

Molecular characterization of Trichomonas gallinae isolates recovered from the Canadian Maritime provinces' wild avifauna reveals the presence of the genotype responsible for the European finch trichomonosis epidemic and additional strains.

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1 Molecular characterization of *Trichomonas gallinae* isolates recovered from the Canadian Maritime provinces' wild avifauna reveals the presence of the genotype 2 responsible for the European finch trichomonosis epidemic and additional 3 strains. 4 Scott McBurnev^{1,2}, Whitnev K, Kellv-Clark^{1,2}, María J, Forzán^{1,2}, Becki Lawson³, Kevin 5 M. Tyler⁴ and Spencer J. Greenwood⁵ 6 7 1- Department of Pathology & Microbiology, Atlantic Veterinary College, University of 8 Prince Edward Island, 550 University Avenue, Charlottetown, PE, C1A 4P3, Canada 9 Prince Edward Island, Canada 10 2- Canadian Wildlife Health Cooperative, Atlantic Region, Atlantic Veterinary College, 11 University of Prince Edward Island, 550 University Avenue, Charlottetown, PE, C1A 12 13 4P3, Canada Prince Edward Island, Canada 🥏 3- Institute of Zoology, Zoological Society of London, Regents Park, London, NW1 4RY, 14 UK 15 4- Norwich Medical School, University of East Anglia, Norwich, NR4 7TJ, UK 16 5- Department of Biomedical Sciences, Atlantic Veterinary College, University of Prince 17 Edward Island, 550 University Avenue, Charlottetown, PE, C1A 4P3, Canada Prince 18 Edward Island, Canada 19 20 **Running title:** Molecular characterization of *T. gallinae* in Canadian wild avifauna 21 Corresponding author: Spencer J. Greenwood, Tel: 902-566-6002, Fax: 902-566-22 0832, e-mail: sgreenwood@upei.ca 23

24 **Summary** (150-200 words)

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26 Finch trichomonosis, caused by *Trichomonas gallinae*, emerged in the Canadian Maritime provinces in 2007 and has since caused ongoing mortality in regional purple 27 finch (Carpodacus purpureus) and American goldfinch (Carduelis tristis) populations. 28 Trichomonas gallinae was isolated from (1) finches and rock pigeons (Columbia livia) 29 submitted for post mortem or live-captured at bird feeding sites experiencing 30 31 trichomonosis mortality; (2) bird seed at these same sites; and (3) rock pigeons live-32 captured at known roosts or humanely killed. Isolates were characterized using internal transcribed spacer (ITS) region and iron hydrogenase (Fe-hyd) gene sequences. Two 33 34 distinct ITS types were found. Type A was identical to the UK finch epidemic strain and was isolated from finches and a rock pigeon with trichomonosis; apparently healthy rock 35 pigeons and finches; and bird seed at an outbreak site. Type B was obtained from 36 37 apparently healthy rock pigeons. Fe-hyd sequencing revealed six distinct subtypes. The predominant subtype in both finches and the rock pigeon with trichomonosis was 38 identical to the UK finch epidemic strain A1. Single nucleotide polymorphisms in Fe-hyd 39 sequences suggest there is fine-scale variation amongst isolates and that finch 40 trichomonosis emergence in this region may not have been caused by a single spill-41 42 over event.

Keywords: *Trichomonas gallinae*, trichomonosis, genotype, ITS, Fe-hydrogenase,
subtype, finch, pigeon

45	Key F	Findings (3-5 bullets of < 90 characters each, including spaces)
46	-	Two T. gallinae ITS sequence types found in the Canadian Maritime provinces'
47		avifauna
48	-	T. gallinae ITS sequence type in Canadian finches identical to UK finch epidemic
49		strain
50	-	Bird seed from an outbreak yielded <i>T. gallinae</i> with the UK finch epidemic strain
51		ITS sequence
52	-	Fe-hyd gene sequencing revealed fine-scale variation with six <i>T. gallinae</i>
53		subtypes
54	-	Fe-hyd subtype of the UK finch epidemic strain was predominant in Canadian
55		finches
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63 Introduction

Trichomonas gallinae is a protozoan parasite which commonly infects the upper 64 digestive tract of columbids (*i.e.*, pigeons and doves) and birds of prey (*i.e.*, eagles, 65 hawks and owls) and less frequently can also infect a variety of other avian taxa 66 including passerines (such as finches and sparrows) (Forrester and Foster; 2009; Amin 67 et al. 2014). In 2005, trichomonosis was first recognized as an emerging infectious 68 69 disease of wild finches in Great Britain (GB) (Pennycott et al. 2005; Lawson et al. 2006). The species affected in the summer/autumn seasonal epidemic were primarily 70 greenfinch (Chloris [Carduelis] chloris) and chaffinch (Fringilla coelebs). Although pre-71 72 existing sporadic reports of disease in free-ranging finches do exist, the 2005-2006 (and on-going) outbreak is the first reported instance of large-scale epidemic mortality due to 73 74 trichomonosis in any passerine species (Lawson et al. 2012). In the years following the initial outbreak in the western and central counties of England and Wales, finch 75 trichomonosis spread to eastern England (2007) and then to southern Fennoscandia 76 (2008) and Germany (2009): epidemiological and historical banding return data 77 supported chaffinch migration as the most likely mechanism of the observed pattern of 78 disease spread (Neimanis et al. 2010; Robinson et al. 2010; Lawson et al. 2011a, 79 80 Peters et al. 2009). The disease range of finch trichomonosis has continued to extend further eastward within continental Europe and had reached Austria and Slovenia by 81 2012 (Ganas et al. 2014). Concurrently, finch trichomonosis spread westward from 82 Britain with finch mortality incidents reported in Northern Ireland from 2006 and in the 83 Republic of Ireland from 2007 (Lawson et al. 2012). 84

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85 In the late summer/early autumn of 2007, trichomonosis was first recognized in the purple finch (Carpodacus purpureus) populations of Nova Scotia, Canada (Forzán 86 et al. 2010). In the following summer and autumn, the Canadian Wildlife Health 87 Cooperative (CWHC), Atlantic region, confirmed additional Canadian mortalities from 88 trichomonosis in the purple finch populations of Nova Scotia and Prince Edward Island 89 (PEI) and in American goldfinch (Carduelis tristis) populations of New Brunswick 90 (Forzán et al. 2010). In 2009, the CWHC confirmed finch trichomonosis incidents in all 91 three Canadian Maritime provinces during the same seasons, and diagnosed the 92 disease in a new species, the pine siskin (Carduelis pinus) (CWHC unpublished data). 93 The diagnosis of trichomonosis in the Canadian Maritime provinces in the summer and 94 autumn of three consecutive years and the infection of multiple finch species not known 95 to be previously affected by the disease, suggests that finch trichomonosis is an 96 emerging disease in this region. Prior to the emergence of trichomonosis in the 97 passerine bird populations of the Canadian Maritime provinces, this disease was not 98 diagnosed in any of the region's wild avian species since the CWHC, Atlantic Region, 99 began collecting diagnostic wildlife health data in 1992. While it is assumed that T. 100 101 gallinae is present in the columbid populations of the Canadian Maritime provinces due to the parasite's ubiquitous distribution in wild pigeon and dove populations worldwide 102 (Amin et al. 2014), to our knowledge reports of T. gallinae in the region's columbid 103 104 populations have not been documented. Lastly, it is noteworthy that the Canadian Maritime provinces represent the eastern limit of North America with closest 105 106 geographical proximity to the UK and that finch trichomonosis emerged in the years 107 immediately subsequent to the onset of epidemic mortality in British finches.

108 The aims of the present study were to firstly investigate the sequence diversity of T. gallinae recovered from finches and columbids from the Canadian Maritime 109 provinces. Secondly, to compare the Canadian T. gallinae sequences with those 110 published from other countries, including isolates from GB. Finally, to provide a 111 description of the temporal, geographical and species-specific variation amongst the 112 113 isolates examined from the Canadian Maritime provinces. Genotyping of isolates was determined by polymerase chain reaction (PCR) and sequencing of the ITS region 114 (5.8S rDNA and flanking internal transcribed spacer regions, ITS1 and ITS2) and the 115 116 hydrogenosomal Fe-hydrogenase (Fe-hyd) gene to evaluate finer scale evolutionary relationships amongst these organisms (Lawson et al. 2011b; Chi et al. 2013). 117

118 Materials and Methods

119 Columbid and passerine capture methods

When suspected finch trichomonosis incidents in the Canadian Maritime 120 121 provinces were reported by members of the public, the CWHC, Atlantic region, facilitated the immediate submission of recently dead passerines by encouraging 122 property owners to submit specimens for a detailed post-mortem examination (PME). 123 124 When PME confirmed trichomonosis (on the basis of gross and microscopic lesions or microscopic lesions alone consistent with trichomonosis with or without a positive 125 culture of *Trichomonas* sp. from upper alimentary tract lesions), the site was visited to 126 live-capture all species of birds present and sample them for Trichomonas sp. by culture 127 (see culture technique below). In addition, food and water sources provided at these 128

sites were individually sampled for *Trichomonas* sp. by culture (see culture techniquebelow).

To investigate the heterogeneity of *T. gallinae* in sympatric columbid populations in PEI, known locations of high columbid population densities were selected for extensive trapping, without pre-existing knowledge of the presence of *T. gallinae* or trichomonosis within these populations.

All birds were captured under a Canadian Wildlife Federation license (permit 135 #SC2707) and Canadian Council on Animal Care guidelines (UPEI protocol #10-020, 136 6003687) by standard methods including mist net, whoosh net, and walk-in box trap. 137 Passerines were captured for this study by using a mist net (Bleitz Wildlife Foundation) 138 139 California, 50D-2 ply mesh, 1¹/₂" mesh, 7' X 42', Stock # 26N-50/2) for two days per location in May-September of 2009 and 2011. Columbid species required extensive 140 time for acclimatization to the box trap, and as a result, columbids were ground trapped 141 142 at each site on multiple days sometimes occurring over a period of several weeks in May-December of 2009 and 2010. A ground box trap (Safeguard single compartment 143 pigeon trap, 28"L x 24"W x 8"H, with eight entry doors, and a capacity to hold up to 30 144 birds) was baited with bird seed for a minimum of 12 days prior to commencing 145 trapping. It is important to note that during the allotted baiting period, all regular 146 supplementary feed sources were removed from the property to ensure the birds fed in 147 the baited area. Mourning dove, another target species, were difficult to capture with the 148 ground box trap so their capture was also facilitated by use of a whoosh-net (Hawkseye 149 150 Nets Virginia Beach, VA, USA - 2 1/8" mesh, 23' whoosh net). Similar to the box-trap

protocol, the area over which the whoosh-net was fired was baited with bird seed for a
minimum of 5 days prior to attempted capture.

Birds captured by all methods were sexed and aged by plumage (hatch year or 153 adult) when possible, weighed, banded and examined for clinical signs consistent with 154 trichomonosis such as the typical oropharyngeal lesions, fluffed up feathers, saliva on 155 the face, food at the commissures of the beak or matted in the feathers of the head or 156 chest and/or reluctance or inability to fly. If *T. gallinae* was isolated from a bird with no 157 clinical evidence of trichomonosis it was designated as "apparently healthy". If T. 158 gallinae was isolated from a bird with clinical signs consistent with trichomonosis, the 159 infection was defined as a clinical case of trichomonosis. Opportunistic sampling was 160 also undertaken for Trichomonas sp. by culture of wild passerines and columbids 161 162 admitted to the Atlantic Veterinary College Teaching Hospital and of rock pigeons that were humanely killed during removal from cattle barns in the winter months on PEI. 163

164 Trichomonad culture

Prior to swabbing live birds, the end of a sterile calcium-alginate cotton-tipped 165 swab with an aluminum shaft (Puritan™ Medical, Fisher Scientific, Canada, catalogue 166 number 22-029-501) was bent into a gentle curve representing ~ 120 ° angle to match 167 168 the natural anatomical curvature of the oral cavity as it opens into the esophagus. The 169 distance between the start of the curve and cotton tip was equivalent to the distance between the oral cavity and crop, and the positioning of this curve depended on the 170 species of bird. Anatomically, bending the swab at the 120° angle facilitated the 171 172 movement of the swab from the oral cavity to the crop. After bending, the swab was

173 moistened with sterile saline and gently inserted into the oral cavity of the bird by pushing the tip against a commissure of the beak. The swab was slowly and gently 174 advanced into the esophagus to the level of the crop while allowing the bird to swallow. 175 Due to the thinness of the esophageal and ingluvial walls in passerine birds this 176 procedure was done with extreme caution and only by experienced individuals to avoid 177 178 iatrogenic damage. Once in the crop, the swab was gently rotated, moved up and down and removed. Care was taken to swab any visible oropharyngeal trichomonosis lesions. 179 The crop and lesions of dead birds were swabbed once the upper digestive tract was 180 181 opened for PME. After collection, all swabs were used to immediately inoculate an InPouch TF[™] test medium kit (BioMed Diagnostics, White City, OR, USA) on-site prior 182 to transport back to the laboratory for incubation at 37°C and daily monitoring for 10 183 days. If the site was not in the province of PEI, the samples were placed in a Hova-184 Bator egg incubator (circulated air model no. 2362N, 20.3 watt, 115 volt AC, 185 G.Q.F.MFG. Co. Inc. Savannah, GA) set at 37°C for transport to the laboratory at the 186 University of Prince Edward Island. 187

Bird seed and water sources at sites experiencing trichomonosis mortality were independently swabbed. The swabs were used to immediately inoculate an InPouch TF[™] test medium kit on-site prior to transport back to the laboratory for incubation at 37°C and daily monitoring for 10 days. If the site was not in the province of PEI, the samples were placed in a Hova-Bator egg incubator for transport to the laboratory at the University of Prince Edward Island as described above.

194 Parasite culture and cryopreservation

195	Parasite cultures were monitored daily using a double chamber hemacytometer,
196	counts of motile trichomonads were performed on both grids and if results did not
197	correlate within 10%, the process was repeated and the average of the four counts was
198	taken instead of the two. Once parasites reached mid-log phase, they were
199	cryopreserved by adding 100 μ l of 100% glycerol to 1ml of the parasite culture. This total
200	volume was subdivided into four separate 500µl aliquots and stored in liquid nitrogen.
201	An additional 1ml aliquot of the original parasite culture was collected to be used for
202	DNA extraction.

203 PCR for ITS region and Fe-hyd gene regions

Trichomonas sp. DNA was obtained from culture isolates using a QIAamp DNA 204 Mini Kit (QIAGEN, Toronto, ON, Canada) as per the manufacturer's instructions for cell 205 cultures. DNA extracts of 42 isolates (Table 1) were examined using PCR protocols 206 specific for the ITS1/5.8S rRNA/ITS2 region (subsequently referred to as the ITS region) 207 and Fe-hyd gene. DNA amplification of the ITS region (~ 300 bp) was performed using 208 trichomonad-specific primers TFR1 (5'-TGCTTCAGTTCAGCGGGTCTTCC-3') and 209 210 TFR2 (5'-CGGTAGGTGAACCTGCCGTTGG-3') (Felleisen 1997) while amplification of the Fe-hyd gene (~ 900 bp) used the primers TrichhydFOR (5'-211 GTTTGGGATGGCCTCAGAAT-3') and TrichhydREV (5'-212 213 AGCCGAAGATGTTGTCGAAT-3') (Lawson et al. 2011b; Chi et al. 2013). Each PCR reaction mixture contained 12.5µl Amplitaq Gold Master Mix (Applied Biosystems, Life 214 215 Technologies, Burlington, ON, Canada), 4.5µl nuclease free water, 2.5µl forward primer 216 $(10\mu M)$, 2.5µl reverse primer $(10\mu M)$ and 3µl of undiluted target DNA and was

217 performed in duplicate. For each reaction, negative controls substituted target DNA with 3µl of nuclease-free water and positive controls used 3µl of T. gallinae DNA (purple 218 finch isolate from Forzán et al. 2010; parasite species confirmed by sequencing the ITS 219 region) and T. gallinae DNA from a British greenfinch (species confirmed by sequencing 220 the Fe-hyd gene) respectively. PCR parameters for the ITS region amplification were 221 94°C for 2 minutes, followed by 40 cycles of 94°C for 30 seconds, 67°C for 30 seconds, 222 72°C for 2 minutes and a final extension at 72°C for 15 minutes. PCR parameters for 223 Fe-hyd gene amplification were 94°C for 15 minutes, followed by 35 cycles of 94°C for 1 224 minute, 66°C for 30 seconds, 72°C for 1 minutes and a final extension at 72°C for 5 225 minutes. PCR amplicons were then examined via 1% agarose gel electrophoresis with 226 ethidium bromide. 227

228 DNA sequencing and phylogeny reconstruction

PCR products were sequenced in both directions at the McGill University and 229 Genome Québec Innovation Centre, Montréal, Québec, Canada. Sequences were 230 aligned with published trichomonad sequences from GenBank using BioEdit (Hall 231 232 1999). Phylogenies were constructed separately for the ITS region and Fe-hyd gene by neighbour-joining (NJ), maximum parsimony (MP) and maximum likelihood (ML) 233 234 methods using MEGA version 6.0 (Tamura et al. 2013). Statistical support for NJ, ML, 235 and MP tree topologies were bootstrap-sampled 1,000 times and support values (%) of NJ, MP and ML analysis were superimposed on the NJ consensus trees. 236 For phylogeny reconstruction using the ITS region, NJ tree evolutionary 237

distances were computed using the Jukes-Cantor method (Jukes and Cantor 1969) and

239 were reported in the units of the number of base substitutions per site. The MP tree was obtained using the Subtree-Pruning-Regrafting algorithm (Nei and Kumar 2000) with 240 search level 1 in which the initial trees were obtained with the random addition of 241 sequences (10 replicates). The ML tree was constructed using Jukes-Cantor 242 substitution model (Jukes and Cantor 1969) as determined by the lowest Bayesian 243 244 Information Criterion (BIC) score and highest Akaike Information Criterion, corrected (AICc) value (Tamura et al. 2013). Initial tree(s) for the heuristic search were obtained 245 by applying the NJ method to a pairwise distance matrix estimated using Maximum 246 247 Composite Likelihood (MCL). For ITS region trees, there were a total of 37 nucleotide sequences using 209 positions in the final dataset. All positions containing gaps and 248 missing data were eliminated. Bootstrap values (1000 replicates) for each NJ, MP and 249 ML trees were computed following Felsenstein (1985). 250

For phylogeny reconstruction based on the Fe-hyd gene, the NJ, MP and ML used the same parameters as for the ITS region sequences including bootstrap replicates (1000). For Fe-hyd trees, there were 15 nucleotide sequences with a total of 803 positions in the final dataset. All positions containing gaps and missing data were eliminated.

256 **Results**

257 *Parasite recovery from columbids, finches and environmental samples*

Forty-two trichomonad isolates were collected between 2009 and 2011 from rock pigeons (n = 12), finch species (n = 29) and bird seed (n = 1) from the Canadian Maritime provinces (Table 1 and Figure 1). Thirty-seven live mourning doves were

captured, swabbed and cultured for this study, and none of these individuals were
positive for *T. gallinae*. Additionally, no water samples were positive for *T. gallinae*. In
the individuals that died of trichomonosis, no gross or microscopic lesions consistent
with another disease being the primary problem (e.g., avipoxvirus infection or
salmonellosis) were identified at post mortem or with histopathology.

266 ITS region sequence and phylogeny

ITS region sequences of 300 nucleotides were derived for the 42 267 trichomonad isolates recovered from finches, rock pigeons and from a bird seed sample 268 in the Canadian Maritime provinces. Two distinct ITS region types were recognized that 269 share 98.5% similarity, (1) Sequence Type A (GenBank: KF214772) was identified in 39 270 T. gallinae isolates collected from American goldfinches (n = 7; 5 apparently healthy 271 individuals and 2 with trichomonosis), purple finches (n = 22; 8 apparently healthy 272 individuals and 14 with trichomonosis), rock pigeons (n = 9; 8 apparently healthy 273 individuals and 1 with trichomonosis) and an aggregate of moist bird seed removed 274 from several birdfeeders and deposited in a compost bin at a site confirmed to be 275 276 experiencing finch trichomonosis (n = 1) and (2) Sequence Type B (GenBank: KF214773) was identified in *T. gallinae* isolates from 3 apparently healthy rock pigeons 277 278 (Table 1).

The ITS region phylogeny confirms that the *T. gallinae* isolates formed a monophyletic assemblage within the trichomonads with two well-supported groups, Type A & B (Figure 2). Type A contains 39 PEI isolates from finches and rock pigeons as well as the bird seed sample (GenBank: KF214772) and also representative isolates

including the UK finch epidemic strain (GenBank: GQ150752) and other isolates from
finches, columbids and raptors from Brazil, Europe, Mauritius, Australia and the USA
(Figure 2). Type B contains an additional three isolates derived from PEI rock pigeons
(GenBank: KF214773) with no evidence of trichomonosis, along with representative
isolates derived from columbids, raptors and a canary from diverse geographic regions
including the USA, Europe and Australia (Figure 2).

289 Fe-hyd gene sequence and phylogeny

290 The Fe-hyd nucleotide sequences (901 nucleotides) were obtained from all finch and rock pigeon isolates from the Canadian Maritime provinces (n=41). Multiple 291 attempts to amplify the Fe-hyd gene from DNA extracted from the bird seed sample 292 (isolate 42) were unsuccessful. Six different Fe-hyd sequence subtypes were 293 discovered that share between 98.1-99.8% similarities. The six Fe-hyd subtypes 294 295 identified in the present study are indicated in Table 1. The first subtype (GenBank: KJ184167) included American goldfinch isolates 1-6, purple finch isolates 8, 11-17 and 296 19-29 and rock pigeon isolates 32, 34-35 and 37-38 that were identical to the clonal UK 297 298 finch epidemic strain (GenBank: JF681136, Lawson et al. 2011b). The second subtype (GenBank: KJ184168) included purple finch isolates 9 and 10, while the third subtype 299 300 (GenBank: KJ184169) included American goldfinch isolate 7 and purple finch isolate 18 301 respectively; each subtype differed by one unique single nucleotide polymorphism (SNP) from the UK finch epidemic strain A1. Similarly, the fourth subtype (GenBank: 302 303 KJ184170) from rock pigeon isolate 39 was identical to an isolate from a Madagascar 304 turtle dove (Streptopelia picturata) from the Seychelles (GenBank: JF681141), while the

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305	fifth subtype (GenBank: KJ184171) from rock pigeon isolate 40 differed by one SNP.
306	The sixth subtype (GenBank: KJ184172) included rock pigeon isolates 33, 36 and 41
307	that were identical to an isolate from a wood pigeon (Columba palumbus) from the UK
308	(GenBank: KC529662).
309	The Fe-hyd phylogeny shows two distinct clusters of sequences. Isolates 1-6, 8,
310	11-17, 19-32, 34-35 and 37-38 (GenBank: KJ184167), isolates 7 and 18 from American
311	goldfinch and purple finch (GenBank: KJ184169) and the isolate from purple finches 9
312	and 10 (GenBank: KJ184168) all grouped with the UK finch epidemic strain A1
313	(GenBank: JF681136). The second cluster contains the two PEI rock pigeons isolates
314	39 and 40 (GenBank: KJ184170 and KJ184171 respectively) in a well-supported (98%
315	by all three phylogeny methods) cluster with an isolate from a Madagascar turtle dove
316	from the Seychelles (A2) (Figure 3).

The three other PEI rock pigeon isolates 33, 36 and 41 (GenBank: KJ184172) 317 grouped with a T. gallinae isolate from a wood pigeon from the UK (C4). These 318 sequences along with the remaining *T. gallinae* Fe-hyd gene sequences show a less 319 cohesive branching structure (Figure 3.). 320

Discussion 321

This study utilised ITS region and the Fe-hyd gene sequencing to investigate the 322 genetic diversity of T. gallinae in finch and columbid populations of the Canadian 323 Maritime provinces following the emergence of finch trichomonosis in this region. 324

The ITS region analysis revealed that two *T. gallinae* sequence types are present in the wild avifauna of the Canadian Maritime provinces. In phylogenies based on ITS region sequence data, *T. gallinae* splits into two very distinct groups as noted by previous authors (Gerhold *et al.* 2008; Sansano-Maestre *et al.* 2009; Grabensteiner *et al.* 2010, Lawson *et al.* 20011b).

All finch isolates in this study, whether they originated from apparently healthy 330 birds or birds with trichomonosis, were identical to the *T. gallinae* Type A that has been 331 previously identified in European finches and is widespread in North American 332 columbids (Gerhold et al. 2008; Lawson et al. 2011b; Girard et al. 2014). Importantly, 333 this same type was identified in nine rock pigeons (eight apparently healthy individuals 334 and one with trichomonosis) (Table 1). Thus, the ITS region sequence typing alone 335 cannot discriminate whether the origin of trichomonosis in finches in the Canadian 336 337 Maritime provinces is a translocation of the European finch strain or is simply the result of contact with infected sympatric columbids. However, because both American 338 goldfinch and purple finch populations in the Canadian Maritimes are considered local 339 340 resident populations with limited distance North-South migrations (mainly associated with weather conditions and food availability) and rock pigeons are non-migratory year-341 round residents, a plausible scenario for transmission between these species at local 342 bird feeding stations is reasonable without requiring movement of the disease from 343 Europe to the Canadian Maritime provinces. 344

A common factor in the emergence of trichomonosis in finches in all geographical locations is that the mortality is identified where large numbers of birds congregate at private birdfeeding and watering stations (Forzán *et al.* 2010; Neimanis *et al.* 2010;

348 Robinson et al. 2010). Therefore, it has been suggested that indirect transmission associated with contaminated bird seed, water bowls, or bird baths plays a role in the 349 epidemiology of this disease (Boal et al. 1999; Neimanis et al. 2010; Robinson et al. 350 2010; Gerhold et al. 2013). In the present study, T. gallinae was not detected in water 351 collected from sites where trichomonosis mortalities were occurring. This was surprising 352 given that Bunbury et al. (2007) were successful in recovering T. gallinae from puddles 353 and Gerhold et al. (2013) found that T. gallinae was able to survive for up to 20 minutes 354 in both distilled and chlorinated water when organic matter (detritus, leaves and soil) 355 356 was present. One caveat to our water sampling success was that property owners undergoing bird mortalities in their backyards became more diligent in cleaning feeders 357 and waterers. Thereby reducing the likelihood of recovering parasites from water 358 samples collected in our study. In support of this fact, the only successful isolation of T. 359 gallinae from bird seed was from a composite sample disposed of in a compost bin at a 360 property experiencing trichomonosis mortality. This isolation supports the experimental 361 evidence that showed T. gallinae can survive in moist grain for 120 hours (Kocan 1969). 362 Furthermore, ITS typing confirmed that the bird seed isolate was Type A, identical to T. 363 gallinae isolates recovered from sick birds on the same property. 364

Interestingly, we also identified three rock pigeons infected with Type B *T*. *gallinae*, a type that has been reported in columbids from the USA, eastern Spain and Austria as well as in raptors from eastern Spain (Gerhold *et al.* 2008; Sansano-Maestre *et al.* 2009). In a prevalence study of *T. gallinae*, Sansano-Maestre *et al.* (2009) examined pigeons and raptors with gross lesions consistent with trichomonosis and apparently healthy birds with no identifiable lesions and found that Type A *T. gallinae*

371 were recovered more frequently from birds with gross lesions of trichomonosis, whereas Type B T. gallinae were recovered from individuals with no lesions, suggesting a 372 relationship between Type A and increased virulence. Sansano-Maestre et al. (2009) 373 also speculated that Type B parasites may be adapted to pigeon hosts as this Type was 374 much more prevalent in pigeons than in raptors. Similar to Sansano-Maestre et al. 375 (2009) study, we found that all Type B isolates were recovered from apparently healthy 376 rock pigeons, and all finch species and rock pigeons with evidence of clinical 377 trichomonosis were infected with Type A. However, it is important to note that while all 378 379 isolates recovered from either finches or pigeons with clinical evidence of trichomonosis were Type A, Type A isolates were also recovered from apparently healthy birds. Also, 380 while rock pigeon isolates were not all Type B, all Type B isolates in our study were 381 382 recovered exclusively from rock pigeons, all of which were apparently healthy individuals. Thus our results are consistent with the hypothesis put forward by Sansano-383 Maestre *et al.* that there may be a relationship between Type A and increased virulence. 384 Through examination of multiple gene regions (ITS region, Fe-hyd gene and 385 small sub-unit rDNA), as well as random amplified polymorphic DNA analyses, Lawson 386 et al. (2011b) examined over 50 isolates obtained from finch trichomonosis cases and 387 found no evidence for multiple strains, concluding that a clonal strain of Type A was 388 responsible for the emergence of epidemic trichomonosis in GB. Lawson et al., (2011b) 389 further speculated that due to the clonal nature of the passerine epidemic strain, it most 390 391 likely recently arose from a bottleneck, such as a single spill-over event (*i.e.*, hostswitching) from columbids to sympatric finches. In the present study, ITS region 392 sequence analysis revealed that all Type A isolates from the Canadian Maritime 393

394 provinces were identical to the UK finch epidemic strain. Furthermore, our examination of the Fe-hyd gene also revealed that several finch and rock pigeon isolates were 395 identical to the UK finch epidemic strain (Lawson et al. 2011b). However, it is equally 396 important that Fe-hyd sequence analysis also revealed single nucleotide polymorphisms 397 amongst some of the Canadian Type A isolates. Based on Fe-hyd nucleotide sequence 398 analysis, four Canadian Type A isolates, including American goldfinch, purple finch and 399 rock pigeon isolates, and one Canadian Type B isolate, only from a rock pigeon, were 400 found to be different from both the clonal UK epidemic strain and the Canadian Maritime 401 402 provinces' isolates similar to the clonal UK epidemic strain mentioned above (see Figures 2 and 3). This suggests divergence not only from the British finch and 403 Seychelles columbid strains they were compared to, but also from each other, indicating 404 that a number of strains of T. gallinae are present in the wild avifauna of the Canadian 405 Maritime provinces. Analysis of the Fe-hyd gene sequences from the Canadian 406 Maritime provinces bird isolates showed that there is fine-scale variation amongst 407 isolates akin to that observed in UK columbid populations. This observation suggests 408 that the emergence of finch trichomonosis in this region may have been caused by 409 410 multiple spill-over events, either from sympatric columbids, another bird species as yet unknown to be infected with the parasite or from virulent T. gallinae developing 411 independently within the Canadian Maritime provinces' finch populations. In support of 412 413 this view a recent paper has reported the presence of the UK finch epidemic subtype A1 in North American columbids (Girard et al. 2014) similar to the findings in this study. 414

Indeed, when historic *T. gallinae* DNA samples were subtyped, the A1 subtype
had also been isolated from Mauritian columbids sampled in 2004 (*unpublished data*)

417 suggesting distribution of this subtype may actually be longstanding and global. Other reports of finch trichomonosis in North America have since emerged in west and east-418 central United States of America (Gerhold 2009) and western Canada (Canadian 419 Cooperative Wildlife Health Centre unpublished data) in 2009. During the winter and 420 spring of 2009, the Southeastern Cooperative Wildlife Disease Study (SCWDS) 421 conducted PMEs on passerines of multiple species, including American goldfinch, 422 house finch (Carpodacus mexicanus), northern cardinal (Cardinalis cardinalis), pine 423 siskin and purple finch, submitted from mortality incidents from the eastern United 424 425 States and found that whilst the majority had salmonellosis, at least 12 birds were suffering from trichomonosis or had concurrent infection with both of these pathogens 426 which result in upper alimentary tract lesions (Hernandez et al. 2013; Gerhold 2009). 427

As with GB, there is evidence of some finch trichomonosis incidents in North 428 429 America prior to the emergence of finch trichomonosis in the Canadian Maritime Provinces in 2007. On the western coast of the USA, Anderson et al. (2002) screened 430 birds for trichomonad parasites on admission to a northern California wildlife 431 rehabilitation facility over a period of four years (2001-2005) and found evidence of a 432 low prevalence of the infection in the house finch (1.7%) with a high case fatality rate 433 (95.5%); these authors hypothesised that the infection may be endemic in this (and 434 other) passerine species in the region. Moreover, an outbreak affecting house finches, 435 house sparrows and American goldfinches, contemporaneous with American mourning 436 dove mortality (Zenaida macroura), occurred in the Midwest (Kentucky, Ohio and 437 Indiana) in the autumn of 2002. A combination of trichomonosis and West Nile virus 438 (WNV) infection was diagnosed as the cause of mortality (estimated total of 200 birds) 439

although the relative importance of these agents was not described (NWHC 2002). In
the summer of 2006, a mixed species mortality incident of circa 200 birds involving
house finches, American goldfinches and a gray catbird (*Dumetella carolinensis*) was
reported to the SCWDS. Eighteen birds were submitted for PME with trichomonosis
confirmed in ten cases and WNV infection detected in one bird (Gerhold 2009).

Various potential routes exist through which the UK finch epidemic strain of T. 445 gallinae could have been introduced to the Canadian Maritime Provinces. Bird migration 446 is believed to be the primary route of spread of the disease within Europe. Large 447 numbers of the finch and columbid species in which trichomonosis has been most 448 frequently diagnosed in GB in recent years have been banded (1960-2012 inclusive) 449 (greenfinch n= 2,107,976, chaffinch n=1,287,396, goldfinch (Carduelis carduelis) n= 450 466,108, siskin (*Carduelis spinus*) *n*=503,097 and collared dove *n*=37,780, wood pigeon 451 452 n=45.823): however, no banded birds of these species have been recovered in North America over that period suggesting international exchange is negligible (Robinson and 453 Clark 2013). Indeed, there are remarkably few exchanges of any British wild bird 454 species recorded with North America, with the most frequent being for seabirds and 455 waders, including the kittiwake (Rissa tridactyla) n=73, Manx shearwater (Puffinus) 456 *puffinus*) *n* =25, knot (*Calidris canutus*) *n* =19, turnstone (*Arenaria interpres*) *n*=14, and 457 fulmar (*Fulmarus glacialis*) n=13; all other species with <10 individual birds recorded as 458 North American band recoveries are seabirds, shorebirds or waterfowl species in which 459 T. gallinae infection has not been recorded (Robinson and Clark, 2013). Collectively, 460 therefore bird migration from Europe is an unlikely route of introduction. Since T. 461 gallinae is not capable of long-term environmental persistence, movement with fomites 462

463 is also an implausible method of parasite translocation. Anthropogenic movement of captive birds, whether deliberate (e.g. cage and aviary birds, game birds, zoological 464 collections) or accidental (e.g. wild bird stowaways or stray racing pigeons) could have 465 occurred; however, there is no available evidence to support or refute this hypothesis 466 further. Collectively, therefore, whilst the emergence of finch trichomonosis in the 467 Canadian Maritime Provinces occurred shortly after the emergence of the disease in GB 468 in time, there is no clear candidate for a plausible route of introduction of the finch 469 epidemic strain of *T. gallinae* from the UK. 470

Instead, there is evidence that favours the hypothesis that finch trichomonosis emerged locally in the Canadian Maritime Provinces, through spillover from sympatric birds; this route is most consistent with the SNPs in Fe-hyd subtypes found amongst the finch and columbid isolates from PEI. The occurrence of endemic finch trichomonosis in western USA (Anderson *et al.* 2009), and other isolated finch mortality incidents due to the disease, indicates that parasite strains with the potential to cause disease in passerines have been present in North America for some time.

Future studies should examine *T. gallinae* isolates using multiple gene regions, or full genome sequencing, in order to provide more detailed information about their genetics which could lead to a better understanding of the epidemiology of avian trichomonosis and the mechanisms of disease emergence.

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Figure 1. Geographical distribution of the sites in the Canadian Maritime provinces where *Trichomonas gallinae* isolates were collected. Superscripts correspond to the birds from which the isolate was recovered: F = finch; P = pigeon; and BS = bird seed. Refer to Table 1 for additional details for each isolate.

611

Figure 2. Neighbour-joining 60% bootstrap-consensus tree based on *Trichomonas* 612 gallinae ITS region sequences. Values at nodes represent the bootstrap percentages 613 from 1,000 replicates for neighbour-joining, maximum parsimony and maximum 614 likelihood respectively. There were a total of 209 positions in the final dataset as all 615 positions containing gaps and missing data were eliminated. GenBank accession 616 617 numbers are given along with host names or isolate designations and country for each trichomonad. Isolates in bold are from birds sampled in the present study. For additional 618 isolate details see Table 1. * indicates UK Finch epidemic strain. 619

620

621 Figure 3. Neighbour-joining 60% bootstrap-consensus tree based on *Trichomonas* gallinae Fe-hydrogenase gene sequences. Values at nodes represent the bootstrap 622 percentages from 1,000 replicates for neighbour-joining, maximum parsimony and 623 maximum likelihood respectively. There were a total of 803 positions in the final dataset. 624 GenBank accession numbers are given beside host names or isolate designations and 625 country for each trichomonad. Isolates in bold are from birds sampled and designated 626 627 into the six Fe-hyd subtypes identified in the present study. For additional isolate details see Table 1. * indicates UK Finch epidemic strain. 628

Table 1. Case data and *Trichomonas gallinae* isolates used for the ITS region and Fehydrogenase (Fe-hyd) gene PCR analyses. The last two digits of the year of collection are indicated as the first two digits of the case number. Bird state (alive or dead) indicates whether the sample was collected in-field from live-sampling or at necropsy. ITS typing and Fe-hyd subtyping results from sequence data are recorded.

Isolate ID	Case no.	Species	Bird state	Trichomonosis Status	Site location	ITS type	Fe-hyd subtype
1	09-01	American goldfinch	Alive	Apparently Healthy	Vernon River, PEI	A	1
2	09-02	American goldfinch	Alive	Apparently Healthy	Vernon River, PEI	A	1
3	09-07	American goldfinch	Alive	Apparently Healthy	Vernon River, PEI	A	1
4	09-14	American goldfinch	Alive	Apparently Healthy	New Haven, PEI	A	1
5	09-28	American goldfinch	Alive	Apparently Healthy	New Haven, PEI	A	1
6	11-116	American goldfinch	Alive [†]	Trichomonosis	Orwell, PEI	A	1
7	11-122	American goldfinch	Alive	Trichomonosis	Orwell, PEI	A	3
8	09-04	Purple finch	Alive ^{††}	Trichomonosis	Vernon River, PEI	А	2
9	09-08	Purple finch	Alive	Trichomonosis	Vernon River, PEI	A	2
10	09-12	Purple finch	Alive	Apparently Healthy	Vernon River, PEI	A	2
11	09-13	Purple finch	Alive	Apparently Healthy	Vernon River, PEI	A	1

12	09-15	Purple finch	Alive	Trichomonosis	Vernon River, PEI	A	1
13	09-24	Purple finch	Alive	Apparently Healthy	New Haven, PEI	A	1
14	11-29	Purple finch	Dead	Trichomonosis	Durham Bridge, NB	A	1
15	11-31	Purple finch	Alive ^{††}	Apparently Healthy	Durham Bridge, NB	A	1
16	11-32	Purple finch	Alive	Apparently Healthy	Durham Bridge, NB	A	1
17	11-46	Purple finch	Alive	Apparently Healthy	Durham Bridge, NB	A	1
18	11-50	Purple finch	Alive ^{††}	Apparently Healthy	Durham Bridge, NB	A	3
19	11-100	Purple finch	Dead	Trichomonosis	Pointe-du- Chêne, NB	A	1
20	11-136	Purple finch	Alive	Apparently Healthy	Pointe-du- Chêne, NB	A	1
21	11-113	Purple finch	Alive	Trichomonosis	Orwell, PEI	А	1
22	11-114	Purple finch	Alive [†]	Trichomonosis	Orwell, PEI	А	1
23	11-115	Purple finch	Alive	Trichomonosis	Orwell, PEI	А	1
24	11-117	Purple finch	Alive	Trichomonosis	Orwell, PEI	А	1
25	11-119	Purple finch	Alive	Trichomonosis	Orwell, PEI	А	1
26	11-120	Purple finch	Alive	Trichomonosis	Orwell, PEI	А	1
27	11-121	Purple finch	Alive	Trichomonosis	Orwell, PEI	А	1
28	11-124	Purple finch	Alive	Trichomonosis	Orwell, PEI	А	1
29	11-146	Purple finch	Dead	Trichomonosis	Montague, PEI	А	1
30	10-08	Rock pigeon	Alive	Apparently	Charlottetown,	А	1

				Healthy	PEI		
31	10-09	Rock pigeon	Alive	Apparently Healthy	Charlottetown, PEI	А	1
32	10-12	Rock pigeon	Alive	Apparently Healthy	Charlottetown, PEI	A	1
33	10-14	Rock pigeon	Alive	Apparently Healthy	Charlottetown, PEI	В	6
34	10-16	Rock pigeon	Alive	Apparently Healthy	Charlottetown, PEI	A	1
35	11-06	Rock pigeon	Alive	Apparently Healthy	New Dominion, PEI	A	1
36	11-07	Rock pigeon	Alive	Apparently Healthy	New Dominion, PEI	В	6
37	11-08	Rock pigeon	Alive	Apparently Healthy	New Dominion, PEI	A	1
38	11-09	Rock pigeon	Alive	Apparently Healthy	New Dominion, PEI	A	1
39	11-12	Rock pigeon	Dead	Trichomonosis	Charlottetown, PEI	A	4
40	11-13	Rock pigeon	Dead	Apparently Healthy	Murray River, PEI	A	5
41	11-151	Rock pigeon	Dead	Apparently Healthy	Charlottetown, PEI	В	6
42	09-BF	Bird seed	N/A	N/A	New Haven, PEI	A	NE ^{††}

[†] died immediately after swabbing (approx. 45 minutes), confirmed trichomonosis as cause of death via post-mortem examination.^{††} died after swabbing (days to weeks), confirmed trichomonosis as cause of death via post-mortem examination

^{††}NE – Not evaluated as the PCR was unsuccessful.



160x120mm (152 x 152 DPI)



160x120mm (152 x 152 DPI)



160x120mm (152 x 152 DPI)