

Pre-copulatory courtship behavior in a solitary bee, *Nomia triangulifera* Vachal (Hymenoptera: Halictidae)

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Summary — Complex courtship behavior occurs both before and during mating of a solitary bee, *Nomia triangulifera*. Males begin emerging before females, and patrol over the emergence site, although some males also fly near sunflower plants, the exclusive food source of the females. The sizes of males patrolling at these two locations, and those collected while mating were not significantly different. Courtship behavior involves use of the fore- and mid-legs, the metasoma (abdomen), antennae, and the indirect flight muscles to produce loud, audible buzzes. The hind legs have expanded tibiae which are used for claspings the female. Females rarely mated more than once when held in cages. Receptive females usually lacked sperm in their spermathecae, while unreceptive females often had sperm.

***Nomia triangulifera* / Halictidae / mating behavior / sexual selection**

INTRODUCTION

Many features of a male's behavior and morphology are assumed to function to induce a female to initiate copulation, and then stimulate her to continue reproductive behavior (eg sperm transport, ovulation), thereby increasing the probability that the copulating male will fertilize her ova (Eberhard, 1985). Such behavior is believed to underlie the evolutionary process of sexual selection by female choice

(eg Darwin, 1871; Richards, 1927; Alcock and Gwynne, 1991; West-Eberhard, 1984). Alternatively, species-specific courtship behavior functions in contexts of reproductive isolation or mate recognition (Verrell, 1988).

Most studies to date have reported little or no pre-copulatory courtship behavior among bees (eg Batra, 1966; Barrows, 1975; Alcock *et al*, 1978; Triplett and Gittins, 1988; Eberhard, 1991). *Nomia* (*Epinomia*) *triangulifera* Vachal, reported on

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here, seems unusual because of its extremely complex pre-copulatory courtship behavior, combined with structural modifications. The genus *Nomia* (Hymenoptera: Halictidae) occurs throughout most regions of the world, and in the New World is represented by 20 species in four subgenera (Moure and Hurd, 1987)*. Most male *Nomia* have highly modified hind legs, metasomal sterna and genitalia, which provide species-specific taxonomic characters (eg Cross, 1958; Michener, 1965; Ribble, 1965). The biological functions of these secondary sexual characters are unknown, and comparative studies of the mating biology of this group promise to illuminate their function. This report, together with previous observations on this species from another locality (Cross and Bohart, 1960), may provide an example of intra-specific divergence in courtship behavior.

MATERIALS AND METHODS

The mating behaviour of *Nomia triangulifera* was studied at large aggregations of nests at a farm on a floodplain along the south bank of the Kansas River between Eudora and Lawrence (Douglas County), Kansas (38°57'30" N, 95°7'30" W). Bees have been nesting at this site at least since 1972 (Shipe, personal communication).

Courtship and mating behavior were studied by direct observation of > 200 mating pairs. Copulating bees were largely unresponsive to nearby movements, could be approached within one cm (a 10X hand-lens was sometimes used), and mating pairs could even be picked up. Mating was either: i) video-taped (3.5 h) using a Panasonic videocamera with close-up lens and tripod (in 1987 and 1988); or ii) filmed (4 h) with a Paillard Bolex 16-mm film camera with an Yvar 150-mm macro-lens and tripod (in 1986).

Mating pairs were picked up and quickly placed on a platform set up in front of the camera. Approximately 7 h of audio recordings were made using a Uher 4000 reel-to-reel tape recorder (tape speed = 19 cm/s), with a standard microphone (which often served as the platform for filming). For analyses, recordings were played through a Krohn-Hite mode 3550 filter (LP = \approx 575 Hz) to a Kay Elemetrics DSP Sonagraph (Model 5500).

To examine size-related patterns of mating behavior, we measured the sizes (inter-tegular distances) of females and their male partners ($N = 20$ males and 20 females), males patrolling at the emergence/nesting site ($N = 44$), and males patrolling at sunflowers ($N = 51$), using a Wild microscope with an ocular micrometer (magnification = 10X). Some males ($N = 21$) were dried to a constant weight at 50 °C (Thermolyne Oven-Incubator), and weighed on a Fisher electronic balance to the nearest 0.001 g; for these males the lengths at the longest distance of the inner face of the expanded hind tibiae were also measured.

The receptivity of females in a cage was studied as follows (after O'Neill and Bjostad, 1987). After being "courted", females exhibited obvious behaviors indicating their sexual receptivity: receptive females elevated their metasoma slightly, and unreceptive females curled the metasoma under their body, and sometimes also bit at the males' legs. Receptivity status of females with otherwise unknown sexual histories was classified as follows: "receptive to 1st male and mated" (= a female copulated with the first introduced male) ($N = 34$); "unreceptive" (= a female refused to mate with the first male) ($N = 18$); and "receptive to 1st male and separated before mating" (= a female was receptive to the first male, but the pair was separated before intromission) ($N = 26$). Females from these classes were individually introduced to 3–5 males in either a 16 x 12 x 12 cm net-covered fish-breeding cage, or a 1 x 0.5 x 1 m enclosure (an emergence trap) to ascertain subsequent receptivity.

Mating pairs were observed in the field, and females were categorized as "receptive but un-

* *Nomia triangulifera* is presently placed in the subgenus *N* (*Epinomia*). In a forthcoming key, Michener *et al* (in preparation) recognize the genus *Dieunomia* for *N* (*Epinomia*) and *N* (*Dieunomia*), and the genus *Nomia* for *N* (*Acunomia*) and *N* (*Curvinomia*). For convenience these subgeneric names are retained in the *Discussion*, although it is likely that some of the groups are artificial.

mated" ($N = 11$) (ie, males were removed before intromission), or "unreceptive" ($N = 14$). Females were placed in vials containing insect preservative (Kahle's solution), and were later dissected and examined for presence of sperm cells in the spermathecae.

Statistical tests were from Sokal and Rohlf (1981), or calculated using Statview on a Macintosh computer.

Voucher specimens of the bees are in the Snow Entomological Museum, and audio recordings are in the Cornell University Library of Natural Sounds; video and audio recording may be obtained from the first author.

RESULTS

N. triangulifera was active from mid- to late-August and into September in eastern Kansas. The populations were protandrous, but after several days the emergence distributions of the sexes overlapped. On all dates there was a heavily male-biased operational sex ratio, both in a year when the secondary sex ratio was near equality, and one when it was male biased (Wcislo, 1992). Males flew in great numbers over the previous year's nest site, which was where receptive females emerged and where they returned to nest if the plant cover was suitably sparse. Males were also abundant on the plant (*Helianthus annuus*) from which females gathered pollen and nectar; males also drank sunflower nectar. Males flying near plants frequently pounced on females, but successful copulations were never observed there. After mating, females dug nests which they provisioned with sunflower pollen and nectar. Approximately 2–3 weeks after initial emergence, males became rare at the nest site and were frequently collected on flowers, while female continued provisioning nests.

Courtship behavior of *N. triangulifera* was also briefly observed at a small aggregation near the levee north of the entrance

to River Front Park, Lawrence, and at a site 5 miles north-west of Lawrence (described in Cross and Bohart, 1960). At both sites courtship behavior was qualitatively similar to behavior described below (Wcislo, unpublished observations).

Most of our observations were of mating pairs on the ground at the emergence site. We do not know the relative abundance of receptive females at the emergence sites and at flowers. Males were never observed successfully mating on flowers ($>N = 32$ attempted matings), even on plants close to the emergence site. Males flying over the emergence/nesting site were not, on average, different in size from males collected at sunflower plants nor from males collected with females *in copula* ($P > 0.5$, with repeated Mann–Whitney *U* tests).

The courtship behavior involved motions of the antennae, all three pairs of legs, the metasoma and associated sternal modifications, and presumably the dorso-ventral flight muscles. Several details did not fully agree with the description of courtship behavior by Cross and Bohart (1960), based upon their limited observations of a Utah population; descriptions by those authors are parenthetically italicized for comparison, indicated by "C and B". Otherwise, descriptions given below agree with those by Cross and Bohart (1960).

In the nesting area males flew rapidly over the ground where females emerged, and frequently alighted to inspect or enter nest entrances and emergence holes. Upon contacting a female, a male rapidly climbed on her, and wrapped his hind legs around the latero-ventral portion of metasomal sterna 1–3 (C and B: "*His hind legs lock beneath the posterior portion of her abdomen*"). The hind tibiae are greatly expanded, and the inner faces are contoured to fit the general shape of the female metasoma. Males with larger inter-tegular distances had larger faces on their inner hind

tibiae (Kendall's $t = 0.52$, $P = 0.002$, $N = 21$). Throughout the courtship, but especially soon after pairing, other males flew at and pounced on the mating pair: sometimes a cluster of up to 8 males crawled over a mating pair. The expanded male tibiae prevented a male from being knocked off a female, or otherwise prevented the pair from being separated. In numerous observations (> 200), a male *in copula* was never displaced from a female by other males. The expanded tibia as well as the femur both have enlarged flexor muscles (Wcislo, unpublished observations).

A male's mid-legs were held lateral to the female's metasoma, nearly parallel to the longitudinal axis of her metasoma, and were directed posteriorly (C and B: "(mid-legs are locked)... between her fore and hind wings and under her propodeum"). The male's forelegs were placed either on the pronotum, or sometimes on the front edge of the female's front wings (C and B: "(the male)... locks this forelegs beneath her mesothorax").

Precopulatory courtship behavior is diagrammed in figure 1. The female was nearly motionless throughout, except for occasionally grooming an antenna; sometimes a (subsequently unreceptive) female walked around the surface with a male mounted on her back. The following time durations are for $\approx 30^\circ\text{C}$.

Both of the male's antennae were slowly moved downward in a motion lasting 1.07 s (SD = 0.1; $N = 9$), and then were rapidly flung upward ($x = 0.33$ s duration (SD = 0.1; $N = 9$)), repeating about once per s, giving the appearance of an asymmetric metronome (C and B: "(there is)... alternate vertical and lateral jerking of the antennae").

The male's first pair of legs rapidly tapped the female's pronotum, quickly followed by rubbing her pleural area or mid-

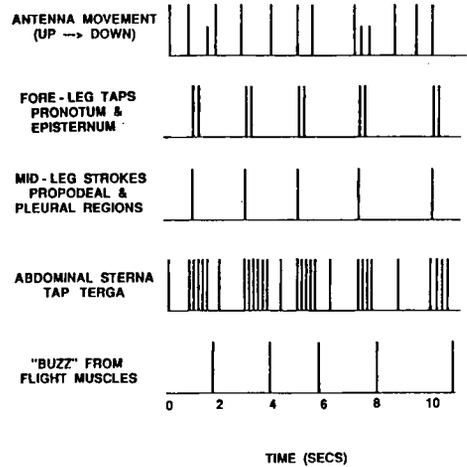


Fig 1. Representation of a portion of the precopulatory behavior of male *Nomia triangulifera* (Note: for the antennal motions, half bars indicate partial motions; for foreleg motions, each bar indicates that the male touches both the pronotal and pleural regions).

legs, and then repeating these movements; this entire sequence is repeated once per 1.3–2.0 s (not reported by C and B).

The extended mid-legs (fig 2, position A) were rotated $\approx 90^\circ$ around the coxae to be brought forward and downward to stroke the female's pleural region (the exact position varied with the position of the male's body) (fig 2, position B); the legs were then contracted and lifted up (fig 2, position C), which again stroked the female; the legs were then counter-rotated and extended to their starting position (fig 2, position A). After courtship has begun, the mid-legs often quivered when they were in the extended "starting" position. This sequence was repeated every 1.3–2.0 s (not reported by C and B).

Male metasomal sternum 5 has a modified pair of protruding stubs which have a contoured and knobbed surface (there are

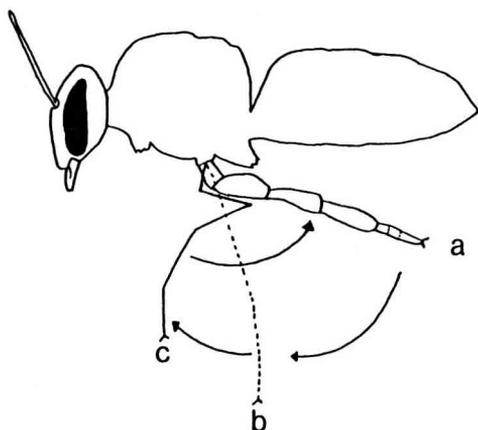


Fig 2. Schematic representation showing the motion of the male mid-leg during courtship (other legs omitted for clarity). The outlined leg (A) indicates the starting position, while the dashed and solid lines (B and C) represent intermediate positions.

species-specific modifications on this sternum in numerous species of *Nomia*; see refs in *Introduction*). A series of 4–6 taps (= one drumming bout) was delivered rapidly in succession by the male's metasoma. These taps were forceful, as evidenced by the displacement of the

female's body following contact; in slow motion it is obvious that each tap did not land in the same place, and instead the male moved his metasoma to the left and to the right of center.

Following a "drumming bout", the male produced a loud "buzz" (600–900 Hz), presumably from the flight muscles (fig 3, bottom) (not reported by C and B); during this time the male had his mandibles open and in contact with the female's scutum or scutellum (not reported by C and B). After these vibrations, the metasoma again tapped the female once or twice, followed by a new drumming bout. This "drum-buzz-taps" sequence was repeated up to 114 times (C and B: the pre-copulatory mating dance lasted "several minutes"). The final sequences increased in intensity, such that drumming bouts were often accompanied by wing fluttering (C and B: during a mating dance a male... "*flutters his wings, at the same time drumming his abdomen rapidly against that of the female*").

After being courted, a female indicated receptivity by slightly raising her metasoma, and the male then fluttered his wings,

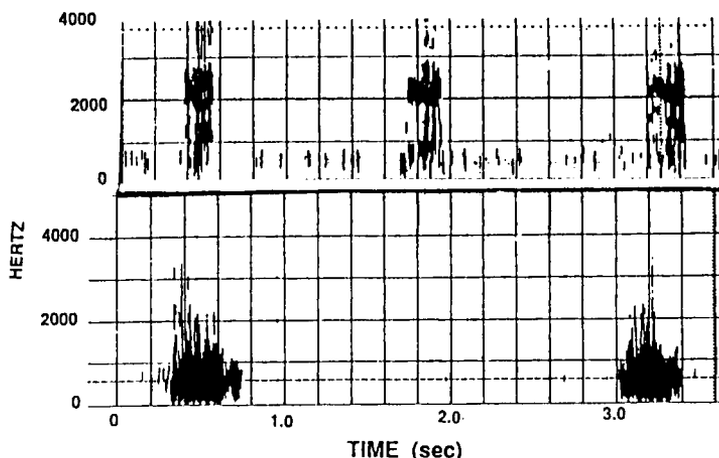


Fig 3. Bottom: representative sonogram of the pre-copulatory vibrations of male *Nomia triangulifera*. Top: representative sonogram of the copulatory vibrations of male *Nomia triangulifera*.

and inserted an inflated membranous eversible sac (Roig, unpublished observations). During copulation the male made no other courtship motions, except for periodic contractions of the abdomen each of which occurred with a loud buzz ($> 1\ 200$ Hz) of higher frequency than the previously described buzzing that occurred before copulation (fig 3, top) (not reported by C and B; these authors reported that during copulation the male "strokes the female with his antennae at a rate of about one stroke per second"). After copulating, a male released his hind legs and then flew away. Unreceptive females, in contrast, curled their metasoma downward and forward, so that the males were unable to insert their genitalia; females often bit at the males' legs until they left.

The duration of precopulatory courtship lasted from 37–149 s ($\bar{x} = 70.58$, $SD = 27.47$, $N = 55$), although our homosexual courtings were much briefer (< 10 s). The duration of actual copulation was 9.95 s ($SD = 4.32$). For mating pairs collected at the same temperatures (± 1 °C), there was a significant positive correlation between the duration of the courtship and the duration of the copulation ($r = 0.64$, $N = 19$). The durations of precopulatory behavior were not different for males with receptive or unreceptive females ($P > 0.2$, Mann–Whitney U test).

Females were allowed to court, or to court and mate, and then given subsequent opportunities to mate again. Table I shows that already unreceptive females, as well as recently mated ones, were not receptive to subsequent males, at least under artificial conditions. Females receptive to the first male, but not allowed to mate with him, were subsequently receptive to a second male. Such bioassays suggest that many females do not mate multiple times, although some do. Dissections of receptive and unreceptive females support this interpretation. Three of 11 pre-

Table I. The relationship between the receptivity of female *Nomia triangulifera* and prior sexual behavior.

Sexual experience with first male	Female mates with second male	
	Yes	No
Mated with 1st male	4	30
Unreceptive to 1st male	3	15
Receptive to 1st male: separated before intromission	22	4

viously mated females (having sperm in their spermathecae) were receptive to males, and 12 of 14 unreceptive females had sperm.

DISCUSSION

The courtship of *Nomia (Epinomia) triangulifera* involves extraordinarily complex mating behavior for bees (see Alcock *et al*, 1978; Eickwort and Ginsberg, 1980), and presumably is necessary for successful intromission. None of the unusual behaviors seen in *N (E) triangulifera* were reported for *N (E) nevadensis* (O'Neill and Bjostad, 1987; O'Neill, personal communication), *N (Acunomia) melanderi* (Johansen and Mayer, 1976), or *N (Curvinomia) tetrazonata tetrazonata* (Wcislo, unpublished observations). Males of *N (Dieunomia) heteropoda* occasionally wave their antennae before intromission, and sometimes move their metasoma in a way which might suggest tapping, but might be reaching motions for the female's genital opening (Wcislo, unpublished observations). In *N(E) nevadensis*, the initial stages of mating take place underground, if at all. "Sweet-smelling" females often emerge from the soil with males mounted on them, having

their hind legs wrapped around the female's anterior metasoma (O'Neill and Bjostad, 1987). Intromission is accompanied by rhythmic pulsations of the male's metasoma, and often the female is released from the leg grasp, as is sometimes true for *N (D) heteropoda* (Wcislo, unpublished observations). The total duration of copulation for *N (E) nevadensis* is about twice as long as for *N (E) triangulifera*. In *N (E) nevadensis* mating is followed by a brief (≈ 22 s, on average) period when the male remains mounted on the female, but has withdrawn his aedeagus (O'Neill and Bjostad, 1987). After intromission, males of *N (E) triangulifera* never remained with their mates.

The use of the fore- or mid-legs for stroking females is not common in the Apoidea, and to date is unknown in other Halictidae (Eberhard, 1991; Low and Wcislo, 1992), including other species of *Nomia*. Structural modifications of male hind legs are well-known in orchid bees (Euglossinae), but they function in chemical communication, as with some male *Centris* bees (eg Coville *et al*, 1986). Expansions of the hind legs occur in many other species of Nomiinae, as well as some other Halictidae (see refs in the *Introduction*), Colletidae (Xeromelissinae) (Toro and Moldenke, 1979), Fideliidae (Rozen, 1970), the anthrophorid genus *Ancyloscelis* (Wcislo, unpublished observations) and in sphecid wasps such as *Dynastus* (Bohard and Menke, 1976). The hind legs of the colletids and fideliids that have been examined contain large flexor muscles and no sign of glandular material (Toro and Magunacelaya, 1987), as for several species of *Nomia* (Wcislo, unpublished observations). Among North American *Nomia*, claspers on the hind legs can be feebly developed (eg *N (E) boharti*, or well-developed (eg *N (D) heteropoda*)), but there is not enough information to correlate the development of such structures with allometry, or with particular mating behavior.

Several analogous elements of the courtship behavior of *N (E) triangulifera* also occur in other Apoidea, including the Halictidae (partial review in Eberhard, 1991). Rhythmic motions of the antennae, for example, are not uncommon, as variations on a "waving", "stroking", or "tapping" theme. Danks (1971), among others, suggested that these motions function to immobilize females. The antennae of *N (E) triangulifera* are not modified other than the usual sex differences in bees, in contrast to the broadly flattened terminal flagellomere of other nomiine bees such as *N (D) heteropoda* or *Spatunomia*. In *N (C) tetrazonata* the antennae are usually motionless before intromission, while in *N (D) heteropoda* there is much individual variation (Wcislo, in preparation). Antennae are rhythmically moved up and down during copulation in the latter two species (Wcislo, unpublished observations) and Cross and Bohart (1960) described antennal stroking behavior for the Utah population of *N (E) triangulifera*.

Wing vibrations or stridulations in courtship occur in some other Halictidae, and various other groups of bees (eg Rozen, 1977; Alcock and Buchmann, 1985; Larsen *et al*, 1986; Tengö *et al*, 1988; Wcislo, unpublished observations) as well as other Hymenoptera (eg Markl, 1977; Sivinski and Webb, 1989). In *N (E) triangulifera* these pre-copulatory vibrations are at a frequency of ≈ 600 Hz.

Curiously, a similar frequency artificially applied to honey bee (*Apis*) combs induces workers nearby to become akinetic (Spangler, 1969 and refs therein). Male *N (D) heteropoda*, produce very low frequency vibrations before copulating, yet provide no further stimulation. Different frequencies may simply be related to body size (Spangler and Buchmann, 1991).

The rapid metasomal drumming was always performed by male *N (E) triangulifera*.

ra, but its function is unknown. Abdominal musculature and morphology of the sternal glands of male *N (A) melanderi* have been studied by Youssef (1969). Females of some species (*N triangulifera* and *N heteropoda*) employ such tapping behavior to collect pollen from the open heads of sunflower with their metasomal sterna (Wcislo, unpublished observations). The use of the metasoma by males during courtship to tap or rub the female metasoma occurs sporadically in some aculeates (eg Cowan, 1986; Wcislo *et al*, 1988). It also sometimes occurs after copulation in *Triepeolus* (Wcislo, unpublished observations), and in *Centris pallida* such post-copulatory behavior reduces females' receptivity to re-mating (Alcock and Buchmann, 1985). Drumming behavior during courtship occurs independently in other orders of insects (eg Plecoptera).

The differences between the behavior of the Kansas bees reported here and those of an Utah population (Cross and Bohart, 1960) are especially striking. If substantiated, they then represent a documented case of intra-specific divergence in copulatory courtship behavior in insects (Eberhard, 1991). Cross and Bohart did not deposit voucher specimens, but Cross (1958), who most recently revised the subgenus *N (Epinomia)*, has no doubt that the species determination is correct (Cross, personal communication). Furthermore, specimens from the Utah locality in the Snow Entomological Museum, University of Kansas, are not obviously different from other *N (E) triangulifera*. The movements of the legs and antennae are very noticeable, as are the precopulatory buzzes, which are audible from up to 1 m. It is, therefore, highly probable that Cross and Bohart would have noticed and reported such unusual behavior for bees, especially since they described the positions of the legs, antennae, etc.

If supported, these intra-specific behavioral differences may support recent ideas

that sexual selection leads to rapid character evolution (refs in *Introduction*). At present, however, there are insufficient data to determine the evolutionary context of such elaborate courtship behavior (eg species recognition or sexual selection – see discussion in Verrell, 1988). Interpopulational differences in courtship behavior have been rarely documented among insects (eg Carson and Bryant, 1979; Spiess and Carson, 1981; Bieman, 1986). Further comparative studies are needed to substantiate intra-specific differences in the courtship behavior of *N (E) triangulifera*. More detailed studies are also needed within populations because results only report information collected when temperatures were hot (> 30°) and relative humidity high. These are the usual weather conditions in Kansas in August; occasionally there are cool, wet years, but no samples have been analyzed. Larsson (1989a,b) has shown that microclimate differentially influences the reproductive success of small and large individuals of *Colletes* (Colletidae).

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Résumé — Comportement de cour précopulatoire chez une abeille solitaire, *Nomia triangulifera* Vachal (Hymenopte-

ra, Halictidae). L'abeille terricole, *Nomia triangulifera*, présente un comportement de cour complexe avant et pendant l'accouplement. Le comportement d'accouplement a été étudié sur de grosses aggrégations dans une ferme du Nord-Est du Kansas, où les abeilles nidifient depuis de nombreuses années. En 1989 et 1990, on a capturé des abeilles qui émergeaient afin de déterminer la phénologie de l'émergence et d'estimer le sex-ratio journalier d'accouplement (nombre de mâles présents / nombre de femelles réceptives). La population est protandre (les mâles apparaissent en premier) et le sex-ratio est toujours très en faveur des mâles (fig 1). Ceux-ci sont légèrement plus petits que les femelles et ils patrouillent en grand nombre au-dessus des endroits où les femelles émergent et autour des plantes où elles récoltent du pollen (*Helianthus annuus*, tournesol). La taille (distance intertégulaire) des mâles qui patrouillent dans ces endroits et de ceux récoltés pendant qu'ils s'accouplaient n'était pas significativement différente. Au champ, les accouplements ont été filmés avec une caméra Bolex 16 mm (4 h), enregistrés sur bande vidéo avec une caméra Panasonic (3,5 h) et des enregistrements sonores ont été faits avec un microphone standard et un magnétophone Uher (7 h). L'analyse de l'ensemble de ces enregistrements montre un comportement de cour extraordinairement complexe pour une abeille. Pendant la cour, le mâle utilise ses pattes antérieures pour tapoter le pronotum de la femelle (fig 1); les pattes médianes caressent la région pleurale de la femelle (fig 2); son metasoma (abdomen) frappe rapidement celui de la femelle; ses antennes oscillent rythmiquement de haut en bas; les muscles du vol produisent un fort bourdonnement audible d'environ 600 Hz de fréquence (fig 3). Les pattes postérieures sont étendues et leur tibia est utilisé pour saisir la femelle. Pendant l'accouplement, le mâle produit une série de

bourdonnements de fréquence plus élevée (environ 1 200 Hz), qui coïncident avec les contractions de son metasoma (fig 3). Des femelles maintenues en cage se sont rarement accouplées plus d'une fois (tableau I). Les femelles réceptives n'avaient généralement pas de sperme dans leur spermathèque, contrairement aux femelles non réceptives qui, elles, en avaient. La comparaison du comportement des mâles de la population du Kansas avec des observations faites précédemment sur une population de l'Utah montre des différences intraspécifiques frappantes dans le comportement de cour de *N. triangulifera*. Si ces résultats sont confirmés lors de nouvelles études, ils constituent le premier cas établi de divergence intraspécifique du comportement d'accouplement chez les insectes.

***Nomia triangulifera* / Halictidae / comportement d'accouplement / sélection sexuelle**

Zusammenfassung — Balzverhalten vor der Kopula bei einer solitären Biene, *Nomia triangulifera* Vachal (Hymenoptera: Halictidae). Vor und während der Paarung einer solitären, bodennistenden Furchenbiene, *Nomia triangulifera*, erfolgt ein komplexes Balzverhalten. Das Paarungsverhalten wurde an großen Ansammlungen auf einer Farm im Nordosten von Kansas studiert, wo die Bienen schon seit vielen Jahren nisten. Die Populationen sind protandrisch (das heißt die Männchen erscheinen zuerst) und das Geschlechtsverhältnis während der Paarung ist immer sehr stark zugunsten der Männchen verschoben. Die Männchen sind etwas kleiner als die Weibchen, und sie patrouillieren in großer Zahl über der Ansammlung, wo die Weibchen schlüpfen und rund um die Pflanzen (*Helianthus annuus*, Sonnenblume), wo diese Pollen sammeln. Die Größe (interteguläre

Distanz) der Männchen, die an diesen beiden Orten patrollierten, war von derjenigen der bei der Paarung gefangenen Männchen nicht signifikant verschieden. Während der Paarung wurden die Bienen im Feld mit einer Bolex 16 mm Kamera gefilmt (4 Stunden), auf Videoband mit einer Panasonic Videokamera aufgenommen (3,5 Stunden) und die dabei hörbaren Geräusche mittels Standardmikrofon und einem Uher Tonbandgerät registriert.

Die Analysen dieser Filme und Bänder ergaben ein für eine Biene äußerst komplexes Balzverhalten. Wie in Abbildung 1 zusammengefaßt, klopf das Männchen während des Balzens mit den Vorderbeinen auf das Pronotum des Weibchens; die Mittelbeine des Männchens streichen über die Pleuralregion des Weibchens, (Abbildung 2); sein Metasoma (Abdomen) trommelt oder klopft rasch auf ihr Abdomen; seine Antennen schwingen rhythmisch auf- und abwärts; und seine Flugmuskeln erzeugen laute, hörbare Tonstöße mit einer Frequenz von ungefähr 600 Hz (Abbildung 3). Die Hinterbeine haben verbreiterte Tibien, die zum Ergreifen des Weibchens benutzt werden. Während der Paarung erzeugt das Männchen eine Serie von Tonstößen höherer Frequenz (bis 1200 Hz), gleichzeitig mit Kontraktionen seines Metasomas (Abbildung 3). Bei Käfighaltung paarten sich Weibchen nur selten mehr als einmal (Tabelle I). Paarungsbereite Weibchen hatten nur selten Samen in ihrer Spermatheka, während bei paarungsunwilligen Weibchen gewöhnlich Samen gefunden wurden.

Vergleiche des Verhaltens der Männchen in der Population von *N. triangulifera* von Kansas mit früheren Beobachtungen an einer Population derselben Art in Utah zeigen Unterschiede im Balzverhalten innerhalb der Art.

Wenn diese Befunde durch noch eingehendere Studien bestätigt werden können, so ist dies der erste dokumentier-

te Fall von Unterschieden im Paarungsverhalten innerhalb ein und derselben Art bei Insekten.

***Nomia triangulifera* / Halictidae / Paarungsverhalten / Sexuelle Selektion**

REFERENCES

- Alcock J, Buchmann SL (1985) The significance of post-insemination display by male *Centris pallida* (Hymenoptera: Anthophoridae). *Z Tierpsychol* 68, 231-243
- Alcock J, Gwynne DT (1991) Evolution of insect mating systems: the impact of selectionist thinking. In: *Reproductive Behaviour of Insects: Individuals and Populations* (Bailey WJ, Ridsdill-Smith J, eds) Chapman, London
- Alcock J, Barrows EM, Gordh G, Hubbard LJ, Kirkendall LL, Pyle D, Ponder TL, Zalom FG (1978) The ecology and evolution of male reproductive behaviour in the bees and wasps. *Zool J Linn Soc* 64, 293-326
- Barrows EM (1975) Mating behavior in halictine bees (Hymenoptera: Halictidae): III. Copulatory behavior and olfactory communication. *Insectes Soc* 22, 307-322
- Batra SWT (1966) The life cycle and behavior of the primitively social bee, *Lasioglossum zephyrum* (Halictidae). *Univ Kansas Sci Bull* 64, 359-423
- Bieman CFM den (1986) Acoustic differentiation and variation in planthoppers of the genus *Ribatodelphax* (Homoptera, Delphacidae). *Netherlands J Zool* 36, 461-480
- Bohart RM, Menke AS (1976) *Sphecoid Wasps of the World*. University of California Press, Berkeley, CA
- Carson HL, Bryant PJ (1979) Change in a secondary sexual character as evidence of incipient speciation in *Drosophila silvestris*. *Proc Natl Acad Sci USA* 76, 1929-1932
- Coville RE, Frankie GW, Buchmann SL, Vinson SB, Williams HJ (1986) Nesting and male behavior of *Centris heithausi* (Hymenoptera: Anthophoridae) in Costa Rica with chemical analysis of the hindleg glands of males. *J Kansas Entomol Soc* 59, 325-336
- Cowan DP (1986) Sexual behavior of eumenid wasps. *Proc Entomol Soc (Wash)* 88, 531-541

- Cross EA (1958) A revision of the bees of the subgenus *Epinomia* in the New World (Hymenoptera: Halictidae). *Univ Kansas Sci Bull* 38, 1261-1301
- Cross EA, Bohart GC (1960) The biology of *Nomia* (*Epinomia*) *triangulifera* with comparative notes on other species of *Nomia*. *Univ Kansas Sci Bull* 41, 761-792
- Danks HV (1971) Biology of some stem-nesting aculeate Hymenoptera. *Trans R Entomol Soc (Lond)* 122, 323-399
- Darwin C (1871) *The Descent of Man and Selection in Relation to Sex* (1986 reprint). Princeton University Press, Princeton
- Eberhard WG (1985) *Sexual Selection and Animal Genitalia*. Harvard University Press, Cambridge
- Eberhard WG (1991) Copulatory courtship and cryptic female choice in insects. *Biol Rev* 66, 1-31
- Eickwort GC, Ginsberg HS (1980) Foraging and mating behavior in Apoidea. *Annu Rev Entomol* 25, 421-446
- Johansen C, Mayer D (1976) Alkali bees: their biology and management for alfalfa seed production in the Pacific Northwest. *Pac Northwest Ext Publ* 155, 1-19
- Larsen O, Gleffe G, Tengö J (1986) Vibration and sound communication in solitary bees and wasps. *Physiol Entomol* 11, 287-296
- Larsson F (1989a) Insect mating patterns explained by microclimatic variables. *J Therm Biol* 14, 155-157
- Larsson F (1989b) Mating patterns in six insect species: effects of weather and population density. *Acta Univ Ups Abstr Upps Dis Sci* 234, 1-31
- Low BS, Wcislo WT (1992) Foretibial plates and the mating behavior of *Crabo cribrellifer* (Hymenoptera: Sphecidae), with a review of male foreleg modifications in Apoidea. *Ann Entomol Soc Am*
- Markl H (1977) Mating behavior and sound production in harvester ants (*Pogonomyrmex*, Formicidae). *Insectes Soc* 24, 191-212
- Michener CD (1965) A classification of the bees of the Australian and South Pacific regions. *Bull Am Mus Nat Hist* 130, 1-362
- Moure JS CMF, Hurd PD Jr (1987) *An Annotated Catalog of the Halictid Bees of the Western Hemisphere* (Hymenoptera: Halictidae). Smithsonian Institution Press, Washington, DC
- O'Neill KM, Bjostad L (1987) The male mating strategy of the bee *Nomia nevadensis* (Hymenoptera: Halictidae): leg structure and mate guarding. *Pan-Pac Entomol* 63, 207-217
- Ribble DW (1965) A revision of the banded subgenera of *Nomia* in America (Hymenoptera: Halictidae). *Univ Kansas Sci Bull* 45, 277-358
- Richards OW (1927) Sexual selection and allied problems in insects. *Biol Rev* 2, 298-360
- Rozen JG Jr (1970) Biology, immature stages, and phylogenic relationships of fideline bees, with the description of a new species of *Neofidelia* (Hymenoptera, Apoidea). *Am Mus Novit* 2427, 1-25
- Rozen JG Jr (1977) Biology and immature stages of the bee genus *Meganomia* (Hymenoptera, Melittidae). *Am Mus Novit* 2630, 1-14
- Sivinski J, Webb JC (1989) Acoustic signals produced during courtship in *Diachasmimorpha* (= *Biosters*) *longicaudata* (Hymenoptera: Braconidae) and other Braconidae. *Ann Entomol Soc Am* 82, 116-120
- Sokal RR, Rohlf FJ (1981) *Biometry*. WH Freeman and Co, New York, NY, 2nd edn
- Spangler HG (1969) Suppression of honey bee flight activity with substrate vibration. *J Econ Entomol* 62, 1185-1186
- Spangler HG, Buchmann SL (1991) Effects of temperature on wingbeat frequency in solitary bee *Centris caesalpinae* (Anthophoridae: Hymenoptera). *J Kansas Entomol Soc* 64, 107-109
- Spieß EB, Carson HL (1981) Sexual selection in *Drosophila silvestris* of Hawaii. *Proc Natl Acad Sci USA* 78, 3088-3092
- Tengö J, Eriksson J, Borg-Karlson AK, Smith BH, Dobson H (1988) Mate-locating strategies and multimodal communication in male mating behavior of *Panurgus banksianus* and *P calceratus* (Apoidea, Andrenidae). *J Kansas Entomol Soc* 61, 388-395
- Toro H, Moldenke A (1979) Revision de los *Xeromellisinæ* chilenos (Hymenoptera, Colletidae). *Ann Mus Hist Nat Valp* 12, 95-182
- Toro H, Magunacelaya JC (1987) Estructura muscular femoral de *Xeromellisinæ* (Hymenoptera: Colletidae). *Acta Entomol Chil* 14, 13-24

- Triplett DC, Gittins AR (1988) Nesting, mating and foraging habits of *Melissodes* (*Melissodes*) *tepida tepida* Cresson in Idaho (Hymenoptera: Anthophoridae). *Proc Entomol Soc (Wash)* 90, 462-470
- Verrell PA (1988) Stabilizing selection, sexual selection and speciation: a view of specific-mate recognition systems. *Syst Zool* 37, 209-215
- Wcislo WT (1992) Attraction and learning in the mate-finding behavior of solitary bees, *Lasioglossum* (*Dialictus*) *figueresi* Wcislo and *Nomia triangulifera* Vachal (Hymenoptera: Halictidae). *Behav Ecol Sociobiol* 31, 139-148
- Wcislo WT, West-Eberhard MJ, Eberhard WG (1988) Natural history and behavior of a primitively social wasp, *Auplopus semialatus*, and its parasite, *Irenangelus eberhardi* (Hymenoptera: Pompilidae). *J Insect Behav* 1, 247-260
- West-Eberhard MJ (1984) Sexual selection, competitive communication, and species-specific signals in insects. *In: Insect Communication* (Lewis TN, ed) Academic Press, New York, NY
- West-Eberhard MJ (1991) Sexual selection and social behavior. *In: Man and Beast Revisited* (Tiger L, Robinson MH, eds) Smithsonian Institution Press, Washington, DC
- Yousseff NN (1969) Musculature, nervous system and glands of metasomal abdominal segments of the male of *Nomia melanderi* Ckll (Hymenoptera, Apoidea). *J Morphol* 129, 59-80