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1	Metabarcoding reveals selective dietary responses to
2	environmental availability in the diet of a nocturnal, aerial
3	insectivore, the European Nightjar (Caprimulgus europaeus)
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21	Many hird analisa are vulnerable to environmental change, or knowledge of their dist
22	Many bird species are vulnerable to environmental change, so knowledge of their diet
23	and its variation can help to understand population status and flexibility to respond to
24	change. Insectivorous species are predicted to have a flexible diet within and between
25	individuals, which can respond to naturally fluctuating prey abundance, thus allowing

26 opportunistic exploitation of available resources. We analysed the diet of a nocturnal, aerial insectivore, the European Nightjar Caprimulgus europaeus, using high-throughput 27 metabarcoding. We quantified diet diversity and composition of 130 faecal samples from 28 29 nests and roosts on a northern breeding site in the UK from 2015 - 2018, and compared differences among individuals and years. Although dominated by moths, diet varied 30 significantly between individual faecal samples and between years and months. Prey 31 species composition varied between years, and was more variable between samples in 32 2017, compared to other years. Faecal samples were significantly more likely to contain 33 moth species with a wingspan of >60mm and less likely to contain moth species of 34 <25mm wingspan. This indicated size-selective foraging, which also varied between 35 months and years. Diet was driven by inter-individual variation, indicating population-36 level flexibility in prey choice. Metabarcoding provided a valuable tool in the exploration 37 of insectivorous diets, but efforts are needed to build comprehensive reference libraries, 38 in order to compile full prey species lists. 39

40

Keywords: avian ecology, diet, metabarcoding, faecal sampling, 16S rRNA gene, moth,
Lepidoptera

Understanding variation in the composition and diversity of diet within a bird 43 population can assist understanding of how a population or species is likely to fare in 44 response to anthropogenic change (Howells et al. 2017). Environmental change is 45 likely to affect certain groups of species more acutely than others; for example, 46 specialist predators reliant on ephemeral insect prey may be more sensitive (Nocera 47 et al. 2012, Stanton et al. 2017). Species that take flying insect prey on the wing 48 49 ('aerial insectivores') are identified globally as being at risk (Nebel et al. 2010, Nocera et al. 2012) due in part to the significant decline in insect populations 50 51 worldwide (Hallmann et al. 2017, van Strien et al. 2019). The diet composition of aerially-feeding, insectivorous birds may reflect localised spatial and temporal 52 variation in prey abundance related to habitat type (Mills et al. 2020) or weather 53 conditions (Imlay et al. 2017). Their diet may also reflect the ease of catching 54 particular prey items, their nutritional content or habitat guality (Garlapow 2007, 55 English 2009, Sharps 2013, Razeng & Watson 2015). 56

Difficulty observing feeding, and in finding and dissecting faecal matter 57 (Mumma et al. 2016, Nielsen et al. 2018) mean that there is little information on the 58 diet of adult, insectivorous birds, especially nocturnal species. Moreover, soft-bodied 59 moths and flies that are primarily taken by this feeding guild are not readily 60 identifiable in faecal matter (Razgour et al. 2011, Trevelline et al. 2016). Recently, 61 rapid development of molecular techniques such as metabarcoding, has allowed 62 researchers to acquire valuable information relevant to species conservation through 63 diet analysis (Gillet et al. 2015, Kress et al. 2015, Gerwing et al. 2016). 64 Metabarcoding facilitates diet analysis using faecal samples, by identifying short 65 sequences from specific genes within the DNA of prey items that remain in the faecal 66 matter. Commonly-used barcodes include the cytochrome c oxidase subunit I (COI) 67

and the 16S rRNA gene, and these can effectively distinguish between species
(Metzker 2010, Alberdi *et al.* 2012, da Silva *et al.* 2019) where the barcodes have
high interspecific but low intraspecific diversity. As well as allowing samples to be
processed in bulk (Taberlet *et al.* 2012), these methods are non-invasive and can be
more comprehensive than visual identification (Ji *et al.* 2013, De Barba *et al.* 2014,
Krehenwinkel *et al.* 2017), aspects that are particularly beneficial for vulnerable or
secretive species (Thalinger *et al.* 2018).

The European Nightjar Caprimulgus europaeus (henceforth, 'Nightjar') is a 75 nocturnal insectivore of conservation concern in the UK (Amber listed - Eaton et al. 76 2015, IUCN classification is least concern, BirdLife International 2016). This species 77 completes an annual migration from central Africa to the UK to breed from May until 78 August, and feeds on arthropods (Cramp 1985, Sharps 2013). Limited diet data have 79 been obtained through stomach content analysis and physical dissection of faeces 80 from European Nightiars and the Common Nighthawk Chordeiles minor (Sierro et al. 81 2001, Sharps 2013, Knight et al. 2018). However, these methods result in a bias 82 towards remaining hard parts of consumed arthropods such as beetle elytra, legs 83 and antennae, which means that methods such as metabarcoding that are capable 84 of identifying partially or fully digested remains are needed. Extracting sufficient DNA 85 from avian faeces is challenging, especially after degradation by exposure to sun, 86 rain and soil microbes (Deagle et al. 2006, Oehm et al. 2011, Kamenova et al. 2018). 87 The high concentration of uric acid (Eriksson et al. 2017) in conjunction with 88 environmental exposure inhibits extraction and amplification of DNA, making the 89 extraction and clean-up stages of metabarcoding, along with choice of primer 90 sequences, critical to success (Alberdi et al. 2017, Carneiro et al. 2020). 91

Here we have analysed the diet of a sample of Nightjar individuals at a key breeding site in the UK. To understand variation in Nightjar diet within this population we used metabarcoding techniques, methods which have had limited application to the diet of this species prior to now (Evens *et al.* 2020). We used metabarcoding to test the following hypotheses:

1) Diet composition and diversity varies among individuals (as in Alberdi *et al.*2020), as individual Nightjars at this site have been shown to vary significantly in
their habitat preferences (Mitchell *et al.* 2020).

2) Females have lower diet diversity than males (da Silva *et al.* 2020), as they
only spend a short time foraging due to higher investment in parental care.

3) Diet diversity and composition varies between years. Nightjars are
constrained in their breeding schedule due to migration, so their short window of
opportunity should coincide with peaks of preferred moth prey. However, weather,
and thus emergence and activity of moths, can vary substantially throughout the
breeding season and between years (Clare *et al.* 2011, 2014).

4) Nightjars select large moths more often than their relative abundance in the
environment, due to both nutritional value (Razeng & Watson 2015), and greater
ease of detecting large moths in low light (Bayne & Brigham 1995).

110

111 METHODS

112 This study took place at the Humberhead levels National Nature Reserve (NNR;

South Yorkshire, UK, comprising Thorne Moor (53.636N, 0.898W) and Hatfield Moor

114 (53.545N, -0.938W). The site is classified as a degraded raised bog, with 28% of the

site covered by bog or marsh, 10% by standing water, and 15% mixed woodland

(JNCC 2015). It is also a European Special Protection Area (SPA) for its breeding 116 Nightjar population (1.9% of the UK population; JNCC 2015). The NNR has 117 undergone substantial habitat management since 2014, primarily the removal of 562 118 km² of birch woodland and an increase in wet habitat through the damming of 119 drainage channels. Both techniques will influence vegetation structure and insect 120 prey availability; selective and patchy clearance of woodland has been shown to 121 122 increase species richness of Lepidoptera (Summerville & Crist 2002), and should also provide an increased amount of breeding space for Nightjars (Sharps 2013). 123 124 Use of wetland habitat (e.g. bogs, mires) by nightjars is little considered in the literature explicitly as a breeding habitat, but there is some indication that they act as 125 suitable foraging areas, containing a high number of invertebrates (English et al. 126 2017, Evens et al. 2017). 127

128

129 Faecal sample collection

Nightjar faecal samples were collected from June to August, 2015 to 2018 (2015: n =130 9, 2016: n = 20, 2017: n = 35, 2018: n = 36) and were obtained from previously 131 located roosts and nests, during the day. Upon approaching the nest or roost, we 132 used binoculars to view the bird in situ. If the bird was not visible, we slowly 133 134 approached the location until the bird flew away, whereupon the sex of the bird was visible (male Nightjars possess bright white spots on primaries 3,4 and 5 whilst 135 females have smaller, beige spots). Nests and roosts of Nightjars and the size and 136 137 shape of their faeces are distinctive and identifiable, as they are discrete curled pellets of between 3 – 8mm in width, found individually or in small clusters (see 138 image in supplementary information). Faeces were placed into sterile, 30ml screw-139 top tubes (Elkay labs, UK) with wooden toothpicks, then labelled with sample 140

number, date, location, and the sex of the bird. Because samples were taken from 141 roosts and nests, some samples at nests may have been from juveniles, which 142 cannot be sexed. Samples from nests were therefore excluded from comparative 143 analyses between years, months and sexes. All samples were included in analyses 144 of prey size. Birds were not individually identifiable in the field, so it is possible that 145 some samples are repeats from the same bird. Roost and nest details were 146 147 documented and these locations are held faithfully by individuals throughout the season, so we consider it unlikely that there is much duplication of individuals within 148 149 the dataset. Where we have repeatedly sampled the same roost, or nest, we document these as likely to be from the same bird. Although defecations at roost and 150 nest sites could potentially be from more than one bird (e.g. the male and the female 151 at the nest), the male only visits the nest for short periods of time, and where we find 152 fresh defecations during the day when the female is on the nest, we associate these 153 with the female. Additionally, males were found by Berry (1979) not to roost with the 154 female at the nest, but separately, often up to 100 metres away. This ties in with 155 other information presented by Berry where strict fidelity to roost sites by individual 156 male Nightjars was found within and between seasons. 157

Fresh faecal samples, identified by their soft consistency, were targeted. Where there were multiple faecal pellets at a location, we took softer, fresher pellets over dry samples. Where there were only drier, older pellets, we took a sample, but visited the site again in the next 2-3 days in order to find a fresher sample, both in order to improve the amount of DNA extracted and to better pinpoint deposition date. Samples were transferred to a -20°C freezer within four hours of collection, and then to a -80°C freezer in batches every seven days, and stored there until processing. All

laboratory work took place at the NERC Biomolecular Analysis Facility (NBAF) within
 the Department of Animal and Plant Sciences, at the University of Sheffield, UK.

167

168 **DNA extraction**

We extracted DNA from faecal samples using the QIAmp DNA Fast Mini Stool Kit 169 (Qiagen, Germany), following the human DNA extraction protocol (Supplementary 170 Material, Appendix I), adapted to suit more highly degraded DNA from bird faeces by 171 a) increasing the amount of sample (300mg vs standard 220mg), b) using the 172 inhibitex buffer provided with the kit at a higher volume than recommended (1.4ml vs 173 standard 1ml), and c) incubating samples for 10 minutes longer than recommended 174 175 after the lysis buffer was added to increase the number of cells broken down from the sample and increase the DNA yield. We quantified DNA concentration using a 176 Fluostar Optima (BMG Labtech, Germany) and kept the DNA extracts in a -20°C 177 freezer until PCR. 178

179

180 Polymerase Chain Reaction (PCR)

Primer sets were selected based on their specificity and their ability to amplify short 181 fragments of degraded DNA (fragments of between 100 - 180 base pairs (bp)), 182 based on data from previous studies (Zeale et al. 2011, Clarke et al. 2014, Alberdi et 183 al. 2018). Two commonly-used primer sets targeting the mitochondrial cytochrome 184 oxidase COI gene: the 'Uniminibar' primer set (Meusnier et al. 2008) and the 'ZBJ' 185 primer set (Zeale et al. 2011) were tested, but not used because they produced 186 inconsistent amplification across different groups of insects (also highlighted in 187 Brandon-Mong et al. 2015), and amplified primer dimer (Brownie et al. 1997), when 188

specific products failed to amplify. We then tested the Ins16S 1Short F and R primer 189 set that amplifies a 156bp fragment of the mitochondrial 16S region (Clarke et al. 190 2014), identified in the literature as providing wider taxonomic coverage across 191 invertebrate orders. The use of these 16S rRNA primers resulted in more consistent 192 amplification levels across all samples. A temperature-gradient PCR from 55 – 70°C 193 (Supplementary Material, Appendix I) with a random subset of faecal samples (n =194 195 4), moth (n = 4), Nightjar control samples (DNA from egg albumen collected at nests, n = 4) and negative water samples (n = 2). The optimal PCR annealing temperature 196 197 was identified as 62°C. This temperature amplified moth tissue positive control samples but not Nightjar DNA. PCR products were visualised on a 1% agarose gel 198 with a 100bp ladder (Invitrogen, Thermo Fischer Scientific, UK) to verify amplification 199 200 success.

201

202 **Reference DNA library**

Although the INS16S 1Short primers provided good taxonomic coverage, there are 203 unfortunately many fewer reference sequences available with which to compare 204 sequence reads (Clarke et al. 2014, Tournayre et al. 2020). To improve the ability to 205 identify sequences from faecal samples, we created a local reference 16S sequence 206 207 library, representative of some of the assemblage present on the site. We extracted DNA from 81 specimens of 80 species of moths and beetles collected from the moth 208 traps, to supplement those already available in the BLAST global database (Altschul 209 210 et al. 1990). Specimens were identified to species based on visual and microscopic examination by LJM, using the criteria of Townsend and Waring (2014). Extractions 211 followed an ammonium acetate precipitation method (Bruford et al. 1998), where a 212 digestion solution, Proteinase K and ammonium acetate were added to each sample 213

of moth legs and incubated overnight, to penetrate the invertebrates' solid chitin-214 based exterior. Extracted insect DNA was Sanger-sequenced using di-dedoxy dye 215 terminators (see Supplementary Material, Appendix I for protocol) using both the ZBJ 216 COI primers (Zeale et al. 2011) and the INS16S 1Short primers (Clarke et al. 2014). 217 Sequences were visualised using an ABI3730 DNA Sequencer (Applied Biosystems, 218 ThermoFischer Scientific, USA). Due to the lack of British 16S moth sequences in 219 220 the NCBI /EMBL /DDJB databases, we verified species identification by comparing the sequenced moth 16S genes against COI sequences, which are more frequently 221 222 present in the BLAST database. Finally, 16S sequences from 78 invertebrate species were submitted to the NCBI GenBank database (sequence accession 223 numbers: MK620910 - MK620988). 224

225

226 Sequencing and bioinformatics

Detailed sequencing preparation and procedures are provided in the Appendices.
Briefly, samples were cleaned, individually labelled, quantified and then pooled for
sequencing. We then used the high-powered computer clusters '*Iceberg*' and '*Sharc*'
at the University of Sheffield, to process and quantify the raw reads generated by the
Illumina Miseq Benchtop Sequencer.

232 Sequences were processed, trimmed and clustered into 'Molecular 233 Operational Taxonomic Units' (MOTUs). The threshold at which sequences are 234 clustered – i.e. their level of similarity – strongly influences how many MOTUs are 235 produced. MOTUs are taxonomic proxies (Hemprich-Bennett *et al.* 2018), and are 236 frequently interpreted as equating to species (da Silva *et al.* 2020). Depending on the 237 level of clustering used however (most commonly between 96% and 99%; Clare *et* *al.* 2016), sequences may be equated to family or order (Clare *et al.* 2016,
Hemprich-Bennett *et al.* 2018, Gordon 2019). We clustered our MOTUs at 97%,
based on our knowledge of the close-relatedness of Lepiodptera species (Rytkonen *et al.* 2019), as well as an understanding that if degradation has taken place, as is
common with DNA within faecal matter, it may be better to take a more conservative approach (Clare *et al.* 2016).

Sequences were then compared to those stored in the NCBI GenBank 244 nucleotide database (including those of the 78 potential prey species that we 245 submitted) using a Nucleotide BLAST (blastn; Altschul et al. 1990). Because of the 246 lack of 16S reference sequences of British Lepidoptera, some MOTUs were 247 automatically assigned to species not present in the UK or to multiple species with 248 an identical level of certainty. We therefore manually assigned such species to 249 genus or family, using the NCBI taxonomy browser. For all diversity analyses we 250 251 therefore used a matrix of MOTUs, rather than species and/or genera because of the described discrepancy in assigning units (as in Razgour et al. 2011). Classifying 252 sequence reads this way allowed us to include all units identified through the 253 clustering process, whether or not we were able to assign them to a known taxa 254 (Clare et al. 2011, Hawlitschek et al. 2018, da Silva et al. 2020). MOTUs that had 255 been assigned to specific non-target species (Homo sapiens, Sus scrofa, Columba 256 palumbus and Akkermansia sp), were removed (da Silva et al. 2020). 257

Finally, we created matrices of final read numbers and presence/absence matrices in R (v. 3.6), which were used to filter out low numbers of reads (Galan 2018), that are likely to represent contamination rather than true sequences. Following De Barba *et al.* (2014), we used our negative control samples to set thresholds for each PCR amplification, sequences with numbers below these

thresholds were considered as noise, and removed (see Supplementary material,
Appendix II for more detail). False positives can occur at multiple stages of the
extraction and PCR procedures (Ficedola *et al.* 2015, Zepeda-Mendoza *et al.* 2016,
Alberdi *et al.* 2018) and caution over such sequences is important (Froslev *et al.*2017).

Sequencing data will be available online upon publication through the
European Nucleotide Archive (ENA), part of the European Molecular Biology
Laboratory (EMBL), as fastq files.

271

272 Statistical analyses

To examine the diet of this population of Nightjars as a whole, we calculated the 273 frequency of occurrence of sequences (FOO) – i.e. the proportion of samples in 274 which each MOTU was found. Conversion of sequence data into a more quantitative 275 estimate of diet, known as relative read abundance (RRA), has flaws (see discussion 276 in Deagle et al. 2019). However, the use of FOO alone can be conservative, as it 277 essentially equates one sequence that registers 10,000 reads, with another that only 278 registers 10 reads, because the only criterion is to be present (Deagle et al. 2019). 279 Thus, we use RRA to calculate all metrics, bearing in mind that numbers of 280 sequence reads do not perfectly translate to percentage biomass (Rytkönen et al. 281 2019). 282

To examine variation in composition between samples, we used multivariate PERMANOVAs (using '*adonis*' in '*vegan*'). These tested for differences in the means of Chao dissimilarity indices (R package '*vegan*', function '*vegdist*'; Oksanen *et al.* 2019) between sex, month and year (McClenaghan *et al.* 2019a). Because it was not

possible to sex chicks, these were removed from the data for calculation of the PERMANOVAs, as were within-sample repeats. Sex, month and year in which the sample was collected, were modelled separately as they could not be included in the same PERMANOVA, because there were missing data from both the sex and month categories. We used Chao dissimilarity indices within the PERMANOVAs in order to account for unknown species, and because they were stable in our unbalanced sample size scenario (Chao *et al.* 2005, Chao & Chiu 2016).

PERMANOVAs are a non-parametric version of a MANOVA and are stable 294 when faced with heterogeneity within groups (Anderson 2017). We used package 295 'pairwise Adonis' (Martinez Arbizu 2020) to conduct among-year pairwise 296 comparisons. As we had some unbalanced sample sizes, we calculated 297 homogeneity of group variances (function 'betadisper' in vegan (Oksanen et al. 298 2019)), which calculates the distance from each sample to the group centroid 299 300 (Anderson, 2006) to estimate compositional variance within groups, using the Chao dissimilarity indices again. We calculated 95% confidence intervals for all tests using 301 Tukey's HSD. 302

For a random subset of the samples, we undertook repeat extraction, PCR 303 and sequencing. Although most variation is attributed to between sample variation 304 305 (Mata et al. 2019), it is possible that due to a large number of variables such as size of the dropping, digestion time and size of the prey items (Oehm et al. 2011), there 306 may be variation within a faecal sample as well (Ando 2020). We compared the 307 within-sample repeats using Chao dissimilarity indices, calculated from RRA as 308 above. The repeat samples are included in the supplementary information file, 309 marked with the suffix 'r'. 310

311 Prey size variation and selection by Nightjars

To understand if size-selective foraging took place, we used moth traps to obtain an 312 indication of flying insect prey availability on the breeding site. We placed 15W 313 actinic light traps (made in the electronics department at the University of York -314 details and photos within the supplementary information), at eight locations within the 315 316 boundary of Thorne and Hatfield Moors (n = 16). Moths were trapped at four sites for one night in each alternate week (i.e. sites 1-4 for one night (e.g. overnight Thursday 317 - Friday) in weeks 1, 3, 5 etc., followed by sites 5-9 for the same night in weeks 2, 4, 318 6 etc.) starting at the beginning of June until the middle of August. Traps were placed 319 at each site within two hours prior to dusk and emptied the following morning within 320 one hour of dawn. Trap locations were identified as grid squares on a map of the 321 site, and the traps placed in a randomly chosen but consistent location on the ground 322 within this square. Although the same grid squares were being sampled (each 323 square was sampled n = 4 times), the two-week interval between sampling 324 occasions should remove some concern about repeat sampling (Truxa & Fiedler 325 2012). The sites chosen for moth trapping included all habitat types present on the 326 nature reserve. Habitat types were classified as woodland (trees with canopy >2 327 metres high), scrub (trees with canopy <2 metres high), heather, wetland (rush-328 329 dominated, <50% standing water), bog (>50% standing water, Eriophorum dominated, bare-ground (bare peat areas with minimal (< 10% vegtation cover) and 330 'off site' – all areas outwith the boundary of the NNR). Habitat structure was also 331 incorporated, so as to include information about habitat manipulation (there were 332 three categories here: unmanaged, machine-manipulated (mass management using 333 flail/digger) or hand-manipulated (selective chainsawing, volunteers with handtools). 334 We recorded numbers of species and individuals, calculated diversity (Inverse 335

Simpson's dominance; Morris *et al.* (2014)) and allocated species to a size category
(in mm; <25, 25-30, 30-40, 40-50, 50-60, or >60) based on Townsend and Waring
(2014).

To explore size composition for the subset of MOTUs identified to species 339 level within each sample, including those fed to chicks, we allocated each species 340 341 identified within the faecal samples to a size category according to mean wingspan, using Townsend and Waring (2014). To test for significant differences in the 342 frequency of occurrence of different size classes between years we used Kruskal-343 Wallis rank sum tests. To test for size selection of moth prey by Nightjars, we used 344 proportional Z-tests to compare the frequency of occurrence of each size class found 345 in the faecal samples, to those found in the traps, between years and months. 346

347

348 **RESULTS**

We retrieved a total of 13,900,616 sequence reads from 141 Nightjar faecal samples 349 using the Illumina MiSeg Benchtop sequencer, which comprised 1631 MOTUs after 350 351 trimming and clustering. After processing and filtering (i.e. removing 'impossible' reads likely to be a result of contamination and degradation: De Barba et al. 2014), 352 we retained 625 unique MOTUs from 130 samples. Some samples (n = 11), 353 contained no reads once filtered, meaning that the amount of non-arthropod 354 contamination was high, as has been found in other studies (Regnaut et al. 2006). 355 65% of MOTUs were identified to species, 5% to genus, 12% to family and 18% only 356 to order (all Lepidoptera). Lepidoptera were found in 99% of samples, followed by 357 Diptera (27%), Coleoptera (9%), Neuroptera (7%) and <1% of both Hemiptera and 358 359 Hymenoptera. Of the sequences identified to species level, the most common moth

species present were Poplar Hawk-moth Lathoe populi (in 43% of samples), Silver Y 360 Autographa gamma (45%), Yellow-tail Euproctis similis (45%), True Lover's Knot 361 Lycophotia porphyrea (49%), Smoky Wainscot Mythimna impura (50.4%), and Large 362 Yellow Underwing Noctua pronuba (52%). Of the Coleoptera we were able to 363 identify, we found Harpalus sp. (Carabidae; 2% of samples), Melanotus villosus and 364 Stenagostus rhombeus (both Elateridae and in 6% of samples). Diptera sequences 365 366 identified belonged to the families Chironomidae, Sciaridae, Culicidae and Tipulidae, but were not identifiable to any more detailed level than that of family. All identified 367 368 species are listed in the Supplementary Information (Appendix III).

369 Variation in composition and diversity among samples

Samples contained an average of 15.6 MOTUs (+/- SD 9.7). Mean Chao dissimilarity of adult samples (2015: n = 8, 2016: n = 17, 2017: n = 27, 2018: n = 27; no chicks and no repeat samples, and a reduced total number of 429 MOTUs) was 0.91 (+/-SD 0.17), which shows relatively high individual variation. Mean Chao dissimilarity of the repeat samples was 0.85 (+/- SD 0.19), therefore making related samples slightly more similar.

No significant differences were found in sample composition between sexes 376 (Female: n = 26, Male: n = 43; $F_{1,68}$: 0.36, P = 0.99), nor between the months in 377 which the samples were collected (June: n = 21; July: n = 32; August: n = 15; F_{2,70}: 378 1.21, P = 0.16). Between-year differences in sample composition were significant 379 $(2015: n = 8, 2016: n = 17, 2017: n = 27, 2018: n = 27; F_{3,75} = 3.3, P = 0.001; r^2:$ 380 381 0.12), supported by a non-significant difference in the distribution of samples around the centroid ($F_{3,75}$: 1.37, P = 0.26; Figure 1), although some caution must be taken 382 due to uneven sample sizes. Specific significant among-year differences in variance 383 were shown between 2016 – 2017 and 2017 -2018 (Table 1). 384

386 Moth species sizes within faecal samples

96% of faecal samples contained moths with a wingspan of between 30 and 40mm and 46% of samples contained the largest moths (>60 mm wingspan). The five most frequent species in the samples all had a mean wingspan of more than 30mm; two of these had a wingspan of more than 50mm (*N. pronuba, L. populi*). The largest moths (>60mm) were present in significantly more samples in 2018 (66%), than all other years (2015: 0%; 2016: 14%; 2017: 36%; X^2 = 27.08, df = 3, *P* <0.0001).

393

394 Size selection in diet samples compared to prey availability

Faecal samples were more likely than moth traps to contain the largest moths (>60mm). This was significant in 2017 (diet samples: 0.4, moth traps: 0.18; X^2 = 4.61, df= 1, *P* = 0.03) and 2018 (diet samples: 0.67, moth traps: 0.34; X^2 = 9.5, df =1, *P* = 0.002; Figure 2). In contrast, moth traps were more likely than faecal samples to contain smaller moths (<25mm), but the difference was only significant in 2018 (diet samples: 0.18, moth traps: 0.49; X^2 = 9.55, df =1, *P* = 0.002; Figure 3).

Across all years, less than 18% of diet samples contained moths with the smallest wingspan (<25mm). In June and August, faecal samples were more likely than moth traps to contain the largest moths (June: 50-60mm: diet samples: 22%; moth traps: 4%, X^2 : 11.66, df =1, P = <0.001. August: >60mm: diet samples: 15%, moth traps: 3%, X^2 : 5.73, df = 1, P = 0.02).

407 **DISCUSSION**

We have demonstrated the use of high-throughput sequencing in investigating the 408 diet of an insectivorous, avian migrant to the UK, joining a growing bank of literature 409 exploring the use of metabarcoding to reduce the bias shown by physical dissection 410 of faeces towards retained, undigested fragments of prey items (Evens et al. 2020, 411 Mills et al. 2020). Our analyses showed strong intra- and inter-annual variation in diet 412 413 richness and size-based prey selection, which varied annually and seasonally. Differences among years in Nightjar diet potentially reflected a response by Nightjars 414 415 to environmental fluctuation influencing the annual life cycle of moth species. There was a dominance of Lepidoptera over other arthropods in Nightjar diet, 416 which has been reported previously (Sierro et al. 2001, Sharps 2013, Evens et al. 417 2020), although this does not hold true for all species of caprimulgid (Knight et al. 418 2018). The prevalence of Lepidoptera in the diet of this population may reflect local 419 availability of this prey type (exceedingly high numbers of larger moths, or very few 420 Coleoptera) and an adjustment of the population to being almost entirely Lepidoptera 421 specialists. It is also possible that due to our molecular optimisation procedures, 422 there has been a lack of amplification of groups other than Lepidoptera. However, 423 previous tests using the primers in our study showed very high taxonomic coverage 424 (>96%) for 11 invertebrate orders, and showed higher success for Coleoptera than 425 for Lepidoptera (Clarke et al. 2014). 426

427 Moths present in the faecal samples were diverse in terms of their habitat 428 preferences and population trends. The most common species found in faecal 429 samples were all habitat generalists (e.g. *N. pronuba, E. similis*). True Lover's Knot, 430 however, is a specialist of heathlands, its larvae feeding exclusively on heather 431 (Townsend & Waring 2007, Thomsen *et al.* 2015). A number of heathland and wet

woodland specialist species (including for example, Oak Eggar Lasiocampa quercus, 432 Map-winged swift Hepialus fusconebulosa and Gold Spot Plusia festucae were 433 caught in the light traps, but most species found in both locations were generalists, 434 found in such widespread habitats as hedgerows, woodlands and gardens. Although 435 some such generalist species have more positive population trends than habitat 436 specialists, large moths such as those preferentially selected by Nightjars, and 437 438 another migrant insectivore the Common Cuckoo Cuculus canorus are, overall, in decline (Coulthard et al. 2019, Mills et al. 2020, Fox et al. 2021). The most 439 440 commonly identified moth species within Nightjar faeces, the Large Yellow Underwing, has increased over the past 50 years by 72%, clearly thriving in 441 generalist habitats (Harrower et al. 2020). Conversely, True Lover's Knot, Silver Y, 442 Yellow-Tail, Poplar Hawkmoth and Smoky Wainscot, the other five most common 443 components of Nightjar faecal matter, have all declined since 1968 by between 1.5 444 and 83% (the latter being the heathland specialist, True Lover's Knot). 445

Samples differed significantly from each other. Although some samples may 446 have been taken from the same individual, intraspecific variation in diet is common 447 and can reflect both resource availability and the avoidance of competition through 448 resource partitioning (Kotler & Brown 1988, Maldonado et al. 2019, Alberdi et al. 449 2020), or differences in habitat selection. We tracked a number of individuals in this 450 population over the same four years of this study, and this tracking work highlighted 451 a lack of overall population preference for any particular habitat type (Mitchell et al. 452 2020). The lack of similarity in diet among samples potentially also indicates a lack of 453 association with a particular habitat for foraging. For species exploiting patchy, 454 ephemeral resources as the Nightjars are here, it is beneficial to be flexible to be 455

able to adjust to prey availability and abundance (Maldonado *et al.* 2019, Szigeti *et al.* 2019).

Contrary to expectation, no significant difference in sample composition or 458 diversity was found between male and female Nightiars (Mata et al. 2016, Knight et 459 al. 2018). We expected to find that females had less diverse diets than males, 460 461 because of their limited foraging opportunity (Houston 1997). However, not all females were incubating during the period when the faecal samples were acquired, 462 so perhaps were not restricted in their foraging duration. Differences between years 463 in sample variance and prey species richness may indicate that the population could 464 respond to variation in prey availability, by taking advantage of emergences of 465 particular species, perhaps including influxes of migratory moths, several species of 466 which (e.g. Silver Y) were identified within our Nightjar samples (Lee & McCracken 467 2005). Annual and seasonal variation in moth community diversity and overall 468 469 abundance have been shown to be related to variation in climate, as well as habitat management (Summerville & Crist 2004, Summerville et al. 2007). 470

By comparing diet and moth trap samples, we examined consumption 471 compared to local availability, bearing in mind that not all the MOTUs identified in the 472 diet samples could be translated into species and that moth trap catches do not 473 474 sample moths representatively (see Truxa & Fiedler 2012, Jonason et al. 2014 for information on bias in trapping and Evens et al. 2020 for use of the same type of 475 trap). It is also pertinent to note that some Nightjars make trips away from their 476 477 nesting areas to separate foraging locations, in this and other studies (Alexander & Cresswell 1992, Evens et al. 2020, Mitchell et al. 2020), which means that 478 depending on habitat type at any distant foraging locations, we will not have 479 completed accounted for all prey available to Nightjars across their individual 480

foraging ranges. Again, their preference for large moths, which should provide higher 481 energy gain per item than small moths is a strategy also recognised in other 482 nocturnal insectivores (Clare et al. 2009, Vesterinen et al. 2016). Twilight foraging 483 limits the time available for capturing prey, which should drive size bias upwards in 484 line with the energy maximization – time minimization rule (Pyke 1984, Schoener 485 2003), and our findings suggest that availability of larger moths might be a key 486 487 aspect of Nightjar foraging strategy and requirements (Araújo et al. 2011, Schrimpf et al. 2012, Vesterinen et al. 2016, Evens et al. 2020). 488

Understanding the demographic consequences of variation in prey availability 489 is important in order to make practical decisions about future management. The 490 habitat management that occurred on our study area has resulted in a reduction in 491 birch woodland, a rich invertebrate prey source. However, the outcome of this is not 492 necessarily negative, if regenerating, herbaceous, understorey vegetation that 493 results from the clearance sustains sufficient numbers of moths (Summerville & Crist 494 2004) and continuing low-intensity management supports large moth species (Mills 495 et al. 2020). Whilst Nightjars may be resilient to variation in prey availability on an 496 annual basis, it remains especially important to encourage habitat types that provide 497 larval food for moths during the Nightjar chick raising period to allow offspring to 498 499 fledge.

500

501 Use of metabarcoding for insectivorous diets

Metabarcoding has made it possible to identify physically unrecognisable remains of
prey specimens for many taxa (e.g. Kartzinel & Pringle 2015, Hawlitschek *et al.*2018, McClenaghan *et al.* 2019a). Despite considerable progress, such methods are

still hindered by a few key challenges. Firstly, contamination of samples with nontarget DNA means that caution must be taken when conducting laboratory work and
the post-sequencing bioinformatics (Zepeda-Mendoza *et al.* 2016). Although one of
the benefits of metabarcoding is that less common prey species can be identified,
the post-sequencing filtering thresholds that must be established to produce
confident results, can sometimes remove rare sequences.

Additionally, some studies have found within-sample variation in MOTU 511 composition and diversity (Alberdi et al. 2018, Mata et al. 2018), although between-512 sample variation was always stronger. We did find lower within-sample variation than 513 between-sample variation, but both measures were higher than in these other 514 referenced studies. We do not know a sufficient amount about Nightjar gut transit 515 time so we are unable to quantify how many specimens or prey capture events were 516 contained in each faecal sample, but high within-sample variation could relate to a 517 518 high number of specimens per defaecation, or may relate more to the methods used to extract the DNA. This does indicate that there are several steps that may be 519 advised in order to ensure maximum information and variation is obtained, such as 520 multiple extracts and PCR replicates of the same sample, as well as repeated 521 sampling from the same individuals across the season, where known (Alberdi et al. 522 2018). The latter would ensure that individual specialisation in diet could be recorded 523 through longitudinal data collection as opposed to a 'snapshot' of an individual's diet 524 (Araujo et al. 2011). Although there is concurrence across general metabarcoding 525 protocols, the specific details such as number of replicates and use of controls are 526 not standardised, which can limit utility and cross-study comparison (Deagle et al. 527 2019, Ando et al. 2020). 528

For our study, by far the biggest of these challenges was a lack of reference 529 sequences for particular groups of species in global databases (such as NCBI 530 Genbank used for BLAST processing). This lack of reference sequences meant that 531 species could not be identified for around 35% of our MOTUs, even after the addition 532 of 78 species of moth to the 16S reference database (compared to 90% of COI 533 sequences identified to species in Evens et al. 2020). This means that despite being 534 535 able to assess sample richness, variation and change between months and years, understanding the ecological information that comes with a species-level 536 assignment, which we are lacking, is invaluable (Hebert et al. 2003). Although we did 537 contribute several dozen 16S sequences to GenBank, we know from a 538 comprehensive compilation of Lepidoptera records on Thorne Moors alone (not both 539 Moors that constitute the Humberhead Peatlands NNR), that there have been 717 540 moth species recorded (382 macro-, 335 micromoths; Moat 2014). This indicates 541 that much more structured, systematic barcoding of the 16S rRNA gene of British 542 moth species needs to take place, to further this type of work. 543

544

545 **Conclusions**

Nightjars showed within-population and annual diet variation, as well as size selection. The population appears able to adjust its diet in response to variation in prey availability, reflecting ecological theory. This has consequences for Nightjars' ability to withstand habitat change, with populations and species exhibiting diverse diets often able to adapt better. For cryptic, insectivorous species such as the Nightjar, metabarcoding provides a very useful tool to identify diet components, despite continuing challenges related to extraction through to sequencing.

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560

561 Data availability statement

562 Raw Fastq sequencing files will be openly-available upon publication through the

563 ENA, part of the EMBL under the primary accession number PRJEB44974.

564 Summary files for statistical analysis are available as an excel file in SOM 4.

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Figure 1. Mean and variance of faecal sample distances to group centroids for each year samples
were collected, produced by the '*betadisper*' function in r. Sample sizes vary between groups and are
presented in the text.



Figure 2. Proportion of each moth size class present in diet samples and moth trap catches, for each year. Differences between faecal samples and moth trap samples were calculated using proportional Z-tests and significance levels are denoted using: ** = <0.05, *** = <0.01. Note: No moth trapping took place in 2015.

Table 1. Pairwise comparisons of among-year differences in mean sample composition using the

898 function pairwise.Adonis in R which takes the Chao dissimilarity values as input. Significant adjusted

P values (< 0.05) in bold.

		Sums of				Р
Pairs	df	Sq	F	R ²	Р	adjusted
2015 vs 2016	1	0.663	1.587	0.065	0.034	0.204
2015 vs 2017	1	0.623	1.452	0.042	0.089	0.534
2015 vs 2018	1	0.872	2.301	0.065	0.006	0.036
2016 vs 2017	1	1.656	4.064	0.088	0.001	0.006
2016 vs 2018	1	2.495	6.775	0.139	0.001	0.006
2017 vs 2018	1	1.061	2.758	0.050	0.006	0.036