A unique mating pattern of Panorpodes kuandianensis (Mecoptera: Panorpodidae)

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Abstract

Sexually reproductive insects exhibit diverse mating behaviors. However, the mating pattern remains unknown for Panorpodes of Panorpodidae to date. In this study, we investigated the mating behavior and copulatory mechanism of the short-faced scorpionfly Panorpodes kuandianensis Zhong, Zhang and Hua, 2011 for the first time. The results show that the male provides a salivary mass as a nuptial gift to the female and starts to copulate with the female in a V-shaped position, then changes to an end-to-end position by temporarily twisting the female abdominal segments VII–IX by 180°. During mating the basal processes and the basal teeth of the gonostyli and the hypandrium are used to obtain copulation and sustain the coupling of genitalia to secure successful sperm transfer. This unique mating pattern is greatly different from that of other Mecoptera reported and is likely evolved as an adaptation in the context of sexual conflict.

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Introduction

The vast majority of sexually reproductive animals propagate through mating. For insects diverse mating positions have evolved to deliver sperm ejaculates, with the female-above pattern as the groundplan (Huber et al., 2007; Huber, 2010), from which all other deviations evolved through rotations or flexions of the terminal abdomen of males (McAlpine et al., 1981; Bickel, 1990; Huber, 2010). The evolution of mating position and other behavioral traits involved in mating are regarded to relate to sexual conflict in insects (Chapman et al., 2003; Parker, 2006).

Most Mecoptera are characterized by their interesting sexual behaviors, including various mating positions, nuptial feeding or forced copulation by modified grasping devices (Thornhill, 1973, 1980, 1981; Thornhill and Sauer, 1991; Sauer et al., 1998; Engqvist and Sauer, 2001, 2003; Byers, 2002; Engqvist, 2007, 2009, 2011; Kock et al., 2009; Gao and Hua, 2013; Zhong and Hua, 2013; Zhong et al., 2015a, b). Of the nine families, five have been studied with respect to their mating behaviors and/or copulatory mechanisms (Cooper, 1974; Thornhill, 1981; Byers and Thornhill, 1983; Thornhill and Sauer, 1991; Ma et al., 2010; Zhong and Hua, 2013; Zhong et al., 2015a, b). The male of Boreidae uses his paired hook-shaped wings to secure the female dorsally to form a typical female-above mating position (Cooper, 1974). In Bittacidae the male temporarily twists his abdomen by 180° to maintain a belly-to-belly mating position to control the nuptial gift (Gao and Hua, 2013). Most males of Panorpidae provide a salivary mass as a nuptial gift and sustain a V-shaped mating
position (Thornhill, 1981; Byers and Thornhill, 1983; Zhong et al., 2015a). The male of *Panorpa liui* Hua, 1997 offers the female only prey rather than salivary mass as a nuptial gift owing to the less-developed salivary glands of males (Ma and Hua, 2011). The species of *Furcatopanorpa* (Panorpidae) (Zhong et al., 2015b) and *Chorista* (Choristidae) (Byers and Thornhill, 1983) adopt an O-shaped mating position for the male lacking a notal organ to help control the female. Similarly in the males of Panorpodidae the notal organ is vestigial or thoroughly lacking (Carpenter, 1953; Byers and Thornhill, 1983; Tan and Hua, 2008; Zhong et al., 2011; Krzeminski and Soszynska-Maj, 2012), but their mating behavior has not been satisfactorily studied thus far.

Panorpodidae is the sister group of Panorpidae (Willmann, 1987; Friedrich et al., 2013), and consists of only 13 extant species in two genera: *Brachypanorpa* Carpenter, 1931 occurring in North America and *Panorpodes* MacLachlam, 1875 almost exclusively distributed in eastern Asia (Pollmann et al., 2008; Hu and Hua, 2016). *Panorpodes colei* Byers, 2005 from western North America is the only exception (Byers, 2005). The male of *Brachypanorpa* does not provide any nuptial gift, and starts mating with the female in a V-shaped mating position at any time of the day but usually in the evening (Carpenter, 1953; Byers, 1997). As far as we know, however, the mating behavior in *Panorpodes* has not been reported to date.

In this study, we investigated the mating behavior and copulatory mechanism of the short-faced scorpionfly *Panorpodes kuandianensis* Zhong, Zhang and Hua, 2011 and found an unusual mating pattern in Mecoptera.

### Material and Methods

#### Insect collection

Adults of *P. kuandianensis* were collected at the Quanshan Forest Park (41°06′N, 125°02′E, elev. 650–800 m), Kuandian County, Liaoning Province in northeastern China from early July to August in 2015 and 2016.

#### Insect rearing

Male and female adults were reared in three gauze cages (40 cm × 40 cm × 60 cm) in the laboratory under natural conditions following Zhong and Hua (2013). Absorbent cotton containing water was replaced each day in a Petri dish. Twigs with leaves were placed in the cages to simulate the habitat of the adults, and were replaced every three days. Each cage was supplied with five drops of honey per day to prevent low nutritional status (Jiang et al., 2014).

#### Mating behavior observation

Ten males and ten females were reared in each cage to investigate their mating behaviors. These adults were observed 24 h a day during 6–10 July 2015 to check the circadian rhythms of mating activity. Then adults were observed every 30 min from daybreak to noon to record the mating behaviors, including mating positions and copulatory processes. Photographs were taken with a Nikon D90 digital camera.

#### Freeze-fixation of copulating pairs

Pairs in copula were frozen through carbon dioxide aerosol spray compressed in hydraulic cans, and were immediately fixed in Carnoy’s solution at room temperature for 24–48 h to stabilize the interactions of their genital structures and preserved in 75% ethanol.

#### Light microscopy and scanning electron microscopy

Fifty adults (30 males and 20 females) fixed in Carnoy’s solution were dissected under a Nikon SMZ1500 microscope. Photographs were taken with a QImaging Retiga-2000R digital camera attached on the microscope and were stacked with the Syncroscopy Auto-Montage software.

For scanning electron microscopy (SEM), the dissected genital structures were cleaned ultrasonically for 90 s, and then dehydrated in a graded ethanol series, freeze-dried for 3 h, coated with gold in a sputter coater, and examined in a Hitachi S-3400N scanning electron microscope (Hitachi, Tokyo, Japan) at 5 kV.

### Results

#### Mating behavior

Thirteen pairs of *P. kuandianensis* were recorded to mate in the morning and three of them were frozen in copulation. The male provided a salivary mass to a nearby female and tried to attract her through vibrating his wings rapidly (Fig. 1a). Then he stretched out his paired gonostyli to grasp the abdominal end of the female, used his hypandrium to seize the female cerci and attempted to establish the connection of the genita-
lia. Meanwhile, the female usually tried to get rid of the male control by wriggling her abdomen. To suppress the female resistance, generally, the male adjusted his posture and seized the female abdomen by the paired basal processes and the basal teeth to couple their genitalia and guarantee the sperm transfer. In the initial phase of copulation, the male randomly stood on one side of the female (six left-sided matings and seven right-sided matings) to sustain a V-shaped position with their genitalia coupled (Fig. 1b). While the female consumed the salivary mass, the male changed the mating position from the V-shape to an end-to-end position (Fig. 1c). The male kept on pulling the female till the end of copulation, and some females (2 of 10) were observed to invert their bodies before terminating copulation (Fig. 1d).

Male and female genitalia

The genital segment (abdominal segment IX, A9) of male *P. kuandianensis* consists of a dorsal epandrium (tergum IX), a ventral hypandrium (sternum IX), and the median genitalia (Figs 2a, b). The genitalia consist of a central aedeagus, a pair of parameres, and a pair of lateral gonopods (Fig. 2a). The gonopods are fused at the base, each comprising a basal gonocoxite and a distal gonostylus (Figs 2a, b).

The epandrium is broad basally and nearly parallel apically. Two short, round-tipped cerci are born on the lateral sides of abdominal segment XI (A11) and extend out from below the epandrium (Fig. 2b).

The hypandrium consists of a long broad basal stalk and a pair of short distal hypovalves (Figs 2e, f). Each
hypovalve has a rounded incrassate process, which is situated basally on the inner margin of the hypovalve and covered with spinules on the dorsal surface (Fig. 2f).

The paired gonocoxites fuse basally to form a U-shaped concavity from the middle part to accommodate the aedeagus centrally (Fig. 2a). Each gonostylus has a prominent lobe-like basal tooth and a well-developed finger-shaped basal process. The elongate basal process bears numerous conical sensilla (Figs 2c, d).

Each paramere is composed of an applanate ventral branch, a strongly-sclerotized dorsal branch with a lateral process arising from the middle part, and a slender basal stalk. The parameres cling to the aedeagus, so that the ventral branches are unable to stretch ventrally from the aedeagus (Figs 2a, 3b).

The aedeagus is an interconnecting organ of the male, and consists of a pair of ventral valves, a pair of dorsal valves, and a central phalotreme. The ventral valves are usually concealed by parameres (Figs 3a, b).

The female genitalia consist mainly of a genital plate and a subgenital plate (Figs 2g, h). The subgenital plate is oblong with two weakly-sclerotized sclerotomes (Fig. 2h). The two sclerotomes curve dorsally to form a genital chamber, inside which the genital plate is situated.

The female genital plate is a short strongly-sclerotized structure, situated at the base of the genital chamber. The orifice of the spermathecal duct is located ventrally at the sub-apical area of the genital plate distal to the sculptured area (Figs 3c, d).

**Coupling of the genitalia**

The claspers and other grasping devices of the male grasp the terminal end of the female abdomen to establish the coupling of the genitalia (Figs 4, 5). The hypovalves with incrassate processes seize the female cerci and cause the cerci to bend dorsad (Figs 4, 5a, b). The finger-like basal processes of the male gonostylus clasp the rotated genital segments of the female (Figs 4a, 5b, d). The paired lobe-like basal teeth seize the rear area of the genital chamber (Figs 4b, 5c).
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Copulatory mechanism

When the female is under control, the male seizes her cerci with his hypovalves and opens her genital chamber. After that, the male inserts his aedeagus into her genital chamber, with the lateral membranes of the female A9 obstructing the paired lateral processes into the genital chamber (Fig. 4b). Then the male adjusts his posture, thus enabling the phallotreme to contact the copulatory pore of the female to deliver his sperm ejaculates into the female spermatheca. During copulation the male keeps the stability of genital connection by using the lobe-like basal teeth of his gonostyli to restrict the movement of the genital plate (Figs 4b, 5c) and utilizing the basal processes of his gonostyli to prevent the female abdomen from shaking off the male control. The coupling genital structures mesh completely when the mating position changes from a V-shape to an end-to-end position.

Torsion of the female abdomen

The abdominal segments of female *Panorpodes kuandianensis* are unique in that the abdominal segments VII and VIII (A7 and A8) and the most part of A9 are usually concealed within abdominal segment VI (A6). In the process of position-change, the terminal abdomen of the female undergoes a rotation clockwise or anti-clockwise (Figs 5c, d). The abdominal segments are recurved upward when the paired male and female...
sustain a V-shape mating position. When the mating position changes to end-to-end, A7 of the female is twisted a very small angle, A8 is twisted ~60° (Figs 4, 5c, b) and A9 is twisted ~120° (Figs 4, 5d). The torsion of the female abdomen results from the deformation of the intersegmental membrane.

Discussion

This research is likely the first attempt to describe the mating behavior and the copulatory mechanism of the genus *Panorpodes* in Panorpodidae. *P. kuandianensis* is unique in mating behavior for the male and female to change the mating position from a V-shape to an end-to-end by temporarily twisting the female abdomen by 180°. Some specialized structures (such as the basal processes and basal teeth) play important roles in obtaining and maintaining a successful mating. This unique mating pattern is remarkably different from that of other Mecoptera studied (Cooper, 1974; Thornhill, 1981; Byers and Thornhill, 1983; Thornhill and Sauer, 1991; Byers, 1997; Ma et al., 2010; Zhong and Hua, 2013; Zhong et al., 2015a, b).

The genital and non-genital grasping devices are usually modified in internally inseminating species of insects to maintain the firm coupling of genitalia between the two sexes (Arnqvist, 1997; Matthews and Matthews, 2010; Chapman, 2013; Zhong et al., 2015a; Richmond et al., 2016), such as the specialized abdominal apparatus in water striders (Arnqvist and Rowe, 2002; Perry and Rowe, 2012) and the notal organ in scorpionflies (Mickoleit, 1971; Thornhill and Sauer, 1991; Zhong and Hua, 2013). The basal processes of gonostyli can restrict the movement of the female genital segments in *P. kuandianensis*, but are used to impede the genital plate from retreating in *Neopanorpa longiprocessa* (Zhong and Hua, 2013). The hypandrium with incrassate processes of *Panorpodes* likely plays a part in stimulating the cerci of the female during mating. A similar phenomenon is also reported in Panorpidae species (Zhong and Hua, 2013; Zhong et al., 2015a).
Flexion and rotation of the abdomen are considered to be adaptations for mating and storing the abdominal segments when not in use (McAlpine et al., 1981), and take place in the male terminal abdomen in the majority of insects. Well-known examples are the males of Diptera, the abdomens of which twist from 90° to 360° around the long axis of the abdomen to adapt the diverse mating positions (McAlpine et al., 1981; Bickel, 1990; Huber et al., 2007). For the reason of abdominal rotation, Bickel (1990) suggests that female sexual selection pressure promotes the rotation of male abdomen and encourages the circumversion of the terminal abdomen to couple in a male-above position, because a male on top of the female may not only grasp his mate more securely and control the timing of disengagement, but is also able to fend off any intruding males or to stimulate the female using his legs and mouthparts. By contrast, however, in P. kuandianensis the torsion occurs in the female. This phenomenon may be correlated with the abdominal morphologies of both sexes. The segments A7–A9 are thick and short in the male, but slender and concealed within A6 in the female (Zhong et al., 2011). In this case, it seems more convenient to rotate the female than the male abdomen. We suggest that during the copulation of insects either the active rotation in the male abdomen or the passive rotation in the female abdomen seems to have evolved to adapt to the sexual conflict, and is mainly beneficial to the male to dominate the mating process.

Fig. 5. Coupling genitalia and torsion of the female abdomen. a, coupling genitalia in lateral view; b, the same, with the hypandrium removed to show the cerci of the female and the basal process of the male; c, the torsion of the female abdominal segments VII–VIII, lateral view, with the epandrium removed to show the subgenital plate of the female and the basal tooth of the male; d, the torsion of the female abdominal segments VII–IX, lateral view. BP, basal process; BT, basal tooth; Ce, cercus; Ep, epandrium; Gc, gonocoxite; Gs, gonostylus; Hv, hypovalve; Pm, paramere; SP, subgenital plate; S6, sternum VI; T6–T9, tergum VI–IX.
The position-changed mating is unusual in Mecoptera (Mickoleit, 1971; Cooper, 1974; Thornhill, 1981; Thornhill and Sauer, 1991; Byers, 1997; Gao and Hua, 2013; Zhong and Hua, 2013; Zhong et al., 2015a, b), but is common in Orthoptera (Alexander and Otte, 1967), Diptera (McAlpine et al., 1981), and Lepidoptera (Scott, 1986; Fänger and Naumann, 1998). The initial side-by-side mating position is sustained by the majority of Lepidoptera species, and changes to end-to-end immediately or shortly after establishing genital contact (Scott, 1986; Fänger and Naumann, 1998). For the possible reasons of the position-changed mating, McAlpine and Munroe (1968) believe that the initial position in Diptera is used to establish the interlock of their genitalia and usually takes place during flight. The secondary position is needed to continue copulation and transfer sperm and frequently occurs while the pair rests on a substrate. The position-changed mating in Panorpodes is likely an effective strategy of the male to dominate the mating and to compensate for the lacking of a notal organ, because the end-to-end mating position is frequently accompanied by the pulling behavior. At the pulling phase, the female is almost helpless, and the locomotion of the coupled pair usually depends on the male. This mating pattern may be best explained as a male strategy against female disengagement by turning away. However, details of the contact of the phalotreme of the male and the copulatory pore of the female have not been observed in this study. It may require subsequent preparations of histological serial sections and 3D reconstructions as applied in the fruit fly (Mattei and Wolfner, 2016) and the bushcricket (Wullf et al., 2015).

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