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2	Hens vary their vocal repertoire and structure when anticipating different types of
3	reward
4	Nicky McGrath ^a , [*] Rebecca Dunlop ^a , Cathy Dwyer ^c , Oliver Burman ^b , Clive J.C. Phillips ^a
5	^a School of Veterinary Sciences, University of Queensland, Gatton, Queensland, Australia
6	^b School of Life Sciences, University of Lincoln, Lincoln, U.K.
7	^c Scotland's Rural College (SRUC), Edinburgh, U.K.
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12	*Correspondence: N. McGrath, School of Veterinary Sciences, University of Queensland,
13	Gatton, 4343, Queensland, Australia
14	E-mail address: nicmcgrath@hotmail.com
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27 Abstract

The vocalizations of nonhuman animals are considered potential indicators of motivational or 28 internal state. In many species, different call types, and structural variation within call types, 29 30 encode information about physical characteristics such as age or sex, or about variable traits such as motivation. Domestic chickens, Gallus gallus, have an elaborate vocal repertoire, 31 enabling investigation into whether reward-related arousal is encoded within their call type 32 and structure. Twelve hens were given a Pavlovian conditioning paradigm using sound cues 33 to signal the availability of two food rewards (mealworms, normal food), one nonfood reward 34 (a container of substrate suitable for dustbathing), and a sound-neutral event, (sound cue, no 35 reward). A muted-neutral treatment (no sound cue, no reward) provided a baseline for vocal 36 behaviour. Sound cues preceded a 15 s anticipation period during which vocalizations were 37 recorded. Hens produced a 'Food call' (previously defined in other studies) in anticipation of 38 all rewards, including the nonfood reward. 'Food calls' and 'Fast clucks' were more prevalent 39 in anticipation of rewards, and most prevalent following the cue signalling the dustbathing 40 substrate, suggesting that this reward induced the most arousal in hens. The peak frequency of 41 'Food calls' made in anticipation of the dustbathing substrate was significantly lower than 42 those made in anticipation of food rewards, potentially reflecting differences in arousal. 43 Vocalizations that reliably indicate hens' motivational state could be used as measures of 44 welfare in on-farm assessment situations. Our study is the first to reveal variation in the 45 frequency-related parameters of the 'Food call' in different contexts, and to show prevalence 46 of different call types in reward and nonreward contexts, which may have implications for 47 welfare assessments. 48

Keywords: laying hens, motivation, referential, rewards, vocalizations.

Animal vocalizations contain information encoded within parameters such as frequency 50 (Townsend, Charlton, & Manser, 2014), amplitude (Gustison & Townsend, 2015; Reichard 51 & Anderson, 2015), rate of production (Clay, Smith, & Blumstein, 2012) duration 52 53 (Dentressangle, Aubin, & Mathevon, 2012) and energy distribution (Linhart, Ratcliffe, Reby, & Špinka, 2015). Some information may be static, relating to age, sex or body size (Briefer & 54 McElligott, 2011; Charlton, Zhihe, & Snyder, 2009). However, variation in these acoustic 55 parameters may also provide 'markers' of internal states (Manteuffel, Puppe, & Schon, 2004; 56 Tallet et al., 2013). Internal states, in this context, refer to states of arousal induced by both 57 internal (e.g. hunger) and external (environmental) stimuli and the interactions between them. 58 Such states exist on a continuum, with arousal levels being in constant flux according to 59 changes in stimuli and internal adjustments to these stimuli (for a detailed discussion see 60 Berridge (2004)). 61

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Flexible traits, such as signaller motivation, can be reflected in vocal frequency, amplitude, 63 duration and rate in both humans (Scherer, 1986) and animals (Briefer, 2012; Taylor & Reby, 64 2010), as shown in meerkats, Suricata suricatta (Hollén & Manser, 2007) and rats, Rattus 65 norvegicus (Knutson, Burgdorf, & Panksepp, 2002). The flexible features of vocalizations 66 67 tend to be subject to certain 'motivation-structural rules' (Morton, 1977). According to this concept, vocalizations produced in one motivational context (e.g. a hostile situation) should 68 vary in structure from vocalizations produced in a very different motivational context (e.g. 69 friendly interactions; Morton, 1977). This theory has been tested in many species including 70 domestic dogs, Canis lupus familiaris (Yin & McCowan, 2004), chimpanzees, Pan 71 troglodytes (Siebert & Parr, 2003), coatis, Nasua nasua (Compton, Clarke, Seidensticker, & 72 Ingrisano, 2001) and elk, Cervus canadensis (Feighny, Williamson, & Clarke, 2006). For 73 most of these animals, the call types commonly produced in hostile contexts are long in 74 duration with a low frequency, whereas in fearful or nonaggressive contexts, short, high-75

frequency, tonal calls are produced (see review by Briefer, 2012). In addition to different call
types, motivation may also be encoded within a sound's structure. For example, the frequency
of particular sounds may be lowered when a signaller is feeling aggressive (Bee & Perrill,
1996).

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Certain animal vocalizations function referentially, passing information about specific 81 environmental stimuli to receivers (Macedonia & Evans, 1993). To fulfil the criteria for 82 functional reference, calls must be elicited by a narrow range of stimuli, and evoke a response 83 in the receivers as if they had experienced the stimuli themselves (Evans, 1997; Macedonia & 84 Evans, 1993; Marler, Evans, & Hauser, 1992). For example, some species of primate produce 85 different call types according to particular threats or predators (Murphy, Lea, & Zuberbühler, 86 2013; Zuberbuhler, Noe, & Sevfarth, 1997), which elicit distinctive adaptive responses in the 87 receivers. Functionally referential vocalizations may also encode motivational information at 88 the same time as being referential (Hollén & Manser, 2007). In these instances, the call rate or 89 structural variation within call types may encode information relating to arousal, motivation 90 or urgency (Clay et al., 2012; Manser, 2001). Townsend and Manser (2013) described a 91 motivational-referential continuum, giving the example of meerkats which produce calls that 92 refer to nearby predators and also deliver information about the urgency of the threat. The 93 'Food call' of the chicken, Gallus gallus (Collias, 1987; Evans & Marler, 1994) has been 94 described as the most rigorous example of food-specific functional reference within terrestrial 95 animals (Clay et al., 2012). This call appears to meet all the criteria of functional reference; 96 there is acoustic specificity between the stimulus (food) and the signal, and playback elicits 97 feeding-related behaviours in receivers (Clay et al., 2012; Townsend & Manser, 2013). It is 98 also likely that the 'Food call' contains motivational information in its acoustic variables. 99

The anticipation of rewards is thought to increase arousal in animals induced by changes in 101 motivational state, and this, in turn, may elicit vocalizations. Rats, for example, produce 102 ultrasonic vocalizations at 50 kHz in anticipation of rewards such as the presence of a play 103 104 partner, during tickling from a familiar human or in response to a cue signalling food (Burgdorf & Panksepp, 2006; Knutson, Burgdorf, & Panksepp, 1998; Panksepp & Burgdorf, 105 106 2000). By contrast, negative stimuli such as the presence of a predator, or a cue signalling an 107 electric shock, tend to elicit a 22 kHz vocalization in rats (Knutson et al., 2002). This knowledge, theoretically, allows us to assess whether a rat is in a 'rewarding' environment, 108 which has implications when determining their welfare. Clear indicators about whether 109 110 animals are experiencing rewarding or nonrewarding environments may guide decisions made within management systems. The anticipation of rewards induced by signals has been linked 111 to 'pleasure-based' (dopaminergic) activity in the brain (Berridge, 1996). Therefore, while 112 stressful environments are known to have deleterious effects on productivity (Broom, 1991), 113 it is reasonable to assume that the reduction of stress through the provision of rewarding 114 115 environments could positively influence animal health and productivity (Boissy et al., 2007). 116

The domestic chicken is a good candidate for the study of vocalizations made in anticipation 117 of rewards for three main reasons. First, the behaviour of chickens in anticipation of rewards 118 has already been well documented (Garland, Castellote, & Berchok, 2015; Moe et al., 2014; 119 Moe, Nordgreen, Janczak, Spruijt, & Bakken, 2013; Moe et al., 2009; Zimmerman, Buijs, 120 Bolhuis, & Keeling, 2011). Second, behavioural data suggest that the motivational state of 121 chickens changes according to the type of reward (Garland et al., 2015; McGrath, Burman, 122 Dwyer, & Phillips, 2016), and therefore it may be possible to link anticipatory behaviour with 123 vocalizations produced in anticipation of rewards. Third, the chicken has a wide and varied 124 vocal repertoire. Between 20 and 25 discrete calls have been documented in various studies 125 (Collias & Joos, 1953; Evans, 1993; Evans & Evans, 1999; Kruijt, 1964; Marx, Leppelt, & 126

Ellendorff, 2001; Woodgush, 1971) including those classed as referential (Evans & Evans,
2007). Interestingly, it seems that motivational information may be encoded within referential
calls. Alarm calls, for example, differentiate between terrestrial threats and aerial predators
(Evans, Evans, & Marler, 1993) while simultaneously encoding the motivational state of the
bird (Kokolakis, Smith, & Evans, 2010).

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Chickens have been shown to modify the rate and numbers of their 'Food calls' in response to 133 different types of food reward (Marler, Dufty, & Pickert, 1986; Wauters, Richard-Yris, Pierre, 134 Lunel, & Richard, 1999). These changes appear to reflect variation in motivational state 135 according to food type, which indicates there is an opportunity to test for motivational 136 information within a referential call. The'Food call' of chickens is described by various 137 authors as having a characteristic appearance consisting of trains of pulsatile calls delivered in 138 a regular temporal pattern, emphasizing low frequencies, and at a rate of 4–10 per s (Collias, 139 1987; Evans & Marler, 1994). They are given by both males and females, and their structure 140 facilitates location of the sender (Hughes, Hughes, & Covalt-Dunning, 1982). No studies have 141 investigated vocalizations produced by chickens in anticipation of other types of reward, or 142 variation in their acoustic parameters. Therefore, the goal of this study was to characterise 143 vocalizations made in anticipation of different types of reward (different food types known to 144 be 'rewarding' to chickens, Bruce, Prescott, & Wathes, 2003, and a substrate suitable for 145 dustbathing). We aimed to investigate whether call parameters varied within the call types 146 according to reward type, and according to whether it was a reward or nonreward. To achieve 147 this, we experimentally induced anticipation of these rewards, using a Pavlovian conditioning 148 paradigm. We tested the hypothesis that chickens would produce specific call types in 149 anticipation of different rewards. We also tested the hypothesis that vocal parameters would 150 differ according to the perceived quality of the reward. These differences would provide 151

152 information on the motivational state of the signaller, and therefore could be used as

153 indicators of baseline welfare.

154 Methods

155 Subjects and Housing

Twelve ISA Brown hens, approximately 18 weeks old, were obtained from the University of 156 Queensland's poultry unit. The hens were housed in groups of three in pens measuring 266 x 157 266 cm and 133 cm high. The floor of the home pen was shredded rubber chip, and each pen 158 contained a metal structure used as a perch (149 x 119 cm and 41 cm high) and two nestboxes 159 (40 x 35 cm and 45 cm high). Food (standard layer pellets) and water were available ad 160 libitum in the home pens. The housing had natural as well as artificial light (the latter on 161 between 0600 and 1800 hours). There was no temperature control, but all experimental work 162 was conducted between 0800 and 1230 hours to standardize the conditions. Hens were 163 individually identifiable to the experimenter (N.M.) by plumage colouring, marking and comb 164 165 size, avoiding the need for individual marking or ringing.

166

167 Ethical note

The methods used in this study were approved by the University of Queensland Animal 168 Ethics Committee (Ref. SVS/314/12). The experiment used a minimum number of hens based 169 on related published studies. The ISA Brown hen is a commonly used strain in production 170 environments, and therefore conclusions about welfare would have a general application. 171 Hens were not marked and the experimenter used natural markings on the hens for individual 172 173 recognition. The procedures used in this experiment could have exposed hens to stress through handling when moving from the home pen to the experimental pen, and when 174 isolated in the experimental pen. The experimenter was trained in handling chickens, and 175 176 therefore any such stress was minimized, and the experimental procedure was, by its nature,

mainly rewarding, with no aversive conditions used. Cleaning was performed routinely every
few days. Hens habituated quickly to these procedures. They were housed in large pens, each
furnished with a perch, a tray of dustbathing substrate and two nestboxes to cater for natural
behaviour. In addition, they could access a cauliflower or broccoli hanging from a piece of
string above the ground as food enrichment.

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183 Treatment Groups

Hens were subjected to a Pavlovian conditioning paradigm as used by Zimmerman et al. 184 (2011) and Moe et al. (2009). In our experiment, an initially neutral stimulus (conditioned 185 stimulus, CS) was repeatedly paired with the presentation of one of three different rewards 186 (mealworm, normal food or dustbathing substrate) or a sound-neutral event (an empty 187 compartment) which served as the unconditioned stimuli (US). We used different sound cues 188 for the conditioned stimuli all 5 s long: 'ring' (ringing of a rotary dial telephone), 'beep' (an 189 alarm-clock style beep), 'buzz' (a buzz sound as in a game show) and 'horn' (an old-190 fashioned car horn sound). A muted neutral (MN) treatment (5 s of 'nothing', no CS or US) 191 was used to control for the effect of sound in the other treatments. The sound cues were 192 played from a computer at a sound pressure level of 75 dBA, measured 1 m from the centre of 193 the experimental chamber. Each of the four sound cues was used to signal the presence of 194 each type of US. Consequently, four cue groups of three hens (from the same home pen) each 195 experienced different combinations of CS and US (Table 1). 196

197 Experimental Apparatus

An experimental pen (200 x 125 cm and 60 cm high) was located in a sound-proofed room adjacent to the room in which the hens were housed. The pen contained two compartments of equal size, a waiting compartment and a reward compartment, separated by a wire-mesh partition and connected by a swing door in the middle of the partition (Fig. 1). The door could be locked and released by increasing or decreasing an electrical current passing through an electromagnet attached to it, and only opened in the direction of the reward compartment.
Three of the four walls of the experimental pen were made of plywood and one was made of
wire-mesh to allow video recordings of both compartments. A lamp, secured to the middle of
the outer wall at 60 cm from the floor, could be operated by the experimenter who sat behind
a screen out of sight of the hens during tests. This light shone into the reward compartment
and was used to highlight the reward and indicate that the door was open.

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The apparatus used for rewards comprised a white food bowl, a tray filled with topsoil/sand mix and the hens' normal feeders, enabling hens to discriminate between reward types. The topsoil/sand mix was chosen after a review of the literature on functional substrates for dustbathing, and its dry crumbly texture suggested that it was suitable for this purpose (Olsson & Keeling, 2005). Duplicates of the white food bowl containing food and a white tray filled with the topsoil/sand mix were put in the home pens 3 days before training started to allow hens to become accustomed to them.

217

218 **Training Procedure**

The training consisted of several phases similar to those used by Zimmerman et al. (2011) 219 220 with adjustments in the length of each phase due to the number of conditioned stimuli used. As a result of the hens learning to enter the reward compartment more slowly, training took 221 place over 25 days (compared with 22 days in Zimmerman et al. 2011). In phase 1 (Days 1 -222 7) hens were trained to use the swing door. This initial training was done in groups of three to 223 increase the speed of training. During the first 2 days the door was kept fully open, and a trail 224 225 of sunflower seeds led through the door into the reward compartment where the white bowl held more sunflower seeds. Each home pen group of three hens was allowed to accustom 226 themselves to moving from the waiting compartment to the reward compartment following 227 the trail of seeds. On the following 5 days, home pen groups were trained to go through the 228

door, the opening width of which was gradually reduced more on each day. During this
period, the hens were food deprived for an average of 2 h and mealworms were placed in the
white bowl in the reward compartment to motivate them to go through the door. Each group
of three hens experienced eight consecutive trials. On the final day, the door was fully closed,
but unlocked, so that the hens had to push through it to gain access to the mealworms.

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In phase 2 (Days 8–13), the hens were individually trained to recognize the specific CS for 235 each of the rewards. The containers containing the dustbathing substrate were removed from 236 the home pens from this time onwards. In this phase, the door was kept unlocked and hens 237 were given 10 min to go through the swing door after their particular CS for mealworm, 238 dustbathing substrate or normal food was played and the light switched on. All hens entered 239 the compartment within the 10 min time limit. After consumption of the reward, the light was 240 switched off and the hens were guided back into the waiting area by the experimenter. Each 241 cue group was trained for one stimulus on one day and each individual experienced three 242 243 consecutive trials. Hence, during the 6 days, each hen experienced six trials of each stimulus with the door unlocked. In phase 3 (Days 14–17), the swing door was locked, and the CS and 244 light signal were made contingent on the behaviour of the hen. Hens were placed individually 245 246 into the waiting compartment and allowed to try to push through the locked door twice before the CS was played, the light was switched on and, simultaneously, the door was unlocked 247 (CS/Light/Door). This procedure was repeated three times and then the CS /Light/Door 248 sequence was performed at random when the hen was not near the door. When the hen went 249 through the door immediately after the CS/light was given in five consecutive trials, the 250 251 training session was ended and hens were returned to their home pen. In phase 4 (Days 18– 19), a trace conditioning procedure (Moe et al., 2009) was used to accustom the hens to an 252 interval between the CS being played and activating the light signalling the door was 253 unlocked. The CS was played for 5 s and the interval between the end of the CS and the light 254

signal was gradually increased from 0 to 15 s over five consecutive trials for each individual 255 hen on each day. Hens were deemed to have successfully reached our criterion when they 256 went through the swing door within 5 s after the light had been switched on. In phase 5 (Days 257 258 20-22), all hens were introduced to their sound-neutral CS. The sound-neutral CS was not introduced in an earlier phase to aid consolidation of learning with regard to going through 259 260 the door after the trace delay. In these sound-neutral trials, nothing happened after the light had been switched on. The light was kept on for 15 s and then switched off. In phase 6 (Days 261 23–25), rewarded (mealworm, dustbathing substrate, normal food) and sound-neutral trials 262 were presented in a randomized order, with each cue being presented at least once to each hen 263 on each day. Hens successfully reached our criterion when they went through the swing door 264 within 5 s on every rewarded trial. Hens learned at different speeds during this training phase; 265 however, there were no consistently 'slow learners' for all phases, and training schedules 266 were adjusted to ensure that all hens progressed to the next stages together, so that there was 267 no bias in exposure to stimuli for individual hens. 268

269

270 **Test Procedure**

For testing, a hen was collected from her home pen and put in the experimental pen. The 271 272 order of testing was determined using an orthogonal Latin square design in which every single condition followed another on two occasions. Each hen received one test session per day on 5 273 consecutive days. Hens were deprived of food for an average of 1.5 h prior to testing, and 274 deprived of a substrate suitable for foraging and dustbathing in their home pens for all 5 days 275 of the test period. Each test session consisted of presentation of each of five stimuli: three 276 277 reward treatments (mealworm, dustbathing substrate, normal food), one sound-neutral and one muted-neutral trial. At the start of each session a hen was allowed to habituate to the 278 experimental pen for 30 s. Then the appropriate CS was given for 5 s, after which behaviour 279 and vocalizations were recorded for 15 s before the light was switched on signalling the door 280

was unlocked. There was no CS in the muted-neutral trial, but behaviour was recorded for 15 281 s from when the trial started. In the mealworm trial, after the CS and the 15 s anticipation 282 period, the door was unlocked and the hen entered the reward compartment and ate the 283 284 mealworm. Then the light was switched off and the hen was ushered gently into the waiting compartment by the experimenter who held the swing-door open. In the normal food trial, the 285 same happened except that the hens were allowed 1 min to feed before the light was switched 286 off and the hen was returned to the waiting compartment. In the dustbathing substrate trial, 287 the same process was followed except that the hens were allowed to dustbathe or forage (with 288 no food present) for 5 min before the light was switched off and the hen was returned to the 289 waiting compartment. If the hens stopped feeding or foraging/dustbathing and walked away 290 from the stimulus, or engaged in other behaviour in other parts of the pen for a continuous 291 period of 10 s, then the trial was ended. In a sound-neutral trial, the CS was given and, after a 292 15 s anticipation period, the light was switched on but the door did not open. In a muted-293 neutral trial, no sound cue was given, and behaviour and vocalizations were recorded for 15 s. 294 295 In all trials, between the end of the trial and the start of a new waiting period, there was an intertrial interval of 10, 20 or 30 s (balanced between hens), to prevent hens from anticipating 296 the start of the next trial, 297

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During the test procedure, vocalizations were recorded using a Sennheiser ME66 condenser
shotgun microphone connected to a Tascam DR100 MkII DAT recorder. Gain settings were
set to High and the rotary dial input gain setting was set at 6. Recordings were made with 24
bit resolution at a sampling rate of 44.1 kHz. The DAT files were transferred to a PC (Dell) to
analyse the vocalizations using Raven Pro: Interactive Sound Analysis Software (Version 1.5,
Cornell Laboratory of Ornithology, Ithaca, NY, U.S.A.).

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306 Acoustic analysis

Spectrograms of recordings were generated using Raven Pro 1.5 (Cornell Laboratory of 307 Ornithology; 1792-sample Hann window, 35.4 Hz filter bandwidth, 5.38 Hz frequency 308 resolution (grid spacing), discrete Fourier transform (DFT) size of 8192 samples and time grid 309 310 hop size 200 samples (88.8% overlap)). Measurements were made in Raven Pro and values were imported into Excel by syllable. A syllable is defined as a sound that makes a 311 continuous impression in time on the spectrogram. A call is defined as a clearly discernible 312 sequence (both audibly and visually) of syllables grouped together. Recordings with 313 extensive background noise (N=94) were excluded from further analysis (Fig. 2). The 314 difference in percentage of calls rejected by treatment was not significant ($\chi^2 = 0.26$, P>0.05). 315 No vocalizations were elicited in 23 of the 300 treatments, and this number was distributed 316 across contexts as follows: dustbathing substrate: 5; mealworm: 6; normal food: 2; sound-317 neutral:3; muted-neutral: 7. 318

319

320 Call Classification

321 The call classification process followed those used in previous studies, including Garland et al. (2015) and Rekdahl, Dunlop, Noad, and Goldizen (2013). There are no fully automated 322 classification procedures that do not include a qualitative component. Manual classification is 323 commonly used as a starting point as this method picks up the subtle features of the sounds 324 that quantitative measures will miss. However, because of these subtle differences, there is a 325 need to test the robustness of a manual classification. In this study, we first manually 326 classified the sounds and then carried out a classification and regression tree analysis (CART) 327 and Random Forest analysis to corroborate this initial classification (see below). A high 328 percentage agreement between the manual and quantitative methods suggests that the manual 329 classification consistently and correctly groups the sounds into types, and therefore is robust 330 and repeatable. A total of 546 calls were manually classified. The data were then grouped by 331

call in Excel and calculations were made for the temporal variables. Frequency variables wereaveraged across calls. All variables measured are detailed in Table 2.

334

335 Descriptive analyses were performed on call types (Appendix Table A1). Example
336 spectrograms of these call types were visually compared with already documented chicken
337 calls (Collias, 1987; Evans and Evans, 2007).

338

Hens have distinct call types within their repertoire, but some calls also blend from one type 339 into another on a continuum of calls. Therefore, we created a 'Mixed' call category to include 340 these calls. The 'Food call' given by chickens was identified by comparing our recordings 341 with published spectrograms (Evans & Evans, 2007; Sherry, 1977). These calls were 342 characterized by a consistent fast rhythm and low pitch (Table A2). Of a total of 60 'Food 343 calls', 24 were made by one hen. To reduce bias, 16 of this hen's food calls (randomly 344 chosen) were removed from the analysis to better balance the number of calls made by 345 346 individuals.

347

To validate the manual classification of calls, we used a nonparametric CART and Random 348 Forest analysis. Applying the CART method, a decision tree was constructed using the rpart 349 package in R, which does not require independence of samples. In a decision tree a set of 350 hierarchical decision rules is created. Each rule can branch into another rule or a terminal 351 category. At each step, the split is made based on the independent variable that results in the 352 largest possible reduction in heterogeneity of the dependent variable; this is called the 'Gini 353 index', which measures impurity or 'goodness of split' (Breiman, Friedman, Stone, & Olshen, 354 1984). The optimal decision tree must contain criteria that classify the data as accurately as 355 possible, without 'overfitting' the data. Therefore, we pruned the tree to minimize 356 misclassification rate, by choosing a complexity parameter that minimizes the cross-validated 357

error. We also conducted a Random Forest analysis using the randomForest package in R 358 (Liaw & Wiener, 2002). This classification tool randomly selects a subset of predictor 359 variables and creates a 'forest', or a collection of decision trees (Breiman, 2001). The 360 361 aggregation of trees evaluates the classification uncertainty of each tree [the out of bag (OOB) error] in addition to ranking the importance of each predictor variable (i.e. the call 362 variables we measured; Breiman, 2001). As Random Forests estimate error internally, no 363 cross-validation is required (Breiman, 2001). Based on the lowest OOB error, the number of 364 predictors randomly selected at a node for splitting was set to three and 1000 trees were 365 grown. Classification success was evaluated using the overall OOB error rate and individual 366 call type errors. 367

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369 Statistical analyses of call types

370

All analyses of call types were performed in R (R CoreTeam, 2013). The prevalence of call 371 372 types used by hens was calculated by treatment. Prevalence of call types was analysed using a generalized linear mixed model (GLMM) within the lme4 package (Bates et al., 2015) with 373 'treatment' as the predictor variable and each call type as the response variable with subject 374 375 ID as the random effect. A mixed model was used to account for the within-bird variance associated with repeated measurements. A negative binomial model was used as the most 376 appropriate method to analyse these data, as these models allow for overdispersed data. 377 Incident rate ratios and 95% confidence levels were extracted, and mean predictions were also 378 checked against observed data to ensure they did not deviate, as deviation would indicate a 379 380 poorly fitting model. We used the function glmmADMB to run all negative binomial models. 381

To determine whether hens varied their call structure in anticipation of different rewards, weanalysed variation of acoustic parameters within call types. Only the call types 'Single

clucks', 'Double clucks', 'Fast clucks', 'Food calls', 'Whines' and 'Gakel calls' were used in 384 this analysis. 'Singing' calls (N=16) were eliminated due to the small number of calls. 385 'Mixed' calls (N=124) were also excluded as their structure was highly variable and therefore 386 387 they were difficult to classify. Many of the acoustic variables were highly correlated. Therefore, we conducted a factor analysis (FA) with varimax rotation using the function 388 'factanal'. Acoustic variables were log transformed to better match hearing perception in 389 390 vertebrates (Cardoso, 2013), and plots were examined for linear relationships between variables. The Kaiser-Meyer-Olkin (KMO) measure of sampling adequacy (MSA) was used 391 to distinguish which variables should be included in the FA. This measure indicates whether a 392 variable belongs to a family of variables psychometrically (Dziuban & Shirkey, 1974). 393 Variables with MSA scores below 0.5 indicate that an item does not belong to a group and 394 may be removed from the FA. Following these criteria, the variable 'Number of syllables' 395 (MSA score of 0.35) was removed from the FA. 396

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398 In a preliminary FA, the model did not converge because variables 'Centre Frequency', 'Q1 frequency' and 'Q3 frequency' had very low levels of uniqueness (less than 0.0000001). 399 These variables were removed, and the models were run on the remaining variables, moving 400 401 from a one-factor up to a four-factor model. 'Call duration' and 'Average Syllable Length' loaded on to separate individual factors and were therefore excluded from the FA and tested 402 as separate responses. The overall KMO test output of 0.69 and Bartlett's test of sphericity 403 (P<0.001) indicated that the FA was justified (McGregor, 1992). Using the package 404 nFactors(), the optimal number of factors as determined by both eigenvalues and parallel 405 analysis was two. 'Bandwidth 90%', 'Delta Frequency' and 'IQR Bandwidth' loaded on to 406 Factor 1, with 'Bandwidth 90%' loading the highest (0.98), and 'Peak', 'Low' and 'High 407 Frequency' loaded on to Factor 2, with 'Peak Frequency' loading the highest (0.96). To 408 409 facilitate interpretation of results, the variables that loaded highest on to each factor,

'Bandwidth 90%' (from Factor 1) and 'Peak Frequency' (from Factor 2), were used for the
subsequent call structure analysis. 'Call Duration' and 'Average Syllable Length' were also
analysed to detect whether hens altered the temporal structure of their calls in anticipation of
different rewards.

414

Variation in acoustic parameters for each call type was analysed using linear mixed-effects 415 416 models (LMMs) from within the lme4 package (Bates et al., 2015). The assumptions of normal distribution, linearity and homoscedasticity of the residuals were checked by visual 417 inspection of residual plots and by Shapiro–Wilks tests. We computed parameter estimates 418 419 using the maximum likelihood method, and the significance of predictor variables was tested using maximum likelihood ratio tests (anova() function in R). Each response variable was 420 modelled separately. For all LMMs the chi-square statistic (γ^2), degrees of freedom and 421 associated P values are reported. (Full results are available in Appendix Table A3.) Post hoc 422 analyses were conducted using the lsmeans package (version 2.20-23) in R, applying the 423 424 Tukey method for individual comparisons of different factor levels within the same model, incorporating an adjustment of P values for multiple comparisons. A Bonferroni correction 425 was applied to account for the number of variables being tested using the same data set, and 426 427 therefore *P* values of and below 0.01 were considered significant.

428

Model selection was carried out using Akaike information criterion (AIC) scores, and by
performing chi-square tests to compare models using the anova() function, with values of *P*<0.05 considered significant (Appendix Table A4). Initial models included 'Preceding</p>
Treatment' as a predictor variable to establish whether there were any carryover effects
between treatments. As there was no significant effect of preceding treatment, this predictor
was left out of the final models. Final models included the following sets of predictor
variables: (1) Treatment only, (2) Treatment and Cue Group, (3) Treatment, Cue Group and

436 Day. Model comparison and AIC scores are detailed in Appendix Table A4. Residual plots

were checked by using the plot() function to check for any patterns in the data. All finalmodel outputs are detailed in Appendix Tables A5–A10.

439

440 **Results**

441 Call classification

Hens produced eight different call types according to both the manual classification and 442 CART analysis, suggesting a high agreement between the two methods and a high probability 443 this call repertoire is 'correct'. These ranged from the very short 'Single cluck', with a call 444 duration of 0.08 s, to a long 'Gakel call' (Zimmerman and Koene, 1998; 3.1 s). 'Single 445 clucks' and 'Double clucks' were short sharp clucks with either one or two syllables. Two 446 other 'cluck' type calls, the 'Fast cluck' and the 'Food call', were structurally similar to the 447 'Single cluck' and 'Double cluck', but consisted of a longer series of short sharp clucks (Fig. 448 6). The 'Fast cluck' was not as consistent in rhythm or pitch as the 'Food call'. 449

450

Hens also produced long, high-pitched calls such as a 'Whine' (Fig. 7), which was
characterized by a wide frequency range (delta frequency = 274 Hz), compared with the
'cluck' calls whose frequency range was from 162.9 to 173.7 Hz (Table A2). 'Gakel calls'
were also identifiable by having a harsh or whiny tone, and a high frequency range (246 Hz).
A small number (*N*=16) of calls identified as 'Singing' (Collias, 1987) also had a high pitch
(644.6 Hz) and frequency range (366.8 Hz). 'Mixed' calls could not be categorized into any
of the other groups due to their high structural variation.

458

All variables listed in Table 2 were available for construction of the CART decision tree, and
the variables CART selected in tree construction were 'Call duration', 'Maximum frequency',
'Maximum syllable length', 'Number of syllables', 'Peak frequency' and 'Average syllable

length'. The tree correctly classified over 69% of calls (Fig. 3). The first branch in the tree was based on call duration, which separated the cluck call types ('Food call', 'Fast', 'Single' and 'Double clucks') from the 'Gakel calls' and 'Whines'. Further branching was heavily influenced on the longer calls by duration of the call and maximum frequency, and on the shorter calls by the number of syllables and syllable length, as well as acoustic frequency parameters. The analysis resulted in eight terminal nodes (call categories), with 'Fast Clucks' being the most prevalent in two terminal nodes.

469

Random Forest correctly classified a high number of calls (OOB estimate error rate was 470 26.6%; Appendix Table A11). This represents a high level of agreement in classification with 471 our manual classification of calls. The most important variables used by Random Forest 472 classification were Call duration and Number of syllables (mean decrease in Gini index = 88 473 and 72, respectively), followed by Maximum syllable length and Average syllable length 474 (mean decrease in Gini index = 62 and 55, respectively; Appendix Table A12). The majority 475 of misclassifications occurred within the 'Singing' category which had a low number of 476 examples, and a high misclassification rate and this increased the measure of error. 'Fast 477 clucks', 'Food calls' and 'Mixed' calls also had a moderate misclassification rate when 478 479 compared with our manual classification. It was expected that 'Mixed' calls would be difficult to classify quantitatively, and this was reflected in the misclassification rate. 'Mixed' calls 480 contained a number of calls that represented different call types blending into others; for 481 example, 'Food calls' may blend into a short 'Whine'. 'Food calls' and 'Fast clucks' have 482 very similar structures. However, the CART analysis correctly classified 80% of 'Food calls'. 483 Only calls that demonstrated the regular temporal and acoustic structure described previously, 484 and that were visibly similar to a spectrogram published by Evans and Evans (2007), were 485 selected as 'Food calls' and analysed further. 486

488 **Prevalence of call types**

The call repertoire of hens varied significantly depending on the type of reward. The prevalence of 'Food calls' and 'Fast clucks' was significantly greater in anticipation of all rewards compared to both neutral treatments (Fig. 4a, b) suggesting that these call types are produced more when hens anticipate a positive event.

493

494 **Differences in call parameters**

As 'Food calls' were prevalent in reward contexts but scarce in neutral treatments (Fig. 4a), call parameters were only compared in anticipation of rewards. Only the peak frequency of 'Food calls' varied in anticipation of rewards, where they were produced at significantly lower peak frequencies in anticipation of the dustbathing substrate compared with the mealworm reward (dustbathing substrate: 411.85 Hz; normal food: 467.55 Hz; mealworm: 487.16; χ^2_2 =8.78, *P*=0.01; Fig. 5).

501

The 90% Bandwidth of 'Fast clucks' was significantly lower during all three reward 502 treatments than during the muted-neutral treatment (sound-neutral: 86.71 Hz; normal food: 503 89.96 Hz; dustbathing substrate: 90.04 Hz; mealworm: 98.86; muted-neutral: 126.81 Hz; 504 χ^2_4 =13.25, P=0.01). Interestingly, 'Fast clucks' were also produced at lower peak frequencies 505 in anticipation of the dustbathing substrate and mealworm rewards, which were presumably 506 high-value rewards, than in the muted-neutral treatment, although the results were not 507 significant at P<=0.01 (dustbathing substrate: 402.66 Hz; mealworm: 403.12; normal food: 508 409.19 Hz; sound-neutral: 56.59 Hz; muted neutral: 548.59 Hz; χ^2_4 =11.30, P=0.02). The 509 above variations suggest that these call types encode information about motivational state 510 according to different types of reward. However, the frequency of 'Double clucks' and 511 'Single clucks' did not vary between treatments (P>0.01), which indicates that shorter calls 512 do not encode information about hens' internal states (see Appendix Tables A9 and A10). The 513

acoustic parameters of 'Whines' and 'Gakel calls' did not vary between treatments (*P*>0.01;
see Appendix Tables A7 and A8).

516

517 **Discussion**

The results of this study show that hens produce 'Food calls', 'Fast clucks', 'Double clucks' and 'Single clucks' when anticipating rewards. Importantly, hens produced 'Food calls' when anticipating a dustbathing substrate, which contradicts the suggestion that 'Food calls' made by chickens are functionally referential (Clay et al., 2012; Evans & Evans, 1999). The peak frequency of 'Food calls' made in anticipation of the dustbathing substrate was 45–75 Hz lower than those made in anticipation of food rewards, potentially reflecting differences in arousal within the same behavioural state.

525

We found that all four call types produced by hens in anticipation of rewards ('Food calls', 526 'Fast clucks', 'Double clucks' and 'Single clucks') are structurally similar calls, with a 527 528 fundamental frequency of around 400 Hz and a distinct first harmonic. Calls were mainly differentiated by the number of syllables, with 'Food calls' being a series of 3–15 syllables, 529 produced at a slightly higher peak frequency than 'Fast clucks'. During the neutral treatments 530 (i.e. no rewards), hens produced very different signals. These were long, high-frequency, 531 tonal 'Whines' and long, distinctive, harsh-sounding 'Gakel calls'. 'Whines' produced during 532 our experiment were very similar to the 'whines of frustration' given by red junglefowl, G. 533 gallus, in contexts when food is shown or offered, but subsequently not released to the birds 534 (Collias, 1987). Similarly, 'Gakel calls' are given by hens in frustrative nonrewarding 535 contexts (Garland et al., 2015; Zimmerman & Koene, 1998; Zimmerman, Koene, & van 536 Hooff, 2000). Hens also produced 'Mixed' calls, which did not have a regular or specific 537 structure. The anticipatory period in our experiment may have induced conflicting arousal 538 states; a drive to acquire the reward may have existed simultaneously with frustration induced 539

by not having instant access to the reward. The grading of calls, where one signal merges into 540 another, occurs in red junglefowl and domestic chickens, particularly under intermediate 541 stimulus conditions (Collias, 1987) or in situations of varying intensity (Konishi, 1963). 542 543 Therefore, it appears that hens' call types indicate the motivational state of the signaller; short, sharp 'clucks' and 'Food calls' tend to be produced in anticipation of rewards 544 (indicating possible excitement) and longer, higher frequency 'Whines' and 'Gakel calls' in 545 frustrative non-rewarding contexts. The vocalizations of other animals have been linked to 546 levels of arousal, and in some cases, valence (Briefer, 2012). Therefore, these findings 547 present an opportunity to explore how hens' vocal communication could be used as markers 548 549 of their welfare in on-farm assessments.

550

An important finding of this study was that 'Food calls' were not solely produced in the 551 context of food, but also given in anticipation of both food and nonfood rewards. This result 552 contradicts claims that they are functionally referential (Clay et al., 2012; Evans & Evans, 553 554 1999). Other studies have indicated that food calling can occur in the absence of food, and either may be associated with stimuli that, in the past, reliably predicted the presence of food 555 (Moffatt & Hogan, 1992; Wauters & Richard-Yris, 2002) or may be separation calls (Hughes 556 et al., 1982; Konishi, 1963). Certainly, there is evidence to suggest that the majority of 'Food 557 calls' made by other species are not food-specific, but rather food-associated, as they do not 558 fulfil the strict criteria for functional reference with regard to either production or perception 559 (for a review, see Clay et al., 2012). In these cases, 'Food calls' may function more as social 560 recruitment calls, to enhance defence, reproductive or social status, or alternatively as a 561 562 method to reduce competition by announcing resource ownership (Clay et al., 2012). Male chicken 'Food calls' appear to be used to attract females by inducing orientation towards the 563 sound source (Evans & Evans, 1999). However, 'Food calls' made by females may function 564 to recruit others, either to share a resource or to ensure extra security (vigilance) when 565

engaged in foraging, feeding or dustbathing (Clay et al., 2012; Townsend, Zöttl, & Manser,
2011). Hence, information appears to be contained within the overall structure of hens' calls
(with hens producing short sharp cluck sounds when the signaller has located a resource).
However, there is no evidence in the literature to suggest that these calls differ between
various resources.

571

Many species of animal vary the structure of specific call types depending on their 572 motivational state. For example, the call structure of specific 'food calls' in great ape species 573 can vary according to the perceived quality of the food item (e.g. bonobos, *Pan paniscus*, 574 Clay & Zuberbühler, 2009; chimpanzees, Kalan, Mundry, & Boesch, 2015; Slocombe & 575 Zuberbühler, 2006). In this study, we found that hens produced 'Food calls' more often, and 576 at lower peak frequencies, when anticipating the dustbathing substrate than the food rewards. 577 'Fast clucks' were also more prevalent in anticipation of the dustbathing substrate, and 578 produced at lower frequencies in anticipation of both the dustbath and mealworm rewards 579 580 than the neutral events. This suggests that, in hens, motivational information is contained within the acoustic structure of different call types, as well as within the call type itself. If we 581 group together 'Food calls' and 'Fast clucks' as call types made in anticipation of rewards, 582 this decrease in frequency may indicate further information relating to motivation, such as the 583 signaller's level of arousal. Behavioural data (McGrath et al., 2016) suggest a hierarchy of 584 arousal state, with increased goal-directed behaviour (pushing and pecking against the door), 585 higher activity levels and more approach behaviours demonstrated in anticipation of a 586 dustbathing substrate than the other rewards, and in anticipation of all rewards than neutral 587 588 treatments. Interestingly, in animals such as baboons, *Papio hamadrayas ursinus* (Rendall, 2003), pigs, Sus scrofa (Puppe, Schön, Tuchscherer, & Manteuffel, 2005) and domestic cats, 589 Felis catus (Yeon et al., 2011), the acoustic frequency of calls increases in parallel with 590 heightened arousal, and this pattern is in accordance with Morton's motivation structural rules 591

(Morton, 1977). However, when approached by humans, tame silver foxes, *Vulpes vulpes*, produce calls at lower peak frequencies (based on the fundamental frequency) than aggressive (untamed) silver foxes (Gogoleva, Volodin, Volodina, Kharlamova, & Trut, 2010), suggesting that a lower peak frequency reflects a positively valenced high arousal state rather than a lowarousal feeling of contentment. Therefore, if we assume that lower acoustic frequency calls reflect a reward-related high arousal state in hens, then call frequency, as well as the call types identified in this study, could be used as a marker of welfare.

599

One potential issue with our study was that the dustbathing substrate could have been viewed 600 601 by hens as an opportunity to forage. Hens were deprived of food for 1.5 h to ensure that they were motivated to eat once they had access to food. As hens dustbathe every 2 days on 602 average (Vestergaard, 1982), we deprived the hens of dustbathing substrate for a longer (24 h) 603 period. Chickens tend not to dustbathe during the morning hours, and the fact that some hens 604 dustbathed immediately after accessing the reward chamber suggests that they were highly 605 606 motivated to do so (testing had ended by 1230 hours on each day). Based on the behavioural data, it seems that they were motivated to use the dustbathing substrate for exactly this 607 purpose rather than for foraging, although the two potential opportunities are not mutually 608 609 exclusive. A dustbathing substrate can be a multifaceted resource for hens, including as a potential nest site in a caged environment (Smith, Appleby, & Hughes, 1993), and scratching, 610 foraging and pecking in litter are not only precursors to dustbathing, but also behaviours that 611 hens are motivated to perform in their own right (Olsson & Keeling, 2005). None of our hens 612 performed nesting behaviour during the trials in our experiment, and based on published 613 614 behavioural results (McGrath et al., 2016) we suggest that anticipation of a dustbathing substrate induced high arousal in hens, indicating they were motivated to dustbathe. 615

Another limitation of this study is that we cannot definitively attribute acoustic variation to 617 any particular valence or emotional state. Indeed, Briefer (2012) emphasized that vocal 618 correlates of positive valence are lacking in the literature, often due to a lack of opportunity to 619 620 record vocalizations in positive situations. Detecting valence is possible if studies use situations of contrasting valence but similar arousal, and studies on dogs (Taylor, Reby, & 621 McComb, 2009) squirrel monkeys, Saimiri sciurius (Fichtel, Hammerschmidt, & Jürgens, 622 2001) and goats, Capra hircus (Briefer, Tettamanti, & McElligott, 2015) have successfully 623 separated the effects of arousal and valence on vocal parameters. Measuring acoustic 624 parameters such as formants, frequency range and frequency modulation (Briefer, 2012; 625 Briefer et al., 2015) in conjunction with behavioural and physiological measurements may 626 provide the most convincing evidence for valence. For example, different ultrasonic call types 627 made by rats have been linked to activation of positive or negative internal states in both 628 signallers and receivers (Brudzynski, 2007; Burman, Ilyat, Jones, & Mendl, 2007). As our 629 study used items that are known to be rewarding to chickens, we can hypothesize that shifts to 630 a lower peak frequency within 'Food calls' and 'Fast clucks' in anticipation of rewards 631 reflects an increase in positively valenced arousal. The function of this affective response, as 632 suggested by Mendl et al. (2010), could be to guide animals towards acquisition of fitness-633 enhancing rewards. Hence, future investigations could focus on how conspecifics perceive 634 these vocalizations. Furthermore, there is scope for studies to simultaneously investigate 635 vocal, behavioural and physiological indicators of arousal and valence in chickens. 636

637

638 <H2>Conclusions

This is the first study to find that hens' vocalizations may contain motivational information in call types and call structure. Moreover, the 'Food call' was not given exclusively in response to cues signalling food rewards, suggesting this call was not functionally referential but more likely to be a 'Reward call'. Further work, using other types of reward, will help determine if

643	this is the case. In addition, confirmation of whether males and females produce acoustically
644	different 'Reward calls' would help to assess the function of these calls. Linking
645	physiological and behavioural responses of hens with changes in vocal parameters when
646	anticipating rewards will further identify whether levels of signaller arousal and valence are
647	encoded within the parameters of their calls. Overall, the results of this study suggest that the
648	vocal behaviour of hens can provide an indicator of whether they are in a rewarding
649	environment. These findings can be developed into a tool that could provide a relatively easy
650	method with which to assess the welfare of hens. In a longer study, this tool could monitor
651	welfare outcomes as a result of environmental improvements that may also enhance health
652	and productivity.
653	
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656	
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Groups	Mealworm	Dustbathing substrate	Normal food	Sound-neutral
Cue group 1 (N=3)	Ring	Beep	Buzz	Horn
Cue group 2 (N=3)	Horn	Ring	Beep	Buzz
Cue group 3 (N=3)	Buzz	Horn	Ring	Beep
Cue group 4 (N=3)	Beep	Buzz	Horn	Ring

Table 1: Hen cue groups and combinations of CS and US

Measurement	Description
Call duration	Time (s) from beginning of first syllable to the end of the last syllable in the sequence
Number of syllables	Number of syllables in the call
Average syllable length	Mean length (s) of syllables in a call
Maximum syllable length	Length of the syllable with the longest duration within the call (s)
Minimum Frequency	The lowest frequency bound of the syllable (Hz)
Maximum Frequency	The highest frequency bound of the syllable in (Hz)
Q1 Frequency	The frequency that divides the fundamental frequency into two frequency intervals containing 25% and 75% of the energy (Hz)
Q3 Frequency	The frequency that divides the fundamental frequency into two frequency intervals containing 75% and 25% of the energy (Hz)
Centre Frequency	The frequency that divides the fundamental frequency into two frequency intervals of equal energy (Hz)
Peak Frequency	The frequency at which the maximum power (dB) occurs within the fundamental frequency component (Hz)
Delta Frequency	Difference between the lowest and upper limits of frequency (Hz)
Bandwidth 90%	Difference between the 5% and 95% frequencies (Hz)
Interquartile (IQR) Bandwidth	Difference between the 1st and 3rd Quartile Frequencies (Hz)

Table 2: Description of the measurements used in the quantitative classification of call types

All acoustic measurements are measured on the fundamental frequency component.

933 934 Table A1

Number of call types made by birds

935

Bird	Food call	Fast cluck	Whine	Gakel call	Singing	Mixed	Double cluck	Single cluck	Total no of calls made
1	1	13	0	7	0	19	6	5	51
2	5	4	4	10	2	3	1	6	35
3	1	5	11	1	2	9	8	11	48
4	3	16	0	0	1	11	10	17	58
5	24	3	1	0	0	9	10	11	58
6	4	6	0	10	0	12	5	5	42
7	2	0	16	5	2	4	0	0	29
8	4	5	9	0	0	16	1	1	36
9	0	4	11	1	0	7	12	16	51
10	6	7	3	2	7	9	1	2	37
11	8	6	2	8	1	11	5	6	47
12	2	6	19	9	1	14	1	2	54
Totals	60	75	76	53	16	124	60	82	

939 940 Table A2

941 Mean values of acoustic parameters by call type

Description	<u>Peak Fre</u> (H	equency Iz)	<u>Min Fre</u> (H	quency z)	<u>Max fre</u> (H	<u>quency</u> z <u>)</u>	<u>Cer</u> Freque	<u>ntre</u> ncy (Hz)	<u>Q1 Frec</u> (Н	<u>uency</u> z)	<u>Q3 Frec</u> (<u>Н</u>	<u>quency</u> z)	<u>Delta Fre</u> (H	equency z)	<u>Bandwid</u> (H	<u>1th 90%</u> z <u>)</u>	<u>IQR Ban</u> (H	idwidth z)
	Mean	SD	Mean	SD	Mean	SD	Mean	SD	Mean	SD	Mean	SD	Mean	SD	Mean	SD	Mean	SD
Single cluck	340.6	112.3	261.4	110.2	433.8	121.2	342.7	111.8	324.3	111.2	361.6	112.2	171.8	39.3	97.9	30.4	37.3	13.9
Double cluck	389.2	122.4	277.2	108.0	516.6	151.3	390.8	122.2	372.0	120.4	410.9	124.7	165.6	33.2	96.4	26.7	38.8	14.3
Fast cluck	411.6	132.8	244.8	125.8	609.9	194.1	412.2	133.7	393.0	130.4	432.2	136.7	162.9	41.3	95.8	29.0	39.2	13.4
Food call	457.6	84.2	318.5	98.8	620.3	118.5	459.1	84.3	438.5	83.8	479.8	84.2	173.7	30.1	101.7	26.8	41.4	14.4
Mixed	474.6	137.6	287.4	138.0	694.6	184.5	474.9	136.9	452.9	135.2	497.5	138.8	192.4	56.4	108.6	38.4	44.5	21.4
Whine	621.4	137.7	401.4	145.4	869.4	148.1	617.9	128.3	591.4	126.9	641.8	130.7	274.0	78.1	130.0	54.4	50.4	25.9
Singing	644.6	107.6	337.2	108.5	937.3	127.3	638.1	100.0	589.2	95.9	679.3	106.4	366.8	123.8	218.7	94.1	90.2	41.5
Gakel call	502.6	96.6	313.7	74.7	733.2	179.0	498.7	93.4	473.4	86.4	521.1	100.4	245.6	99.5	120.5	58.6	47.9	34.0

Table A3	
Vocal parameters: Ismeans, 95% confidence limits and significance levels	

Treatment		Dus	sty substra	ate	٦	Mealworm		N	ormal food	1	So	und-neutra	al	Mu	uted-neutra	al			
Vocalization type	Parameter	Lsmean	Lower CL	Upper CL	Lsmean	Lower CL	Upper CL	Lsmean	Lower CL	Upper CL	Lsmean	Lower CL	Upper CL	Lsmean	Lower CL	Upper CL	X ²	X² df	Ρ
Food call	PF	411.85ª	383.86	441.88	487.16 ^b	451.36	525.80	467.55 ^{ab}	431.98	506.04							8.78	2	0.01
	BW 90%	95.24	90.44	100.29	96.44	90.14	103.18	94.70	88.15	101.73							0.04	2	0.98
	SL	0.10	0.09	0.11	0.10	0.09	0.11	0.10	0.09	0.11							0.22	2	0.89
	CD	0.86	0.77	0.96	1.05	0.92	1.21	1.02	0.88	1.18							2.04	2	0.36
Fast cluck	PF	402.66ª	372.47	435.29	403.12 ^a	369.26	440.08	409.19 ^{ab}	374.72	446.84	456.59 ^{ab}	412.44	505.46	548.59 ^b	490.62	613.40	11.30	4	0.02
	BW 90%	90.04 ^a	85.06	95.32	98.86 ^{ab}	92.08	106.14	89.96ª	83.74	96.63	86.71 ^a	79.33	94.78	126.81 ^b	114.72	140.18	13.25	4	0.01
	SL	0.09	0.08	0.10	0.09	0.09	0.10	0.10	0.09	0.11	0.08	0.07	0.09	0.09	0.08	0.10	5.34	4	0.25
	CD	0.76	0.69	0.84	0.66	0.59	0.75	0.73	0.65	0.83	0.80	0.69	0.92	0.87	0.74	1.02	3.51	4	0.48
Whine	PF	574.61	530.93	621.87	628.35	578.13	682.94	630.57	585.21	679.44	644.15	600.14	691.38	642.80	597.85	691.12	4.01	4	0.40
	BW 90%	128.11	112.75	145.55	112.89	97.99	130.04	118.50	105.47	133.14	119.12	107.88	131.53	129.27	116.26	143.74	1.25	4	0.87
	SL	0.42	0.34	0.51	0.36	0.29	0.46	0.47	0.39	0.57	0.47	0.39	0.56	0.56	0.46	0.68	5.70	4	0.22
	CD	1.38	1.19	1.60	1.52	1.31	1.77	1.47	1.29	1.68	1.58	1.38	1.80	1.49	1.30	1.70	1.64	4	0.80
Gakel call	PF				481.99	450.38	515.82	502.85	470.60	537.31	474.48	444.98	505.95	484.51	455.36	515.52	1.41	3	0.70
	BW 90%				96.74	82.85	112.95	113.47	97.66	131.84	106.75	92.64	123.00	113.82	99.45	130.27	1.62	3	0.65
	SL				0.61	0.49	0.76	0.65	0.53	0.81	0.70	0.57	0.86	0.76	0.63	0.93	1.61	3	0.66
	CD				2.42	2.03	2.88	2.52	2.13	2.99	3.12	2.67	3.64	2.62	2.26	3.03	2.36	3	0.50
Double cluck	PF	351.20	320.54	384.80	369.98	333.79	410.10	456.35	406.47	512.35	460.03	402.85	525.33	406.20	350.10	471.29	10.95	4	0.03
	BW 90%	93.46	87.60	99.72	88.15	81.43	95.42	95.82	86.69	105.90	96.32	85.09	109.03	95.29	82.55	109.98	0.97	4	0.91
	SL	0.09	0.09	0.10	0.08	0.07	0.08	0.09	0.08	0.11	0.09	0.08	0.11	0.11	0.09	0.13	7.04	4	0.13
	CD	0.33	0.31	0.36	0.27	0.25	0.30	0.32	0.28	0.36	0.29	0.25	0.33	0.32	0.27	0.38	3.62	4	0.46
Single cluck	PF	331.84	303.89	362.36	310.08	277.22	346.83	350.65	315.74	389.43	289.94	262.10	320.74	343.98	311.70	379.60	4.86	4	0.30
	BW 90%	89.12	84.13	94.40	83.44	76.11	91.47	99.37	91.49	107.92	89.04	82.34	96.28	111.01	102.84	119.84	8.78	4	0.07
	SL	0.09	0.08	0.10	0.09	0.08	0.10	0.09	0.08	0.10	0.07	0.06	0.08	0.08	0.07	0.09	7.92	4	0.09
	CD	0.09	0.08	0.10	0.09	0.08	0.10	0.09	0.08	0.10	0.07	0.06	0.08	0.08	0.07	0.09	7.91	4	0.09

944 PF = Peak Frequency, BW 90% = Bandwidth 90%, SL = syllable length, CD = Call duration. Bold indicates significant *P* values.

- 945 946 947 948
- Table A4 Comparison of the different linear mixed-effects models (LMMs) for behavioural responses

Parameter	Call Type	Fixed variables in model	Comparison	df	AIC	X ²	X ² df	P-value
Peak Frequency	Fast cluck	Treatment		7	-110			
		Treatment + CG	1 vs 2	10	-109	5.62	3	NS
		Treatment + CG+ Day	2 vs 3	14	-106	4.30	4	NS
Bandwidth 90%	Fast cluck	Treatment		7	-113			
		Treatment + CG	1 vs 2	10	-112	4.60	3	NS
		Treatment + CG+ Day	2 vs 3	14	-106	1.56	4	NS
Average syllable length	East cluck	Treatment		7	-110			
Average synable rength	I dot cluck	Treatment + CG	1 vs 2	, 10	-110	6.46	3	NS
		Treatment + CG+ Day	2 vs 3	14	-116	14.28	4	0.01
Duration	Fast cluck	Treatment + CG	1 1 2 2	7 10	-47	1 1 2	2	NS
		Treatment + CG+ Dav	2 vs 3	10	-43	4.12	4	NS
		,						
Peak Frequency	Food call	Treatment		7	-99		_	
		Treatment + CG	1 vs 2	10	-98 02	5.71	3	NS
		freatment + CG+ Day	2 VS 3	14	-95	2.75	4	112
Bandwidth 90%	Food call	Treatment		7	-74			
		Treatment + CG	1 vs 2	10	-69	1.84	3	NS
		Treatment + CG+ Day	2 vs 3	14	-67	5.87	4	NS
Average svilable length	Food call	Treatment		7	-57			
siege synasic icigui		Treatment + CG	1 vs 2	10	-53	1.70	3	NS
		Treatment + CG+ Day	2 vs 3	14	-45	0.56	4	NS
Duration	Food	Treatment		-	20			
Duration	Food call	Treatment + CG	1 vs 7	/ 10	-20 -19	<u>4</u> 47	а	NS
		Treatment + CG+ Day	2 vs 3	14	-11	0.17	4	NS
		,						
Peak Frequency	Gakel call	Treatment		7	-134			
		Treatment + CG	1 vs 2	10	-129	0.63	3	NS
		freatment + CG+ Day	2 VS 3	14	-125	2.33	4	112
Bandwidth 90%	Gakel call	Treatment		7	-30			
		Treatment + CG	1 vs 2	10	-27	3.79	3	NS
		Treatment + CG+ Day	2 vs 3	14	-22	2.78	4	NS
Average syllable length	Gakel call	Treatment		7	4			
werdge syndole rength	Guker cull	Treatment + CG	1 vs 2	10	2	7.98	3	0.05
		Treatment + CG+ Day	2 vs 3	14	5	4.38	4	NS
				_	_			
Duration	Gakel call	Treatment + CG	1 vs 2	7 10	-5 0	1.87	з	NS
		Treatment + CG+ Day	2 vs 3	14	3	4.44	4	NS
Peak Frequency	Whine	Treatment	1	7	-162	0.65	2	NG
		Treatment + CG	1 vs 2	10 14	-157	0.65	3	NS
		Treatment + CO+ Day	2 3 3	14	-157	1.52	4	145
Bandwidth 90%	Whine	Treatment		7	-47			
		Treatment + CG	1 vs 2	10	-42	1.38	3	NS
		Ireatment + CG+ Day	2 vs 3	14	-38	3.69	4	NS
Average syllable length	Whine	Treatment		7	-1			
<u> </u>		Treatment + CG	1 vs 2	10	1	4.24	3	NS
		Treatment + CG+ Day	2 vs 3	14	1	8.05	4	NS
Duration	Whine	Treatment		7	01			
Durdtion	whine	Treatment + CG	1 vs 7	7 10	-81 -80		з	NS
		Treatment + CG+ Day	2 vs 3	14	-86	13.41	4	0.01
		-						
Peak Frequency	Single cluck	Treatment	4	7	-102	0.01	2	
		Ireatment + CG	1 vs 2	10	-96 -05	0.91	3	
		rreatment + CG+ Day	2 42 3	14	-95	0.09	4	142
Bandwidth 90%	Single cluck	Treatment		7	-104			
		Treatment + CG	1 vs 2	10	-102	4.54	3	NS
		Treatment + CG+ Day	2 vs 3	14	-98	3.27	4	NS
Average syllable length	Single cluck	Treatment		7	-67			
werde synaple length	Jungle Cluck	Treatment + CG	1 vs 2	, 10	-58	2.19	3	NS
		Treatment + CG+ Day	2 vs 3	14	-60	9.90	4	0.04
	C	T		_				
Duration	Single cluck	Treatment + CC	1.46.2	7	-62	2 10	2	NC
		Treatment + CG+ Dav	1 vs 2 2 vs 3	10	-58	2.19 9.90	4	0.04
				•				
Peak Frequency	Double cluck	Treatment		7	-87			

950 The table shows Akaike information criterion (AIC	950	The table shows	Akaike information	criterion	(AIC)
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958 Table A5

959 Food call: final model output from LMM

Model	Estimate	SE	t	Pr(> <i>t</i>)
Peak Frequency~Treatment+1 CG:Bird				
<i>df</i> (5),AIC -78.6				
(Intercept) Dustbathing substrate	2.615	0.0288		
Mealworm	0.073	0.0231	3.16	0.00
Normal food	0.055	0.0256	2.16	0.04
Bandwidth 90%~Treatment+1 CG:Bird				
<i>df</i> (5),AIC -66.5				
(Intercept) Dustbathing substrate	1.979	0.0203		
Mealworm	0.005	0.0340	0.16	0.87
Normal food	-0.002	0.0351	-0.07	0.94
Syllable Length~Treatment+1 CG:Bird				
<i>df</i> (5),AIC -53.9				
(Intercept) Dustbathing substrate	-1.008	0.0299		
Mealworm	0.010	0.0367	0.28	0.79
Normal food	-0.011	0.0397	-0.27	0.79
Call Duration~Treatment+1 CG:Bird				
<i>df</i> (5),AIC -17.5				
(Intercept) Dustbathing substrate	-0.066	0.0433		
Mealworm	0.089	0.0633	1.41	0.17
Normal food	0.075	0.0670	1.12	0.27

Table include model degrees of freedom and AICs. 'Estimate' refers to the estimates of the mean of the dependent variable in the model. The first estimate given is the mean of the intercept, with the rest of the estimates in each model being the difference from the intercept. The estimates refer to the values given using a log-transformed dependent variable.

Table A6

Fast cluck: final model output from LMM

Model	Estimate	SE	t	Pr(> <i>t</i>)
Peak Frequency~Treatment+1 CG:Bird				
<i>df</i> (7),AIC -109.7				
(Intercept) Muted neutral	2.739	0.0464		
Dustbathing substrate	-0.134	0.0425	-3.16	0.00
Mealworm	-0.134	0.0435	-3.08	0.00
Sound neutral	-0.080	0.0490	-1.63	0.11
Normal food	-0.127	0.0460	-2.77	0.01
Bandwidth 90%~Treatment+1 CG:Bird				
<i>df</i> (7),AIC -113.4				
(Intercept) Muted neutral	2.103	0.0413		
Dustbathing substrate	-0.149	0.0432	-3.44	0.00
Mealworm	-0.108	0.0452	-2.39	0.02
SN	-0.165	0.0505	-3.27	0.00
Normal food	-0.149	0.0471	-3.17	0.00
Syllable				
Length~Treatment+CG+Day+1 CG:Bird				
<i>df</i> (14),AIC -116.2				
(Intercept) Muted neutral	-1.050	0.0506		
Dustbathing substrate	-0.008	0.0395	-0.20	0.84
Mealworm	0.003	0.0410	0.06	0.95
Sound neutral	-0.047	0.0461	-1.02	0.31
Normal food	0.045	0.0433	1.03	0.31
CG2	0.020	0.0460	0.44	0.67
CG3	0.036	0.0513	0.69	0.50
CG4	0.123	0.0437	2.82	0.02
Day2	-0.032	0.0323	-1.00	0.32
Day3	-0.082	0.0314	-2.61	0.01
Day4	-0.092	0.0374	-2.45	0.02
Day5	0.033	0.0351	0.93	0.36
Call Duration~Treatment+1 CG:Bird				
df(7),AIC -47.1				
(Intercept) Muted neutral	-0.059	0.0660		
Dustbathing substrate	-0.062	0.0664	-0.93	0.35
Mealworm	-0.118	0.0688	-1.72	0.09
Sound neutral	-0.040	0.0771	-0.51	0.61
Normal food	-0.075	0.0722	-1.04	0.30
	0.070	0.0722	1.01	0.00

Table include model degrees of freedom and AICs. 'Estimate' refers to the estimates of the mean of the dependent variable in the model. The first estimate given is the mean of the intercept, with the rest of the

estimates in each model being the difference from the intercept. The estimates refer to the values given using a log-transformed dependent variable. 988 989 990

992 Table A7

993 Whine: final model output from LMM

994

Model	Estimate	SE	t	Pr(> <i>t</i>)
Peak Frequency~Treatment+1 CG:Bird				
<i>df</i> (7),AIC -161.93				
(Intercept) Muted neutral	2.808	0.0295		
Dustbathing substrate	-0.049	0.0265	-1.84	0.07
Mealworm	-0.010	0.0284	-0.35	0.73
Sound neutral	0.001	0.0221	0.04	0.97
Normal food	-0.008	0.0245	-0.34	0.73
Bandwidth 90%~Treatment+1 CG:Bird				
df(7),AIC -47.0				
(Intercept) Muted neutral	2.112	0.0424		
Dustbathing substrate	-0.004	0.0591	-0.07	0.95
Mealworm	-0.059	0.0646	-0.91	0.37
Sound neutral	-0.036	0.0500	-0.71	0.48
Normal food	-0.038	0.0555	-0.68	0.50
Svllable Length~Treatment+CG+1 CG:Bird				
df(7).AIC -1.2				
(Intercept) Muted neutral	-0.252	0.0765		
Dustbathing substrate	-0.129	0.0773	-1.67	0.10
Mealworm	-0.185	0.0831	-2.23	0.03
Sound neutral	-0.077	0.0646	-1.19	0.24
Normal food	-0.074	0.0716	-1.03	0.31
Call Duration~Treatment+CG+Day+1 CG:Bird				
df(14).AIC -85.7				
(Intercept) Muted neutral	0.270	0.0781		
Dustbathing substrate	-0.032	0.0413	-0.78	0.44
Mealworm	0.010	0.0449	0.22	0.83
Sound neutral	0.026	0.0363	0.72	0.47
Normal food	-0.004	0.0392	-0.10	0.92
CG2	-0.156	0.1542	-1.01	0.32
CG3	-0.203	0.0838	-2.43	0.04
CG4	-0.209	0.0881	-2.37	0.04
Day2	0.086	0.0437	1.96	0.05
Day3	0.065	0.0492	1.33	0.19
Day4	-0.021	0.0372	-0.57	0.57
Day5	0.089	0.0412	2.15	0.03

995

Table include model degrees of freedom and AICs. 'Estimate' refers to the estimates of the mean of the

997 dependent variable in the model. The first estimate given is the mean of the intercept, with the rest of the 998 estimates in each model being the difference from the intercept. The estimates refer to the values given

999 using a log-transformed dependent variable.

1003 Table A8

1004 Gakel call: final model output from LMM

Model	Estimate	SE	t	Pr(> <i>t</i>)
Peak Frequency~Treatment+1 CG:Bird				
<i>df</i> (7),AIC -133.9				
(Intercept) Muted neutral	2.685	0.0253		
Mealworm	-0.002	0.0215	-0.11	0.92
Sound neutral	-0.009	0.0190	-0.48	0.63
Normal food	0.016	0.0213	0.76	0.45
Bandwidth 90% ~Treatment+1 CG:Bird				
<i>df</i> (7),AIC -29.6				
(Intercept) Muted neutral	2.056	0.0547		
Mealworm	-0.071	0.0604	-1.17	0.25
Sound neutral	-0.028	0.0534	-0.52	0.60
Normal food	-0.001	0.0594	-0.02	0.98
Syllable Length~Treatment+1 CG:Bird				
<i>df</i> (10),AIC 1.8				
(Intercept) Muted neutral	-0.279	0.0921		
Mealworm	-0.096	0.0793	-1.21	0.23
Sound neutral	-0.037	0.0701	-0.52	0.60
Normal food	-0.066	0.0774	-0.85	0.40
CG2	0.044	0.1511	0.29	0.78
CG3	0.513	0.1460	3.51	0.00
CG4	0.090	0.1166	0.77	0.46
Call Duration~Treatment+1 CG:Bird				
<i>df</i> (7),AIC -4.6				
(Intercept) Muted neutral	0.418	0.0590		
Mealworm	-0.034	0.0799	-0.43	0.67
Sound neutral	0.076	0.0707	1.08	0.29
Normal food	-0.016	0.0776	-0.21	0.84

Table include model degrees of freedom and AICs. 'Estimate' refers to the estimates of the mean of the dependent variable in the model. The first estimate given is the mean of the intercept, with the rest of the estimates in each model being the difference from the intercept. The estimates refer to the values given using a log-transformed dependent variable.

1021 1022 Table A9 Single cluck: final model output from LMM

Model	Estimate	SE	t	Pr(> <i>t</i>)
df(/),AIC -101.5	2 5 2 7	0.0406		
(Intercept) Muted neutral	2.537	0.0406	0.44	0.66
	-0.016	0.0358	-0.44	0.66
	-0.045	0.0475	-0.95	0.35
Sound neutral	-0.074	0.0417	-1.78	0.08
Normal lood	0.008	0.0450	0.185	0.85
Bandwidth 90% ~Treatment+1 CG:Bird				
<i>df</i> (7),AIC -103.8				
(Intercept) Muted neutral	2.045	0.0309		
Dustbathing substrate	-0.095	0.0374	-2.55	0.01
Mealworm	-0.124	0.0482	-2.57	0.01
Sound neutral	-0.096	0.0431	-2.22	0.03
Normal food	-0.048	0.0448	-1.074	0.29
Svilable				
Length~Treatment+CG+Dav+1 CG:Bird				
df(14).AIC -59.7				
(Intercept) Muted neutral	-1.167	0.0623		
Dustbathing substrate	0.053	0.0468	1.13	0.26
Mealworm	0.046	0.0597	0.77	0.44
Sound neutral	-0.073	0.0531	-1.38	0.17
Normal food	0.045	0.0556	0.806	0.42
CG2	0.011	0.0541	0.194	0.85
CG3	-0.009	0.0683	-0.14	0.89
CG4	0.043	0.0666	0.65	0.53
Day2	-0.003	0.0530	-0.06	0.95
Day3	0.175	0.0690	2.531	0.01
Day4	0.025	0.0452	0.543	0.59
Day5	0.104	0.0506	2.06	0.04
Call Duration~Treatment+1 CG·Bird				
df(10) AIC				
(Intercent) Muted neutral	-1 167	0 0623		
Dustbathing substrate	0 023	0.0023	1 1 7	0.26
Mealworm	0.055	0.0400 0 0507	1.12 0.77	0.20
Sound neutral	0.040 _0 072	0.0397	0.77 _1 22	0.44 0 17
Normal food		0.0551	-1.20 U 8∪⊏	0.17
(G2	0.043	0.0550		0.42 0.95
CG3	-0 000	0.0241	_0.195	0.00
CG4	-0.009	0.0005	-0.14	0.09
	0.043		0.00	0.52
Dayz	-0.003	0.0531	-0.06	0.95

Day3	0.175	0.0691	2.53	0.01
Day4	0.025	0.0453	0.543	0.59
Day5	0.104	0.0506	2.05	0.04

1025 Table include model degrees of freedom and AICs. 'Estimate' refers to the estimates of the mean of the 1026 dependent variable in the model. The first estimate given is the mean of the intercept, with the rest of the

1026 estimates in each model being the difference from the intercept. The estimates refer to the values given1028 using a log-transformed dependent variable.

1035 Table A10

1036 Double cluck: final model output from LMM

1037

Model	Estimate	SE	t	Pr(> t)
Peak Frequency~Treatment+1 CG:Bird				
<i>df</i> (7),AIC -82.4				
(Intercept) Muted neutral	2.609	0.0608		
Dustbathing substrate	-0.063	0.0562	-1.12	0.27
Mealworm	-0.041	0.0544	-0.75	0.46
Sound neutral	0.054	0.0650	0.83	0.41
Normal food	0.051	0.0636	0.795	0.43
Bandwidth 90% ~Treatment+1 CG:Bird				
<i>df</i> (7),AIC -80.1				
(Intercept) Muted neutral	1.979	0.0572		
Dustbathing substrate	-0.008	0.0603	-0.14	0.89
Mealworm	-0.034	0.0609	-0.56	0.58
Sound neutral	0.005	0.0727	0.06	0.95
Normal food	0.002	0.0674	0.036	0.97
Syllable				
Length~Treatment+CG+Day+1 CG:Bird				
<i>df</i> (14),AIC -58.2				
(Intercept) Muted neutral	-0.990	0.0666		
Dustbathing substrate	-0.075	0.0695	-1.08	0.28
Mealworm	-0.162	0.0721	-2.25	0.03
Sound neutral	-0.088	0.0865	-1.02	0.31
Normal food	-0.070	0.0767	-0.916	0.36
CG2	-0.039	0.0425	-0.927	0.36
CG3	0.136	0.0491	2.77	0.01
CG4	0.041	0.0592	0.69	0.49
Call Duration~Treatment+1 CG:Bird				
<i>df</i> (10),AIC -61.0				
(Intercept) Muted neutral	-0.493	0.0676		
Dustbathing substrate	0.016	0.0707	0.22	0.83
Mealworm	-0.069	0.0707	-0.98	0.33
Sound neutral	-0.051	0.0846	-0.60	0.55
Normal food	-0.007	0.0792	-0.085	0.93

1038

Table include model degrees of freedom and AICs. 'Estimate' refers to the estimates of the mean of thedependent variable in the model. The first estimate given is the mean of the intercept, with the rest of the

estimates in each model being the difference from the intercept. The estimates refer to the values givenusing a log-transformed dependent variable.

1045 Table A11

1046 Random Forest confusion matrix and classification error for each call type

							·)[··		
Call type	Double cluck	Fast cluck	Food call	Gakel call	Mixed	Singing	Single cluck	Whine	Classification error
Double cluck	59	0	0	0	0	0	1	0	0.017
Fast cluck	0	49	13	0	13	0	0	0	0.347
Food call	0	17	33	0	10	0	0	0	0.450
Gakel call	0	0	0	38	6	1	0	8	0.283
Mixed	0	19	11	3	74	3	0	14	0.403
Singing	0	0	0	0	7	6	0	3	0.625
Single cluck	0	0	0	0	0	0	82	0	0.000
Whine	0	0	0	6	10	0	0	60	0.211

The OOB error rate was 26.56%. The first column lists call types from our subjective classification by
name, and the following columns show the number of each call type the initial call types were classified
into by Random Forest. The final column shows the classification error rate for each call type.

1055 1056 Table A12Gini index showing importance of predictor variables in the Random Forest analysis.

Variable	Mean decrease in Gini				
Call Duration	87.52				
Max Syllable Length	61.59				
Number of Syllables	72.42				
Average syllable length	54.83				
Peak Frequency	19.69				
Minimum Frequency	22.31				
Maximum Frequency	27.06				
Centre Frequency	20.22				
Q1 Frequency	19.57				
Q3 Frequency	20.21				
Frequency Range	25.26				
Bandwidth	17.56				
IQR Bandwidth	16.72				

1059

- 1062 List of Figures.
- 1063 Fig.1. The experimental pen. L shows where the lamp was positioned.
- 1064 Fig. 2. Number of calls made and number of calls analysed by treatment.
- 1065 Fig.3. Classification and regression tree of vocalizations made by chickens in anticipation of rewards. The
- tree shows the variables used at each split, with the criteria (<,>,=). Terminal nodes indicate the prediction
- 1067 of call type for that partition and the number of correctly classified call types out of the total in that
- subgroup. ('Food C' = Food call, 'Fast C' = Fast cluck, 'Double C' = Double cluck, 'Single C' = Single
- 1069 cluck, 'Duration' = Call Duration, 'Timesyll' = Average syllable length, 'Numsyll' = Number of syllables,
- 1070 'Peak' = Peak Frequency, 'Max' = Maximum Frequency)
- 1071
- 1072 Fig. 4. Model predictions of call rate per hen for eight different call types in anticipation of five different
- treatments. (a) Food calls and clucks. (b) Other calls. SN: sound-neutral; MN: muted-neutral. Means are
- shown with lower and upper confidence limits. P<0.05; P<0.01; P<0.01; P<0.01; P<0.001: call rates that were
- 1075 significantly higher than MN.
- 1076
- 1077 Fig. 5. Variation in the peak frequency of 'Food calls' by reward. Central dot indicates mean values and1078 horizontal lines show SEs.
- 1079
- 1080 Fig. 6 Spectrograms of (a) 'Food call', (b) 'Fast cluck', (c) 'Single cluck' and (d) 'Double cluck'. A key
- 1081 within the 'Food call' spectrogram indicates the syllable length and call duration.
- 1082
- 1083 Fig. 7. Example spectrograms of (a) 'Whine' and (b) 'Gakel calls'.