

# Relative Effects of Juvenile and Adult Environmental Factors on Mate Attraction and Recognition in the Cricket, Allonemobius socius

Authors: Olvido, Alexander E., Fernandes, Pearl R., and Mousseau, Timothy A.

Source: Journal of Insect Science, 10(90): 1-17

Published By: Entomological Society of America

URL: https://doi.org/10.1673/031.010.9001

The BioOne Digital Library (<a href="https://bioone.org/">https://bioone.org/</a>) provides worldwide distribution for more than 580 journals and eBooks from BioOne's community of over 150 nonprofit societies, research institutions, and university presses in the biological, ecological, and environmental sciences. The BioOne Digital Library encompasses the flagship aggregation BioOne Complete (<a href="https://bioone.org/subscribe">https://bioone.org/archive</a>), the BioOne Complete Archive (<a href="https://bioone.org/archive">https://bioone.org/archive</a>), and the BioOne eBooks program offerings ESA eBook Collection (<a href="https://bioone.org/esa-ebooks">https://bioone.org/esa-ebooks</a>) and CSIRO Publishing BioSelect Collection (<a href="https://bioone.org/csiro-ebooks">https://bioone.org/esa-ebooks</a>)

Your use of this PDF, the BioOne Digital Library, and all posted and associated content indicates your acceptance of BioOne's Terms of Use, available at <a href="https://www.bioone.org/terms-of-use">www.bioone.org/terms-of-use</a>.

Usage of BioOne Digital Library content is strictly limited to personal, educational, and non-commmercial use. Commercial inquiries or rights and permissions requests should be directed to the individual publisher as copyright holder.

BioOne is an innovative nonprofit that sees sustainable scholarly publishing as an inherently collaborative enterprise connecting authors, nonprofit publishers, academic institutions, research libraries, and research funders in the common goal of maximizing access to critical research.



## Relative effects of juvenile and adult environmental factors on mate attraction and recognition in the cricket, Allonemobius socius

Alexander E. Olvido<sup>1,3a\*</sup>, Pearl R. Fernandes<sup>2b</sup>, and Timothy A. Mousseau<sup>3c</sup>

<sup>1</sup>Division of Science, Gainesville State College (Oconee campus), 1201 Bishop Farms Parkway, Watkinsville, Georgia, 30677, USA

<sup>2</sup>Division of Science, Mathematics and Engineering, University of South Carolina Sumter, Sumter, South Carolina, 29150, USA

<sup>3</sup>Department of Biological Sciences, University of South Carolina – Columbia, Columbia, South Carolina, 29208, USA

#### **Abstract**

Finding a mate is a fundamental aspect of sexual reproduction. To this end, specific-mate recognition systems (SMRS) have evolved that facilitate copulation between producers of the mating signal and their opposite-sex responders. Environmental variation, however, may compromise the efficiency with which SMRS operate. In this study, the degree to which seasonal climate experienced during juvenile and adult life-cycle stages affects the SMRS of a cricket, Allonemobius socius (Scudder) (Orthoptera: Gryllidae) was assessed. Results from two-choice behavioral trials suggest that adult ambient temperature, along with population and family origins, mediate variation in male mating call, and to a lesser extent directional response of females for those calls. Restricted maximum-likelihood estimates of heritability for male mating call components and for female response to mating call appeared statistically nonsignificant. However, appreciable "maternal genetic effects" suggest that maternal egg provisioning and other indirect maternal determinants of the embryonic environment significantly contributed to variation in male mating call and female response to mating calls. Thus, environmental factors can generate substantial variation in A. socius mating call, and, more importantly, their marginal effect on female responses to either fast-chirp or long-chirp mating calls suggest negative fitness consequences to males producing alternative types of calls. Future studies of sexual selection and SMRS evolution, particularly those focused on hybrid zone dynamics, should take explicit account of the loose concordance between signal producers and responders suggested by the current findings.

Keywords: heritability, maternal effect, mating call, Orthoptera, phonotaxis, reaction norm

**Abbreviations: SMRS**, specific-mate recognition systems; *crat*, chirp rate; *ppc*, pulses per chirp; *cdur*, chirp duration; *dfrq*, dominant frequency; *init*, initial choice; *Ingr*, loitering behavior; **MTDFREML**, multiple trait derivative-free restricted maximum likelihood

Correspondence: a\* aolvido@gsc.edu, b pefernan@uscsumter.edu, c mousseau@sc.edu, \* Corresponding author Associate Editor: H. Fred Nijhout was editor of this paper.

Received: 28 May 2008, Accepted: 11 June 2009

**Copyright:** This is an open access paper. We use the Creative Commons Attribution 3.0 license that permits unrestricted use, provided that the paper is properly attributed.

ISSN: 1536-2442 | Vol. 10, Number 90

#### Cite this paper as:

Olvido AE, Fernandes PR, Mousseau TA. 2010. Relative effects of juvenile and adult environmental factors on mate attraction and recognition in the cricket, *Allonemobius socius*. *Journal of Insect Science* 10:90 available online: insectscience.org/10.90

#### Introduction

A well-supported idea for mate discrimination is avoidance of unfit hybrids. In birds, for example, genetically based courtship display traits may have evolved to prevent heterospecific mating that likely yields hybrid offspring with lower-thanaverage fitness (Dobzhansky 1937). While appearing to explain many contemporary reproductive patterns of isolation, Dobzhansky's theory of isolating mechanisms sheds little light on sexual selection and its possible role in speciation, especially in the common case where related species overlap in their geographic distributions (Andersson 1994).

In contrast, the specific-mate recognition concept puts forth the notion that secondary sexual traits evolved to promote the pairing of compatible genotypes, e.g. coordination signal and of mating intraspecific responders (Paterson 1985). Hence, behaviors such as courtship displays can signal sexual readiness to genetically compatible individuals in a population, with the probable result that such matings produce high-fitness offspring. Unlike the avoidance-of-unfit-hybrids explanation, recognition specific-mate provides conceptually straight-forward framework for studying sexual selection within a species and initiation of the speciation process (Andersson 1994).

The southern ground cricket, *Allonemobius socius* (Orthoptera: Gryllidae), presents opportunities for investigating the evolution of a specific-mate recognition system (SMRS). A small (less than 2 cm in anterior-posterior length) terrestrial insect, *A. socius* inhabits fields and woodlands throughout the southeastern region of North America (Howard and Furth 1986).

Allozyme studies indicated A. socius forms part of a complex of two sister species meeting in a hybrid zone from southern New Jersey through Illinois (~40° latitude) (Howard and Furth 1986; Howard and Waring 1991). While genital morphology and mating behavior prove useful in characterizing other insects (Walker 1957; Lloyd 1984; Bonduriansky 2001), no such clear distinctions exist between A. socius and its more northern congener, A. fasciatus (Howard and Furth 1986; Veech et al. 1996). Currently, the only quick and reliable method to distinguish the two species in the wild is collecting by geographic site (Howard and Furth 1986; DJ Howard New Mexico State University (Las Cruces, NM), personal communication).

Given its widespread North American distribution, A. socius varies substantially in development rate, morphology, reproductive behavior (Mousseau and Roff 1989, 1995). Northern A. socius populations produce one generation per year, while more southern populations produce two or more (Howard and Furth 1986; Walker and Masaki 1989; Mousseau and Roff 1989; Mousseau 1991). Like in other gryllids, A. socius males stridulate, or rub their forewings together, to emit a chirp-like mating call that functions as a signal to nearby females of the male's readiness to mate. While the immediate effects of ambient temperature on insect mating calls are well-known, i.e. since Brooks (1882) and Dolbear (1897), only recently have researchers begun to explore environmental variation during the juvenile stages of the life cycle might shape reproductive behavioral reaction norms, e.g. degree to which male mating call varies systematically with ambient temperature

(Whitesell and Walker 1978; Olvido and Mousseau 1995; Grace and Shaw 2004).

Environmental variation might also affect female responses to male mating call. Olvido and Wagner (2004) showed that A. socius females preferred experimentally manipulated long-chirp mating calls, i.e. those with above-average chirp duration, and paid surprisingly little attention to variation in chirp rate, i.e. the number of chirps per seconds in a mating call. Since chirp duration and other components of A. male mating call vary with socius temperature and rearing environment (Olvido and Mousseau 1995), it is quite possible that female preferences also vary with ambient temperature and/or rearing environment.

In this study, the effects of genetic and nongenetic factors on the SMRS of A. socius were examined as a continuing effort to assess, ultimately, their contributions to this species' persistent hybridization with A. fasciatus. the hypothesis of environmentally mediated phenotypic and genetic coupling between male mating call and relative preference of females for mating call was tested by, first, rearing split broods of A. socius juveniles in different laboratory "seasonal" environments and, later, analyzing their sex-specific behaviors across different ambient temperatures. If, indeed, the current A. fasciatus-A. socius hybrid zone is maintained primarily by A. socius females migrating northward into A. populations fasciatus and mating preferentially with A. fasciatus males, then a weakly evolved SMRS would be expected in A. socius, i.e. weak or absent phenotypic and/or genetic coupling between A. socius mating call and female relative preference for mating call, all else being equal. However, if the A. fasciatus-A. socius hybrid zone persists mainly for reasons

other than promiscuous *A. socius* females mating with heterospecifics, then significant phenotypic and/or genetic coupling between *A. socius* mating call and female relative preference for mating call would be expected.

#### **Methods**

#### Field collection and animal husbandry

Cricket stocks were derived from individuals collected from two sites located approximately 170 km apart (in Columbia and Travelers' Rest, South Carolina, USA). Gravid field-caught females were housed singly and allowed to oviposit ad libitum in cheesecloth. Field-caught juveniles were reared in several small-group cages and maintained in the laboratory at 31° C with a 15-hr daily light cycle, or DLC. From each maternal line (established either from fieldcaught gravid females or from singly paired males and females that had matured under laboratory conditions). the following generation was reared exclusively at 31° C and a 15-hr DLC to minimize any confounding genotype-by-environment interactions subsequent phenotypic in analyses.

For the second laboratory-reared generation (i.e. two generations removed from the field), one-half of a brood of newly hatched nymphs was reared exclusively under spring-like conditions (24° C, 11-hr DLC) small-group several cages simultaneously, the other half exclusively under summer-like conditions (31° C, 15-hr DLC) in several small-group cages. As in previous generations. females were separated from males before the penultimate juvenile stage to assure virginity of subjects in the ensuing behavioral assays. Adult crickets were maintained temperatures at and photoperiods of their respective "juvenile"

environments through all experimental trials.

#### Measuring cricket sexual behaviors

Each male cricket was allowed 5-20 minutes to acclimate to a particular ambient temperature before its continuously produced mating call was recorded at that temperature in an echo-dampened chamber. In total, 424 males (1-5 weeks from adult emergence) were recorded calling at 24°, 28°, and 31° C. A haphazardly chosen 10second sample of each mating call was later imported to a desktop computer as an 8-bit, 22-kHz WAV file and analyzed with "Spectrogram 2.2" (©1994, RS Horne) and "Wave for Windows 2.03" (@1993, Turtle Beach Systems, www.turtlebeach.com ) graphical analysis software. The four mating call components of interest were chirp rate (crat, number of chirps per second), pulses per chirp (ppc, number of acoustic pulses per chirp), chirp duration (cdur, number of seconds from beginning to end of each chirp), and dominant frequency (dfrq, frequency of oscillation at peak power spectral density).

Which mating call to use in phonotaxis trials was determined beforehand via analysis principal components PRINCOMP procedure in SAS/STAT) of the four mating call components mentioned above. For each population, the two unmanipulated mating calls having their first principal component (= PC1, which explained 54% of total variation in mating call) most closely match the mean PC1 of males calling at 24° C and at 31° C ("cold" and "hot" songs, respectively) were chosen. Analysis of eigenvectors indicated that crat and cdur both loaded equally well on PC1. but from opposite directions (-0.505 for and 0.508 for *cdur*), whereas crat eigenvectors for ppc and dfrq were lower (0.382 and -0.386, respectively). Thus, the

chosen stimuli seemed to differ mainly in chirp rate and chirp duration, which were also strongly correlated with each other (r = -0.4989).

Each singly tested female cricket (1-5 weeks after adult emergence) experienced simultaneous playback of "hot" and "cold" call stimuli (standardized to 70 db SPL at approximately 1 m from each speaker) through two three-minute trials — once at 24° C and, several days later, again at 31° C (after a 5- to 20-minute acclimation period for each trial). For each test subject at each ambient temperature, left-right orientation of speakers was randomized in the square (1.44-m<sup>2</sup> floor area) anechoic observation chamber such that a left-corner speaker would broadcast a "hot" mating call in one trial before being switched in a subsequent trial with the other speaker broadcasting a "cold" mating call at the adjacent corner. Given the initially large sample size (> 300 females) and labor-intensive nature of measuring phonotaxis, 3-7 days were allowed between repeated observations. Positive phonotaxis was scored when the test subject approached within 30 cm of a speaker. The freely accessible space between speakers (approximately 45 cm in width) assured that female phonotaxis was measured independently of either call stimulus. Through 1086 playback trials videotaped under low-intensity red light (to minimize any confounding visual cues), 300 females yielded score-able phonotaxis.

Female response to call stimuli was characterized in two different ways. First, phonotaxis was quantified as the inverse of time elapsed (in seconds) for a female test subject to approach its first speaker, i.e. initial choice (*init*): Positive initial-choice scores indicated attraction to the speaker broadcasting (exclusively) a "hot" mating call, while negative scores indicated

attraction to one broadcasting (exclusively) a "cold" mating call, regardless of relative orientation of speakers. Initial choice, thus, appeared to measure a female *A. socius*'s general degree of attentiveness to "hot" and "cold" variations of intraspecific mating call.

Alternatively, female phonotaxis quantified as the difference in time (in seconds) a female test subject loitered between the two speakers playing their respective call stimuli simultaneously: Positive loitering behavior scores indicated greater time spent within 30 cm of the speaker broadcasting exclusively the "hot" mating call, and negative scores indicated greater time spent within 30 cm of the speaker broadcasting exclusively the "cold" regardless mating call, relative orientation of speakers. Thus, loitering behavior (lngr) appeared to measure relative strength of directional preference of females for either the "hot" or "cold" mating call stimulus.

As a matter of clarification, the measurement of relative preferences presumed that, given the audible and quantifiable differences in "hot" versus "cold" call stimuli, our female test subjects would prefer one or the other call stimulus type. The main issue was assessing how malleable a female's relative preference might be, as relative preference may reflect a female's absolute preference for male calls of a specific chirp rate, chirp duration, etc. Assessing the malleability of female absolute preferences or the relative importance for individual mating call components will require a far greater investment of resources than was possible in the current work.

To discount a possible left- versus rightside walking bias in *A. socius* females, each test subject was allowed to wander in the observation chamber without playback of call stimuli. For each test subject, silent trials were conducted in random order with respect to phonotaxis trials and also under red-light conditions, but temperature (25-27° C). The magnitude of walking side bias was quantified as a proportion of the total time within a threeminute observation period that a test subject loitered within 30 cm of either silent speaker  $[(t_{LEFT} - t_{RIGHT})/(t_{LEFT} + t_{RIGHT})].$ Positive scores indicated for left-side wandering bias, and negative scores indicated right-side wandering bias. Females failing to approach either silent speaker within the alloted three minutes were excluded from further analysis (sample size in silent trials,  $n_{TRIAL0}$ , was 114 females).

#### Phenotypic analyses of adult behaviors

To minimize bias from genotype-byenvironment interactions, all phenotypic analyses of reproductive behavior was limited to sexually mature crickets from the laboratory-reared second generation. Borrowing a technique from a van der Waerden normal scores analysis (Conover 1999), all raw variates were transformed to their normalized ranks before performing analysis-of-variance, or ANOVA, testing (see also Olvido and Wagner 2004). Hence, the phenotypic analyses explicitly satisfy two fundamental assumptions of ANOVA models, namely that all treatments for a given factor share a common mean (here, overall  $\mu = 0$ ) and have comparable variance (i.e. overall s<sup>2</sup> approximates unity).

To facilitate analysis of female relative preference for mating call, and as dictated by the repeated-measures design, a crossnested ANOVA was created with one repeated factor:

$$\begin{split} Y_{ijklm} &= \mu_{.....} + Adu_i + Juv_j + Pop_k + Fam_{l(k)} + Ind_{m(kl)} \\ &\quad + (Adu \times Juv)_{ij} + (Adu \times Pop)_{ik} + (Adu \times Fam)_{il(k)} + (Adu \times Ind)_{im(kl)} \\ &\quad + (Juv \times Pop)_{jk} + (Juv \times Fam)_{jl(k)} + (Juv \times Ind)_{jm(kl)} \\ &\quad + (Adu \times Juv \times Pop)_{ijk} + (Adu \times Juv \times Fam)_{ijl(k)} + (Adu \times Juv \times Ind)_{ijm(kl)} + \epsilon_{(ijklm)} \end{split}$$

where Yiiklm indicates the female trait of interest, Adui indicates adult ambient temperature (repeated factor; fixed effect), Juv<sub>i</sub> indicates rearing environment (fixed effect), Popk indicates population origin (random effect), Fam<sub>l(k)</sub> indicates withinpopulation family origin or maternal line (random effect), and Ind<sub>m(kl)</sub> indicates the individual test subject nested within population and family (random effect), and  $\mu_{\text{....}}$  and  $\epsilon_{\text{(iiklm)}}$  indicate the common mean and error term, respectively. Having had recorded female behaviors only once at each ambient temperature, an ANOVA model was constructed that lacked an independent error term. Interaction terms were considered random when involving at least one random main effect. Since repeated-measures ANOVA requires that each test subject complete both phonotaxistemperature trials, data from only 160 females ( $\times$  2 ambient temperatures = 320 repeated observations) of the total 300 test subjects were analyzed.

Female phonotaxis was analyzed without age as a covariate because an earlier study showed consistency in call stimulus preference through 90% of the adult life of *A. socius* females: Three-week old females preferring long-chirp calls still preferred those same calls at 17 weeks of age, with only a marginal decline in time spent near speakers broadcasting those preferred calls (Olvido and Wagner 2004).

The same cross-nested, repeated-measures ANOVA was applied to the analysis of male mating calls. As required of repeated-measures designs, only data from males

completing all temperature treatments ( $N = 266 \text{ males} \times 3 \text{ ambient temperatures} = 798 \text{ observations}$ ) were analyzed.

The GLM procedure was used in SAS/STAT to obtain information ANOVA degrees of freedom and Type III mean squares, from which observed Fvalues were calculated. (Tables 3 and 4 contain explicit description of each F test.) For each observed F-value through the distribution functions, p-values obtained in Stat-SAK 2.14 (©1986, GE Dallal).

#### **Pedigree Analysis**

A series of Fortran-77 programs, known collectively as "Multiple Trait Derivative-Free Restricted Maximum Likelihood," or simply "MTDFREML" (Boldman et al. 1995), were run to evaluate mixed-model equations that partition phenotypic variance specifically into additive-genetic and other model variance components. The particular mixed-model equations model applied is known as the full animal model with no covariates for inbreeding of offspring with dams (most similar to Model 4 in Ferreira et al. (1999)), and can be expressed as:

$$y = X\beta + Zu + S_1v + Wm + S_2n + e$$

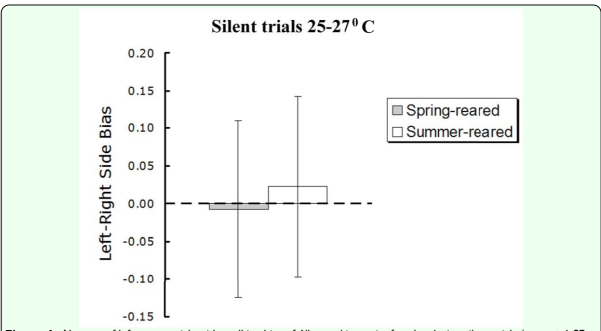
where y represents a vector of observations (for a single trait),  $\beta$  is a vector of fixed effects (which include population origin, adult ambient temperature, and juvenile rearing environment), u indicates a vector of random animal (direct) effects, v indicates a vector of random animal permanent environmental effects, m indicates a vector of random maternal

(indirect) genetic effects, *n* indicates a vector of random maternal permanent environmental effects, *e* indicates a vector of random residual effects, and X, Z, S<sub>1</sub>, W, and S<sub>2</sub> represent association matrices for fixed, random direct, animal permanent environmental, random indirect, and maternal permanent environmental effects, respectively. (An association matrix ties performance data to particular test subjects and treatment groups, as well as specifies familial relationships.)

To document how juvenile rearing environment might affect estimates of heritability and proportional other variances, separate MTDFREML analyses were performed on spring- and summerreared crickets (even though both rearing groups shared the same pedigree). The pedigree was composed of 1.362 individuals (including grandsires, granddams, sires, and dams) and a total of 752 individuals from the second laboratoryreared generation (446 males and 300 females recorded multiple times) in the cumulative performance data set that was sub-sampled to obtain separate quantitativegenetic parameter estimates from springsummer-reared groups. versus Each MTDFREML session stopped when variance of the simplex algorithm, Var (- $2\log\Lambda$ ), reached 1 x  $10^{-6}$ . We presumed convergence at a global maximum when both the simplex values and heritability from point estimates consecutive MTDFREML sessions remained unchanged at the second decimal place (LD Van Vleck, communication, USDA-ARS, personal University of Nebraska at Lincoln). For each proportional variance, the 95% confidence interval was approximated as twice the REML standard error of the point and determined estimate. significance when that interval excluded zero. Ferreira et al. (1999) and Boldman et (1995)provide more descriptions of MTDFREML programs and their correct implementation. The pedigree and phenotypic data file is available in Appendix 1 online, or upon request to the first author.

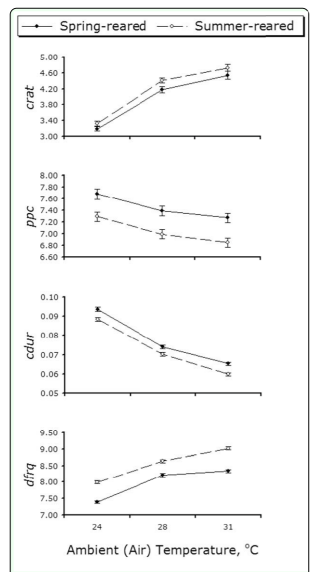
#### Results

Visual analysis of results from the silent



**Figure 1.** Absence of left- versus right-side walking bias of *Allonemobius socius* females during silent trials (mean  $\pm$  1 SE, in seconds). A positive score indicates that a female spent a greater proportion of the three-minute trial period wandering near the right-side corner of the observation chamber ( $n_{TRIALO} = 114$  females). See Methods for full description of measurement protocols. High quality figures are available online

trials indicated no left- or right-side walking bias in female *A. socius* used in this study. Regardless of rearing environment, *A. socius* females were as likely to wander near the left corner of the observation chamber as the right corner: walking scores for both spring- and summer-reared females hovered near zero (Figure 1).



**Figure 2.** Environmental effects on *Allonemobius socius* male mating call (mean ± 1 SE). Full-sibling males were reared in paired treatments as juveniles exclusively under "spring" versus "summer" conditions. Each male's mating call was recorded only once in each of three adult ambient temperatures (n = 266 males). Traits codes are *crat*: chirp rate (i.e. number of chirps per second) of male mating call; *ppc*: number of acoustic pulses per chirp of male mating call; *cdur*: chirp duration in seconds of male mating call; *dfrq*: dominant frequency in kilohertz of male mating call. See Methods for full description of trait codes and measurement protocols. High quality figures are available online.

#### Factors affecting male mating call

Adult ambient temperature clearly affected all four components of male mating call. Chirp rate and dominant frequency of mating call tended to increase with increasing ambient temperature (Figure 2), though two- and three-way interactions with population origin, rearing environment, family origin, and individual suggest non-linear ambient temperature effects on these two male traits (crat and dfrq in Table 1, construction of F-test is given in Table 3). On the other hand, chirp duration and number of pulses per chirp significantly varied only with temperature (Adu on *cdur* and *ppc* in Table 1), indicating that chirp duration and pulses-per-chirp tend to decline uniformly as ambient temperature increases (Figure 2).

Compared with adult ambient temperature, juvenile rearing environment considerably less effect on male mating call. Only pulses per chirp registered significant variation due to juvenile rearing environment (Juv on ppc in Table 1, construction of F-test in Table 3). The twointeractions of juvenile rearing environment with adult ambient temperature and with population origin (Adu x Juv and Juv x Pop) appeared statistically non-significant, which supports consistently higher number of pulses per chirp in spring- versus summer-reared males (Figure 2). Moreover, the significant two-way interaction of juvenile rearing environment and individual on dominant frequency and chirp rate indicated that individuals within families varied nonlinearly across the two rearing environments with respect to these two traits (Juv x Ind on dfrq and crat in Table 1, construction of F-tests in Table 3).

 Table 1. Summary of observed F values from statistical analyses of male mating call and female call preference.

Source of Variation		Mati	Relative			
Source of Variation	crat	ррс	cdur	dfrq	init	Ingr
Among-Subjects Effects						
Juvenile environment (Juv)	0.36	1136.36*	2.61	8.58†		
Population origin (Pop)	0.13	11.33**	18.74***	8.21**	2.66	2.78
Juv x Pop	1.28	0.01	1.91	1.09		
Family origin (Fam)	1.93***	1.82***	1.48*	1.38*	0.92	1.10
Juv x Fam	1.22	0.74	17.48*	1.39	3.90	6.67
Individual (Ind)	1.66	1.73	1.85	2.77*	0.87	0.40
Juv x Ind	8.79**	2.48	0.21	120.98***	0.36	0.46
Within-Subject Effects						
Adult environment (Adu)	51.71*	59.36*	937.31**	388.31**	3.05	112.51‡
Adu x Juv	3.07	3.34	2.82	4.39		
Adu x Pop	3.36*	0.20	0.67	0.97	8.39**	0+
Adu x Juv x Pop	0.56	0.07	0.05	0.05		
Adu x Fam	0.97	1.03	0.95	0.93	1.20	0.87
Adu x Juv x Fam	3.37*	1.45	1.38	52.51***	1.02	0.62
Adu x Ind	4.7*	1.22	1.39	40.93***	0.81	0.46

Degrees of freedom and Type III mean squares for each observed F value are given in Tables 3 and 4.

All traits were transformed to their normalized ranks before analysis.

Please refer to Methods for full description of trait codes and measurement protocols.

Numbers in red indicate statistically significant effect.

---, not available; \*P<0.05; \*\*P<0.01; \*\*\*P<0.001; †P=0.104; ‡P=0.060

Table 2. Restricted maximum likelihood estimates of model variance components.

	nobs	V <sub>P</sub>	h²+SE	m²+SE	c <sub>A</sub> <sup>2</sup> +SE	CM2+SE	e²+SE
"Spring"-Reared							
crat	447	77.028	0.00 <u>+</u> 0.708	0.24 <u>+</u> 0.019	0.00 <u>+</u> 0.361	0.00 <u>+</u> 0.004	0.71 <u>+</u> 0.067
ррс	447	0.839	0.49 <u>+</u> 0.725	0.04 <u>+</u> 0.003	0.00 <u>+</u> 0.369	0.00 <u>+</u> 0.364	0.51 <u>+</u> 0.057
cdur	447	104.229	0.00 <u>+</u> 0.698	0.20 <u>+</u> 0.016	0.18 <u>+</u> 0.357	0.00 <u>+</u> 0.000	0.58 <u>+</u> 0.063
dfrq	447	0.255	0.54 <u>+</u> 0.536	0.01 <u>+</u> 0.000	0.00 <u>+</u> 0.281	0.00 <u>+</u> 0.266	0.47 <u>+</u> 0.053
init	211	6415.209	0.00 <u>+</u> 0.482	0.01 <u>+</u> 0.001	0.01 <u>+</u> 0.296	0.00 <u>+</u> 0.239	0.97 <u>+</u> 0.176
Ingr	211	1842.556	0.00 <u>+</u> 0.623	0.05 <u>+</u> 0.005	0.00 <u>+</u> 0.358	0.00 <u>+</u> 0.000	0.95 <u>+</u> 0.168
"Summer"- Reared							
crat	645	104.998	0.00 <u>+</u> 0.293	0.04 <u>+</u> 0.002	0.15 <u>+</u> 0.159	0.03 <u>+</u> 0.145	0.79 <u>+</u> 0.052
ррс	645	1.235	0.00 <u>+</u> 0.428	0.11 <u>+</u> 0.007	0.21 <u>+</u> 0.225	0.06 <u>+</u> 0.214	0.63 <u>+</u> 0.051
cdur	645	140.068	0.00 <u>+</u> 0.385	0.09 <u>+</u> 0.005	0.22 <u>+</u> 0.203	0.00 <u>+</u> 0.196	0.66 <u>+</u> 0.051
dfrq	645	0.440	0.00 <u>+</u> 0.559	0.24 <u>+</u> 0.278	0.40 <u>+</u> 0.289	0.00 <u>+</u> 0.000	0.36 <u>+</u> 0.036
init	254	4148.592	0.01 <u>+</u> 0.567	0.03 <u>+</u> 0.003	0.00 <u>+</u> 0.328	0.00 <u>+</u> 0.000	0.98 <u>+</u> 0.135
Ingr	254	2341.948	0.03 <u>+</u> 0.481	0.03 <u>+</u> 0.239	0.00 <u>+</u> 0.268	0.00 <u>+</u> 0.000	0.97 <u>+</u> 0.127
All Environments							
crat	1092	93.665	0.00 <u>+</u> 0.258	0.08 <u>+</u> 0.004	0.10 <u>+</u> 0.137	0.05 <u>+</u> 0.130	0.77 <u>+</u> 0.041
ррс	1092	1.063	0.15 <u>+</u> 0.306	0.01 <u>+</u> 0.001	0.18 <u>+</u> 0.162	0.01 <u>+</u> 0.153	0.60 <u>+</u> 0.039
cdur	1092	124.968	0.03 <u>+</u> 0.275	0.07 <u>+</u> 0.138	0.21 <u>+</u> 0.146	0.00 <u>+</u> 0.000	0.64 <u>+</u> 0.040
dfrq	1092	3.591	0.27 <u>+</u> 0.324	0.00 <u>+</u> 0.000	0.27 <u>+</u> 0.172	0.04 <u>+</u> 0.002	0.41 <u>+</u> 0.031
init	465	4449.282	0.02 <u>+</u> 0.189	0.02 <u>+</u> 0.001	0.00 <u>+</u> 0.142	0.00 <u>+</u> 0.000	0.98 <u>+</u> 0.107
Ingr	465	2119.176	0.02 <u>+</u> 0.243	0.03 <u>+</u> 0.002	0.00 <u>+</u> 0.164	0.00 <u>+</u> 0.122	0.97 <u>+</u> 0.099

 $n_{\mbox{\scriptsize OBS}},$  number of single and repeated observations

V<sub>P</sub>, total phenotypic variance

h2, narrow-sense heritability

m<sup>2</sup>, maternal genetic effect

cA2, animal permanent environmental effect

cm², maternal permanent environmental effect

e2, residual environmental effect

Point estimates in red are significantly different from zero.

Within each population, significant effects of family origin were detected on all four call components, though "family effects" on chirp duration indicated a two-way interaction with juvenile rearing environment (Juv x Fam on *cdur* in Table 1, construction of F-tests in Table 3). Quantitative-genetic analysis revealed that, aside from unexplained (residual) variance sources, maternal genetic factors accounted for an appreciable proportion of this effect, particularly in males reared under "spring-like" conditions (m<sup>2</sup> column in Table 2).

None of the male calling song components showed significant heritability (h<sup>2</sup>). Point estimates reached as high as 54% of total phenotypic variation, as in the case for dominant frequency (*dfrq*) in spring-reared males, but were obscured by large standard errors (Table 2).

## Factors affecting female response to mating call

Adult ambient temperature, as a main factor, had no consistent effect on female relative preference. The mostly positive *init* scores across the two ambient (adult) temperatures indicated that females initially associated with the "hot" call stimulus (Figure 3, upper panel). However, variation

in a test subject's initial choice across the two adult temperature treatments was not statistically significant (Adu on init in Table 1, construction of F-tests in Table 4). The interaction significant between adult ambient temperature and population origin (Adu x Pop on *init* in Table 1, construction of F-tests in Table 4) indicated that females the two sample populations approached the "hot" call stimulus in a nonlinear and inconsistent manner.

Similarly, at both 24° C and 31° C, test subjects from either rearing environment tended to spend more time near the "cold" call stimulus than the alternative stimulus: The majority of *lngr* scores were negative, with such loitering behavior appearing more pronounced for summer-reared females (Figure 3, lower panel). However, the effect of ambient temperature on *lngr* scores was only marginally significant (Adu on *lngr* in Table 1, construction of F-tests in Table 4).

There was no effect of population or family origins on female relative preferences (Table 1, construction of F-tests in Table 4). Quantitative-genetic analyses on either *init* or *lngr* scores indicated no significant variance other than that from maternal

Source of			Type III	F-Test Terms			
Variation	DF	crat	ррс	cdur	dfrq	Numerator	Denominator
Adult environment							
(Adu)	2	93.0417	6.4331	167.1210	83.8956	MS <sub>Adu</sub>	MS <sub>Adu*Pop</sub>
Juvenile environment							
(Juv)	I	0.5843	13.4525	3.7793	9.1469	MS <sub>Juv</sub>	MS <sub>Juv*Pop</sub>
Population origin (Pop)	I	0.1596	22.8136	14.2592	9.6447	MS <sub>Pop</sub>	MS <sub>Fam</sub>
Family origin (Fam)	130	1.2687	2.0136	0.7608	1.1753	MS <sub>Fam</sub>	MS <sub>Ind</sub>
Individual (Ind)	113	0.6570	1.1069	0.5151	0.8486	MS <sub>Ind</sub>	MSAdu*Juv*Ind
Adu x Juv	2	0.6838	0.1589	0.0380	0.0643	MS <sub>Adu*Juv</sub>	MS <sub>Adu*Juv*Pop</sub>
Adu x Pop	2	1.7993	0.1084	0.1783	0.2161	MS <sub>Adu*Pop</sub>	MS <sub>Adu*Fam</sub>
Adu x Fam	260	0.5357	0.5553	0.2672	0.2222	MS <sub>Adu*Fam</sub>	MS <sub>Adu*Ind</sub>
Adu x Ind	226	0.5532	0.5389	0.2808	0.2389	MS <sub>Adu*Ind</sub>	MS <sub>Adu*Juv*Ind</sub>
Juv x Pop	ı	1.6208	0.0118	1.4499	1.0666	MS <sub>Juv*Pop</sub>	MS <sub>Juv*Fam</sub>
Juv x Fam	15	1.2687	0.8173	0.7608	0.9814	MS <sub>Juv*Fam</sub>	MS <sub>Juv*Ind</sub>
Juv x Ind	3	1.0358	1.0990	0.0435	0.7060	MS <sub>Juv*Ind</sub>	MSAdu*Juv*Ind
Adu x Juv x Pop	2	0.2229	0.0476	0.0135	0.0146	MS <sub>Adu*Juv*Pop</sub>	MS <sub>Adu*Juv*Fam</sub>
Adu x Juv x Fam	30	0.3965	0.6417	0.2790	0.3064	MS <sub>Adu*Juv*Fam</sub>	MS <sub>Adu*Juv*Ind</sub>
Adu x Juv x Ind (Error)	10	0.1178	0.4426	0.2027	0.0058	MSAdu*Juv*Ind	

genetic factors (m<sup>2</sup> column for *init* and *lngr* in Table 2) and apart from large residual variation (e<sup>2</sup> column in Table 2). However, these maternal genetic effects were small, accounting for no more than 5% of total phenotypic variance for either female trait, i.e. m<sup>2</sup> < 0.05 for either *init* or *lngr* score in either environment (Table 2).

#### **Discussion**

As the initial step in a complex reproductive repertory, female response towards male sexual signals provides the impetus for assortative mating and, hence, evolution of SMRS (Kirkpatrick 1982). While mate recognition in A. socius and other orthopterans likely involves other sensory modalities (Tregenza and Wedell 1997; Mullen et al. 2007), females recognize species-specific generally acoustic signals (Walker 1957; Olvido and Wagner 2004), though it remains less clear why between-species mating still occurs (Andersson 1994; Marler and Ryan 1997) or how exactly sexual choosiness and premating isolation might evolve within a species (Etges et al. 2007). More troubling, perhaps, is the finding from this study that different methods of quantifying female preference for call stimuli appear to yield diametrically different results (note the

mostly positive *init* scores, suggesting relative preference for "hot" call stimulus across ambient temperatures, and the mostly negative *lngr* scores, suggesting relative preference for "cold" call stimulus across ambient temperatures (Figure 3) further illustrating the complexity of interpreting female responses to male mating call. Nonetheless, it is clear that environmental variation affects pre-mating behaviors in this species, and might explain in part the naturally occurring hybridization between *A. socius* and its more northern congener, *A. fasciatus*.

## **Environmental effects on male calling behavior**

Though causing an apparent shift in mating call reaction norms, rearing environment, when compared with ambient temperature, had a small effect on the *A. socius* mating call. Across different taxa, variation in juvenile characteristics often translates to increased variation in the adult stage (Dingle 1996; Raff 1996; Walker 2000; Hebets 2003). And given the well-established correlation between ambient temperature and chirp rate of cricket mating calls (since Brooks 1882; Dolbear 1897; and until this study) (Figure 1), significant effects of juvenile environment were detected on only one of the four mating call

Source of Variation	Type III MS			F-Test Terms		
Source of Variation	DF	init	Ingr	Numerator	Denominator	
Adult environment (Adu)	I	19.0824	0.1312	MS <sub>Adu</sub>	MS <sub>Adu*Pop</sub>	
Juvenile environment						
(Juv)	I	0.4614	0.0872	MS <sub>Juv</sub>	MS <sub>Juv*Pop</sub>	
Population origin (Pop)	I	1.6348	2.7899	MS <sub>Pop</sub>	MS <sub>Fam</sub>	
Family origin (Fam)	86	0.6149	1.0033	MS <sub>Fam</sub>	MS <sub>Ind</sub>	
Individual (Ind)	58	0.6665	0.9121	MS <sub>Ind</sub>	MS <sub>Adu*Juv*Ind</sub>	
Adu x Juv	I	0.0625	4.1321	MS <sub>Adu*Juv</sub>	MS <sub>Adu*Juv*Pop</sub>	
Adu x Pop	I	6.2636	0.0012	MS <sub>Adu*Pop</sub>	MS <sub>Adu*Fam</sub>	
Adu x Fam	86	0.7465	0.9095	MS <sub>Adu*Fam</sub>	MS <sub>Adu*Ind</sub>	
Adu x Ind	58	0.6197	1.0426	MS <sub>Adu*Ind</sub>	MS <sub>Adu*Juv*Ind</sub>	
Juv x Pop	0			MS <sub>Juv*Pop</sub>	MS <sub>Juv*Fam</sub>	
Juv x Fam	6	1.0814	1.2353	MS <sub>Juv*Fam</sub>	MS <sub>Juv*Ind</sub>	
Juv x Ind	I	0.2774	0.1852	MS <sub>Juv*Ind</sub>	MS <sub>Adu*Juv*Ind</sub>	
Adu x Juv x Pop	0			MS <sub>Adu*Juv*Pop</sub>	MS <sub>Adu*Juv*Fam</sub>	
Adu x Juv x Fam	6	0.7791	1.4050	MS <sub>Adu*Juv*Fam</sub>	MS <sub>Adu*Juv*Ind</sub>	
Adu x Juv x Ind (Error)	I	0.7626	2.2659	MS <sub>Adu*Juv*Ind</sub>		

components: pulses per chirp (ppc in Table 1) of summer-reared males appeared consistently lower than in their springreared full siblings (Figure 2). Failure to detect more widespread and pronounced juvenile environmental effects is most likely due to using a weak statistical test. A single degree of freedom in a repeatedmeasures ANOVA appears insufficient to detect relatively subtle adult phenotypic variation caused by variation in juvenile environment. It is also possible that any effect of juvenile environment may have "decayed" over the adult lifespan of the test subjects (Grace and Shaw 2004). Future studies should consider including more than experimental levels of juvenile environment and a more precise accounting for adult age.

juvenile environment did That fundamentally alter the shape of mating call reaction norms suggests physiological constraints on calling behavior, perhaps a reflection of expressed physiology-related genes that maintain the nature of a population- or even species-specific mating signal. Past studies on this and other gryllids have reported significant heritability of mating call components (Hedrick 1988; Webb and Roff 1992; Mousseau and Howard 1998), indicating a genetic basis for variation in such signaling traits. The current study, however, failed to detect heritable variation in all four mating call components (Table 2). Given the apparent absence of additive genetic variation in mating call components in this study, any attempt at estimating heritability of reaction norms for mating call seemed pointless. Future investigation into heritability of reaction norms will require far larger sample sizes and more extensive pedigrees than those reported here.

Family origin appears to be another major factor in male calling behavior, as it was significant for all four male mating call components (Table 1). The significant interaction of family origin and rearing environment on chirp duration (Juv x Fam on *cdur* in Table 1) suggested that, in terms of variation in chirp duration, males from different maternal lines respond non-uniformly to variation during their juvenile stages.

The quantitative genetic analysis revealed some "maternal genetic effects" (Table 2) on male mating call, i.e. heritable traits of mothers that shape offspring environment, which in turn affects male mating call. Given the absence of direct parental care in A. socius, any maternal genetic effect on mating call must be of a remote nature, e.g. maternal genes that affect the embryonic environment, including maternal gene products transferred into eggs (Yamashita 1996; Saino et al. 2005) and/or maternal choice of oviposition substrate (to the extent that oviposition behaviors determined). genetically At present, however, it is not clear how the embryonic environment of A. socius might normally affect behavior manifested in its other lifecycle stages. Further investigation to quantify A. socius egg "quality" and maternal oviposition behavior certainly seems warranted.

## **Environmentally mediated variation in female phonotaxis**

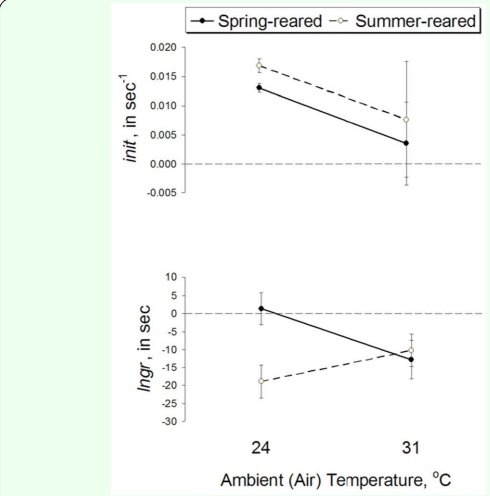
Ambient temperature can affect *A. socius* female response to male mating call. Aside from their increased locomotory activity at higher temperatures, free-walking females tended to move toward the fast-chirp call stimulus much more quickly when observed at 24° C than at 31° C (Figure 3, top panel), though the difference in approach times between these two temperature treatments

was not statistically significant (Table 1). And then, on any given trial, a female may abandon its first-chosen stimulus - a previously undocumented behavior - to spend significantly more time in the vicinity of the other, lower chirp-rate call (Figure 3, bottom panel).

Why *A. socius* females would show such a level of "acoustic promiscuity" is not clear. Searching for prospective mates, while apparently not physiologically costly to *A. socius* females, may prove costly in terms of increased predation risk (Walker and Masaki 1989). For example, most of the field collections prior to sorting and species identification contained various ground-dwelling spiders (presumably, *Lycosa* sp.)

as by-catch. These large and fast-moving spiders seem capable of preying successfully on females that phonotactically locate prospective mates, though such predatory behavior was not observed when spiders were in accidentally prolonged confinement in the field-collected *A. socius* colonies.

Similarly, attraction to both fast-chirp and long-chirp stimuli in *A. socius* females may reflect a series of decisions that females make about a prospective mate. Perhaps females assess mate quality based on chirp rate (slow- versus faster-chirp mating calls) before further assessing mate quality based on chirp duration (short- versus long-chirp mating calls). Females may be attracted to multiple components of male mating call



**Figure 3.** Environmental effects on call preference traits (mean  $\pm$  1 SE) of *Allonemobius socius* females. Relative preference of each female was scored once at each ambient temperature (n = 160 females): a positive score indicates female preference for the "summer-like" (or "hot") male mating call, while a negative score indicates preference for the "spring-like" (or "cold") male mating call. Trait codes are *init*: strength of initial association (i.e. inverse number of seconds spent walking toward a stimulus) made by females for "hot" versus "cold" male mating call and *lngr*: net directional preference in seconds of females for "hot" versus "cold" male mating call. See Methods for full description of trait codes and measurement protocols. High quality figures are available online.

because different male traits provide complementary independent and/or information about fitness benefits (Wagner and Basolo 2007 and references therein). An earlier study of A. socius preference functions (Olvido and Wagner 2004) indicated that females generally associate with longer-chirp stimuli (i.e the "springlike" mating call stimulus in this study) and were less apt to associate with stimuli varying only in chirp rate. To the best of our knowledge, however, no previous study explored stimulus response in terms of the rate at which test subjects approach a given call stimulus, and thus cannot explain the lack of consistency between acoustic preference measured as approach behavior and acoustic preference measured as association behavior. In short, neither ambient temperature nor rearing environment explains why A. socius females approached the faster-chirp call stimulus sooner than the slow-chirp call and later preferred to associate with the longerchirp call stimulus over the short-chirp alternative (Figure 3). Future studies should investigate more closely the relationship between female stimulus approach and association behaviors, as well as identify the relative importance of components, i.e. beyond chirp rate and chirp duration (Olvido and Wagner 2004) of A. socius mating call.

## On hybridization between A. socius and A. fasciatus

The persistence of natural hybrids produced from matings between *A. socius* and its more northern congener, *A. fasciatus*, continues to puzzle biologists. If intraspecific matings result in the highest possible offspring fitness (e.g. Groot et al. 2005), then why do *A. socius* females continue to mate with closely related heterospecifics? Furthermore, conspecific sperm precedence (Howard and Waring

1991) along with high population numbers, abundance of mobile individuals, many capable of long-distance flight in the wild (AO, personal observation), and widespread distribution (Marshall 2004) all indicate potential selection for intraspecific matings and selection against interspecific matings. So, why don't *A. socius-A. fasciatus* hybrids disappear from natural populations (Britch et al. 2001)?

One plausible explanation is that contemporary selection cannot yet suppress behaviors that lead to interspecific matings, at least in A. socius. An earlier study of individual preference functions established unequivocally the importance of the chirp structure of A. socius mating calls. Female A. socius responded positively to variation in A. socius mating calls and did not associate with the mating call typical of a sympatric trilling species, Allonemobius tinnulus (Olvido and Wagner 2004). The current study suggests that A. socius females normally approach conspecific mating calls, but will likely leave an A. socius male for a nearby A. fasciatus male, which produces a longer-chirp mating call (as estimated from Mousseau and Howard 1998). Thus, the current findings suggest that relatively promiscuous or confused A. socius females initiate interspecific matings and subsequently produce most of the naturally occurring hybrids, though similar promiscuity or confusion in A. fasciatus females expanding their range southward into A. socius populations cannot be ruled out. The northward range expansion of A. socius (Britch et al. 2001) seems more consistent with the former idea, however. Future studies should compare mating-call preference functions of A. socius females with those of *A. fasciatus* females.

#### **Acknowledgements**

We thank Sejal Shah, Greg Cauthen, Emmet Maas, and David Perry for helping with cricket husbandry and collecting behavioral data. Dale Van Vleck (USDA-ARS) provided invaluable consultation and technical support for MTDFREML. Ken Fedorka. Eilleen Lawson, LaReesa Wolfenbarger, John McCarty, Fred Nijhout, and several anonymous reviewers greatly improved earlier presentations of this work. This work was supported by National Science Foundation (U.S.A.) research grants to T.A.M., and by an Animal Behavior Society grant, Ford Foundation pre-doctoral fellowship, and National Science Foundation (U.S.A.) post-doctoral fellowship to A.E.O.

#### References

Andersson MB. 1994. *Sexual Selection*. Princeton University Press.

Boldman KG, Kriese LA, Van Vleck LD, Van Tassell CP, Kachman SD. 1995. *A Manual for use of MTDFREML. A Set of Programs to Obtain Estimates of Variances and Covariances* [Draft]. United States Department of Agriculture, Agricultural Research Service.

Bonduriansky R. 2001. The evolution of male mate choice in insects: A synthesis of ideas and evidence. *Biological Reviews* 76: 305-339.

Britch SC, Cain ML, Howard DJ. 2001. Spatio-temporal dynamics of the *Allonemobius fasciatus-A. socius* mosaic hybrid zone: A 14-year perspective. *Molecular Ecology* 10: 627-638.

Brooks MW. 1882. Influence of temperature on the chirp of the cricket. *Popular Science Monthly* 20: 268.

Conover WJ. 1999. *Practical Nonparametric Statistics*, 3rd edition. John Wiley and Sons.

Dingle H. 1996. *Migration: The biology of life on the move*. Oxford University Press.

Dobzhansky Th. 1937. Genetics and the origin of species. Columbia University Press.

Dolbear AE. 1897. The cricket as a thermometer. *The American Naturalist* 31: 970-971.

Etges WJ, de Oliveira CC, Gragg E, Ortíz-Barrientos D, Noor MAF, Ritchie MG. 2007. Genetics of incipient speciation in *Drosophila mojavensis*. I. Male courtship song, mating success, and genotype x environment interactions. *Evolution* 61: 1106-1119.

Ferreira GB, MacNeil MD, Van Vleck LD. 1999. Variance components and breeding values for growth traits from different statistical models. *Journal of Animal Science* 77: 2641-2650.

Grace JL, Shaw KL. 2004. Effects of developmental environment on signal-preference coupling in a Hawaiian cricket. *Evolution* 58: 1627-1633.

Groot AT, Horovitz JL, Hamilton J, Santangelo RG, Schal C, Gould F. 2005. Experimental evidence for interspecific directional selection on moth pheromone communication. *Proceedings of the National Academy of Sciences U.S.A.* 103: 5858-5863.

Hebets EA. 2003. Subadult experience influences adult mate choice in an arthropod: Exposed female wolf spiders prefer males of a familiar phenotype. *Proceedings of the National Academy of Sciences U.S.A.* 100: 13390-13395.

Hedrick AV. 1988. Female choice and the heritability of attractive male traits: An empirical study. *The American Naturalist* 132: 267-276.

Howard DJ, Furth DG. 1986. Review of the *Allonemobius fasciatus* (Orthoptera, Gryllidae) complex with the description of two new species separated by electrophoresis, songs, and morphometrics. *Annals of the Entomological Society of America* 79: 472-481.

Howard DJ, Waring GL. 1991. Topographic diversity, zone width, and the strength of reproductive isolation in a zone of overlap and hybridization. *Evolution* 45: 1120-1135.

Kirkpatrick M. 1982. Sexual selection and the evolution of female choice. *Evolution* 36: 1-12.

Lloyd JE. 1984. On deception, a way of all flesh, and firefly signaling and sytematics. In: Dawkins R, Ridley M, editors. *Oxford Surveys in Evolutionary Biology* Vol. 1. pp. 48-54. Oxford University Press.

Marler CA, Ryan MJ. 1997. Origin and maintenance of a female mating preference. Evolution 51: 1244-1248.

Marshall JL. 2004. The *Allonemobius-Wolbachia* host-endosymbiont system: Evidence for rapid speciation and against reproductive isolation driven by cytoplasmic incompatibility. *Evolution* 58: 2409-2425.

Mousseau TA. 1991. Geographic variation in maternal-age effects on diapause in a cricket. *Evolution* 45: 1053-1059.

Mousseau TA, Howard DL. 1998. Genetic variation in cricket calling song across a hybrid zone between two sibling species. *Evolution* 52: 1104-1110.

Mousseau TA, Roff DA. 1995. Genetic and environmental contributions to geographic variation in the ovipositor length of a cricket. *Ecology* 76: 1473-1482.

Mousseau TA, Roff DA. 1989. Adaptation to seasonality in a cricket: Patterns of phenotypic and genotypic variation in body size and diapause expression along a cline in season length. *Evolution* 43: 1483-1496.

Mullen SP, Mendelson TC, Schal C, Shaw KL. 2007. Rapid evolution of cuticular hydrocarbons in a species radiation of acoustically diverse Hawaiian crickets (Gryllidae: Trigonidiinae: Laupala). *Evolution* 61: 223-231.

Olvido AE, Mousseau TA. 1995. Effect of rearing environment on calling-song plasticity in the striped ground cricket. *Evolution* 49: 1271-1277.

Olvido AE, Wagner Jr WE. 2004. Signal components, acoustic preference functions and sexual selection in a cricket. *Biological Journal of the Linnean Society* 83: 461-472.

Paterson HEH. 1985. The recognition concept of species. In: Vrba ES, editor. *Species and Speciation*. pp. 21-29. Transvaal Museum.

Saino N, Ferrari RP, Romano M, Martinelli R, Lacroix A, Gil D, Møller AP. 2006. Maternal allocation of androgens and antagonistic effects of yolk androgens on sons and daughters. *Behavioral Ecology* 17: 172-181.

Tregenza T, Wedell N. 1997. Definitive evidence for cuticular pheromones in a cricket. *Animal Behaviour* 54: 979-984.

Veech JA, Benedix Jr JH, Howard DJ. 1996. Lack of mating call displacement between two closely related ground crickets. *Evolution* 50: 1982-1989.

Wagner Jr WE, Basolo AL. 2007. The relative importance of different direct benefits in the mate choices of a field cricket. *Evolution* 61: 617-622.

Walker Jr TJ. 2000. Pulse rates in the songs of trilling field crickets (Orthoptera: Gryllidae: Gryllus). *Annals of the Entomological Society of America* 93: 565-572.

Walker Jr TJ. 1957. Specificity in the response of female tree crickets (Orthoptera, Gryllidae, Oecanthinae) to mating calls of the males. *Annals of the Entomological Society of America* 50: 626-636.

Walker TJ, Masaki S. 1989. Life cycle. In: Huber F, Moore TE, Loher W, editors. *Cricket Behavior and Neurobiology*. pp. 1-42. Cornell University Press.

Webb KL, Roff DA. 1992. The quantitative genetics of sound production in *Gryllus firmus*. *Animal Behaviour* 44: 823-832.

Whitesell JJ, Walker Jr TJ. 1978. Photoperiodically determined dimorphic mating calls in a katydid. *Nature* 274: 887-888.

Yamashita O. 1996. Diapause hormone in the silkworm, *Bombyx mori*: Structure, gene expression, and function. *Journal of Insect Physiology* 42: 669-679.