To females of a noctuid moth, male courtship songs are nothing more than bat echolocation calls

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It has been proposed that intraspecific ultrasonic communication observed in some moths evolved, through sexual selection, subsequent to the development of ears sensitive to echolocation calls of insectivorous bats. Given this scenario, the receiver bias model of signal evolution argues that acoustic communication in moths should have evolved through the exploitation of receivers’ sensory bias towards bat ultrasound. We tested this model using a noctuid moth Spodoptera litura, males of which were recently found to produce courtship ultrasound. We first investigated the mechanism of sound production in the male moth, and subsequently the role of the sound with reference to the female’s ability to discriminate male courtship songs from bat calls. We found that males have sex-specific tymbals for ultrasound emission, and that the broadcast of either male songs or simulated bat calls equally increased the acceptance of mated males by the female. It was concluded that females of this moth do not distinguish between male songs and bat calls, supporting the idea that acoustic communication in this moth evolved through a sensory exploitation process.

Keywords: bat predation; courtship ultrasound; receiver bias; tymbal

1. INTRODUCTION

Moths have evolved ears that detect the echolocation calls of insectivorous bats (Roeder 1962; Minet & Surlykke 2003). Subsequently, some groups of moths have evolved sound-producing organs for intraspecific acoustic communication and/or defensive responses against predators (Spangler 1988; Conner 1999; Greenfield 2002). In wax moths, males produce ultrasonic clicks with tegular tymbal organs to attract females, while they show a freeze response to bat echolocation calls (Spangler 1988; Greenfield & Weber 2000). In tiger moths, both sexes emit ultrasonic clicks with metathoracic tymbals in response to bat calls to jam the echolocation of or to deter bats from attacking (Minet & Surlykke 2003; Barber & Conner 2006). Some tiger moths produce ultrasonic clicks for sexual communication as well (Conner 1999). These examples indicate that acoustically communicating moths have gained the ability to distinguish the ultrasonic sounds of bats from those of conspecifics based on characteristics of the sounds, pulse structure for instance (Greenfield & Weber 2000; Fullard et al. 2007).

The receiver’s sensory bias for detecting a predator and/or prey drives the evolution of signals in animal communications (e.g. vibratory displays in water mites, sword ornaments in swordtail fish and calling song frequency in túngara frogs: see ‘receiver bias model’ in Ryan 1998; Greenfield 2002). Ultrasonic communication in moths has also been supposed to have developed through exploitation of the ability to detect ultrasound for avoiding predation by bats (Endler & Basolo 1998; Conner 1999; Greenfield & Weber 2000). However, the receiver bias model has not been experimentally verified in acoustic communication in moths, because the signal receivers in the moths examined had already evolved the ability to distinguish singing mates from echolocating bats, as described above (Surlykke & Fullard 1989; Greenfield & Weber 2000; Skals & Surlykke 2000; Fullard et al. 2007). If we can find a case where the receivers do not discriminate between the sounds of bats and those of a mate, it would present direct evidence for the process of sensory exploitation.

We recently reported acoustic sexual communication in the common cutworm Spodoptera litura (Nakano et al. 2009). However, the mechanism of sound production and the ability of receivers to discriminate sounds have not been studied. Here, we present the mechanism of ultrasound production, and the receivers’ responses to male ultrasound playbacks and simulated bat calls in S. litura.

2. MATERIAL AND METHODS

Using 1–2-day-old S. litura moths, we conducted two behavioural tests in the last 2 h of the scotophase, in which this moth shows high mating activity (methods in the electronic supplementary material; Nakano et al. 2009). First, to verify the occurrence of acoustic communication, we examined the effect of the muting of males on mating success. For muting, we punctured the tymbals on the male metathorax using fine forceps (methods in the electronic supplementary material). A sham operation was conducted by puncturing the mesothoracic coxae. For all surgical operations, male moths were anaesthetized by CO2 and treated under a stereomicroscope 1 day before the tests.

Next, to examine the ability of the female to discriminate between male songs and bat calls, the mating success of sham-operated males, which generate ultrasound, was compared with that of mated males with the broadcast of (i) male songs, (ii) background noise, or (iii) simulated bat calls from a loudspeaker set 20 cm from the female (methods and figure S1 in the electronic supplementary material). The male songs and bat calls were broadcast at the natural levels, 70±4 dB SPL and 100±1 dB SPL respectively, at the point of female moth (Surlykke & Kalko 2008; Nakano et al. 2009). Two different types of echolocation calls of moth-eating bats were simulated; one was ultrasonic clicks of short frequency-modulation (FM) pulses (e.g. calls of Eptesicus fuscus), and the other was those of long constant-frequency (CF) pulses (e.g. calls of Rhinolophus ferrumequinum) in the search-approach phase, which elicit evasive responses in moths including S. littoralis, a congener of S. litura (Skals et al. 2005) (see methods in the electronic supplementary material).
Males used in this experiment were deafened to eliminate effects of broadcast sound stimuli on their behaviour. In all behavioural tests, a single unmated male was introduced into a cubic mesh cage (18 × 18 × 18 cm) housing 5–10 intact virgin females. Multiple females were housed in the cage so that at least one female would be releasing sex pheromones during the experiment. The male usually readily flew up to one of the females, and started courting. We defined the female mate acceptance as completion of copulation, which occurred subsequent to the cessation of her locomotion. The observation of mating behaviour was continued until the female accepted the male or rejected him by flying away. The pair observed was then removed from the cage, and the experiment was continued with a new male. The ultrasound emitted from males and the loudspeaker was continuously monitored with an ultrasound detector (model D240x, Pettersson Elektronik AB, Uppsala, Sweden).

3. RESULTS

Male *S. litura* emitted ultrasounds during courtship in the proximity of a female (figure 1; electronic supplementary material, movie S1). The male moth possesses a striated cuticular membrane, i.e. tymbal organ, on the ventral side of the metathorax (figure 2). Destruction of this organ deprived the male of the ability to produce ultrasound. The ultrasonic song emitted from the tymbal consisted of clicks of short duration (0.1–0.2 ms), and showed active/passive modulation cycles divided by a silent gap (figure 1). Females do not have tymbals on the thorax, and did not emit ultrasound during mating behaviour (the electronic supplementary material, movie S1).

Behavioural experiments using muted males demonstrated the significance of courtship ultrasound for mating success. Most of the females, 95–100%, accepted singing males for mating, whereas only 40% per cent accepted muted males (Fisher's exact probability test for male treatments, *p* < 0.0001) (table 1). Sham-operated males producing natural ultrasound had 100 per cent mating success (table 2). Playback of male ultrasounds completely restored the muted males’ mating success to 100 per cent in comparison with playback of noise (56% acceptance) (table 2; Fisher's exact probability test for sound stimuli, *p* < 0.0001). Bat calls were also effective; females showed high mate acceptance (91 and 100%) when simulated FM or CF bat echolocation calls were broadcast (table 2).

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**Table 1.** Effect of male ultrasound on mating success. (Mating success of muted males was significantly lower than that of intact or sham-operated males.)

<table>
<thead>
<tr>
<th>male treatment</th>
<th>n</th>
<th>% matingb (no. of pairs mated)</th>
</tr>
</thead>
<tbody>
<tr>
<td>intact</td>
<td>21</td>
<td>100 (21)</td>
</tr>
<tr>
<td>tymbal punched</td>
<td>17</td>
<td>41 (7)</td>
</tr>
<tr>
<td>sham-operated</td>
<td>21</td>
<td>95 (20)</td>
</tr>
</tbody>
</table>

*a*All females were intact;  
bFisher’s exact probability test, *p* < 0.0001;  
cmesothoracic coxae were punctured instead of tymbals.

**Table 2.** Effect of synthesized ultrasound on mating success. (Broadcasts of male ultrasound, FM (frequency-modulation) bat calls, and CF (constant-frequency) bat calls increased the mating success of muted males.)

<table>
<thead>
<tr>
<th>sound stimuli</th>
<th>sound level (dB SPL)</th>
<th>n</th>
<th>% matingb (no. of pairs mated)</th>
</tr>
</thead>
<tbody>
<tr>
<td>natural male ultrasound</td>
<td>70</td>
<td>13</td>
<td>100 (13)</td>
</tr>
<tr>
<td>playback of male ultrasound</td>
<td>70</td>
<td>20</td>
<td>100 (20)</td>
</tr>
<tr>
<td>playback of background noise</td>
<td>22</td>
<td>18</td>
<td>56 (10)</td>
</tr>
<tr>
<td>simulated FM bat calls</td>
<td>100</td>
<td>22</td>
<td>91 (20)</td>
</tr>
<tr>
<td>simulated CF bat calls</td>
<td>100</td>
<td>18</td>
<td>100 (18)</td>
</tr>
</tbody>
</table>

*a*All males tested here were deafened to eliminate the effect of sound stimuli on their behaviour;  
bFisher’s exact probability test, *p* < 0.0001;  
cmesothoracic coxae were punctured as a sham operation.
4. DISCUSSION

Males of *S. litura* have evolved a tymbal organ (figure 2), which looks similar to the tymbals found in many arctiid tiger moths (Barber & Conner 2006). The tymbal of *S. litura*, however, differs in its location, the ventral side of the metathorax, versus the lateral side in arctiids. Also, the tymbals are found in both sexes in arctiids but only in males in *S. litura*. Both sexes of tiger moths show phonoresponses to bat calls, i.e. they react to bat calls by producing sound (Barber & Conner 2006), but neither sex of intact *S. litura* shows a phonoresponse (data not shown). These findings suggest that male *S. litura* have developed a sound-producing mechanism not for defence, but for sexual communication.

The male ultrasound may be a signal used for mate recognition by receptive females, because muted males were accepted at a higher rate when the male song was played back. Hence, we considered the possibility that the female distinguishes male ultrasound from bat calls, as reported for females of wax moths, *Galleria mellonella* (Barber & Weber 2000; Skals & Surlykke 2000). In playback experiments, however, simulated bat calls had the same effect as the male courtship song (table 2). These results suggest that females of *S. litura* do not discriminate between male songs and bat calls.

Eared moths freeze to evade echolocating bats because stationary moths have a better chance to be detected by bats (Roeder 1962; Spangler 1988; Greenfield & Weber 2000; Minet & Surlykke 2003). We recently showed that the ultrasonic songs of the male *Ostrinia furnacalis* render the females motionless, and argued that this is the same response as that to ultrasonic bat calls (Nakano et al. 2010). Therefore, the finding presented here supports the idea that by emitting ultrasounds, males deceive the female into making a freeze response during the male’s courting. Subsequently, the singing male can copulate with the stationary female with much less difficulty (Nakano et al. 2008, 2010). Consequently, the female may incidentally select ultrasonic-producing males over silent males by accepting a significantly higher proportion of singing males for mating. These reactions by both genders would have contributed to the establishment of sexual communication in moths. In conclusion, we have demonstrated that females of *S. litura* do not distinguish between male songs and bat calls, consistent with the idea that acoustic communication in this moth evolved through a sensory exploitation process.

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