

Divergent Mechanisms of Acoustic Mate Recognition Between Closely-Related Field Cricket Species (*Teleogryllus spp.*)

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1 ABSTRACT

2 Effective recognition of conspecific mating signals reduces the risk of maladaptive
3 hybridisation. Dissecting the signal recognition algorithms that underlie preferences is a
4 useful approach for testing whether closely related taxa evaluate the same or different signal
5 features to achieve mate recognition. Such data provide information about potential
6 constraints and targets of selection during evolutionary divergence. Using a series of mate
7 choice trials, we tested whether closely-related, but genetically and phenotypically divergent,
8 field cricket species (*Teleogryllus oceanicus* and *Teleogryllus commodus*) use shared or
9 distinct recognition algorithms when evaluating acoustic male calling songs. These species
10 overlap in sympatry, show premating isolation based on female discrimination of male
11 calling songs, yet are capable of producing hybrid offspring. Unexpectedly, female selectivity
12 for features of male song differed between the two species. We found that the two species use
13 a combination of shared and unique signal filtering mechanisms, and we characterised how
14 information about male carrier frequency, pulse rate and temporal patterning is integrated to
15 achieve song recognition in each species. These results illustrate how comparatively few,
16 simple modifications in key components of signal recognition algorithms can lead to striking
17 interspecific discrimination among closely related taxa, despite apparent signal complexity.
18 The finding that some steps during signal recognition and filtering are shared between the
19 species, while others differ, can help to identify behavioural traits targeted by selection
20 during evolutionary divergence.

Keywords: **acoustic communication, divergence, female preference, mate recognition, reproductive isolation, sexual selection, speciation, *Teleogryllus***

22 INTRODUCTION

23
24 The decision-making processes that animals use to evaluate and select among potential mates
25 can have an important influence on the evolutionary outcome of sexual selection (Bateson,
26 1983). For example, mismatches between populations in sexually-selected traits and
27 preferences can generate reproductive isolation and promote speciation (West-Eberhard,
28 1983; Greenfield, 2002; Coyne & Orr, 2004; Mendelson & Shaw, 2005; Safran et al., 2013;
29 Shaw & Mendelson, 2013). Understanding how individuals recognise different male signals
30 is therefore a fundamental goal of sexual selection research (Bateson, 1983; Andersson, 1994;
31 Ritchie, 2007; Chenoweth & McGuigan, 2010), and theoretical models of sexual selection in
32 systems with female choice have predicted a key role for female responsiveness, preference
33 and discrimination of such signals (Lande, 1981; Bateson, 1983; Mead & Arnold, 2004;
34 Andersson & Simmons, 2006). Understanding the mechanistic bases of mating preferences
35 and decision-making behaviours can help to answer questions about their function and
36 evolution. For example, work on the genetic basis of mate choice in drosophilid fruit flies has
37 illustrated an evolutionary link between ecological and mating traits (Chung et al., 2014),
38 studies of the zebra finch *Taenopygia guttata* have clarified neural architecture that might
39 control species difference in song preferences (MacDougall-Shackleton, Hulse, & Ball,
40 1998), and characterising perceptual tuning in the acoustically-signalling anuran
41 *Physalaemus pustulosus* has shown how pre-existing sensory biases can facilitate evolution
42 via sexual selection (Ryan et al., 1990).

43 One way to study the neurophysiological mechanisms underlying mate recognition is
44 to treat the decision-making process as a computational algorithm, or series of operations
45 used to evaluate incoming signals and transform that evaluation into a behavioural action
46 such as a mating response (Ronacher, Hennig, & Clemens, 2015). Filters are integral

47 components of such signal processing algorithms, and in animals, signal filters represent
48 traits of the organism that exclude irrelevant information contained in incoming signals to
49 focus reception upon important signal features. In acoustically-signalling organisms, for
50 example, species can differ in the physical or mechanical properties of structures used to
51 receive sounds, such as tympana, providing peripheral filtering of signals, and the central
52 nervous system can also filter incoming signals depending on the configuration of neural
53 pathways (Greenfield, 2002).

54 By designing tests that manipulate male signal components and assess female
55 responses, it is possible to gain insight into which signal features females attend, which are
56 filtered out, how different signal features might be traded off against one another during
57 assessment, and which ones are possible targets of sexual selection (Kostarakos, Hartbauer,
58 & Römer, 2008; Hedwig, 2006; Hennig, 2009; Henni, Heller, & Clemens, 2014). Much work
59 examining signal recognition algorithms underlying female choice has focused on evaluations
60 that females make among potential mating partners of the same species, and this has taken the
61 form of measuring female preference functions (Wagner, 1998). However, it is less clear
62 whether closely related taxa that risk coming into contact and producing low fitness hybrids
63 use the same, different, or more complex algorithms when faced with the challenges of mate
64 recognition. For instance, closely related species in the treefrog genus *Hyla* have been found
65 to distinguish conspecific from heterospecific calls using different sets of temporal call
66 features, reflecting divergence in signal recognition algorithms (Schul & Bush 2002). In
67 addition to clarifying similarities and differences in the neural mechanisms underlying mate
68 recognition in related species, such data can inform likely targets of sexual selection and
69 constraints during the evolution of reproductive isolation and reinforcement (Coyne & Orr,
70 2004).

71 We tested whether the algorithms and filters underlying mate recognition differ
72 between two closely related field cricket species, *Teleogryllus oceanicus* and *T. commodus*,
73 which are a classic system in the study of acoustic signalling and reproductive isolation (e.g.
74 Hoy & Paul, 1973; Hoy, 1974). These crickets are firmly established as separate species, and
75 both attract mates using long-range acoustic signals that are clearly distinguishable at the
76 phenotypic level (Otte & Alexander, 1983). Both species inhabit coastal regions of Australia,
77 with *T. oceanicus* in the north and *T. commodus* in the south, and their distributions overlap
78 for several hundred kilometres along the central eastern seaboard (Fig. 1a) (Otte &
79 Alexander, 1983). The species readily hybridise in the lab, though hybrid females are almost
80 always infertile, providing an unusual, reciprocal exception to Haldane's Rule (Hogan &
81 Fontana, 1973; Moran, Ritchie, & Bailey, *in press*). Despite their known ability to interbreed,
82 hybridisation is thought to be rare or absent in the wild (Hill, Loftus-Hills, & Gartside, 1972,
83 though see Otte & Alexander 1983).

84 Long-range male advertisement songs of Australian *Teleogryllus* are unusual owing to
85 a patterning complexity not normally observed in grylline crickets: the songs consist of two
86 stereotyped elements, or phonemes: a higher-amplitude pulse train we refer as the “chirp”
87 followed by a series of shorter, lower-amplitude pulses we refer to as “trills” (Figs. 1b, c).
88 Both species also produce a similarly-structured, short-range courtship song which functions
89 to release female mounting behaviour (Balakrishnan & Pollack, 1996), but here we focus on
90 the long-range attraction signal given its known contribution to premating isolation (Hill,
91 Loftus-Hills, & Gartside, 1972, Bailey & MacLeod 2014). Both species exhibit this two-part
92 calling song pattern, although a distinguishing feature between them is that in *T. oceanicus*,
93 the lower-amplitude trills following the initial chirp are comprised of paired pulses (with
94 occasional triplets or, less frequently, higher pulse number trills), whereas the lower
95 amplitude trills of *T. commodus* are comprised of a smaller number of longer-duration trill-

96 like elements composed of a greater number of pulses (Fig. 1b, c). Average carrier
97 frequencies are also higher for *T. oceanicus* (ca. 5 kHz) than for *T. commodus* (ca. 4 kHz)
98 (Bailey & Macleod, 2014). This system therefore provided an opportunity to test whether
99 recognition algorithms underlying female mate choice for conspecific vs. heterospecific
100 songs rely on differential filtering of the same acoustical traits of male calling song, or
101 whether females have diverged in the traits that their signal filters target. Put another way, are
102 females of both species selective for the same or different song features when exerting
103 preference?

104 Previous work illustrated the importance of pattern recognition for conspecific female
105 phonotaxis in *T. commodus*, and suggested that a different balance of peripheral versus
106 central nervous processing contributes to conspecific song recognition in each species
107 (Hennig & Weber, 1997). After validating this result, we developed tests to examine the
108 overall selectivity for con- and heterospecific song patterns and test the contributions of
109 carrier frequency, pulse rate during chirps and trills, and trill pattern composition to song
110 recognition and selectivity. We expected that both species use a combination of spectral and
111 temporal filters (Hennig & Weber, 1997), but given that frequency differences may not
112 definitively identify *T. commodus* and *T. oceanicus*, temporal patterns of song envelopes
113 were expected to play an important role. Two main findings provide insight into divergence
114 of mate recognition algorithms. First, closely related taxa do not necessarily employ the same
115 filter types to differentiate individuals of the other taxon, and second, the strong divergence in
116 mate recognition that this causes can reflect relatively few, minor shifts in the way signals are
117 processed by the nervous system.

118

119 **METHODS**

120

121 *Cricket Rearing*

122

123 We used laboratory-reared adults from two allopatric populations located near
124 Townsville, QLD (*T. oceanicus*) and Moss Vale, NSW (*T. commodus*). Otte & Alexander
125 (1983) reported a single recording of *T. commodus* calling song during a field survey near
126 Townsville. However, that specimen's reported carrier frequency was consistent with *T.*
127 *oceanicus* (4.6 kHz compared to an average of 3.65 kHz for *T. commodus* reported by Otte &
128 Alexander (1983)), and we observed no *T. commodus* in the field or among the laboratory-
129 reared offspring of field-caught individuals from Townsville (Moran & Bailey, 2013). We
130 therefore considered the populations used in this study to be allopatric. Prior to testing, the
131 populations had been reared separately in common-garden conditions in the lab for at least
132 one generation to mitigate maternal effects that could reflect field conditions. Stock crickets
133 were kept in 16L translucent plastic containers at ca. 25 °C on a photo-reversed 12h:12h
134 light:dark cycle. They were fed Supa Rabbit Excel Junior and Dwarf Rabbit nuggets *ad*
135 *libitum*, and provisioned with cardboard egg cartons and moistened cotton pads. Sexually
136 mature adult females (7 days or older) were tested.

137

138 *Female Phonotaxis Tests*

139

140 Female phonotaxis responses were tested using a trackball system and a series of
141 artificially-constructed song playbacks. Setup of the trackball and its operation followed
142 Dahmen (1980) and Hedwig & Poulet (2004). The general protocol we used for phonotaxis
143 assays has been described in detail elsewhere (e.g. Blankers, Hennig & Gray, 2015; Hennig,
144 Blankers & Gray, 2016), so here we summarise the approach and highlight key differences in
145 our experiments.

146 Females were suspended in a walking position over a hollow Styrofoam ball (100 mm
147 diameter, weighing 1.2 to 1.8 g) positioned within a 50 x 50 x 50 cm box lined with acoustic
148 foam. The ball floated on an airstream and its movements were recorded from the bottom by
149 an optical sensor (Agilent ADNS-2051), or by two laterally-focused sensors (ADNS-5050,
150 Avago Technologies) positioned perpendicular to one another. Each channel had a sampling
151 rate of 10 kHz and signal was processed through an A/D-board (PCI-6221, National
152 Instruments, Texas) with Labview v.7.1 or v.9 software. This enabled us to record
153 longitudinal and lateral movements of the trackball when crickets responded during
154 playbacks.

155 Playbacks with the required carrier frequencies and pulse characteristics (1 ms rise
156 and fall) were constructed using LabView 7.0 and transmitted as described in Hennig,
157 Blankers and Gray (2016). Briefly, songs were played back at 80 dB through two Piezo
158 Horntweeter PH8 loudspeakers 25 cm away and 45° to the left and right of the trackball's
159 upper surface. Speakers were calibrated by playing a 1s tone matching the required carrier
160 frequency and assessing with a Brüel and Kjaer sound level meter and a condenser
161 microphone on a fast reading relative to 2×10^{-5} Pa (Brüel and Kjaer 2231 and 4133,
162 respectively). Test sessions were run at 25 ± 1 °C, and for each, we performed one 45 s silent
163 control at the beginning, one 45 s continuous tone control at the end, a positive control at the
164 beginning and a positive control at the end (Fig. 2a, b), plus the 8 focal test signals in
165 randomized order. Parameter values for test signals are provided in the figure captions for
166 each species. Signal presentations were separated by 10 s silent intervals. Silent and tone
167 controls allowed us to monitor and adjust for female motivation and selectivity. Positive
168 controls represented the most attractive combination of song elements for each species (Fig.
169 2d: Positive controls for *T. oceanicus*: 5.0 kHz and TP1: chirp duration: 275 ms, pulse rate
170 during chirp: 16 pulses per second, pps, pulse duty cycle 0.6; trill duration 960 ms composed

171 of double pulses at pulse periods of 40 ms and 80 ms. *T. commodus* 4.0 kHz and TP3: chirp
172 duration: 320 ms, pulse rate during chirp 18 pulses per second, pps, pulse duty cycle 0.65;
173 trill duration 700 ms at a pulse rate of 35 pps followed by a pause of 200 ms). Here we
174 consider female selectivity as the degree to which females discriminate trait values to which
175 they respond most strongly (cf. ‘preference window’ in Butlin (1993), ‘discrimination’ in
176 Bailey (2008), and ‘tolerance’ in Fowler-Finn & Rodríguez (2011)).

177

178 *Phonotaxis Response Scores*

179

180 We calculated phonotaxis scores (*PS*) of 9–32 females for each species, for each 45 second
181 test pattern, using females’ longitudinal forward (*X*) and lateral sideward deviations (*Y*)
182 towards the playback. Both *X* and *Y* were normalised to the attractive controls, and female
183 response relative to the two speakers was averaged to obtain a robust measure of response
184 strength. The *PS* was calculated using the formula:

$$186 \quad PS = \left[\frac{\left(\frac{X_T}{\bar{X}_{CP1,2}} \right) + \left(\frac{|Y_T|}{\bar{Y}_{CP1,2}} \right)}{2} \right] \times [sgn(Y_T)]$$

185

187 where X_T and Y_T represent the forward (*X*) and lateral (*Y*) walking components during the
188 test, and $\bar{X}_{CP1,2}$ and $\bar{Y}_{CP1,2}$ represent forward (*X*) and lateral (*Y*) walking components
189 averaged over positive controls at the beginning (*CP1*) and end (*CP2*) of a test session.
190 Multiplication by the sign of the lateral walking component, $sgn(Y_T)$ (equivalent to turns
191 away from the active speaker), ensured that the overall *PS* could obtain negative values.
192 Negative scores and scores larger than 1 could thus be obtained, although *PS* typically ranged
193 between 0 and 1. For example, $PS < 0$ could result if females turned away from the active

194 speaker, and $PS > 1$ could result if during a test, females exhibited a turning response
195 stronger than that which they exhibited during the control stimulus. In some of the presented
196 data, responses of females were high but did not reach scores of 1.0 (e.g. Figs. 3c, 4a and 5a,
197 b). This reduction was most likely due to suboptimal combinations of the large number of
198 parameters that describe the song patterns of these species. If female PS to the initial positive
199 control of a test session fell below 0.5, the session was aborted. Females were also excluded
200 from further analysis if their final positive control PS was less than 50% of their initial
201 positive control PS , or if they were highly responsive during silent and tone controls,
202 although the latter occurred infrequently (Fig. 2a, b).

203 Statistical comparisons of the turning responses to test patterns were performed using
204 paired t-tests. Statistical significance was assessed at $\alpha = 0.05$. Unless otherwise specified,
205 means and standard errors of the data are presented, and sample sizes (n) for each test series
206 are given in the figure captions. Degrees of freedom (df) were calculated as $df = 2(n - 2)$.
207 R v. 2.15.2 was used in construction of the map in Fig. 1 (R Core Team 2012; Becker &
208 Wilks 2013a,b).

209

210 **RESULTS**

211

212 *Interspecific Variation in Female Selectivity*

213

214 Females of both species were tested for their ability to discriminate con- and
215 heterospecific song types. As illustrated in Fig. 1, song structure is distinct between these
216 species (see also: Otte & Alexander, 1983; Hennig & Weber, 1997; and Table S1 in Bailey &
217 Macleod, 2014). For this test, song patterns were constructed that exhibited an initial chirp

218 section typical for the respective species, plus a trill part that mimicked the song pattern with
219 respect to pulse rate (TP1 for *T. oceanicus*, TP3 for *T. commodus* in Fig. 2d). Additionally,
220 females were tested with patterns representing a fusion of otherwise separated trill pulses to
221 longer blocks of sound (TP2, TP4 in Fig. 2d). The latter two test patterns were expected to be
222 indicative of potential differences in selectivity for the trill part between both species. Each
223 test pattern was presented at the con- and heterospecific carrier frequency (4.0 and 5.0 kHz in
224 Fig. 2c). *T. commodus* females were highly selective for carrier frequency and temporal
225 patterning elements, whereas *T. oceanicus* females were less selective for temporal pattern
226 features (Fig. 2c). For instance, *T. oceanicus* accepted all test patterns, provided they were
227 broadcast at 5.0 kHz. *T. oceanicus* responses were attenuated at 4.0 kHz. In contrast, females
228 of *T. commodus* only responded if both the carrier frequency and the temporal pattern
229 corresponded to the conspecific song. This distinction illustrates that *T. commodus* females
230 only showed strong responses to song models with the lower species-specific 4 kHz carrier
231 frequency when they were presented with an appropriate species-specific pulse pattern,
232 whereas *T. oceanicus* females responded strongly to species-specific 5 kHz frequency
233 playbacks regardless of the pulse pattern presented. Females of *T. commodus* were therefore
234 more selective for the temporal pattern than females of *T. oceanicus* (Fig. 2).

235

236 *Components of female selectivity*

237

238 In a further series of tests, females of both species were exposed to test patterns
239 designed to dissect the contribution to the selectivity observed before of carrier frequency,
240 pulse rates in chirp and trill, and trill composition (Fig. 2c). As predicted, responses to carrier
241 frequency were differently tuned in the two species. *T. commodus* females showed a peak
242 response to calling songs at 4.0 kHz, whereas *T. oceanicus* females preferred songs 4.5 kHz

243 or higher in frequency (Fig. 3a). Female responses for pulse rate during chirps were broadly
244 similar, with only a small difference in the most preferred pulse rate (*T. oceanicus*: 12 pps, *T.*
245 *commodus* 16 -18 pps, Fig. 3b). However, *T. commodus* showed selectivity for a specific
246 pulse rate of 32 pulses per second during the trill portion of the calling song, whereas *T.*
247 *oceanicus* females only responded if the pulse rate during the trill part was the same as during
248 the chirp part, that is at 12 pps (Fig. 3c, c.f. *T. oceanicus* in Fig. 3b). *T. commodus* thus
249 exhibits different pulse rate selectivity for the two song phonemes, requiring two pulse rate
250 filters, whereas the most preferred pulse rate (12 pps) is the same for each phoneme in *T.*
251 *oceanicus*, for which a single pulse rate filter suffices. The addition of a separate filter for
252 pulse rate selectivity suggests higher sensitivity to temporal pattern properties of calling song
253 for *T. commodus* females than for *T. oceanicus*. Indeed, the preferred pulse rate of 12 pps by
254 *T. oceanicus* in Fig. 3c indicated that females did not require a trill part for recognition and
255 that the pulse rate of the chirp part alone sufficed.

256 The contribution of the trill composition in terms of pulses per trill and trill duration
257 indicated broadly similar responses in both species (Fig. 4). *T. oceanicus* females accepted
258 trills built from two or more pulses, whereas *T. commodus* accepted trills built from three
259 pulses or more (Fig. 4). Longer trills were accepted by both species equally readily, although
260 only females of *T. commodus* appeared to be selective for a particular pulse rate during this
261 part (Fig. 3c).

262 To examine whether *T. oceanicus* simply ignored features of the trill part or whether
263 they exhibited selectivity to other temporal cues, females were tested with patterns that varied
264 the pulse duty cycle. Such patterns exhibit different amounts of sound energy independent of
265 a particular pulse rate as illustrated in Fig. 5c, as the duty cycle is calculated from the pulse
266 duration divided by the pulse period. *T. oceanicus* females exhibited a strong selectivity for

267 all patterns with a pulse duty cycle higher than 0.5, which corresponded to patterns with high
268 sound energy as they contained pulses longer than the pauses in between (Fig. 5).

269

270 DISCUSSION

271

272 The origin and maintenance of mating barriers is a fundamental requirement for
273 speciation to occur in situations where diverging populations could hybridise, or when
274 secondary contact occurs between closely related taxa (Coyne & Orr, 2004). Divergence in
275 signalling and mate recognition traits facilitates the establishment of such barriers. While
276 changes in signalling traits and mate recognition at the phenotypic level have been well-
277 characterised in a number of systems, less is known about whether the underlying
278 physiological mechanisms that control such mate recognition are shared or not in such taxa.
279 Because signals are typically multi-component and complex, divergence could occur as a
280 result of changes in the same filtering mechanism in different species, such that different
281 values of the same signal trait are preferred, or by establishment of new filters such that
282 divergent taxa are tuned to different signal traits. We found a mixture of both scenarios in *T.*
283 *oceanicus* and *T. commodus*, which we can illustrate by separating the different filter
284 components of the processing algorithm much like a flow diagram (Fig. 6).

285 Our dissection of mate recognition algorithms in *Teleogryllus* showed that females of
286 both species attended to frequency differences and showed sharply tuned filters that almost
287 perfectly match the documented differences in carrier frequency of conspecific male calling
288 songs, consistent with prior reports (Hennig & Weber, 1997; Bailey & Macleod, 2014). The
289 majority of known examples of acoustic species recognition in insects, and particularly
290 crickets and other ensiferan insects, focus on temporal patterning of male advertisement
291 songs (e.g. Ritchie, 1991; Mendelson & Shaw, 2005; Meckenhäuser, Hennig, & Nawrot

292 2013; Kostarakos & Hedwig, 2015), and a longstanding assumption about the evolution of
293 cricket calling songs is that there is unlikely to be significant variation in carrier frequency
294 among closely related taxa, due to the mechanical constraints imposed by physical features of
295 male forewings used in song production (Alexander, 1962). For example, neural recordings
296 of responses to courtship song in a gryllid from the western hemisphere, *Gryllus assimilis*,
297 indicate the importance of temporal song patterning compared to carrier frequency, with
298 female auditory neurons exhibiting a broad frequency response spectrum ranging from 3.5
299 kHz to 14.5 kHz (Vedenina & Pollack, 2012), and early perceptual models for discrimination
300 of acoustic signals in *T. oceanicus* suggested that the main frequency-based distinction this
301 species makes is of a categorical nature, between low frequency and ultrasound (Wytenbach,
302 May & Hoy, 1998).

303 Nevertheless, our results confirm that both *T. oceanicus* and *T. commodus* share
304 frequency filters, with the result that females of both species filter incoming male signals as a
305 function of those signals' dominant carrier frequency. Selectivity for frequency indicated that
306 peak female responses were only approximately 1kHz apart. However, this selectivity
307 matches observed differences in frequency differences of males, both from these populations
308 (Moran & Bailey, 2013) and from other populations of the same species (Bailey & Macleod,
309 2014). Such a shift in the frequency filter does not necessarily require evolutionary change in
310 complex neural architecture or physiological processes, and could be underpinned by simple
311 size scaling differences that have arisen during the evolutionary history of these two species.
312 For example, a meta-analysis of 58 species of tettigoniids, an ensiferan group in which males
313 sing using a forewing file and scraper mechanism, uncovered significant overall covariance
314 between body size and carrier frequency (Montealegre-Z, 2009). *Teleogryllus commodus* are
315 larger than *T. oceanicus* on average, and if male forewing structures and tympanal hearing
316 organs scaled with body size in a correlated manner during divergence, corresponding

317 frequency filters in females of each species could be selectively tuned to the dominant carrier
318 frequency produced by conspecific males.

319 Both cricket species appear to share another filter, by which the pulse rate of the chirp
320 portion of the song is evaluated. Pulse rate selectivity can arise from only a small network of
321 neurons, in which the property of a rebound oscillation plays a crucial role (Weber &
322 Thorson, 1989; Pollack, 2000; Clemens & Hennig, 2013; Schöneich, Kostarakos, & Hedwig,
323 2015). Notably, the preference functions for this song component were very similar in the
324 two species (Fig. 3b, 6). This similarity is consistent with previous reports suggesting that
325 pulse rate during the chirp is under stabilising selection in both species (Hennig & Weber,
326 1997). In contrast with the chirp filter, the species differ in selectivity of the trill portion of
327 the song. *T. oceanicus* females appear to be unselective towards the trill pattern (Figs. 2, 3
328 and 4), but they preferred trill patterns with longer pulses and shorter pauses (Fig. 5). Taken
329 together, this is indicative of duty cycle selectivity favouring patterns with higher energy. The
330 particular timing of pulses as given by a pulse rate did not appear relevant, which contrasted
331 distinctly with *T. commodus* females (Fig. 3c). Thus, female selectivity for pulse rate within
332 the trill portion of calling song highlights a key difference between the species: *T. commodus*
333 females are more highly selective of trill patterning, focusing on temporal aspects of trill
334 pulses such as pulse rate, whereas *T. oceanicus* females attend to the pulse duty cycle of the
335 trill irrespective of the patterning (Fig. 2, Fig. 6). *T. commodus* appears to be the rarer species
336 in sympatry (Moran & Bailey, 2013), and it enters a diapause in more southern populations
337 (Otte & Alexander, 1983). Both scenarios might favour enhanced female selectivity in *T.*
338 *commodus* females: rarity would increase the chances of maladaptive hybridization, and
339 introgression of genes that reduce or eliminate the tendency to enter diapause would be
340 detrimental to *T. commodus* females.

341 The integration of similar signal recognition algorithms based on frequency filters

342 with a different mechanism based on discrimination of pulse rate during the trill portion of
343 the song contrasts with recent findings in several gryllids producing either short, chirp-like
344 phonemes (Hennig, Blankers, & Gray, 2016) or long, trill-like phonemes (Blankers, Hennig,
345 & Gray, 2015). The latter species show identical computational algorithms for evaluating
346 acoustic signals based on pulse pattern and chirp/trill features. (Blankers, Hennig, & Gray,
347 2015; Hennig, Blankers, & Gray, 2016). Nevertheless, these species differ in their preference
348 for a particular pulse rate or chirp/trill duty cycle. Some gryllid species show a transition
349 from a pulse rate filter to a pulse duty cycle filter, consistent with what we have observed in
350 *Teleogryllus* (Fig. 5) (Hennig, Blankers, & Gray, 2016). Our behavioural experiments cannot
351 resolve how the algorithmic flow of information during phonotaxis or particular filter
352 component is implemented in terms of physiological or neural activity. Nevertheless,
353 physiological recordings from sensory cells in the tympanic ear (Imaizumi & Pollack, 1999)
354 and brain neurons sensitive to pulse rate (Schöneich, Kostarakos & Hedwig, 2015) support
355 the proposed sequential processing steps and filter properties illustrated in Fig. 6.

356 There are several illustrative differences between song pattern recognition in the
357 gryllids mentioned above versus *Teleogryllus*, which suggest a more general, taxonomically-
358 widespread pattern underlying the evolution of signals and signal recognition during
359 diversification. For example, most gryllids produce a series of pulses grouped into chirps or
360 trills, which are separated by variable durations of silence (Blankers, Hennig, & Gray, 2015;
361 Hennig, Blankers, & Gray, 2016). In contrast, the *Teleogryllus* species we studied produce
362 calling songs with a greater number of phonemes, as in the chirp and trill part (Fig. 1),
363 although *Teleogryllus* species with simpler song patterns are known (Rothbart & Hennig,
364 2012). The tendency toward additional song pattern elements, or phonemes, can be even
365 greater in other ensiferan taxa; certain species of the Tettigoniid genus *Amblycorypha*
366 produce some of the most complex acoustic signals of any insect, with varied arrangements

367 of up to four phonemes (Walker & Dew, 1972). A tempting prediction is that the signal
368 recognition filters required to process complex incoming signals will be correspondingly
369 complex, and may therefore provide a larger target for selection or drift to modify (Fig. 6)
370 (Hebets & Papaj, 2005).

371 Despite the phenotypic differences in song recognition and apparently larger number
372 of filters required for mate recognition in *Teleogryllus* (Fig. 6), the filters themselves are in
373 principle similar or even identical to those described in other crickets. This observation
374 suggests that the apparently derived situation in *Teleogryllus* builds on existing schemes of
375 pattern recognition. Two important transitions are worth highlighting: first the duplication of
376 a pulse rate filter, and second, the transformation of a pulse rate filter to a duty cycle filter
377 (Fig. 6), the latter of which appears complicated at first but can be simply achieved by small
378 changes of the filter template used for song recognition (Hennig, Heller, & Clemens, 2014).
379 These observations also suggest that recognition of a complex song pattern such as the trill
380 portion of *T. oceanicus* calling song (Fig. 1) does not necessarily evolve because of a more
381 complex filter, but may arise in response to a relatively simple duty cycle filter (Figs. 5, 6).
382 The combined effects of multiple, simple filters thus provide a parsimonious explanation for
383 the multitude of different ways in which species-identifying signals can diverge alongside
384 recognition mechanisms for those signals. In *T. oceanicus* and *T. commodus*, divergence in
385 signal recognition appears to have arisen from a combination of different filters applied to the
386 same signal features, plus the modification of filters to target distinct signal features. Changes
387 in decision algorithms must ultimately reflect measurable physical changes in the structure or
388 neural connections within the organism, and our results are consistent with the idea that such
389 divergence will follow an evolutionary “path of least resistance”: apparent signal recognition
390 complexity can arise from few, basic decision algorithms.

391

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393

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404

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541 **Figure captions:**

542

543 **Figure 1.** Cricket ranges and signals. (a) Approximate Australian distribution of *T. oceanicus*
544 (light grey), *T. commodus* (dark grey), and region of sympatry (stripes). , based on Otte &
545 Alexander (1983) and Moran & Bailey (2013). Locations of populations used in this study
546 are indicated with arrows. Our field and laboratory observations are consistent with these
547 being pure-species, allopatric populations (see main text for details). Male calling song
548 diagrams are based on Bailey & Macleod (2014) and illustrate song features of interest for (b)
549 *T. oceanicus* and (c) *T. commodus*. Different authors have historically used different
550 terminology to describe elements of *T. oceanicus* calling song. Those employed in the present
551 study are indicated with larger font, while alternative terms for the same song features are
552 indicated with smaller font in parentheses to ease comparison with prior work.

553

554 **Figure 2.** Female selectivity for male calling song models that varied in carrier frequency and
555 temporal patterning. Phonotaxis scores are shown for *T. oceanicus* females (black bars, n =
556 15) and *T. commodus* females (open bars, n = 13). (a,b) Female response to positive
557 (attractive stimuli) and negative controls (unattractive stimuli) during a test session (CP1, 2
558 positive controls at the beginning and end of a test session, CS: silent control, CT tone
559 control). (c,d) Females were presented with test patterns (shown in (d)) similar to a *T.*
560 *oceanicus* (TP1,2) or *T. commodus* (TP3,4) calling song. Each test pattern was presented at
561 4.0 and 5.0 kHz, corresponding to the carrier frequency of the song of both species.
562 Responses in (c) marked with ‘#’ were not significantly different from the positive controls in
563 (a) and (b), and the response marked with ‘*’ was significantly ($p < 0.05$, t-test) different from
564 the negative controls in (a) and (b). Means and standard errors are presented in (a)-(c).

565

566 **Figure 3.** Preferences (means and standard errors) for calling song features exerted by
567 females of each species: (a) carrier frequency (*T. oceanicus* n = 15; *T. commodus* n = 13),
568 (b) pulse rate in the chirp portion of the song, holding trill pulse rate constant (*T. oceanicus* n
569 = 9; *T. commodus* n = 10), and (c) pulse rate in the trill portion of the song, holding chirp
570 pulse rate constant (*T. oceanicus* n = 12; *T. commodus* n = 41). Pulse rates are given in pulses
571 per second. Response levels higher than 0.7 were not significantly different from the positive
572 controls, response levels below 0.3 were not significantly different from the negative controls
573 (c.f. Fig. 2A, B). Test patterns in (a) corresponded to conspecific songs as in Fig. 2D (TP1 for
574 *T. oceanicus* and TP3 for *T. commodus*). Test patterns in (b) corresponded to continuous
575 pulse trains with variable pulse rate for *T. oceanicus* and variable pulse rate during the part
576 with a continuous pulse train during the trill for *T. commodus* as for TP3 in Fig. 2D. Test
577 patterns in (c) had a constant chirp part as TP1 and TP3 in Fig. 2D and a continuous trill part
578 with variable pulse rate as TP3 in Fig. 2D. Typical trait values for the calling song signal of
579 both species are available from Bentley & Hoy (1972), Hill, Loftus-Hills & Gartside (1972),
580 and Hennig & Weber (1997). (For *T. commodus*/*T. oceanicus*, respectively: carrier
581 frequency: 3.5-3.8//4.5-4.9, pulse rate within chirp: 19-20//15-16, pulse rate within trill: 31.5-
582 31.6//24-26).

583

584 **Figure 4.** Preferences for overall trill composition. (a) Phonotaxis scores (means and standard
585 errors) for *T. oceanicus* (n = 11) and *T. commodus* females (n = 23). (b) Diagram of test
586 patterns in which the number of pulses was varied during the trill portion, thereby changing
587 the trill duration. Pulse periods were set to 40 ms, and pulse periods between groups of pulses
588 were set to 80 ms. Phonotaxis scores higher than 0.3 were significantly different from the
589 negative controls (p < 0.05, c.f. Fig. 2a,b).

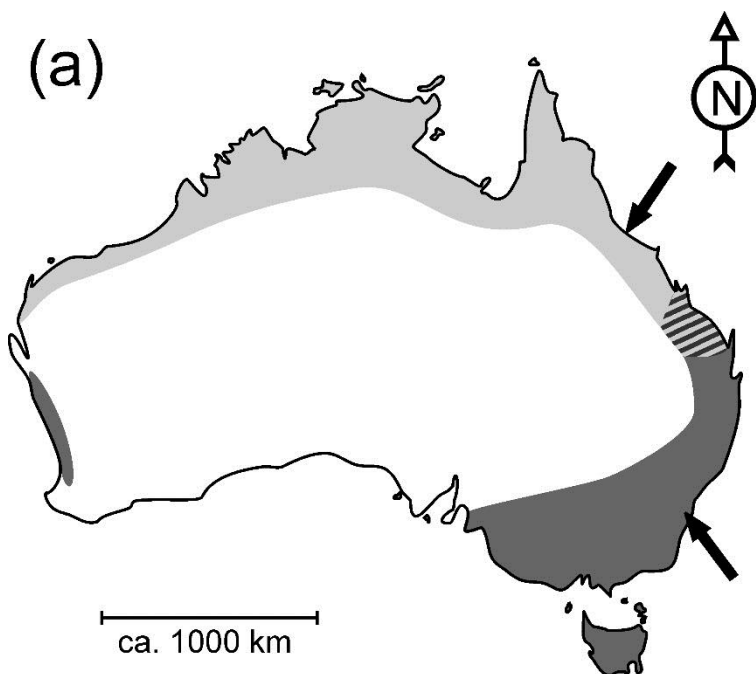
590

591 **Figure 5.** (a) Selectivity for temporal cues during the trill part containing more sound energy
592 by females of *T. oceanicus*. Numbers refer to test patterns in (c). The open circle at the center
593 refers to phonotaxis score to the positive control pattern. Diameter of circles indicates
594 strength of phonotaxis score which was 1.0 for the positive control. (b) Females exhibit
595 selectivity for the pulse duty cycle in the trill portion of the calling song (means and standard
596 errors are presented). The curves correspond to transects through (a) from upper left to lower
597 right at different pulse periods, as indicated. (c) Selected test patterns, as indicated in (a), with
598 a constant chirp part (TP1 in Fig. 2D) and a varied trill section. Numbers to the right refer to
599 the pulse duty cycle (pdc) of each pattern. Response levels higher than 0.3 were significantly
600 different from the negative controls ($p < 0.05$, c.f. Fig. 2a,b).

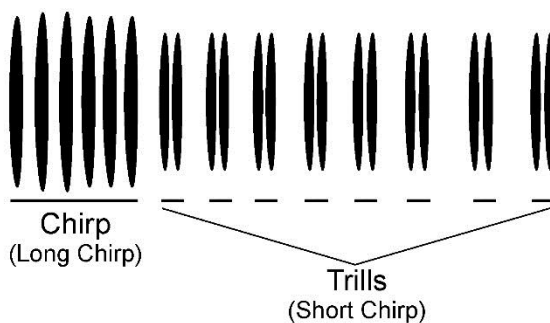
601

602 **Figure 6.** Flow diagram describing differential processing for processing for signal
603 recognition in *Teleogryllus* species. (Top): representative song signals for each species. (First
604 recognition level): sensitivity to carrier frequency given by the frequency response of the
605 tympanic ear and sensory cells depicted as tuning curves. (Middle level): processing of the
606 pulse pattern within the phonemes of chirp and trill depicted by sensory templates for pulse
607 rate and integration of sound energy for duty cycle evaluation. (Bottom level): integration of
608 processing across time scales of both phonemes of chirp and trill. Common filters for carrier
609 frequency of the song are differently tuned in the two species, leading to quantitative
610 differences in female responses (grey boxes: brown lines indicate preferences for lower
611 carrier frequencies by *T. commodus*, blue lines for higher carrier frequencies by *T.*
612 *oceanicus*). Both species also share similar filters for the pulse rate during the chirp portion
613 (grey boxes: black rectangles indicate sound pulses, brown lines (*T. commodus*) and blue lines
614 (*T. oceanicus*) indicate sensory templates with rebound properties that will respond best to
615 the given pulse rate in the chirp pattern (Schöneich, Kostarakos, & Hedwig, 2015)). A

616 qualitative difference is a more selective pulse rate filter in *T. commodus* for pulse rates
617 during the trill part of a song, while *T. oceanicus* remain largely unselective for the trill
618 pattern provided sound energy remains high (i.e. sensitivity for high duty cycle, yellow
619 boxes: filters for trill pulse rate are symbolised by a rebound oscillation of the sensory
620 template, filters for pulse duty cycle by an integration). Separate streams of information about
621 chirp and trill features are finally integrated similarly for song recognition and discrimination
622 in both species. In aggregate, while females of both species might employ similar algorithms
623 to process incoming signals on the basis of carrier frequency and chirp pulse rate (grey
624 boxes), they show divergent filter properties for the trill part (yellow boxes), for which *T.*
625 *commodus* females are more selective.

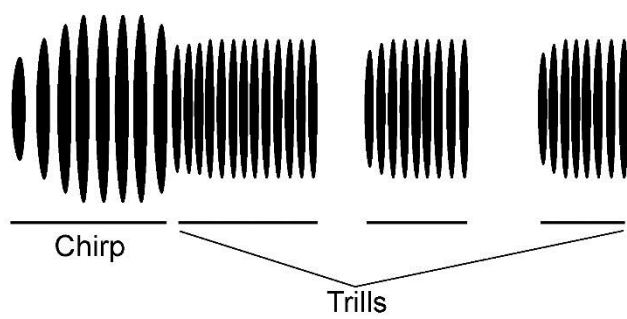


(b) *Teleogryllus oceanicus*



ca. 100 ms

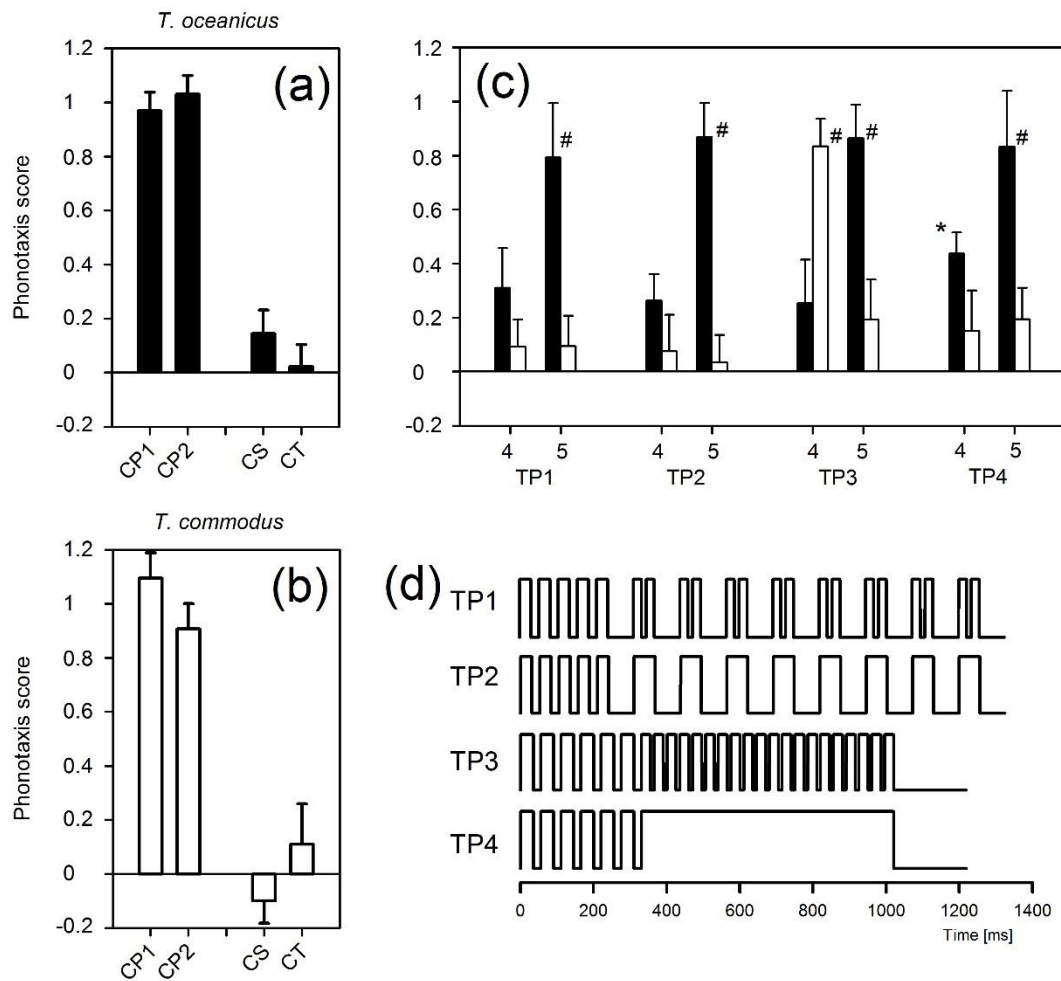
(c) *Teleogryllus commodus*



626

627 **Figure 1**

Fig. 2



628

629 **Figure 2**

Fig. 3

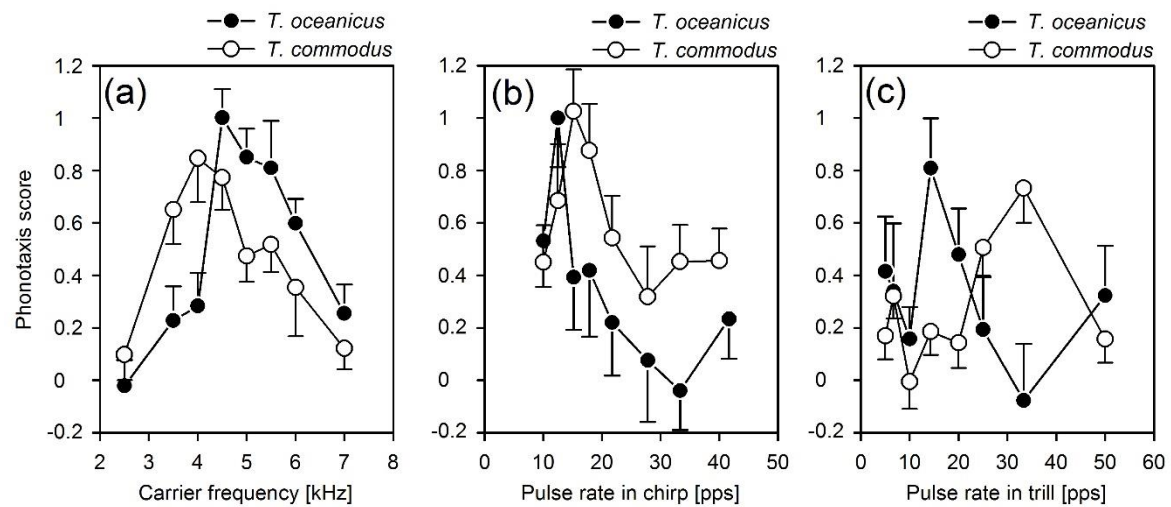
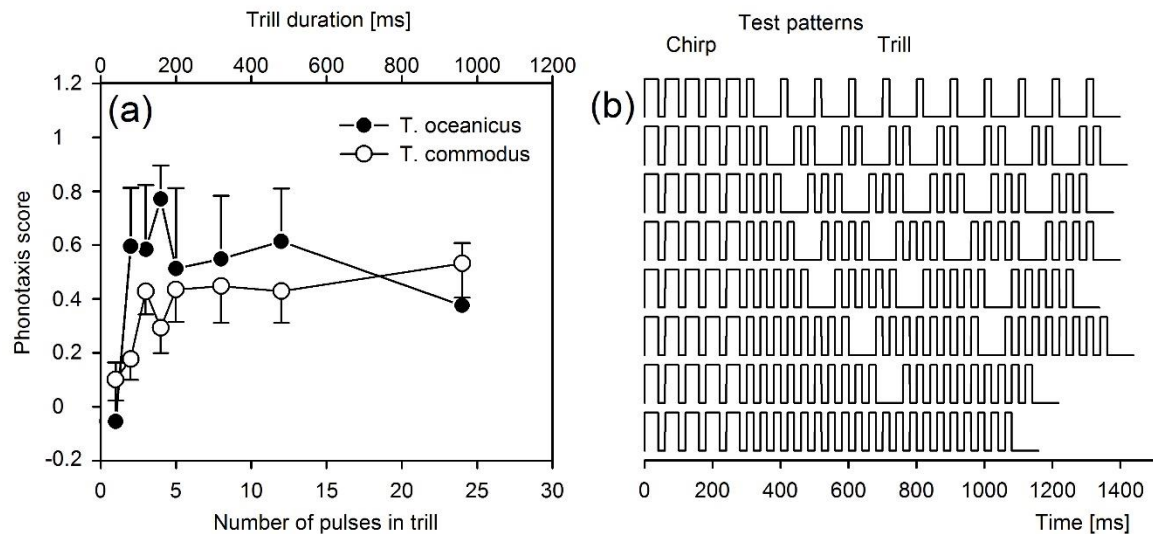
632 **Figure 3**

Fig. 4



633

634 **Figure 4**

Fig. 5

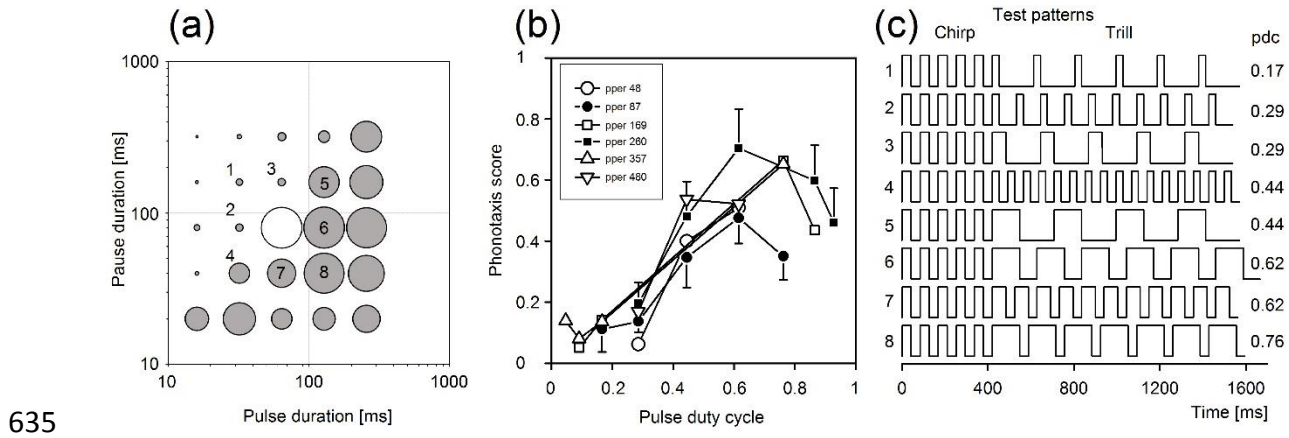
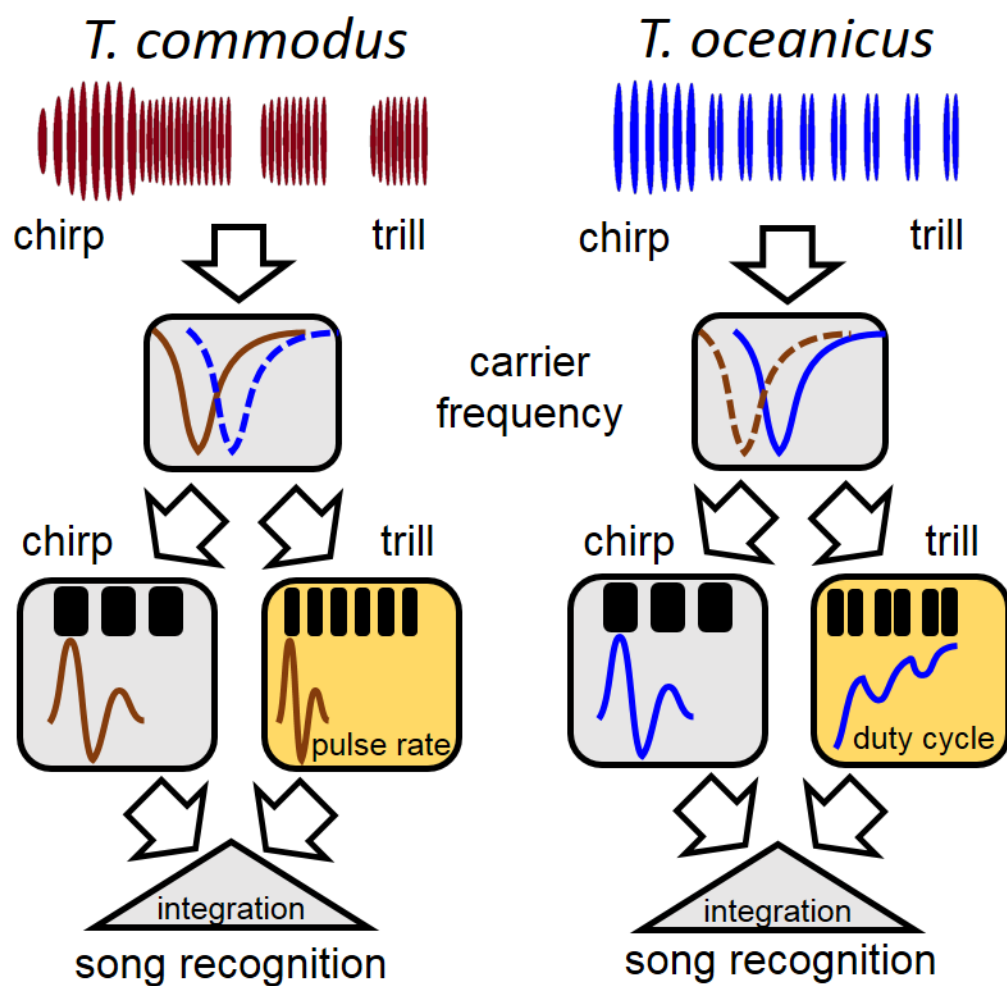


Fig. 6



637

638 **Figure 6**