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1	Environmental DNA (eDNA) metabarcoding of pond water as a tool to
2	survey conservation and management priority mammals
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23	

24 Abstract

25

26 Environmental DNA (eDNA) metabarcoding can identify terrestrial taxa utilising aquatic habitats 27 alongside aquatic communities, but terrestrial species' eDNA dynamics are understudied. We evaluated eDNA metabarcoding for monitoring semi-aquatic and terrestrial mammals, 28 29 specifically nine species of conservation or management concern, and examined 30 spatiotemporal variation in mammal eDNA signals. We hypothesised eDNA signals would be 31 stronger for semi-aquatic than terrestrial mammals, and at sites where individuals exhibited 32 behaviours. In captivity, we sampled waterbodies at points where behaviours were observed 33 ('directed' sampling) and at equidistant intervals along the shoreline ('stratified' sampling). We 34 surveyed natural ponds (N = 6) where focal species were present using stratified water 35 sampling, camera traps, and field signs. eDNA samples were metabarcoded using vertebrate-36 specific primers. All focal species were detected in captivity. eDNA signal strength did not differ 37 between directed and stratified samples across or within species, between semi-aquatic or 38 terrestrial species, or according to behaviours. eDNA was evenly distributed in artificial 39 waterbodies, but unevenly distributed in natural ponds. Survey methods deployed at natural 40 ponds shared three species detections. Metabarcoding missed badger and red fox recorded by 41 cameras and field signs, but detected small mammals these tools overlooked, e.g. water vole. 42 Terrestrial mammal eDNA signals were weaker and detected less frequently than semi-aquatic 43 mammal eDNA signals. eDNA metabarcoding could enhance mammal monitoring through 44 large-scale, multi-species distribution assessment for priority and difficult to survey species, and provide early indication of range expansions or contractions. However, eDNA surveys need high 45

46 spatiotemporal resolution and metabarcoding biases require further investigation before47 routine implementation.

- Key-words: camera traps, field signs, lentic, monitoring, semi-aquatic mammals, terrestrial
 mammals
- 51

52 1. Introduction

53

54 Mammals are a highly threatened taxon, with 25% of species at risk of extinction globally due 55 to harvesting, habitat degradation/loss, non-native species or perception as pests (Visconti et 56 al., 2011). Most species lack long-term, systematic monitoring, with survey efforts biased 57 towards rare species (Massimino, Harris, & Gillings, 2018). Data deficiency prevents robust 58 estimation of mammalian range expansions/declines and population trends (Bland, Collen, 59 Orme, & Bielby, 2015). Therefore, effective and evidence-based strategies for mammal 60 conservation and management are urgently needed (Mathews et al., 2018).

61 Many mammals are nocturnal and elusive thus monitoring requires non-invasive, 62 observational methods such as camera traps and field signs, e.g. footprints, scat (Caravaggi et 63 al., 2018; Harris & Yalden, 2004; Kinoshita et al., 2019; Sadlier, Webbon, Baker, & Harris, 2004). Camera trapping is cost-efficient, standardised, reproducible, and produces data suited to site 64 65 occupancy modelling, but only surveys a fraction of large, heterogeneous landscapes. Trap 66 placement can substantially influence species detection probabilities, and traps often miss 67 small species (Burton et al., 2015; Caravaggi et al., 2018; Ishige et al., 2017; Leempoel, Hebert, 68 & Hadly, 2019). Field sign surveys are inexpensive, but resource-intensive for broad geographic 69 coverage (Kinoshita et al., 2019; Sadlier et al., 2004). Species can have similar footprints and 70 scat, increasing the potential for misidentification (Franklin et al., 2019; Harris & Yalden, 2004). 71 Mammal survey methods can be species-specific, thus multiple methods are necessary for 72 large-scale, multi-species monitoring schemes (Massimino et al., 2018; Sales et al., 2019).

73

Environmental DNA (eDNA) analysis is a recognised tool for rapid, non-invasive, cost-

74 efficient biodiversity assessment across aquatic and terrestrial ecosystems (Deiner et al., 2017). 75 Organisms transfer genetic material to their environment via secretions, excretions, gametes, 76 blood, or decomposition, which can be isolated from environmental samples (Thomsen & 77 Willerslev, 2015). Studies using eDNA analysis to target specific semi-aquatic and terrestrial 78 mammals have employed PCR or quantitative PCR (qPCR) (e.g. Franklin et al., 2019; Lugg, 79 Griffiths, van Rooyen, Weeks, & Tingley, 2017; Rodgers & Mock, 2015; Thomsen et al., 2012; 80 Williams, Huyvaert, Vercauteren, Davis, & Piaggio, 2018). eDNA metabarcoding can screen 81 entire communities using PCR combined with high-throughput sequencing (Deiner et al., 2017; 82 Thomsen & Willerslev, 2015), but mammalian assessments are uncommon (Klymus, Richter, 83 Thompson, & Hinck, 2017; Kinoshita et al., 2019; Leempoel et al., 2019; Sales et al., 2019; Ushio 84 et al., 2017). Tropical mammal assemblages have been obtained by metabarcoding invertebrate 85 blood meals (e.g. Tessler et al., 2018) and salt licks (Ishige et al., 2017), but samples from the 86 physical environment have tremendous potential to reveal mammal biodiversity over broad spatiotemporal scales (Sales et al., 2019; Ushio et al., 2017). 87

88 In aquatic ecosystems, eDNA metabarcoding has predominantly been applied to 89 characterise fish (e.g. Evans et al., 2017; Hänfling et al., 2016; Lawson Handley et al., 2018; 90 Valentini et al., 2016) and amphibian (e.g. Bálint et al., 2018; Valentini et al., 2016) 91 communities. However, mammals also leave eDNA signatures in water that metabarcoding can 92 detect (Harper et al., 2019; Klymus et al., 2017; Sales et al., 2019; Ushio et al., 2017). Ponds in 93 particular provide drinking, foraging, dispersive, and reproductive opportunities for semiaquatic and terrestrial mammals (Klymus et al., 2017). Samples from these waterbodies could 94 95 uncover biodiversity present in the wider environment (Deiner et al., 2017; Harper et al., 2019).

96 Drinking is a major source of eDNA deposition due to the release of saliva, but mammals may 97 also swim, wallow, urinate or defecate in water (Rodgers & Mock, 2015; Ushio et al., 2017; 98 Williams et al., 2018). Furthermore, arboreal mammals may use ponds less than semi-aquatic 99 and ground-dwelling species, non-territorial mammals may visit ponds less than territorial 100 species, and group-living species may deposit more eDNA than solitary species (Williams et al., 101 2018). Despite evidence for eDNA deposition by semi-aquatic and terrestrial mammals in 102 freshwater ecosystems, little is known about the influence of mammal behaviour on the 103 distribution and strength of the eDNA signal left behind (defined here as proportional read 104 counts).

105 In this study, we conducted two experiments under artificial and natural conditions to 106 evaluate eDNA metabarcoding of pond water as a tool for monitoring semi-aquatic, ground-107 dwelling, and arboreal mammals of conservation or management concern. The first 108 experiment, carried out on nine focal species housed at two wildlife parks, examined the role of 109 sampling strategy, mammal lifestyle, and mammal behaviour on eDNA detection and signal 110 strength under artificial conditions. Mammal eDNA detection is expected from enclosure water that is frequently used by individuals for drinking, swimming and bathing. We hypothesised 111 112 that: (1) eDNA would be unevenly distributed, thus directed sampling would yield stronger 113 eDNA signals (i.e. higher proportional read counts) for mammals than stratified sampling; (2) 114 semi-aquatic mammals would have stronger eDNA signals than ground-dwelling or arboreal 115 mammals; and (3) mammal behaviours involving water contact would generate stronger eDNA 116 signals. The second experiment validated eDNA metabarcoding against camera trapping and 117 field sign searches for mammal identification at natural ponds, and investigated spatiotemporal

variation in mammal eDNA signals. Mammal eDNA detection is unpredictable at natural waterbodies that can be extensive, subject to environmental fluctuations, and used rarely or not at all by individuals. We hypothesised that: (1) eDNA metabarcoding would detect more mammals than camera trapping or field signs; (2) semi-aquatic mammals would be readily detected and their eDNA evenly distributed in ponds in comparison to terrestrial mammals; and (3) temporal sampling would reveal that terrestrial mammal eDNA is detectable for short periods in comparison to fully aquatic vertebrates.

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127 **2. Materials and methods**

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129 2.1 Study species

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131 We studied nine mammal species that are the focus of European conservation or management 132 (Mathews et al., 2018): European water vole (Arvicola amphibius), European otter (Lutra lutra), 133 Eurasian beaver (Castor fiber), European hedgehog (Erinaceus europaeus), European badger 134 (Meles meles), red deer (Cervus elaphus), Eurasian lynx (Lynx lynx), red squirrel (Sciurus 135 vulgaris), and European pine marten (Martes martes). Water vole, otter, red squirrel, pine 136 marten and hedgehog are UK Biodiversity Action Plan species (Joint Nature Conservation 137 Committee, 2018). Water vole, otter, and beaver are semi-aquatic, red squirrel and pine 138 marten are arboreal, and the other species are ground-dwelling. Badger and red deer live in 139 groups whereas the other species are predominantly solitary.

141 **2.2 Experiment 1: eDNA detection and signal strength in artificial systems**

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Behavioural observation and eDNA sampling were conducted between 18th – 21st September 143 2017 at Wildwood Trust (WT), Kent, England, and $10^{th} - 11^{th}$ October 2017 at Royal Zoological 144 Society of Scotland (RZSS) Highland Wildlife Park (HWP), Kingussie, Scotland. Sixteen categories 145 146 of behaviour were defined based on potential contact with waterbodies and species lifestyle, 147 and the frequency and duration of behaviours recorded (Table 1, Appendix A: Table A1). The 148 number of individuals in each enclosure was recorded alongside waterbody size (Table 2). 149 Beaver, lynx, red deer, and red squirrel were present at both wildlife parks, whereas other 150 captive species were only present at WT. Each species was observed for one hour on two 151 separate occasions except nocturnal mammals (badger and beaver), which were observed 152 overnight using camera traps (Bushnell Trophy Cam Standard, Bushnell Corporation, KS, USA). 153 One camera trap per enclosure was positioned perpendicular to the ground (1 m height, 2 m 154 from shoreline) to capture water and shoreline. Cameras took 30 s videos (1920 x 1080) when 155 triggered (30 s interval between triggers) at high sensitivity. Behavioural observation was not 156 undertaken for WT water voles as animals were under guarantine or HWP red squirrels as 157 individuals were wild. Photos of waterbodies in animal enclosures are provided in Appendix B.

Water samples were collected from enclosures within 3 hrs of the second behavioural observation period. Up to six directed or stratified samples were collected, but sample number waried by species according to waterbody size and observed behaviours (Tables A1, A2). Enclosure drinking containers were also sampled and classed as 'other' samples. Bathing and

drinking bowls were sampled where enclosures contained no artificial waterbodies (WT water vole, red squirrel, and hedgehog). The HWP beaver enclosure was empty for 24 hrs before sampling. Water was sampled from a RZSS Edinburgh Zoo (EZ) enclosure containing beavers and classed as 'other'. A sample was collected from a water bath in the HWP woods to capture wild red squirrels and classed as 'other'.

167 Directed samples (2 L surface water taken approximately where behaviours were 168 observed) were collected before stratified samples (2 L surface water [8 x 250 ml pooled 169 subsamples] taken at equidistant points [access permitting] around the waterbody perimeter) 170 to minimise disturbance to the water column and cross-contamination risk. Samples were 171 collected using sterile Gosselin[™] HDPE plastic bottles (Fisher Scientific UK Ltd, UK) and 172 disposable gloves. A field blank (1 L molecular grade water [MGW]) was taken into each species 173 enclosure, opened, and closed before artificial water sources were sampled. Samples (n = 80)174 collected from WT and HWP were transported alongside field blanks (n = 13) in sterile 175 coolboxes with ice packs to the University of Kent (UoK) and EZ respectively, where ice was 176 added to coolboxes.

Samples and blanks were vacuum-filtered within 6 hrs of collection in a UoK wet laboratory and within 24 hrs of collection in an EZ staff room. Surfaces and equipment were sterilised before, during, and after set-up in temporary work areas. Surfaces and vacuum pumps were wiped with 10% v/v chlorine-based commercial bleach (Elliott Hygiene Ltd, UK) solution. Non-electrical equipment was immersed in 10% bleach solution for 10 minutes, followed by 5% v/v MicroSol detergent (Anachem, UK), and rinsed with purified water. Up to 500 ml of each 2 L sample was vacuum-filtered through sterile 0.45 μm mixed cellulose ester

184 membrane filters with pads (47 mm diameter; Whatman, GE Healthcare, UK) using Nalgene™ 185 filtration units. One hour was allowed for each sample to filter and a second filter used if 186 clogging occurred. A filtration blank (1 L MGW) was processed during each filtration round (n =187 12), and equipment sterilised after each filtration round. After 500 ml had filtered or one hour 188 had passed, filters were removed from pads using sterile tweezers, placed in sterile 47 mm 189 petri dishes (Fisher Scientific UK Ltd, UK), sealed with parafilm (Sigma-Aldrich Company Ltd, 190 UK), and stored at -20 °C. The total water volume filtered per sample was recorded for 191 downstream analysis (Table A2; Fig. A1).

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193 **2.3 Experiment 2: eDNA detection and signal strength in natural systems**

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195 At three sites where focal species were present based on cumulative survey data, we selected two ponds (range 293-5056 m², average 1471 m²) within 4 km of each other. The Bamff Estate 196 197 (BE), Alyth, Scotland, was selected for beaver, otter, badger, red deer, and red squirrel, but roe 198 deer (Capreolus capreolus) and red fox (Vulpes vulpes) were also present. Otter, water vole, and 199 badger were present at Tophill Low Nature Reserve (TLNR), Driffield, East Yorkshire, alongside 200 American mink (Neovison vison), stoat (Mustela erminea), weasel (Mustela nivalis), rabbit 201 (Oryctolagus cuniculus), brown hare (Lepus europaeus), red fox, roe deer, and grey squirrel 202 (Sciurus carolinensis). We selected Thorne Moors (TM), Doncaster, South Yorkshire, for red deer 203 and badger, but stoat, weasel, red fox, roe deer, and Reeve's muntjac (Muntiacus reevesi) were 204 also present. Camera traps (Bushnell Trophy Cam Standard/Aggressor, Bushnell Corporation, 205 KS, USA) were deployed at TM (one per pond) and BE (three per pond) one week prior to eDNA sampling and collected once sampling was completed. At TLNR, camera traps (two to three per
pond) were deployed one day before a 5-day period of eDNA sampling and collected one week
after sampling was completed. Camera traps were positioned perpendicular to the ground (1 m
height, 0.3-1 m from shoreline) to capture water and shoreline. Cameras took three
photographs (5 megapixel) when triggered (3 s interval between triggers) at high sensitivity.

Ten stratified samples were collected from the shoreline of each pond (TM: 17th April 211 2018; BE: 20th April 2018; TLNR: 23rd – 27th April 2018) and a field blank (1 L MGW) included as 212 in Experiment 1. TLNR ponds were sampled every 24 hrs over 5 days to investigate 213 214 spatiotemporal variation in mammal eDNA signals. TM and TLNR samples were transported on 215 ice in sterile coolboxes to the University of Hull (UoH) eDNA facility, and stored at 4 °C. BE 216 samples were transported in sterile coolboxes with ice packs to BE accommodation. Surfaces 217 and equipment were sterilised before, during, and after set-up as in Experiment 1. Samples (n =218 140) and field blanks (n = 14) were vacuum-filtered within 4 hrs of collection as in Experiment 1 219 with minor modifications to maximise detection probability as follows. The full 2 L of each 220 sample was vacuum-filtered where possible, two filters were used for each sample, and 221 duplicate filters were stored in one petri dish at -20 °C. A filtration blank (1 L MGW) was processed during each filtration round (n = 21). The total water volume filtered per sample was 222 223 recorded (Table A3).

225 2.4 DNA extraction

227	DNA was extracted within 2 weeks of filtration at the UoH eDNA facility using the Mu-DNA
228	water protocol (Sellers, Di Muri, Gómez, & Hänfling, 2018). The full protocol is available at:
229	https://doi.org/10.17504/protocols.io.qn9dvh6. Duplicate filters from samples in Experiment 1
230	were lysed independently and the lysate from each loaded onto one spin column. As more
231	samples were collected in Experiment 2, duplicate filters were co-extracted by placing both in a
232	single tube for bead milling. An extraction blank, consisting only of extraction buffers, was
233	included for each round of DNA extraction (n = 17). Eluted DNA (100 µl) was stored at -20 °C
234	until PCR amplification.
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236	2.5 eDNA metabarcoding
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237 238	Our eDNA metabarcoding workflow is fully described in Appendix A. Briefly, we performed
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238 239 240	nested metabarcoding using a two-step PCR protocol, where Multiplex Identification (MID) tags were included in the first and second PCR for sample identification (Kitson et al., 2019). The first
238 239 240 241	nested metabarcoding using a two-step PCR protocol, where Multiplex Identification (MID) tags were included in the first and second PCR for sample identification (Kitson et al., 2019). The first PCR amplified eDNA in triplicate with published 12S ribosomal RNA (rRNA) primers 12S-V5-F (5'-
238 239 240 241 242	nested metabarcoding using a two-step PCR protocol, where Multiplex Identification (MID) tags were included in the first and second PCR for sample identification (Kitson et al., 2019). The first PCR amplified eDNA in triplicate with published 12S ribosomal RNA (rRNA) primers 12S-V5-F (5'- ACTGGGATTAGATACCCC-3') and 12S-V5-R (5'-TAGAACAGGCTCCTCTAG-3') (Riaz et al., 2011).
238 239 240 241 242 243	nested metabarcoding using a two-step PCR protocol, where Multiplex Identification (MID) tags were included in the first and second PCR for sample identification (Kitson et al., 2019). The first PCR amplified eDNA in triplicate with published 12S ribosomal RNA (rRNA) primers 12S-V5-F (5'- ACTGGGATTAGATACCCC-3') and 12S-V5-R (5'-TAGAACAGGCTCCTCTAG-3') (Riaz et al., 2011). Harper et al. (2018) validated these primers <i>in silico</i> for UK vertebrates, and found 91/112

247 plate; n = 16) were exotic cichlid (*Maylandia zebra*) DNA (0.05 ng/µl), and PCR negative controls 248 (two per PCR plate; n = 16) were MGW (Fisher Scientific UK Ltd, UK). PCR products were pooled 249 to create sub-libraries (Fig. A2) and purified with Mag-BIND® RxnPure Plus magnetic beads 250 (Omega Bio-tek Inc, GA, USA), following the double size selection protocol established by 251 Bronner et al. (2009). Ratios of 0.9x and 0.15x magnetic beads to 100 μ L of each sub-library 252 were used. Eluted DNA (30 µL) was stored at -20 °C until the second PCR could be performed. 253 The second PCR bound pre-adapters, MID tags, and Illumina adapters to the sub-libraries. PCR 254 products were purified with Mag-BIND® RxnPure Plus magnetic beads (Omega Bio-tek Inc, GA, 255 USA), following the double size selection protocol established by Bronner et al. (2009). Ratios of 256 0.7x and 0.15x magnetic beads to 50 µL of each sub-library were used. Eluted DNA (30 µL) was 257 stored at 4 °C until quantification and normalisation. The library was purified again, quantified 258 by qPCR using the NEBNext[®] Library Quant Kit for Illumina[®] (New England Biolabs[®] Inc., MA, 259 USA), and fragment size (330 bp) and removal of secondary product verified using an Agilent 260 2200 TapeStation and High Sensitivity D1000 ScreenTape (Agilent Technologies, CA, USA). The 261 library (220 eDNA samples, 27 field blanks, 33 filtration blanks, 17 extraction blanks, 16 PCR 262 negative controls, and 16 PCR positive controls) was sequenced on an Illumina MiSeq[®] using a MiSeq Reagent Kit v3 (600-cycle) (Illumina, Inc, CA, USA). Raw sequence reads were 263 264 demultiplexed using a custom Python script. metaBEAT v0.97.11 (https://github.com/HullUni-265 bioinformatics/metaBEAT) was used for quality trimming, merging, chimera removal, clustering, 266 and taxonomic assignment of sequences against our UK vertebrate reference database (Harper 267 et al., 2018) which contains sequences for 103 UK mammals. Taxonomic assignment used a 268 lowest common ancestor approach based on the top 10% BLAST matches for any query that 269 matched a reference sequence across more than 80% of its length at minimum identity of 98%.270

271 **2.6 Data analysis**

272

273 Analyses were performed in R v.3.4.3 (R Core Team, 2017). The total unrefined read counts (i.e. 274 raw taxonomically assigned reads) per sample were calculated and retained for downstream 275 analyses. Assignments were corrected: family and genera containing a single UK species were 276 reassigned to that species, species were reassigned to domestic subspecies, and 277 misassignments were corrected, e.g. Lynx pardinus and Lynx lynx. Manual reassignment 278 duplicated some metaBEAT assignments thus the read count data for these assignments were 279 merged. Taxon-specific sequence thresholds (i.e. maximum sequence frequency of each taxon 280 in PCR positive controls) were used to mitigate cross-contamination and false positives (Table 281 A4, Fig. A3), and remnant contaminants and higher taxonomic assignments removed excluding 282 the following genera. Anas (Dabbling ducks) was retained because potential for hybridisation 283 reduced confidence in species-level assignments, and Emberiza (Buntings) and Larus (White-284 headed gulls) were retained because reference sequences were missing for several common 285 species. Dataset refinement is fully described in Appendix A. Taxonomic assignments remaining 286 in the refined dataset were predominantly of species resolution and considered true positives. 287 We split the refined dataset by Experiment 1 (artificial waterbodies) and Experiment 2 (natural 288 ponds). Proportional read counts for each species were calculated from the total unrefined 289 read counts per sample. Our proportional read count data were not normally distributed 290 (Shapiro–Wilk normality test: W = 0.915, P < 0.001), thus we used a Mann-Whitney U test to

291 compare the median proportional read count of stratified and directed samples across species.

292 We employed binomial Generalized Linear Mixed-effects Models (GLMMs) with the logit 293 link function using the package glmmTMB (development version; Brooks et al., 2017) for the 294 following tests. First, we compared the eDNA signals from stratified and directed samples for 295 each mammal species using a hierarchical model including sample type nested within species 296 (fixed) and wildlife park (random) as effects. We tested the influence of species lifestyle on 297 mammal eDNA signals using a model with species lifestyle (fixed) and species nested within 298 wildlife park (random) as effects. Using directed samples, we tested the influence of behaviour 299 on mammal eDNA signals using two hierarchical models, including species nested within 300 wildlife park (random) and specific (e.g. swimming, drinking) or generic (i.e. water contact 301 versus no water contact) behaviour(s) respectively (fixed) as effects. We assessed model fit 302 using diagnostic plots and performed validation checks to ensure model assumptions were met 303 and overdispersion was absent (Zuur, Ieno, Walker, Saveliev, & Smith, 2009).

304 For Experiment 2, we qualitatively compared mammal presence-absence records 305 generated by eDNA metabarcoding, camera trapping, and field signs. TLNR ponds were 306 sampled every 24 hrs for 5 days, thus proportional read counts were averaged across days for 307 comparison to BE and TM ponds (sampled once each). We qualitatively compared the 308 distribution and persistence of eDNA signals between semi-aquatic and terrestrial mammals 309 using tile plots and heat maps of the unaveraged proportional read counts for identified species 310 at TLNR over the 5-day period. All figures were produced using the package ggplot2 v3.0.0 311 (Wickham, 2016).

3. Results

3.1 eDNA metabarcoding

The sequencing run generated 47,713,656 raw sequence reads, of which 37,590,828 remained following trimming, merging, and length filter application. After removal of chimeras and redundancy via clustering, the library contained 21,127,061 sequences (average read count of 64,215 per sample including controls), of which 16,787,750 (79.46%) were assigned a taxonomic rank. Contamination (Fig. A4) was observed in the field blanks (badger, beaver, lynx, pine marten, red squirrel, and water vole) as well as in the filtration and extraction blanks (human [Homo sapiens] and cichlid). PCR negative controls were contaminated to different extents with human, cichlid, beaver, and pine marten as well as non-focal species. After threshold application, contaminants remaining in eDNA samples included Gentoo penguin (Pygoscelis papua), reindeer (Rangifer tarandus), cichlid, and human. The refined dataset contained 59 vertebrate species, including six amphibians, 10 fish, 19 birds, and 24 mammals (Table A5).

3.2 Experiment 1: eDNA detection and signal strength in artificial systems

333 All nine focal species were detected in captivity, of which seven were detected in all water

334 samples taken from their respective enclosures. HWP red deer were not detected in 2 of 5 335 stratified samples, and WT hedgehog was not detected in 1 of 2 drinking bowl samples (Fig. 1). 336 'Other' samples (neither directed nor stratified) were excluded from further comparisons, thus 337 hedgehog, red squirrel, and water vole were omitted in downstream analyses. Across species, 338 stratified samples (0.406) had a higher median proportional read count than directed samples 339 (0.373), but this difference was not significant (Mann-Whitney U test: U = 1181.5, P = 0.829). Proportional read counts for directed and stratified samples did not significantly differ (χ^2_6 = 340 0.364, P = 0.999) within species either (Fig. 2a; GLMM: θ = 0.168, χ^2_{53} = 8.915, P = 1.000, 341 pseudo- R^2 = 39.21%). Otter proportional read counts were lower than other species, but not 342 343 significantly so. Similarly, species lifestyle (semi-aquatic, ground-dwelling, arboreal) did not influence (χ^2_2 = 0.655, *P* = 0.721) proportional read counts (Fig. 2b; GLMM: θ = 0.213, χ^2_{61} = 344 13.002, *P* = 1.000, pseudo- R^2 = 11.85%). Proportional read counts did not differ (χ^2_{11} = 1.369, *P* 345 = 0.999) according to specific behaviours exhibited by species (Fig. 3a; GLMM: θ = 0.355, χ^2_{31} = 346 11.013, P = 0.999, pseudo- $R^2 = 9.17\%$). Likewise, generic behaviour (i.e. water contact versus no 347 water contact) did not influence (χ^2_{11} = 0.002, P = 0.964) proportional read counts (Fig. 3b; 348 GLMM: θ = 0.217, χ^2_{41} = 8.897, *P* = 1.000, pseudo-*R*² = 8.50%). 349

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351 **3.3 Experiment 2: eDNA detection and signal strength in natural systems**

352

At natural ponds, eDNA metabarcoding, camera trapping, and field signs all detected beaver, red deer, and roe deer. Camera traps (Fig. 4) and field signs recorded red fox and badger when eDNA metabarcoding did not (Fig. 5). However, eDNA metabarcoding revealed small mammals 356 missed by cameras and field signs, including water vole, water shrew (Neomys fodiens), bank 357 vole (Myodes glareolus), common shrew (Sorex araneus), brown rat (Rattus norvegicus), rabbit, 358 grey squirrel, and common pipistrelle (Pipistrellus pipistrellus). We observed mice or vole 359 footprints at BE Pond 1, but could not ascertain species. Fig. 5 summarises mammals recorded 360 by different methods at each site with reference to cumulative survey data. Notably, only 361 beaver was found at the same ponds by all methods. Although methods shared species at site 362 level, species were not always detected at the same pond. Detection rates for species captured 363 by at least one survey method are summarised in Table A6.

364 Sampling of natural ponds revealed spatial patterns in eDNA detection and signal 365 strength. eDNA from non-domestic terrestrial mammals (i.e. mammals excluding dog [Canis 366 lupus familiaris], pig [Sus scrofa domesticus], sheep [Ovis aries] and cow [Bos taurus]) was 367 unevenly dispersed compared with semi-aquatic mammals (Fig. A5). Semi-aquatic beaver and 368 water vole were detected in at least 90% and 60% respectively of water samples (n = 10) 369 collected from single ponds, albeit water shrew was only detected in 10% of samples. Non-370 domestic terrestrial mammals were routinely detected in <20% of water samples collected from 371 a pond and left relatively weak eDNA signals. Overall, beaver was the most consistently 372 detected mammal with the highest proportional read counts. However, the strongest and most 373 evenly distributed signals belonged to amphibians, particularly common frog (Rana temporaria) 374 and great crested newt (Triturus cristatus) (Fig. A5).

TLNR samples collected over a 5-day period (D01-05) revealed that mammal detection heavily depends on the spatial and temporal resolution of eDNA metabarcoding surveys (Fig. A6). Mammal eDNA signals in pond water were ephemeral, often disappearing within 24-48 hrs

of initial detection, as opposed to amphibians that were detected for multiple days and whose eDNA signal increased in strength. The majority of semi-aquatic or terrestrial mammals were only detected in a single sample on each day.

381

382

383 **4. Discussion**

384

385 We have demonstrated the potential of eDNA metabarcoding for monitoring conservation and 386 management priority mammals, but species detection rates are variable. Our experiments have 387 validated this molecular approach and provided new insights that will inform the development 388 and application of mammal eDNA metabarcoding. Sampling strategy, mammal lifestyle, and 389 mammal behaviour did not influence eDNA detection and signal strength in captivity, but all 390 played vital roles in natural ponds. Although semi-aquatic and terrestrial mammals were 391 detected from pond water, their eDNA signals were temporary and weak in comparison to 392 aquatic amphibians and fishes. Nonetheless, this suggests that eDNA is representative of 393 contemporary and local mammal diversity.

394

395 **4.1 Influence of sampling strategy and mammal behaviour on eDNA detection**

396

In Experiment 1, all nine focal species were detected in captivity, and seven were detected in all
 water samples taken from their respective enclosures. This demonstrates that our method can
 successfully detect a variety of mammals from pond and drinking water. Surprisingly, we found

400 that neither sampling strategy nor mammal lifestyle nor mammal behaviour influenced eDNA 401 detectability and signal strength in captivity. This included behaviours associated with eDNA 402 deposition, e.g. swimming, drinking, urination, and defecation (Rodgers & Mock, 2015; Ushio et 403 al., 2017; Williams et al., 2018). Enclosures were permanently occupied and artificial 404 waterbodies likely saturated with eDNA, which possibly masked behavioural signals. Modest 405 replication may have limited experimental power, preventing patterns being detected 406 statistically. Nonetheless, our results show that mammal contact with water enables eDNA deposition and detection. 407

408 Unsurprisingly, given the nature of wild mammal interactions with natural systems 409 versus those in captivity, Experiment 2 results highlight the challenges of mammal eDNA 410 detection. We recorded 17 mammals using three monitoring tools, comparable to the 17 411 mammals expected from cumulative survey data despite discordance. Field signs and camera 412 trapping detected red fox and badger where eDNA metabarcoding did not, but eDNA 413 metabarcoding identified water vole and other small mammals missed on camera or with 414 ambiguous field signs, i.e. mice, voles, shrews. Importantly, camera trap deployment period, 415 height, and positioning may have influenced small mammal detection by this method 416 (Caravaggi et al., 2018). Ishige et al. (2017) achieved comparable mammal detection at salt licks 417 with eDNA metabarcoding and camera trapping, but species presence was inconsistent 418 between salt licks surveyed. Using multi-species occupancy modelling for three mammal 419 species, Sales et al. (2019) observed water-based eDNA metabarcoding provided comparable 420 detection probabilities to conventional survey methods and actually outperformed camera 421 trapping. Similarly, Leempoel et al. (2019) found soil-based eDNA metabarcoding identified the

same mammals as camera trapping as well as small mammals rarely seen on camera, albeit the
methods differed between sites. Our own results echo all three studies, where despite some
inconsistencies, eDNA metabarcoding enhanced species inventories and identified smaller,
cryptic taxa.

426 Notably, no survey method captured semi-aquatic otter despite presence at study sites 427 and successful detection in eDNA metabarcoding studies of UK ponds (Harper et al., 2019), 428 lakes (Hänfling et al., 2017), and rivers/streams (Sales et al., 2019). Captive otter also had a 429 weaker eDNA signal than other semi-aquatic mammals studied here. Lower eDNA detection 430 rates for otter, badger, and red fox may stem from species' ecologies (Sales et al., 2019). These 431 mammals are wide-ranging (Gaughran et al., 2018; Thomsen et al., 2012) and may not readily 432 release DNA in water. Otters often spraint on grass or rock substrata outside water and use 433 latrines associated with caves and dens (Ruiz-Olmo & Gosálbez, 1997). As terrestrial mammals, 434 red fox and badger must drink from or enter ponds for eDNA deposition to occur (Rodgers & 435 Mock, 2015; Ushio et al., 2017; Williams et al., 2018). Otter, badger, and red fox detection may 436 require greater spatiotemporal resolution of eDNA sampling. This is reinforced by other eDNA 437 metabarcoding studies where mammal detection was highly variable across sites surveyed 438 (Ishige et al., 2017; Klymus et al., 2017; Leempoel et al., 2019; Sales et al., 2019; Ushio et al., 439 2017). False negatives may instead be symptomatic of metabarcoding bias, but this is unlikely in 440 our study (section 4.2).

eDNA from other semi-aquatic mammals was evenly distributed, being found in most or all samples collected on fine spatial scales within natural ponds, whereas terrestrial mammal eDNA was highly localised and detected in few (<20%) samples. Mammal eDNA signals varied

444 temporally, being detectable for two consecutive days maximum. Depending on the species, 445 mammal eDNA may be spatially and temporally clumped in lentic ecosystems due to the nature 446 and frequency of water contact. Unless non-domestic mammals exhibit behaviours involving 447 prolonged water contact (e.g. swimming, wallowing), they may only be detected at drinking 448 sites (Klymus et al., 2017; Ushio et al., 2017; Williams et al., 2018). Conversely, domestic 449 mammals may have elevated detection rates in ponds due to high occurrence of these 450 waterbodies in agricultural landscapes as well as eDNA transport by rainfall and run-off (Staley 451 et al., 2018). eDNA detection and persistence are further influenced by group size, where eDNA 452 from multiple individuals endures for longer periods in water than eDNA from single individuals 453 (Williams et al., 2018). Detailed investigations incorporating biotic (e.g. population size, body 454 mass, behaviour) and abiotic (e.g. temperature, pH, rainfall) factors are needed to understand 455 the longevity of mammal eDNA signals in aquatic ecosystems (Rodgers & Mock, 2015; Sales et 456 al., 2019; Williams et al., 2018).

Our two experiments have shown that sampling strategy influences mammal eDNA 457 458 detection. Mammal eDNA was evenly distributed in closed, artificial waterbodies, but locally distributed in open, natural ponds. Captive mammal enclosures contained one species 459 460 (excluding HWP red deer) and a drinking container(s) and/or small waterbody (range 0.01-162 m^2 , mean 27.4 m^2). Some enclosures housed more individuals of a species than others, thereby 461 462 increasing eDNA deposition and detection probability (Williams et al., 2018). Wild mammals 463 have an array of freshwater habitats at their disposal and can hold vast territories. Therefore, 464 rates of pond visitation and eDNA deposition are more irregular (Klymus et al., 2017; Ushio et 465 al., 2017), possibly leading to between-sample variation (Williams et al., 2018).

467 **4.2** Accounting for false positives and false negatives in metabarcoding

468

469 eDNA metabarcoding has potential for inclusion in mammal monitoring schemes (section 4.3), 470 but like existing monitoring tools, may produce false negatives or false positives. Our process 471 controls identified low-level contamination at all stages of metabarcoding, but primarily during 472 sampling or PCR (Appendix A). We applied taxon-specific sequence thresholds to our data to 473 mitigate false positives as in Harper et al. (2019). Remnant contaminants were cichlid 474 (laboratory), Gentoo penguin (environment), reindeer (environment), and human (environment/laboratory). Gentoo penguin is housed at EZ and was identified from EZ beaver 475 476 enclosure water. The WT red squirrel and reindeer enclosures are in close proximity. DNA 477 transport by wildlife (e.g. waterfowl [Hänfling et al., 2016]) and park staff/visitors may explain 478 this environmental contamination. Human DNA was present across process controls 479 corresponding to artificial and natural waterbodies. Human DNA may be amplified and 480 sequenced instead of focal species, potentially resulting in false negative detections for rare 481 and/or less abundant species. Human DNA blocking primers can prevent this bias, but may 482 impair PCR amplification efficiency (Klymus et al., 2017; Ushio et al., 2017; Valentini et al., 483 2016). Sequence thresholds are one method of accounting for contamination in metabarcoding 484 datasets, but this is a topic that warrants deeper investigation aimed at researching and 485 refining standardised methods for false positive identification and mitigation, e.g. the R 486 package microDecon (McKnight et al., 2019).

487

In our study, eDNA metabarcoding produced false negatives for otter, badger, and red

488 fox at natural ponds. We selected a 12S metabarcode designed to amplify vertebrate DNA (Riaz 489 et al., 2011). One of four fox reference sequences (NCBI Accession: KF387633.1) possessed one 490 mismatch to the forward primer, and one of three otter reference sequences (NCBI Accession: 491 EF672696.1) possessed one mismatch to the reverse primer. These mismatches did not occur 492 within the first or last four bases of either primer sequence, and there were no primer 493 mismatches with the badger reference sequences (Harper et al., 2018). Therefore, amplification 494 bias was not responsible for these false negatives. DNA from aquatic and more abundant 495 species may have overwhelmed otter, badger, and red fox DNA during amplification and 496 sequencing, i.e. species-masking (Kelly, Port, Yamahara, & Crowder, 2014; Klymus et al., 2017). 497 Species-masking may also arise from use of proportional read counts as an index of eDNA signal 498 strength. High proportional read counts for a species may translate to a weak eDNA signal if the 499 total mammalian eDNA concentration is highly variable between samples or lower than the 500 total eDNA concentration for other taxonomic groups in a sample. Metabarcoding primers 501 targeting mammals (Ushio et al., 2017) or multi-marker (e.g. 12S, 16S, COI) investigations 502 (Evans et al., 2017; Hänfling et al., 2016; Kelly et al., 2014; Klymus et al., 2017) may improve 503 mammal detection in systems with competition from non-target aquatic species and where 504 total mammalian eDNA concentration varies between samples. Similarly, more biological and 505 technical replication may improve species detection probabilities (Evans et al., 2017; Lawson 506 Handley et al., 2019; Sales et al., 2019; Valentini et al., 2016). Importantly, otter also had lower 507 qPCR detection than amphibians and fish (Thomsen et al., 2012). A metabarcoding and qPCR 508 comparison (e.g. Harper et al., 2018; Lacoursière-Roussel, Dubois, Normandeau, & Bernatchez, 2016) would confirm whether poor amplification efficiency for otter arises from technical bias 509

510 or species ecology, and whether eDNA metabarcoding can reliably monitor otter alongside the 511 wider mammalian community.

512

513 **4.3 Scope of eDNA metabarcoding for mammal monitoring**

514

515 Mammal population assessments are hindered by lack of data and systematic monitoring for 516 many species (Mathews et al., 2018). Distribution and occupancy data are poor for most 517 species, with ongoing survey effort biased toward rare species. Surveys heavily rely on citizen 518 science and casual records (Massimino et al., 2018). Tools that provide standardised, systematic 519 monitoring of mammal populations are needed (Mathews et al., 2018). Despite issues inherent 520 to metabarcoding for biodiversity monitoring (Deiner et al., 2017), this tool has enormous 521 potential to enhance mammal monitoring, conservation, and management. eDNA 522 metabarcoding generates distribution data for multiple species, whether rare, invasive, or 523 abundant, and could track conflicting species simultaneously, e.g. water vole, American mink, 524 and otter (Bonesi & Macdonald, 2004) or red squirrel, grey squirrel, and pine marten (Sheehy, 525 Sutherland, O'Reilly, & Lambin, 2018).

eDNA metabarcoding can rapidly survey multitudes of aquatic sites at landscape-scale where camera traps might be resource-intensive, cost-inefficient, and susceptible to theft/damage (Ushio et al., 2017). Field signs require volunteer time and skill (Sadlier et al., 2004) to be employed at comparable spatial scales to eDNA metabarcoding which could provide accurate data for species misidentified from field signs, e.g. mice and voles, otter and mink (Franklin et al., 2019; Harris & Yalden, 2004). However, camera traps and field signs both

recorded species that eDNA metabarcoding missed. Therefore, eDNA metabarcoding is complementary and should be incorporated into, not replace, existing monitoring schemes (Leempoel et al., 2019; Sales et al., 2019). This tool could be most effective in mammal monitoring if deployed at the edges of known species distributions, in areas where species presence is unknown, and in areas with isolated species records (Mathews et al., 2018).

537

538 **4.4 Recommendations for mammal survey using eDNA metabarcoding**

539

540 Water-based eDNA metabarcoding shows great promise for mammal monitoring encompassing conservation and management priority species (Sales et al., 2019). However, there are factors 541 542 to be considered when designing and conducting mammal eDNA surveys that may not be 543 problematic for surveys of fishes or amphibians. Mammal eDNA detection probabilities from 544 natural ponds will likely be high when areas with dense populations are studied, but rigorous 545 sampling strategies will be required to track mammals in areas sparsely populated by 546 individuals. Multiple ponds must be sampled repeatedly, and samples taken at multiple 547 locations within ponds without pooling to enable site occupancy inferences. Importantly, we 548 sampled natural ponds in spring but sampling in other seasons may produce different results, reflective of species' ecologies (Lawson Handley et al., 2019). To account for differential 549 550 mammal visitation rates and maximise eDNA detection probabilities, we recommend that 551 researchers and practitioners using eDNA metabarcoding for mammal monitoring channel their 552 efforts into extensive sampling of numerous waterbodies in a given area over prolonged 553 timescales. Water-based eDNA appears to be indicative of contemporary mammal presence,

554	with most mammal eDNA signals lost within 1-2 days. Therefore, eDNA metabarcoding could
555	provide valuable mammalian community "snapshots" that may not be obtained with other
556	survey methods (Ushio et al., 2017). Different sample types (e.g. water, soil, snow, salt licks,
557	feeding traces, faeces, hair, and blood meals) may also offer new insights to mammal
558	biodiversity (Franklin et al., 2019; Ishige et al., 2017; Kinoshita et al., 2019; Leempoel et al.,
559	2019; Sales et al., 2019; Tessler et al., 2018; Ushio et al., 2017).

561

562 Data accessibility

563

Raw sequence reads have been archived on the NCBI Sequence Read Archive (Study:
SRP164740; BioProject: PRJNA495011; BioSamples: SAMN10195928 - SAMN10196255; SRA
accessions: SRR7986451 - SRR7986778). Jupyter notebooks, R scripts and corresponding data
are archived online (https://doi.org/10.5281/zenodo.2561415).

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569

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571

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583

584

585 Author contributions

586

587 L.R.H, B.H, and L.L.H conceived and designed the study. A.I.C and M.G coordinated sampling at 588 Wildwood Trust and RZSS Highland Wildlife Park respectively. L.R.H, C.D.M, C.J.M, and T.L 589 collected and filtered water samples. A.L, T.L, and T.B helped select natural ponds to be 590 surveyed using eDNA, camera trapping, and field signs, and provided camera traps for the 591 study. L.R.H, A.L, and T.L deployed camera traps, which were then collected and footage 592 analysed by L.R.H. L.R.H processed samples in the laboratory with advice from C.D.M and A.M. 593 D.S.R sequenced the final library. L.R.H completed bioinformatic processing of samples, and 594 subsequent data analysis. L.R.H wrote the manuscript, which all authors contributed critically to 595 drafts of and gave final approval for publication.

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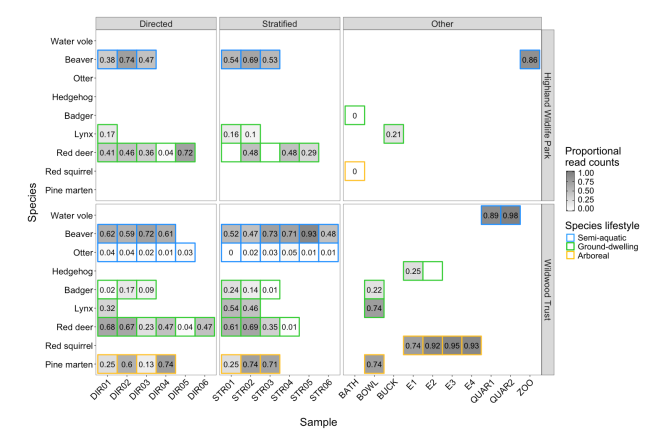
760 **Table 1.** Ethogram used to catalogue mammal behaviours that occur in or near artificial 761 waterbodies in captive enclosures. Importantly, this ethogram was designed to catalogue 762 mammal behaviours potentially leading to eDNA deposition. Therefore, it may not be 763 comparable to ethograms typically used in study of captive animals.

764

Behaviour	Definition			
Swimming	Mammal completely submerged in and moving through waterbody			
	using limbs			
Bodypart in water	Mammal partially submerged in waterbody, e.g. foot or tail ir water			
Drinking	Water taken into mouth and swallowed by mammal			
Feeding	Food taken into mouth and swallowed by mammal in or near waterbody, e.g. otter and fish			
Scratching	Bodypart or external object in enclosure used by mammal to relieve itch near waterbody			
Urinating/scent-marking	Liquid excretion passed by mammal in or near waterbody			
Pooing	Solid excretion passed by mammal in or near waterbody			
Sniffing	Air visibly drawn through nose of mammal to detect a smell around waterbody, possibly involving contact with water			
Standing	Mammal motionless in or near waterbody			
Walking	Mammal moving around waterbody at a regular pace by lifting and setting down each foot in turn, never having both feet off the ground at once			
Running	Mammal moving around waterbody at a speed faster than a walk never having both or all the feet on the ground at the same time			
Vocalising	Mammal producing sound while in or near waterbody			
Grooming	Mammal cleaning fur or skin with its tongue while in or near waterbody			
Resting	Mammal lying down or sitting in or near waterbody			
Other	Behaviour exhibited in or near waterbody that does not conform to			
	other categories, e.g. chasing tail			
Not visible	Mammal moved to part of enclosure not visible to the observer			

- **Table 2.** Summary of focal species studied at wildlife parks and their lifestyle. The number of
- 767 individuals present and waterbody size in enclosures is provided.

Site	Species	Lifestyle	Enclosure	Number of individuals	Waterbody size (m ²)
/ildwood Trust	European otter (Lutra lutra)	Semi-aquatic	1	2	162
European water vole (Arvicola amphibius)		Semi-aquatic	1	4	0.09
		2	1	0.09	
	European beaver	Semi-aquatic	1	2	100
(Castor fiber) European hedgehog (Erinaceus europaeus)	(Castor fiber)		2	1	100
		Ground- dwelling	1	1	0.04
	(Erinaceus europaeus)		2	2	0.04
	European badger (<i>Meles meles</i>)	Ground- dwelling	1	4	1.73
	Red deer (Cervus elaphus)	Ground- dwelling	1	8	100
	Eurasian lynx (<i>Lynx lynx</i>)	Ground- dwelling	1	2	2
	Red squirrel (Sciurus vulgaris)	Arboreal	1	2	0.01
			2	3	0.01
			3	3	0.01
			4	2	0.01
European pine marten	Arboreal	1	1	2	
	(Martes martes)		2	1	0.375
lighland Wildlife Park	Red squirrel (Sciurus vulgaris)	Arboreal	NA	NA	0.25
	Eurasian lynx (<i>Lynx lynx</i>)	Ground- dwelling	1	8	2
	European beaver (Castor fiber)	Semi-aquatic	1	2	50
	Red deer (Cervus elaphus)	Ground- dwelling	1	30	NA



771

772 **Figure 1.** Heatmap showing proportional read counts for eDNA samples (n = 81) from 773 Experiment 1. The heatmap is faceted by sample type (directed, stratified or other) and wildlife 774 park (Highland Wildlife Park or Wildwood Trust). Each cell represents an individual sample 775 taken from an enclosure containing the focal species in that row. Directed (DIR01-DIR06) and 776 stratified (STR01-STR06) samples were collected for each species from artificial waterbodies. 777 Samples were also collected from drinking containers (E1, E2, E3, E4, BOWL, BUCK), water vole (QUAR1, QUAR2) and RZSS Edinburgh Zoo beaver (ZOO) enclosures, and a water bath (BATH) in 778 779 RZSS Highland Wildlife Park woods. The maximum proportional read count for each cell (i.e. 780 sample) is 1, if all reads from a particular sample belonged to the focal species. Cells containing 781 0 represent samples with proportional read counts less than 0.01 whereas empty cells are 782 samples with proportional read counts of exactly 0.

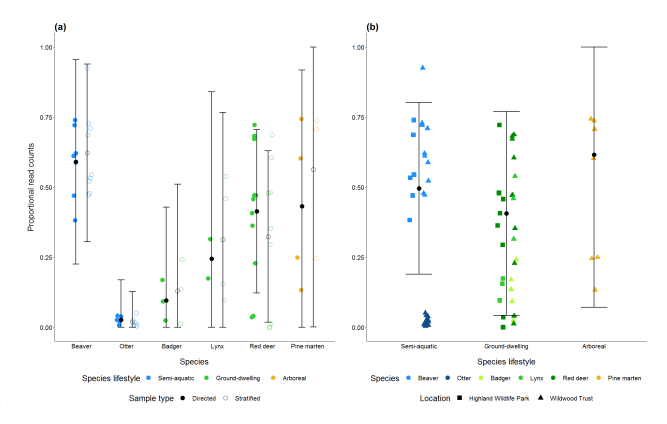




Figure 2. Relationships predicted by the binomial GLMMs between proportional read counts and sample type nested within species (a) or species lifestyle (b) for Experiment 1. The observed data (coloured points) are displayed against the predicted relationships (black points with error bars) for each species (a) or species lifestyle (b). Points are shaped by sample type (a) or wildlife park (b), and coloured by species lifestyle. Error bars represent the standard error around the predicted means.

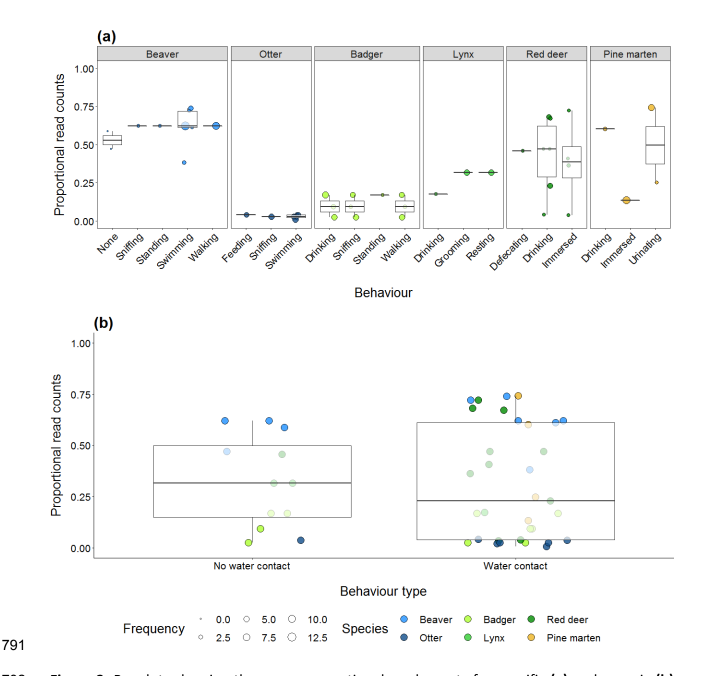


Figure 3. Boxplots showing the mean proportional read counts for specific (a) and generic (b) behaviour(s) exhibited by focal species in Experiment 1. Boxes show 25th, 50th, and 75th percentiles, and whiskers show 5th and 95th percentiles. Points are coloured by species lifestyle, and each point in (a) represents a directed sample sized by frequency of behaviour. The behaviour 'none' for beaver represents occurrences of beaver in water but out of view of camera traps.

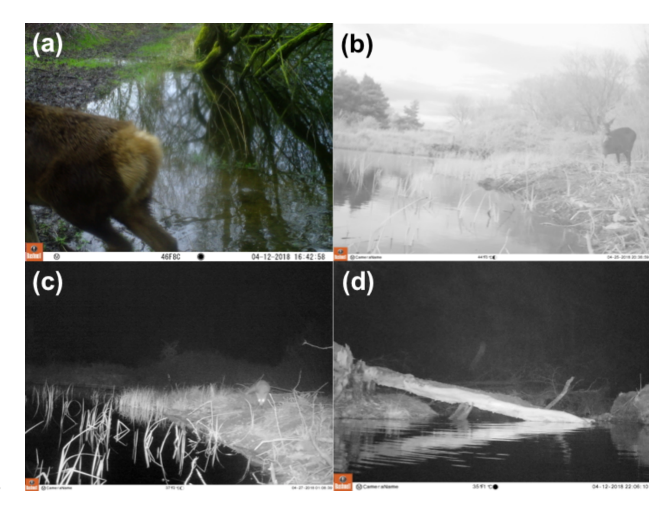


Figure 4. Exemplar camera trap photos taken at natural ponds where focal species were
present in Experiment 2. Red deer was recorded at Thorne Moors (a), roe deer (b) and red fox
(c) were recorded at Tophill Low Nature Reserve, and beaver was recorded at the Bamff Estate
(d).

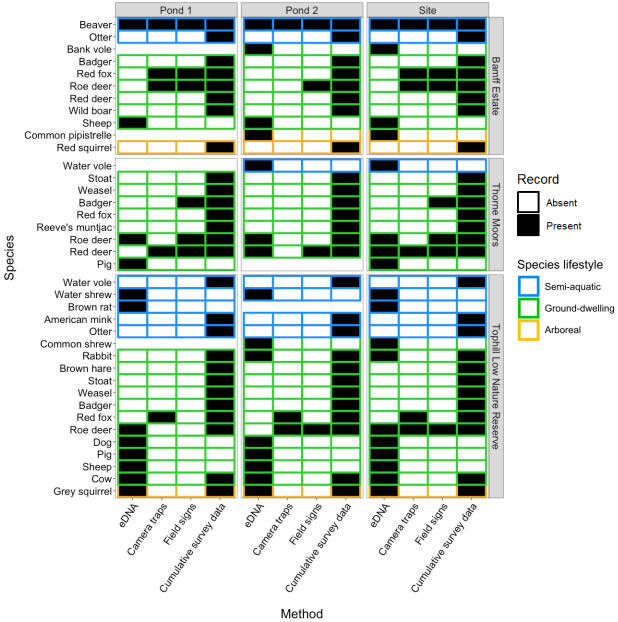


Figure 5. Tile plot showing species presence-absence at individual pond and site-level as 805 indicated by field signs, camera trapping, and eDNA metabarcoding in Experiment 2. Surveys 806 807 were performed at sites where focal species presence was confirmed by cumulative survey 808 data.