

1 ***Theileria lestoquardi* displays reduced genetic diversity relative to sympatric**

2 ***Theileria annulata* in Oman**

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20 **Abstract**

21 The Apicomplexan parasite *Theileria lestoquardi* and *T. annulata* *T. annulata* the causative agents
22 of theileriosis in small and large ruminants, are widespread in Oman, in areas where cattle, sheep
23 and goats co-graze. Genetic analysis can provide insight into the dynamics of the parasite and the
24 evolutionary relationship between species. Here we identified ten genetic markers (micro- and mini-
25 satellites) spread across the *T. lestoquardi* genome, and confirmed their species specificity. We then
26 genotyped *T. lestoquardi* in different regions in Oman. The genetic structures of *T. lestoquardi*
27 populations were then compared with previously published data, for comparable panels of markers,
28 for sympatric *T. annulata* isolates. In addition, we examined two antigens genes in *T.*
29 *annulata* (*Tams1* and *Ta9*) and their orthologues in *T. lestoquardi* (*Tlms1* and *Tl9*).

30 The genetic diversity and multiplicity of infection (MOI) were lower in *T. lestoquardi* ($He=0.64$ –
31 0.77) than *T. annulata* ($He=0.83$ – 0.85) in all populations. Very limited genetic differentiation was
32 found among *T. lestoquardi* and *T. annulata* populations. In contrast, limited but significant linkage
33 disequilibrium was observed within regional populations of each species. We identified eight *T.*
34 *annulata* isolates in small ruminants; the diversity and MOI were lower among ovine/caprines
35 compared to bovine. Sequence diversity of the antigen genes, *Tams1* and *Ta9* in *T.*
36 *annulata* ($\pi=0.0733$ and $\pi=0.155$ respectively), was 10-fold and 3-fold higher than the
37 orthologous *Tlms1* and *Tl9* in *T. lestoquardi* ($\pi=0.006$ and $\pi=0.055$, respectively).

38 Despite a comparably high prevalence, *T. lestoquardi* has lower genetic diversity compared to
39 sympatric *T. annulata* populations. No evidence of differentiation among populations of either
40 species. In comparison to *T. lestoquardi*, *T. annulata* has a larger effective population size. While,
41 genetic exchange and recombination occurs in both parasite species, the extent of diversity, overall,
42 is less for *T. lestoquardi*. It is, therefore, likely that *T. lestoquardi* evolved from an ancestor of
43 present day *T. annulata* and that this occurred either once or on a limited number of occasions.

44 **Keywords:**

45 *Theileria lestoquardi*, *Theileria annulata*; population genetics, evolution, host species jump, Oman

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52 1. Introduction

53 *Theileria lestoquardi* is a highly pathogenic ovine and caprine parasite and is considered to be the
54 only *Theileria* species of economic significance in small ruminants (Leemans *et al.*,2001; Li *et*
55 *al.*,2014). The parasite is transmitted by *Hyalomma anatolicum anatolicum*, which is common in
56 South-eastern Europe, Northern Africa, Southern Russia and the Middle East. However,
57 distribution of *T. lestoquardi* is limited compared to the range of its vector. Although *T. lestoquardi*
58 has been shown to be antigenically closely related to *T. annulata* (Leemans *et al.*,1997), it has been
59 reported as being incapable of infecting cattle (Leemans *et al.*,1999). Conversely, it is known that *T.*
60 *annulata* can infect sheep; experiments in sheep indicate that *T. lestoquardi* infection protects
61 against subsequent *T. annulata* infection (Leemans *et al.*,1999) and although prior infection with *T.*
62 *annulata* does not prevent infection from *T. lestoquardi* sporozoites, it does protect against the
63 major clinical effects. However, these experiments were carried out on limited numbers of animals
64 with a very limited number of parasite genotypes, and the actual transmission dynamics in the field
65 are unknown.

66 *Theileria lestoquardi* was first reported in sheep in Sudan and Egypt (Littlewood,1916), and later
67 detected in sheep and goats in other countries of the Middle East such as Algeria
68 (Lestoquard,1927), Turkey (Baumann,1939), Iraq (Khayyat *et al.*,1947), Iran (Hooshmand-Rad *et*
69 *al.*,1976; Hawa,1981) as well as India (Raghvachari,1959) and Serbia (Dschunkovsky *et al.*,1924).
70 A previous study in Oman demonstrated a high level of theileriosis-attributed mortality in a local
71 sheep breed (Tageldin *et al.*,2005). This confirmed previous individual case reports and outbreak
72 records of a pathogenic species of *Theileria* in sheep and goats in Oman (Annual Reports
73 VRC2004-2006) (MOAF,2008). These reports indicated that in Oman, sheep, in general, were
74 significantly more at risk of clinical theileriosis than cattle and goats, and this has been attributed to
75 a higher tick infestation of sheep. However, the relative distribution of the major pathogenic
76 species of *Theileria* (*T. lestoquardi* and *T. annulata*) is not yet known in the Sultanate of Oman.
77 Thus, there is currently no information on the prevalence of *T. lestoquardi* in different regions in

78 Oman and nothing is known regarding the *T. lestoquardi* population structure. In contrast, a recent
79 survey demonstrated that *T. annulata* is widely distributed across the country and is comprised of a
80 highly genetically diverse, inter-breeding population (Al-Hamidhi *et al.*,2015).

81 Genetic analysis of parasite populations can provide important information about the epidemiology
82 of disease and may facilitate the development of rational control approaches. Polymorphic genetic
83 markers have been developed for some species of *Theileria*, e.g. *T. annulata* and *T. parva* (Oura *et*
84 *al.*,2003; Weir *et al.*,2007), however, such tools are not yet available for the small ruminant
85 *Theileria* species parasites, *T. lestoquardi* and *T. ovis*. Micro- and mini-satellites are considered as
86 highly appropriate molecular markers for population genetics applications. Their high mutation rate
87 and Mendelian mode of inheritance make them particularly useful for the study of both fine and
88 broad-scale population genetic structure (Abdelkrim *et al.*,2009). Common applications include
89 assessing genetic diversity, degree of population inbreeding, bottleneck effects, gene flow and
90 migration rates, the assignment of population of origin and parental lineages (Goldstein *et al.*,1999).

91 The present study included the development of micro- and mini-satellite genotyping for
92 *T. lestoquardi* and their application to investigate the genetic diversity of parasite populations from
93 four regions in Oman. The extent of diversity and population structure of *T. lestoquardi* was then
94 compared to available published data on sympatric *T. annulata* populations for three of the four
95 regions. We aimed to gain an understanding of whether local gene flow and genetic diversity differs
96 between these two species in an area of similar prevalence and distribution of tick species. We also
97 investigated the hypothesis that *T. lestoquardi* is a relatively recently evolved species that has
98 diverged from the more ancient cattle parasite species, *T. annulata*, following a host species jump to
99 small ruminants.

100 **2. Materials and method**

101 **2.1. Parasite material and DNA preparation**

102 Blood samples (n = 1,454) were collected from clinically healthy sheep and goats in four
103 governorates of Oman: Batinah (n = 584), Dhira (n = 357), Sharqia (n = 369) and Dakhiliya

104 (n = 144) (Figure 1). The climate across these regions is hot and dry throughout the year, with 3-4
105 months (Oct to Feb) of relatively moderate temperatures (below 30 °C).

106 For comparison of diversity and population structure, genotyping data representing 97 *T. annulata*
107 isolates from Batinah (n = 21), Dhira (n = 57) and Sharqia (n = 19) derived from cattle co-grazed on
108 the same farms as the sheep/goats that provided *T. lestoquardi* isolates was utilised. These were
109 previously genotyped with a set of *T. annulata* specific micro- and mini-satellites (Al-Hamidhi *et*
110 *al.*,2015).

111 **2.2. Identification of specific *T. lestoquardi* micro- and mini-satellite sequences**

112 A draft sequence of the *T. lestoquardi* genome has been generated (Weir *et al.*, unpublished). To
113 identify micro- and mini-satellite loci specific for *T. lestoquardi*, sequence contigs were screened
114 using the tandem repeat finder program (Benson,1999). A filtration pipeline was used to identify a
115 subset of high-value loci, which could be tested using a panel of available stocks and isolates.
116 Filtration included discarding repeat regions greater than 500 bp in length and those that possessed
117 insufficient flanking sequence for primer design. The remaining sequences were ranked, based on
118 the fidelity of the repeat within each region (> 70 % fidelity) and the number of repeats. A subset of
119 28 loci with conserved repeat motifs was then derived.

120 **2.3. PCR amplification of specific micro- and mini-satellite loci**

121 Primers were designed to unique sequence flanking each repeat and used to amplify DNA purified
122 from a panel of stocks (*T. lestoquardi*, *T. annulata* and *T. ovis*) and field isolates to test marker
123 specificity and polymorphism. In addition, to test for marker sensitivity, serial dilutions of *T.*
124 *lestoquardi* DNA were generated and PCR performed with each primer set and sample.

125 PCR was carried out in a total reaction volume of 20 µl using conditions described previously (Al-
126 Hamidhi *et al.*, 2015). Thermocycler parameters were as follows: denaturation at 94 °C for 5
127 minutes, 32 cycles at 94 °C for 30 seconds, 42-55 °C for 30 seconds, and 65 °C for 30 seconds,
128 followed by a final extension step of 5 minutes at 65 °C. Amplified products were observed on a 2

129 % ethidium bromide pre-stained agarose gel and their size determined with reference to either a
130 1 kb or 100 bp DNA ladder.

131 To identify length polymorphism down to the level of 1 base pair (bp), PCR products were
132 denatured and then capillary electrophoresed in an ABI3130 xl Genetic Analyser (Applied
133 Biosystems, UK). DNA fragment sizes were determined relative to ROX-labeled GS500 size-
134 standards (Applied Biosystems) using GeneMapper software (Applied Biosystems). For all loci and
135 DNA samples, fragment size (i.e. peak position) was determined to two decimal places. Analysis of
136 the distribution of fragment sizes facilitated the creation of 'fixed bins' of variable size to score
137 alleles. Since these loci represent genomic regions encoding hypothetical proteins, variation among
138 allele sizes was assumed to be in steps of three base pairs or multiples thereof.

139 The single or predominant allele for each of the ten selected loci was utilised to compute allele
140 frequencies. Each of the markers selected for further analysis was shown to represent a different
141 single-copy locus based on genome data and PCR fragments amplified from *T. lestoquardi* (Lahr)
142 DNA. Since *Theileria* parasites are haploid, the presence of one or more additional alleles at a
143 particular locus was interpreted as a co-infection with one or more genetically distinct genotypes.
144 An additional allele was scored if the peak was at least one-third the height of the predominant
145 allele (highest peak) on the electropherogram traces, a method that has been widely used in
146 previous studies (Anderson *et al.*,1999). In this way, the predominant allele at each locus was
147 identified for each sample and the data combined to generate a multi-locus genotype (MLG),
148 representing an estimate of the most abundant genotype in each sample, as described previously
149 (Weir *et al.*,2007).

150 The MLG dataset was then used to measure population genetic indices such as heterozygosity,
151 linkage disequilibrium and population differentiation. Since *Theileria* is haploid and heterozygosity
152 cannot be observed directly, the estimated heterozygosity was calculated using the predominant
153 allele dataset for each marker and averaged across all ten loci.

154 **2.4. Sequence analysis of *Tams1/Tlms1* and *Ta9/TL9* orthologues in *T. annulata* and *T.***
155 ***lestoquardi***

156 *Theileria annulata* and *T. lestoquardi* isolates were obtained from the same farms in Sharqia and
157 Dhira, since a high level of *Theileria* infection had been detected in animals from each region. The
158 PCR products for *Tams1/Tlms1* and *Ta9/TL9* genes were generated and cloned using the Topo
159 sequencing vector. DNA from 5 purified colonies representing each isolate was sequenced by
160 ABI3130 xl Genetic Analyser (Applied Biosystems, UK). The obtained nucleotide sequence was
161 confirmed by via the NCBI BLAST web interface (<http://www.ncbi.nlm.nih.gov/>), and nucleotide
162 sequences translated to amino acid sequences using MEGA4 software (Tamura *et al.*,2007).
163 Nucleotide and translated amino acid sequences were aligned with the corresponding reference
164 gene from the *T. annulata* and *T. lestoquardi* genome sequence using MEGA V. software.
165 Sequence polymorphism and diversity was estimated using DnaSP version 5.0 (Librado *et al.*,2009)
166 by calculating the total number of polymorphic sites (S); the average pair-wise nucleotide diversity
167 (π), the average number of nucleotide differences (k) and haplotype diversity combinations for all
168 divergent sequences. The HKY+G mutational model applied was chosen using jmodeltest
169 (<http://jmodeltest.org>). The tree for nucleotide sequence of *Tams1/Tlms1* gene was constructed
170 using a PhyML 3.0. software (Guindon S.,2010), and visualize using archaeopteryx software
171 (<https://sites.google.com/site/cmzmasek/home/software/archaeopteryx>).

172 **2.5. Data analysis**

173 The Excel Microsatellite toolkit (Bowcock *et al.*,1994) was used for a similarity comparison of
174 MLGs. Genetic diversity parameters were calculated for the entire population using GenAlex v6.5
175 (Peakall *et al.*,2012) by determining the number of alleles per locus (A) and the expected
176 heterozygosity (Dschunkovsky *et al.*,1924). Allelic diversity was determined using the formula for
177 ‘unbiased heterozygosity’, the equivalent of diploid expected, also named as haploid genetic
178 diversity, $H_e = [n/(n-1)][1-\sum p^2]$ where n is the number of isolates and p the frequency of each

179 different allele at a locus (Anon,1996). Expected heterozygosity ranges between 0 and 1, with
180 values close to 1 reflecting high genetic diversity levels in a population.

181 To determine whether the *T. lestoquardi* and *T. annulata* populations in different regions comprised
182 a single panmictic population with a high degree of genetic exchange, linkage disequilibrium (LD),
183 i.e. the non-random association of alleles among loci was quantified using the standard index of
184 association (I^S_A). Each region was analysed separately and then the samples were pooled and
185 analysed as a single set. Both I^S_A and the variance data were calculated using the program LIAN,
186 version 3.5 (Haubold *et al.*,2000). This software tests for independent assortment of alleles by
187 determining the number of loci at which each pair of MLGs differs, and from the distribution of
188 mismatch values, a variance V_D (the variance of the number of alleles shared between all pairs of
189 haplotypes observed in the population) is calculated which is then compared with the variance
190 expected for linkage equilibrium (LE), termed V_e . The null hypothesis that $V_D = V_e$ is tested by
191 either a Monte Carlo simulation or a parametric method and the results provide 95 % confidence
192 limits, which are denoted L_{MC} and L_{PARA} , respectively. If there is limited or no association between
193 alleles at different loci, indicating panmixia, a value close to zero is obtained for the I^S_A , whereas if
194 association is detected at a value significantly greater than 0, LD is indicated (Haubold *et al.*,2000).

195 The variance of pair-wise difference (V_D) between the data and that predicted for panmixia (V_e) and
196 L were calculated in order to test the hypothesis of panmixia. To test whether the populations in
197 each region were genetically differentiated, the reduction in heterozygosity for sub-populations
198 compared to the overall population, Wright's fixation index (F_{ST}) (Brown,1970) value was
199 calculated.

200 As some of the loci are located in regions near or inside coding genes, we have conducted a F_{ST}
201 outlier tests for detect loci that might have being under selective pressure. These tests were
202 conducting using the algorithm included in F_{ST} concept (Beaumont MA *et al.*,1996), using the java
203 based software user-friendly Mcheza (Antao *et al.*,2011).

204 Principal co-ordinate analysis (PCoA), a multivariate analysis also known as multidimensional
205 scaling (MDS), was used to investigate the genetic relationships between the isolates MLGs. A F_{ST}
206 based genetic distances matrix was used to calculate the PCoA, which the results can be plotted to
207 visualize the genetic relationships between individuals and/or populations. This analysis was
208 calculate using Genalex V6, excel plugin software (Peakall *et al.*,2012).

209 **2.6. Multiplicity of infection**

210 Multiplicity of infection was defined as the “presence of multiple genotypes per isolate” by the
211 detection of more than one allele at a locus, when minor peaks were >33% the height of the
212 predominant allele present. The mean number of alleles across ten selected loci in each sample was
213 calculated and this index value was used to represent the multiplicity of infection within each
214 sample. The overall mean for the index value for each sample was then calculated to provide the
215 average multiplicity of infection for each region.

216 **3. Results**

217 **3.1. Identification and evaluation of *T. lestoquardi* micro- and mini-satellites**

218 A panel of twenty-eight repeat-containing single-copy loci were initially identified by screening the
219 draft genome of *T. lestoquardi* with repeat finder (Benson,1999). These loci represented 13 micro-
220 satellite (motif size 3 - 6 bp) and 15 mini-satellite (motif size 9 - 24 bp) markers. Of the 28, only ten
221 loci had flanking sequence suitable for designing primers specific for *T. lestoquardi*; the other 18
222 were either flanked with sequence common to *T. annulata* and *T. lestoquardi* or the flanking
223 sequences were too short to allow primer design. The ten selected loci consisted of four micro-
224 satellites (TL_MS07, TL_MS13, TL_MS19 and TL_MS16) and six mini-satellites (TL_MS05,
225 TL_MS281, TL_MS280, TL_MS18, TL_MS04 and TL_MS25). The characteristics of these loci
226 are summarised in supplementary Table 1 and supplementary Table 2. Eight of the ten loci are
227 located in exons; one is in an intron and another in an intergenic region. The genes associated with
228 or flanking these loci are all annotated as hypothetical proteins with orthologues present in the
229 *T. annulata* genome (Pain *et al.*,2005).

230 PCR of the selected ten loci generated amplicons of the predicted size with *T. lestoquardi* DNA, but
231 no product was obtained with *T. annulata* and *T. ovis* template DNA, demonstrating that the
232 selected markers were specific for *T. lestoquardi*. The selected marker primer sets were then used to
233 genotype 36 DNA samples representing *T. lestoquardi* field isolates, after the presence of *T.*
234 *lestoquardi* DNA was confirmed by PCR-RFLP of the 18S rRNA locus. Each DNA sample/marker
235 combination produced an amplicon. Variation in amplicon size among isolates was observed for
236 each marker, confirming these loci as being polymorphic and thus informative for population
237 analysis. The differences in allele size for each marker ranged from 3 to 9 bp and agreed well with
238 the motif size of each marker. A subset of DNA samples showed evidence of more than one allele
239 at one or more loci, indicating the presence of multiple genotypes in a number of animals.

240 **3.2. Prevalence and multiplicity of infection of *T. lestoquardi* relative to *T. annulata* across** 241 **three regions in Oman**

242 The ten micro- and mini-satellites were then used to analyse *T. lestoquardi* populations in four
243 regions in Oman. The extent of diversity and population structure of *T. lestoquardi* were then
244 compared to *T. annulata* using previously published mini- and micro-satellite data for 97 isolates
245 from cattle, obtained from three of the four regions where cattle and small ruminants co-graze.

246 Of the 1,688 blood samples collected [1454 small ruminant and 234 bovine], a total of 190/1454
247 (13 %) and 97/234 (41 %) were positive for *T. lestoquardi* or *T. annulata* parasites, respectively, as
248 detected by PCR/RFLPs and/or PCR/RLB (Al-Fahdi *et al.*, 2015). The difference in prevalence of
249 either species across the different regions was not significant (chi squared test, $P > 0.05$). However,
250 the prevalence of *T. annulata* in cattle was significantly higher than that of *T. lestoquardi* in small
251 ruminants.

252 Genotyping data generated from the ten micro- and mini-satellite markers for each of the two
253 species showed significantly greater MOI among *T. annulata* cattle isolates (ranging between 2.9
254 and 3.2) than the *T. lestoquardi* small ruminant isolates (1.49 to 1.63) (t test, $P < 0.001$) (Table 1).
255 Similarly, the *T. annulata* dataset had a significantly larger proportion of multiple infections

256 (52 %), with more than one allele at one or more loci, than the *T. lestoquardi* dataset (44 %) (Chi-
257 squared test, 1 df: P = 0.0045).

258 **3.3. Relative diversity of mini- and micro- satellite markers**

259 All ten markers for *T. lestoquardi* were found to be polymorphic, with the number of alleles for
260 each marker ranging from four, for TL_MS25, to 22 for TL_MS280. The average of number of
261 alleles per marker was 12.6. Broadly similar allele frequencies were observed for each marker in
262 each region, as the example of TL_07 in Figure 2 shows, while a limited number of private alleles
263 specific to sub-populations from each region were observed (Table 2). Three markers revealed a
264 lower level of diversity (H_e range 0.121-0.441), compared to higher levels observed for the
265 remaining seven (H_e range 0.548-0.867) (Table 2). The average heterozygosity identified within
266 each of the four geographical regions was found to be moderate, ranging from 0.637 within Sharqia
267 to 0.575 in Batinah (Table 2).

268 The extent of gene diversity among *T. lestoquardi* isolates was compared to that of *T. annulata*
269 cattle isolates obtained from the same sites. Genetic diversity was consistently higher for
270 *T. annulata*, where the estimate of diversity within each region (H_e range 0.820 to 0.854) was
271 similar to the average of combined diversity in all regions ($H_e = 0.836$) (Table 3), consistent with
272 little or no differentiation between sub-populations.

273 **3.4. Comparative analysis of sequence diversity of antigen genes**

274 We assessed the extent of diversity of two antigen genes in *T. annulata*: the immunodominant
275 merozoite/piroplasm surface antigen of *T. annulata* (*Tams1*) (Shiels *et al.*,1995) and *Ta9* which
276 encodes peptides recognised by CD8⁺ T cells from immune animals (MacHugh *et al.*,2011). The
277 level of sequence diversity in Omani isolates was then compared to that of the orthologous genes in
278 sympatric *T. lestoquardi* isolates (*Tlms1* and *TI9*).

279 Partial sequence of *Tams1* and *Tlms1* were obtained from *T. lestoquardi* (38 isolates) and
280 *T. annulata* (36 isolates) from the same region in Oman. For *Tams1*, 144 nucleotide site
281 polymorphisms were found among aligned *T. annulata* sequences in comparison with the reference

282 genome sequence strain (Ankara, C9), while only 19 polymorphisms were detected across the *T.*
283 *lestoquardi* sequences (Table 4).

284 Nucleotide alignment of *Tlms1* revealed eight haplotypes with haplotype diversity of 0.649 among
285 *T. lestoquardi* sequences. However, 20 haplotypes were identified for the *T. annulata* orthologue
286 *Tams1*, with a haplotype diversity (Hd) of 0.968 (Table 4). The overall nucleotide diversity (π) for
287 *T. annulata* ($\pi = 0.0733$) *Tams1* was 10-fold higher than that computed for the *Tlms1* sequences
288 ($\pi = 0.006$) and the average number of pair-wise nucleotide differences (k) was 3.902 and 45.832 in
289 *T. lestoquardi* and *T. annulata*, respectively (Table 4). Thus, these results demonstrate that
290 nucleotide diversity of the major merozoite/piroplasm surface antigen gene is significantly higher in
291 *T. annulata* than in its *T. lestoquardi* orthologue, based on analysis of a similar number of
292 sympatric isolates. This difference in sequence diversity between alleles representing the two
293 orthologues was illustrated by the generation of a phylogenetic tree Figure 3. Clearly, the branch
294 lengths are longer within the *T. annulata* tree, indicating more diversity/distant relationship between
295 sequences. In addition, the sequences for both species, as might be predicted, show clear separation,
296 with the *T. lestoquardi* indicated as branching/evolving from a common ancestor of the *T. annulata*
297 sequences.

298 For the *Tl9/Ta9* comparison, 9 and 23 distinct sequences were obtained from a similar number of *T.*
299 *lestoquardi* and *T. annulata* isolates, respectively. Haplotype number and Hd was 7 and 0.9, for *T.*
300 *lestoquardi* 10 and 0.978 for *T. annulata*, respectively (Table 4). However, nucleotide diversity (π)
301 was 3-fold higher for *T. annulata* ($\pi = 0.155$) sequences compared to that of *T. lestoquardi*
302 ($\pi = 0.055$). Thus, the results for *Ta9/Tls9* reflect those of *Tams1/Tlms1*, demonstrating that two
303 antigen genes selected for analysis have higher diversity in *T. annulata* than in their *T. lestoquardi*
304 orthologues (Table 4), and this consistent with the results of the micro- and mini-satellites.

305

306 **3.5. Genetic diversity of *T. annulata* isolated from small ruminants in Oman**

307 Eight *T. annulata* isolates collected from small ruminants were genotyped using the ten published
308 *T. annulata* micro- and mini-satellites and compared to the *T. annulata* genotyping results from the
309 bovine isolates (Al-Hamidhi *et al.*,2015). Similar to bovine-derived isolates, each of the small
310 ruminant isolates was found to carry multiple genotypes, with several alleles identified at one or
311 more loci. However, the mean MOI was lower compared to that obtained for bovine isolates
312 (average of 2.9 in small ruminants compared to 3.27 in bovine), but this difference was not
313 significant. Six private alleles were observed, for the small ruminant isolates, on four loci (one
314 allele each for Ts12 and Ts9 and two alleles each for Ts6 and Ts8). Due to the small number of
315 isolates from small ruminants, genetic differentiation between *T. annulata* genotypes derived from
316 the different host species could not be estimated.

317 **3.6. Linkage disequilibrium analysis**

318 To assess whether *T. lestoquardi* parasites in the study regions undergo random mating with a high
319 level of genetic exchange, the extent of LD at pairs of loci was measured using the standard index
320 of association (I^S_A). Low, yet significant LD was found when each region was treated as a single
321 population and a low overall I^S_A value of 0.0264 was obtained. A V_D value (2.28) greater than L
322 (1.98) was calculated indicating LD (Table 5). However, when each regional population was treated
323 separately Dhira and Batinah showed (I^S_A) close to zero with pair-wise variance (V_D) less than the
324 critical L value, indicating that those two populations were in LE. The above inconsistencies was
325 likely a reflection of variation in effective population size (N_e) and sub-population structure,
326 Wahlund effect (Waples *et al.*,2011). This agrees with the small effective population size among *T.*
327 *lestoquardi* in different regions, which ranged between 2.71 and 3.73 compared to a higher N_e seen
328 among *T. annulata*, which ranged between 6.96 and 8.46 (Table 3). For *T. annulata*, a lower but
329 significant LD score was found for two of the three populations, whereas linkage equilibrium was
330 evident in the Batinah population (Table 5).

331 **3.7. Detection of possible selection on mini- and microsatellites**

332 The F_{ST} outlier test conducted to detect departures from neutrality found in four loci showing low
333 F_{ST}/H_e , with significant statistical support to be classified as lower threshold outliers (Table 6).
334 Loci showing low F_{ST} are often under balancing selection, as this process forces alleles to maintain
335 heterozygosity and lower differentiation across populations under the same environments.

336 **3.8. Population structuring**

337 A low level of F_{ST} was detected between each pair of the four *T. lestoquardi* populations (Table 7),
338 as well as between pair-wise combinations of the three *T. annulata* populations, indicating a lack of
339 differentiation between regional populations. A low level of differentiation between regional
340 parasite populations is supported by Principal Coordinate Analysis (PCoA) (Figure 4A and B).
341 PCoA demonstrated no evidence of regional structuring for either species, with haplotypes
342 distributed throughout the main cluster independent of geographic origin.

343 **4. Discussion**

344 Small ruminant theileriosis is a major problem in Oman, as it is a leading cause of morbidity and
345 mortality and is associated with significant economic loss. To establish innovative control measures
346 and assess their effectiveness, information on the extent of genetic diversity and population
347 structure of *T. lestoquardi* is desirable. It is also of interest to investigate how Apicomplexan
348 parasites may evolve by adapting to novel host species, and to determine whether such events occur
349 at low or high frequency. The *T. annulata*/*T. lestoquardi* relationship provides a good model for
350 this, as biological and molecular phylogenetic data suggest that *T. lestoquardi* has most likely
351 evolved from an ancestral *T. annulata* infection of small ruminants (Leemans, *et al.*, 1998; Katzer *et*
352 *al.*, 1998; Schttinger *et al.*, 2000) generating a parasite species that manifests acute pathology in
353 susceptible hosts. In this study we investigated these questions by developing and validating a set of
354 ten micro- and mini-satellites markers specific for *T. lestoquardi* and used them in a comparative
355 analysis of *T. lestoquardi* and *T. annulata* parasites in four regions of Oman.

356 Although micro- and mini-satellites representing *T. annulata* (Weir *et al.*, 2007) and *T. parva* (Oura
357 *et al.*, 2003) have previously been identified and characterised, this study is the first to report similar

358 markers for estimating genetic diversity within and between isolates of *T. lestoquardi*. The present
359 study describes the development of a panel of ten *T. lestoquardi*-specific markers, which are
360 distributed over the four chromosomes. The ten loci showed considerable diversity within the
361 studied populations with seven having an excess of high H_e . Together these markers represent a
362 useful tool for analysing *T. lestoquardi* populations in the field, as they negate any issues of co-
363 infection with related *Theileria* species and can provide an estimate of the level of genetic diversity
364 and divergence within and between populations.

365 The markers revealed a high level of genetic diversity, a limited degree of linkage disequilibrium
366 and an absence of differentiation across different *T. lestoquardi* populations in Oman. However, the
367 extent of diversity among *T. lestoquardi* isolates was much lower than observed within *T. annulata*
368 isolates in three regions where the two species co-exist. The mean H_e index for *T. lestoquardi*
369 isolates in each site ranged from 0.575 to 0.637, lower than that observed among *T. annulata* in
370 Oman (H_e ranged between 0.819 and 0.854) (Al-Hamidhi *et al.*,2015) and other endemic countries
371 (Weir *et al.*,2011), as well as that reported for *T. parva* in Zambia (Muleya *et al.*,2012). The higher
372 level of genetic diversity in the *T. annulata* population may be the result of genetic recombination
373 over an extended period of time compared to *T. lestoquardi*, which may have emerged more
374 recently. Whether the higher MOI of *T. annulata* in the cattle population is simply a reflection of
375 increased diversity in this parasite population is difficult to gauge. However, given the high level of
376 identity at the 18S rRNA locus between the *T. lestoquardi* and *T. annulata* (Schnittger *et al.*,2000),
377 it is most probable that *T. lestoquardi* has evolved from an ancestral cattle-infective parasite related
378 to present day *T. annulata* and that parasite speciation occurred as the parasite adapted to the small
379 ruminant host. A similar conclusion on host switching and parasite speciation has been made,
380 following analysis of mitochondrial genome sequences, for primate malaria parasites among several
381 species that live in sympatry (Escalante *et al.*,1998).

382 The above hypothesis is also consistent with the greater diversity of two antigen genes in
383 *T. annulata* compared to that of their orthologues in *T. lestoquardi*. Indeed, construction of

384 phylogenetic trees from sequence data from each antigen gene clearly shows separation of
385 sequences representing either species with no indication that any *T. lestoquardi* sequence showed a
386 closer relationship to *T. annulata* than the rest of the dataset (Figure 3). Taken together the data
387 indicate that the jump from the ancestral species that allowed adaptation to small ruminants is not a
388 frequent event, and may have only happened on a limited number of occasions, involving a low
389 number of genotypes. Thus, much of the pre-existing diversity in the cattle population would not
390 have been carried over into the *T. lestoquardi* population. Whether speciation events linked to host
391 adaptation of vector-borne Apicomplexan parasites are generally infrequent is unknown. However,
392 a study of the evolution of *Plasmodium falciparum* concluded that the jump of the ancestral parasite
393 from gorillas may have resulted from a single cross-species transmission event (Liu *et al.*,2010).
394 These studies may indicate that, while evolution of new pathogenic Apicomplexan species after
395 transmission to a novel host has occurred on a number of occasions (Arisue *et al.*,2015), the
396 probability of this happening on a frequent basis is not high.

397 We identified eight *T. annulata* isolates in small ruminants (seven ovine and one caprine), and
398 found that the average number of alleles and MOI were slightly lower (2.9 vs 3.27) in the
399 ovine/caprine isolates than in bovine isolates of *T. annulata*. This preliminary data suggests
400 *T. annulata* is less well adapted to sheep than *T. lestoquardi* and that establishing *T. annulata*
401 infection is more difficult in ovine cells than bovine. Exactly how competent small ruminants are in
402 the transmission of *T. annulata* in the field is unknown; however the weight of evidence to date
403 does not suggest they play a major role in the epidemiology of tropical theileriosis in comparison to
404 cattle. With the common ancestor of *T. annulata* and *T. lestoquardi* presumed to be a cattle parasite,
405 essentially similar to modern day *T. annulata*, adaptive changes promoting establishment and
406 transmission capability in small ruminants would likely have developed as *T. lestoquardi*
407 established as a species. However, whether the most recent common ancestor shared *T. annulata*'s
408 inability to produce piroplasms in small ruminants (Li *et al.*,2014) is impossible to say and the
409 degree of each host-species adaptation in the intervening time is unknown.

410 The high diversity of antigen genes *Tams1* and *Ta9* (Table 4) in *T. annulata* has been proposed to
411 confer a selective advantage to parasite genotypes by facilitating evasion from a protective immune
412 response (Wang *et al.*,2014). This and genetic diversity, in general, could promote a more
413 widespread distribution and survival of this species, even in the face of various control strategies.
414 However, it should be noted that although less divergent, a stable transmissible endemic population
415 of *T. lestoquardi* exists in a number of countries of the Middle East and Africa.

416 The high genetic diversity of bovine *T. annulata* populations in Oman compared to that detected
417 among sympatric *T. lestoquardi* is consistent with the multiplicity of infection data (Table 1). MOI
418 is a prerequisite for cross-mating and recombination among different parasite genotypes in the tick
419 vector midgut, and thus the generation of novel recombinant genotypes. The proportions of animals
420 harbouring multiple infections were similar for each species; however the mean MOI values
421 differed considerably, being two-fold higher for *T. annulata*. This cannot be attributed to variation
422 in density of infection between the two species, as PCR detection can favor the most abundant
423 genotypes existing at high parasitaemia compared to those at low levels, as all samples were
424 collected from animals not showing clinical signs. MOI could result from inoculation of multiple
425 clones from one infected tick or multiple ticks infected with distinct parasite genotypes feeding on a
426 single bovine (superinfection). The former is expected to happen more readily in *T. annulata* due to
427 the high level of diversity displayed by infected bovine isolates. Whatever the cause of MOI, the
428 higher multiplicity of *T. annulata* genotypes could sustain a high rate of cross-mating and
429 recombination in the tick vector, which in turn would result in increased genetic diversity in the
430 bovine host, as demonstrated for the human malaria parasite *P. falciparum* (Babiker *et al.*,1994;
431 Conway *et al.*,1999). In addition, 98% of the adult ticks collected from examined cattle and sheep
432 were *H.anatolicum*: thus, it appears unlikely that transmission by different tick vectors could
433 account for the differences in diversity of the two *Theileria* species.

434 The significant LD seen among some populations of both species contrasts with the expected high
435 levels of out-crossing; however the LD is essentially mild and is only a slight departure from

436 panmixia. LD can be influenced by demography and/or selection events. Diverse factors, other than
437 the extent of inbreeding including the recombination rate, the local parasite effective population
438 size and population differentiation (Hill and Babiker, 1995; Hill *et al.*,1995). Similar to other
439 vector-transmitted parasites, *T. lestoquardi* genotypes are not randomly distributed, but rather the
440 population is fragmented, with individual host animals supporting a sub-population of genotypes.
441 Similarly, effect can also be achieved by undergoing selection. Individual carrying a genotype that
442 positively affects its fitness, will be selected and increase in frequency in the population, at the
443 expenses of the less “fitted” genotypes which will be erased, reducing the number of possible
444 genotype combination available in a given population. The moderate LD values observed in the *T.*
445 *lestoquardi*, can be explained by undergoing balancing selection. In this type of selection, several
446 genotypes bring similar advantages to the individuals and therefore the frequency of those
447 genotypes tends to be even. In other words, balancing selection promotes diversity (heterozygosity)
448 rather than positive selection that promotes fixation (homozygosity). However, co-uptake of sexual
449 stages of closely related genotypes by the feeding tick may result in non-random mating and
450 consequently LD. Assuming random pairing of male and female gametes, the frequency of cross-
451 mating equals the probability these gametes are sampled from different genotypes carried in a
452 single animal. The probability of inbreeding can be related to the numbers of genotypes detected
453 per infection, assuming that all blood form parasites are represented in the gametocyte population
454 (Hill *et al.*,1995). It has been shown that a small number of genetically related parasites in the
455 vertebrate host can generate significant linkage disequilibrium (Anderson *et al.*,2000). Thus, the
456 observed LD and lower level of genetic diversity in *T. lestoquardi* relative to *T. annulata*, does not
457 necessarily indicate the absence of a broadly panmictic population structure or reduced levels of
458 genetic recombination.

459 Very low levels of genetic differentiation were detected between *T. lestoquardi* populations in the
460 four sites in Oman with most pair-wise F_{ST} values being less than 0.04. This is consistent with the
461 analysis of sympatric *T. annulata* populations which also show a low level of population

462 differentiation (Al-Hamidhi *et al.*,2015). The results suggest a rate of genetic exchange and gene
463 flow between parasites in different parts of the country, sufficient to allow the population to remain
464 homogenous and to overcome genetic drift through geographical and genetic isolation. It is likely
465 that homogenisation of the population is underpinned by movement of infected/infested animals
466 from one area to another. It would be of interest to examine *T. lestoquardi* populations in
467 neighboring countries in the region, where theileriosis is also a major problem, to determine how
468 closely related these populations are. This could determine whether control measures, based on
469 vaccine or drug therapy should be implemented separately or if a regional policy can and should be
470 adopted.

471 In conclusion, the present study compared genotypic and population diversity between sympatric *T.*
472 *lestoquardi* and *T. annulata* in Oman. *Theileria annulata* populations were shown to be consistently
473 more diverse and hosts displayed a greater MOI. These results provide an insight into the evolution
474 of *T. lestoquardi*, reinforcing the hypothesis that it has diverged from ancestral *T. annulata* and
475 evolved following adaption to small ruminant hosts, potentially via a single cross-species adaptive
476 event. Further work investigating the molecular basis that promoted host adaptation and speciation
477 of *T. lestoquardi* is warranted, together with investigation of whether a reduced level of antigenic
478 diversity impacts on transmission efficiency of *T. lestoquardi* relative to *T. annulata* in the field.

479

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634 **Figure 1:** Locations of sample collection sites in Oman for *T. lestoquardi* and *T. annulata*.

635 **Figure 2:** The frequency of *T. lestoquardi* alleles in the four governorates of Oman for the
 636 representative marker TL07. The size of each allele (in bp) is given on the x- axis

637 **Figure 3:** ML phylogenetic tree showing relationships between *Tams1* and *Tlms1* alleles.
 638 Evolutionary distances were computed using the Maximum Composite Likelihood method and are

639 in units of the number of base substitutions per site. All positions containing gaps and missing data
640 were eliminated from the dataset. TA (*T. annulata*), TL (*T. lestoquardi*), and TP (*T. parva*).
641 **Figure 4:** A) Principal Coordinates Analysis of *T. lestoquardi* from four regions in Oman. B)
642 Principal component analysis of *T. annulata* from three regions in Oman. The amount of variation
643 explained by each axis is shown as a percentage of the overall variation.

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Table 1: Prevalence and multiplicity of infection of small ruminant *T. lestoquardi* and bovine *T. annulata* populations in three regions in Oman

Species	Region	No. of animals samples	No. of infected animals (%)	No. of infected animals with multiple genotypes (%)	Mean MOI (SD)
<i>T. annulata</i>					
	Batinah	78	21 (26.9)	21 (100)	3.3 (1.0)
	Dhira	120	57 (47.5)	57 (100)	2.9 (0.8)
	Sharqia	36	19 (48.7)	19 (100)	3.3 (0.7)
<i>T. lestoquardi</i>					
	Batinah	584	57 (9.8)	53 (93)	1.63 (0.33)
	Dhira	357	52 (14.6)	52 (100)	1.65 (0.30)
	Dakhiliya	144	25 (17.4)	24 (96)	1.64 (0.30)
	Sharqia	369	56 (15.2)	53 (95)	1.49 (0.26)

660 **Table 2:** Allelic diversity and heterozygosity at ten micro- and mini-satellite loci from 190 *T. lestoquardi* isolates in Oman

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Region	n	TL_MS05		TL_MS18		TL_MS281		TL_MS04		TL_MS07		TL_MS13		TL_MS16		TL_MS19		TL_MS280		TL_MS25		Average H_e
		H_e	Private alleles	H_e	Private alleles	H_e	Private alleles	H_e	Private alleles	H_e	Private alleles	H_e	Private alleles	H_e	Private alleles	H_e	Private alleles	H_e	Private alleles	H_e	Private alleles	
Batinah	57	0.823	1	0.531	2	0.805	0	0.395	0	0.736	1	0.741	1	0.342	0	0.595	1	0.679	3	0.102	0	0.575
Dakhiliya	25	0.830	0	0.550	1	0.720	0	0.527	0	0.690	0	0.730	0	0.477	0	0.717	2	0.570	1	0.227	1	0.604
Dhair	52	0.850	2	0.521	0	0.824	3	0.115	0	0.705	0	0.847	0	0.419	0	0.632	0	0.816	4	0.111	0	0.584
Sharqia	56	0.882	2	0.581	2	0.858	5	0.410	2	0.655	0	0.815	5	0.510	5	0.733	4	0.816	4	0.105	1	0.637
Overall		0.867		0.548		0.839		0.359		0.705		0.796		0.441		0.665		0.760		0.121		0.610

H_e : gene diversity; heterozygosity

662 **Table 3:** Estimates of genetic diversity of *T. lestoquardi* and *T. annulata* populations in
 663 three regions in Oman

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Species	Region	n	H_e	N_e
<i>T. lestoquardi</i>	Batinah	57	0.575	2.904
	Dhira	52	0.584	3.394
	Sharqia	56	0.637	3.736
	Dakhiliya	25	0.604	2.714
<i>T. annulata</i>				
	Batinah	21	0.854	6.967
	Dhira	57	0.820	8.460
	Sharqia	19	0.833	6.153

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672 **Table 4:** Estimates of genetic diversity of antigen genes among *T. lestoquardi* and *T.*

673 *annulata* isolates

Antigen gene	<i>Tl9</i>	<i>Ta9</i>	<i>Tlms1</i>	<i>Tams1</i>
Parasite species	<i>T. lestoquardi</i>	<i>T. annulata</i>	<i>T. lestoquardi</i>	<i>T. annulata</i>
Polymorphic sites (S)	33	134	19	144
Average number of nucleotide differences (k)	18.600	52.270	3.902	45.832
Nucleotide diversity (π)	0.055	0.155	0.006	0.073
Haplotype diversity (Hd)	0.900	0.978	0.649	0.968

674

675 **Table 5:** Linkage equilibrium among *T. lestoquardi* and *T. annulata* populations in Oman

Species	Region	I^S_A	V_D	L_{MC}	L_{PARA}	Linkage
<i>T. annulata</i>	Batinah	0.0028	1.1174	1.3	1.277	LE
	Dhira	0.0219	1.5537	1.4497	1.4378	LD
	Sharqia	0.0337	1.5627	1.4568	1.4342	LD
	Total	0.0169	1.3841	1.2896	1.2835	LD
<i>T. lestoquardi</i>	Batinah	0.0102	2.1347	2.2212	2.2073	LE
	Dhira	0.0018	1.7015	1.872	1.8559	LE
	Sharqia	0.0752	2.9698	2.0328	2.0174	LD

	Dakhiliya	0.0462	3.0053	2.6308	2.5871	LD
	Total	0.0264	2.284	1.9792	1.975	LD

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677 **Table 6:** Outlier loci outputs Heterozigosity and F_{ST} obtained by using Mcheza (DFDIST

678 algorithm). Statistical significance was obtained as Simulated $F_{ST} < \text{sample } F_{ST}$

Locus	<i>He</i>	F_{ST}
TL_MS05	0.139	-0.026**
TL_MS18	0.191	-0.020**
TL_MS281	0.139	-0.021**
TL_MS04	0.202	0.032
TL_MS07	0.379	0.111
TL_MS13	0.252	-0.016
TL_MS16	0.188	0.016
TL_MS19	0.188	-0.021**
TL_MS280	0.188	0.013
TL_MS25	0.187	0.104

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** Significant at $P < 0.01$

680 **Table 7:** Pair-wise F_{ST} value between *T. lestoquardi* and *T. annulata* populations in

681 Oman

		Batinah	Dhira	Dakhiliya
<i>T. annulata</i>	Batinah	0.0		
	Dhira	0.0257		
	Sharqia	0.0201	0.0266	ND
<i>T. lestoquardi</i>	Batinah	0.0		
	Dhira	0.0256		
	Dakhiliya	-0.0013	0.0457	
	Sharqia	0.0227	0.0281	0.0232

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