Response to Kroodsma's critique of banded wren song performance research

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1 A critical review of a popular scientific theory, large or small, is something we applaud because, if well executed, it stimulates discussion and progresses science. However, such a review needs 2 3 to be balanced, objective, informed and logical, especially if it concludes that a well-supported 4 theory is flawed. Unfortunately, Kroodsma's current criticism (Kroodsma, 2016) of the birdsong 5 performance literature suffers from the same weaknesses as his earlier criticism of song 6 repertoire use in sexual selection (Byers & Kroodsma, 2009), despite the fact that he has been 7 alerted to those mistakes (Collins, de Kort, Perez-Tris, & Telleria, 2011). Those weaknesses 8 include outright errors and misrepresentations, highly selective citation of the literature, and 9 convoluted logic (sensu Podos, 2016). Here we would like to take this opportunity to redress the 10 specific issues he raised with respect to our work on the banded wren (*Thryophilus pleurostictus*) 11 and by doing so, illustrate how his criticism is flawed as a result of the above weaknesses, his 12 restricted definition of "song performance", and a misunderstanding of the song system of the banded wren. 13

14 Banded wren males possess song repertoires of approximately 25 distinct song types, 15 which are largely shared with other males in their neighborhood. The terminal trills of these song 16 types vary in their trill note rate, frequency bandwidth, and vocal deviation. (Maximal observed 17 trill rates and maximal observed frequency bandwidths are inversely related in many songbirds. 18 defining a negatively-sloped upper limit line on a trill-rate versus frequency bandwidth plot; the 19 perpendicular distance of a given trill from this line is its vocal deviation.) Moreover, the trill 20 notes themselves vary greatly in shape and complexity. Most of our research on this species has 21 focused on the use of these song types in male-male territorial interactions (Burt & Vehrencamp, 22 2005; Hall, Illes, & Vehrencamp, 2006; Molles, 2006; Molles & Vehrencamp, 1997, 2001; Trillo 23 & Vehrencamp, 2005; Vehrencamp, 2001; Vehrencamp, Ellis, Cropp, & Koltz, 2014;

24 Vehrencamp, Hall, Bohman, Depeine, & Dalziell, 2007). We have shown that males negotiate their territorial boundaries primarily by varying short-term song-type diversity and switching rate 25 26 to indicate their propensity to approach, stand their ground, or retreat from a territorial rival. 27 Males also frequently song type-match each other during aggressive encounters. This primary 28 role of song type choice does not rule out the possibility that subtle details of song structure also 29 play a role and provide additional types of information about the sender, for both male and 30 female receivers. The type-matching behaviour of countersinging males provides ample 31 opportunities for receivers to compare their performances on the same song type, as proposed by 32 Logue & Forstmeier (2008) for repertoire species.

33 Trade-off coding (where two components of a signal are negatively correlated such that 34 extreme values of one tends to inhibit extreme values of the other) sets up the potential for 35 receivers to exert selective pressure on combinations that reveal useful information about the 36 sender. This idea has been around for over two decades (Bradbury & Vehrencamp, 2011; Hebets 37 & Papaj, 2005; Podos, 1997, 2016; Wells & Taigen, 1986). Whenever one observes a negative 38 correlation between two signal components, it is worth testing this trade-off hypothesis (Podos' 39 hypothesis #1) by looking to see whether receivers pay attention to alternative combinations of 40 those components, and if so, whether individual variation in these combinations is associated 41 with sender condition, context, or reproductive success (Podos' hypothesis #2). This is how 42 science proceeds: repeated testing of an hypothesis' predictions in a variety of species using a 43 variety of experimental approaches. The trills of many bird songs are an obvious candidate for 44 testing this hypothesis, because they are highly precise vocal utterances that have evolved under 45 selective pressure from receiver responses. We examined several aspects of performance in the 46 banded wren, not limited to vocal deviation as Kroodsma has restricted himself to here, but also

the individual components of vocal deviation, trill rate and frequency bandwidth, along with trill
note consistency, in multiple experimental and correlational studies. We have obtained consistent
evidence that performance does matter, even in a repertoire species.

50 Our first indication that performance components affected male responses was obtained 51 by Illes, Hall & Vehrencamp (2006), building on the Ballentine, Hyman, & Nowicki (2004) 52 study, where songs modified to have faster or slower trill rates were simultaneously presented to 53 territorial males. We found that most subjects initially approached the fast stimulus, but if they 54 were exposed to a broader bandwidth (lower vocal deviation) trill they subsequently spent less 55 time close to the fast speaker. Kroodsma disparages the design, execution, analysis, results, and 56 conclusions of this study, and chides the many researchers who have cited the paper. We show 57 below that each of his criticisms is incorrect or misinformed.

58 First, Kroodsma states that the playback should have been conducted with blind 59 observers. The experimental design consisted of a two speaker set-up each broadcasting a 60 separate stimulus. The observers were not informed about which speaker broadcasted which 61 stimulus. Nevertheless, as acknowledged for the Cramer & Price (2007) study, an acute observer 62 might discern which was which by listening and we would have had to deafen the observers to 63 exclude this possibility. Second, we are surprised that an experienced ornithologist expresses 64 doubt about the possibility of tracking movements of birds in their tropical deciduous forest 65 habitat. We always had three observers for these trials, and they were all extraordinarily adept women with keen and experienced observational skills. The birds usually sang and called during 66 the trials, further revealing their location. Third, the pseudoreplication criticism is a red herring. 67 68 Each subject's stimulus exemplars were uniquely prepared from a different base song, and we 69 used a wide variety of song and trill types and source males for the base song, thus preventing

70 pseudoreplication. Fourth, our modification of trill rate involved increasing or decreasing the silent gap between trill notes to a similar degree, so both alternative stimuli had an equivalently 71 72 altered note duration to silent interval ratio. The minimal difference in trill duration was unlikely 73 to be important, as individual birds naturally vary the number of notes in the trill as well as the 74 silent gap between notes. Moreover, the theoretically more intense signal had the shorter 75 duration, a conservative experimental design strategy. Fifth, we did examine the tendency for 76 trill performance components to vary in a consistent way among song types within males in the 77 Vehrencamp, Yantachka, Hall & De Kort (2013) study, and we found largely consistent 78 differences related to male age. Sixth, Kroodsma appears unable to consider that subjects that 79 initially approached the fast stimulus would subsequently spend less time close to the speaker if 80 the stimulus was a broad bandwidth (low vocal deviation) song. The only logical conclusion for 81 this result was that the subjects were more strongly repelled by the repeated playback of higher 82 performance trills. This repelling effect was subsequently verified by de Kort, Eldermire, Cramer, & Vehrencamp (2009), as discussed below. The approach – negotiate – retreat sequence 83 84 is typical of banded wren interactions (Vehrencamp et al., 2014). We know that the birds are still 85 interested in the stimulus when they back off because they keep singing, albeit from a distance. 86 Nuanced responses may be typical of two-speaker playback to rival male subjects (Reichert, 87 2011). Seventh, Kroodsma asks how we know over what time frame songs should have an effect. 88 We had conducted numerous playback experiments on this species, with stimulus durations 89 ranging from 1-2 minutes and post playback periods ranging from 3-5 minutes, by which time all 90 subjects had usually left the area. We had also observed many natural interactions between males 91 lasting from about 2-10 min. In Vehrencamp et al. (2014) we quantified such interactions and 92 found that interactions from start to finish ranged from 26 to 828 sec with a median of 161 sec.

93 So the Illes et al. experimental protocol, with a playback period averaging 86 sec and a post playback period equal to 180 sec, was well within the typical time frame of approach and 94 95 withdrawal for male-male encounters in the banded wren. Finally, Kroodsma argues that we 96 should have corrected all of the statistical tests in the entire results section with a Bonferroni 97 multiple comparisons procedure. It is commonly acknowledged that the Bonferroni correction is 98 far too harsh (i.e., Moran, 2003; Narum, 2006); the False Discovery Rate correction is superior in 99 reducing Type II errors, and we have done this correction in our papers where multiple variables 100 were tested and presented in tables. In the Illes et al. paper, our analyses were generated from 101 three independent datasets and addressed completely different questions, thus they should not be 102 combined as Kroodsma proposes. Some of the tests related to the plavback experiment were 103 presented to examine and dispel potential confounding effects. The remaining few tests 104 addressed specific hypotheses and were not part of a multivariate fishing expedition to find the 105 most significant variables. We think that a multiple comparison correction was not needed here. 106 In addition, we presented power analyses and effect sizes for our tests, and these revealed 107 stronger effects than the p-values indicate. Our study therefore cannot be criticized on the basis 108 of unreasonable claims of unnatural stimuli, pseudoreplication, incompetent observers, and 109 faulty statistics.

As skeptical scientists in search of the truth, we set out to further examine the interesting results in Illes et al. (2006) with another playback experiment that manipulated only the frequency bandwidth of trills (de Kort, Eldermire, Cramer, & Vehrencamp, 2009). Contrary to Kroodsma's claim, this study *was* conducted with observers blind to the bandwidth treatments, and the differences could not be detected by the observers. We separately presented three alternative bandwidth stimuli to subjects, and expressly quantified multiple measures of male

116 response to address the significance of nuanced retreat responses. The results strongly confirmed 117 the earlier study: subjects avoided high-performance stimuli but continued to sing and call from a 118 distance; approached and negotiated with matching songs to the median-performance stimuli; 119 and approached quickly but showed a lower vocal response to the low-performance stimuli. 120 Another pair of experiments explored male responses to songs of different trill note 121 consistency (de Kort, Eldermire, Valderrama, Botero, & Vehrencamp, 2009). One experiment 122 used natural songs of the same type and from the same male in his first year versus in his second 123 or third year, and the second experiment compared songs of first year birds to the same song 124 manipulated to have greater note consistency. Both experiments found stronger responses to the 125 more consistent song stimuli. As mentioned earlier, we showed in Vehrencamp et al. (2013) that 126 trill note consistency of all measured song types increased in males from their first to their 127 second and third year, and then plateaued or decreased slightly for older birds. Male age is not 128 only associated with territorial defense experience, which could be assessed during territorial 129 encounters, but multiple lines of evidence also suggested that females avoided mating with or 130 divorced first year males and preferred older and more consistently singing males as extra-pair 131 partners (Cramer, Hall, de Kort, Lovette, & Vehrencamp, 2011).

Kroodsma's criticism of the language in the first sentence of the abstract of (Vehrencamp et al., 2013) is totally unwarranted. Abstract wording is necessarily brief. We merely stated the general theoretical proposition being tested in our study, a standard protocol for scientific articles. Kroodsma doesn't appear to question the results, which showed that trill note consistency and frequency bandwidth increase with male age. We also discovered that trill note rate for a given song type increases during playback experiments in relation to the male's level of aggressive response, a result that has now been found in other species (Funghi, Cardoso, &

Vehrencamp et al. 8

Mota, 2015; Linhart, Jaska, Petruskova, Petrusek, & Fuchs, 2013). Thus this aspect of
performance seems to provide cues to receivers about a rival's immediate aggressive motivation.
We did not find any associations with male survival or our measure of body condition. Our
results and interpretations were not biased by any desire to support or disprove the hypothesis,
and in several instances we offered alternative hypotheses where appropriate.

144 Commenting further on this paper, Kroodsma writes (his italics): "According to the 145 scatterplot of trill rate and bandwidth for banded wrens (Fig. 11), relatively few songs are 146 difficult to execute as defined in this performance context, because most songs fall far from the 147 upper bound on the graph. Every male 'willingly' learns many 'low-performance', easy-to-148 execute songs in order to have particular song types in his repertoire, as if performance did not 149 *matter*, as if there were no selection for difficult-to-execute songs as claimed in this paper". 150 Repertoire species such as the banded wren use contrasting song types to emphasize switching 151 rates, short-term diversity, and matching during territorial interactions. But Kroodsma has 152 conveniently ignored another component of banded wren trills: their varied and complex note 153 shapes as mentioned earlier. Trill note consistency is a third axis of performance in this species, 154 and we showed in this paper (Vehrencamp et al., 2013, see supplementary online material) that 155 consistency and vocal deviation trade-off (are negatively correlated) within male and song type. 156 Thus song types far from the trill rate vs. bandwidth upper limit are not necessarily easy to 157 execute, as they may have a complex shape that is difficult to repeat consistently.

Kroodsma surprisingly concludes that "we await good answers to the question... of what information listeners extract about singers from their songs (beyond species identification)". In fact, there is a growing body of data showing that aspects of vocal performance, including trill rate, vocal deviation, trill note and song consistency, and call rate/call duration trade-offs, do

162	provide useful information to receivers in some species and are associated with reproductive
163	benefits in many birds, mammals, anurans, and crickets (e.g., Botero, et al., 2009; Byers, Akresh,
164	& King, 2015; Funghi et al., 2015; Linhart et al., 2013; Pasch, George, Campbell, & Phelps,
165	2011; Petruskova et al., 2014; Reichert & Gerhardt, 2012; Sprau, Roth, Amrhein, & Naguib,
166	2013; Wagner, Beckers, Tolle, & Basolo, 2012; Welch, Smith, & Gerhardt, 2014). Our studies
167	have contributed to this body of knowledge, specifically by demonstrating the existence of cues
168	to age and aggressive motivation, along with the strategic use of song type use patterns to
169	indicate approach and retreat during territorial negotiations.
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