Corazonin Signaling Is Required in the Male for Sperm Transfer in the Oriental Fruit Fly Bactrocera dorsalis

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Corazonin (Crz) is a widely distributed neuropeptide (or neurohormone) in insects with diverse physiological functions. The present study aimed to reveal the functions of Crz and its receptor (CrzR) in the regulation of sexual behavior and fertility in male Bactrocera dorsalis. Tissue-specific expression analyses showed that the BdCrz transcript was most abundant in the central nervous system (CNS), and the BdCrzR transcript was most abundant in both the fat body and CNS. Immunochemical localization confirmed that three pairs of Crz-immunoreactive neurons are located in the dorsolateral protocerebrum region of male adult brain. Importantly, RNAi-mediated Crz knockdown lengthened mating duration in males, and knockdown of Crz or CrzR strongly decreased male fertility in the following 3 days, while the courtship behavior and mating efficiency were not affected. The reduced number of sperm in the reproductive organs of mated females indicated that Crz knockdown in males reduced sperm transfer. The findings of this study indicate that Crz contributes to the reproductive physiology of the oriental fruit fly B. dorsalis by regulating sperm transfer in male adults.

Keywords: corazonin, male adult, reproduction, sperm transfer, mating duration

INTRODUCTION

The oriental fruit fly, Bactrocera dorsalis (Hendel) (Diptera: Tephritidae), is a widely distributed quarantine polyphagous pest in tropical and subtropical areas of the world. Due to the increased invasiveness and potential influence on food production, B. dorsalis has been one of the most destructive pest of fruit industries in many countries (Clarke et al., 2005; Stephens et al., 2007). Because of the rapid development of insecticide resistance and particular way of damage, this insect became resurgence and difficult to control (Chen et al., 2013; Huang et al., 2015), and therefore, new insecticide targets are urgently needed. Insect neuropeptide signaling systems have been regarded as a novel type of pest management (Allstein, 2001; Boonen et al., 2009; Allstein and Nässel, 2010).

Insect neuropeptides are essential physiological regulators for various biological events such as development, metabolism, and behavior. They also play a key role in the regulation of sexual behavior and reproduction (Allstein and Nässel, 2010). The peptide corazonin (Crz) was first identified as a cardioactive peptide in the American cockroach Periplaneta americana (Veenstra, 1989). Subsequent identification of the Crz in other insect species revealed two major members of the Crz family: [Arg7]-Crz and [His7]-Crz (Lee et al., 2008). In our previous study, we have identified and characterized the [His7, Ser11]-Crz isoform (pQTFQYSHGWTSamide) in B. dorsalis
(Hou et al., 2017b), which is different from the [Arg7]-Crz (pQTFQYSRGWTNamide) as reported in several Drosophila species (Veenstra, 1994; Choi et al., 2005), and also different isoforms in other insects have been reported so far (Predel et al., 2007; Hou et al., 2017b). The Crz receptor (CrzR) of B. dorsalis is a typical Class A GPCR member with seven transmembrane domains, showing a high sequence similarity to the Crz receptor in Drosophila (DmCrzR) (Hou et al., 2017b).

Molecular and histochemical studies have demonstrated that the distribution of Crz in neurosecretory neurons of the central nervous system (CNS) seems to be conserved among different insect species (Predel et al., 2007; Lee et al., 2008). Various functions have been found for physiological and behavioral actions in different experimental systems adopted in diverse insect groups. For example, a significant role for Crz in modulating myostimulatory activities on the heart muscle of the cockroach P. americana (Veenstra, 1989) and the hyperneural muscle of stick insect Carausius morosus (Predel et al., 1999), while the myostimulatory activity of Crz was not found in other insects (Predel et al., 1994; Sláma et al., 2006; Hillyer et al., 2012). Physiological studies with the moth Manduca sexta suggested that Crz signaling initiated the ec dys behavior by inducing the release of pre-ec dysis and ec dysis-triggering hormones from the Inka cells (Kim et al., 2004). In D. melangaster, ablation of Crz neurons affected the metabolism and lead to a differential lifespan under stress, which showed a strong dependence on sex (Zhao et al., 2010). More importantly, silencing of Crz neurons affected the mating behavior and blocked the transfer of sperm and seminal fluid in D. melanogaster (Taylor et al., 2012). Peptide injection to the early transitioning ants demonstrated that Crz stimulates hunting and inhibits dueling in the ant Harpegnathos saltator (Gospocic et al., 2017). Crz and CrzR have also been identified using a heterologous functional assay in Aedes aegypti (Oryan et al., 2018). In our earlier study, the silencing of CrzR by RNAi effectively disrupted the larval-pupal transition and led to a delay in pupariation in B. dorsalis (Hou et al., 2017b). Crz-type signaling is also present in other invertebrates, including other protostomes (annelids and mollusks) and deuterostomes (cephalochordates, hemichordates, and echinoderms) (Zandawala et al., 2017). For example, Kavanaugh and Tsai (2016) have characterized a Crz-type signaling system in Aplysia and interestingly the Crz-type receptor is expressed in reproductive organs (small hermaphroditic duct, and ovotestis). A Crz-type signaling system has also been characterized in the starfish Asterias rubens, which causes contraction of cardiac stomach, apical muscle, and tube foot preparations (Tian et al., 2016, 2017). Another recent publication showed, however, that Crz had no effect on heart activity, blood glucose levels, lipid mobilization, or pigment distribution in chromatophores in the crab Carcinus maenas (Alexander et al., 2017).

To gain further insight into the function of Crz in B. dorsalis, in the present study, we have extended our previous work with the larval stages (Hou et al., 2017b) now with the adult stage to investigate the sexual behavior regulation of the Crz/CrzR signaling pathway by RNAi. Our data may contribute to explore its potential as a novel insecticide target.

**MATERIALS AND METHODS**

**Test Insects**

Bactrocera dorsalis was originally collected from Dongguan, Guangdong province, China, and reared as described previously (Chen et al., 2013). Shortly, the flies were reared at 27 ± 1°C, 70 ± 5% relative humidity with a photoperiod of 14 h light: 10 h dark. Shortly after the adult emergence, virgin flies sexed and kept isolated on a standard medium until they were used for experiments. In this study, we used the 9-day-old virgin flies in most experiments.

**Sample Preparation, RNA Isolation, and cDNA Synthesis**

Tissues from the CNS, fat body, midgut, hindgut, Malpighian tubules, and testes were excised from 9-day-old adults to determine the tissue-specific expression pattern. At least 15 individuals were dissected as one sample for tissue-specific analysis with four replications. The adults were chilled on ice for 30 min and dissected under a stereomicroscope (Olympus SZX12, Tokyo, Japan). The samples were isolated on ice, placed in a 2.0 mL of diethyl pyrocarbonate (DEPC)-treated centrifuge tube containing RNA storage reagent (Tiangen, Beijing, China), and immediately frozen in liquid nitrogen and stored at −80°C.

Total RNA was extracted from each sample using the TRizol reagent (Invitrogen, Carlsbad, CA, United States) and treated with DNase I (Promega, Madison, WI, United States) to prevent potential genomic DNA contamination. First-strand cDNA was synthesized using the GoScript Reverse Transcription System (Promega) according to the manufacturer’s instructions.

**Quantitative Real-Time PCR (qPCR)**

Primers used in this study (Table 1) were synthesized by Invitrogen (Shanghai, China). qPCR was performed on an ABI 7500 Real-Time PCR System (Applied Biosystems, Foster City, CA, United States). The reaction mixtures: cDNA template (0.5 μL), of each primer (0.5 μL, 0.2 mM), Novostar-SYBR Supermix (5 μL, Novoprotein, Shanghai, China) and of ddH₂O (3.5 μL). Reaction conditions: an initial denaturation at 95°C for 2 min, followed by 40 cycles of 95°C for 15 s, 60°C for 30 s. At the end of each qPCR, a melting curve analysis from 60 to 95°C was generated to rule out the possibility of primer-dimer formation. The data were normalized to the stable reference gene α-tubulin (GenBank accession no. GU269902) based on previous evaluations (Shen et al., 2010). The relative expression levels were calculated using the 2^−ΔΔCt method (Livak and Schmittgen, 2001).

**Immunohistochemistry**

For immunohistochemical detection of Crz in B. dorsalis, we used the polyclonal Crz antibody (against D. melanogaster Crz, pQTFQYSRGWTNamide) that was a gift from Dr. Jan Veenstra.
(Université de Bordeaux, France; Veenstra and Davis, 1993) with use of our protocol as previously published (Hou et al., 2017b). In brief, the brains from 9-day old males were dissected in chilled PBS (pH 7.4) and then tissues were fixed overnight at 4°C in fresh 4% paraformaldehyde in PBS. After washing for 3×5 min in PBS with 0.5% Triton X-100 (PBST), the tissues were incubated with primary antibody (1:1000 diluted in PBST) for 2 days at 4°C and then washed in 3×5 min in PBST. Tissues were incubated overnight at 4°C in Alexa 488-conjugated goat anti-rabbit IgG antibody (1:1000 in PBST). The samples were washed in 2×5 min in PBST. Then, samples were mounted on a clean slide with 100% glycerol. Images were captured in a confocal microscope Zeiss LSM780 (Zeiss, Jena, Germany).

RNAi Bioassay
Primers (Table 1) with T7 promoter sequences were used to amplify a fragment of BdCrz, BdCrzR, or GFP (CA58789) for the double-stranded RNA (dsRNA) synthesis. The dsRNA was transcribed by Transcript Aid T7 High Yield Transcription Kit (Thermo Scientific, Lithuania) following the manufacturer’s protocol. The dsRNA was quantified using a NanoPhotometer (Implen, Germany) and the integrity was confirmed by a 1% agarose gel electrophoresis.

For the RNAi experiments, 9-day-old adult virgin flies were used. Then 1.2 µg (in 300 nL) of dsRNA (Crz-dsRNA, CrzR-dsRNA or GFP-dsRNA) was injected into the body cavity of the 9-day-old males directly between the second and third abdominal segments using a Nanoject II Auto-Nanoliter Injector (Drummond Scientific, Broomall, PA, United States). Injected flies were kept on an artificial diet under the conditions described above. The RNAi efficiencies of Crz and CrzR were measured by qPCR as described above, and analyzed by an independent Student’s t-test.

### Table 1 | Primer sequences used in this study.

| Target        | Direction | Sequence 5’ to 3’         |
|---------------|-----------|---------------------------|
| BdCrz (qPCR)  | Forward   | TTGCCGAAATGCTCCAACAAA     |
|               | Reverse   | CATAATCGTGGTCCTCGGCGC     |
| BdCrzR (qPCR) | Forward   | TGCTCAOOGTCACCTACATT      |
|               | Reverse   | TCACAAAAGACAGTCGACAGC     |
| α-Tubulin (qPCR) | Forward     | CGCATCTGAGTGGATAACG       |
|               | Reverse   | GGCCCAACAGATGCTGAGA       |
| BdCrz (dsRNA) | Forward   | taataagactactataggTCTTCCTCATTTACGTGTCTCT    |
|               | Reverse   | taataagactactataggGATTTACCATTTACGTGTCTCTCAG |
| BdCrzR (dsRNA) | Forward     | taataagactactataggTTACAAAATCGAC CGCAACGC |
|               | Reverse   | taataagactactataggGGCTGGTGAATTGC ATCGCC |
| GFP (dsRNA)   | Forward   | taataagactactataggGAGTTCTTGTTGAA TTAGATG |
|               | Reverse   | taataagactactataggTTTGTTGTTCTCCTAGGATG |

Behavioral Assays
The mating behavior assays were performed as in our previous report (Hou et al., 2017a). All assays were performed in the mating arenas (a small round inverted cup with 6 cm in diameter and 10 cm high) under standard laboratory conditions. Virgin females and native males were collected separately on the day of adult emergence. Subsequently, 9-day-old male flies were injected with dsRNA. Twenty-four hours after the dsRNA injection, the treated males were crossed to the normal females individually and reared in the mating arenas. Each pair of flies was transferred to the mating chamber at the beginning of the light photoperiod to ensure adequate circadian time for acclimation. Mating behavior was observed every 30 min over a 10 h-dark photoperiod or until the male dismounted the female following mating. For the behavior assay, 30 pairs of flies were used per treatment with three biological replicates. Courtship behavioral sequence was based on the previous reports (Poramarcom, 1988), which is including the steps from wing vibration to attempting copulation. The percentage of males displaying courtship behavior, the percentage of mating pairs and the copulation duration were recorded. Furthermore, the numbers of eggs laid by successfully mated flies were recorded at 24 h-intervals for 3 days following mating.

DAPI Straining
After the completion of mating behavior, the reproductive organs (spermatheca) of mated females were dissected in PBS. Then the tissues were gently grinded and stained with DAPI (4’,6-diamidino-2-phenylindole, 2 µg/ml) (Sigma, St. Louis, MO, United States) for 15 min. A fluorescence microscope Zeiss LSM780 (Zeiss, Jena, Germany) was used for images capture and cells counting.

Statistical Analysis
Data on copulation durations were analyzed by a non-parametric Mann–Whitney U test, and these on mating and fertility by an independent Student’s t-test or one-way ANOVA, both in SPSS 16.0 software (SPSS, Chicago, IL, United States, 2008). Results are presented as mean ± SE (standard error), and statistical significance is assumed for p < 0.05.

RESULTS
Tissue-Specific Transcript Profiling
We analyzed the spatial distribution of BdCrz and BdCrzR in different organs of the adult by qPCR. As shown in Figure 1, BdCrz was mainly expressed in the CNS, and no detectable qPCR products were determined in the other tissues tested (as the Ct values with the other tissues were >35) (Figure 1A). The transcript of BdCrzR was highly accumulated in the fat body, and at medium levels in the CNS. Low-level expressions were observed in testis, malpighian tubules, and gut (Figure 1B).

Localization of BdCrz in the Brain
In an attempt to gain insight into the localization of BdCrz in the brain of male adults of B. dorsalis, whole mount
immunohistochemistry was carried out with a rabbit antibody against *D. melanogaster* mature Crz. Figure 2 shows a representative image of 20 different samples. As the result indicated, three pairs of adult Crz-immunoreactive neurons were observed in a cluster in the dorsolateral region of the protocerebrum (DLP).

**Knockdown of BdCrz or BdCrzR by RNAi**

To better understand the function of Crz signal pathway on mating behavior, we injected specific dsRNA against *BdCrz* and *BdCrzR* in 9-day-old adult flies. The results showed that the knockdown efficiencies were significant for *BdCrz* and *BdCrzR* after the dsRNA injection. Compared with the control groups injected with dsGFP, the gene silencing efficacies for *BdCrz* mRNA levels were about 60% at 24 and 48 h post-injection of dsCrz, and these for *BdCrzR* with dsCrzR injection reached 50 and 57% at 24 and 48 h post-injection, respectively (Figure 3).

**Effects of BdCrz or BdCrzR Knockdown on Mating Behavior**

The effects of RNAi-mediated *BdCrz* and *BdCrzR* knockdown on mating behavior were analyzed. Consequently, the courtship behavior and mating frequency of the Crz- or CrzR-silenced males were normal as compared with the control (Figures 4A,B). By contrast, the silencing of Crz resulted in a dramatically increased copulation duration in males. In the control group, the copulation duration of dsGFP-treated males typically lasted 7 h, while the copulation duration of dsCrz-treated males increased with more than 40% to 10 h. Nevertheless, CrzR silencing had no effect on the copulation duration in males (Figure 4C). In addition, it was striking that the silencing of either Crz or CrzR caused a similar defect in fecundity: the egg-production in the respective partner female was significantly less than that in the control group (i.e., females that had mated with dsGFP-injected males) in the followed 3 days (Figures 4D,E). Indeed, the numbers of eggs per female after mating with a male injected with dsCrz or dsCrzR were reduced by 80–90% compared to the control.

**Effects of BdCrz or BdCrzR Knockdown on Sperm Transfer**

To clarify the effects of Crz signaling pathway on sperm transfer in *B. dorsalis*, the reproductive organs of females, that had
copulated with dsGFP-, dsCrz-, or dsCrzR-injected males, were dissected after the mating was finished and then we did a DAPI staining of sperm cells. The results showed that BdCrz or BdCrzR knockdown caused a marked lower number of sperm cells in the spermatheca compared to the control group (Figure 5).

**DISCUSSION**

Crz and its receptor are widespread and highly conserved in insects, but no universal function has been described so far, while its function in other basal taxa is unknown also. In the present study, we described the tissue-specific expression patterns of BdCrz and BdCrzR in the male adult of the oriental fruit fly. Furthermore, we have delineated a neural function of Crz pathway in the sexual behavior and male fertility in B. dorsalis by RNAi.

The tissue-specific expression analysis of BdCrz revealed that mRNA accumulation was greatest for the CNS of the male adult B. dorsalis, and no detectable qPCR products existed in other tissues. Immunohistochemistry showed that three pairs of Crz neurons are located in the DLP region.
FIGURE 5 | Effect on sperm transfer of BdCrz or BdCrzR knockdown in B. dorsalis. The DAPI staining of sperm cells in the spermatheca of mated females with a male injected with dsGFP (A), dsCrz (B), or dsCrzR (C). Scale bars = 100 µm. (D) Number of sperm cells in the spermatheca of mated females with a male injected with dsGFP, dsCrz, or dsCrzR. Data are means ± SE based on three independent experiments. *p < 0.05, t-test.

of the adult CNS. These results seem to be equivalent to the data obtained in D. melanogaster and Musca domestica (Choi et al., 2005; Lee et al., 2008; Sha et al., 2012). A previous study demonstrated the production of Crz in the larval CNS of B. dorsalis, specifically by a group of three neurons in the dorsolateral (DL) region and eight pairs in the ventral nerve cord (VNC) (Hou et al., 2017b). In D. melanogaster, both the VNC and dorso-medial (DM) groups of Crz neurons of the larva die via programmed cell death during metamorphosis, whereas the DL neurons that are involved in the regulation of trehalose metabolism persist to adulthood and form a cluster of neurons, that are called DLP neurons (Choi et al., 2006; Lee et al., 2008; Kubrak et al., 2016). Holometabolous insects display substantial changes in morphology and behavior from juveniles to adults, whereas reproduction is a primary task of the adult for survival (Lee et al., 2008). Here, our results may also indicate the distinct fates of Crz neurons during CNS metamorphosis in B. dorsalis.

Previously, we found that BdCrzR are located in both the CNS and epitracheal gland in the larval stage of B. dorsalis (Hou et al., 2017b). Here, qPCR analysis demonstrated that BdCrzR transcript was most abundant in the fat body and CNS of the male adult of B. dorsalis. Consistent with this pattern, expression of CrzR occurred both in the heads and bodies with no sexual difference in M. domestica (Sha et al., 2012). In the Drosophila adult, high expression levels of CrzR were substantiated in the fat body, heart, and salivary glands, and a low level was determined in the CNS (Kubrak et al., 2016). In A. aegypti adults, quantitative spatial expression analysis revealed CrzR transcript in a variety of organs including head, thoracic ganglia, primary reproductive organs (ovary and testis), as well as male carcass (Oryan et al., 2018). These data suggest the multiple functions of the Crz signaling pathway in the adult fly. In addition, the developmental expressions of BdCrz and BdCrzR mirrored each other properly, which may indicate biologically relevant interactions between the Crz and its receptor (Hou et al., 2017b). A purpose of the present study was to test the sexual function of Crz signaling.
pathway in *B. dorsalis*. Our results showed that RNAi-mediated Crz knockdown lengthened the mating duration and decreased the fertility in males, and CrzR knockdown also decreased the numbers of egg in mated flies, while the behavioral sequence and the success of mating with Crz or CrzR knockdown were the same as the control flies. Injection of Crz in transitioning *H. saltator* individuals suppressed expression of vitellogenin in the brain and inhibited gamergate behaviors, such as dueling and egg deposition (Gospocic et al., 2017). Given the enrichment of CrzR in the primary and secondary sex organs of adult mosquitoes of *A. aegypti*, Crz signaling may play a role in regulating the reproductive biology (Oryan et al., 2018). In *Drosophila*, the silencing of Crz neurons resulted in infertility and extended copulation duration, and these two phenotypes were specific to males (Tayler et al., 2012). Our findings seem to be in accordance with the phenotypes obtained above. As reported by Acebes et al. (2004), copulation involves the transfer of sperm and seminal fluid, including substances that change female fertility and postmating behavior. In the present study, *BdCrz* or *BdCrzR* knockdown decreased the numbers of sperm cells in the reproductive organs (spermatheca) of the mated females, which is indicating that Crz signaling pathway modulates the reproduction via sperm transfer in *B. dorsalis*. In *Drosophila*, it has been reported that Crz controls the transfer of sperm and seminal fluid and copulation duration, thereby coupling the timing of these two processes for transfer of sperm and seminal fluid (Tayler et al., 2012).

A Crz-type peptide-receptor signaling pathway has also been identified in other protostomian invertebrates (e.g., annelids and mollusks) and in deuterostomian invertebrates (e.g., echinoderms), as demonstrated by phylogenetic analysis of genome/transcriptome sequence data and receptor deorphanization (Tian et al., 2016; Zandawala et al., 2017). In the starfish *A. rubens*, pharmacological experiments revealed that Crz causes contraction of cardiac stomach, apical muscle, and tube foot preparations, which are indicative of its physiological roles (Tian et al., 2017). The evidence of a reproductive role has been reported in *A. californica*, where the long form of the Crz receptor is expressed in parts of the ovotestis (Kavanaugh and Tsai, 2016).

Despite the fact that the complex mechanism of how the Crz signaling system is controlling the copulation physiological and behavioral programs remains to be elucidated, the present study indicated that Crz signaling pathway in the male adult is closely related to the mating duration and reproductive physiology in *B. dorsalis*. Indeed our data showed a strong loss (>80–90%) of oviposition in the recipient female adult. In conclusion, we believe this approach may be a new target that is different from all ovicidal insecticide activities reported in the female body, useful for pest control as *B. dorsalis* via development of Crz antagonists.

**AUTHOR CONTRIBUTIONS**

Q-LH, E-HC, H-BJ, YP, J-JW, and GS conceived the study and participated in its design. Q-LH and E-HC wrote the original draft and performed all of the experiments with the help of X-QL, S-FY, and P-JY. J-JW and GS provided the materials. Q-LH, E-HC, H-BJ, and GS analyzed the data. H-BJ, YP, J-JW, and GS edited and reviewed the paper.

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