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- 1 Utilizing the attached hard ticks as pointers to the risk of infection by *Babesia* and *Theileria*
- 2 species in sika deer (*Cervus nippon yesoensis*), Japan
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24 Abstract

25 Ticks are hematophagous ectoparasites that have a significant impact on their animal hosts. 26 Moreover, along with mosquitoes, they are the main arthropod vectors of disease agents in 27 domestic animals, wildlife and humans. To investigate the occurrence and prevalence of 28 piroplasmids in the ticks, DNA was extracted from 519 hard ticks collected from 116 hunted 29 Hokkaido sika deer (Cervus nippon yesoensis). The success of the DNA extraction was 30 confirmed by touchdown PCR targeting the mitochondrial 16S rDNA gene of ticks. Touchdown 31 PCR and reverse line blot (RLB) hybridization targeting the 18S rRNA gene were used to detect 32 14 piroplasm species. All hard ticks parasitizing Hokkaido sika deer were identified as belonging to the genera Ixodes and Haemaphysalis. A total of 163 samples (31.4%) were positive for 33 34 Babesia and Theileria spp. among different tick species according to RLB hybridization. The 35 DNA from 27.0% (140/519), 10.6% (55/519), 1.7% (9/519), 0.6% (3/519), 0.4% (2/519), and 36 0.4% (2/519) of ticks hybridized to the *Theileria* sp. Thrivae, *Theileria capreoli*, *Babesia* 37 divergens-like, Babesia sp. (Bab-SD), Babesia microti U.S., and B. microti Hobetsu 38 oligonucleotide probes, respectively. The partial sequencing and phylogenetic analyses of the 39 18S rRNA gene confirmed the RLB hybridization results. Further investigations are needed to 40 reveal the epidemiology and respective vectors of these pathogens. 41 **Keywords** 42 Babesia, Hard ticks, Reverse line blot, Sika deer, Theileria

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47 Introduction

Ticks are hematophagous ectoparasites that have significant impacts on their animal hosts, such as by causing anemia and direct skin tissue damage. Moreover, along with mosquitoes, they are the main arthropod vectors of disease agents in domestic animals and humans (Dantas-Torres et al. 2012; Ferrolho et al. 2016; Masatani et al. 2017). Two major families of ticks, Ixodidae (hard ticks) and Argasidae (soft ticks), which include more than 900 tick species, have been recorded globally (Chen et al. 2014). In Japan, 47 tick species have been reported, of which 21 parasitize humans (Takano et al. 2014).

The incidence of tick-borne diseases (TBDs) and the emergence of novel TBD agents have been increasing worldwide during the past two decades (Kernif et al. 2016; Piesman and Eisen 2008). Hence, it is important to investigate the prevalence of tick-borne pathogens to develop effective control strategies against TBDs. Tick-borne apicomplexan parasites, such as *Babesia* and *Theileria* spp., are of great importance worldwide, as they cause diseases in domestic and wild animals. In addition, some *Babesia* spp., such as *Babesia microti*, *Babesia divergens*, and *B. divergens*-like, are considered zoonotic pathogens (Andersson et al. 2017).

In Japan, many studies have identified several *Babesia* and *Theileria* spp. in domestic animals (Fujinaga 1981; Kim et al. 2004; Minami and Ishihara 1980; Ota et al. 2009; Sivakumar et al. 2012), wild animals (Elbaz et al. 2017; Ikawa et al. 2011; Inokuma et al. 2004; Watanabe et al. 2016) and in humans (Wei et al. 2001). These studies have provided information about the potential vector hosts for *Babesia* and *Theileria* spp. in Japan. For example, *Haemaphysalis longicornis* and *Ixodes ovatus* act as vectors for *Babesia ovata* (Sivakumar et al. 2014), and *Ixodes persulcatus, Haemaphysalis megaspinosa, Haemaphysalis douglasi*, and *Ixodes ovatus*

act as vectors for *Theileria orientalis* (Yokoyama et al. 2012); zoonotic *Babesia* spp. such as *B. microti* U.S. and Hobetsu strains have been detected in *I. ovatus* and *I. persulcatus*, respectively
(Zamoto-Niikura et al. 2012), and *B. divergens*-like in *I. persulcatus* (Zamoto-Niikura et al.
2018). Other studies have identified piroplasm species from ticks feeding on animals (Inokuma

73 et al. 2003; Iwakami et al. 2014; Masatani et al. 2017; Tateno et al. 2015).

74 Moreover, the same tick may be co-infected with different piroplasm species (Abdallah et 75 al. 2017) or various other tick-borne pathogens that may have important implications for 76 transmitting pathogens to the animal host (Moutailler et al. 2016). Disease severity may be 77 enhanced if humans or animals are bitten by co-infected ticks, as reported for concurrent 78 babesiosis and Lyme diseases (Golightly et al. 1989; Grunwaldt et al. 1983). This could also 79 have serious consequences for the diagnosis and treatment of TBDs (Chen et al. 2014). 80 Sika deer (*Cervus nippon*) is a native cervid species in Japan (Ohdachi et al. 2009) that 81 has recently expanded in range and number due to environmental and climate changes 82 (Takatsuki 2009). Moreover, sika deer are one of the most important reproductive hosts for 83 several tick species at all developmental stages, such as *H. longicornis*, *Haemaphysalis* 84 flava, Haemaphysalis yeni, H. megaspinosa, I. ovatus, and Amblyomma testudinarium, which 85 have been collected from western and central Japan (Inokuma et al. 2002; Tsukada et al. 2014). 86 Although several piroplasm species have been detected in sika deer from Japan (Elbaz et al. 87 2017; Inokuma et al. 2004; Watanabe et al. 2016; Zamoto-Niikura et al. 2014), few molecular 88 epidemiological studies have been conducted to detect piroplasm species in host-feeding ticks 89 collected from sika deer. Two Theileria spp. have been reported in feeding Haemaphysalis spp. 90 collected from sika deer in southern Japan (Masatani et al. 2017). One of these *Theileria* spp. is

closely related to *Theileria sp.* Thrivae and the other to *T. capreoli*, both of which were detected
previously in sika deer (Elbaz et al. 2017; Watanabe et al. 2016).

93 In our previous study, four piroplasm species were identified in Hokkaido sika deer blood 94 samples: B. divergens-like, a novel Babesia species (here referred to as : Babesia sp. Bab-SD), 95 Theileria sp. Thrivae, and T. capreoli (Elbaz et al. 2017). In addition, Theileria sp. (sika 1) was 96 detected in sika deer collected from Hokkaido, Japan (Shibata et al. 2018) and was found to be 97 different from T. orientalis. Theileria sp. Thrivae is a non-pathogenic Theileria sp. that has been 98 reported from sika deer in Japan (Watanabe et al. 2016). A closely related *Theileria* sp. has been 99 detected in China and named "Theileria cervi" (Liu et al. 2016). However, T. cervi that was 100 reported from white-tailed deer and elk (Cervus canadensis) in USA and Canada (Chae et al. 101 1999) is genetically separated from *Theileria* sp. Thrivae and *T. cervi* of Japan and China, 102 respectively.

Many of tick-borne pathogens (TBPs) are zoonotic (Moustafa et al., 2016); for examples, human granulocytic anaplasmosis, human babesiosis and human monocytic ehrlichiosis are caused by zoonotic tick-borne pathogens and both wildlife and domestic animals act as reservoir or amplifying hosts (Dantas-Torres et al. 2012). Thus, detecting pathogens in ticks collected from animals is crucial to predict their spread and emergence and to protect humans and animals. Hence, the present study investigated the occurrence and prevalence of piroplasmids in ticks collected from sika deer in Hokkaido.

- 110 Material and methods
- 111

Tick collection and identification

A total of 519 ticks were collected from 116 Hokkaido sika deer (*Cervus nippon yesoensis*)
from May to November 2012, which were hunted for nuisance control procedures in two

114 different regions on the Shiretoko Peninsula in Hokkaido Prefecture (Shari and Shibetsu). All

115 collected ticks were either engorged with blood or questing on the surface of the sika deer (non-

116 engorged). The collected ticks were identified under a microscope according to their

117 morphological characteristics (Yamaguti et al. 1971). Subsequently, the ticks were separated into

- 118 groups according to species and stage and stored in 70% ethanol at -20°C until DNA extraction.
- 119

DNA extraction

120 Total DNA was extracted from ticks individually using the Wizard ®Genomic DNA

121 purification Kit (Promega, Madison, WI, USA) per the manufacturer's instructions with

122 modifications (Taylor et al. 2013). The whole ticks were crushed in a 50/50 mixture of the

123 included cell and nuclear lysis solutions supplemented with 17.5 µl proteinase K, followed by

124 incubation at 60°C for 90 min. DNA concentration and quality were assessed by

125 spectrophotometry (NanoDrop 2000, Thermo Fisher Scientific, Yokohama, Japan). The DNA

126 was stored at -20° C until analysis.

127 Touchdown polymerase chain reaction (PCR) to validate tick DNA extraction

128 The success of DNA extraction was confirmed by touchdown PCR targeting the mitochondrial 129 16S rDNA gene (mt-rrs) of ticks. Touchdown PCR was conducted using the PCR System 9700 130 (Applied Biosystems, Foster City, CA, USA), KOD-Plus-Neo high fidelity DNA polymerase kit 131 (Toyobo Co. Ltd., Tokyo, Japan), and Mt-rrs1 (5'- CTG CTC AAT GAT TTT TTA AAT TGC 132 TGT GG-3') and Mt-rrs2 (: 5'-CCG GTC TGA ACT CAG ATC AAG TA-3') primers (Ushijima 133 et al. 2003) to amplify a 401–416 bp fragment. The 25 μ l PCR mixture consisted of 2.5 μ l 10× 134 KOD-Plus-Neo buffer, 2.5 µl dNTPs (2 mM), 1.5 µl 25 mM MgSO₄, 0.75 µl each primer (10 135 pmol/µl), 0.5 µl KOD-Plus-Neo DNA polymerase, 15.5 µl molecular biology-grade water, and 1 µl DNA. The PCR conditions were 94°C for 2 min, a touchdown step of 10 cycles at 94°C for 10 136

137 s, 65°C for 30 s, and 68°C for 30 s with the annealing temperature decreasing by 1°C every cycle. 138 This was followed by 40 cycles of 98°C for 10 s, 55°C for 30 s, 68°C for 30 s, and a final extension 139 of 68°C for 7 min. The PCR products (5 μ l) were mixed with 1 μ l loading buffer (Nippon Gene 140 Co. Ltd., Tokyo, Japan) and examined by 1% agarose gel electrophoresis with ethidium bromide 141 staining, followed by visualization using a UV illuminator.

142

Touchdown PCR and reverse line blot (RLB) hybridization

143 Touchdown PCR targeting the 18S rRNA gene of Babesia and Theileria spp. was performed 144 using the PCR System 9700, KOD-Plus-Neo high fidelity DNA polymerase kit, and the RLB-F2 145 (5'-GAC ACAGGG AGG TAG TGA CAA G-3') and RLB-R2 (biotin-5'-CTA AGA ATT TCA 146 CCT CTG ACA GT-3') primer pair (Gubbels et al. 1999; Matjila et al. 2004). The PCR mixture 147 was used as described above to amplify a 460–540 bp segment. One microliter of molecular grade 148 water and genomic DNA samples of B. divergens-like and Theileria sp. Thrivae were used as negative and positive controls for Babesia and Theileria spp., respectively. The PCR conditions 149 150 were 94°C for 2 min, a touchdown step of 10 cycles at 94°C for 10 s, 67°C for 30 s, and 68°C for 151 30 s with a 1°C decrease in annealing temperature every cycle. This was followed by 40 cycles at 152 98°C for 10 s, 57°C for 30 s, 68°C for 30 s, and a final extension at 68°C for 7 min. The PCR 153 products were examined by 1% agarose gel electrophoresis with ethidium bromide staining 154 followed by visualization using a UV illuminator.

The PCR products were subjected to RLB hybridization as described previously (Kong and Gilbert 2006; Moustafa et al. 2016) with modifications. Briefly, a 15×15 cm Biodyne C membrane (Pall Life Science, Ann Arbor, MI, USA) was activated with 20 ml 16% 1-ethyl-3-(3dimethyl-amino-propyl) carbodiimide (Sigma Aldrich, St. Louis, MO, USA) for 10 min at room

159 temperature. The membrane was washed gently with MilliQ water for 2 min and placed in the 160 Miniblotter MN45 (Immunetics, Boston, MA, USA). This study included 19 oligonucleotide 161 probes: 17 previously published probes (Elbaz et al. 2017) and 2 probes designed in this study. 162 The designed probes (T. capreoli 5'-ATACGAGTTTTTGCATTGTG-3' and Babesia sp. (Bab-163 SD) 5'-GTTGGCTTTTCTTATTACTTTGA-3') linked to a C6 amino linker were obtained from 164 Sigma Aldrich (Tokyo, Japan) and reconstituted as 100 pmol/µl solutions by using Tris-EDTA 165 (TE) buffer (PH 8.0) (Nippon Gene, Japan). Ten microliters of each 100 pmol/µl probe solution 166 were diluted in 0.5 M NaHCO₃ to a final volume of 170 μ l. The openings in the miniblotter were 167 filled with 150 µl each diluted oligonucleotide, and the membrane was incubated for 5 min at room 168 temperature. The membrane was inactivated in 250 ml 0.1 M NaOH with gentle shaking for 169 exactly 8 min at room temperature, then washed with prewarmed 250 ml $2\times$ SSPE/0.1% SDS at 170 60° C. Finally, the membrane was sealed in a plastic bag with 15 µl 20 mM EDTA for future use. 171 Ten microliters of each PCR product were diluted in $2 \times SSPE/0.1\%$ SDS to a final volume of 170 172 µl, denatured by heating at 100°C for 10 min, and immediately cooled on ice. The denatured 173 mixtures were allowed to hybridize to the membrane at 60°C for 1 h. Afterwards, the membrane 174 was washed twice in 250 ml 2× SSPE/0.5% SDS at 52°C for 10 min and then incubated with 175 diluted peroxidase-labeled NeutrAvidin (Thermo Fisher Scientific, Waltham, MA, USA) at 42°C 176 for 45–60 min. The NeutrAvidin was removed by washing twice with 250 ml 2× SSPE/0.5% SDS 177 at 42°C for 10 min and with $2 \times$ SSPE twice at room temperature for 5 min. The membrane was 178 exposed to 15 ml Immobilon[™] Western Chemiluminescent HRP Substrate (Millipore, Tokyo, 179 Japan) for 5 min and photographed using the Ez-Capture MG/ST (ATTO Corp. Tokyo, Japan). To 180 remove the PCR products, the membrane was washed twice in pre-warmed 1% SDS at 90°C for 30 min and washed with 250 ml 20 mM EDTA. The membrane was sealed in a plastic bag with
15 ml 20 mM EDTA and stored at 4°C for future reuse.

183

Cloning and sequencing

184 The RLB results were confirmed by cloning and sequencing 12 randomly selected positive 185 PCR products. A 460-540 bp fragment was amplified from the 18S rRNA gene of piroplasm 186 species using the KOD-Plus-Neo high fidelity DNA polymerase kit with the abovementioned 187 RLB-F2 and RLB-R2 primers and PCR conditions. The PCR products were examined by 188 electrophoresis, and the band corresponding to 460–540 bp was excised for purification using the 189 NucleoSpin[®] Gel and PCR clean up kit (Macherey-Nagel, Düren, Germany). The purified PCR 190 products were attached to dA nucleotides by mixing $0.9 \ \mu l \ 10 \times KOD$ -Plus-Neo buffer, $0.9 \ \mu l \ 2$ 191 mM dNTPs, 0.54 µl 25 mM MgSO₄, 6.66 µl each purified PCR product, and 1 µl 10× A-192 attachment mix (Toyobo) at 60°C for 10 min. One microliter of each dA-attached product was 193 cloned into the pMD20 T-vector (Takara Bio, Otsu, Japan). Five colonies per sample were selected 194 and screened by EmeraldAmp MAX PCR (Takara Bio) as described in the manufacturer's manual. 195 The purified PCR products were sequenced using the Big-Dye Terminator version 3.1 Cycle 196 Sequencing Kit and purified using the BigDye® XTerminatorTM Purification Kit (Applied 197 Biosystems). Subsequently, the purified products were sequenced using the ABI PRISMTM 310 198 genetic analyzer (Applied Biosystems). BLASTn searches were performed to compare the 199 sequences to reference sequences recorded in GenBank for identification.

200 Phylogenetic analysis

201 The 18S rRNA partial sequences were examined by BLASTn search against the *Babesia* and 202 Theileria spp sequences in GenBank and submitted to the DNA Data Bank of Japan (DDBJ). The 203 obtained sequences were aligned in MEGA software version 7 with representative 18S rRNA gene 204 sequences of the previously identified Babesia and Theileria spp. in the GenBank to know its 205 taxonomic position. The "Find Best DNA/Protein Models" option in MEGA 7 was used to select 206 the most suitable substitution model for the phylogenetic analysis. The phylogenetic trees were 207 constructed using the Maximum Likelihood method based on the Tamura-Nei model (Tamura and 208 Nei 1993) using the gamma distribution in the same software and a full 18S rRNA gene sequence 209 for *Plasmodium falciparum* (XR_002966679) was used as an outgroup. Node support was assessed 210 using 1,000 bootstrap replicates (Kumar et al. 2016).

211 Statistical analysis

Linear mixed models (LMM) was performed to investigate differences in the prevalence of *Theileria* sp. Thrivae and *T. capreoli*. We fitted LMM with predictive variables (tick species, stage, and status) in presence of sampling location as a random effect variable. We used "lmer" function in R package lme4 to perform the LMM. The correlations among the detected piroplasm species were investigated by the chi-square test. The statistical analyses were performed using SPSS version 21.0 (SPSS Inc., Chicago, IL, USA). A P-value < 0.05 was considered significant.

- 218 Results
- 219

Identifying the tick species and detecting tick DNA

220 The collected ticks were identified as belonging to the genera *Ixodes* and *Haemaphysalis*.

221 The *Ixodes* ticks included 330 (63.6%), 40 (7.7%), 11 (2.1%) and 110 (21.2%) *I. persulcatus*

nymph, larvae, adult and *I. ovatus* adults, respectively. The remaining collected ticks including
15 (2.9%), 11 (2.1) and 2 (0.4%) were identified as *H. japonica* adult, nymph and larvae,
respectively. Of these ticks, 70 (18.4%) and 66 (60%) of the collected *I. persulcatus* and *I. ovatus* were engorged with blood; while 311 (81.6%), 44 (40%) and 28 (100%) of the collected *I. persulcatus*, *I. ovatus* and *H. japonica* were non-engorged, respectively.

227 Detection and identification of *Babesia* and *Theileria* spp. in tick samples by 228 touchdown PCR/RLB hybridization

229 Touchdown PCR and RLB hybridization, used to detect a partial fragment of the 18S rRNA

230 gene of *Babesia* and *Theileria* spp., showed that 163 of the 519 (31.7%) DNA samples examined

231 were positive for *Babesia* and *Theileria* spp. including *Theileria* sp. Thrivae, *T. capreoli*, *B.*

divergens-like, Babesia sp. (Bab-SD), B. microti U.S., and B. microti Hobetsu (Table 1).

Furthermore, the prevalence of *Theileria* sp. Thrivae was the highest among the detected

234 piroplasms and was detected mostly in adult *I. ovatus*.. In addition, the highest prevalence

235 *Theileria* sp. Thrivae was found in the nymph stage of *H. japonica* (45.5%; 5/11) and adult of *I.*

236 *persulcatus* (36.4%; 4/11). (Table 1).

237 In addition, *T. carpreoli* was found in all tick species and stages except *H. japonica* larvae, and

the highest prevalence were observed in the adult stages of *I. ovatus*: 22.7% (25/110) and *H.*

239 *japonica*: 26.7% (4/15). A total of 7.5% (3/40) and 9.1% (1/11) of the DNA samples extracted

from engorged I. persulcatus larvae and adult I. persulcatus were positive for T. capreoli,

respectively.

The highest prevalence of *B. divergens*-like were 6.7% (1/15) in *H. japonica*. The *Babesia* sp. (Bab-SD) was detected only in *Ixodes* spp., with prevalence of 0.3% (1/330) and

1.8% (2/110) in *I. persulcatus* nymphs and *I. ovatus* adults, respectively. In addition, *B. microti*

245 U.S. and Hobetsu were identified in *Ixodes* spp., with prevalence of 0.5% (2/381) in *I*.

246 *persulcatus* and 1.8% (2/110) in *I. ovatus*, respectively.

Of the 163 ticks positive for piroplasm species, the co-infection rate was 26.4% (43/163) and

included 39 double, 3 triple, and 1 quadruple infection cases. The prevalence of double co-

249 infection were 79.1% (34/43), 9.3% (4/43), and 2.3% (1/43) between *Theileria* sp. Thrivae and

250 T. capreoli, Theileria sp. Thrivae and B. divergens-like, and T. capreoli and B. divergens-like,

respectively (Table 2). Of the three ticks with triple co-infection, two were infected with

252 *Theileria* sp. Thrivae, *T. capreoli*, and *B. divergens*-like, and the third tick was infected with

253 Theileria sp. Thrivae, Babesia sp. (Bab-SD), and B. microti (Hobetsu type). In addition, only one

tick was infected with four pathogens: Theileria sp. Thrivae, T. capreoli, the Babesia sp. (Bab-

SD), and *B. divergens*-like. The highest co-infection rate, 53.5% (23/43), was detected in adult

ticks, whereas a 41.9% (18/43) co-infection rate was found in nymph ticks.

DNA sequencing and molecular characterization

258 The sequencing results yielded 12 partial 18S rRNA gene sequences (Table 3), which were 259 classified as six piroplasm species: two Theileria spp. and four Babesia spp. (Figs. 1 and 2). The 260 sequences from I. persulcatus (LC336717–LC336718), I. ovatus (LC336719), and H. japonica 261 (LC336720) were identical to those from the uncultured *Theileria* spp. (LC215385–LC271213) 262 and *Theileria* sp. Thrivae (AB981972), which were previously detected in sika deer in Japan and 263 are 99% similar to *Theileria* sp. sequences (LC169094 and LC169091) found in *Haemaphysalis* 264 sp. collected from sika deer and vegetation, respectively. Our sequences obtained from I. 265 persulcatus (LC336721 and LC336724), a H. japonica (LC336722), and I. ovatus (LC336723)

266	were identical to each other and were identical to the uncultured Theileria spp. sequence
267	(LC271206) from sika deer in Japan and 99% similar to sequences of <i>T. capreoli</i> (KJ188219)
268	from red deer in China and of Theileria sp. (LC169093) from a feeding Haemaphysalis sp.
269	detected on sika deer in Japan.
270	The obtained sequence (LC336725) was identical to that of the Babesia sp. (Bab-SD)
271	(LC215388), whereas the sequence LC336727 was identical to a <i>B. divergens</i> -like sequence
272	(LC215387) detected in sika deer from Japan. The sequences (LC336726 and LC336728) were
273	clustered with B. microti. The first sequence was identical to that of B. microti Hobetsu
274	(AB050732) from a field rodent (Apodemus speciosus), while the other sequence was identical
275	with that of the B. microti U.S. strain (LC127372) from I. persulcatus in Japan.
276	Prevalence of <i>Theileria</i> spp. according to tick variables
277	In this study, LMM test was used to analyze the results of Theileria spp., but not for Babesia
278	spp. because of the low number of positive Babesia spp. samples. The prevalence of Theileria
279	sp. Thrivae and <i>T. capreoli</i> were significantly higher in the engorged ticks than in the non-
280	engorged ones (LMM: $P < 0.001$ and $P < 0.05$), respectively (Table 4 and Fig. 3). Additionally, a
281	significant positive correlation between Theileria sp. Thrivae and T. capreoli was detected by

282 chi-square test (P < 0.01) (Table 5).

283 **Discussion**

Ticks are obligate blood feeders and important vectors of a large number of pathogens infecting animals and humans (Azmi et al. 2016). Despite that many studies have reported piroplasm species from sika deer in Japan (Elbaz et al. 2017; Inokuma et al. 2004; Shibata et al.

2018; Watanabe et al. 2016; Zamoto-Niikura et al. 2014), only few published reports are available
on piroplasm species from feeding ticks detected on sika deer in Hokkaido (Masatani et al. 2017).

Here, a total of six piroplasms, *Theileria* sp. Thrivae, *T. capreoli*, *B. divergens*-like, *Babesia* sp. (Bab-SD), *B. microti* U.S., and *B. microti* Hobetsu, were identified. Hokkaido sika
deer are potential hosts for *Theileria* sp. Thrivae, *T. capreoli*, *B. divergens*-like, and the *Babesia*sp. (Bab-SD) (Elbaz et al. 2017). However, *B. microti* strains have not been reported in
Hokkaido sika deer but were detected previously in rodents; therefore, the ticks infected with *B. microti* strains in this study may have been infected during previous life stages when those ticks
fed on other animal hosts.

296 Our results revealed *Theileria* sp. Thrivae and *T. capreoli* are the most prevalent species in 297 the ticks examined, and both organisms were detected in all examined stages of *Ixodes* nymphs 298 and adults of *H. japonica*. This observation is in agreement with the high prevalence of *Theileria* 299 sp. Thrivae in sika deer, indicating the persistence of infection in sika deer (Elbaz et al. 2017). 300 Recently, a high prevalence of *Theileria* spp. in sika deer (97.8%; 89/91) was detected in Hokkaido, 301 Japan (Shibata et al. 2018). Here, most of the ticks positive for *Theileria* spp. were engorged with 302 sika deer blood, which may have caused this high prevalence. Although the prevalence of T. 303 *capreoli* in Hokkaido sika deer is unknown, the results of this study suggest that sika deer could 304 be the main animal host for this piroplasm species in the Hokkaido ecosystem. Moreover, the 305 finding that the sequences from ticks collected in Kyushu and Hokkaido were almost identical 306 suggests that *Theileria* sp. Thrivae and *T. capreoli* are distributed throughout Japan (Masatani et 307 al. 2017). However, this study cannot confirm that these tick species are the vector hosts to the 308 detected pathogens, because the blood meal from infected host could be the source of the positive

309 results. It is required to examine salivary glands of tick species to confirm that these ectoparasites310 act as epidemiological vectors.

311 The prevalence of *Theileria* sp. Thrivae and *T. capreoli* in *H. japonica* in this study are 312 higher than the previously reported rates of 6.3% (2/32) and 12.5% (4/32) (Masatani et al. 2017), 313 respectively. This is possibly due to the higher sensitivity of the RLB technique compared with 314 conventional PCR assays. In addition, most of ticks that were examined in this study were 315 engorged with sika deer blood, which can lead to an increase in the prevalence of *Theileria* species. 316 Interestingly, both *Theileria* spp. were detected in the larval stage of *I. persulcatus*; however, 317 *Theileria* spp. are not known to be transmitted via a transovarian route (Bhattacharyulu et al. 1975). 318 Positivity for *Theileria* spp. was likely because all larvae were engorged with infected sika deer 319 blood.

The detected *Babesia* spp., *B. divergens*-like, *B. microti* U.S., and *B. microti* Hobetsu are zoonotic pathogens. The highest rate of *B. divergens*-like recovery was from an *I. persulcatus* nymph in this study and was in accordance with a previously published report on questing *I. persulcatus* (1.3%; 11/845) from Japan (Zamoto-Niikura et al. 2018).

In this study, the *Babesia* sp. (Bab-SD) was detected in *Ixodes* spp. only. Surprisingly, the *Babesia* sp. (Bab-SD) was found in a non-engorged *I. persulcatus* nymph, suggesting that the larvae of this tick were infected before molting into nymphs, either from an infected reservoir host or by transovarian transmission. However, not all larvae examined in this study were infected by the *Babesia* sp. (Bab-SD); thus, further study on this *Babesia* sp. is needed to understand its life cycle.

330 Sika deer are not hosts for *B. microti*; however, this study found that *Ixodes* spp. feeding 331 on sika deer were positive for B. microti strains. Furthermore, the B. microti U.S. and Hobetsu 332 types were detected in *I. persulcatus* and *I. ovatus*, respectively. This result is consistent with 333 previous studies reporting that I. persulcatus and I. ovatus are vectors for B. microti U.S. and B. 334 *microti* Hobetsu in Japan and Russia, respectively (Katargina et al. 2011; Rar et al. 2011; Rar et 335 al. 2014; Zamoto-Niikura et al. 2012). The prevalence of B. microti U.S. type in I. persulcatus in 336 this study was in accordance with the rates of those previously published reports from Japan and 337 Russia (Katargina et al. 2011; Rar et al. 2011; Rar et al. 2014; Zamoto-Niikura et al. 2012).

338 The partial 18S rRNA gene sequences were identified as belong to two *Theileria* spp. and 339 four Babesia spp. The two Theileria spp. clustered with Theileria sp. Thrivae and T. capreoli, 340 which were detected in sika deer in Japan. The *Babesia* spp. sequences were associated with four 341 branches. The first branch clustered with *B. divergens*-like sequences, the second branch with the 342 Babesia sp. (Bab-SD) sequences detected in sika deer in Japan (Elbaz et al. 2017), and the 343 remaining two with *B. microti* sequences, one with a previously published *B. microti* U.S. sequence 344 from *I. persulcatus* in Japan (Zamoto-Niikura et al. 2016) and the other with *B. microti* Hobetsu 345 type detected in a field rodent (Apodemus speciosus) (Wei et al. 2001).

The prevalences of both *Theileria* spp. detected in this study were significantly higher in the engorged ticks than in non-engorged ones, which could be due to the high infection rates of both *Theileria* spp. in sika deer blood, indicating the persistence of infection in sika deer.

Co-infection of ticks with more than one pathogen is not uncommon, and one tick can be infected with up to five pathogens (Eshoo et al. 2014; Moutailler et al. 2016; Reis et al. 2011). Multiple pathogens in individual ticks increases the risk of co-transmission to, and co-infections 352 in, humans and animals (Moutailler et al. 2016). In this study, a significant co-infection correlation 353 was shown between *Theileria* sp. Thrivae and *T. capreoli*, which may have occurred because both 354 *Theileria* spp. have similar life cycles. This was supported by the phylogenic analysis showing that 355 the two Theileria spp. belong to the same Theileria spp. cluster (Inokuma et al. 2008; Li et al. 356 2014). The association between *Theileria* and *Babesia* spp. reported in double, triple, or quadruple 357 co-infection is likely due to shared reservoir hosts or co-feeding during different life stages. 358 Additionally, the highest number of co-infection cases was observed in adult ticks, which may be 359 because of the greater opportunity of adult ticks to receive more pathogens from host feeding via 360 transstadial transmission.

361 Conclusion

362 Ticks and TBDs constitute a growing burden for human and animal health. This study 363 revealed that ticks collected from sika deer are exposed to a high diversity of piroplasm species 364 and demonstrated the ecology of these pathogens endemic to Hokkaido. The sika deer is an 365 amplifying host for Theileria sp. Thrivae, T. capreoli, B. divergens-like, and Babesia sp. (Bab-366 SD). In addition, zoonotic TBPs, such as B. divergens-like and B. microti U.S. and Hobetsu, were 367 detected in the ticks of this study. Because this is the first report of piroplasm species detected in 368 feeding ticks collected from sika deer in Hokkaido, further investigations are needed to reveal the 369 epidemiology and respective vectors of these pathogens.

370 Competing interests

The authors declare that they have no competing interests.

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376 English check

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	Tick	samples				Detected	l pirolplasm sp	p.		
Species	Stage	N Engorged	o.tested Non-engorged	<i>Theileria</i> sp. Thrivae	T. capreoli	B. divergens-like	Babesia sp. (Bab-SD)	B. microti U.S.	<i>B. microti</i> Hobetsu	Co-infection
	Larvae	15	25	8 (20%)	3 (7.5%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	2
I	Nymph	53	277	74 (22.4%)	21 (6.4%)	7 (2.1%)	1 (0.3%)	1 (0.3%)	0 (0.0%)	18
1. persuicatus	Adult	2	9	4 (36.4%)	1 (9.1%)	0 (0.0%)	0 (0.0%)	1 (10%)	0 (0.0%)	1
	Sub total	70	311	86 (22.6%)	25 (6.6%)	7 (1.8%)	1 (0.3%)	2 (0.5%)	0 (0.0%)	21
I. ovatus	Adult	66	44	45 (40.9%)	25 (22.7%)	1 (0.9%)	2 (1.8%)	0 (0.0%)	2 (1.8%)	20
	Sub total	66	44	45 (40.9%)	25 (22.7%)	1 (0.9%)	2 (1.8%)	0 (0.0%)	2 (1.8%)	20
	Larvae	0	2	0 (0.0%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	0
Haemaphysalis	Nymph	0	11	5 (45.5%)	1 (9.1%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	0
japonica	Adult	0	15	4 (26.7%)	4 (26.7%)	1 (6.7%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	2
	Sub total	0	28	9 (32.1%)	5 (17.9%)	1 (3.6%)	0 (0.0%)	0 (0.0%)	0 (0.0%)	2
Total			519	140 (27%)	55 (10.6%)	9 (1.7%)	3 (0.6%)	2 (0.4%)	2 (0.4%)	43

Table 1 The prevalence of piroplasm species in ticks collected from Hokkaido sika deer by RLB

- - -

Co- infection Piroplasms I. persulcatus I. ovatus H. japonica **39@**1 Theileria sp. Thrivae and T. capreoli Double infection Theileria sp. Thrivae and B. divergens like T. capreoli and B. divergens like Theileria sp. Thrivae, T. capreoli and B. divergens like Triple infection Theileria sp. Thrivae, Babesia sp. (Bab-SD) and B. microti Hobetsu Quadruple infection Theileria sp. Thrivae, T. capreoli, Babesia sp. (Bab-SD) and B. divergens like 395

391 Table 2. A summary of co-infections between piroplasm species in ticks collected from Hokkaido sika deer by RLB

Table 3 Samples of sequencing

Number	Sample I.D.	Accession number	RLB results
1	130308 I.P.N	LC336717	
2	130304 I.P.A	LC336718	- Theileria on Thrivee
3	120619D3 I.O.A	LC336719	- <i>Theneria</i> sp. Thirvae
4	120618D21 H.N	LC336720	_
5	120925D2 I.P.L	LC336721	
6	120610D14 H.A	LC336722	_
7	120219D13 I.O.A	LC336723	- Theileria capreoli
8	120219D3 I.P.N	LC336724	
9	12012D3 I.O.A	LC336725	Babesia sp. (Bab-SD)
10	120618D13 I.O.A	LC336726	Babesia microti Hobetsu
11	120928N2 I.P.N	LC336727	Babesia divergens-like
12	130305 I.P.A	LC336728	Babesia microti U.S.

409 I.P.N: I. persulcatus nymph, I.P.A: I. persulcatus adult, I.P.L: I. persulcatus larvae, I.O.A: I. ovatus adult,

410 H.N: *H. japonica* nymph and H.A: *H. japonica* adult

Table 4. The final models for the prevalence of *Theileria* sp. Thrivae and *T. capreoli* in ticks collected from Hokkaido sika deer.

		Р	revalence	of Theiler	<i>ia</i> sp. Thr	ivae		Prevalen	ce of <i>Thei</i>	leria cam	reoli	410
Random	Predictors	Estimate	Std. Err	df	t value	$\frac{1}{\Pr(> t)}$	Estimate	Std. Err	df	t value	Pr(> t)	417
	(Intercept)	0.44	0.12	40.98	3.68	0.00 ***	0.29	0.07	141.83	3.92	0.00 **	41/ *
	Species_I. ovatus	0.04	0.11	512.22	0.38	0.71	-0.04	0.07	434.92	-0.47	0.64	118
Location	Species_I. persulcatus	-0.11	0.10	509.42	-1.11	0.27	-0.08	0.07	503.20	-1.14	0.26	410
Location	Stages_Larvae	-0.07	0.12	512.99	-0.57	0.57	-0.10	0.08	512.99	-1.21	0.23	419
	Stages_Nymph	0.06	0.10	512.67	0.58	0.56	-0.09	0.07	499.98	-1.20	0.23	117
	Status_Non-engorged	-0.18	0.05	512.25	-3.82	0.00 ***	-0.08	0.03	512.60	-2.36	0.02 *	420

			T. capreoli		434
		Positive	Negative	Total	435
<i>Theileria</i> sp.	Positive	37*	103	140	_+
Thrivae	Negative	18	361	379	436
	Total	55	464	519	
					437

Table 5 Number of Ticks infected with *Theileria* sp. Thrivae and/or *T. capreoli*.

438 * Correlation between *Theileria* sp. Thrivae and *T. capreoli* was significant positive (P<0.01) using chi-square (χ^2) test in SPSS

446 Fig. 1 Phylogenetic tree based on nucleotide 18S rRNA gene sequences of *Babesia* spp. from 447 GenBank. Sequences from the present study are labeled in bold. The DNA sequences from this 448 study was deposited to DDBJ as accession number from LC336725 to LC336728. Phylogenetic 449 trees were constructed using the Maximum Likelihood method based on the Tamura-Nei model 450 (Tamura and Nei 1993). Initial tree(s) for the heuristic search were obtained automatically by 451 applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using 452 the Maximum Composite Likelihood (MCL) approach, and then selecting the topology with 453 superior log likelihood value. A discrete Gamma distribution was used to model evolutionary 454 rate differences among sites (5 categories (+G, parameter = 0.2599)). The analysis involved 22 455 nucleotide sequences. All positions containing gaps and missing data were eliminated. There was 456 a total of 174 positions in the final dataset. Evolutionary analyses were conducted in MEGA7 457 (Kumar et al. 2016).

458 Fig. 2 Phylogenetic tree based on nucleotide 18S rRNA gene sequences of *Theileria* spp. from 459 GenBank. Sequences from the present study are labeled in bold. The DNA sequences from this 460 study was deposited to DDBJ as accession number from LC336717 to LC336724. Phylogenetic 461 trees were constructed using the Maximum Likelihood method based on the Tamura-Nei model 462 (Tamura and Nei 1993). Initial tree(s) for the heuristic search were obtained automatically by 463 applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using 464 the Maximum Composite Likelihood (MCL) approach, and then selecting the topology with 465 superior log likelihood value. A discrete Gamma distribution was used to model evolutionary rate differences among sites (5 categories (+G, parameter = 0.3910)). The analysis involved 31 466 467 nucleotide sequences. All positions containing gaps and missing data were eliminated. There was

468 a total of 177 positions in the final dataset. Evolutionary analyses were conducted in MEGA7

- 469 (Kumar et al., 2016).
- **Fig.** 3 Descriptive analysis of the prevalence of *Theileria* sp. Thrivae and *T. capreoli* in (a)
- 471 engorged and non-engorged ticks and (b) the developmental stages of ticks

Declarations

- **Funding** (This study was supported in part by the Program for Leading Graduate Schools
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- **Conflicts of interest/Competing interests** (The authors declare that they have no competing
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- **Ethics approval** (Not applicable)
- **Consent to participate** (Not applicable)
- **Consent for publication** (Not applicable)
- 482 Availability of data and material (The DNA sequence data were registered at the DNA
- 483 DataBank of Japan (DDBJ) under the accession numbers LC336717 LC336728)
- **Code availability** (Not applicable)

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0.050

	LC336722	
	AB012189 Theileria sp. Sika Deer Japan	
	LC336723	
	61 LC336724	
	LC336721	
e	51 LC169093 <i>Theileria</i> sp. Ticks Japan	
	LC271206 <i>Theileria</i> sp. Sika Deer Japan	
	LC271197 <i>Theileria</i> sp. Sika Deer Japan	
9'	7 KJ188208 Theileria capreoli Deer China	
	- KU958547 Theileria capreoli Ticks Hungary	
	KX470616 Theileria sp. Ticks Slovakia	
	KX470614 Theileria sp. Ticks Slovakia	
LC336720		
	LC336719	
	LC336718	
	LC336717	
	LC215385 Theileria sp. Sika deer Japan	
	LC271213 Theileria sp. Sika deer Japan	
99	LC169094 <i>Theileria s</i> p. Ticks Japan	
	LC169091 Theileria sp. Ticks Japan	
	AB981972 Theileria sp. Thrivae Sika deer Japan	
	HQ184411 Theileria cervi Sika deer China	
	EU126895 Theileria sp. Sika deer Japan	
	99 – HQ895985 Theileria parva African buffalo South Africa	
	KT736498 <i>Theileria annulata</i> Cattle India	
	99 DQ104611 Theileria buffeli China	
HM538205 <i>Theileria buffeli</i> Cattle China		
AB016074 Theileria sergenti Cattle Japan		
XR696404 Theileria orientalis Cattle Japan		
XR696411 Theileria orientalis Cattle Japan		
	XR 002966679 Plasmodium falciparum	

0.050



Fig. 3