

Please cite the Published Version

Shutt, Jack D ^(D), Nicholls, James A, Trivedi, Urmi H, Burgess, Malcolm D, Stone, Graham N, Hadfield, Jarrod D and Phillimore, Albert B (2020) Gradients in richness and turnover of a forest passerine's diet prior to breeding: a mixed model approach applied to faecal metabarcoding data. Molecular Ecology, 29 (6). pp. 1199-1213. ISSN 0962-1083

DOI: https://doi.org/10.1111/mec.15394

Publisher: Wiley

Version: Accepted Version

Downloaded from: https://e-space.mmu.ac.uk/625378/

Usage rights: O In Copyright

Additional Information: This is an Author Accepted Manuscript of a paper accepted for publication in Molecular Ecology, published by and copyright Wiley.

Enquiries:

If you have questions about this document, contact openresearch@mmu.ac.uk. Please include the URL of the record in e-space. If you believe that your, or a third party's rights have been compromised through this document please see our Take Down policy (available from https://www.mmu.ac.uk/library/using-the-library/policies-and-guidelines)

1	Title: Gradients in richness and turnover of a forest passerine's diet prior to
2	breeding: a mixed model approach applied to faecal metabarcoding data
3	
4	Running title: Pre-breeding dietary gradients of a passerine
5	
6	Authors: Jack D Shutt ^{1‡} , James A Nicholls ^{1,2‡} , Urmi H Trivedi ³ , Malcolm D Burgess ^{4,5} ,
7	Graham N Stone ¹ , Jarrod D Hadfield ^{1#} , Albert B Phillimore ^{1#*}
8	
9	Affiliations:
10	1 Institute of Evolutionary Biology, The University of Edinburgh, The King's Buildings,
11	Edinburgh, EH9 3FL, UK;
12	2 Current address: Australian National Insect Collection, CSIRO, Clunies Ross Street, Acton,
13	ACT 2601, Australia;
14	3 Edinburgh Genomics, The University of Edinburgh, The King's Buildings, Edinburgh, EH9
15	3FL, UK;
16	4 Centre for Research in Animal Behaviour, University of Exeter, Exeter, EX4 4QG, UK;
17	5 RSPB Centre for Conservation Science, The Lodge, Sandy, Bedfordshire, SG19 2DL, UK
18	
19	‡ and # Contributed equally.
20	Correspondence: Albert Phillimore albert.phillimore@ed.ac.uk
21	Orcid ID's: JDS 0000-0002-4146-8748, MDB 0000-0003-1288-1231, ABP 0000-0002-6553-
22	1553
23	

24 Abstract

Little is known about the dietary richness and variation of generalist insectivorous 25 species, including birds, due primarily to difficulties in prey identification. Using 26 27 faecal metabarcoding we provide the most comprehensive analysis of a passerine's diet to date, identifying the relative magnitudes of biogeographic, habitat and temporal 28 29 trends in the richness and turnover in diet of Cyanistes caeruleus (blue tit) along a 39site, 2° latitudinal transect in Scotland. Faecal samples were collected in 2014-15 from 30 31 adult birds roosting in nestboxes prior to nest building. DNA was extracted from 793 samples and we amplified COI and 16S minibarcodes. We identified 432 molecular 32 33 operational taxonomic units (MOTUs) that correspond to putative dietary items. Most dietary items were rare, with Lepidoptera being the most abundant and taxon-rich prev 34 order. We present a statistical approach for estimation of gradients and inter-sample 35 variation in taxonomic richness and turnover using a generalised linear mixed model. 36 We discuss the merits of this approach over existing tools and present methods for 37 38 model-based estimation of repeatability, taxon richness and Jaccard indices. We find that dietary richness increases significantly as spring advances, but changes little with 39 elevation, latitude or local tree composition. In comparison, dietary composition 40 41 exhibits significant turnover along temporal and spatial gradients and among sites. Our study shows the promise of faecal metabarcoding for inferring the macroecology of 42 food webs, but we also highlight the challenge posed by contamination and make 43 44 recommendations of laboratory and statistical practices to minimise its impact on inference. 45

46

48 Keywords

Beta diversity, avian/bird, Jaccard, insectivore, prey, repeatability, blue tit, Cyanistes
caeruleus

51

52

53 Introduction

54

55 Insectivorous passerine birds in temperate environments tend to be dietary generalists feeding on a broad range of invertebrate taxa (Betts, 1955; Cholewa & Wesołowski, 56 57 2011). There is potential for the diet of such generalists to vary over geographic gradients, among habitats and seasonally within a year. Such dietary variability within 58 generalist species is poorly understood and could have profound ecological 59 consequences. Spatial variation in resource availability has implications for 60 geographic patterns in population density, breeding productivity and the degree to 61 which local adaptation in resource use may evolve. Seasonal variation in resource 62 consumption has implications for the optimal scheduling of life history events, such 63 as reproduction (Charmantier et al., 2008; Durant et al., 2005) and seasonal movements 64 65 (Thorup et al., 2017).

66

67 Spatiotemporal trends in diet will arise from a combination of underlying trends in 68 invertebrate resource availability and the prey preferences of the consumer. Species 69 richness – or α -diversity – of temperate invertebrate taxa generally decreases with 70 increasing latitude (Baselga, 2008) and peaks at mid-elevations (Beck et al., 2017) in 71 the summer months (Thomsen et al., 2016). Within forests, invertebrate richness can 72 vary among tree taxa by more than an order of magnitude, and in the UK has been found to be highest on Salix, Ouercus and Betula (Kennedy & Southwood, 1984; Shutt, 73 Burgess, & Phillimore, 2019). In addition to changes in species richness, species 74 75 composition may change from one community to the next, which is quantified as β diversity (Baselga, 2010; Whittaker, 1972). While there is evidence that forest 76 77 invertebrate communities show turnover over biogeographic gradients (Novotny & 78 Weiblen, 2005) and among host tree taxa (Murakami, Ichie, & Hirao, 2008), the 79 relative magnitude of turnover along different gradients has received scant attention 80 (Novotny & Weiblen, 2005). Whether diet mirrors these gradients in resource 81 availability will largely depend on how much prey selection by the consumer departs from random. 82

83

Forest-dwelling hole-nesting insectivorous birds, such as blue tits (Cyanistes 84 caeruleus), have been subject to decades of intensive study (C. M. Perrins, 1979). 85 While the diet of nestlings has proven relatively straightforward to quantify, either via 86 videos/cameras at the nest (Samplonius, Kappers, Brands, & Both, 2016), or neck 87 collars on nestlings (Burger et al., 2012), much less is known about the diet of adults 88 (but see Cholewa & Wesołowski, 2011; J. A. Gibb, 1954). The paucity of information 89 about adult diet arises because these birds often forage high in trees on small prey 90 items. To date most of our taxonomic information on adult tit diet has been derived 91 from dissections of the gizzard and gut contents of euthanised birds (Betts, 1955), a 92 method that precludes the identification of soft-bodied dietary items, has relatively 93 94 poor taxonomic resolution (e.g. order or family level) and is destructive. These studies reveal that tits consume various insects (including Lepidoptera, Hemiptera, Diptera, 95

96 Coleoptera, Hymenoptera) and spiders, as well as some plant matter in winter (Betts,

97 1955; Cholewa & Wesołowski, 2011; Cramp & Perrins, 1993).

98

99 The advent of next-generation sequencing and faecal DNA metabarcoding now provides a non-destructive means of obtaining diet information at a fine taxonomic 100 101 resolution (Pompanon et al., 2012; Symondson, 2002; Taberlet, Coissac, Pompanon, 102 Brochmann, & Willerslev, 2012). Where invertebrates comprise a large proportion of 103 the diet, DNA barcodes from the rapidly evolving cytochrome oxidase I (COI) 104 mitochondrial gene have become the standard and allow identification to species-level 105 in many cases (Kress, García-Robledo, Uriarte, & Erickson, 2015). To date, most published faecal metabarcoding studies have examined variation in mammalian diet 106 107 (Bohmann et al., 2011; Clare, Symondson, Broders, et al., 2014; Clare, Symondson, & Fenton, 2014; Razgour et al., 2011). In comparison to mammals in general, and bats 108 in particular, application of faecal metabarcoding for inference of the diet of avian 109 110 insectivores is a small but rapidly growing field. Progress has been hampered by the challenge of extracting and successfully amplifying dietary DNA from avian faeces 111 112 (Jedlicka, Sharma, & Almeida, 2013; Vo & Jedlicka, 2014). As such, avian faecal 113 metabarcoding studies have sampled small numbers of individuals and/or locations (Table 1) and the latter limitation has precluded detailed analysis of the drivers of 114 115 spatial or temporal variation in the diet of avian insectivores (for an exception see 116 Sullins et al., 2018).

117

118 To date the statistical tools employed by the nascent metabarcoding field have largely 119 borrowed from community ecology. In some studies the objective has been to describe 120 the diet composition of a taxon such that statistical analysis may be unnecessary (De Barba et al., 2014). Metabarcoding studies that focus on patterns in taxon richness 121 commonly apply a two-step analysis, first using rarefaction to quantify diversity at a 122 123 focal sampling level and then using a statistical model to examine variation in taxon richness among samples (Quéméré et al., 2013). Studies interested in how taxonomic 124 125 composition varies among samples have tended to rely on pairwise metrics, such as 126 the Jaccard index, and non-parametric methods, such as PERMANOVA and the 127 Mantel test (Alberdi, Aizpurua, Gilbert, & Bohmann, 2018; Mata et al., 2019; 128 Trevelline, Nuttle, Hoenig, et al., 2018). Generalised linear mixed models (GLMMs) 129 and their extensions provide a method for including structure in the data collection and multiple predictors into an analysis (Warton et al., 2015), but few studies have utilised 130 131 them in diet metabarcoding to date (for exceptions see Mata et al., 2019; Nichols, 132 Åkesson, & Kjellander, 2016).

133

134 Here we employ faecal metabarcoding using COI minibarcodes to infer the diet of an insectivorous woodland passerine, the blue tit, in early spring along a 220 km transect 135 in Scotland (Appendix 1 Fig. S1). We have three main aims: (i) to quantify dietary 136 137 taxon richness and composition at the molecular operational taxonomic unit (MOTU) level; (ii) to quantify the magnitude of changes in both measures along gradients of 138 time (day of year), latitude, elevation and tree taxon composition; and (iii) to quantify 139 140 gradients in the contributions that six key invertebrate orders (Araneae, Coleoptera, Diptera, Hemiptera, Hymenoptera, Lepidoptera) make to diet. We show that by 141 142 applying a GLMM to presence/absence data it is possible to estimate changes in taxon richness and turnover among points and along gradients. We also demonstrate how 143

this mixed model approach can be used to estimate repeatability and control for sometypes of systematic contamination.

146

147

148 Material and Methods

149

150 Field data collection

151 Fieldwork was conducted during the springs of 2014 and 2015 at 39 predominantly 152 deciduous woodland sites that together comprise a 220km latitudinal transect in 153 Scotland (Shutt, Bolton, Benedicto Cabello, Burgess, & Phillimore, 2018). At each 154 site there were six Schwegler 1B 26mm-hole nestboxes distributed at approximately 155 40m intervals. From mid-March the base of each nestbox was lined with greaseproof paper – with the aim of slowing DNA degradation (Oehm, Juen, Nagiller, Neuhauser, 156 & Traugott, 2011) - which was replaced when damaged or heavily soiled, and 157 158 removed at the onset of nest building or once a bird had attempted removal. Each nestbox was inspected on alternate days and faeces on the greaseproof paper were 159 160 removed with sterilised tweezers (after use they were wiped with lab tissue and 161 flamed), with up to a maximum of three faeces collected in a 2mL Eppendorf tube containing pure ethanol. The total number of faeces in a nestbox was recorded 162 (excluding 129 samples from early 2014). Samples were stored at -18°C within a day 163 of collection and transferred to a -20°C freezer at the end of each spring. Faecal 164 samples were collected from 35 of the 39 sites from 19 March in 2014 and 18 March 165 166 in 2015 until nest building, giving a median sampling range of 20 days per site in 2014 and 24 days in 2015 (Appendix 1 Table S1). 167

168

Latitude (site range 55.98 – 57.88°N) and elevation (10 – 433m) were obtained for
each nestbox (Shutt et al., 2018). Site-level habitat metrics were derived from surveys
of numbers of trees of different genera belonging to three size classes (based on girth
at breast height) within 15m radius of each nestbox, as described in Shutt et al. (2018).
The site-level habitat variables we considered were total foliage, tree diversity
(Simpson's index), the amount of oak foliage and the amount of birch foliage (Shutt

175 et al., 2018).

176

177 Molecular protocol

We balanced sampling across nestboxes and dates as far as possible by imposing an upper limit of 10 samples per nestbox per year and where this maximum was exceeded we subsampled such that we maximised the range of dates per nestbox. If multiple faeces (n = 2 - 3) were present within a sample tube, part of each individual scat was used for the DNA extraction with the aim of sampling a broader range of diet. This protocol resulted in processing of 793 of a total of 959 faecal samples.

184

Thirty samples were processed in duplicate to allow us to estimate technical repeatability. The selected samples were evenly distributed throughout the sampling period, including samples from multiple sampling locations in both 2014 and 2015. The faeces for each of the 30 duplicated samples were evenly divided into two and DNA extractions were performed on each subsample; although each subsample contained sections from along the entire length of the original faeces, the faeces was not completely homogenised before subsampling. Each duplicate extraction was subsequently treated as an independent sample for all downstream processes. All aspects of the laboratory protocol (DNA extraction, PCR amplifications, PCR cleanup, sequencing on a MiSeq run) were performed at different times using different aliquots of reagents for each replicate within a pair of subsamples. In addition we included 24 controls (including extraction negatives, PCR negatives and *Dryocosmus israeli* as a non-native invertebrate PCR positive).

198

199 DNA was extracted from faecal samples using the QIAamp DNA Stool Mini kit, 200 following the protocol for pathogen detection with a few custom modifications 201 designed to improve DNA vields (see online protocol for details; 202 dx.doi.org/10.17504/protocols.io.ve6e3he). Three loci were targeted for amplification through PCR - the standard animal barcoding gene (COI), a secondary barcoding gene 203 204 to detect invertebrate prey DNA and confirm the faecal sample originated from a blue tit and no other hole-roosting or -nesting passerine (16S rRNA), and a standard plant 205 206 barcoding gene (rbcL) (see online protocol for further details; dx.doi.org/10.17504/protocols.io.2jdgci6). Given that DNA from dietary items is 207 expected to be very degraded, the primers used amplified a small 'minibarcode' region 208 209 of each gene (184-220 base pairs). Invertebrate primer sets were validated to ensure that they would amplify DNA from the expected range of invertebrate taxa (two orders 210 211 of arachnids, isopods, nine insect orders).

212

We followed a two-stage PCR process, firstly to amplify the target regions, then secondly to add indexed Illumina adaptors to the amplicons from each sample. Amplicons were multiplexed into three pools, each containing between 273 and 276

samples (inclusive of 30 replicate samples) and 8 controls (3x PCR positives, 3x PCR
negatives and 2x extraction negatives; a total of 24 controls across the whole
experiment). Each pool was sequenced on an Illumina MiSeq, using 150 bp pairedend reads.

220

221 **Bioinformatics processing**

222 Sequencing reads were initially de-multiplexed into sets corresponding to individual 223 faecal samples using the index combinations present within the adaptor sequences 224 using bcl2fastq (version v2.17.1.14). Reads were then de-multiplexed using fastq-225 multx from ea-utils (version1.1.2-537) with parameter '-m 2' into sets corresponding to each locus using the locus-specific primer sequences present at the beginning of 226 227 each read. Adaptor sequences, primer sequences and poor quality base calls were then removed using cutadapt (version 1.8.3) with parameters: '-m 50', '-q 30', '-f fastq', 228 229 leaving only sequence corresponding to the targeted gene regions. Subsequent 230 processing of the sequences applied the UPARSE pipeline (initially developed for 16S 231 metabarcoding of bacteria, (Edgar, 2013)) to data for each locus separately.

232

The first step in the bioinformatics pipeline was to merge the paired reads derived from either end of the sequenced fragment. This process was successful for all COI and rbcL reads and many 16S reads; 16S reads derived from avian DNA did not overlap, but comparison with known blue tit 16S sequences indicated that these reads could be combined by adding four "N"s between the forward and reverse reads to produce a composite sequence of the correct length (hereafter referred to as fused reads). Reads were then filtered to ensure that within a locus they were all of the same length; this

240 process removed possible pseudogenes incorporating insertions/deletions from the 241 coding COI and rbcL data. The rbcL data were not used for subsequent analyses in this 242 study, and 16S data were only used to confirm the faeces were derived from blue tits. 243 The set of filtered COI sequences was then used for two purposes. Firstly, the set of 244 unique sequences present within the full data set derived from all samples was 245 determined, with counts made of their frequencies. Unique sequences represented by 246 only a single read were removed as they most likely represent sequencing errors. The 247 unique sequences were then clustered into molecular operational taxonomic units 248 (MOTUs), grouping sequences together that had an identity of 98% or more. The most 249 frequently occurring sequence within each MOTU was designated as the reference 250 sequence for that MOTU. The second use of the filtered reads involved mapping them 251 back to this reference set of MOTU sequences on a sample by sample basis, allowing 252 a mismatch of up to 2% between filtered reads and a reference sequence, to provide a more accurate assessment of the frequency of each MOTU within each faecal sample. 253 254 The taxonomic identity of MOTUs was determined using a BLAST search of the 255 reference set of MOTU sequences against public databases (GenBank and BOLD).

256

257 Quality control and MOTU refinement

Our analysis plan from this point on was pre-registered (osf.io/xgvm8). Some aspects of our methods deviate from what was outlined in the pre-registration (see table S2 in appendix 1 for an explanation of the motivation for these departures). We tested whether samples were from blue tits by verifying the presence of blue tit fused 16S sequences. The highest number of blue tit 16S reads from the 24 control samples was 58 and as a precaution all faecal samples that yielded fewer than 100 blue tit 16S reads were excluded from further analyses as they were not conclusively confirmed to be blue tit faeces (n = 9). Of the remaining avian faecal samples, blue tit was the commonest of the fused 16S MOTU in all but one sample, but this sample still had sufficient (n = 1465) blue tit reads to confirm its identity. No other avian DNA was present in any sample.

269

270 COI reads were checked from control samples to confirm the presence of positive 271 control species and provide a baseline for background contamination. All nine PCR 272 positive control samples contained MOTUs attributable to Dryocosmus israeli (range 273 of reads = 7796 - 19115) and no more than 16 reads of any other MOTU identified as belonging to the Metazoan kingdom. Eight out of nine PCR negative controls 274 275 contained no more than 19 reads of any MOTU. The ninth was highly contaminated and contained 6798 reads arising from more than 20 MOTUs. Therefore, we checked 276 for contamination along rows or columns within plates by estimating Spearman's 277 278 correlations in the number of MOTU reads between samples in neighbouring cells in the same PCR column or row. The row containing the contaminated negative sample 279 280 was found to have a substantially higher mean level of within row correlation (r = 0.37) 281 than other row and column correlations (mean r = 0.04). This was considered to be most likely a systematic contamination event and this row (n = 11 focal samples + 1 282 negative control) was excluded from all analyses. In addition, closer inspection of the 283 contaminated plate revealed two wells (both faecal samples) in the neighbouring row 284 to the contamination event containing very similar MOTUs with the contaminated row 285 286 and these were also removed from further analysis. Of the six extraction negative controls, four contained no MOTU at a higher read frequency than 3. The remaining 287

two contained contamination (maximum reads = 10037 and 1611) but on further inspection there was no evidence for this being systematic. As there were few cases where a control (positive or negative) had > 20 reads for any non-target MOTU, we adopted 20 reads as the cut-off for identifying MOTU presence.

292

293 The above steps reduced the number of samples from 847 to 824 (772 focal) containing 294 2524 MOTUs. All MOTUs with fewer than 20 reads in any single sample were 295 removed as possible false positives (remaining n = 1432 MOTUs). All MOTUs 296 without any BLAST match, or identified as environmental contamination, were 297 removed (remaining n = 1323). Then, a full taxonomy was obtained for each remaining 298 MOTU and taxonomic reduction of the dataset began to eliminate non-prey items. 299 Firstly, only MOTUs belonging to the Metazoan kingdom were considered possible prey items (remaining n = 1078). Then, all MOTUs not belonging to the phyla 300 Annelida, Arthropoda and Mollusca were discarded (remaining n = 1005). Finally, all 301 302 mites in the dataset of orders Astigmata (11), Mesostigmata (56), Oribatida (1), 303 Siphonoptera (2) and Trombidiformes (24) were removed, as they were likely to be 304 ectoparasites rather than actively foraged prey (remaining n = 911). For the MOTU 305 identification we required that the percentage match was at least 90% (remaining n =785). Taxa identified to an identification match of 90% or more are considered correct 306 307 to a minimum of order level, and this is the level that is important to the analyses in 308 this study. Several MOTUs identified as 'Arachnida sp' were removed on finding that these MOTUs were most closely matched to fungi (remaining n = 778). All 309 310 Dryocosmus (positive control) and waxworm (Galleria mellonella – from a feeding experiment in 2014 that provided 10 waxworms in a plastic cup adjacent to two 311

312 nestboxes per site) MOTUs were removed (remaining n = 757). Then, all remaining MOTUs belonging to the same best-hit taxon were merged (remaining n = 432). 313 Finally, due to the importance of Lepidoptera to tit diet we assessed the biological 314 315 plausibility of Lepidoptera identifications, which was possible due to comprehensive 316 UK occurrence data for this order (Sterling & Parsons, 2012; Waring & Townsend, 317 2017). Nineteen of 131 Lepidopteran MOTUs assigned species names were reassigned 318 to a British species when this species was within a 1% match of a geographically 319 implausible top hit. We assigned species status to taxa with a 99% or greater identity 320 match with the BLAST hit and a histogram of identity matches is provided (Fig S2).

321

322 Statistical analyses

Analyses focussed on the presence/absence of MOTUs in a sample, as read numbers are not considered a reliable measure of the amount of a MOTU in a sample due to biases in primer binding and amplification (Clare, 2014; Yu et al., 2012). Control samples were excluded from analyses. DNA within a sample was often derived from multiple faeces, and the effect of this on MOTU presence was controlled for by including number of faeces as a four-level categorical fixed effect (1, 2, 3, unknown).

To examine geographic, habitat and temporal variation in blue tit diet (Shutt, Nicholls, et al., 2019), we included the presence or absence of each MOTU in each sample as the response variable in a Bayesian generalized linear mixed model (GLMM) with a probit error structure (Hadfield, 2010). This analysis excluded the replicate samples (for reasons discussed in Appendix 2). The effects of year and number of faeces in the sample (treated as categorical) and the effects of ordinal date, latitude, elevation, total 336 foliage, birch foliage, oak foliage and tree diversity (treated as continuous) were treated as fixed. These fixed effects quantify trends in dietary richness. After 337 338 accounting for these trends, variation in richness amongst sites, nest-boxes, days 339 within year (categorical) and faecal samples were modelled by fitting each term as 340 random. MOTU effects were fitted as random in order to capture differences amongst 341 MOTUs in their overall prevalence. Variation in the prevalence of individual MOTUs 342 amongst sites, nest-boxes, days within years (categorical) and faecal samples was 343 modelled by interacting each term with MOTU. In the core model we also allowed the 344 prevalence of individual MOTUs to vary with ordinal date, latitude and elevation 345 effects, again by interacting each term with MOTU to form random regressions. The 346 three slope terms were allowed to covary with each other and the main MOTU effect (the intercept). We also include plate by MOTU random interaction term to control for 347 any plate-wide contamination by particular MOTUs present. To estimate and correct 348 349 for any tendency for contamination of rows or columns within a plate we ran an 350 additional model with row (within plate) and column (within plate) interacted with 351 MOTU as random terms and this is the main model that we present in the results.

352

In addition to the core model, we also ran four additional models, each of which allowed the prevalence of individual MOTUs to vary across one of the four habitat variables. The additional random slope terms were allowed to covary with the original three slope terms and the intercept. However, because of the length of time that the core model took to run (three months) we excluded the day within year term and its interaction with MOTU. The importance of these effects are minor relative to other terms in the model (day in year variance = 0.003, day in year:MOTU variance = 0.036, table S4A) and the interaction in particular contributed a lot to computation time
because with 91 days and 432 MOTUS there are nearly 40,000 effects. All models
were run for 260,000 iterations, with the first 60,000 removed as burn-in and thinning
every 100. These models took two months to run on an iMac 10.13.6 with 3.4 Ghz
Intel core i7, 16GB RAM and 4 cores.

365

366 To examine trends in the presence/absence of prey orders in blue tit diet, the dataset was reduced down to presence/absence of the six most common orders (Araneae, 367 368 Coleoptera, Diptera, Hemiptera, Hymenoptera and Lepidoptera), termed 'focal 369 orders', which together comprise over 91% of all prey taxa identified. A similar 370 GLMM to that described above was then employed, but with focal order and date, 371 latitude, elevation and tree diversity individually and interacted with focal order as fixed effects. Site, nest-box, day and faecal samples were fitted as random main effects 372 and as random interactions with focal order. These models were run for 195,000 373 374 iterations, with the first 45,000 removed as burn-in and thinning every 75.

375

376 To assess the repeatability of the approach we used a similar analysis to that described 377 above with the presence/absence of each MOTU as a response for the faecal samples for which extraction, PCR and metabarcoding had been replicated (29 samples x 432 378 MOTUs). Fixed effects were year and the number of faeces in the sample, both as 379 380 factors, with random terms limited to MOTU, faecal sample ID, faecal sample ID by MOTU interaction, extraction sample ID and residual. This model was run for 13 381 382 million generations with the first 3 million removed as burn-in and thinning every 5000. 383

All numeric predictor variables in all analyses were scaled to have a mean of 0 and a variance of 1 to provide direct comparability of results. We used parameter expanded priors for the variances such that the marginal priors on all variances followed a scaled (1000) $F_{1,1}$ distribution. Traces of posteriors were visually inspected to check for convergence and adequate sampling. For the main model, the effective sample sizes (ESS) were a bit low for some variances (< 500), but in all cases the ESS were adequate

391 to provide a reliable point estimate (>100) even if in some instances the accuracy of 392 the credible intervals is poor. As a test of model adequacy we conducted posterior predictive simulations to assess whether key features of the data were captured (Fig. 393 394 S3). We opted to use an MCMCglmm approach rather than much faster numerical 395 integration approaches, such as lme4 (Bates, Maechler, & Bolker, 2012) or glmmTMB 396 (Brooks et al., 2017), because posterior predictions revealed that parameter estimates from MCMCglmm provided an accurate description of the data, whereas those from 397 398 lme4 and glmmTMB were highly inconsistent (Appendix 2). Additional simulations 399 confirmed that parameter estimates from lme4 were highly biased, most likely because 400 with rare-outcome data the approximations used for integrating over the random 401 effects break down.

402

In order to get a quantitative understanding of how α and β diversity change across different levels of biological organisation (e.g., nestbox or site) and as a function of continuous biogeographic variables (e.g., elevation or tree diversity) we develop a framework for *focussing* repeatability metrics at the appropriate biological level (see Appendix 2). The two-way dichotomy into between-group and within-group that

408 forms the basis of standard repeatability calculations (see Nakagawa & Schielzeth, 2010 for a review) can be seen as a special case. The quantities required for these 409 410 calculations also appear in many indices developed by ecologists to quantify similarity 411 in community structure. Given this, we show how such indices can also be derived 412 directly from a GLMM which has the advantages that credible intervals can easily be 413 computed, incomplete sampling is naturally dealt with (Chao, Chazdon, Colwell, & 414 Shen, 2006) and changes in the indices as a function of differences in a continuous 415 variable (such as latitude) can be handled. The main disadvantage of the approach is 416 that between-species correlation structures may typically be richer than what a fitted 417 GLMM assumes, such that variation in community structure may be greater than the 418 model allows. However, posterior predictive checking allows model inadequacies to 419 be detected, and richer correlation structures are available, for example through a phylogeny (Hadfield & Nakagawa, 2010) or through factor analysis (Niku, Hui, 420 421 Taskinen, & Warton, 2019; Warton et al., 2015).

422

In Appendix 2 we also present methods for using model outputs to generate expectations for the taxon richness of a faecal sample and Jaccard index (often used in studies of β -diversity) that quantifies the similarity of facecal samples. This allows us to relate model coefficients back to effect sizes that are more often used in community ecology. However, as the Jaccard index captures both turnover and community nestedness (Baselga, 2010), in the results we mainly use repeatability to quantify turnover.

430

433

434 **Read quality**

The three MiSeq runs combined generated 34.04 million raw paired-end reads, of which 9.8 million were classified as COI amplicons after de-multiplexing based on the primer sequences. Amplicons for 16S and rbcL were also generated, but our diet analysis focuses only on COI. 8.9 million merged sequences passed all the quality filters. Out of these, 8.7 million sequences were retained after alignment against the reference OTU sequences.

441

442 **Diet Composition**

443 After identifying samples that tested positive for blue tit 16S DNA, excluding nonprey taxa and collapsing similar sequences, we identified 432 prey MOTUs across 444 772 faecal samples. Of these MOTUs, 57% could be matched to candidate species on 445 446 the basis of > 99% sequence identity and a voucher/reference specimen identified to species level. A further 4% were >99% matched and therefore identifiable to species 447 448 level, but lacking a reference initially identified to species level. The remainder of 449 MOTUs are not identifiable to species level but are diagnostically distinct dietary items at minimum within the order identified by the best hit. (Appendix 1 Fig. S2, 450 Table S3). The mean number of MOTUs per sample was 5.06, with mode = 3, 451 452 median = 5 and range = 0 - 20. The MOTU abundance distribution was highly rightskewed, with 42.4% recorded in only one sample and 74.3% recorded in five or 453 454 fewer samples (Fig. 1A).

456 Only 15 MOTUs were recorded in more than 50 samples (five Lepidoptera, four Hemiptera, three Diptera and one each of Collembola, Coleoptera and Hymenoptera). 457 458 Eleven of these MOTUs were identified to species level, with Argyresthia goedartella 459 (Lepidoptera: Yponomeutidae) most common (34.6% of samples, Fig. 1A inset). Most of these species are associated with resources available early in spring (Table S4), such 460 461 as catkins on birch (Betula pendula/pubescens) or alder (Alnus glutinosa) or buds of 462 birch or sycamore (Acer pseudoplatanus). We also found winter moth (Operophtera 463 brumata) in 27 samples (3.5%), the larvae of which comprise a major component of 464 nestling diet later in the spring but were not known to occur in the diet in early spring. 465

Eighteen invertebrate orders were encountered in at least one sample, with Insecta
contributing 86.1% of MOTUs. Within insects, MOTUs matched to the order
Lepidoptera were the most commonly recorded (present in 73.6% of samples, Fig. 1B)
and taxon-rich (131 taxa, Fig. 1C). Other commonly recorded orders were Hemiptera,
Diptera, Hymenoptera, Coleoptera, Araneae and Collembola.

471

472 Technical Repeatability

The value of faecal metabarcoding as a tool to infer diet depends on how reliable it proves to be and a key measure of this is repeatability. Our protocol included 30 paired replicate extractions from a different portion of the same faecal sample (although note that the sample was not homogenised prior to extraction), 29 of which remained after quality control and which we used to estimate technical repeatability (Appendix 1 Table S5G). The repeatability estimate is highly sensitive to the quantity being measured (measurand), the definition of within and between group, the reference 480 population and whether it is considered on the latent or data scale (Appendix 2). The technical repeatability of a MOTU within a faeces (with only faeces and faeces:MOTU 481 482 contributing to the between-group variance) had a posterior mode of 0.305 (95% 483 credible interval = 0.223 - 0.408) on the data (0,1) scale and 0.783 (0.712 - 0.845) on the latent (threshold) scale. Variation in MOTU richness at the sample level was 484 485 reasonable (0.325 (0.118 - 0.770)) but the richness of samples within faeces are not 486 strongly correlated and so the technical repeatability of richness for a faeces is low 487 (Data; 0.003 (0 - 0.714), Latent; 0.003 (0 - 0.676)). However, the credible intervals 488 are large, and the main analysis (see below) shows non-zero correlations between the 489 richness of faeces from the same nestbox suggesting the true technical repeatability of richness must be non-zero. 490

491

492 Dietary MOTU richness

We used a generalized linear mixed model (GLMM) with a binary (threshold) response 493 494 to examine the predictors of MOTU presence. From the main effects we can gain insights into how dietary MOTU richness (related to a-diversity) varies across time 495 496 and space. Day of year predicted a small but significant increase in dietary richness 497 over the course of the spring (b = 0.082 (0.024 - 0.135), Fig. 2C), with the expected number of MOTUs per faecal sample increasing from 1.981 to 3.933 from the first to 498 last date (Table 2). For elevation (b = -0.022 (-0.131 - 0.104)) and latitude (b = 0.058499 500 (-0.015 - 0.144)) gradients in dietary richness were non-significant (Fig. 2A-B, Table 2), as were the metrics describing among-site variation in woodland habitat (total 501 502 foliage, foliage diversity, amount of oak, amount of birch, Table S5B). The repeatability of species richness within nestboxes at a site was moderate (Data; 0.140 503

(0.041-0.264), Latent; 0.158 (0.046-0.296), Appendix 2) but we found little evidence
that richness varied among sites or among days within a year (after controlling for the
linear increase). The effect of including more than one faeces in the sample was
positive, but non-significant.

508

509 Dietary MOTU turnover

510 The probability of being present in a sample varied substantially across MOTUs 511 (variance on probit scale = 0.574 (0.475 - 0.696), Appendix 1 Table S5B). From the 512 interactions between MOTU identity and other terms we can gain insights into how 513 the probability of sampling individual MOTUs changes over time and space, providing 514 a measure of turnover and its significance. There was significant among MOTU 515 variation in the slope of presence/absence on day of year, elevation and latitude 516 (Appendix 1 Table S5B, Fig. 2D-F), with MOTU turnover more pronounced over elevation and day of year. However, the predicted repeatabilities for MOTUs in faeces 517 518 sampled at the same elevation (but at different sites) were rather low (Data; 0.002 (0.001-0.003), Latent; 0.041 (0.028-0.059)). Due to the substantial between- faeces 519 520 and between nest-box variation in MOTU presence the repeatability for the site-level 521 probability of a MOTU at the same elevation was higher (Data; 0.066 (0.041 - 0.095)), Latent; 0.148 (0.106 - 0.215)), but still modest. The effect of date was similarly low 522 and even within nestboxes the repeatability of a MOTU in faeces from the same day 523 was small (Data; 0.002 (0.001-0.004), Latent; 0.081 (0.057-0.114)). See Appendix 2 524 for further analysis of repeatabilities. As an alternative measure of how environmental 525 526 variables affect community composition we calculated the expectation for the Jaccard index and standardised Jaccard index (Appendix 2) between two sites at (i) the mean 527

and (ii) sampled extremes of latitude, elevation and day of year (Table 2). For all three
environmental variables communities are less similar (lower Jaccard index) at the
extremes than they are at the mean, but this effect is most pronounced for elevation
and day of year.

532

533 We considered among-MOTU variation in the relationship between the four 534 continuous habitat variables and probability of occurrence in four additional models 535 (Tables S5C-F). For three habitat metrics (total foliage, tree diversity and oak 536 availability) among-MOTU variation in habitat slopes was small and non-significant, 537 implying no discernible MOTU turnover along these gradients. The slope of MOTU 538 presence/absence on birch availability exhibited significant among-MOTU variation, 539 but turnover along this gradient is less than found for biogeographic and temporal gradients (Appendix 1 Table S5F, Fig. S4) indicating a weak relationship. 540

541

The variance in the MOTU identity by site effects was large (0.474 (0.394 - 0.551)), 542 revealing that even after controlling for biogeographic trends in turnover gradients 543 there is substantial MOTU turnover among sites (Table S5B). Indeed, the 544 545 biogeographic and habitat variables in aggregate only explained a small fraction of the between site variance (Data; 0.101 (0.069 - 0.142), Latent; 0.236 (0.174 -0.296), 546 Appendix 2). The total within-site (due to both biogeographic variation and random 547 548 site variation) repeatability was small if assessed at the level of faeces (Data; 0.023 (0.016 - 0.029), Latent; 0.275 (0.242 - 0.306)) but larger if assessed at the level of 549 550 nestboxes (Data; 0.270 (0.223 - 0.334), Latent; 0.568 (0.520 - 0.628)). This arises 551 because of the considerable variance amongst faeces within a nestbox. The variance

in MOTU identity by nestbox effects was comparable to the site effects (0.434 (0.376 -0.529)), but the within-nestbox repeatability at a single site was small (Data; 0.016 (0.012 - 0.022), Latent; 0.275 (0.247 - 0.312)), again because of the large betweenfaeces variance. The within-nestbox repeatability across sites (where site and nestbox effects contribute to the between group variance) was greater (Data; 0.069 (0.057 –

- 557 0.083), Latent; 0.474 (0.445 0.502)).
- 558

Interactions between MOTU and plate, plate-row and plate-column were also significant (Appendix 1 Table S5B), which may reflect within plate contamination. However, our placing of samples on the plate in the order in which samples were collected in the field (spatially and temporally structured) could also contribute to this signature if there is spatiotemporal structure in MOTU presence/absence that is not accounted for by the day of year:MOTU and site:MOTU terms.

565

566 Order level trends

Lepidoptera showed a significant increase in probability of occurrence with increasing latitude (b = 0.236 (0.044 - 0.430)) and elevation (b = 0.309 (0.073 - 0.583), Fig. 3AB, Appendix 1 Table S6). Other than Lepidoptera, only Diptera also showed a significant increase with latitude (b = 0.252 (0.058 - 0.446)). Hymenoptera showed a significant increase in probability of occurrence with increasing elevation (b = 0.319(0.061 - 0.557)), with positive trends also apparent for Diptera, Hemiptera and Coleoptera.

The probability of sampling a hemipteran increases very steeply through time over the course of the spring (b = 0.422 (0.259 - 0.590)), with significant positive relationships also apparent for Lepidoptera (b = 0.174 (0.006 - 0.341)) and Coleoptera (b = 0.269(0.113 - 0.424)) (Fig. 3C). Increasing site level tree diversity had a significant positive effect on the probability of sampling Diptera (b = 0.344 (0.095 - 0.586)) and a significant negative effect on the probability of sampling Hymenoptera (b = -0.283 (-0.528 - -0.037), Fig. 3D).

582

583

584 **Discussion**

585

We demonstrate that faecal metabarcoding can provide deep insights into the diet of a generalist woodland bird, and provide the first in-depth analysis of the natural diet of a passerine bird prior to breeding. We show that across Scottish woodlands in early spring - when overall food availability is low - blue tits are able to locate and harvest over 400 prey taxa. Further, we show strong temporal patterns in the taxonomic richness and composition of the invertebrate prey items.

592

593 **Diet Composition**

594 Our findings on blue tit diet composition broadly agree with previous work on this 595 species (Betts, 1955; J. Gibb & Betts, 1963). As for previous faecal metabarcoding 596 studies on generalist insectivores (Clare, Fraser, Braid, Fenton, & Hebert, 2009; 597 Jedlicka, Vo, & Almeida, 2016; Sedlock, Krüger, & Clare, 2014), we found most 598 dietary taxa to be rare. The six most common orders were also detected using

- 599 morphology-based identification of gizzard contents by Betts (1955). For a fuller
- discussion of the commonest taxa see the extended discussion in Appendix 1.
- 601

602 One surprise in our data was the prevalence of winter moth early in the spring. The larvae of this species are one of the main foods provisioned to nestling tits (Betts, 1955; 603 604 C. Perrins, 1991) and whilst they are the most common spring Lepidopteran larvae on 605 our transect, their availability peaks in late May/early June (Shutt, Burgess, et al., 606 2019), and so we did not anticipate finding them in the diet in March/April. A post hoc 607 analysis (GLMM with threshold response, site and nestbox effects as random and year 608 effects as fixed) revealed that the probability of occurrence in a sample increases 609 significantly in the days running up to the site-average first egg laying date (b = 0.039, CI = 0.023 - 0.055), from around a 2% chance at 30 days prior to laying to 17% at the 610 611 average site-level blue tit first egg date. This increase in the incidence of winter moth 612 in the diet most likely corresponds with a change in the availability of early instar 613 larvae, rather than eggs, which would be available throughout the period (Waring & Townsend, 2017). This finding raises the possibility that tits might use early instars of 614 615 winter moth and other foliar caterpillar larvae as a cue of when to breed.

616

617 Dietary Richness and Turnover

The biogeographic variables that we considered, latitude and elevation, had no significant effect upon dietary MOTU richness, but a significant effect upon dietary turnover. This reveals that whilst the total richness of prey eaten may not vary geographically (see also the very low site variance), the taxa comprising the diet vary along biogeographic clines (more so over elevation than latitude) and also from site to site, as revealed by the significant site by MOTU interaction component. These findings are consistent with those from faecal metabarcoding of insectivorous bats (Clare, Symondson, Broders, et al., 2014; Sedlock et al., 2014) and could indicate local dietary specialisation. However, we suspect that a more likely explanation for this apparent specialisation is that it arises from patterns in prey availability (V. Moran & Southwood, 1982) and that the birds are flexible in their prey.

629

630 The increase in dietary MOTU richness as spring progresses parallels seasonal 631 increases in the abundance and availability of herbivorous insects in European forests 632 (Bale et al., 2002; Southwood, Wint, Kennedy, & Greenwood, 2004). Whilst dietary richness generally increases during spring, some taxa become less likely to occur and 633 634 others more so, arising from the distinct phenologies of individual prey taxa (Forrest, 2016; Southwood et al., 2004). All of the main orders showed a tendency toward 635 increasing as spring progressed, though on the data scale the increase was steepest for 636 637 Hemiptera, which may be attributable to a pronounced spring phenology in the abundance of aphids on buds and leaves (Bell et al., 2015). 638

639

The habitat indices that we consider were non-significant predictors of blue tit dietary richness, and MOTU turnover along such gradients was much weaker than estimated for the biogeographic and temporal variables. One potential explanation for our low estimate of turnover along such habitat gradients is that most invertebrate prey species may not be entirely restricted to a particular tree species. Alternatively, perhaps our 'territory' based habitat metrics are inadequate measures of the availability of different tree species to each bird at this time. At face value our results are consistent with the 647 greater importance of larger-scale geographic clines (i.e. latitude, elevation) as 648 determinants of prey presence/absence, presumably because they act as a proxy for 649 other environmental variables that limit invertebrate distributions, such as temperature. 650 However, substantial spatial turnover remained even after controlling for 651 spatiotemporal gradients, which suggests that there are important drivers of prey 652 turnover that we have overlooked.

653

654 Model based inference of richness and turnover

655 Describing and explaning temporal and geographical variation in components of 656 diversity is a mainstay of community ecology (Dornelas et al., 2014; Li et al., 2018; 657 Magurran, 2013). α -diversity can be calculated for the sampled community scale (be 658 that a location or point in time), which has made its statistical analysis relatively straightforward. In comparison, β -diversity is often calculated as a pairwise 659 similarity between communities (Koleff, Gaston, & Lennon, 2003), and where 660 661 multiple communities are considered the non-independence of comparisons presents a challenge to statistical inference (Baselga, 2010). In an important development 662 663 Baeten et al. (2014) explained how a generalized linear model with taxon 664 presence/absence as a binomial response could be used to estimate changes in richness and turnover between points and crucially determine statistical significance. 665 Here we have extended their framework to a generalized linear mixed model and we 666 show that the interaction of taxon (MOTU) with categorical (random intercepts) and 667 668 continuous (random slopes) variables estimates turnover between points (in space or 669 time) and along gradients, respectively. We also show that it is possible to predict the Jaccard index (measure of β -diversity) between a pair of communities sampled at 670

671 points in space or time as a measure of effect size (Appendix 2). The principal benefits of this new model-based approach over existing pair-wise approaches are 672 673 that (i) it allows estimation of confidence intervals and p values for turnover and 674 richness along gradients and among samples without such calculations being complicated by non-independence; (ii) hierarchical structure in the sampling can be 675 676 included, and turnover can be assessed at each level explicitly taking into account 677 heterogeneity in sampling effort at lower levels; (iii) multiple covariates can be 678 included; (iv) inferences can be made including or excluding a control for taxon 679 abundance and (v) model based inference of repeatability is possible (see Appendix 680 2). The model coefficients can also be used to derive predictions of the total number 681 of taxa in a community and the Jaccard index (or alternative β -diversity metric) between communities. Our model is defined in the context of the probability of a 682 taxon being present in a faecal sample, and as the number of samples (n) increases 683 total taxon richness is predicted to increase monotonically (with a decelerating 684 685 slope), such that when $n = \infty$, every taxon will be present. There are similarities between this curve and rarefaction curves that are often used to standardise for 686 687 heterogeneity in sampling in ecology (Gotelli & Colwell, 2011), with both methods 688 requiring inference of the probability of each taxon being in a sample. In addition, the Jaccard index will increase monotonically and with an accelerating function with 689 increasing species richness (Appendix 2) and monotonically and with a decelerating 690 691 function with sampling effort. Given that community level diversity metrics are highly sensitive to the choice of n, we suggest that when using our framework an n =692 693 1 represents the most natural level at which to report community-level metrics (see Appendix 2) and requires no extrapolation. 694

696 A limitation of our approach is that by imposing a parametric correlation structure on 697 the data, that correlation structure is relatively simple and probably doesn't catch the 698 full complexity of species associations. For example, if there was a patchily distributed species of herb on which three prey taxa were specialised on, then these 699 700 three species would co-occur with higher probability than our model would suggest. 701 Rectifying these problem would require a) identifying the herb that generates these 702 correlations, measuring its prevalence and incorporating that data into the model b) 703 use more complex correlation structures to be modelled in situations where the 704 number of taxa is large (Runcie & Mukherjee, 2013; Warton et al., 2015) or c) 705 develop sandwich type estimators (Huber, 1967; Zeger, Liang, & Albert, 1988) that 706 would allow robust inferences to be made even when unmodelled correlations exist. 707

708 Methodological Considerations

709 In this study we have demonstrated that faecal metabarcoding can provide a robust and 710 powerful method for assessing passerine diet, allowing greater sample sizes and 711 taxonomic resolution than direct assessment (Betts, 1955). Inclusion of positive and 712 negative controls and repeat samples are part of the standard laboratory practice (Alberdi et al., 2018) – though few previous metabarcoding studies have included any 713 714 of these (but see De Barba et al., 2014; Jedlicka et al., 2016) - and have proven 715 invaluable in informing this work. Our protocol yielded fourteen MOTUs for the 716 positive control taxon, suggesting that the 2% divergence rule of thumb used in early 717 barcoding studies to group conspecific COI barcode sequences in Metazoa (Hebert, 718 Cywinska, & Ball, 2003 and http://www.barcodinglife.com) is likely to produce 719 spurious taxa, potentially misleading naïve analyses and underlining the necessity for subsequent quality control steps. Negative controls (extraction and PCR) allowed us 720 to identify a case of systematic contamination and also informed our cut-off number 721 722 of reads (but see Deagle et al., 2018 for a critique of thresholds). After strict removal of samples that appeared likely to have been affected by systematic contamination, 723 724 some residual contamination on plates was evident and we were able to control for this 725 to some degree by including row:MOTU, column:MOTU and plate:MOTU as random 726 terms. We recommend that future studies adopt the plate:MOTU random term and 727 randomise samples across plates, such that samples from a single year, site or time of 728 year do not all appear on one plate. Although the maximum number of taxa in a sample was high (n = 20), PCR competition and the methodological maximum reads per 729 730 metabarcoding plate presumably place a limit on detecting very rare dietary items. 731 Reducing the number of target loci (three in this study, see methods) or level of multiplexing (i.e. the number of samples per sequencing run) could increase the reads 732 733 available per locus per sample and increase detectability. However, reducing multiplexing may come at an increased financial cost for sequencing. 734

735

From our repeat samples we were able to estimate technical repeatability and several measures of biological repeatability (Appendix 2). Repeatability of MOTU presence/absence was rather low, consistent with low repeatability estimates found by another study that subsampled avian faecal samples (Jedlicka et al., 2016). An implication is that if the focus of an avian faecal metabarcoding study is on the detection of the presence/absence of a specific taxon, then multiple repeat DNA extractions, amplifications and metabarcoding runs are advisable. Homogenisation of faecal samples prior to DNA extraction may increase both the ability to detect a
particular taxon and repeatability given the possible heterogeneity within single faeces.

746 Conclusion

Using a metabarcoding approach, we reveal the diet of a generalist passerine at a finer resolution than any previous study and quantify dietary richness and turnover across space and time. At the scale of our study, blue tit dietary richness increases as spring progresses, but is unaffected by latitude, elevation and habitat, whilst dietary turnover is most pronounced over temporal (day of year) and elevational gradients.

752

753

754 Acknowledgements

755

756 The authors thank Irene Benedicto Cabello and Ed Ivimey-Cook for assistance in the

757 field, Mark Blaxter for molecular advice, two anonymous reviewers for comments on

the ms and Andrés Baselga for discussion of beta-diversity. We are indebted to the

- 1759 landowners who allowed us access to their land. JDS was funded by a NERC
- doctoral training studentship (NE/1338530), ABP by a NERC Advanced fellowship
- 761 (NE/I020598/1) and JDH by a Royal Society URF.
- 762

763

764 **References**

Alberdi, A., Aizpurua, O., Gilbert, M. T. P., & Bohmann, K. (2018). Scrutinizing key steps
for reliable metabarcoding of environmental samples. *Methods in Ecology and Evolution*, 9(1), 134-147.

768	Baeten, L., Warton, D. I., Van Calster, H., De Frenne, P., Verstraeten, G., Bonte, D.,
769	Eriksson, O. (2014). A model-based approach to studying changes in compositional
770	heterogeneity. Methods in Ecology and Evolution, 5(2), 156-164.
771	Bale, J. S., Masters, G. J., Hodkinson, I. D., Awmack, C., Bezemer, T. M., Brown, V. K.,
772	Farrar, J. (2002). Herbivory in global climate change research: direct effects of rising
773	temperature on insect herbivores. <i>Global change biology</i> , 8(1), 1-16.
774	Baselga, A. (2008). Determinants of species richness, endemism and turnover in European
775	longhorn beetles. <i>Ecography</i> . 31(2), 263-271.
776	Baselga, A. (2010). Partitioning the turnover and nestedness components of beta diversity.
777	Global ecology and biogeography, 19(1), 134-143.
778	Bates D. Maechler, M. & Bolker, B. (2012) Ime4: Linear mixed-effects models using S4
779	classes
780	Beck, J. McCain, C. M., Axmacher, J. C., Ashton, L. A., Bärtschi, F., Brehm, G., Fiedler,
781	K (2017) Elevational species richness gradients in a hyperdiverse insect taxon: a
782	global meta-study on geometrid moths Global ecology and biogeography 26(4)
783	
784	Bell I R Alderson I Izera D Kruger T Parker S Pickup I Harrington R (2015)
785	Long term phenological trends species accumulation rates aphid traits and climate:
705	five decades of change in migrating onlyide. Journal of Animal Ecology, 84(1), 21, 24
/00 797	Potte M M (1055) The feed of titmice in only woodland. The Journal of Animal Feelers
101	Deus, M. M. (1955). The food of thinke in oak woodiand. The Journal of Animal Ecology,
700	202-525. Debugun V. Manadiam A. Naar C. I. Desmussion M. Zaela M. D. Clana E
709	M. T. D. (2011). Molecular dist analysis of two A frican free toiled hets (Molecular)
790	W. 1. F. (2011). Molecular diet analysis of two African free-tailed dats (Molossidae) using high throughput sequencing $PLoS One 6(6)$, $o21441$
791	using high unoughput sequencing. $FLos One, o(0), e21441$.
792	Diooks, M. E., Klisteliseli, K., Vali Delitielii, K. J., Magilussoli, A., Berg, C. W., Nielseli, A.,
793	for zero infloted concretized linear mixed modeling. The D journal 0(2), 278,400
794	Durger C. Delskii E. Esve T. Leeksenen T. Mägi M. Märd P
795	Durger, C., Derskii, E., Leva, T., Laaksonen, T., Magi, M., Manu, K., Doui, C. (2012).
790	contrate change, bleeding date and nesting diet. Now temperature differentiany affects
709	seasonal changes in pied hycatcher diet depending on nabhat variation. <i>Journal of</i>
790	Animal Ecology, $01(4)$, $920-950$. doi: $10.1111/J.1505-2050.2012.01908.x$
/99	Chao, A., Chazdon, K. L., Colwell, K. K., & Shen, I. J. (2006). Abundance - based similarity
800	indices and their estimation when there are unseen species in samples. <i>Biometrics</i> , $(2(2), 2(1, 27))$
801	02(2), 301-3/1.
802	Charmantier, A., Miccleery, K. H., Cole, L. K., Perrins, C., Kruuk, L. E. B., & Sneidon, B. C.
803	(2008). Adaptive Phenotypic Plasticity in Response to Climate Change in a wild Bird
804	Population. Science, $320(5877)$, $800-803$. doi:10.1126/science.1157174
805	Cholewa, M., & wesolowski, I. (2011). Nestling food of European hole-nesting passerines:
806	do we know enough to test the adaptive hypotheses on breeding seasons? Acta
807	Unithologica, 40(2), 105-116.
808	Clare, E. L. (2014). Molecular detection of trophic interactions: emerging trends, distinct
809	advantages, significant considerations and conservation applications. Evolutionary
810	applications, /(9), 1144-115/.
811 912	Clare, E. L., Fraser, E. E., Braid, H. E., Fenton, M. B., & Hebert, P. D. (2009). Species on the
812 912	menu of a generalist predator, the eastern red dat (Lasturus borealis): using a
813	Clara E. L. Symon day, W. O. Dradaw, H. Eshieval, E. Erssen, E. E. Madkauzia, A.
014 015	Uare, E. L., Symondson, W. U., Broders, H., Fabianek, F., Fraser, E. E., MacKenzie, A.,
813 916	Martinez- Nunez, F. (2014). The diet of Myotis lucifugus across Canada: assessing
810 917	Ioraging quality and diet variability. <i>Molecular ecology</i> , $23(15)$, $3618-3632$.
ð1/	Clare, E. L., Symondson, W. O., & Fenton, M. B. (2014). An inordinate fondness for beetles?
818	variation in seasonal dietary preferences of night-roosting big brown bats (Eptesicus
819	1uscus). <i>Molecular ecology</i> , 23(15), 3633-3647.

820 Cramp, S., & Perrins, C. M. (1993). Handbook of the Birds of Europe the Middle East and 821 North America. Oxford. UK: OUP. 822 Crisol-Martínez, E., Moreno-Moyano, L. T., Wormington, K. R., Brown, P. H., & Stanley, D. 823 (2016). Using next-generation sequencing to contrast the diet and explore pest-824 reduction services of sympatric bird species in macadamia orchards in Australia. PLoS 825 One, 11(3), e0150159. De Barba, M., Miquel, C., Boyer, F., Mercier, C., Rioux, D., Coissac, E., & Taberlet, P. (2014). 826 827 DNA metabarcoding multiplexing and validation of data accuracy for diet assessment: application to omnivorous diet. *Molecular Ecology Resources*, 14(2), 306-323. 828 Deagle, B. E., Thomas, A. C., McInnes, J. C., Clarke, L. J., Vesterinen, E. J., Clare, E. L., . . . 829 830 Eveson, J. P. (2018). Counting with DNA in metabarcoding studies: how should we convert sequence reads to dietary data? Molecular ecology. 831 Dornelas, M., Gotelli, N. J., McGill, B., Shimadzu, H., Moyes, F., Sievers, C., & Magurran, 832 833 A. E. (2014). Assemblage time series reveal biodiversity change but not systematic 834 loss. Science, 344(6181), 296-299. 835 Durant, J. M., Hjermann, D. Ø., Anker-Nilssen, T., Beaugrand, G., Mysterud, A., Pettorelli, 836 N., & Stenseth, N. C. (2005). Timing and abundance as key mechanisms affecting 837 trophic interactions in variable environments. Ecology Letters, 8(9), 952-958. 838 Edgar, R. C. (2013). UPARSE: highly accurate OTU sequences from microbial amplicon 839 reads. Nature methods, 10(10), 996. 840 Forrest, J. R. (2016). Complex responses of insect phenology to climate change. Current 841 opinion in insect science, 17, 49-54. 842 Gibb, J., & Betts, M. M. (1963). Food and food supply of nestling tits (Paridae) in Breckland 843 pine. The Journal of Animal Ecology, 489-533. 844 Gibb, J. A. (1954). Feeding ecology of tits, with notes on treecreeper and goldcrest. *Ibis*, 96(4), 845 513-543. Gotelli, N. J., & Colwell, R. K. (2011). Estimating species richness. Biological diversity: 846 847 frontiers in measurement and assessment, 12, 39-54. Hadfield, J. D. (2010). MCMC Methods for Multi-Response Generalized Linear Mixed 848 849 Models: The MCMCglmm R Package. Journal of Statistical Software, 33, 1-22. 850 Hadfield, J. D., & Nakagawa, S. (2010). General quantitative genetic methods for comparative 851 biology: phylogenies, taxonomies and multi-trait models for continuous and 852 categorical characters. Journal of evolutionary biology, 23, 494-508. 853 Hebert, P. D., Cywinska, A., & Ball, S. L. (2003). Biological identifications through DNA 854 barcodes. Proceedings of the Royal Society of London B: Biological Sciences, 270(1512), 313-321. 855 Huber, P. J. (1967). The behavior of maximum likelihood estimates under nonstandard 856 857 conditions. Proceedings of the fifth Berkelev symposium on mathematical statistics 858 and probability, I(1), 221-233. 859 Jedlicka, J. A., Sharma, A. M., & Almeida, R. P. (2013). Molecular tools reveal diets of 860 insectivorous birds from predator fecal matter. Conservation Genetics Resources, 861 5(3), 879-885. 862 Jedlicka, J. A., Vo, A.-T. E., & Almeida, R. P. (2016). Molecular scatology and high-863 throughput sequencing reveal predominately herbivorous insects in the diets of adult 864 and nestling Western Bluebirds (Sialia mexicana) in California vineyards. The Auk, 865 134(1), 116-127. Kennedy, C., & Southwood, T. (1984). The number of species of insects associated with 866 British trees: a re-analysis. The Journal of Animal Ecology, 455-478. 867 868 King, R., Symondson, W., & Thomas, R. (2015). Molecular analysis of faecal samples from birds to identify potential crop pests and useful biocontrol agents in natural areas. 869 870 Bulletin of entomological research, 105(3), 261-272. 871 Koleff, P., Gaston, K. J., & Lennon, J. J. (2003). Measuring beta diversity for presence-872 absence data. Journal of Animal Ecology, 72(3), 367-382.

873	Kress, W. J., García-Robledo, C., Uriarte, M., & Erickson, D. L. (2015). DNA barcodes for
874	ecology, evolution, and conservation. Trends in Ecology & Evolution, 30(1), 25-35.
875	Li, Y., Evans, N. T., Renshaw, M. A., Jerde, C. L., Olds, B. P., Shogren, A. J., Pfrender,
876	M. E. (2018). Estimating fish alpha-and beta-diversity along a small stream with
877	environmental DNA metabarcoding. <i>Metabarcoding and Metagenomics</i> , 2, e24262.
878	Magurran A E (2013) Measuring hiological diversity: John Wiley & Sons
879	Mata V A Rebelo H Amorim F McCracken G F Jarman S & Beia P (2019) How
880	much is enough? Effects of technical and biological replication on metabarcoding
881	dietary analysis Molecular acology 28(2) 165 175
887	Moran A I Prosser S W & Moran I A (2010) DNA metabarooding allows non invasive
002	Morall, A. J., Flossel, S. W., & Morall, J. A. (2019). DNA includincoding allows non-invasive
003	(Schembergersfer) Deep L 7 (50)
884	(Selasphorus rufus). PeerJ, $/$, e0596.
885	Moran, V., & Southwood, I. (1982). The guild composition of arthropod communities in trees.
886	The Journal of Animal Ecology, 289-306.
887	Murakami, M., Ichie, T., & Hirao, T. (2008). Beta-diversity of lepidopteran larval
888	communities in a Japanese temperate forest: effects of phenology and tree species.
889	Ecological research, 23(1), 179-187.
890	Nakagawa, S., & Schielzeth, H. (2010). Repeatability for Gaussian and non-Gaussian data: a
891	practical guide for biologists. <i>Biological Reviews</i> , 85(4), 935-956.
892	Nichols, R. V., Åkesson, M., & Kjellander, P. (2016). Diet assessment based on rumen
893	contents: A comparison between DNA metabarcoding and macroscopy. PLoS One,
894	11(6).
895	Niku, J., Hui, F. K., Taskinen, S., & Warton, D. I. (2019). gllvm: Fast analysis of multivariate
896	abundance data with generalized linear latent variable models in r. Methods in Ecology
897	and Evolution, 10(12), 2173-2182.
898	Novotny, V., & Weiblen, G. D. (2005). From communities to continents: beta diversity of
899	herbivorous insects. Paper presented at the Annales Zoologici Fennici
900	Oehm I Juen A Nagiller K Neuhauser S & Traugott M (2011) Molecular scatology:
901	how to improve prev DNA detection success in avian faces? <i>Molecular ecology</i>
902	resources 11(4) 620-628
902	Perring C (1991) Tits and their caternillar food supply <i>Ibis</i> $133(s1)$ 49-54
00/	Perrins, C. M. (1971). This and then eacephian rood supply. <i>Tots</i> , 155(51), 47-54.
005	Dompanon E. Deagle P. E. Symondson W. O. Brown D. S. Jarman S. N. & Taberlet P.
905	(2012) Who is pating what dist apparent using next parametical apparent.
900	(2012). Who is eating what, the assessment using next generation sequencing. Malaxier analysis $2l(8)$, 1021-1050
907	Molecular ecology, 21(8), 1951-1950.
908	Quemere, E., Hiberi, F., Miquei, C., Lhuiller, E., Kasolondraide, E., Champeau, J.,
909	Gautier, L. (2013). A DNA metabarcooling study of a primate dietary diversity and
910	plasticity across its entire fragmented range. <i>PLoS One</i> , 8(3), e589/1.
911	Razgour, O., Clare, E. L., Zeale, M. R., Hanmer, J., Schnell, I. B., Rasmussen, M., Jones,
912	G. (2011). High-throughput sequencing offers insight into mechanisms of resource
913	partitioning in cryptic bat species. Ecology and Evolution, $1(4)$, 556-570.
914	Runcie, D. E., & Mukherjee, S. (2013). Dissecting high-dimensional phenotypes with
915	Bayesian sparse factor analysis of genetic covariance matrices. Genetics, 194(3), 753-
916	767.
917	Rytkönen, S., Vesterinen, E. J., Westerduin, C., Leviäkangas, T., Vatka, E., Mutanen, M.,
918	Orell, M. (2019). From feces to data: A metabarcoding method for analyzing
919	consumed and available prey in a bird-insect food web. <i>Ecology and evolution</i> , 9(1),
920	631-639.
921	Samplonius, J. M., Kappers, E. F., Brands, S., & Both, C. (2016). Phenological mismatch and
922	ontogenetic diet shifts interactively affect offspring condition in a passerine. <i>Journal</i>
923	of Animal Ecology, 85(5), 1255-1264.
-	\mathcal{J}

924	Sedlock, J. L., Krüger, F., & Clare, E. L. (2014). Island bat diets: does it matter more who you
925	are or where you live? Molecular ecology, 23(15), 3684-3694.
926	Shutt, J. D., Bolton, M., Benedicto Cabello, I., Burgess, M. D., & Phillimore, A. B. (2018).
927	The effects of woodland habitat and biogeography on blue tit Cyanistes caeruleus
928	territory occupancy and productivity along a 220 km transect. Ecography, 41, 1967-
929	1978.
930	Shutt, J. D., Burgess, M. D., & Phillimore, A. B. (2019). A spatial perspective on the
931	phenological distribution of the spring woodland caterpillar peak. American
932	Naturalist, 194. E000.
933	Shutt, J. D., Nicholls, J. A., Trivedi, U. H., Burgess, M. D., Stone, G. N., Hadfield, J. D., &
934	Phillimore, A. B. (2019). MOTU presence absence data. Retrieved from:
935	https://doi.org/10.5061/dryad.hhmggnkd3
936	Southwood, T. R. E., Wint, G. W., Kennedy, C. E., & Greenwood, S. R. (2004). Seasonality
937	abundance, species richness and specificity of the phytophagous guild of insects on
938	oak (Ouercus) canopies. European Journal of Entomology. 101(1), 43-50.
939	Sterling, P., & Parsons, M. (2012). Field guide to the micro-moths of Great Britain and
940	Ireland: British Wildlife Publishing
941	Sullins, D. S., Haukos, D. A., Craine, J. M., Lautenbach, J. M., Robinson, S. G., Lautenbach,
942	L D Sandercock B K (2018) Identifying the diet of a declining prairie grouse
943	using DNA metabarcoding <i>The Auk</i> : Ornithological Advances 135(3) 583-608
944	Symondson W (2002) Molecular identification of prev in predator diets. <i>Molecular ecology</i>
945	11(4) 627-641
946	Taberlet P Coissac E Pompanon F Brochmann C & Willersley E (2012) Towards
947	next-generation biodiversity assessment using DNA metabarcoding Molecular
0/8	acology 21(8) 2015 2050
0/0	Thomsen D F. Jargensen D S. Bruun H H. Dedersen J. Bijs-Nielsen T. Jonko K.
050	Karsholt O (2016) Desource specialists lead local insect community turnover
950	Kaisholi, O. (2010). Resource specialists lead local insect community turnover
951	associated with temperature–analysis of an 18- year full- seasonal record of moths and
952	There K. Tetter A. D. Willeman M. Klenner D. H. Steen there D. Vere M. I.
933	Inorup, K., Iøurup, A. P., Willemoes, M., Klaassen, K. H., Strandberg, K., Vega, M. L.,
954	kanbek, C. (2017). Resource tracking within and across continents in long-distance
933	Travelling D. K. Lette, C. Marshell, L. C. Nuttle, T. & Darten, D. A. (2016). Melecular
930	Irevenine, B. K., Laua, S. C., Marshall, L. C., Nutter, I., & Porter, B. A. (2010). Molecular
957	Waterthrush (Derlagio motocillo) The Auto 122(2) A15 A28
938	Travelling D. K. Nuttle T. Heggin D. D. Drawwer, N. L. Darten D. A. & Lette S. C.
939	(2019) DNA metal-and diag of nextline frace reveals meridian of a metal and and
900	(2018). DNA metabarcoding of nesting feces reveals provisioning of aquatic prey and
961	resource partitioning among Neotropical migratory songbirds in a riparian nabitat.
962	Uecologia, 18/(1), 85-98.
963	I revelline, B. K., Nuttle, I., Porter, B. A., Brouwer, N. L., Hoenig, B. D., Steffensmeier, Z.
964	D., & Latta, S. C. (2018). Stream acidification and reduced aquatic prey availability
965	are associated with dietary shifts in an obligate riparian Neotropical migratory
966	songbird. PeerJ, b , $e > 141$.
96/	Vo, A. I., & Jedlicka, J. (2014). Protocols for metagenomic DNA extraction and Illumina
968	amplicon library preparation for faecal and swab samples. <i>Molecular ecology</i>
969	resources, 14(6), 1183-119/.
9/0	waring, P., & Townsend, M. (2017). Field guide to the moths of Great Britain and Ireland:
9/1	BIOOMSbury Publishing.
972	warton, D. I., Blanchet, F. G., O'Hara, R. B., Ovaskainen, O., Taskinen, S., Walker, S. C., &
973	Hui, F. K. (2015). So many variables: joint modeling in community ecology. <i>Trends</i>
974	in Ecology & Evolution, $30(12)$, $766-779$.
11/15	-3370, 44 , 1 , 11 , (1177) , 12 , 1 , 1 , 1 , 1 , 1 , 1 , 1 , 1

975 Whittaker, R. H. (1972). Evolution and measurement of species diversity. *Taxon*, 213-251.

- Yu, D. W., Ji, Y., Emerson, B. C., Wang, X., Ye, C., Yang, C., & Ding, Z. (2012). Biodiversity
 soup: metabarcoding of arthropods for rapid biodiversity assessment and
- 977 soup: metabarcoding of artifiopods for rapid biodiversity assessment an 978 biomonitoring. *Methods in Ecology and Evolution, 3*(4), 613-623.
- 279 Zeger, S. L., Liang, K.-Y., & Albert, P. S. (1988). Models for longitudinal data: a generalized
 980 estimating equation approach. *Biometrics*, 1049-1060.

982 Data Accessibility Statement

- 983 The MOTU presence/absence data for COI are available
- 984 from https://doi.org/10.5061/dryad.hhmgqnkd3. The data and model outputs used in
- 985 appendix 2 are available from
- 986 <u>https://github.com/allyphillimore/faecalmetabarcoding-adults</u>
- 987
- 988

989 Author contributions

JDS, JAN, ABP and JDH were the main contributors to study conceptualization and methodology, with JDS and ABP responsible for fieldwork, JAN responsible for designing and conducting the molecular work and JDH designing the statistical methods, developing the theory and writing Appendix 2. JDS and UHT contributed to data curation. Statistical analysis was conducted by JDS, ABP and JDH. ABP was responsible for project administration and ABP and JDH for funding acquisition. JDS wrote the original draft, and all authors contributed to further writing and editing.

997

Tables and Figures

Table 1. Sampling and laboratory protocols employed by published faecal barcoding
studies focusing on the invertebrate component of diet. An entry of 'None' means that
while steps may have been taken in the study, no specific method was detailed.

Number of study species (most common species)	Total number of samples (maximum number of samples per species)	Number of sites (region)	Controls	Measures taken to assess repeatability	Reference
l (Lesser Prairie- Chicken)	314	4 (Kansas and Colorado, USA)	None	None	(Sullins et al., 2018)
l (Western Bluebird)	210	3 (neighbouring vineyards, California, USA)	None	Ten faeces subsampled.	(Jedlicka et al., 2016)
3 (Wood Thrush)	137 (51)	1 (Pennsylvania , USA)	PCR negatives and positives	None	(Trevelline , Nuttle, Hoenig, et al., 2018)
1 (Louisiana Waterthrush)	130	2 (Arkansas and Pennsylvania, USA)	None	None	(Trevelline , Latta, Marshall, Nuttle, & Porter, 2016)
1 (Louisiana Waterthrush)	92	3 (headwater streams, Pennsylvania, USA)	None	None	(Trevelline , Nuttle, Porter, et al., 2018)
(Rufous hummingbird)	30	1 (Vancouver Island, Canada)	1 x extraction negative	None	(A. J. Moran, Prosser, & Moran, 2019)
13 (Lewin's Honeyeater)	82 (29)	1 (Bundaberg, Australia)	Extraction negatives	PCR run twice to test amplification repeatability	(Crisol- Martínez, Moreno- Moyano, Wormingt on, Brown, & Stanley, 2016)

l (Western Bluebird)*	16	2 (neighbouring vineyards, California, USA)	None	Faecal sample was subdivided and run on two extraction kits.	(Jedlicka et al., 2013)
4 (Blue tit, Great Tit, Willow Tit)	14 (4)	2 (Oulu and Kuusamo, Finland)	Extraction negative	None	(Rytkönen et al., 2019)
3 (Sedge Warbler)‡	11 (6)	3 (South Wales, UK)	None	None	(King, Symondso n, & Thomas, 2015)

1005 ‡ Study employed Sanger sequencing rather than metabarcoding.

Table 2. Expectations for the MOTU richness of - and Jaccard indices between samples of communities at (i) the same and (ii) extreme points along latitude,
elevation and day of year gradients. Expectations are calculated for a random sample,
nestbox, day and site averaging over variation in other predictor variables (for further
details see Appendix 2). Expectations were generated for 2014 and a single faecal
sample.

1013

Predictor	Sampling	MOTU	MOTU	MOTU	Jaccard	Standardised
	position	richness at	richness at	richness at	index	Jaccard index ¹
		mean	minimum	maximum		
Latitude	Mean	2.339			0.011	5.255 (4.238 - 6 296)
		(1.665 -			0.015)	0.290)
		3.271)				
Latitude	Extremes		2.607 (1.476 - 3.831)	2.213 (1.266 - 3.444)	0.009 (0.006 - 0.012)	3.569 (2.776 - 4.423)
Elevation	Mean	2.340			0.012)	5.966 (4.814 -
		(1.669 -			0.017)	7.179)
		3.277)				
Elevation	Extremes		2.435 (1.524 - 3.647)	2.138 (1.084 - 3.700)	0.008 (0.005 - 0.011)	2.344 (1.727 - 3.047)
Day of	Mean	2.464			0.013	5.922 (4.811 -
year		(1.655 - 3.252)			(0.008 - 0.017)	7.103)
Day of	Extremes		1.981 (1.352 - 2.848)	3.933 (2.459 - 5.603)	0.007 (0.005 -	1.973 (1.431 - 2.592)
year					0.010)	

1014

1015 ¹ The standardised Jaccard index is the ratio of the observed index to that expected if the same number 1016 of species were sampled at random from two communities (see Appendix 2). It will tend to be > 1 as 1017 common/widespread species will be over-represented in both communities. The expectation for the 1018 Jaccard index and standardised index for two samples taken entirely at random from the transect is

- 0.01 (0.007 0.014) and 4.727 (3.891 5.732), respectively, and these values can be taken as a
- 1020 baseline that captures the effect of common/widespread species on measures of community similarity.



1021

1022 Fig. 1 A Histogram of the abundance distribution of prey MOTUs. Inset details the 1023 most prevalent MOTUs identified to species level (those recorded in more than 50 1024 samples), with the number of samples they were recorded in. B Relative abundance of prey orders in the spring diet of blue tits. C Number of MOTUs within prey orders 1025 (families comprising > 10 MOTUs are highlighted individually within their respective 1026 1027 orders). In **B** and **C** orders within Insecta (left) are split from orders within other classes 1028 (right). Images are used to indicate taxonomic order rather than the life-stage or species 1029 that is preyed upon.



1032Fig. 2 Dietary richness (A - C) and turnover (D - F) along latitudinal (A, D),1033elevational (B, E) and temporal (C, F) gradients. In A - C the solid black lines indicates1034the model prediction of dietary MOTU occurrence (related to richness), with the solid1035orange area illustrating the 95% credible intervals in the slope. In D - F the green lines1036correspond to the 95% upper and lower bounds of the estimated distribution of among-1037MOTU slopes. The wider the difference between the upper and lower line the greater1038the turnover along the gradient. Predictions are made from the core model (Table S4B).



Fig. 3 Model predictions for the occurence of six prey orders across A. latitude, B.
elevation, C. day of year and D. tree diversity. Predictions are made based on the
intercept of the model reported in Table S5.