAATTTCTT         |

2.4. Phylogenetic Analysis and Identification

The CncC and Nrf2 protein sequences were obtained from the NCBI web server (https://www.ncbi.nlm.nih.gov/, accessed on 20 June 2022) and aligned with the sequences generated in the present study using the Clustal website (https://www.ebi.ac.uk/Tools/msa/clustalo/, accessed on 20 June 2022). To infer evolutionary relationships, the neighbor-joining method was used to construct a phylogenetic tree in MEGA11.0.1 software with 1000 bootstrap replicates.

2.5. Reverse-Transcription Quantitative PCR Analysis

RNA was extracted from different tissues, including the brain, antennae, hemolymph, foregut, midgut, hindgut, Malpighian tubules, and fat body. TRIzol® reagent (Invitrogen, Carlsbad, CA, USA) was used for RNA isolation. The extracted RNA was purified using the phenol/chloroform method and dissolved in RNase-free water. The purity of the extracted RNA was assessed spectrophotometrically by measuring the OD260/280 ratio, where an OD260/280 of 1.8–2.0 indicated good-quality RNA. RNA integrity was evaluated via electrophoresis on a formaldehyde agarose gel. The RNA (1 µg) was then reverse transcribed to cDNA using the PrimeScript™ RT reagent Kit with gDNA Eraser (Takara, Otsu, Japan) according to the manufacturer’s instructions. Biosynthesized cDNA was used as a template in RT–qPCR conducted on a C1000 Touch thermal cycler (Bio-Rad Laboratories, CA, USA) with TB Green Premix Ex Taq II (Tli RNase H Plus) (Takara Bio, Otsu, Japan). The thermal cycling conditions were as follows: 95 °C for 30 s, 40 cycles at 95 °C for 5 s, and 60 °C for 34 s. RT–qPCR was conducted with three technical and five biological replications. α-Tubulin and RPL were used as reference genes for gene expression analysis in B. dorsalis due to their expression stability.

2.6. Statistical Analysis

CncC gene transcript levels were quantified using the $2^{-\Delta\Delta CT}$ method [15]. Experimental data were analyzed and plotted using SPSS 26.0 and GraphPad Prism 9.0. BdCncC expression pattern analysis was performed by one-way analysis of variance (ANOVA) with Duncan’s multiple range test. A probability value of $p < 0.05$ was considered statistically significant.
3. Results

3.1. Molecular Cloning and Sequencing of BdCncC

PCR amplification of BdCncC was carried out using the specific primers designed using the whole insect cDNA of *B. dorsalis* as the template, and electrophoresis showed that the amplified target fragment was consistent with the length of the target gene (Figure 1B). The results of NCBI Blast on the cloned sequence showed that it was the CncC gene of *B. dorsalis*. The analysis of the sequencing results using DNAMAN9.0 showed that the full-length *B. dorsalis* BdCncC gene was 3494 bp and encoded 1161 amino acids. The ORF of BdCncC was predicted to be 3378 bp long, encoding 1125 amino acids, according to NCBI ORF Finder, and the amino acid sequence was obtained by translation. The obtained ORF amino acid sequence was analyzed using NCBI Conserved Domains for the conserved structural region of BdCncC; the protein contains a conserved bZIP superfamily structural domain in which the conserved structural domain is located between amino acids 910–977, with an E-value of $1.09 \times 10^{-32}$. The conserved structural domain includes 14 DNA-binding sites and 19 dimer interfaces (Figure 1A,C). SMART was used to analyze the functional structural domains of the BdCncC protein, which includes two conserved functional domains: basic region and leucine zipper (BRLZ) domains (Figure 1D).

**Figure 1.** Functional domain prediction based on the BdCncC amino acid sequence. (A, D) Schematic illustration of BdCncC; (B) electrophoresis of BdCncC PCR products; M: DNA Marker DL5000; C: control group; 1: PCR product; (C) amino acid sequence structure of BdCncC, where red letters represent the conserved regions of the BdCncC protein, including the basic region domain and leucine zipper—yellow highlighting indicates the DNA-binding site, and green highlighting indicates the dimer interface site.

3.2. Analysis of Biological Information of BdCncC

The molecular weight of the BdCncC protein was predicted by ProtParam to be 121.42 kDa, indicating that it is a large protein; the isoelectric point (isoelectric point) was 5.17, indicating that it is an acidic protein; the instability index (II) was 47.11 (>40), indicating that it is unstable; and the aliphatic index was 61.15 (<90), indicating that it is a water-soluble protein. The highest Score value was 2.011 at amino acid position 886, and the lowest Score value was $-3.611$ at amino acids 1104 and 1105, whereas the grand average of hydropathicity (GRAVY) was $-0.784$, indicating that it is a hydrophilic protein (Figure 2A). PSORT II was used to predict the subcellular localization of BdCncC in nuclear,
cytoskeletal, cytoplasmic, and mitochondrial fractions (Table 2). By applying PSORT II, the tertiary structure of *BdCncC* was predicted using TF MaB (PDB chain ID: 2wty:1A) as a template, and the Ramachandran favored value was 98.93%, indicating that the predicted 3D structure was plausible (Figure 2B,C). Signal peptide prediction using SignalP revealed no signal peptide in *BdCncC*, indicating that it cannot be secreted extracellularly (Figure 2D). Transmembrane prediction of the *BdCncC* protein using TMHMM predicted a transmembrane helix number of zero, indicating that it is not a transmembrane protein (Figure 2E).
3.3. Sequence and Phylogenetic Tree of BdCncC

The evolutionary tree constructed by comparing the sequences of insect CncC and Nrf2 from other species showed that CncC and Nrf2 were sister groups (Figure 3A). The phylogenetic tree of the CncC protein sequences of dipteran insects showed that CncC proteins were relatively conserved, and the amino acid sequence of B. dorsalis BdCncC presented the closest affinity to the amino acid sequence of B. tryoni cap ‘n’ collar (Figure 3B). The evolutionary tree results suggest that BdCncC was a member of the CNC family. And the cladograms and the high similarity of the conserved regions in the multiple sequence alignment results proved the conservativeness of CncC (Figures S1 and S2).

Figure 3. Phylogenetic tree of BdCncC generated with the neighbor-joining method. (A) Phylogenetic tree of CncC and Nrf2; (B) phylogenetic tree of dipteran CncC sequences. The numbers along the branches indicate bootstrap support from 1000 replicates; (▲) represent CncC in B. dorsalis.

Table 2. Subcellular localization prediction of BdCncC.

| Predicted Location | BdCncC |
|--------------------|--------|
| Nuclear            | 87%    |
| Cytoskeletal       | 4.3%   |
| Cytoplasmic        | 4.3%   |
| Mitochondrial      | 4.3%   |

3.4. Temporal and Spatial Expression Patterns of BdCncC

The results of the analysis of BdCncC expression levels in different tissues and developmental stages of B. dorsalis using qRT-PCR showed that BdCncC expression was highest in the adult Malpighian tubule, followed by the antennae, foregut, and midgut, and then the brain, hemolymph, hindgut, and fat body (Figure 4A, F(7, 32) = 25.016, p < 0.001). Among different developmental stages, BdCncC was highly expressed mainly in adults, 7-day larvae, and eggs, and the highest expression was found in mature males. The expression levels were relatively low in 1-day and 4-day larvae and pupae (Figure 4B, F(10, 42) = 9.292, p < 0.001).
3.4. Temporal and Spatial Expression Patterns of BdCncC

The results of the analysis of BdCncC expression levels in different tissues and developmental stages of *B. dorsalis* using qRT–PCR showed that BdCncC expression was highest in the adult Malpighian tube, followed by the antennae, foregut, and midgut, and then the brain, hemolymph, hindgut, and fat body (Figure 4A, \( F(7, 32) = 25.016, p < 0.001 \)). Among different developmental stages, BdCncC was highly expressed mainly in adults, 7-day larvae, and eggs, and the highest expression was found in mature males. The expression levels were relatively low in 1-day and 4-day larvae and pupae (Figure 4B, \( F(10, 42) = 9.292, p < 0.001 \)).

![Figure 4](image)

**Figure 4.** Expression profiles of the *BdCncC* gene in different tissues and developmental stages of *B. dorsalis*. (A) Expression profiles of the *BdCncC* gene in different tissues, and different colors indicate different tissue; (B) Expression profiles of the *BdCncC* gene in different developmental stages, and different colors indicate different developmental stages. Means and SEs from five biological replicates are shown. The different letters above the bars indicate significant differences in gene expression levels among treatments (\( p < 0.001 \), one-way ANOVA test).

4. Discussion

The CncC TF protein family was originally studied in vertebrates, invertebrates, and postnatal animals. This TF is thought to be a regulator associated with antioxidant and detoxification genes [5]. In this study, the cDNA sequence of the *B. dorsalis* BdCncC TF was cloned, and the complete ORF sequence was obtained, showing a sequence length of 3378 bp and encoding a total of 1125 amino acids. The predicted protein molecular weight was 121.42 kDa. The conserved structural region of BdCncC was also analyzed, and it was found to include a conserved bZIP superfamily structural domain. Phylogenetic tree analysis showed that the amino acid sequences of *B. dorsalis* BdCncC were very similar to those of *B. tryoni*, indicating that they are more evolutionarily homologous to each other than other sequences in this family. Spatiotemporal expression analysis showed that BdCncC was expressed at every developmental stage of *B. dorsalis* and that its expression was highest in adults. Since the field control of *B. dorsalis* is usually performed by spraying insecticides on adults; the high expression of BdCncC in adults may contribute to the metabolism of insecticides or phytochemicals in *B. dorsalis* and, thus, enhance resistance. The tissue distribution of a gene is usually related to its function. BdCncC expression was highest in the adult Malpighian tube. The Malpighian tube is a multifunctional organ capable of playing roles in compound excretion, insect metabolism, and accumulated metal of harmful compound detoxification [16]. Higher detoxification enzyme activity is also associated with the malleolus [17,18]. Therefore, the overexpression of BdCncC in the detoxification organ of *B. dorsalis* may play an important role in insecticide detoxification, and further
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studies can be conducted to reveal the function of CncC in different developmental stages and tissues of insects.

In summary, we identified and analyzed the structural features of BdCncC in B. dorsalis. Numerous studies have shown that insects increase their resistance to drugs by regulating the expression of detoxification enzyme genes [19–21]. A study showed a significant increase in transcript levels of BmCncC and enhanced activity of GST and CYP450 after 24 h of octreotide treatment. [22]. In Drosophila, the expression of the GSTd7, GSTd2, CYP6A2, and CYP6A8 genes is significantly downregulated, and sensitivity to malathion is increased after silencing CncC [6]. The RNAi knockdown of CncC decreases CYP6DA2 gene expression and increases the sensitivity of cotton aphids to cotton cotyledon phenol [23]. In Spodoptera litura, reactive oxygen species (ROS) activate the CncC TF to regulate the expression of the cytochrome P450 gene CYP6AB12, which mediates resistance to λ-cyhalothrin [24]. Therefore, it is hypothesized that CncC may mediate resistance in B. dorsalis by regulating the expression of multiple detoxification genes. Studying the functions of detoxification genes and the binding sites of TFs is important for understanding how BdCncC regulates the expression of resistance genes in B. dorsalis, and further studies on this topic are needed.

Supplementary Materials: The following supporting information can be downloaded at: https://www.mdpi.com/article/10.3390/insects13090785/s1, Figure S1: Multiple sequence alignment of conserved regions of insect CncC and Nrf2 from other species; Table S1: Sequences of insect CncC and Nrf2 from other species for the phylogenetic analysis; Figure S2: Multiple sequence alignment of conserved regions of CncC of diptera; Table S2: Sequences of diptera CncC for the phylogenetic analysis.

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Institutional Review Board Statement: The B. dorsalis fruit flies used in this study are fruit pests in China. All experiments were in line with Chinese law and ethics guidelines.

Data Availability Statement: The data presented in this study are available upon request from the corresponding author.

Conflicts of Interest: The authors declare no conflict of interest.

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