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20 Abstract

Ticks (Acari: Ixodida) transmit a greater variety of pathogens than any other blood-feeding
group of arthropods. While numerous microbes have been identified inhabiting Australian
Ixodidae, some of which are related to globally important tick-borne pathogens, little is
known about the bacterial communities within ticks collected from Australian wildlife. In this
study, 1,019 ticks were identified on 221 hosts spanning 27 wildlife species. Next-generation

26 sequencing was used to amplify the V1-2 hypervariable region of the bacterial 16S rRNA 27 gene from 238 ticks; Amblyomma triguttatum (n=6), Bothriocroton auruginans (n=11), 28 Bothriocroton concolor (n=20), Haemaphysalis bancrofti (n=10), Haemaphysalis bremneri 29 (n=4), *Haemaphysalis humerosa* (n=13), *Haemaphysalis longicornis* (n=4), *Ixodes antechini* 30 (n=29), Ixodes australiensis (n=26), Ixodes fecialis (n=13), Ixodes holocyclus (n=37), Ixodes 31 *myrmecobii* (n=1), *Ixodes ornithorhynchi* (n=10), *Ixodes tasmani* (n=51) and *Ixodes* 32 trichosuri (n=3). After bioinformatic analyses, over 14 million assigned bacterial sequences 33 revealed the presence of recently described bacteria 'Candidatus Borrelia tachyglossi', 'Candidatus Neoehrlichia australis', 'Candidatus Neoehrlichia arcana' and 'Candidatus 34 35 Ehrlichia ornithorhynchi'. Furthermore, three novel Anaplasmataceae species were identified 36 in the present study including; a Neoehrlichia sp. in I. australiensis and I. fecialis collected 37 from quenda (Isoodon fusciventer) (Western Australia), an Anaplasma sp. from one B. 38 concolor from echidna (Tachyglossus aculeatus) (New South Wales), and an Ehrlichia sp. 39 from a single *I. fecialis* parasitising a quenda (WA). This study highlights the diversity of 40 bacterial genera harboured within wildlife ticks, which may prove to be of medical and/or 41 veterinary importance in the future. 42

Keywords: Microbiome; ticks; Ixodida; wildlife; marsupials; Anaplasmataceae

45 1. Introduction

46 Current estimates suggest that approximately 17% of all infectious diseases of humans are 47 vector-borne (Rinker et al., 2016) and global trends show that vector-borne diseases (VBDs) 48 are rising at a rapid rate (Jones et al., 2008; Morens and Fauci, 2012). The complex interplay 49 between pathogen, vector, host(s) and the environment make VBDs particularly challenging 50 to understand. In addition, factors such as climate change (Ostfeld and Brunner, 2015), land

use (Ferrell and Brinkerhoff, 2018), feral animal populations (Merrill et al., 2018) and the
microclimate within a landscape (Dobson et al., 2011) can further influence the prevalence
and distribution of VBDs.

54

55 Ticks (Acari: Ixodida) comprise a group of haematophagous (blood feeding) arthropods with 56 over 900 species described globally (Guglielmone et al., 2014; Mans et al., 2019). Ticks are 57 known to transmit various pathogens; however, they also harbour a range of endosymbiont 58 and commensal species (Špitalská et al., 2018). The epidemiology of recognised tick-borne 59 diseases (TBDs) in the northern hemisphere demonstrates that wildlife serve as sentinels and 60 can be used to monitor the presence and distribution of tick-borne pathogens (TBPs). 61 Importantly, research has shown that some wildlife species act as dilution hosts for certain 62 TBPs, whereas others may act as amplification hosts (LoGiudice et al., 2003). 63 64 The complexity of TBDs means that studies are increasingly shifting away from isolated 65 species-specific studies toward ecosystem-based, collaborative research (Estrada-Peña et al., 66 2013; Pfäffle et al., 2013). Metabarcoding provides an informative molecular tool to 67 characterise the bacterial diversity in ticks. Worldwide, next-generation sequencing (NGS)-68 based analyses have been applied to a range of tick species that are important from medical 69 and veterinary perspectives, including Amblyomma americanum (Ponnusamy et al., 2014), 70 *Ixodes ricinus* (Bonnet et al., 2014), and *Rhipicephalus microplus* (Andreotti et al., 2011). 71

Metabarcoding studies of the bacterial microbiome of Australian ticks have been reported
only recently, with the first bacterial profiling by NGS conducted on the human-biting tick *Ixodes holocyclus* (Gofton et al., 2015a). These authors identified a highly abundant
endosymbiont '*Candidatus* Midichloria mitochondrii' (CMm) and after blocking the

76 amplification of this organism, a greater bacterial diversity was revealed, including a number 77 of novel microbes (Gofton et al., 2015a,b). Critically, in contrast to many parts of the world where multiple TBPs have been elucidated within well-studied tick-host-environment 78 79 ecologies, there is a relative dearth of such information available for Australia. With this in 80 mind, the aims of our study were to survey the bacterial communities present in ticks 81 collected from Australian wildlife and to investigate their genetic relatedness to 'taxa of 82 interest', i.e. tick-associated pathogenic and endosymbiotic organisms (Parola and Raoult, 83 2001; Mediannikov and Fenollar, 2014; Sumrandee et al., 2016). 'Taxa of interest' in the 84 present study were defined as genera within alphaproteobacteria, gammaproteobacteria and 85 spirochaetes known to be transmitted by ticks in other parts of the world, specifically; 86 Anaplasma, Bartonella, Borrelia, Coxiella, Ehrlichia, Francisella, Midichloria, 87 Neoehrlichia, Rickettsia and Rickettsiella. (Ahantarig et al., 2013; Vayssier-Taussat et al., 88 2015; Bonnet et al., 2017; de la Fuente et al., 2017). 89 90 2. Materials and methods 91 92 2.1 Sample collection and identification 93 1,019 ticks were sourced opportunistically from wildlife in Australia by veterinarians, 94 veterinary nurses, wildlife carers, researchers and via submissions from members of the 95 public, and preserved in 70% ethanol before being shipped to Murdoch University, Western 96 Australia (WA), for identification. Ticks were identified morphologically to life stage and 97 species using keys and species' descriptions (Roberts, 1970; Barker and Walker, 2014). 98 Details of sample collection are available in Supplementary File S1.

99

100 2.2 DNA Extraction

101 A sub-sample of ticks (*n*=238) was chosen for DNA extraction and bacterial profiling. Ticks 102 were selected to represent as many tick species, hosts, geographical regions and life stages as possible within the study. Prior to DNA extraction, individual ticks were surface-sterilised in 103 10% hypochlorite solution, rinsed in 70% ethanol and DNA-free PBS, and then air-dried. 104 105 Total genomic DNA (gDNA) was extracted using the Qiagen DNeasy Blood and Tissue kit 106 (Qiagen, Germany) following the manufacturer's recommendations with the following 107 modifications; ticks were placed in a 2 mL safe lock Eppendorf tube with a 5 mm steel bead, frozen in liquid nitrogen for 1 minute and homogenised by shaking at 40 Hz in a Tissue Lyser 108 LT (Qiagen, Germany). The volume of elution buffer AE was adjusted to 200 µL for 109 110 engorged females, 100 µL for unengorged adults and 40 µL for nymphs. A double elution 111 was carried out to increase gDNA yield for unengorged adults and nymphs. Sterile and DNA-112 free equipment and tubes were used for each step. Extraction reagents blank (EXB) controls 113 were performed alongside tick extractions to assess background bacterial communities.

114

115 2.3 16S amplification and library preparation for NGS sequencing

Amplicons targeting a 250-320 bp product of the V1-2 hypervariable region of the 16S rRNA 116 117 (16S) gene were generated using the primer pair 27F-Y (Gofton et al., 2015b) and 338R 118 (Turner et al., 1999). Previous research identified that paralysis tick, *I. holocyclus*, has a 119 highly abundant bacterial species ('Ca. M. mitochondrii'), which masks the diversity of bacteria in 16S metabarcoding studies (see Gofton et al. 2015a). Therefore, a blocking primer 120 121 developed by Gofton et al. (2015a) was used to inhibit the amplification of 'Ca. M. 122 mitochondrii' in I. holocyclus ticks. Depending on tick life stage and level of engorgement, 3-10 µM of the CMm blocking primer was added to the PCR reaction. Amplicon PCRs were 123 124 conducted in 25 µL reactions each containing 1 X Buffer (KAPA Biosystems, USA), 1.5 mM 125 MgCl₂, 0.4 µM of each forward and reverse primer with MiSeq adapters, 0.4 mg/mL BSA

126 (Fisher Biotech, Australia), 0.25 mM dNTPs (Fisher Biotech, Australia), 0.5 U KAPA Taq 127 (KAPA Biosystems, USA), and 2.0 µL of undiluted genomic DNA. Samples underwent the following thermal cycling conditions; initial denaturation at 95°C for 5 mins, followed by 35 128 cycles of denaturation at 95°C for 30 s, annealing at 62°C (CMm blocking primer present) or 129 55°C (CMm blocking primer absent) for 30 s, and extension at 72°C for 45 s with a final 130 131 extension at 72°C for 5 mins. Libraries were then prepared using the Nextera XT DNA 132 library preparation kit in 25 μ L reactions following manufacturer's recommendations. Libraries were purified using Agencourt[®] AMPure[®] XP PCR purification beads (Beckman 133 134 Coulter Life Sciences, USA) and pooled in equimolar amounts. Final libraries were then 135 quantified using the Qubit[®] 2.0 Fluorometer (Thermo Fisher, Australia). Libraries were 136 sequenced on an Illumina MiSeqTM using v2 chemistry (2x250 paired end). 137 Extraction reagents blanks (n=12) and PCR no-template controls (NTC; n=7) were included 138 in all stages of the workflow. All pre- and post-PCR procedures were performed in physically 139 separate dedicated laboratories and sterile protocols were maintained through library 140 141 preparation in order to minimise amplicon contamination. 142 143 2.4 Bioinformatics and statistical analysis 144 Raw fastq files were downloaded from the Illumina BaseSpace Sequence Hub for analysis in 145 a custom pipeline using USEARCH (Edgar, 2010). Raw paired-end sequences were merged

in USEARCH v10, a minimum of 50 nucleotide (nt) overlap and maximum number of

147 mismatches increased to 15 nt due to long overlap of paired-end sequences. Only sequences

148 with perfect primer sequences were retained, primer sequences and distal bases were trimmed

using USEARCH v8.0. Sequences were quality filtered in USEARCH v10, allowing a <1%

150 expected error rate and singletons were discarded (Edgar and Flyvbjerg, 2015). Sequences

151 were then clustered into operational taxonomic units (OTUs) of 97% similarity using the 152 UPARSE algorithm (Edgar, 2013) in USEARCH v10. Taxonomy was assigned in QIIME2 v2018.4 using the q2-feature-classifier (Bokulich et al., 2018) with reference to a trained 153 154 Greengenes database (DeSantis et al., 2006) (release May 2013) using the primer pair 27F-Y/ 155 338R. Taxonomic assignments were confirmed using NCBI MegaBLAST (Morgulis et al., 156 2008) on a random subsample of OTUs and in the case of tick-associated microbes, GenBank 157 accession numbers and percentage identity of top hits were recorded for 'taxa of interest'. The profiles from EXB controls and NTCs were first assessed to ensure quality of sampling 158 159 and absence of tick-associated bacteria. The following criteria were used to assess inclusion 160 of sequences: all OTUs that appeared exclusively in controls were removed; OTUs that had a 161 higher relative sequence abundance in controls compared to tick samples were removed; and OTUs were removed that appeared in over half of controls (i.e. at least eight) that had a 162 163 taxonomic identity associated with environmental bacteria (e.g. members of the phyla 164 Acidobacteria and Cyanobacteria). In addition, potential cross-contamination during library 165 preparation or 'cross-talk' at the sequencing level was assessed by inspecting the presence of 166 expected tick-associated bacteria (e.g. members of the obligate intracellular bacterial families 167 Anaplasmataceae and Midichloriaceae) in controls. The profiles of EXB controls and NTCs 168 were then removed bioinformatically from associated samples to eliminate background 169 bacteria.

170

Data analysis and visualisation was carried out in RStudio (RStudio Team, 2015) using
packages metacoder (Poisot et al., 2017), phyloseq (McMurdie and Holmes, 2013) and vegan
(Oksanen et al., 2019). Alpha diversity of samples was measured using observed OTUs,
Chao1 index, Shannon index and Simpson index. Removal of samples with a low sequencing
depth (<1000 assigned OTUs after data filtering), did not significantly alter alpha diversity

176 measurements among tick species (data not shown). Rarefaction curves for samples were calculated to assess sequencing depth based on observed number of OTUs. Principal 177 coordinate analysis (PCoA) on weighted unifrac dissimilarity measurements was used to 178 179 assess the differences in microbial composition between tick species. Investigation into 'taxa 180 of interest' warranted a more rigorous assessment of sequence number than was required for 181 diversity measures in order to avoid any potential cross-contamination and machine cross-182 talk (Dong et al., 2017; Wang et al., 2017), as such only samples with >100 sequences were 183 considered positive. Taxonomic assignment to 'taxa of interest' was assessed based on NCBI 184 MegaBLAST top hit of named organism. Where precent identity was $\leq 97\%$, the terminology 185 of nearest named Genus-like was employed. Due to sample collection biases in the present 186 study, prevalence data of these microbes were not considered statistically relevant. 187 Nucleotide sequences from 'taxa of interest' were aligned by MUSCLE (Edgar, 2004) using 188 default parameters and aligned sequences were then imported into MEGA7 (Kumar et al., 189 2016) with the most appropriate nucleotide substitution model chosen based on the lowest Bayesian Information Criterion (BIC) score. Evolutionary histories were inferred using the 190 191 Neighbour-Joining method based on the Tamura 3-parameter model (Tamura, 1992). 192 Bootstrap analysis was conducted using 10,000 replicates to assess the reliability of inferred 193 tree topologies. 194

3. Results

196

197 *3.1 Tick-host associations*

198 Ticks were collected from 221 wildlife hosts including 24 native and three introduced species
199 (Fig. 1, Supplementary File S1). Hosts recorded in the present study included; agile
200 antechinus (*Antechinus agilis*), brown antechinus (*Antechinus stuartii*), spotted-tail quoll

201 (Dasyurus maculatus), water rat (Hydromys chrysogaster), quenda (Isoodon fusciventer),

202 northern brown bandicoot (Isoodon macrourus), western grey kangaroo (Macropus

- 203 *fuliginosus*), eastern grey kangaroo (*Macropus giganteus*), red-necked wallaby
- 204 (Notamacropus rufogriseus), red kangaroo (Osphranter rufus), platypus (Ornithorhynchus
- 205 anatinus), eastern barred bandicoot (Perameles gunnii), long-nosed bandicoot (Perameles
- 206 nasuta), sugar glider (Petaurus breviceps), eastern ring-tailed possum (Pseudocheirus
- 207 peregrinus), black fruit-bat (Pteropus alecto), fruit-bat sp. (Pteropus sp.), bush rat (Rattus
- 208 fuscipes), black rat (Rattus rattus), Tasmanian devil (Sarcophilus harrisii), wild pig (Sus
- 209 scrofa), short-beaked echidna (Tachyglossus aculeatus), rufous-bellied pademelon
- 210 (*Thylogale billardierii*), short-eared brush-tailed possum (*Trichosurus caninus*), common
- 211 brush-tailed possum (Trichosurus vulpecula), wombat (Vombatus ursinus), red fox (Vulpes

212 *vulpes*), and swamp wallaby (*Wallabia bicolor*). Female ticks were the dominant life stage

- 213 recorded (n=547) followed by nymphs (n=319) and males (n=153). Ticks were received from
- animals in the Northern Territory (NT) (n=16), Queensland (QLD) (n=249), New South
- 215 Wales (NSW) (*n*=316), Tasmania (TAS) (*n*=136), Victoria (VIC) (*n*=44) and Western
- 216 Australia (WA) (n=258). Together, Bothriocroton concolor (n=123), Ixodes australiensis

217 (n=210), I. holocyclus (n=173) and Ixodes tasmani (n=184) accounted for over two-thirds of

all ticks submitted.

219

220 3.2 16S rRNA bacterial profiling

A sub-sample of 238 ticks and 19 controls underwent 16S NGS profiling. A total of 23.9
million raw paired-end sequences were generated on the Illumina MiSeq. 17.9 million
sequences were retained after merging, and subsequent quality filtering yielded 14.8 million
sequences for clustering and taxonomic assignment. A total of 4,864 OTUs (average length
299 bases) were retained after background profiles were removed. After removal of

226 background sequences, a total of 14,328,059 bacterial sequences were assigned to tick 227 samples. Despite a high number of OTUs, only 1,535 OTUs had greater than 100 total sequences from tick samples. Tick samples had an average of 60,201 assigned sequences (see 228 229 Supplementary File S2). Amblyomma triguttatum had the highest median alpha diversity as measured by the observed number of OTUs and the chao1 index, whereas Ixodes antechini 230 231 had the highest median alpha diversity as measured by the Shannon and Simpson indexes 232 (Fig. 2). Rarefaction analysis of the sequence depth shows that the observed number of OTUs plateaued at 50,000 sequences (Fig. 3). After the removal of sequences from controls, eight 233 234 phyla were retained. An ordination plot of OTUs (Fig. 4) show that bacteria belonging to the 235 Proteobacteria phylum were the most abundant and diverse taxa classified, followed by 236 Firmicutes and Actinobacteria. Bacterial families identified in tick species, represented as 237 relative number of sequences in Fig. 5, show 26 dominant taxa. While bacterial composition 238 varied between tick species, sequences from the members of Coxiellaceae, Francisellaceae, 239 and Rickettsiaceae families (Phylum: Proteobacteria) were the most abundant. Although sample sizes varied among tick species, beta diversity analysis incorporating abundance 240 241 (sequence number) and taxonomic relatedness, showed evidence of different bacterial communities among tick species (Supplementary File S3). 242

243

244 3.3 Presence of 'taxa of interest'

In total, 37 OTUs were identified as 'taxa of interest' in the present study. Their taxonomic
identity and abundance (number of sequences) in each sample is available in Supplementary
File S4 (see Supplementary File S5 for fasta file of sequences). Phylogenetic analysis of the
'taxa of interest' is represented in Fig. 6, showing tick species and Australian states and
territory, where each OTU was identified.

251 A proposed novel Anaplasma sp. (OTU230, MK814412, 96.3% identity) was identified in a

single *B. concolor* (1/20) collected from (ex) echidna from NSW. A second *Anaplasma*

253 bovis-like OTU (OTU312, JN862824, 97.3% identity) was identified in Haemaphysalis

254 *bancrofti* (3/10) ex red-necked wallaby and long-nosed bandicoot and *Haemaphysalis*

humerosa (n = 1/13) tick ex long-nosed bandicoot, all from NSW.

256

257 OTU5509 was assigned to the genus Bartonella and had a top BLAST hit of Bartonella

258 *bacilliformis* (LN624026, 92.1% identity). OTU5509 was identified in two samples,

259 however, was present in an extremely low number of sequences (2) in each case. Both

samples were *I. fecialis* collected from two different quenda in WA. The low number of

261 sequences means that in the case of statistical analysis, this OTU would have been filtered

out. While there was no sufficient match to this sequence, it is noted that in many instances

native Australian *Bartonella* species are lacking sequence information for this region of the
16S gene (V1-2).

265

Two OTUs (OTU14, CP025785, 100% identity; and OTU4629, CP025785, 100% identity)
were identified as '*Ca*. B. tachyglossi'. Sequences were identified in *Bothriocroton concolor*(8/20) ex echidnas from QLD & NSW and *H. humerosa* (1/13) ex northern brown bandicoot
from QLD.

270

A *Coxiella*-like organism (OTU12, CP032542, 95.5% identity) was identified exclusively in *B. concolor* (19/20) ex echidna from NSW and QLD. This was the second most abundant
OTU identified in *B. concolor*, accounting for ~20% of the overall sequences.
A *Coxiella* sp. (OTU15, KC170757, 100% identity) was identified in *Haemaphysalis*

275 *longicornis* (3/4) ex red fox from NSW and, OTU represented ~61.4% of the overall assigned

276 bacterial sequences.

277

278 A novel *Ehrlichia* sp. (OTU33, AY309970, 96.3% identity) was identified in *I. fecialis* (1/13)

279 ex quenda from WA. Other Ehrlichia OTUs (OTU16 and 1632) represented sequences from

280 'Ca. Ehrlichia ornithorhynchi' in Ixodes ornithorhynchi (6/10) ex platypus QLD and these

sequences accounted for the majority of sequences (~50.2%) from *I. ornithorhynchi*.

282

283 A Francisella endosymbiont (OTU5, AF001077, 98.0% identity) was identified in H.

284 humerosa (11/13) ticks ex northern brown bandicoots from NT and QLD, and

285 Haemaphysalis bremneri (2/4) ex possum (species unknown) in QLD. OTU5 was the most

abundance sequence in *H. bremneri* and *H. humerosa*, representing 27.3% and 52.4% of the

assigned sequences, respectively. A second *Francisella* endosymbiont (OTU10, AB001522,

288 99% identity) was identified in *H. bancrofti* (8/11) ex red-necked wallabies from QLD and

289 NSW, and ex a long-nosed bandicoot in NSW; and in a single *H. humerosa* (1/13) ex red-

290 necked wallaby from NSW. OTU10 was the most abundance sequences in *H. bancrofti*

representing 27.0% of the assigned sequences. OTU40 (AF001077, 97.6% identity) was also

identified as a *Francisella*-like endosymbiont from *A. triguttatum* (6/6) ex red kangaroo from

293 WA. It was highly abundant in these ticks, accounting for 76.9% of the overall sequences.

294

295 Midichloria (OTU6, FM992372, 100% identity) was identified in I. holocyclus (20/36) ex

long-nosed bandicoots from NSW and QLD, *I. fecialis* (1/13) ex red-necked wallaby NSW,

297 *H. bancrofti* (1/11) ex long-nosed bandicoot NSW, and *H. humerosa* (1/13) ex long-nosed

bandicoot NSW. Midichloria (OTU7, FM992373, 100% identity) was identified in I.

299 holocyclus (15/36) ex long-nosed bandicoots from NSW and QLD, I. fecialis (1/13) ex red-

300 necked wallaby NSW and *H. bancrofti* (1/11) ex long-nosed bandicoot NSW. Overall a total

of 15 ticks had both OTU6 and OTU 7, which included *I. holocyclus* (13/36) ex long-nosed

302 bandicoots from NSW and QLD, *I. fecialis* (1/13) ex red-necked wallaby NSW and *H.*

303 *bancrofti* (1/11) ex long-nosed bandicoot NSW. While the CMm blocking primer was

304 incorporated into the PCRs conducted on *I. holocyclus, Midichloria* sequence were still

305 observed in 2/9 females, 3/6 males and 16/21 nymphs. *Midichloria* was the most abundant

sequence from *I. holocyclus* accounting for 36.1% of assigned sequences.

307

308 Three Neoehrlichia OTUs were identified. 'Ca. Neoehrlichia arcana' (OTU25, KT203914,

309 99.3% identity) was identified in eight ticks, I. tasmani (4/51), I. holocyclus (2/36), H.

310 *bancrofti* (1/11) and *H. humerosa* (1/13), ex long-nosed bandicoots NSW and QLD. '*Ca.*

311 Neoehrlichia australis' (OTU29, KT803957, 100% identity) was identified in 10 ticks, *I*.

312 *holocyclus* (8/36) and *I. tasmani* (2/51), ex long-nosed bandicoots NSW and QLD. Three

ticks (*I. tasmani* and *I. holocyclus*) were co-infected with both '*Ca*. N arcana' and '*Ca*. N.

australis'. A novel Neoehrlichia species (OTU31, MG670107, 97.3% identity) was identified

in *I. fecialis* (2/13) and *I. australiensis* (2/26) ex quenda from WA, and *I. antechini* (2/29) ex

316 brown antechinus from NSW.

317

318 Due to the conserved nature of the *Rickettsia* genus at the 16S gene, reported previously 319 (Roux and Raoult, 1995; Stothard and Fuerst, 1995), resolution to species level was not 320 possible using the sequences obtained in the present study. Rickettsia sp. (OTU2, KF318168, 99.7% identity) had a widespread distribution among samples and was detected in 61 ticks 321 322 (B. concolor, H. bancrofti, H. humerosa, I. antechini, I. australiensis, I. holocyclus and I. 323 tasmani) from three states and one territory (NSW, NT, QLD, WA). Rickettsia was the most abundant sequence from *B. concolor* and *I. antechini* ticks accounting for 63.3% and 12.0% 324 325 of the assigned sequences, respectively

326

327 A Rickettsiella-like sp. (OTU1, LC388767, 92.7% identity) was identified in I. australiensis 328 (22/26) and was the most abundant sequence accounting for 76.8% of assigned sequences. A 329 Rickettsiella-like sp. (OTU3, EU430251, 95.1% identity) was identified in I. tasmani (11/51) 330 ex Tasmanian devils and eastern barred bandicoot from TAS. A Rickettsiella-like sp. (OTU4, 331 U97547, 98.4% identity) was identified in, I. tasmani (36/51) ex bandicoots, possum and a 332 sugar glider from NSW, TAS and QLD; I. australiensis (1/26) ex western grey kangaroo 333 from WA; and I. holocyclus (1/36) ex long nosed bandicoot NSW. OTU4 accounted for 334 30.2% of the overall sequences assigned for *I. tasmani*.

335

336 4. Discussion

337

Currently there is a scarcity of detailed information about the life cycles of Australian ticks. 338 339 however it is generally assumed that, as with other hard tick species around the world, the 340 majority will exhibit a three-host life cycle (Barker and Walker, 2014). Despite these limitations, the opportunistic sampling strategy used in this study provides an economical 341 342 way to survey a wide range of tick and wildlife fauna across a range of geographical areas 343 across Australia. The paucity of larvae from this data set is also a limitation of this type of 344 sampling. Tick larval stages, which may indicate the presence of transovarially-transmitted 345 organisms (Kwan et al., 2017), are difficult to see with the naked eye, and in situations where 346 collection of ticks is not the main objective, they are easily overlooked (Lydecker et al., 347 2019). Furthermore, the opportunistic nature of the sampling precluded assessment of the 348 infestation rates of the various hosts. In future, if TBPs of humans and the wildlife 349 themselves are identified, further extensive surveys of tick-pathogen-wildlife ecologies will 350 require more targeted and comprehensive approaches in order to gather sufficient relevant

351 epidemiological data.

352

As with previous studies (Swei and Kwan, 2016; Zolnik et al., 2016) alpha diversity 353 354 measures were highly variable both within and between tick species. This may be driven by a 355 variety of factors such as; starting material (i.e. size/life stage of tick), extraction methods, 356 library preparation, normalisation of DNA input concentrations for sequencing, batch 357 sequencing effects and bioinformatics (Greay et al. 2018). Within the present study, 358 sequencing depth was likely most impacted by sample input type and normalisation of DNA 359 concentrations, as all samples went through the same library preparation and bioinformatic 360 analysis. Overall, bacterial diversity of ticks started to plateau by 50,000 sequences however, 361 it is noted that in some cases this plateau was not achieved and deeper sequencing would be required in order to confidently characterise the full suite of bacterial taxa present. This was 362 363 most evident in A. triguttatum, B. auruginans, I. trichosuri and I. ornithorhynchi and 364 therefore we recommend a minimum of 100,000 sequences per sample for future bacterial 365 16S amplicon studies in these species to be confident that the complete bacterial community 366 has been sampled. Studies that investigate the shift of the core microbiome (i.e. most 367 abundant bacteria taxa) relative to a given parameter may not require the same depth of 368 sequencing, as seen in previous studies investigating the effect of temperature (Thapa et al., 369 2019) and life stage (Andreotti et al., 2011).

370

After bioinformatic analysis, including stringent quality filtering, a large number of OTUs remained with relatively low number of reads. This has been noted in previous studies of the tick microbiome (Budachetri et al., 2016; Zolnik et al., 2016) and in the broader field of highthroughput microbiome studies (Pollock et al., 2018). Depending on bioinformatic analysis and quality filtering, the number of OTUs and therefore diversity can vary greatly among

studies (Greay et al., 2018). In addition, current practice in microbiome studies is to
normalise count data, however models show that this can oversimplify the data (McMurdie
and Holmes, 2014).

379

380 Ordination analysis demonstrated that tick species was the strongest predictor of bacterial 381 composition. Beta-diversity analysis showed that the common marsupial tick, *I. tasmani*, 382 exhibited a variable bacterial composition. This diversity may be explained by the wide geographic distribution of this tick in Australia, its own high genetic diversity (Burnard and 383 384 Shao, 2019), and its ability to parasitise many marsupial species. In the present study, 385 bacterial profiling of *I. tasmani* included ticks from three states (NSW, QLD, TAS) and seven 386 host species. With this in mind, we suggest that future studies should combine careful 387 taxonomic status identity with data on microbial communities of *I. tasmani*. Although sample 388 sizes were limited in 'host-specialist' ticks B. auruginans (wombats), H. bremneri (possums), I. antechini (antechinus) and I. ornithorhynchi (platypus), these specimens showed less 389 390 diversity of bacterial communities between samples.

391

An important caveat on the relative diversity of bacterial communities in the present study 392 393 was the use of a blocking primer inhibiting the amplification of 'Ca. M. mitochondrii'. The inclusion of this blocking primer it vital to explore the full bacterial community of *I*. 394 holocyclus (see Gofton et al., 2015), however it does impact the analysis and interpretation of 395 396 the data. Due to the inhibition of this bacteria, there is an inherent bias in the bacterial 397 community, and comparisons in alpha and beta-diversity measures must account for this. In the case of highly abundant organisms, the use of a blocking primer assay will likely not 398 399 completely inhibit amplification, and thus a reduced level of the organisms may be still detected, as was the case in the present study. In this instance, the use of diversity measures 400

401 that rely on presence/absence data will not be affected by this granted there is still some level 402 of detection (i.e. number of species observed for alpha diversity and Jaccard index for betadiversity). The use of more advanced, and usually preferred, diversity indexes may be 403 404 impacted by the manipulation of bacterial composition, and care must be taken when 405 comparing results with other studies (Greay et al., 2018). Despite these limitations, the use of 406 blocking primers has shown to be vital in the context of investigating known tick-borne 407 pathogens and novel related taxa from tick samples. In the present study we have 408 characterised the bacterial communities of ten tick species for the first time. This information 409 provides the first step of focusing future research and where this established technique should 410 be applied.

411

Members of the Proteobacteria represented the most diverse and abundant (as determine by
number of sequences) in Australian hard ticks analysed in the present study. This finding is
consistent with previous microbiome studies from tick species, such as *Ixodes scapularis*(Sperling et al., 2017; Thapa et al., 2019), *Ixodes persulcatus* (Zhang et al., 2014; Kurilshikov
et al., 2015) and *Amblyomma americanum* (Fryxell and DeBruyn, 2016). Additionally the
identification of novel bacterial taxa in native Australian ticks species is consistent with
recent findings (Gofton et al., 2015b; Panetta et al., 2017).

419

Currently in Australia, the ecology, epidemiology, and incidence of human TBDs remains
largely a matter of conjecture, having received little scientific study compared with many
other parts of the world (Graves and Stenos, 2017). Despite significant national interest,
including a federal government senate inquiry (Radcliffe et al., 2016), the prevailing
scientific opinion concludes that Lyme borreliosis (caused by *B. burgdorferi* sensu lato), for
example, is absent from Australia (Chalada et al., 2016; Irwin et al., 2017). In the unique

426 Australian environment, long isolated in geological terms, it is likely that unidentified tick-427 host life cycles have evolved, given the endemic wildlife (including tick fauna) present 428 (Long, 2017; Beati and Klompen, 2019). It is possible that these cycles may contribute to 429 zoonotic illness when humans encroach these sylvatic ecologies and become exposed to native ticks. Indeed, it is well documented that in endemic areas in Europe and North 430 431 America, the causative agent of Lyme borreliosis is readily identified in wildlife (such as 432 white-tailed deer, Odocoileus virginianus, and white-footed mice, Peromyscus leucopus) as well as their natural tick species (Bosler et al., 1984). The present study therefore is one of 433 the first to explore the concept that Australian wildlife ticks should be a promising source to 434 435 identify exotic and novel TBPs, and we focussed our search towards taxa known to be 436 associated with TBDs overseas ('taxa of interest'), namely Anaplasma, Bartonella, Borrelia, 437 Coxiella, Ehrlichia, Francisella, Midichloria, Neoehrlichia, Rickettsia and Rickettsiella. 438

A new genetic variant of *Anaplasma bovis* was recently described from questing A. 439 440 triguttatum ticks in WA and the nearby Barrow Island (Gofton et al., 2017). In the present 441 study we report a range expansion of a genetically similar A. bovis along the east coast of 442 Australia from two widespread Haemaphysalis species. As suggested by Gofton et al., 443 (2017), this finding further supports the hypothesis that it is likely that endemic A. bovis 444 genotypes exist in sylvatic cycles within native Australian ticks and wildlife fauna. Further research on the phylogenetic position of these A. bovis sequences is needed to understand 445 446 their likely evolutionary history and relatedness to A. bovis genotype Y11 identified in 447 Western Australia. While three other species of Anaplasma (A. marginale, A. centrale and A. platys) have been introduced to Australia (Rogers and Shiels, 1979; Callow, 1984; Angus, 448 449 1996), the present study did not identify any of these species in ticks from wildlife. This is likely due to the absence of the cattle tick (R, (B)) australis) and the brown dog tick (R). 450

451 *sanguineus*) specimens from wildlife hosts in the present study. A species of *Anaplasma*

452 identified exclusively in *B. concolor* from echidnas sheds more light on the diverse range of

453 microbes that have been described from this specialist tick (Loh, 2018).

454

Although the presence of a *Bartonella* sp. was identified by only four sequences, its absence 455 456 in control samples means that this likely represents a true finding in female *I. fecialis*, which 457 were parasitising different individuals of the same host (quenda) in south-west WA. Recent 458 studies have shown that Australian marsupials and native rodents harbour a range of distinct 459 Bartonella species; 'Ca. Bartonella antechini' has been identified in ticks (I. antechini) and 460 fleas (Acanthopsylla jordani) from mardo (Antechinus flavipes) in south-west WA 461 (Kaewmongkol et al., 2011c), 'Ca. Bartonella woyliei' in woylie ticks (I. australiensis) and fleas (Pygiopsylla hilli) in south-west WA, and 'Ca. Bartonella bandicootii' in fleas 462 463 (Pygiopsylla tunneyi) from western barred bandicoots (Perameles bougainville) on Bernier 464 and Dorre Island (Kaewmongkol et al., 2011a). Reports of Bartonella spp. occurring outside 465 south-west WA include *Bartonella australis* ex eastern grey kangaroos (*Macropus giganteus*) 466 (Fournier et al., 2007). Molecular detection of Bartonella DNA from Australian ticks has also 467 been reported in ticks (I. tasmani) parasitising koalas (Phascolarctos cinereus) from Philip 468 Island, VIC (Vilcins et al., 2009b). Additional studies on *Bartonella* in Australia wildlife include those by Gundi et al. (2009), Kaewmongkol et al. (2011a) and Dybing et al. (2016). 469 Due to the limited size and region of the 16S amplicon generated in the present study, there 470 471 were insufficient relevant reference sequences available from other Australian Bartonella 472 species for comparison. A study by Kaewmongkol et al. (2011b) into flea-derived Bartonella from native and introduced Australian species suggests co-evolution of marsupial hosts, their 473 474 fleas and the Bartonella species.

476 'Ca. Borrelia tachyglossi' was identified in 8/20 B. concolor ticks from echidnas in QLD and NSW, an anticipated finding given previous research into this organism (Loh et al., 2016). 477 Importantly however, the present study provides the first evidence of 'Ca. B. tachyglossi' 478 479 sequences from a female *H. humerosa* tick parasitising a northern brown bandicoot from 480 QLD. In addition, no Bothriocroton ticks were present in the extraction batch of the positive 481 H. humerosa sample, and there was no evidence of Borrelia sequences from any controls. 482 Despite the wide geographical range of *H. humerosa* ticks, the restricted finding of '*Ca*. B. tachyglossi' from QLD supports Loh et al. (2016), suggesting a restricted geographical 483 484 distribution along the east coast of Australia. 485 486 In tick microbiome studies overseas, Coxiella spp. are commonly identified (Khoo et al.,

487 2016; Machado-Ferreira et al., 2016). Additionally, recent studies on Australian ticks have

488 also identified *Coxiella* spp. in *A. triguttatum* (Cooper et al., 2013; Gofton et al., 2015b), *B.*

489 auruginans (Vilcins et al., 2009c), I. holocyclus (Cooper et al., 2013) and R. sanguineus

490 (Oskam et al., 2017). Interestingly the present study did not identify the widespread presence

491 of *Coxiella* spp. with only two unique OTUs identified from *B. concolor* and *H. longicornis*,

492 both of which appear to be host specific to those tick species, and the causative agent of Q-

493 Fever (*C. burnetii*) was not identified from ticks in the present study.

494

Ehrlichia sequences (*Ca.* E. ornithorhynchi') from platypus ticks, *I. ornithorhynchi*, were
recently described (Gofton et al., 2018). In the present study, *Ca.* E. ornithorhynchi' was
exclusively observed within *I. ornithorhynchi*, suggesting this genetically distinct *Ehrlichia*species has a unique and host-specific relationship with the platypus. During this study, a
potentially novel species of *Ehrlichia* was identified in an *I. fecialis* female tick from a
quenda in WA. BLAST results show it is a close relative to an *Ehrlichia* sp. detected in

501 *Haemaphysalis* ticks from Japan (Inokuma et al., 2004). No sequences from the recently

502 described '*Ca*. E. occidentalis' (Gofton et al., 2017) were identified in the present study,

503 however it is noted *A. triguttatum* were only represented by six samples.

504

505 *Francisella*-like endosymbionts have been widely reported in ticks overseas, such as

506 Dermacentor spp. (Scoles, 2005), Dermacentor occidentalis (Gurfield et al., 2017),

507 Haemaphysalis longicornis (Wang et al., 2018) and Hyalomma rufipes (Szigeti et al., 2014),

and *Francisella* sequences have previously been identified in *A. fimbriatum* ticks from

509 reptiles in the Northern Territory of Australia (Vilcins et al., 2009a). In the present study, a

510 *Francisella*-like endosymbiont was identified in 100% of *A. triguttatum* ticks, and in a high

511 proportion. The high prevalence and relative abundance of the *Francisella*-like organisms in

512 *A. triguttatum* may be of medical relevance with respect to this common human-biting tick,

513 particularly given the recent report of Francisella bacteraemia in WA (Aravena-Román et al.,

514 2015). Importantly, as noted previously for *I. holocyclus* ticks, organisms in lower abundance

515 may be masked by abundant endosymbionts, unless samples are sequenced deeply and/or a

516 blocking primer is used (Gofton et al., 2015a). Therefore, future bacterial profiling studies of

517 *A. triguttatum* should take these factors into consideration.

518

519 While '*Ca*. M. mitochondrii' has been documented in *I. holocyclus*, the present study

520 provides the first report of '*Ca*. M. mitochondrii' within additional species of native

521 Australian ticks (I. fecialis, H. bancrofti and H. humerosa). Despite this expansion among

522 tick species, identification of 'Ca. M. mitochondrii' remains confined to NSW and QLD as

523 previously reported (Gofton et al., 2015a). 'Ca. M. mitochondrii' has been detected overseas

in laboratory models (Cafiso et al., 2019), wildlife (Serra et al., 2018), and humans

525 (Mariconti et al., 2012). Despite the use of a CMm blocking primer by Gofton et al. (2015a)

in the library preparation of *I. holocyclus*, the bacteria was still identified in the majority of
ticks (21/36); however its abundance was greatly reduced to an average of 26.1% of reads per
sample in comparison to previous research demonstrating a relative abundance of 98.2%
(Gofton et al., 2015a).

530

531 A recent addition to the Anaplasmataceae family, 'Ca. Neoehrlichia mikurensis' was first 532 isolated from wild rats and their ticks (*Ixodes ovatus*) in Japan (Kawahara et al., 2004). In Europe, the hedgehog (Erinaceus spp.), a common peri-urban dweller, has been shown to 533 play an important role in the life cycle of 'Ca. N. mikurensis' (Földvári et al., 2014; Jahfari et 534 535 al., 2017). A high proportion of bandicoot ticks were infected with Neoehrlichia spp. in this 536 study, suggesting that these marsupials may play an important role in the life cycle of this 537 bacterium on the Australian continent. Bandicoots also frequently inhabit gardens and have 538 relatively close contact with humans (Carthey and Banks, 2012). Closer study of these marsupials may provide vital information about the life cycle of these microbes and the 539 540 potential risk for human infection.

541

Rickettsia spp. are among the only currently recognised TBPs affecting people in Australia. 542 543 however, as previously outlined, the conserved nature of the 16S gene in *Rickettsia* precludes rigorous species delimitation from the relatively short sequences generated in this dataset. In 544 addition to the known human pathogens (R. australis and R. honei) (Graves and Stenos, 545 546 2017), the presence of *Rickettsia* has been demonstrated from a variety of Australian tick 547 species. Rickettsia gravesii has been described from A. triguttatum (Li et al., 2010; Abdad et al., 2017) and Rickettsia sequences have been previously identified in I. tasmani ex 548 549 Tasmanian devils (Vilcins et al., 2009c) and Amblyomma fimbriatum from reptiles (Vilcins et 550 al., 2009a).

551

552 The diversity and significance of the *Rickettsiella* genus remains largely unknown globally. 553 Phylogenetic analysis of sequence data places the genus within the Coxiellaceae family (Fournier and Raoult, 2005; Leclergue and Kleespies, 2012). While the presence of these 554 555 organisms has been well documented in tick species around the world and in some cases, they 556 have been identified as pathogenic to arthropods (Kurtti et al., 2002; Leclerque et al., 2012), disease causation within the vertebrate hosts remains unknown at the present time. A study 557 558 by Vilcins et al. (2009c) identified Rickettsiella in I. tasmani ex koalas from Phillip Island, 559 just off the coast of south eastern Australia. A genetically similar sequence was identified in 560 the present study (OTU3) from *I. tasmani* exclusively in Tasmania. 561 562 While this is the first study to characterise the bacterial communities within native Australian 563 wildlife ticks, it is apparent that much research is still required in order to better understand the tick-associated microbial life cycles and their ecologies, let alone elucidating their 564 565 potential for transmission and pathogenicity in vertebrate hosts, including humans. 566 Furthermore, whilst cost-effective, ticks were collected opportunistically resulting in an 567 inherent geographical bias, confining sampling largely to urban areas along the east coast of the Australian continent. Nevertheless, this area is also where most humans receive tick bites, 568 569 although reliable data about this is also lacking. In addition, the present study suggests that 570 'taxa of interest' are largely restricted to a combination of geographical location and tick 571 species. For example, 'Ca. Midichloria' was identified in H. humerosa and I. fecialis from

572 NSW only and was absent in samples from NT for *H. humerosa* and WA and TAS for *I*.

573 fecialis.

574

575 The analysis of ticks removed from wildlife hosts comes with the inherent complication of a

576 blood meal. The host blood meal has been recently shown to influence both the tick microbiome composition and the presence of pathogens (Landesman et al., 2019; Swei and 577 Kwan, 2017). Further targeted studies are needed to assess the source of microbes with 578 579 respect to a tick or host origin (Irwin et al., 2018). In addition, limitations in the current 580 sequencing technologies mean that high-throughput methods favour short amplicons which 581 may not be able to accurately discriminate bacterial species, as is the case with Rickettsia in 582 particular (Gihring et al., 2012). Furthermore, important caveats to consider in metabarcoding microbial diversity analysis include PCR efficiency (including primers used and length of 583 target amplicon), variations in 16S gene copy number (Ahn et al., 2012), sequencing depth, 584 585 machine cross-talk, bioinformatic analysis and taxonomic assignment. While there have been 586 some recent attempts to correct these biases (Kembel et al., 2012; Rosselli et al., 2016), their 587 application to uncharacterised metagenomic samples remains limited. While 16S 588 metabarcoding continues to be vital in bacterial biodiversity discovery (Carpi et al., 2011; 589 Tessler et al., 2017), future molecular studies incorporating metagenomics and 590 metatranscriptomics, to detect actively expressed genes (Cabezas-Cruz et al., 2018; Greay et 591 al., 2018), will be useful in characterising the full suite of micro-organisms present in 592 Australian ticks. A multi-disciplinary approach incorporating cell culture, in vitro tick studies 593 and morphological techniques (e.g. fluorescence microscopy) will also be needed to assess 594 the potential for transmissibility and pathogenicity of these novel tick-associated organisms. 595

596 5. Conclusions

597

598 With over 75% of emerging human infectious pathogens originating from wildlife (King,
599 2014), surveillance methods that target these species are important for investigations of
600 emerging and exotic infectious disease. Research into the interactions of wildlife hosts, ticks

601 and pathogens in Europe and North America continues to highlight the complexity of these 602 dynamic systems (Ostfeld et al., 2018; Tomassone et al., 2018). Results from the present 603 study build on recent research into Australian tick-associated microbes, further highlighting 604 the diversity of organisms present that appear related, yet distinct from their overseas 605 counterparts. With the evolutionary history of Australia's unique tick species (Beati and 606 Klompen, 2019) and wildlife fauna, it is likely that reports of their tick-associated microbes 607 (potential pathogens) will continue to reveal taxonomic differences from those described in 608 the northern hemisphere. As such, future research on emerging tick-borne zoonoses should 609 include methods that are able to detect novel micro-organisms in humans and reservoirs and 610 include a variety of sample types such as blood, tissue (e.g. skin, spleen etc.) and tick from 611 both the host and environment (questing). As anthropogenic changes to the environment 612 continue to grow in Australia, a greater emphasis on wildlife disease surveillance is critical to 613 ensure the early detection of potential infectious diseases affecting humans, livestock, companion animals and wildlife (Woods et al., 2019). Advances in pathogen detection and 614 615 characterisation are greatly enhanced by collaboration; the authors advocate for continued 616 multidisciplinary efforts between health professionals, researchers, land managers and local 617 communities.

618

619 Supporting Information

620

621 Data availability

Associated metadata output from bioinformatic analysis and subsequent data visualisation is
available on FigShare repository https://doi.org/10.6084/m9.figshare.c.4608803.v1. Nextgeneration sequencing data can be accessed from NCBI Sequence Read Archive under
BioProject: PRJNA559059 (BioSample accession numbers: SAMN12512474 –

626 SAMN12512711).

627

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639

640 Compliance with ethical standards

- 641 This study was conducted under the compliance of the Australian Code for the Responsibility
- 642 Conduct of Research (2007) and Australian Code for the Care and Use of Animals for
- 643 Scientific Purposes, 2013. Tick collection was carried out opportunistically with the approval
- 644 from the Murdoch University Animal Ethics Committee.
- 645
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648 Legends to Figures

- **Fig. 1.** Chord diagram showing tick-wildlife associations recorded in the present study. The wildlife hosts recorded are represented on the lower half of the plot, and tick species across the top. Thickness of bar relative to number of ticks. Host records that could not be assigned to species level are not represented in this plot. A complete list of records is available in
- 653 Supplementary File S1. Introduced wildlife species are denotes with an asterisk. Silhouette
- 654 wildlife images sourced from phylopic.org.
- 655 Fig. 2. Alpha diversity of bacterial communities in ticks parasitising Australian wildlife
- 656 measured as; observed number of operational taxonomic units (OTUs), Chao1 index,
- 657 Shannon index and Simpson index. Tick species abbreviated to; A. triguttatum (A. tri), B.
- 658 aurugians (B. aur), B. concolor (B. con), H. bancrofti (H. ban), H. bremneri (H. bremneri),
- 659 H. humerosa (H. hum), H. longicornis (H. lon), I. antechini (I. ant), I. australiensis (I. aus), I.
- 660 fecialis (I. fec), I. holocyclus (I. hol), I. myrmecobii (I. myr), I. ornithorhynchi (I. orn), I.
- 661 tasmani (I. tas), and I. trichosuri (I. tri).
- 662 Fig 3. Rarefaction plot of 16S rRNA bacterial sequences clustered into 97% operational
- taxonomic units (OTUs) from ticks parasitising Australian wildlife.
- 664 Fig 4. Non-metric multidimensional scaling (NMDS) plot of operational taxonomic units
- 665 (OTUs) based on Bray-Curtis dissimilarity matrix. Taxa filtered to only display OTUs that
- 666 were present at least twice in >10% of tick samples.
- 667 Fig 5. Stacked barplot of bacterial composition (shown at family level) from Australian ticks
- 668 represented as percentage of assigned sequences. Taxa at the family level that represented
- 669 < 10% of the relative sequences within each tick species were grouped as "Low abundant".
- 670 Sequences not able to be accurately assigned to family taxa are displayed as "Unclassified".
- 671 Fig 6. Neighbour-joining phylogenetic tree displaying taxa of interest prevalence and

672 distribution among tick samples. Geographic data relating to tick collection is represented by

- 673 state and territory; New South Wales (NSW), Northern Territory (NT), Queensland (QLD),
- 674 Tasmania (TAS) and Western Australia (WA). Evolutionary histories were inferred based on
- 675 the Tamura 3-parameter model with bootstrap analysis (10,000 replicates) (bootstrap values
- 676 > 60 are displayed). Tick samples were considered positive for taxa of interest if >100
- 677 sequences present. Information on number of sequences, taxonomic identity (as inferred from
- 678 NCBI MegaBLAST analysis with nucleotide database), and sequences can be found in
- 679 supplementary information (Supplementary File S4 & S5).
- 680 Supplementary Files

681 Supplementary File S1. Sample metadata of tick-wildlife records identified from the present682 study.

683 Supplementary File S2. Sequence information from 16S rRNA bacterial profiling of ticks

on the Illumina MiSeq platform after bioinformatic analysis; (a) number sequences assigned

to 97% operational taxonomic units (OTUs) and (b) number of sequences per sample.

686 Supplementary File S3. Principal coordinate analysis (PCoA) plot of the weighted UniFrac

687 distance matrix. Plots show beta diversity of tick bacterial communities, displayed by tick

688 species and life stage. Operational taxonomic units (OTUs) that represented <100 sequences

689 in a tick sample were removed.

690 Supplementary File S4. Number of sequences relating to the taxa of interest in tick samples.

691 The top hit from NCBI BLAST results showing GenBank accession number, percent identity

and E-value are given, and final evaluation of taxonomic assignment.

693 Supplementary File S5. Fasta sequences for operational taxonomic units (OTUs) used in the694 taxa of interest investigation.













Proteobacteria

Anaplasmataceae Bradyrhizobiaceae Burkholderiaceae Caulobacteraceae Comamonadaceae Coxiellaceae Enterobacteriaceae Francisellaceae Midichloriaceae Oxalobacteraceae Pseudomonadaceae Rickettsiaceae Sphingomonadaceae

Spirochaetes

Spirochaetaceae

Unclassified

Low abundant (<10%)

