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1 **The protozoan parasite *Trichomonas gallinae* causes adult and nestling mortality in a**  
2 **declining population of European Turtle Doves, *Streptopelia turtur*.**

3

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13

14 **RUNNING TITLE:** Mortality in European Turtle Doves.

15

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26

27 **SUMMARY**

28 Studies incorporating the ecology of clinical and sub-clinical disease in wild populations of  
29 conservation concern are rare. Here we examine sub-clinical infection by *Trichomonas*  
30 *gallinae* in a declining population of the European Turtle Dove and suggest caseous lesions  
31 cause mortality in adults and nestlings through subsequent starvation and/or suffocation. We  
32 found a 100% infection rate by *T. gallinae* in adult and nestling Turtle Doves (n=25) and  
33 observed clinical signs in three adults and four nestlings (28%). Individuals with clinical signs  
34 displayed no differences in any skeletal measures of size but had a mean 3.7% reduction in  
35 wing length, with no overlap compared to those without clinical signs. We also identified *T.*  
36 *gallinae* as the suggested cause of mortality in one Red-legged Partridge although disease  
37 presentation was different. A minimum of four strains of *T. gallinae*, characterised at the  
38 ITS/5.8S/ITS2 ribosomal region, were isolated from free-living Turtle Doves, but all birds  
39 (Turtle Doves and the Red-legged Partridge) with clinical signs carried a single strain of *T.*  
40 *gallinae*, suggesting that parasite spill over between Columbidae and Galliformes is a  
41 possibility that should be further investigated. Overall, we highlight the importance of  
42 monitoring populations for sub-clinical infection rather than just clinical disease.

43

44 **KEYWORDS:** disease, feeding ecology, supplementary food, necropsy, PCR.

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52 **KEY FINDINGS**

- 53 • First known case of mortality in adult and nestling Turtle Doves from trichomonosis.
- 54 • 100% infection rate by *T.gallinae* in Turtle Doves with clinical signs in 28% of birds.
- 55 • Birds with clinical signs had 3.7% shorter wing lengths: no variance in skeletal assays.
- 56 • A recommendation that parasite spill over between Columbidae and Galliformes
- 57 should be further investigated.
- 58 • A recommendation to monitor populations for both clinical and sub-clinical infection
- 59 to better understand disease threats to populations of conservation concern.

60

61 **INTRODUCTION**

62 The avian disease trichomonosis has a global distribution and widespread infection potential  
63 and is now considered a major contributing factor to the regulation and even decline of avian  
64 populations (Stabler 1954; Krone *et al.* 2005; Forrester and Foster 2008; Robinson *et al.*  
65 2010; Amin *et al.* 2014). In recent years, trichomonosis has undergone a European spread as a  
66 consequence of avian migration from the UK and has been linked to widespread declines in  
67 finch (Fringillidae) populations (Robinson *et al.* 2010; Lawson *et al.* 2011b, 2012; Lehikoinen  
68 *et al.* 2013; Ganas *et al.* 2014). This recent trichomonosis epizootic reported in finches is  
69 thought to have resulted from parasite spill over of one clonal strain of *Trichomonas gallinae*  
70 from Columbidae to new host species at shared communal garden feeding stations (Lawson *et*  
71 *al.* 2012; Ganas *et al.* 2014). Within the UK, *T. gallinae* has recently been reported within four  
72 species of Columbidae (Lennon *et al.* 2013).

73

74 Trichomonosis can result in death by suffocation and/or starvation due to necrotic  
75 ulcerations/lesions (Stabler 1954). However, host susceptibility and parasite virulence vary,  
76 and hosts often show no clinical signs unless they are nestlings or are infected with a  
77 pathogenic strain (BonDurant and Honigberg 1994; Bunbury *et al.* 2008b; Sansano-Maestre *et*  
78 *al.* 2009; Robinson *et al.* 2010). The trichomonad life cycle has no intermediate host and  
79 transmission occurs horizontally through mutual courtship feeding, or vertically via transfer  
80 of crop milk from adults to nestlings, as well as indirectly through shared food and water  
81 sources (Stabler 1954; Kocan 1969).

82

83 The European Turtle Dove *Streptopelia turtur* (hereafter referred to as ‘Turtle Dove’) is a sub-  
84 Saharan migrant, the populations of which have undergone sustained declines in abundance  
85 and contractions in range. At a pan-European level, Turtle Doves declined by 73% between

86 1980 and 2010 (PECBMS 2012). In the UK, declines of 93% were recorded between 1970 and  
87 2010 (Eaton *et al.* 2012), with a coinciding 51% reduction in range (Balmer *et al.* 2013).

88

89 Turtle Dove population declines on the UK breeding grounds have been attributed to a  
90 reduction in breeding productivity (Browne and Aebischer 2004), accompanied by a  
91 concurrent dietary switch from 'natural' arable plant seeds to anthropogenic food resources  
92 such as grain piles in farmyards (Browne and Aebischer 2003). The dietary switch and the  
93 reduction in breeding attempts may reflect diminished availability of any food rather than  
94 quality alone. This change in feeding behaviour increases the potential for interactions  
95 between the main UK species of Columbidae and other granivorous farmland birds, including  
96 introduced game birds known to be carriers of *T. gallinae* (Pennycott 1998; Hofle *et al.* 2004).

97

98 Limited information is available about the infection rate of the *T. gallinae* parasite in free-  
99 living Turtle Doves, though Muñoz (1995) found an infection rate of 50% in Spain. Lennon *et*  
100 *al.* (2013) found a high incidence of trichomonad parasite infection(86%) in Turtle Doves on  
101 breeding grounds in the UK ; as high as or higher than in any s resident species of  
102 Columbidae.

103

104 Here we describe mortality in adult and nestling Turtle Doves caused by a single strain of the  
105 protozoan parasite *T. gallinae*, strongly suggested through gross necropsy and subsequent  
106 isolation, culture and sequencing of extracted parasites. We also cultured *T. gallinae* parasites  
107 from artificial food and water sources, suggesting likely routes of transmission.

108 **MATERIALS AND METHODS**

109 Birds were sampled during May – July 2012 on farms in East Anglia, UK at three sites in  
110 Essex (Tolleshunt D'Arcy: 51° 77'N, 0° 79'E; Marks Tey: 51° 88'N, 0° 79'E; and Silver End: 51°  
111 85'N, 0° 62'E) and one in Norfolk (Hilgay: 52° 56'N, 0° 39'E). Sites were baited with either  
112 Wheat *Triticum spp.*, Oil Seed Rape *Brassica napus*, or a standard wild bird seed mix (Maize  
113 *Zea mays L.*, Sunflower *Helianthus annuus*, Pinhead Oatmeal *Avena sativa*, Wheat, Red Dari  
114 *Sorghum L.*, Red and Yellow Millet *Panicum miliaceum*, Hempseed *Cannabis sativa* and Canary  
115 seed *Phalaris canariensis*) in areas where farmers regularly provided supplementary food or  
116 grain tailings, known to be an increasingly important constituent of Turtle Dove diet in the  
117 UK, especially in the early breeding season (Browne and Aebischer 2003). Adults were caught  
118 at each site with either whoosh nets or mist nets (Redfern and Clark 2001). Individuals  
119 displaying clinical symptoms of trichomonosis (feathering around the beak matted, wet and  
120 discoloured by regurgitated saliva) were caught at two of the sites in Essex (Tolleshunt D'Arcy  
121 and Marks Tey), approximately 18 km apart.

122

123 Every bird captured was ringed with a British Trust for Ornithology (BTO) individually  
124 numbered leg ring, weighed with a digital balance (Satrue, Taiwan, ± 0.1g) and standard  
125 morphometrics were recorded (wing length ± 0.5mm with a slotted rule, tarsus length ± 0.1  
126 mm and head-beak length ± 0.1 mm with Vernier callipers; Redfern and Clark 2001). The oral  
127 cavity, throat and crop of each bird were also swabbed using an individual sterile viscose  
128 swab, which was then used to inoculate an individual InPouch culture kit (Biomed  
129 Diagnostics, Oregon, USA). Culture kits were incubated at 37°C for 3 – 7 days in order to give  
130 the protozoan parasites sufficient time to culture (Bunbury *et al.* 2005) before isolating  
131 parasites using a standard procedure (further detailed in Lennon *et al.*, 2013). Samples were  
132 then frozen until subsequent analysis.

133

134 In June and July 2012, we also equipped all captured adult Turtle Doves caught with tail-  
135 mounted Pip3 radio-tags (Biotrack, Dorset, UK) weighing 1.7g (<1.5% of body mass), to help  
136 in locating nests. Some of these birds showed clinical symptoms of trichomonosis (see above)  
137 but none appeared lethargic or had any apparent difficulty breathing, and all flew strongly  
138 upon release. Turtle Dove nests were found by monitoring the movements of radio-tagged  
139 birds and cold-searching suitable habitat known to contain territorial males. Nests were  
140 monitored every 2-3 days and when nestlings reached 7 days old, they were ringed, weighed  
141 and were swabbed using the same procedure as for adults, taking particular care not to  
142 damage to the oro-pharyngeal lining.

143

144 When fresh carcasses of adults (n=2) or nestlings (n=2) were found (i.e. those displaying no  
145 or minimal signs of autolysis), a swab of the oral cavity, throat and crop was taken (as  
146 described above), and any fly eggs or maggots present were removed. The carcasses were  
147 then stored in newspaper and kept at 4°C until gross necropsy could be performed (within 48  
148 hours of being found). A further three nestling carcasses that we couldn't examine post  
149 mortem due to significant fly damage were swabbed for trichomonosis. A moribund Red-  
150 legged Partridge *Alectoris rufa* was also found at one site, and whilst it did not exhibit  
151 diagnostic clinical symptoms of trichomonosis (it was sat in the middle of the farmyard,  
152 unresponsive to stimuli with closed eyes and 'fluffed up' feathers), the bird was retrieved for  
153 necropsy, since it had shared a feeding site with adult Turtle Doves showing clinical signs of  
154 the disease.

155

156 All investigative gross necropsies were carried out by JES following a standard simplified  
157 protocol as previously described (van Riper & van Riper 1980; Cooper 2004; Bunbury *et al.*  
158 2008b) involving both external and internal observation, taking samples from any lesions  
159 found for subsequent DNA analysis and the documentation of findings. Clinical signs of

160 trichomonosis in gross necropsy can include swollen head and eyes and yellow caseous  
161 necrotic lesions predominantly found within the oral cavity, pharynx and upper digestive  
162 tract (Stabler 1954; Bunbury *et al.* 2008b).

163

164 All carcasses except one were found at the Tolleshunt D'Arcy site in Essex. Thus swabs were  
165 collected from one feeding site and three water sources at this site (stagnant pools in artificial  
166 containers); to determine whether associated food or water sources might be an  
167 environmental source of *T. gallinae* parasites (Kocan 1969).

168

169 Total genomic DNA was extracted from isolated parasites and all trichomonad lesions with a  
170 DNeasy Blood and Tissue Kit (Qiagen, Hilden, Germany) following the manufacturer's  
171 instructions. DNA extractions were verified with a Nanodrop ND-1000 Spectrophotometer  
172 (Thermo Scientific, Wilmington, USA), to determine DNA concentration.

173

174 An optimised PCR protocol was used with published primers (Gaspar da Silva *et al.* 2007) to  
175 amplify the ITS1/5.8S/ITS2 ribosomal region. PCR reactions were performed in 50 µl  
176 volumes containing 10 µl of extracted DNA with 0.6µM of both primers TFR1 and TFR2,  
177 0.8mM dNTPs, 0.5 units GoTaq Hot Start Polymerase (Promega, Madison, USA), and 1.5mM  
178 MgCl<sub>2</sub>. The thermal profile included an initial denaturation at 94°C for 5 min, then 36 cycles of  
179 94°C for 1 min, 65°C for 30 sec and 72°C for 1 min, and a final extension at 72°C for 5 min. PCR  
180 reactions were run on a GeneAmp PCR System 9700 (Applied Biosystems, Foster City, CA)  
181 with three previously identified positives from Columbiformes and one negative control of  
182 molecular water. Each sample was run a total of three times to confirm the presence or  
183 absence of parasites. PCR products were electrophoresed on a 0.8% agarose gel stained with  
184 ethidium bromide in 0.5x TBE buffer. The presence of a 400bp band when amplified products

185 were observed under UV light indicated a positive sample. All positive samples were  
186 sequenced by Source BioScience (Nottingham, UK).

187

188 The ITS1/5.8S/ITS2 ribosomal region of DNA is highly conserved in *Trichomonas* spp., with a  
189 low rate of mutation (Grabensteiner *et al.* 2010), thus any sequences differing in one or more  
190 base pairs were considered to be distinct strains. We used a combination of BioEdit (Hall  
191 2005) and 4Peaks (Griekspoor and Groothuis 2006) to trim, manually align, and assess  
192 forward and reverse sequences for each PCR product for sequencing. As strain length can  
193 influence the closest matching Genbank sequence (authors, pers. obs.), all sequences from this  
194 study were initially aligned with each other in order to identify unique sequences. The  
195 longest of each unique sequence was then queried in the NCBI-BLAST database (Altschul *et al.*  
196 1997).

197

198 To establish whether adults with clinical signs of trichomonosis differed in weight, wing  
199 length, or skeletal measures of size (head-beak length and tarsus length) to apparently  
200 healthy birds, we used general linear models in R (R Core Team 2014). All morphometric  
201 variables were normally distributed, so we designated each in turn as the response variable in  
202 a GLM with gaussian error distributions, and used t values to determine any association  
203 between clinical signs and morphometrics. All birds included in the analysis were adults (i.e.  
204 hatched the previous calendar year or before), with fully grown wings and not in active wing  
205 moult, and we also included in the analysis morphometrics from apparently healthy birds  
206 (that all tested positive for infection by the *T. gallinae* parasite: Lennon *et al.*, 2013; Dunn,  
207 unpubl data) captured during 2011 (n=7) and 2013 (n=14) and measured by JCD. A subset of  
208 birds was subsequently sexed by behavioural observations (through a combination of  
209 observations of purring males, and nest attendance, whereby male Columbiformes incubate  
210 during the middle of the day, and females overnight and during early morning and evening;

211 e.g. Thorsen *et al.*, 2004) but we did not include sex in the statistical model due to incomplete  
212 data.

213

## 214 **RESULTS**

215 Oral swabs were obtained from 18 adult and seven nestling Turtle Doves during May - July  
216 2012 (n=25; for full details of data collected from each bird see Table 1). In total 13 nests  
217 were monitored, eight of which were depredated prior to hatching. Of the five nests  
218 monitored to nestling stage (full details in Table 1); three nestlings from three nests were  
219 subsequently found dead. *T. gallinae* parasites were cultured from swabs taken from all  
220 nestlings post-mortem, although a full necropsy could only be carried out on two of these due  
221 to the state of decomposition and autolysis. One additional very small nestling (18.9 g  
222 compared to mean weight of  $75.77 \pm 3.82$  g at 7 days (n=11, including data from 2011; Dunn  
223 unpubl. data) disappeared, and was assumed to have died. A further two nests had three  
224 nestlings between them which were monitored to 7 days old: one nestling was depredated  
225 prior to fledging but the remaining two fledged successfully.

226

227 Swabs taken from all 25 Turtle Doves tested positive for *T. gallinae*. Of these, three adults  
228 showed clinical signs of trichomonosis, with regurgitated saliva staining the feathering  
229 around the beak. A subset of 12 adults, including two of these clinically affected birds (the  
230 third was caught in May, prior to the start of radio-tagging) were radio-tagged, flew strongly  
231 upon release, and were subsequently relocated. Only the clinically affected birds are  
232 considered further here. Bird 20 (Table 1) was relocated alive on the ground ~90 m from the  
233 capture site at approximately 09:00 on the day following capture (at 19:00). The bird  
234 appeared to be gasping for breath, made no attempt to escape capture by hand and died  
235 shortly afterwards. Bird 21 (Table 1) was relocated ~190 m from the capture site at

236 approximately 10:00 on the morning following capture (at 16:30). We believe that this bird  
237 was predated as the carcass had been plucked, making it likely that a raptor was responsible.  
238 However, it was impossible to distinguish with certainty between predation and post mortem  
239 scavenging. Individuals with clinical signs were lighter and had shorter wings (Table 2),  
240 showing no overlap with non-indicative individuals (Fig 1a). There was no difference in other  
241 skeletal measures of size (Fig 1b; Table 2).

242

243 Gross necropsies were carried out on five independent individuals as detailed in Table 1.

244 Both Turtle Dove nestlings displayed clinical signs of trichomonosis with a swollen head and  
245 eyes and visible lesions in the buccal cavity and oropharynx (Figures 2a and 2b). One adult  
246 female Turtle Dove was severely emaciated with caseous lesions found blocking the  
247 oropharynx (preventing the bird from swallowing any seed) the location and extent of which  
248 can be seen in Figure 2c. We were unable to suggest cause of death for the second adult  
249 turtle carcass recovered due to the paucity of remains. In contrast to the Turtle Doves  
250 examined, the Red-legged Partridge had no visible lesions within the buccal cavity or upper  
251 respiratory tract, although an oral swab taken from the dead bird tested positive for *T.*  
252 *gallinae* parasites. A caseous trichomonosis lesion was found to have originated within the  
253 proventriculus, grown through the wall and fused to a lobe of the liver resulting in the  
254 necrosis of the connecting tissue and discolouration (Figure 2d).

255

256 Sequences in both directions were obtained from the 25 individuals screened; however,  
257 sequence quality from 6 individuals was too poor to give meaningful data (Table 1). Two  
258 identical sequences were obtained from lesions and oral swabs from three individuals (IDs  
259 20, 22 and 26: Table 1). Overall four distinct sequences were obtained; the most common  
260 sequence (JN007005.1: 100% query coverage and 100% max identity) was isolated from 16  
261 individuals, including all birds displaying clinical signs, all dead Turtle Doves (adults and

262 nestlings), and the Red-legged Partridge (Table 1). Three sequences were isolated from water  
263 sources and one sequence from a feed site, which all matched Genbank sequence JN007005.1  
264 (100% query coverage and 100% max identity; Table 1). Sequences from two individuals  
265 matched sequence FN433475.1 (100% query coverage and 100% max identity), and  
266 sequences isolated from one individual each matched Genbank sequences AJ784785.1 (99%  
267 query coverage and 98% max identity) and FN433473.1 (99% query coverage and 100% max  
268 identity; Table 1).

269

## 270 **DISCUSSION**

271 We report the first confirmation of mortality in free-living Turtle Doves with clinical signs of  
272 trichomonosis. We found a 100% rate of infection by *T. gallinae* in the 25 live Turtle Doves  
273 screened during 2012. This is higher than during the previous year (n=14; Lennon *et al.*  
274 2013), and combined with previous data gives an overall infection rate of 95% (n=39) across  
275 sites separated by up to 120 km. The only two individuals apparently negative for *T. gallinae*  
276 infection were two nestlings from the same nest in 2011 (Lennon *et al.* 2013).

277

278 The overall rate of *T. gallinae* infection appear unusually high when compared to other  
279 Columbidae (e.g. 19% in Spotted Dove *Streptopelia chinensis* and 59% in Zebra Dove *Geopelia*  
280 *straita*, Bunbury *et al.* 2007; 5.6% in Mourning Doves *Zenaida macroura*, Schulz *et al.* 2005;  
281 34.2% in wintering Wood Pigeons *Columba palumbus*, Villanúa *et al.* 2006), with only Rock  
282 Pigeons *Columba livia* documented as having similarly high rates of infection (92%: Sansano-  
283 Maestre *et al.* 2009). Sub-clinical infection can impact on survival: for example, Pink Pigeons  
284 testing positive for *T. gallinae* infection were 13% less likely to survive for a further two years  
285 after screening than those testing negative (Bunbury *et al.* 2008a). Usually, only a very small  
286 percentage of individuals infected by *T. gallinae* display clinical signs (e.g. 0.37% of

287 Columbidae, Sansano-Maestre *et al.* 2009; 1.9% of Pink Pigeons, Bunbury *et al.* 2008a).

288 However, we report clinical signs in 28% of individuals infected by *T. gallinae* parasites (three  
289 adults and four nestlings).

290

291 We found a minimum of four strains of *T. gallinae* within Turtle Doves: as we only sequenced  
292 the ITS/5.8S/ITS2 region, we acknowledge that we may be observing more than one strain  
293 that is genetically different at other functional genes. However, for clarity we subsequently  
294 refer to each of our four strains as a single strain. All fatal cases of trichomonosis were linked  
295 to the same strain of *T. gallinae* found at our study sites in both Turtle Doves and  
296 Woodpigeons (Lennon *et al.* 2013), which was also isolated from the only Turtle Dove  
297 showing clinical signs during 2011 (a nestling that was predated prior to fledging; Lennon *et*  
298 *al.* 2013). This strain falls within the same clade as *T. gallinae* strain A (Lawson *et al.* 2011a;  
299 Lennon *et al.* 2013; Chi *et al.* 2013) and is identical at the ITS/5.8S/ITS2 region to the  
300 causative agent of the finch trichomonosis epizootic (Robinson *et al.* 2010; Ganas *et al.* 2014).

301 The clade contains strains found in Columbidae worldwide, raptors in Spain, and finches in  
302 the USA and UK, suggesting inter- and intra-specific transmission. Further PCR work is  
303 required to determine whether or not this strain is identical to the epizootic strain reported in  
304 finches (Robinson *et al.* 2010; Lawson *et al.* 2011), by examining other functional genes such  
305 as the iron hydrogenase gene (Lawson *et al.* 2011a; Lennon *et al.* 2013).

306

307 Necropsies carried out on intact Turtle Dove carcasses (one adult, two nestlings) confirmed  
308 trichomonosis as the cause of death and identified large oropharyngeal lesions. Molecular  
309 testing of DNA extracted from the lesions confirmed the gross necropsy diagnoses. . Adult 20  
310 was severely emaciated, but in contrast adult 21 had substantial muscle reserves over the  
311 sternum suggesting that this bird might have been at an earlier stage of infection, although the  
312 paucity of remains did not allow us to establish this with any certainty. The observation of

313 clinical trichomonosis in adult and nestling Turtle Doves is, to our knowledge, the first  
314 suggestion of mortality associated with trichomonosis caseous lesions in this species. Whilst  
315 we did not screen for other pathogens and cannot rule out the possibility of co-infection  
316 increasing susceptibility to *T. gallinae*, the final cause of death was believed to be due to *T.*  
317 *gallinae* lesions. Controlled experimental infections in the absence of co-infecting pathogens  
318 would be necessary to confirm trichomonosis as causing mortality.

319  
320 That individuals showing clinical signs of disease were considerably lighter than those  
321 without is not unexpected: *T. gallinae* lesions constrict the oesophagus and prevent affected  
322 birds from ingesting food, resulting in decreased weight. However, the difference in wing  
323 lengths is marked, with no overlap between the wing lengths of individuals with and without  
324 clinical signs, and a mean 3.47% reduction in the wing length of individuals with clinical signs  
325 compared to those without. Our sample size of birds showing clinical signs is small, and thus  
326 our results should be treated with some caution. There were no differences in any skeletal  
327 measures of size, suggesting that infection may impact upon wing length during moult on  
328 wintering grounds in Africa through competition for energetic resources, rather than smaller  
329 birds simply being more susceptible to infection. Such a mechanism has been proposed  
330 previously in other host-parasite systems, with *Haemoproteus* and *Plasmodium* spp. (Marzal *et*  
331 *al.* 2013), *Haemoproteus* spp. (Dunn *et al.* 2013), *Leucocytozoon* spp. (Hatchwell *et al.* 2001)  
332 and *Trypanosoma* spp. (Rätti *et al.* 1993) posited to reduce feather length through competition  
333 for host resources during moult. Turtle Doves are Europe's only sub-Saharan migrant  
334 Columbidae and undergo a partial post-breeding moult prior to migration, completing their  
335 moult on the African wintering grounds (Baker 1993). Thus, individuals with clinical signs  
336 during summer 2012 may have acquired infections on, or en route to/from, their wintering  
337 grounds, or even during the previous breeding season, highlighting the need to further

338 understand the dynamics of *T. gallinae* infection throughout the annual cycle of migratory  
339 species.

340

341 The finding of a moribund Red-legged Partridge, and subsequent suggestion of the same  
342 strain of *T. gallinae* causing markedly different pathology (through isolation of the parasite  
343 from the lesion) is interesting. Previous work had discounted the possibility of parasite  
344 spillover between Columbidae and introduced Galliformes such as Red-legged Partridges and  
345 common Pheasants *Phasianus colchicus* (Lennon *et al.* 2013), as Galliformes tend to be  
346 infected by *T. gallinarum*, which is genetically distinct from *T. gallinae* (e.g Pennycott 1998).  
347 However, our findings suggest that such a parasite spillover may potentially occur. This  
348 suggests that screening of Galliformes may be worthwhile in order to establish whether  
349 parasite spillover between Columbidae and Galliformes – and potentially Passerines - is a  
350 possible occurrence at shared food resources such as game bird feeders or grain spills in  
351 farmyards. Such parasite transfer may occur potentially through a similar mechanism to that  
352 suggested by Lawson *et al.* (2012) for the putative parasite spillover from Columbidae to  
353 Passerines.

354

355 The same predominant single strain of *T. gallinae* isolated from the moribund Turtle Doves  
356 and Red-legged Partridge was also isolated from both a farmyard grain pile and three artificial  
357 water sources at one of our sites. Food and water sources have previously been postulated as  
358 potential vectors for transfer of *T. gallinae* parasites (Kocan 1969), although Bunbury *et al.*  
359 (2007) found no positive grain samples, and only 2 out of 15 water samples were positive for  
360 trichomonads. Whilst speculative, the unusually wet summer of 2012 may have allowed  
361 parasites to survive for longer on damp grain piles (Kocan 1969; Erwin *et al.* 2000) meaning  
362 that individual birds may have been subjected to high and repeated doses of *T. gallinae*  
363 parasites from repeat visits to infected food and water sources. Further work should examine

364 the survival of parasites in food and water sources in these settings to gauge natural infection  
365 rates in relation to the density of potential hosts, and weather-related factors.

366

367 Turtle Dove populations in NW Europe have been declining for decades and continue to do so.  
368 Whilst a previous intensive study of this species on UK breeding grounds found no evidence of  
369 disease-related issues (S. Browne, pers. 160mm.), no historic data on infection rates are  
370 available. The species has also undergone a dietary switch in the UK, from the seeds of arable  
371 plants (Murton *et al.* 1964) to anthropogenic seed resources such as grain piles in farmyards  
372 (Browne and Aebischer 2003). Food stress can decrease immune function (Lindström *et al.*  
373 2005) and induce chronic stress in birds (Clinchy *et al.* 2004), potentially increasing  
374 susceptibility to infection and the likelihood of clinical signs and this possibility cannot be  
375 negated within this system. More likely, however, is that the dietary switch undergone by this  
376 species has led to an increased risk of intra- and inter-species transference of directly and  
377 indirectly-transmitted parasites and pathogens, such as *T. Gallinae*, at a restricted number of  
378 food resources shared by birds feeding at high densities (e.g. Höfle *et al.* 2004; Lawson *et al.*  
379 2012).

380

381 Historically, the anti-protozoal dimetridazole, or Emtryl, was widely used as a prophylactic  
382 feed additive for game birds reared for sporting purposes, however, since its withdrawal,  
383 concerns have been raised about the potential impacts of motile protozoans on a wide range  
384 of species, mostly captive-reared birds (Dernburg *et al.* 2005; Callait-Cardinal *et al.* 2007). To  
385 our knowledge, no literature is available examining any trends in infection rates of  
386 trichomonads in captive-reared game birds during the period since Emtryl withdrawal,  
387 although Lennon *et al.* (2013) found higher rates of trichomonad infection in Columbidae on  
388 farms with game bird feeding than on farms without, and Höfle *et al.* (2004) suggest that the  
389 supplementary feeding of game birds constitutes a risk factor for the appearance of

390 trichomonosis outbreaks in wild birds. We suggest that the potential for parasite transfer  
391 from non-native game birds to rapidly declining native species is worthy of further  
392 investigation. Supplementary feeding of game and wild birds, especially during the late  
393 winter period when seed food is scarce, is widespread. Although turtle doves are summer  
394 migrants and therefore not present in Europe during the winter, given the results presented  
395 here, and the recent finch trichomonosis epizootic (Robinson *et al.* 2010), we suggest  
396 stringent hygiene precautions when deploying supplementary food are needed throughout  
397 the year to reduce the risk of disease transmission. These include strict adherence to  
398 guidelines to only distribute enough food to match consumption, ensure a fresh supply of food  
399 is maintained without leaving seed unconsumed and rotating feeding sites. (e.g. Natural  
400 England 2012).

401

402 Our work highlights the importance of continued monitoring of *T. gallinae* infection in Turtle  
403 Doves and of monitoring sub-clinical infection in free-living populations rather than relying  
404 on morbidity and mortality reports alone, particularly for species where the population status  
405 gives cause for conservation concern. Further work should address the epidemiology of  
406 infection, as well as establishing any sub-clinical impacts of infection that may impact on  
407 ecological parameters such as reproductive success. *T. gallinae* is thought to be a population-  
408 limiting factor in the Pink Pigeon, despite observed pathogenicity being low (Bunbury *et al.*  
409 2008a). Unless Turtle Dove feeding ecology changes to allow a reduction in infection rates,  
410 parasite infection may potentially amplify the existing reduction in reproductive output and  
411 either hasten the ongoing population decline or prevent population recovery. Greater uptake  
412 of measures that provide abundant and accessible food (e.g. fallows, seed mixes or cultivated,  
413 uncropped margins), which are available in many European agri-environment schemes,  
414 would provide birds with more dispersed feeding opportunities and thus potentially reduce  
415 disease transmission.

416

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424

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564

565 Table 1. Summary of data collected from individual birds.

ID	Outcome	Species	Age	<i>T. gallinae</i> source <sup>a</sup>	Post mortem	
1- 16	Live	Turtle Dove	Adults	1	No	JN007005.1 (n=8) FN433475.1 (n=1) FN433473.1 (n=1) AJ784785.1 (n=1) No sequence (n=5)
17 - 18	Live	Turtle Dove	Nestling (nest 1)	1	No	JN007005.1 (n=1) FN433475.1 (n=1)
19	Predated	Turtle Dove	Nestling (nest 2)	2	No	JN007005.1
20	Died	Turtle Dove	Adult	1, 4	Yes	JN007005.1
21	Predated/Died	Turtle Dove	Adult	1	Yes	JN007005.1
22	Died	Turtle Dove	Nestling (nest 3)	3, 4	Yes	JN007005.1
23	Disappeared (assumed died)	Turtle Dove	Nestling (nest 3)	2	No	No sequence
24	Died	Turtle Dove	Nestling (nest 4)	3	No	JN007005.1
25	Died	Turtle Dove	Nestling (nest 5)	3	Yes	JN007005.1
26	Died	Red legged partridge	Adult	3,4	Yes	JN007005.1

566

567 <sup>a</sup> *T. gallinae* source: 1: swab collected from crop, throat and oral cavity whilst alive; 2: swab  
568 collected from oral cavity only; 3: swab collected post mortem; 4: DNA extracted directly from  
569 lesion.

570

571

572 Table 2. Summary of morphometrics for adult Turtle Doves with and without clinical signs of  
573 trichomonosis.

Measurement	Mean $\pm$ 1 SE		Statistics		
	Clinical signs (n=3)	No clinical signs (n=31)	t	df	p
<b>Weight (g)</b>	<b>121.40 <math>\pm</math> 2.93</b>	<b>161.06 <math>\pm</math> 1.92</b>	<b>-6.276</b>	<b>1</b>	<b>&lt;0.001</b>
<b>Wing length (mm)</b>	<b>172.67 <math>\pm</math> 0.83</b>	<b>179.45 <math>\pm</math> 0.59</b>	<b>-3.493</b>	<b>1</b>	<b>0.001</b>
Head-beak length (mm)	46.57 $\pm$ 0.92	46.23 $\pm$ 0.15	0.623	1	0.538
Tarsus length (mm)	23.23 $\pm$ 1.17	23.52 $\pm$ 0.19	-0.416	1	0.680

574

575

576 Figure 1. a) Wing length and weight distributions and b) head-beak and tarsus length  
577 distributions from adult turtle doves with clinical signs compared to female, male and  
578 unsexed adults with no clinical signs.

579 a)

580

581 b)

582

583 Figure 2. Photographs from post-mortems of A) nestling Turtle Dove 25, B) nestling Turtle  
584 Dove 22, C) adult Turtle Dove 20, and D) Red-legged Partridge 26. Arrows show  
585 oropharyngeal lesions in Turtle Doves and a lesion originating in the proventriculus in the  
586 Red-legged Partridge.

587 A.

588

589 B.

590

591 C.

592

593 D.

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