

1 **Combining bioacoustics and occupancy modelling for improved monitoring of**  
2 **rare breeding bird populations**

3 **AUTHOR DETAILS**

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13 **DECLARATION OF AUTHORSHIP**

14 CA conceived the ideas, designed methodology; collected and analysed the data. CA led the writing of the  
15 manuscript, with MG contributing to occupancy modelling methods and development of the text. Both  
16 authors contributed critically to the drafts and gave final approval for publication.

17

## 18 HIGHLIGHTS

- 19 • Bioacoustic recording is used to generate occupancy and detectability estimates
- 20 • Rare heathland breeding birds varied in their occupancy between 0.68 and 0.13
- 21 • Detectability varied from 0.74 to 0.20, and was affected by habitat
- 22 • Bioacoustics can be used to provide improved data over traditional survey methods

## 23 ABSTRACT

24 Effective monitoring of rare and declining species is critical to enable their conservation, but can often be  
25 difficult due to detectability or survey constraints. However, developments in acoustic recorders are  
26 enabling an important new approach for improved monitoring that is especially applicable for long-term  
27 studies, and for use in difficult environments or with cryptic species.

28 Bioacoustic data may be effectively analysed within an occupancy modelling framework, as  
29 presence/absence can be determined, and repeated survey events can be accommodated. Hence, both  
30 occupancy and detectability estimates can be produced from large, coherent datasets. However, the most  
31 effective methods for the practical detection and identification of call data are still far from established.  
32 We assessed a novel combination of automated clustering and manual verification to detect and identify  
33 heathland bird vocalizations, covering a period of six days at 44 sampling locations

34 Occupancy ( $\Psi$ ) and detectability ( $p$ ) were modelled for each species, and the best fit models provided  
35 values of: nightjar  $\Psi=0.684$ ,  $p=0.740$ , Dartford warbler  $\Psi=0.449$   $p=0.196$  and woodlark  $\Psi=0.13$   $p=0.996$ .  
36 Including environmental covariates within the occupancy models indicated that tree, wetland and heather  
37 cover were important variables, particularly influencing detectability.

38 The protocol used here allowed robust and consistent survey data to be gathered, with limited fieldwork  
39 resourcing, allowing population estimates to be generated for the target bird species. The combination of

40 bioacoustics and occupancy modelling can provide a valuable new monitoring approach, allowing  
41 population trends to be identified, and the effects of environmental change and site management to be  
42 assessed.

## 43 **KEYWORDS**

44 Acoustic ecology, autonomous recorder, bird survey, heathland, occupancy model.

45

## 46 **1. INTRODUCTION**

### 47 **1.1 Bioacoustics for Biodiversity Monitoring**

48 Biodiversity monitoring is central to nature conservation, allowing species status to be evaluated or  
49 assessments to be made of biological responses to environmental changes (Pereira & Cooper, 2006).

50 Long-term monitoring of designated nature conservation sites is particularly needed to identify population  
51 trends and inform management planning efforts, especially in the context of factors such as climate  
52 change and habitat loss/severance (Noss, 1990; Furnas & Callas, 2015). However, existing monitoring  
53 practices and protocols are often sub-optimal, especially in terms of unbiased spatial coverage, sampling  
54 effort optimization, the statistical use of the data, and the lack of repeated sampling (Schmeller et al.,  
55 2012).

56 We assessed the potential to improve the existing monitoring methods currently used on sites that are  
57 internationally important for their breeding bird populations. The most common methods for monitoring  
58 of bird numbers and distributions are transect or point count surveys by human observers. These have  
59 recognised disadvantages, such as observer bias, the availability of skilled/experienced surveyors  
60 (Brandes, 2008; Celis-Murillo et al., 2009; Rempel et al., 2005; Sedláček et al., 2015), and the infrequent  
61 and short-term nature of survey visits (Shonfield & Bayne, 2017; Zwart et al., 2014). In response to these

62 issues, passive acoustic monitoring is increasingly being used as an alternative monitoring technique. This  
63 method uses automated recording units, which can be deployed in the field for days or weeks at a time to  
64 capture animal sounds. The advantages of this approach include the production of a standardised, long-  
65 duration, permanent dataset and record of species identification, which can be repeatedly analysed and  
66 subject to validation by independent reviewers (Abrahams & Denny, 2018; Celis-Murillo et al., 2009;  
67 Rempel et al., 2005). Automated recorders can be synchronized to occur simultaneously across large  
68 spatial extents, reducing temporal variability in studies (Brandes, 2008; Furnas & Callas, 2015;  
69 MacKenzie & Nichols, 2004), and offering large data volumes at low cost and with little resourcing  
70 requirement (Acevedo & Villanueva-Rivera, 2006; Hill et al., 2018; Holmes et al., 2014; Zwart et al.,  
71 2014). Due to potential benefits such as these, the use of automated recorders has increased significantly  
72 over the last ten years (Shonfield & Bayne, 2017), and some researchers have advocated the use of  
73 automated recorders instead of expert personnel for conducting surveys (Darras et al., 2018; Rempel et  
74 al., 2005; Brandes, 2008; Zwart et al., 2014).

75 There are potential barriers to the widespread uptake of passive acoustic monitoring for bird surveys.  
76 These include the need for specific expertise and the increased time required for post-processing  
77 compared to some traditional surveys (Banner et al., 2018; Knight et al., 2017), together with the costs of  
78 equipment (Beason et al., 2018; Farina et al., 2014; Hill et al., 2018). However, open source or low-cost  
79 recording devices are being produced and post-processing methods are constantly improving – although  
80 automated species identification, including machine-learning approaches, is still in development  
81 (Acevedo et al., 2009; Salamon et al., 2016). For fieldwork, a practical disadvantage is the fact that  
82 acoustic monitoring does not allow the collection of visual clues which can sometimes be vital for the  
83 identification of cryptic/quiet species, or for assessing abundance (Klingbeil & Willig, 2015; Sedláček et  
84 al., 2015). In some cases, the use of audio recording units has resulted in detection of fewer species and  
85 detection at shorter distances than human observers (Holmes et al., 2014; Yip et al., 2017), but the  
86 potential for longer term data capture with recording units means that this constraint can normally be

87 addressed by longer deployment times (Darras et al., 2018; Sedláček et al., 2015; Shonfield & Bayne,  
88 2017; Zwart et al., 2014). However, microphone performance and maintenance needs to be considered as  
89 part of the planning of fieldwork campaigns (Turgeon et al., 2017; Yip et al., 2017).

## 90 **1.2 Occupancy Models**

91 Alongside the technological advances in bioacoustics, there has been a dramatic recent increase in the  
92 development and application of occupancy models that explicitly incorporate species detectability (Furnas  
93 & McGrann, 2018; MacKenzie & Nichols, 2004; MacKenzie et al., 2002; MacKenzie et al., 2006). The  
94 presence/absence of a species in a sample can be used to calculate occupancy ( $\Psi$ ) - the proportion of an  
95 area, or number of sites, occupied by a species. The frequency with which a species is repeatedly recorded  
96 at each sampling site can also be used to assess detectability ( $p$ ), to allow for the estimation of, and  
97 correction for, imperfect detection (Banner et al., 2018; MacKenzie et al., 2002; MacKenzie et al., 2006).  
98 The ability to factor these two parameters into assessments allows improved estimates of populations and  
99 greater understanding of ecological patterns such as species/habitat relationships (MacKenzie et al.,  
100 2006).

101 Despite the clear potential and utility of combining bioacoustic techniques and occupancy models, only a  
102 few studies have united these methodological developments to model the population status of a range of  
103 vocal species (Yates & Muzika 2006; Furnas & Callas 2015; Kalan et al. 2015; Campos-Cerqueira &  
104 Aide 2016; Stiffler et al. 2018; Wood et al., 2019). This study, therefore, provides an important additional  
105 case-study in new geographical, habitat and spatiotemporal contexts. Furthermore, it also addresses one of  
106 the most critical questions in this area of study - how to most effectively extract useful information from  
107 acoustic recorders to feed into the occupancy models and allow population estimates to be generated.

108 Although fine-grained data can be gained from acoustic recorders, a significant benefit of the occupancy  
109 modelling approach in field studies is that it relies only on presence/absence data, rather than metrics of  
110 abundance such as counts of individuals (MacKenzie et al., 2006). This is normally much easier to

111 determine, requiring less interpretation in the field/lab, and counteracting the potential for inter-observer  
112 or inter-survey error (MacKenzie et al., 2006). Although some information is perhaps lost by this  
113 approach, data accuracy may be gained as, for rare species, it can be very difficult to correctly estimate  
114 abundance during surveys, whereas estimation of occupancy may still be possible with a high level of  
115 confidence (Campos-Cerqueira & Aide, 2016; Mackenzie & Royle, 2005). Finally, occupancy and  
116 abundance will be linked in most populations, and at small spatial scales and with territorial species,  
117 occupancy may be regarded as equivalent to population size and can be used for investigating population  
118 dynamics or spatial variation (MacKenzie et al., 2006; Royle & Nichols, 2003; Furnas & Callas, 2015;  
119 Campos-Cerqueira & Aide, 2016; Wood et al., 2019).

### 120 **1.3 Heathland Bird Monitoring**

121 Our study was conducted on European nightjar *Caprimulgus europaeus*, woodlark *Lullula arborea* and  
122 Dartford warbler *Sylvia undata*. These three birds are specialists of lowland heathland habitats, and are  
123 rare and declining species considered to be of international conservation importance (Clark & Eyre,  
124 2012). Despite significant legal and policy protection, however, their breeding site habitats are threatened  
125 by air pollution, urban development, inappropriate management and recreational disturbance (Fagúndez,  
126 2013; Mallord et al., 2007).

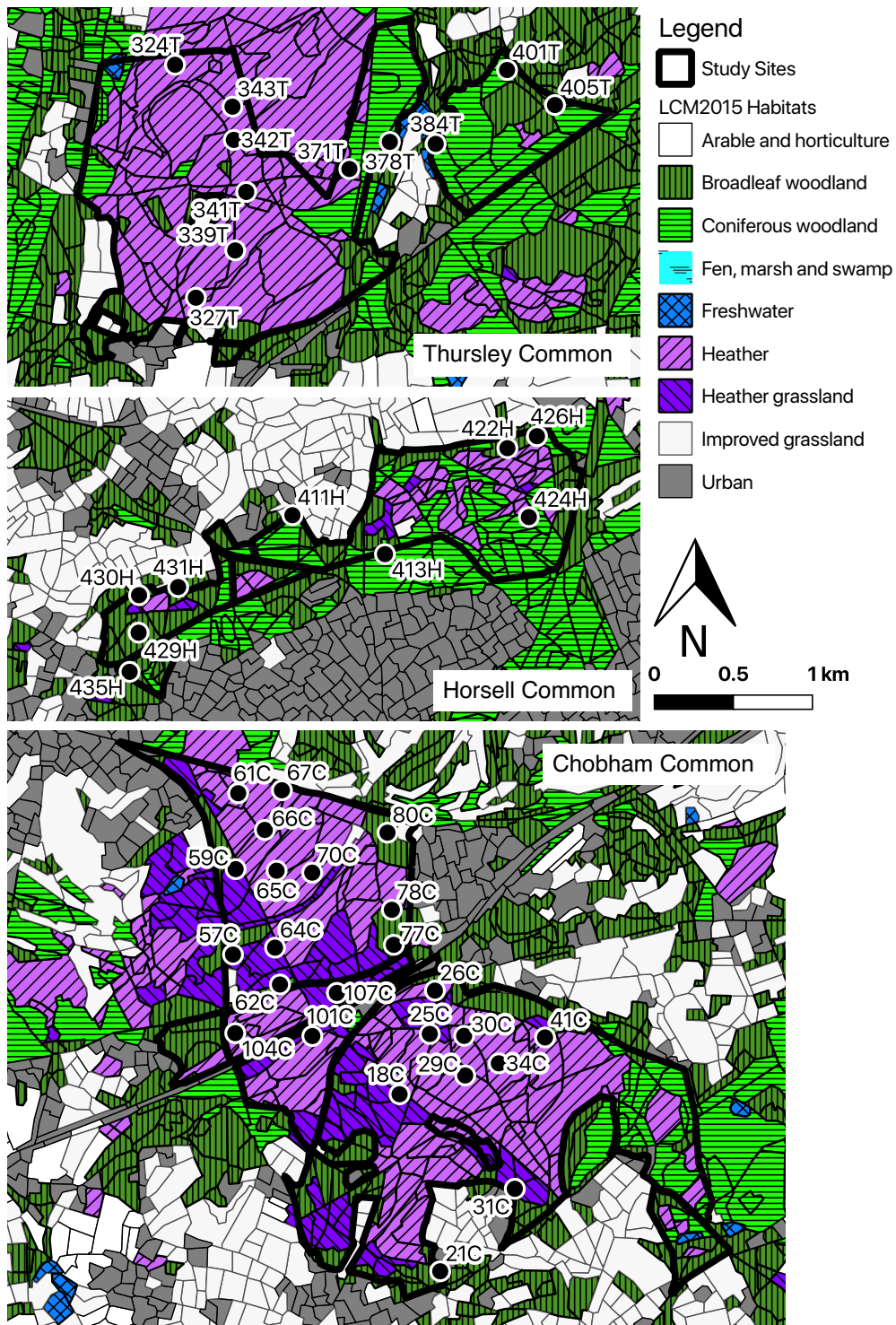
127 Monitoring a variety of bird species, with differing behaviours, over extensive heathland sites, presents  
128 significant challenges for conservation managers. In particular, a number of different surveyors are  
129 inevitably involved in the surveys used for monitoring the target species. Inter-observer differences are  
130 therefore likely to produce variations in data, particularly with nocturnal nightjar surveys, where it is hard  
131 to differentiate individuals and accurately map territories (Liley & Fearnley, 2014). Automated recorders,  
132 used by themselves or in conjunction with existing methods, have great potential to reduce bias and  
133 variability in survey results and account for the effects of detectability between sites and surveys, to  
134 produce more reliable and consistent population estimates.

135 Our goal in this study is to establish effective methods for combining bioacoustic techniques and  
136 occupancy models in the monitoring of rare breeding bird populations. We capture an acoustic dataset and  
137 demonstrate how to efficiently process recordings to detect and identify species vocalizations within this,  
138 using a novel clustering technique. We then analyse the acoustic data to estimate occupancy and  
139 detectability for the three target species, using single-species, single-season occupancy models, and  
140 combine this with environmental covariates, to determine the effects of habitat on model outputs. This  
141 provides useful occupancy and detectability estimates for the target species, highlighting the potential for  
142 bioacoustic methods to be used as an alternative or complement to current monitoring practices, with  
143 benefits in terms of consistent, verifiable and permanent field data.

## 144 **2. MATERIALS AND METHODS**

### 145 **2.1 Study Area**

146 We conducted the study on parts of the Thames Basin Heaths SPA and the Wealden Heaths SPA. These  
147 are two large, internationally important, nature conservation sites in southern England, made up of 18  
148 heathland sites of varying size and character. These sites comprise a mix of dry and wet heath vegetation,  
149 with mire, bog, waterbodies, permanent grassland, scrub and blocks of woodland (Figure 1). Together,  
150 they cover a total of 12,199 ha, of which 5,702 ha is classified as lowland heath (Clark & Eyre, 2012).  
151 Within this overall context, we gathered data at three heathland sites to which access could be readily  
152 gained: Chobham Common, Horsell Common and Thursley Common, which together cover an area of  
153 992 ha.



154  
155

Figure 1. Land Cover Map 2015 habitat data and acoustic sampling site locations.

156



## 157 2.2 Acoustic Monitoring

158 We used Wildlife Acoustics SongMeter SM2 recorders, equipped with a single mono omnidirectional  
159 microphone to record audio data (see Supplementary Information: Appendix 1). These automated  
160 recording units were programmed to record a 1 minute audio sample every ten minutes (i.e. one minute  
161 on, nine minutes off), from two hours before sunrise, until three hours after, and then from one hour  
162 before sunset until two hours after. Daily sampling therefore took place within a 5 hour period at dawn,  
163 and 3 hours at dusk. The units were deployed at a single sample site for a period of six days during May-  
164 June 2018, so that each site had 288 minutes of recording. The audio samples were all recorded as .wav  
165 files onto an SD card, at 48kHz sampling rate and 16-bit depth (Abrahams, 2018). All microphones were  
166 calibrated to ensure comparable sensitivity and performance before deployment (Turgeon et al., 2017;  
167 Yip et al. 2017).

168 Sample locations were defined across the study area by using GIS to place a regular 250 m point grid  
169 across the three heathland sites. It was considered that this would be a sufficient distance for recordings to  
170 be independent of each other, and relevant to the territory sizes of the species being studied. From the 166  
171 possible grid points, 48 were randomly selected, stratified to the relative area of each heathland site, to  
172 provide 9 sampling sites at Horsell Common, 15 at Thursley Common, and 24 at Chobham Common. As  
173 16 recorders were available for the study, the 48 sampling sites were divided into three sessions of field  
174 recording: 26-31 May, 5-10 June, 16-21 June. The sites were randomly assigned to one of the three  
175 survey sessions, so that 3 sites at Horsell Common, 5 at Thursley Common, and 8 at Chobham Common  
176 would be sampled at each session. Despite differences in date, all site samples were treated equally as  
177 individual samples within a single season. A closure assumption was therefore made that bird  
178 distribution, population size and density did not change over the course of the three survey sessions.

179 All sites were given an identification code consisting of a number and site suffix of H, T or C (Figure 1).  
180 Field placements matched the GIS locations as closely as features on the ground would allow. During the  
181 deployments, one recorder failed to record evening sessions repeatedly (at three sampling sites), and

182 another suffered battery failure on one occasion. These failures were all at Thursley Common (sites 315T,  
183 319T, 332T, 391T) and the sites were removed from the dataset, leaving 44 sampling locations.

### 184 **2.3 Audio Data**

185 The audio recordings taken from the field were analysed using a semi-automated system to identify target  
186 species vocalizations (termed ‘phrases’) in the recordings. Kaleidoscope Pro 4.3.2 software (Wildlife  
187 Acoustics, 2017) was first employed, using its cluster analysis method with default settings  
188 (<https://www.wildlifeacoustics.com/images/documentation/Kaleidoscope-Pro-5-User-Guide.pdf>). This  
189 process analysed the time and frequency characteristics of the recorded audio files, using Hidden Markov  
190 Models, to search for sounds within a 1500-7000Hz frequency band and of 2-20 seconds duration, with a  
191 maximum inter-syllable gap of 1 second - creating each as an individual new .wav file. The analysis  
192 process grouped similar phrases in the recordings (e.g. the song of a particular bird species) into clusters  
193 based on their sound characteristics. After the automated clustering was complete, the phrases detected by  
194 the software were manually reviewed by listening to playback and by the visual inspection of  
195 spectrograms to classify the presence/absence of the target species in each phrase.

### 196 **2.4 Environmental Data**

197 In order to investigate the influence of habitat on occupancy and detectability at each of the study sites,  
198 we obtained data from a combination of satellite and terrestrial mapping sources. The proportion of  
199 Broadleaf trees, Coniferous trees, Heather and Heather grassland within 100m of each sample site was  
200 calculated from Land Cover Map 2015 (LCM2015) vector data, accessed from the Centre for Ecology  
201 and Hydrology (Rowland et al., 2017). Distance to the nearest road was calculated based on Ordnance  
202 Survey OpenMap-Local vector data (OS data © Crown copyright and database right 2018). We also used  
203 pre-processed satellite data from Copernicus Pan-European High Resolution Layers (HRL;  
204 <https://land.copernicus.eu/pan-european/high-resolution-layers>) representing Tree Cover Density (TCD),  
205 Water and Wetness (WAW) and Imperviousness (IMD) at a 20m resolution. The Tree Cover Density

206 (forest) HRL provides the level of tree cover in a range from 0-100% for each pixel.. The Water and  
207 Wetness HRL shows the occurrence of water and wet surfaces over the period from 2009 to 2015, on a  
208 scale from (1) permanent water, to (4) temporary wetness. The Imperviousness degree IMD captures the  
209 spatial distribution of artificially sealed (i.e. urbanized/road) areas. We used Zonal Statistics to summarise  
210 these measures for each sampling site, to produce the sum of all pixel values within a 100m radius of the  
211 site. All spatial analyses were performed in QGIS (QGIS Development Team, 2018). Weather was  
212 represented in our environmental variables by ‘derived 24hr sun duration’ from the weather station at  
213 Wisley, Surrey (Ref. src\_id 719/DCNN 5237, WGS84 51.3108, -0.47634), accessed from BADC  
214 (badc.nerc.ac.uk). Other weather variables were unavailable from this source as records for the survey  
215 period were sparse.

## 216 **2.5 Occupancy Models**

217 The occupancy of each of the three target species was modelled separately using a single-species, single-  
218 season modeling approach with observation and habitat covariates (Furnas & Callas, 2015; MacKenzie et  
219 al., 2002; MacKenzie et al., 2006; Stiffler et al., 2018), using established protocols with the ‘Unmarked’  
220 package in R (Fiske & Chandler, 2011; R Core Team, 2013; RStudio Team, 2015). The acoustic data was  
221 summarised to day-level temporal resolution of presence/absence, to produce a detection history at each  
222 sampling site comprising 6 replicate surveys. The naive occupancy for each species was checked and  
223 confirmed to be  $>0.1$ , so that detection histories were not too sparse to fit single-species models. We first  
224 created null models, without covariates, to represent equal probability of detection and/or occupancy  
225 across all survey sites and days. We then developed models including covariates representing the areas of  
226 different habitat types within 100m of the sampling location (from LCM2015 and Copernicus data), and  
227 distance to the nearest road (as shown in Table 2). We anticipated that detection probability might change  
228 over the course of the survey period (Campos-Cerqueira & Aide, 2016; Furnas & McGrann, 2018) due to  
229 seasonal and weather reasons, and used Julian day of survey and 24-hour sun duration to represent this  
230 information. All variables were scaled and centered around zero prior to analysis. The broadleaf and

231 coniferous covariates were excluded as these duplicated the TCDsum habitat type, and the LCM2015 data  
232 were more zero-inflated than the Copernicus data. IMDsum was also rejected as the data were very  
233 sparse. Covariates were applied first to the detection parameter, before the occupancy parameter. Each  
234 model was inspected to check estimates, standard errors and convergence. All models tested are listed in  
235 Table 2.

236 We assessed model fit using Akaike's Information Criterion (AIC), ranking and comparing models based  
237 on AIC relative differences between the top ranked model and each other model ( $\Delta AIC$ ) and AIC  
238 weights. We considered models with  $\Delta AIC < 2$  to be equally supported (Burnham & Anderson, 2002) and  
239 combined these by applying model averaging using the MuMIn package in R (Barton, 2018), to estimate  
240 occupancy and detection for each species. Initially, models without occupancy covariates were fitted to  
241 select the most appropriate covariates for detection. These covariates were then retained for all candidate  
242 models when occupancy covariates were added. The models generated for each species were used to  
243 assess occupancy levels at the study sites, define potential habitat areas and calculate provisional  
244 population estimates.

## 245 **3. RESULTS**

### 246 **3.1 Clustered Audio Segments**

247 Kaleidoscope clustering of the complete audio dataset detected 28,775 phrases as individual .wav files, an  
248 average of 109 phrases per site/day. Each phrase included bird vocalizations and other sounds. With a  
249 mean duration of 6 seconds (range 2-20.9 sec), the clustered phrases comprised 48 hours of audio - 23%  
250 of the total recorded dataset. The phrases were grouped into 55 clusters by the software.

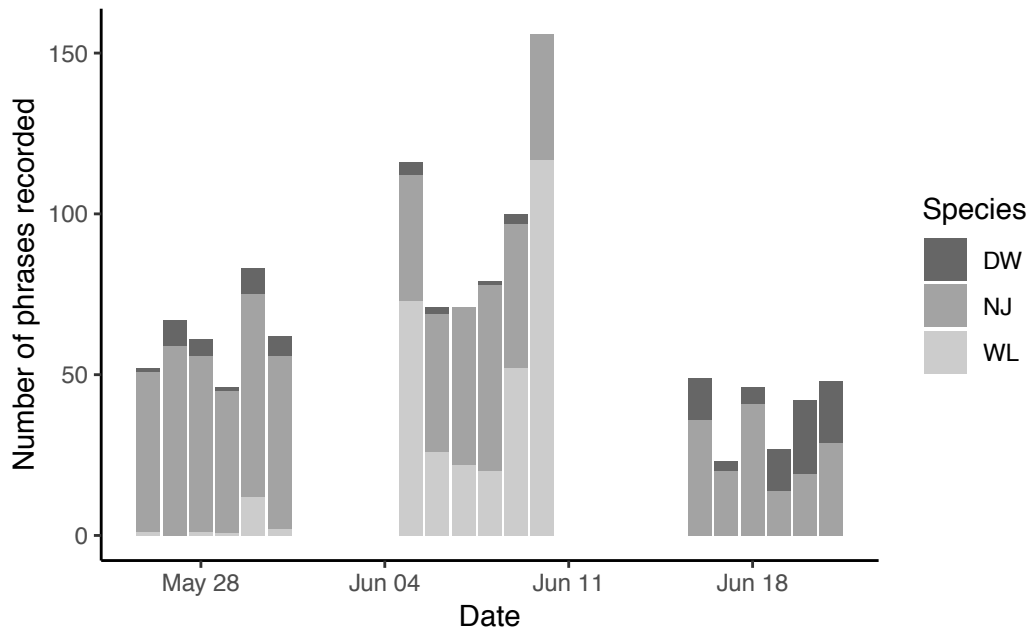
251 Manual review of all the clustered phrases identified the three target species in the dataset, with 757  
252 phrases across 30 sites having vocalizations of nightjar, 327 of woodlark at 7 sites, and 115 of Dartford  
253 warbler at 14 sites. This gave a total of 1,199 phrases recorded for the three target species. Nightjar and

254 Dartford warbler were recorded at all three SPA sites, but woodlark was only recorded at Chobham and  
255 Thursley Commons.

256 **3.2 Patterns in Activity**

257 The total number of phrases recorded per day across all sampling sites varied from 1,974 on 30 May to  
258 1,145 on 17 June. The daily number of phrases was relatively even between recording sessions 1 and 2,  
259 but declined for session 3 in mid-June. This pattern was matched somewhat by the daily numbers of target  
260 species vocalizations (Figure 2). Nightjar and Dartford warbler vocalizations were recorded throughout  
261 all three recording sessions, but woodlark was mostly confined to the early June session only - although  
262 this is likely to be related to presence at the sites being sampled at that time, rather than any reason to do  
263 with seasonal timing.

264 The most vocally active sites were 61C and 70C (north Chobham) for nightjar, 29C and 25C (south  
265 Chobham) for woodlark, and 339T and 343T (central Thursley) for Dartford warbler - see locations at  
266 Figure 1. Significant numbers of calls were not recorded for any species at the Horsell Common sites.



267

268 *Figure 2. Number of target species recorded per day across all sampling sites, for Dartford warbler*  
269 *(DW), nightjar (NJ), and woodlark (WL).*

270

### 271 **3.3 Environmental Parameters**

272 The recorders were placed in habitats that varied from open heath to mature forest (Figure 1). Thursley  
273 Common can be divided into a western part, dominated by Heather, with the eastern part being  
274 Coniferous and Broadleaved woodland. Chobham Common is a mosaic of Heather and Heather  
275 grassland, with Coniferous and Broadleaved woodland around its fringes. This site has a much larger  
276 cover of WAW than the two other sites. Horsell Common is mostly Coniferous and Broadleaved  
277 woodland, with patches of Heather at its eastern end. The means and ranges of the GIS-measured  
278 environmental parameters are listed in Table 1.

279

| Habitat variable           | Mean value | Range   | Units                      |
|----------------------------|------------|---------|----------------------------|
| TCDsum                     | 2570       | 0-6209  | Sum of % per pixel         |
| WAWsum                     | 36.8       | 0-252   | Sum of 1-4 index per pixel |
| Distance to Road (HubDist) | 351        | 29-961  | Metres                     |
| Heather                    | 14459      | 0-31318 | Sum of pixels              |
| Heather grassland          | 4204       | 0-31060 | Sum of pixels              |

280 Table 1. Measured habitat parameters (n=44 sampling sites)

281

### 282 3.4 Occupancy Modelling

283 Naive occupancy was calculated for each species, based on the presence of the species across all 44  
284 sample sites in the study. The naive occupancy values, equal to the proportion of sites with positive  
285 detections, were 0.68 for nightjar, 0.32 for Dartford warbler and 0.16 for woodlark.

286 Models incorporating covariates on the detection and occupancy parameters were generated for each  
287 species (Table 2). Two models for nightjar had equal support ( $\Delta AIC < 2$ ) and so were averaged to produce  
288 covariate estimates. The averaged model included Julian date (JULIAN), Tree Cover Density (TCDsum)  
289 and Water and Wetness (WAWsum) as detectability covariates with no covariates acting on occupancy.  
290 The best fit model for nightjar (NJmdet3), with an AICwt of 53%, indicates an occupancy of 0.684 (SE  
291 0.071) with a detectability of 0.740 (SE 0.035), varying only slightly from the null model ( $\Psi=0.682$ ,  
292  $p=0.733$ ).

293 There were four favoured models for Dartford warbler, including the null model, with TCDsum,  
294 WAWsum, and distance to road (HubDist) featuring on the detectability parameter. Heather grassland  
295 was the only indicator for occupancy. The averaged model for Dartford warbler used only distance to  
296 road as a detectability covariate, with no covariates acting on occupancy. The best-fit model for Dartford  
297 warbler (DWmdet5), with an AICwt of 36%, indicates an occupancy of 0.449 (SE 0.107), with a  
298 detectability of 0.196 (SE 0.053), an increase from the null model occupancy of 0.382 (SE 0.091), but  
299 decrease in detectability from 0.258 (SE 0.057).

300 Woodlark had two favoured models, sharing Julian date, WAWsum, distance to road, Heather and  
301 Heather grassland as detectability covariates, and WAWsum, Heather and Heather grassland for  
302 occupancy covariates. The averaged model for woodlark had five significant covariates, and again, these  
303 were all on the detection parameter. Julian date, WAWsum and Heather were all positively related to  
304 detectability, while distance to road and Heather grassland were negative indicators. For woodlark, the  
305 best-fit model (Wlmocc2), with an AICwt of 59%, indicated an occupancy of 0.13 (SE 0.117), lower

306 than the null model figure of 0.162 (SE 0.056), and a detectability of 0.996 (SE 0.012), which varied  
 307 substantially from the null model detectability of 0.491 (SE 0.081).

| Model                   | Formula   | AIC    | ΔAIC  | AICwt |
|-------------------------|---|--------|-------|-------|
| <b>Nightjar</b>         |   |        |       |       |
| NJmdet3                 | ~JULIAN + TCDsum + WAWsum ~ 1   | 259.62 | 0.00  | 0.528 |
| NJmocc3                 | ~JULIAN + TCDsum + WAWsum ~ TCDsum  | 260.64 | 1.02  | 0.317 |
| NJmocc2                 | ~JULIAN + TCDsum + WAWsum ~ TCDsum + HubDist  | 262.33 | 2.70  | 0.136 |
| NJmocc1                 | ~JULIAN + TCDsum + WAWsum ~ TCDsum + WAWsum + HubDist +<br>Heather + HeatherGrass                           | 267.64 | 8.02  | 0.010 |
| NJm0                    | ~1 ~ 1  | 267.79 | 8.17  | 0.009 |
| <b>Dartford Warbler</b> |   |        |       |       |
| DWmdet5                 | ~TCDsum + HubDist ~ 1   | 157.11 | 0.00  | 0.364 |
| DWmocc3                 | ~HubDist + TCDsum ~ HeatherGrass  | 158.19 | 1.08  | 0.212 |
| DWmdet4                 | ~TCDsum + WAWsum + HubDist ~ 1  | 158.40 | 1.29  | 0.191 |
| DWm0                    | ~1 ~ 1  | 159.00 | 1.89  | 0.142 |
| DWmocc2                 | ~HubDist + TCDsum ~ WAWsum + HeatherGrass   | 160.06 | 2.95  | 0.083 |
| DWmocc1                 | ~HubDist + TCDsum ~ TCDsum + WAWsum + HubDist + Heather +<br>HeatherGrass                                   | 164.89 | 7.79  | 0.007 |
| <b>Woodlark</b>         |   |        |       |       |
| WLmocc2                 | ~JULIAN + WAWsum + HubDist + Heather + HeatherGrass ~ WAWsum +<br>Heather + HeatherGrass                    | 69.31  | 0.00  | 0.593 |
| WLmocc3                 | ~JULIAN + WAWsum + HubDist + Heather + HeatherGrass ~ WAWsum +<br>HeatherGrass                              | 70.75  | 1.44  | 0.288 |
| WLmocc1                 | ~JULIAN + WAWsum + HubDist + Heather + HeatherGrass ~ TCDsum +<br>WAWsum + HubDist + Heather + HeatherGrass | 73.10  | 3.79  | 0.089 |
| WLmdet3                 | ~JULIAN + WAWsum + HubDist + Heather + HeatherGrass ~ 1   | 75.29  | 5.98  | 0.030 |
| WLm0                    | ~1 ~ 1  | 100.55 | 31.24 | 0.000 |

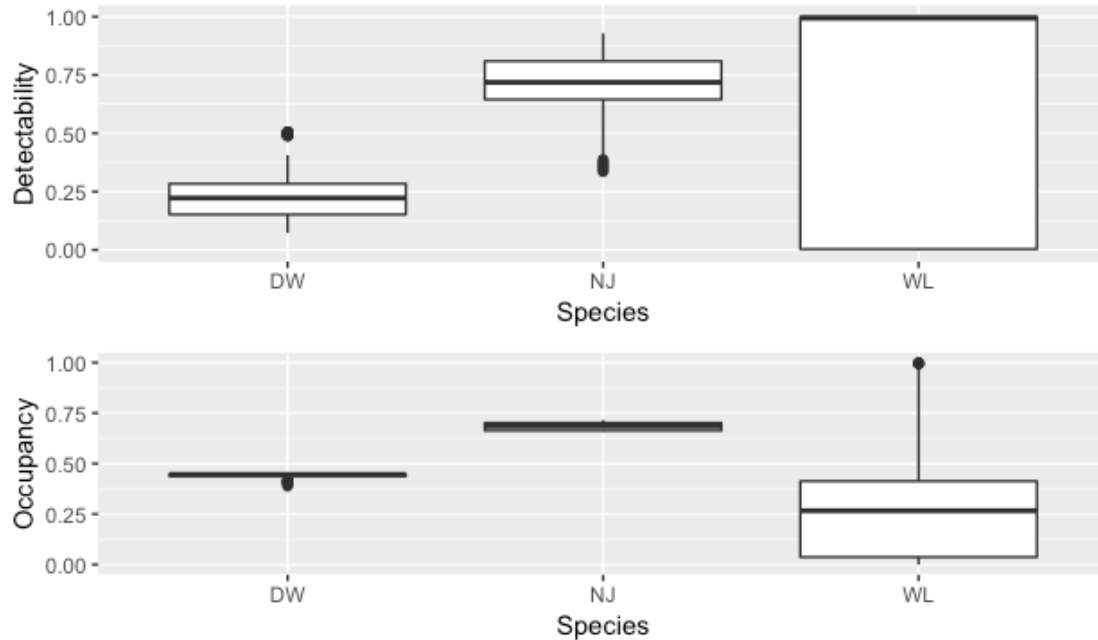
308 Table 2 Model selection list for all species - with detectability and occupancy covariates

309

310 Predicted occupancy varied little between sampling sites for nightjar and Dartford warbler (Figure 3), as  
 311 only single covariates were acting on these species - TCDsum and Heather grassland respectively.

312 Woodlark occupancy predictions varied more widely due to the number of habitat covariates acting on the  
 313 models for this species - including WAWsum, Heather and Heather grassland. Detectability predictions  
 314 were sensible for nightjar and Dartford warbler, but highly polarised to 0-1 in the models for woodlark,  
 315 due to the small number of positive sampling sites.





316

317 *Figure 3. Model-averaged predicted occupancy and detectability across all sampling sites, for Dartford*  
 318 *warbler (DW), nightjar (NJ), and woodlark (WL).*

319

320 Our results can be used to provide a baseline for assessing the population of the three heathland bird  
 321 species studied. We assumed that occupancy is a good surrogate for abundance (MacKenzie & Nichols,  
 322 2004) and that we could quantify the relative abundances of the bird species, based on the proportion of  
 323 sampling sites in which they were recorded to be present. Given the separation distances between recorder  
 324 locations in this study, it is considered reasonable to assume that each occupied sampling site represented  
 325 a separate territory/pair. Using the occupancy estimates from the null models for the three species we can  
 326 calculate that the areas of occupied habitat for each species, from a total 992 ha, are: nightjar 676 ha,  
 327 Dartford warbler 379 ha, woodlark 161 ha (Table 3). Combining these habitat areas with published  
 328 breeding densities of 0.074-0.078 males/ha for nightjar (Berry, 1979; Conway et al., 2007), 0.32-0.42  
 329 pairs/ha for Dartford warbler (Bibby & Tubbs, 1975), and 0.05 pairs/ha for woodlark (Langston et al.,  
 330 2007; Sitters et al., 1996), gives estimated population levels of: nightjar 51 males, Dartford warbler 140  
 331 pairs, and woodlark 8 pairs (Table 3).

| Species          | Occupancy (SE) | Occupied habitat (90% CI) | Density ha <sup>-1</sup> | Pairs (90% CI) |
|------------------|----------------|---------------------------|--------------------------|----------------|
| Nightjar         | 0.682 (0.0702) | 676 ha (562-791)          | 0.075                    | 51 (42-59)     |
| Dartford warbler | 0.382 (0.0914) | 379 ha (230-528)          | 0.37                     | 140 (85-195)   |
| Woodlark         | 0.162 (0.0562) | 161ha (69-252)            | 0.05                     | 8 (3-13)       |

332 Table 3. Calculated areas of occupied habitat, based on intercept-only occupancy estimates

333

## 334 4. DISCUSSION

### 335 4.1 Bioacoustic Approach

336 To our knowledge, this is the first study in Europe to combine bioacoustic survey with occupancy  
337 modelling. It is also the first in the UK to undertake a large scale survey for multiple bird species using  
338 automated recorders. It therefore expands the geographic scope of case studies for these methods, and  
339 applies them in a new habitat, beyond the American forested ecosystems in which most previous studies  
340 have been located (Furnas & Callas, 2015; Campos-Cerqueira & Aide, 2016; Furnas & McGrann, 2018;  
341 Wood et al., 2019).

342 We used species detection data from six repeated days of recording at 44 sampling sites, combining this  
343 with environmental covariates to estimate occupancy and detectability for three bird species. Our results  
344 show that the bioacoustic approach can be used effectively for the survey and monitoring of heathland  
345 bird populations. Although we included models where habitat covariates could influence occupancy in  
346 our candidate sets, the ‘best’ models for each species suggested that the habitat variables were not  
347 important indicators of occupancy at the scale studied. This is possibly due to the fact that the study areas  
348 were all lowland heathland sites, generally suitable for the study species, and so the distribution of  
349 individuals was likely to relate to micro-habitat features that were not detectable at the scale of the field

350 survey, satellite and map data applied. The satellite data used was at 20m pixel size, but the average size  
351 of the LCM polygons was 2.4 ha, equivalent to 87 m radius. Although the covariate data was sampled at a  
352 similar scale (100 m radius) to previous studies (Furnas & Callas, 2015; Campos-Cerqueira & Aide,  
353 2016), these were landscape-scale surveys less dependent on small habitat features to differentiate plots.  
354 Thus, we would agree with the finding of Niedballa et al. (2015), that both the spatial scale of habitat  
355 covariate data, and the radius sampled around survey sites, can affect the fit of occupancy models. Higher  
356 resolution data is needed for a site-based scale of assessment, if habitat covariates are to be included in  
357 analyses. For future studies, this should be gained from either field survey or high-resolution  
358 aerial/satellite imagery, such as the 5m resolution RapidEye imagery used by Niedballa et al (2015).

359

360 Identification of species vocalizations is commonly done either by complete manual analysis or,  
361 increasingly, by the use of automated recognizers, which require the *a priori* compilation and analysis of  
362 a large library of known species vocalizations (Knight et al., 2017; Shonfield & Bayne, 2017). Our  
363 analysis workflow included automated clustering of the acoustic data set, followed by manual validation  
364 of candidate vocalizations of the target species (Abrahams & Denny, 2018). This process has two  
365 benefits. Firstly, the automated clustering identified signals, that may be target bird species, but filtered  
366 out noise. In the current study, this allowed 77% of the total acoustic dataset to be filtered out, before  
367 identifications were attempted, significantly reducing the later workload in manually reviewing data for  
368 target species vocalizations. The second benefit of the analysis approach taken here, was that the manual  
369 validation step helped to minimize false-positive detections (Campos-Cerqueira & Aide, 2016;), which  
370 are often a significant issue with automated species identification systems (Zwart et al., 2014; Salamon et  
371 al., 2016). Misclassification errors such as this violate a major assumption of most occupancy models, and  
372 can lead to substantial errors in occupancy estimates (MacKenzie et al., 2006; Banner et al., 2018). The  
373 issue can potentially be addressed by complete manual identification of all recordings, but this is highly  
374 time-consuming, while the hybrid automated/manual approach taken here reduced the workload in the

375 manual review stage to less than a quarter of what it would have been. The corollary is that the data  
376 rejected by the automated clustering may contain target species vocalizations, and hence false-negatives  
377 may result. However, with the summation of the detailed call data down to daily presence/absence at each  
378 site, the potential loss of some target species phrases is considered unlikely to significantly affect the  
379 occupancy and detectability estimates derived from the modelling (Shonfield et al., 2018). The combined  
380 use of automated clustering and manual verification is therefore recommended as a valid approach for  
381 identification in bioacoustic studies.

## 382 **4.2 Spatial Sampling Design**

383 In bioacoustic studies with static sampling locations, the layout of recorder placements is of high  
384 importance. For occupancy modelling especially, the distance between sampling sites should be relevant  
385 to the territory size of the taxa being recorded (Niedballa et al., 2015), while also ensuring that the  
386 detection process is independent at each site by preventing overlap between the recording radius around  
387 each recorder. While this distance is variable, for many bird species the effective recording radius of most  
388 detectors is in the region of 50 m - although this is dependent on microphone model, variability and  
389 condition (Furnas & Callas, 2015; Turgeon et al., 2017; Yip et al., 2017). Within our study, the closest  
390 spacing between sampling sites was set by the ~250 m sampling grid. The mean nearest neighbour  
391 distances of the recorder sites were 316 m for Chobham, 346 m for Horsell, and 329 m for Thursley  
392 (range 202-703). Due to the sampling sites being spread across three survey sessions, the mean nearest  
393 neighbour distances between recorders in each session were 608 m, 466m and 508m.

394 For nightjar, a threshold of 350 m distance between registrations has been proposed to differentiate  
395 between male territories (Conway et al., 2007), while Stiffler et al (2018) applied a minimum spacing of  
396 400 m for recording wetland birds. The spacing of the recorders within the current study related well to  
397 these studies, and as a result, there can be a reasonable confidence that there was no double-counting for  
398 the bird species being studied. A 250 m sampling grid, as set out in the draft protocol of Abrahams (2018)  
399 is therefore considered to be appropriate for future studies, although additional refinement of detector

400 placement may be warranted to maximise coverage of sites, dependent on the vocal and territorial  
401 characteristics of the species being studied. For example, recent research has indicated that, for a desired  
402 threshold of detection efficiency, careful selection of optimised placements based on topography,  
403 vegetation and weather patterns, may be most efficient (Piña-Covarrubias et al., 2018).

#### 404 **4.3 Temporal Sampling Design**

405 In any occupancy study, the balance between the number of sites and number of sampling events  
406 differentially affects the accuracy and precision of the occupancy and detectability estimates. We  
407 recorded for six days at 44 sites, which we considered likely to balance fieldwork resourcing with  
408 sufficient sample site density. This was a longer deployment time than the two-three days used by Furnas  
409 & Callas (2015) and Stiffler et al. (2018), and equivalent to that employed by Campos-Cerqueira & Aide  
410 (2016) and Wood et al. (2019). For rare species with a high probability of detection (i.e. woodlark for this  
411 study) the required survey effort should maximize the number of sites covered, while for common species  
412 with low detection (i.e. Dartford warbler) the most efficient sampling approach is to increase the number  
413 of survey occasions (Mackenzie & Royle, 2005). With the low occupancy for woodlark found here, it is  
414 likely that an increased number of sampling sites (and lower number of survey days if necessary) would  
415 be likely to improve the modelling results (Mackenzie & Royle, 2005; Banner et al., 2018). This modified  
416 sampling approach would, however, have to be considered in terms of its costs/benefits, taking into  
417 account the potential effects on Dartford warbler modelling and increased fieldwork time or equipment  
418 requirements.

#### 419 **4.4 Detectability**

420 Using the null models, without covariates, we estimated detectability as 0.73 for nightjar, 0.49 for  
421 woodlark and 0.26 for Dartford warbler. The national Breeding Bird Survey (BBS) (Johnston et al., 2014)  
422 found a much lower detectability of 0.30 for nightjar, which is perhaps unsurprising, due to the  
423 difficulties with surveying this species within a standard (mostly daytime) survey method. However, the

424 BBS detectability estimates of 0.47 for woodlark and 0.37 for Dartford warbler are similar to those found  
425 in this bioacoustic study. In this comparison, nightjar is much better detected by acoustic recorders (as  
426 found by Zwart et al., 2014), but Dartford warbler less so, while detectability for woodlark is matched.

427 Taking detectability into account during traditional bird surveys requires repeated visits across the season.  
428 The time often occurring between site visits may then invalidate the assumption that detection probability  
429 remains constant across the survey events. The protocol used in this study enabled six days of back-to-  
430 back recording, simultaneously at 16 sites, minimising the risk that detection probability would change  
431 between sampling events. This would have been difficult to achieve without the use of automated  
432 recorders. The greater number of survey replicates achievable with the bioacoustics approach is therefore  
433 able to improve occupancy and detection estimates (MacKenzie et al., 2006; Stiffler et al., 2018).

434 We found that survey date, combined with habitat characteristics, explained detectability and improved  
435 the performance for some of the species models generated here, similar to the finding of Furnas & Callas  
436 (2015). Wetland (WAWsum) was a positive parameter on detectability for all three species, and woodland  
437 (TCDsum) was also positive for nightjar, as was Heather for woodlark. The probability of detecting a  
438 species during a bioacoustic survey is a function of both the probability of it vocalizing and the recorder  
439 detecting the call. The vocalization rates of many birds vary due to age, sex, breeding status, time of day,  
440 and seasonal variation (Campos-Cerqueira & Aide, 2016; Furnas & McGrann, 2018). As a consequence,  
441 both survey timing and the number of visits need to accommodate species vocalizing behavior to ensure  
442 accurate detection, particularly for species with sporadic vocalization patterns (La & Nudds, 2016). Age  
443 and sex-specific variation in vocalization rates cannot be accounted for easily when using automated  
444 recorders, but our methods allowed for the other variation factors, as we sampled over a relatively short  
445 period of time during the breeding season, and sampled over a wide timeframe every day, thereby  
446 minimising the potential for seasonal and diurnal variation in call rates. Our results, together with those of  
447 Johnston et al. (2014), showing how detection probability varies by species, should be considered in

448 decisions about study design when planning to survey birds using automated recorders or traditional  
449 methods.

#### 450 4.5 Occupancy

451 We calculated occupancy as 0.682 for nightjar, 0.382 for Dartford warbler and 0.162 for woodlark,  
452 showing that nightjar is widespread across the study sites, while woodlark has a much more restricted  
453 distribution. This is in line with other survey data for the sites, collected by traditional survey methods  
454 (J.Eyre & J.Clark; D. Boyd pers. comms.), and previous occupancy studies (Furnas & Callas, 2015;  
455 Campos-Cerqueira & Aide, 2016; Wood et al. 2019). Although the occupancy figures provide a  
456 population estimate in themselves, they could potentially be used to generate an estimate of the number of  
457 pairs, as the common measure for population size. We did this provisionally, using a combination of  
458 habitat area and previously recorded breeding densities to give the following numbers: Dartford warbler  
459 140, nightjar 51 and woodlark 8.

460 The occupancy modelling indicated a positive relationship between nightjar and TCDsum. This  
461 corresponds to associations with woodland found in previous studies (Bright et al., 2007; Conway, 2010].  
462 The negative relationship between Dartford warbler and Heather Grassland was surprising, as this species  
463 is generally associated with dry-humid heath, and gorse, sometimes with a grassy component (Bibby &  
464 Tubbs, 1975). Woodlark occupancy was positively related to Heather Grassland, and negatively to  
465 WAWsum and Heather. These results are more expected, as nest sites for this species are generally found  
466 in tall/dense heather or grass (Mallord et al., 2007), while foraging sites have short grass and bare ground  
467 (Conway et al., 2009).

468

469 **5. CONCLUSION**

470 Our study demonstrates the suitability of the bioacoustics approach to identify the distributions and assess  
471 the populations of target bird species on heathland study areas. Occupancy and detectability estimates  
472 were produced, taking into account imperfect detection. If carried out on a regular basis, this method  
473 could provide a valuable new approach for monitoring of population levels and favourable conservation  
474 status. For future studies in this setting, and with these species, methods might be improved by increasing  
475 the number of sample sites at which recording takes place. This approach would be likely to improve the  
476 modelling for woodlark, but would need to be balanced against potential effects on models for the other  
477 two species studied.

478 The field of conservation biology is continuously adopting improved, cheaper and more easily available  
479 technologies. In the near future, automated interpretation of recordings using machine learning methods  
480 will become increasingly viable, allowing effective identification of a range of bird species (Brandes,  
481 2008; Acevedo & Villanueva-Rivera, 2009; Knight et al., 2017; Shonfield & Bayne, 2017, Stowell et al.,  
482 2019). The permanent nature of bioacoustic recordings will allow these ongoing developments in call  
483 analysis and automated identification to be used to re-analyse previously collected data, perhaps alongside  
484 new recordings (Shonfield & Bayne, 2017; Stiffler et al., 2018). The use of bioacoustics will, therefore,  
485 be indispensable for conducting long-term and potentially continuous monitoring over large spatial scales,  
486 aiding understanding of the ongoing effects of threats and management practices on bird populations on  
487 heathland and in other environments.

488



489 **AUTHORS' CONTRIBUTIONS**

490 CA conceived the ideas, designed methodology; collected and analysed the data. CA led the writing of the  
491 manuscript, with MG contributing to establishment of occupancy modelling methods and development of  
492 the text. Both authors contributed critically to the drafts and gave final approval for publication.

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495 **DATA ACCESSIBILITY**

496 Data, metadata and R script has been archived at Mendeley Data

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## 660 **SUPPLEMENTARY INFORMATION: APPENDIX 1**

661 Kaleidoscope 4.3.2 software settings

662 File parameters:

663 • No subdirectories

664 • No split to max duration

665 • Split channels—yes.

666

667 Signal parameters:

668 • Signal of interest 1500–7000 Hz

669

670 • Duration 2–20 s

671 • Maximum inter syllable gap 1 s

672

673 Scan and cluster recordings:

674 • Max distance 1.0

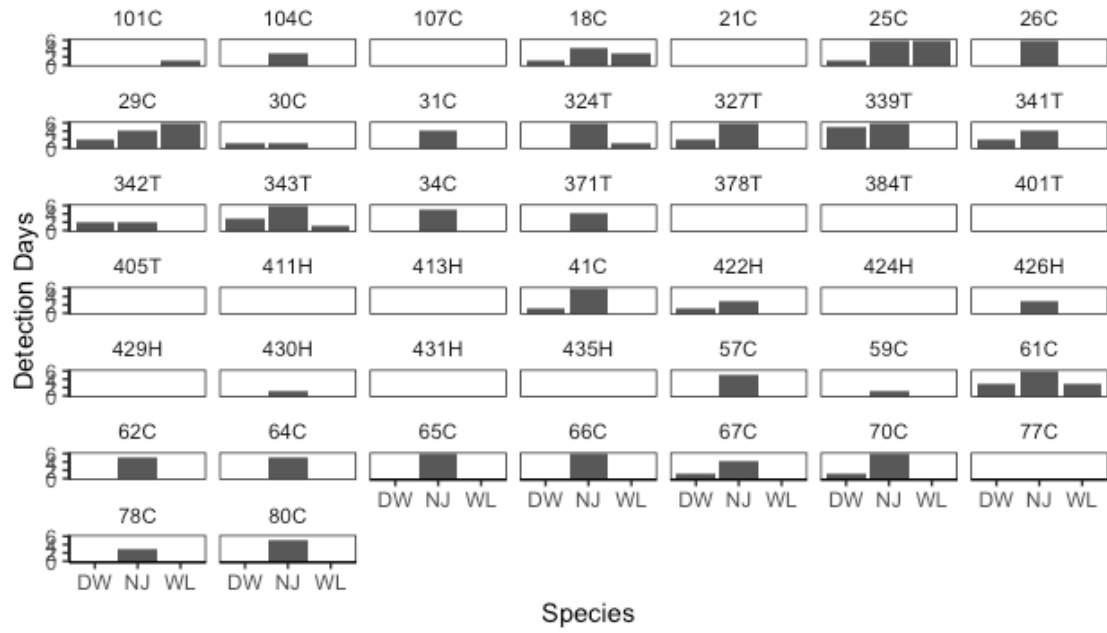
675 • FFT window 5.33 ms

676 • Max states 12

677 • Max distance for building clusters 0.5

678 • Max clusters 500

679



680 Figure 4. Number of detection days for each species at each site.

681

682