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SEXUAL SELECTION OF MALE SONG IN FREE-LIVING SAGEBRUSH CRICKETS, CYPHODERRIS STREPITANS

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ABSTRACT

Measures of lifetime mating success in the sagebrush cricket, Cyphoderris strepitanus, have revealed that most males succeed in obtaining only 1 mating, while many males fail to attract a female at all and a small minority mate 2 to 4 times. Relative to their abundance in the population, virgin males have a greater likelihood of obtaining a mating than non-virgin males have of securing additional matings, a phenomenon known as the virgin male mating advantage. Previous studies of sagebrush crickets have focused primarily on determining the proximate mechanisms responsible for the virgin male mating advantage, but little work has been done to identify the factors that influence male attractiveness in the first place. Because song plays a central role in mate attraction, it’s likely that variability in song parameters among males could account for the observed differences in mating success. Song is an energetically costly signal to produce and could serve as an honest indicator of male quality. Consequently, males that are able to invest greater amounts of energy into singing should be more attractive to females. In a previous field season, we recorded and analyzed the songs of virgin and non-virgin males and indeed found some evidence that females prefer males which invest greater energy into calling. In the present study, we synthesized artificial sagebrush cricket songs and directly measured female song preference with an arena playback experiment. Females were shown to consistently prefer song characteristics that would require greater energy expenditure by males. Males that sing with long pulse duration, long train duration, and at an intermediate dominant frequency were found to be highly attractive to females.

INTRODUCTION

The sagebrush cricket, Cyphoderris strepitanus, occurs in high-altitude sagebrush meadows and is one of seven known surviving species in the once highly diverse family Haglidae (Morris and Gwynne 1978, Kumala et al. 2005). Males emerge from their burrows at sunset and ascend the sagebrush where they sing late into the night, which functions as their primary means of attracting a female (Snedden & Sakaluk 1992, Snedden & Irazusta 1994). Copulation is initiated when a female climbs onto the dorsum of a male. While the male attempts to transfer a spermatophore, the female feeds on the male’s fleshy hind wings. This mating behavior makes the sagebrush cricket an ideal system for studies of sexual selection because it allows the mating status of males to be readily determined in the field by simply examining their hind wings for damage. Virgin males have pearly-white hind wings, while the hind wings of non-virgins are visibly wounded with melanized scars (Figure 1).

Figure 1. The hindwings of non-virgin males (left) are damaged by females during mating. Virgin males (right) have pearly-white, undamaged hindwings. Note: the forewings were removed.
Relative to their abundance in the population, virgin males have a greater likelihood of obtaining a mating than non-virgin males have at securing additional matings—a phenomenon known as the virgin male mating advantage (Morris et al. 1989, Sakaluk and Ivy 1999). The reduced mating success of non-virgin males is a consequence of the wing wounds inflicted by females, which are accompanied by hemolymph loss (Sakaluk, et al. 2004) and an energetically costly immune response (Leman et al. 2009). Males that have recently mated have less energy available for singing and exhibit at least a short-term reduction in calling (Leman et al. 2009).

While previous studies on sagebrush crickets have focused on determining the proximate mechanism of the virgin male mating advantage, a more basic question has not been addressed: what factors influence male mating success in the first place? Measures of sagebrush cricket lifetime mating success have revealed that the median number of matings is 1, with many males failing to attract a mate at all and a very small minority of males obtaining 2 to 4 matings (Morris et al. 1989). Because song is required for mate attraction (Snedden & Sakaluk 1992), it likely plays a central role in determining mating success. The variable mating success of males could be explained by differences in song characteristics. Previous studies of acoustic Orthoptera have indeed shown that song features can influence mate attractiveness (Bentsen, et al. 2006). What song characteristics, if any, influence attractiveness of sagebrush cricket males?

Acoustic signaling by Orthopterans is thought to impose high energetic costs on males (Prestwich & Walker 1981, Hoback & Wagner 1997, Hack 1998, Bailey et al. 1993). Consequently, song may serve as an honest indicator of male quality (Zahavi 1975, Burk 1988). Males that are able to gain access to limited resources and efficiently convert those resources into attractive sexual signals are likely to be of superior quality and should have greater reproductive success (Holzer et al. 2003). Females may impose directional (linear) sexual selection on males by choosing mates that exhibit the greatest song-producing endurance. Additionally, females may also exert stabilizing (non-linear) selection on males if their auditory system is tuned to only a narrow range of sound frequencies. Males that produce song outside of this frequency range may fail to attract a female.

In a previous field season (Ower et al. 2009), we recorded the songs of virgin and non-virgin males and analyzed sexual selection on five focal song characters: pulse duration, interpulse duration, train duration, intertrain duration, and dominant frequency (Figure 2). Sagebrush crickets sing by stridulating their forewings together. During wing closure, the plectrum (analogous to a guitar pick) strikes the teeth on the file of the opposing wing. Each wing closure results in sound pulse, and multiple wing closures in quick succession produce a train of pulses (Figure 2). Singing with longer pulse durations (and greater energy investment), seemingly were more attractive because they had higher mating success (Ower et al. 2009). Contrary to our predictions, males that engaged in long bouts of energetic singing (long train durations or continual wing movement) apparently were...
unattractive because they tended to be virgins. This
could be a result of the virgin male mating advantage: unmated males, having not incurred the costs of wing
wounding, may have had greater resources to sustain
long trains of song.

To determine which song characters are
relevant to mate attraction it was necessary to
conduct a second field experiment that directly
measured female song preference in arena choice
trials.

**METHODS**

The attractiveness of sagebrush cricket song
was assessed by synthesizing artificial songs and
conducting an arena playback experiment. Four
arenas were setup at a field site neighboring the UW-
NPS research station. The UW-NPS research station
has no nearby populations of sagebrush crickets,
which gave us an acoustically controlled field
environment for conducting audio playback choice
trials. Aluminum flashing was placed around the
arena perimeters (2.0 x 0.3 x 0.1 m) to contain the
females. Speakers were positioned at the terminals of
the arena at a height within the natural perching range
of males singing in sagebrush meadows (Figure 3).

Female sagebrush crickets were captured
from Deadman’s Bar and were transported back to
the UW-NPS research station, where they were
placed into dirt-filled cages and provisioned with a
diet of apple, bee pollen, and cat chow. On
subsequent evenings, phonotaxis trials were
conducted by placing a caged female at the center of
an arena (Figure 3). Control and manipulated signals
were broadcast from the left and right terminals of
the arena. After 5 minutes of acclimation, the cage lid
was very carefully removed, so as not to disturb the
female. If a female approached within 0.3 m of the
manipulated signal it was scored 1 for attractive or
otherwise 0 for unattractive if the female preferred
the control signal.

**Artificial Song Synthesis**

For each trial, the control and manipulated
signal was randomly assigned to either the left or
right speaker. The control was a composite signal,
representing the mean trait values of each of the five
focal song characters (Figure 2). The manipulated
signals were generated by drawing an independent,
random value from the known normalized
distributions of each of the five focal song characters
(Figure 2). This was advantageous, because it
eliminated correlations between song characters that
can make the selection analyses difficult to interpret.
Random selection of each focal song character,
independent of the others, also created songs that
might be biophysically impossible for males to
produce, allowing us to determine if female
preference could drive sexual selection further than
what might be naturally possible. Finally, it provided
manipulative rather than descriptive evidence of
sexual selection. All songs were artificially
synthesized using Adobe Audition 3.

**Statistical Analysis**

The standard multiple regression method of
selection analysis was used to estimate linear and
nonlinear selection gradients for each of the five
focal song characters (Lande & Arnold 1983). Prior

<table>
<thead>
<tr>
<th></th>
<th>( \beta )</th>
<th>PD</th>
<th>IPD</th>
<th>DF</th>
<th>TD</th>
<th>ITD</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pulse Duration (PD)</td>
<td>-0.017</td>
<td>0.233</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>InterPulse Duration (IPD)</td>
<td>0.091</td>
<td>-0.005</td>
<td>-0.136</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Dominant Frequency (DF)</td>
<td>0.084</td>
<td>0.106</td>
<td>-0.182</td>
<td>-0.359*</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Train Duration (TD)</td>
<td>0.159</td>
<td>0.078</td>
<td>0.057</td>
<td>0.257*</td>
<td>0.104</td>
<td></td>
</tr>
<tr>
<td>InterTrain Duration (ITD)</td>
<td>-0.076</td>
<td>-0.153</td>
<td>-0.153</td>
<td>-0.141</td>
<td>-0.100</td>
<td>0.046</td>
</tr>
</tbody>
</table>

Table 1. \( \beta \) (linear) and \( \gamma \) (nonlinear) coefficients from the multiple regression selection analysis. \( p \leq 0.05 = \ast, p \leq 0.01 = \ast\ast, p \leq 0.001. \)
to conducting the analysis, song characters were standardized to have a mean of 0 and standard deviation of 1. Separate regression models were fit to estimate the vector of $\beta$ gradients (which represent linear selection) and matrix of $\gamma$ gradients (which represent both nonlinear and correlational selection). Because nonlinear selection is frequently underestimated due to the effects of correlational selection, canonical analysis was performed (Blows & Brooks 2003). Canonical analysis identifies the major axes of the fitness surface and rotates the data to eliminate correlations (Blows & Brooks 2003). Regression models were fit to the rotated data, in which linear selection is represented by the $\theta$ gradients and nonlinear selection by the $\lambda$ gradients. To visualize the fitness surfaces, thin-plate splines were fitted to the data (Green & Silverman 1994). All statistical analyses were done using the R statistical environment with the fields package (R Development Core Team 2011, Furrer, et al. 2011).

### PRELIMINARY RESULTS

Significant convex selection was found on dominant frequency (Table 1). There was also significant positive correlational selection between dominant frequency and train duration. None of the $\beta$ gradients, representing linear selection, were found to be significantly different from 0 (Table 1).

Canonical analysis rotated the data to eliminate correlations that can result in an underestimation of nonlinear selection. Two major axes (M1 & M5) had significant nonlinear selection (Table 2). The $\lambda$ gradient for the M1 axis was positive, indicating a concave fitness surface. Pulse duration and train duration had the strongest influence along the M1 axis, with more minor contributions from dominant frequency and intertrain duration (Table 2). Along the M5 axis, the $\lambda$ gradient was very significantly negative, indicating a convex fitness surface (Table 2). Dominant frequency contributed very heavily to the M5 axis, with relatively minor contributions from interpulse duration, train duration, and intertrain duration.

The fitness surface that was fitted to the significant M1 and M5 major axes formed a saddle (Figure 4). The most attractive songs—found at the highest peak along the M1 axis—had larger M1 values and tended to have long pulse duration and long train duration (Figure 4). Along the M5 axis, the fitness surface was more attractive, while songs at lower elevations were less attractive.

### Table 2. $\theta$ (linear) and $\Lambda$ (nonlinear) coefficients from canonical analysis. p ≤ 0.05 = *, p ≤ 0.01 = **, p ≤ 0.001.

<table>
<thead>
<tr>
<th>Selection</th>
<th>M</th>
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</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>PD</td>
<td>IPD</td>
<td>DF</td>
<td>TD</td>
<td>ITD</td>
<td>$\theta$</td>
<td>$\lambda$</td>
</tr>
<tr>
<td>M1</td>
<td>0.539</td>
<td>0.038</td>
<td>0.375</td>
<td>0.674</td>
<td>-0.335</td>
<td>0.159</td>
<td>0.181*</td>
</tr>
<tr>
<td>M2</td>
<td>0.693</td>
<td>-0.265</td>
<td>-0.033</td>
<td>-0.205</td>
<td>0.637</td>
<td>-0.120</td>
<td>0.134</td>
</tr>
<tr>
<td>M3</td>
<td>-0.461</td>
<td>-0.566</td>
<td>0.288</td>
<td>0.451</td>
<td>0.426</td>
<td>0.020</td>
<td>0.011</td>
</tr>
<tr>
<td>M4</td>
<td>-0.088</td>
<td>0.649</td>
<td>-0.329</td>
<td>0.464</td>
<td>0.497</td>
<td>0.069</td>
<td>-0.060</td>
</tr>
<tr>
<td>M5</td>
<td>-0.093</td>
<td>0.433</td>
<td>0.817</td>
<td>-0.291</td>
<td>0.230</td>
<td>-0.046</td>
<td>-0.299***</td>
</tr>
</tbody>
</table>
songs with intermediate dominant frequency—or M5 values that were closer to zero—were more attractive, while songs with very low or high dominant frequency were highly unattractive (Figure 4).

Males that sing at a dominant frequency deviating significantly from the population mean of 12.7 kHz have a much lower likelihood of successfully attracting a mate. The female auditory system is perhaps tuned to a narrow frequency range, which would impose stabilizing selection on dominant frequency. Males that sing outside of the hearing range of females presumably would not succeed in attracting a mate and would have a fitness value of 0.

Females preferred songs with long pulse duration and long train duration, and producing these signals would demand greater energy expenditure by males. Non-virgin males are at a competitive disadvantage, due to the damage inflicted by the hindwing feeding behavior of females during copulation. In addition to having lost hemolymph, the wounds on their hindwings may become infected, requiring a costly immune system response that would consume energy which might otherwise have been available for singing to attract another mate (Leman, et al. 2009).

The virgin male mating advantage explains why males that sung in long, energetic bouts of song (long train duration) were counterintuitively found to be unattractive in our previous field experiment (Ower, et al. 2009). Virgin males can afford to invest more energy into calling. The virgin males that sung with long train duration likely would have obtained at least one mating later in the field season.

Non-virgin males that are able to obtain additional matings could be more proficient in converting resources into song through a more efficient metabolic system, superior calling musculature, or a better song producing apparatus. They could, perhaps, also have superior immune systems. More research is needed to determine which factors allow a relatively small number of non-virgin males to obtain multiple matings, when they are at a competitive disadvantage with virgin males.

**ACKNOWLEDGEMENTS**

Due to challenging field conditions from a very late spring followed by heavy, frequent rainstorms, we could not have captured enough female sagebrush crickets without the help of Chuck Collis, Katie Storms and their students Alex Breckenridge, Tanner Fortune, Lily Gage, and Sarah McAfee from Clayton High School in St. Louis, MO. We’d like to thank the University of Wyoming and National Park Service for accommodating us with such a wonderful research station. Funding for this project was provided by the National Science Foundation and the Beta Lambda Chapter of the Phi Sigma Biological Honors Society.

**LITERATURE CITED**


