Description of the female, nymph and larva and mitochondrial genome, and redescription of the male, of *Ixodes barkeri* Barker, 2019 (Acari: Ixodidae) from the short-beaked echidna, *Tachyglossus aculeatus*, with a consideration of the most suitable subgenus for this tick

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Abstract

Background

Ixodes barkeri, a tick with a distinctive ventrolateral horn-like projection on palpal segment 1, was described in 2019 from two male ticks from the Wet Tropics of Far North Queensland, Australia. However, females lie at the core of the taxonomy and subgenus classification of Ixodes, hence we sought specimens of female ticks, successfully recovering females, plus nymphs and larvae. Mitochondrial genomes are also desirable additions to the descriptions of species of ticks particularly with regard to subgenus systematics. So, we sequenced the mt genomes of I. barkeri Barker, 2019 and the possible relatives of I. barkeri that were available to us (I. australiensis Neumann, 1904, I. fecialis Warburton & Nuttall, 1909, and I. woyliei Ash et al. 2017) with a view to discovering which if any of the subgenera of Ixodes would be most suitable for I. barkeri Barker, 2019.

Results

The female, nymph, larva, and mitochondrial genome of Ixodes barkeri Barker, 2019 are described for the first time and the male of I. barkeri is redescribed in greater detail than previously. So far, I. barkeri is known only from a monotreme, the short-beaked echidna, Tachyglossus aculeatus (Shaw, 1792), from the highland-rainforests of Far North Queensland, Australia.

Conclusions

Our phylogeny from entire mitochondrial genomes indicated that I. barkeri, and indeed I. woyliei Ash et al. 2017, another tick that was described recently, are best placed in the subgenus Endopalpiger Schulze, 1935.

Background

Ixodes barkeri Barker, 2019 was described from two male ticks collected from the short-beaked echidna Tachyglossus aculeatus (Shaw, 1792). One of these ticks was from rainforest of the Peeramon Scrub, Atherton Tableland, Far North Queensland (FNQ) whereas the other male was from an unknown locality in the vicinity of the Atherton Tableland. Ixodes barkeri is distinctive among Australian ticks, especially for its ventrolateral palpal projection which is found only in an echidna tick from Papua New Guinea tick, I. zaglossi Kohls, 1960. Ixodes zaglossi, however, has syncoxae whereas I. barkeri does not and the ventrolateral palpal projection in I. barkeri is much bigger [1]. However, this extraordinary species, I. barkeri, could not be placed within a higher taxonomic framework, such as the subgenus classification of Ixodes, largely due to the lack of females. Since [1], we have acquired other specimens of I. barkeri: 6 males, 5 females, 34 nymphs and 2 larvae (Table 1, Fig. 1), allowing us to describe the female, nymph, larvae for the first time, to redescribe the male in greater detail to compare and contrast the morphology of I. barkeri with other species of Australasian Ixodes (sensu [2]). We have also described the mitochondrial genome of I. barkeri, enabling inferences on phylogenetic relationships of this species with others in the genus.

The subgenera of Ixodes are morphologically ambiguous and in need of further refinement and testing with genetic data. In this regard, mitochondrial (mt) genomes have been remarkably instructive about the evolutionary
history (phylogeny) of ticks (e.g. [3, 4, 5, 6, 7]). Thus, we sequenced the mt genomes of *I. barkeri* and its possible relatives that were available to us (*I. australiensis* Neumann, 1904, *I. fecialis* Warburton & Nuttall, 1909, and *I. woyliei* Ash *et al.* 2017) with a view to discovering which if any of the subgenera of *Ixodes* would be most suitable for *I. barkeri* Barker, 2019. [8] and [1] were not able to place *I. woyliei* and *I. barkeri* in a subgenus, respectively.

## Methods

### Material examined

Only field-collected ticks were available for study. The specimens were from the Barker and Barker Collection at the University of Queensland (Qld), the Queensland Museum (QM), the Australian National Insect Collection (ANIC), and the Field Museum of Natural History, Chicago, Illinois, USA (FM) (Table 1).

### Microscopy methods

Ticks were studied using a stereoscopic microscope (Nikon SMZ800N, Nikon Corporation, Tokyo, Japan and Olympus SZX16, Olympus Corporation, Tokyo, Japan), compound microscope (Olympus BX53, Olympus Corporation, Tokyo, Japan) and a scanning electron microscope (JEOL JSM6610LV, JEOL Ltd., Tokyo, Japan). An ocular micrometre was used to measure ticks. Measurements are in millimetres for the adults, micrometres for the juveniles, and are given as the range followed by the mean and the number of specimens measured (n) in parentheses. Colour digital images were taken with a Canon 6D camera (Canon Corporation, Tokyo, Japan).

Adobe Photoshop® software was used to correct images for broken legs and other damaged parts of the tick, and to polish the image.

### Sequencing and assembly of mitochondrial genomes

Mitochondrial genomes were sequenced and assembled in two ways. First, the mt genome of *I. barkeri* Barker, 2019 was sequenced at Novogene Singapore and then assembled at the University of Queensland (UQ) by the protocol of [5]. DNA was prepared by us at the University of the Sunshine Coast and the University of Qld. Groups of ticks were cut in half and then incubated at 56°C for 62 hours with Proteinase K to lyse the cells. The QIAGEN DNeasy Blood and Tissue kit was used to extract genomic DNA. The amount of DNA recovered was measured with Nanodrop and Qbit instruments. Groups of ticks that yielded more than 200 nanograms (ng) of DNA were sent to Novogene Singapore for *de novo* library construction and next-generation Illumina sequencing. Groups of ticks with less than 200 ng were combined with DNA from a different organism, usually the host, to reach the minimum threshold of 200 ng of DNA required by Novogene Singapore. At Novogene Singapore, DNA was sonicated to fragment the DNA, then fragments were end-polished, A-tailed and ligated with Illumina adaptors. DNA fragments were amplified with PCR, using P5 and P7 oligos, to create genomic libraries which were purified with AMPure XP system. The Illumina Novaseq 6000 sequencing platform was used to generate two giga-bases of nucleotide sequence data (PE 150). *De novo* contig assemblies of Illumina sequences were then constructed by us in Geneious Prime [9] by the default assembler of Geneious Prime. Blast-searches of contigs revealed mt genes of ticks; these gene sequences were then assembled until entire mt genomes had been assembled.
Second, the mt genomes of *I. australiensis*, *I. fecialis* and *I. woylie* were sequenced at the Hokkaido University, Japan, and then assembled at the University of Queensland by the protocol of [5]. DNA was extracted from ticks with the NucleoSpin® DNA Insect (Macherey-Nagel, Germany). Entire mt genomes were then amplified in two overlapping fragments (long-range and short PCRs). Long-range PCR was used to amplify fragments that comprised about 12-13 kb of the mitogenome with the universal primers: mtG_K23 (5′-TCCTACATGATCTGATGTYAGACCG-3′) and mtG_K25 (5′-AAAATTCTAGGTTCTTTGTCC-3′) or mtG_K26 (5′-ACGGCGATATGTRCATATTTTAGAC-3′). Short PCRs were then used to amplify 1.5-2.5 kb of mt genomes with genus-specific primers. PrimeSTAR® GXL DNA Polymerase (Takara-Bio, Shiga, Japan) was used to amplify the long mt PCR products whereas Tks Gflex™ DNA Polymerase (Takara-Bio) was used to amplify short mt gene fragments as well as nuclear rRNA genes. PCR conditions for PrimeSTAR® GXL DNA Polymerase were: 45 cycles of 98 °C for 10 s, 60 °C for 15 s, and 68 °C for 10 min. PCR conditions for Tks Gflex™ DNA Polymerase were: 94 °C for 60 s, 45 cycles of 98 °C for 10 s, 55 °C for 15 s, 68 °C for 60s, and a final extension of 68 °C for 5 min. PCR products were examined on 1.2% agarose gels stained with Gel-Red™ (Biotium, Hayward, CA). PCR products were purified with a NucleoSpin Gel and PCR Clean Up Kit (Takara-Bio). Illumina sequencing libraries were constructed from the PCR fragments from the long-range and short PCR reactions with the Nextera DNA Library Prep Kit (Illumina, Hayward, CA) and were sequenced with the Illumina MiSeq platform with the MiSeq reagent kit v3 for 600 cycles.

**Annotation of mitochondrial genomes**

Mitochondrial genomes were annotated with Geneious Prime. Protein-coding genes were identified by searches with BLAST [10] for open reading-frames. Regions between protein-coding genes were searched with BLAST [10] to find rRNA genes, tRNA genes and control regions. The tRNA that we expected to find but were not found with BLAST were found with the aid of the tRNAscan-SE Search Server v1.21 [11] and the MITOS Web Server [12]. The nucleotide sequences of tRNA genes were confirmed by studying the putative secondary structure of transcripts, as implemented in Geneious Prime [9].

**Phylogenetic methods**

Phylogenies were inferred by both Maximum Likelihood (ML) and Bayesian Inference (BI) methods implemented in the RAXML-HPC2 v 8.2.12 [13] and MrBayes v3.2.2 [14], respectively. JmodelTest2 v2.1.6 [15] was used to find the optimal substitution model for the nucleotide dataset. The GTR + G + I model was found to be the best fit for our dataset. In all ML and BI runs (experiments), genes were partitioned. Rapid-bootstrapping of 1,000 replicates of our data was executed in RAXML-HPC2 v 8.2.12 [13]. There were two simultaneous BI runs: 10 million generations sampled every 1000 MCMC steps. For every BI run, four MCMC chains (three heated and one cold) were executed. The first 25% of steps were discarded as burn-in. Tracer v 1.5 [16] was used to observe the effective sample size (ESS) and convergence of independent runs. Phylogenetic trees were displayed in FigTree v 1.4.4 [17]. Branch support in the phylogenetic trees generated by RAXML-HPC2 v 8.2.12 [13] and MrBayes v 3.2.2 [14] was assessed by the bootstrap values and posterior probability values, respectively. All phylogenies were inferred through the CIPRES Science Gateway v.3.3 [18]. *Ixodes pavlovskyi* Pomerantzev, 1946, a species from the lineage of the “other *Ixodes* (sensu [2]) was the out-group.

**Results**
Systematics

Family Ixodidae Murray, 1877

Genus Ixodes Latreille, 1795

Subgenus Endopalpiger Schulze, 1935

Ixodes barkeri Barker, 2019

**Male.** [Based on 6 specimens, 3 measured - refer to Table 1; Figs. 2, 3, 8] Idiosoma (Fig. 2) broadly elongate-oval with broadly rounded posterior margin, widest slightly posterior to mid-length; length from apices of scapulae to posterior body margin 2.17–2.59 mm (2.42 mm), width 1.53–1.72 (1.61), ratio 1.42–1.56 (1.49). Lateral groove distinct. Consuctum (Figs. 2, 8) length 2.12–2.56 (2.38), width 1.37–1.48 (1.44), ratio 1.54–1.73 (1.65); laterally and postero-laterally with a distinct narrow non-sclerotised band; scapulae short, blunt; lateral carinae absent; cervical grooves indistinct; dense moderately large punctations evenly distributed over conscutum, except for pseudoscutum area; pseudoscutum with indistinct punctations; setae moderately dense, very short (c. 0.01) and indistinct (Fig. 2). Venter plate outlines as illustrated (Fig. 2); median plate: length 1.06–1.18 (1.10), width 0.74–0.84 (0.81), ratio 1.26–1.43 (1.37); adanal plate: length 0.74–0.82 (0.79), width 0.42–0.48 (0.45), ratio 1.67–1.95 (1.77); anal plate: 0.54–0.60 (0.58), width 0.32–0.36 (0.34), ratio 1.67–1.71 (1.69). All ventral plates with dense, moderately large punctations (Figs. 1, 2). Genital aperture (Fig. 2C) located at level of posterior margin of coxae II; posterior margin of genital apron deeply concave. Ventral setae (Fig. 2) moderately dense, very short, evenly distributed on all plates; length of setae on median plate c. 0.01. Anal groove (Fig. 2C) straight anteriorly and open posteriorly. Spiracular plate (Fig. 2A) broadly oval, longer than wide, length 0.34–0.42 (0.38), width 0.28–0.34 (0.31), ratio 1.21–1.25 (1.23).

Gnathosoma (Fig. 3, 8) length from palpal apices to cornual apices dorsally 0.45–0.48 (0.46), width between lateral projection of palpal segments I 0.43–0.48 (0.46), ratio 0.98–1.04 (1.01). Dorsal basis capituli (Fig. 3A) length from medial insertion of palpal segment I to cornual apices 0.21–0.25 (0.23), width 0.32–0.35 (0.33), ratio width to length 1.41–1.52 (1.46), subrectangular, posterior margin nearly straight; cornua long, ratio total length of basis capituli, including cornua, to cornual length 4.83–5.78 (5.39), triangular with narrowly rounded apex. Ventral basis capituli (Fig. 3C) subrectangular; lateral margins with slight constrictions at mid-length; auriculae absent; short converging ridges in auricular areas. Palpi (Fig. 3B) short, length dorsally (segments II and III) 0.23–0.24 (0.23), maximum width (in dorsolateral plane) 0.18–0.19 (0.19), ratio 1.22–1.29 (1.26), length of palpal segment I ventrally 0.16 (n = 1), maximum width ventrally 0.17 (n = 1), ratio 0.94 (n = 1); segment I greatly enlarged, greatest dimension in anteromedian-posterolateral direction; dorsally segment I subrectangular; ventrally segment I subtriangular, posterior margin with very long and narrow spur with sharply pointed apex; segments II and III fused together with indistinct suture between them, narrower proximally and abruptly widening to broadly rounded apex. Hypostome (Fig. 3C) length 0.18–0.19 (0.18), width 0.08–0.10 (0.09), ratio 1.75–2.10 (1.95); club-shaped, widening to broadly rounded apex with medial indentation; base of hypostome at level of base of palpal segment I; dental formula 3/3, basal half of hypostome without denticles, denticles sharply pointed.
Legs moderately long, slender. Coxae (Fig. 2E, 9): coxae I-IV with long and narrow external spur with narrowly rounded to sharply pointed apex; spur on coxae I-III subequal, spur on coxa IV nearly twice shorter those on coxae I-III; coxae I-IV without syncoxae. Trochanter I with long, triangular spur with sharply pointed apex; trochanters I-IV with long, narrow, with sharply pointed apex spur ventrally. Tarsus I: length 0.56–0.60 (0.58); tarsus IV length 0.53–0.58 (0.55); tarsi only slightly humped subapically.

**Female.** [Based on 1 to 5 specimens - refer to Table 1; Figs. 4, 5, 9] Idiosoma (Fig. 4A, 9) length from scapular apices to posterior body margin in moderately engorged specimen 4.1 (n = 1), width in moderately engorged 2.5 (n = 1), ratio 0.16 (n = 1), broadly suboval, widest approximately at mid-length. Scutum (Fig. 4, 9) length 1.05–1.18 (1.13; n = 4), width 1.40–1.58 (1.49; n = 4), ratio 0.73–0.79 (0.75; n = 4); lateral margins diverging for approximately 2/3 of scutum length, broadly rounded posteriorly; lateral carinae lacking; cervical grooves shallow; dense, small punctations evenly distributed throughout scutum; setae (Fig. 4) relatively sparse, very short (c. 0.005), indistinct and nearly equal to those on alloscutum, distributed as figured. Alloscutum (Fig. 4) as illustrated; setae of alloscutum (Fig. 4C) numerous, evenly distributed, very short, length of setae in central field c. 0.01, indistinct. Venter (Fig. 4E) as illustrated; genital aperture (Fig. 4E) medial to coxae III; genital groove (Fig. 4E) well developed; anal groove (Fig. 4E) oval with open posterior margin; ventral setae numerous, length of preanal setae c. 0.01, evenly distributed. Spiracular plates (Fig. 5A) length 0.27–0.32 (0.30; n = 4), width 0.39–0.47 (0.44; n = 4), ratio 0.68–0.70 (0.69; n = 4); broadly oval; marginal row of perforations in groove anteriorly.

Gnathosoma (Fig. 5B) length from palpal apices to cornual apices dorsally 0.60–0.70 (0.65; n = 4), width between lateral projection of palpal segments I 0.66–0.74 (0.71; n = 4), ratio 0.89–0.95 (0.92; n = 4). Dorsal basis capituli (Fig. 5B) length from medial insertion of palpal segment I to cornual apices 0.22–0.25 (0.24; n = 4), width 0.51–0.55 (0.54; n = 4), ratio width to length 2.20–2.32 (2.26; n = 4), subrectangular, posterior margin nearly straight; cornua short, ratio of total length of basis capituli (including cornua) to cornual length 17.50–20.00 (19.12; n = 4), triangular with broadly rounded apex; subcircular porose areas poorly distinct, not indented, separated by distance nearly equal to their own width, several openings arranged in discrete punctations. Ventral basis capituli (Fig. 5C) subrectangular; lateral margins with slight constrictions at mid-length; auriculae absent; short converging ridges in auricular areas. Palpi (Fig. 5B) short, length dorsally (segments II and III) 0.41–0.50 (0.46; n = 4), maximum width (in dorsolateral plane) 0.21–0.25 (0.23; n = 4), ratio 1.85–2.10 (1.95; n = 4), length of palpal segment I ventrally 0.31–0.36 (0.34; n = 4), maximum width ventrally 0.19–0.20 (0.20; n = 4), ratio 1.64–1.75 (1.68; n = 4); segment I greatly enlarged, greatest dimension in anteromedian-posterolateral direction; dorsally segment I subrectangular with convex medial margin; ventrally segment I subtriangular, posterior margin with long moderately narrow spur with narrowly rounded apex; segments II and III fused together with indistinct suture between them, narrower proximally and abruptly widening to broadly rounded apex. Hypostome (Fig. 5C) length 0.36–0.40 (0.38; n = 4), width 0.12–0.16 (0.14; n = 4), ratio 2.46–2.85 (2.68; n = 4); club-shaped, widening to broadly rounded apex with medial indentation; base of hypostome approximately at level of base of palpal segment II; dental formula 4/4 (few rows may be 3/3), basal half of hypostome without denticles, denticles sharply pointed.

Legs moderately long, slender. Coxae (Figs. 5E, 9): coxae I-IV with moderately long and narrow external spur with narrowly rounded apex; spur on coxae I-III subequal, spur on coxa IV nearly twice shorter those on coxae I-III; coxae I-IV without syncoxae. Trochanter I with moderately long, triangular spur with sharply pointed apex; trochanters I-IV without spur ventrally. Tarsus I: length 0.66–0.76 (0.73; n = 4); tarsus IV length 0.57–0.66 (0.63; n = 4); tarsi only slightly humped subapically.
Nymph. [Based on 34 specimens - refer to Table 6] Scutum (Fig. 6A) length 515–520 (518; n = 2), width 690–700 (695; n = 2), ratio 0.74–0.75 (0.75; n = 2); lateral margins diverging for approximately half of scutum length, broadly rounded posteriorly; posterolateral margin with slight indentations; lateral carinae lacking; cervical grooves shallow; moderately dense, small punctations evenly distributed throughout scutum; setae (Fig. 6A) relatively sparse, very short: length in central field of scutum 10 (n = 2), indistinct and nearly equal to those on alloscutum, distributed as figured. Setae of alloscutum numerous, evenly distributed, very short, length of setae in central field 10–13 (11; n = 2), indistinct. Anal groove oval with open posterior margin; ventral setae numerous, evenly distributed. Spiracular plates (Fig. 6B) broadly oval; marginal row of perforations in groove anteriorly.

Gnathosoma (Fig. 6C) length from palpal apices to cornual apices dorsally 285 (n = 2), width between lateral projection of palpal segments I 375–380 (378; n = 2), ratio 0.75–0.76 (0.76; n = 2). Dorsal basis capituli (Fig. 6C) length from medial insertion of palpal segment I to cornual apices 108 (n = 2), width 238–240 (239; n = 2), ratio width to length 2.21–2.23 (2.22; n = 2), subrectangular; posterior margin nearly straight; cornua moderately long, triangular with narrowly rounded apex. Ventral basis capituli (Fig. 6D) subrectangular; lateral margins with slight constrictions at mid-length; auriculae absent; short converging ridges in auricular areas. Palpi (Fig. 6C) short, length dorsally (segments II and III) 190–193 (191; n = 2), width 73–85 (79; n = 2), ratio 2.24–2.66 (2.45; n = 2), length of palpal segment I ventrally 140–150 (145; n = 2), maximum width ventrally 125–130 (128; n = 2), ratio 1.12–1.15 (1.14; n = 2); segment I greatly enlarged, greatest dimension in anteromedian-posterolateral direction; dorsally segment I subrectangular with convex medial margin; ventrally segment I subtriangular, posterior margin with long moderately broad spur with narrowly rounded apex; segments II and III fused together with indistinct suture between them, narrower proximally and abruptly widening to broadly rounded apex. Hypostome (Fig. 6D) length 150 (n = 1), width 73 (n = 1), ratio 2.07 (n = 1); club-shaped, widening to broadly rounded apex with medial indentation; base of hypostome approximately at level of base of palpal segment II; dental formula 3/3 (few basal rows 2/2), basal half of hypostome without denticles, denticles sharply pointed.

Legs moderately long, slender. Coxae (Fig. 6F): coxae IV with external spur; spur on coxae I–III moderately long, nearly subequal; spur on coxa I with broadly rounded apex, spur on coxae II and III with narrowly rounded to sharply pointed apex; spur on coxa IV very short; coxae I–IV with syncoxae occupying approximately 1/3, 1/4, 1/5 and 1/6 respectively of coxal width. Trochanters I–IV without spur ventrally. Tarsus I: length 320–335 (328; n = 2); tarsus IV length 298–310 (304; n = 2); tarsi only slightly humped subapically.

Larva. [Based on 2 specimens - refer to Table 1; Figs. 7, 10] Scutum (Fig. 7A) length 238 (n = 1), width 360 (n = 1), ratio 0.66 (n = 1); lateral margins diverging for approximately half of scutum length, broadly rounded posteriorly; posterolateral margin with slight indentations; lateral carinae lacking; cervical grooves shallow; setae 3 pairs, length of Sc1 11 (n = 1); length of Sc4 14 (n = 1). Dorsal setae of alloscutum undetermined in number since we only had larvae that were engorged to examine: it was impossible to confidently count and associate setae of the idiosoma dorsally and ventrally. Length of Cd1 17 (n = 1), length of Md1 23 (n = 1). Ventral setae undetermined number; 1 pair on anal valves; 3 pairs of sternals, length of St1 19 (n = 1); 2 pairs of preanals, length of Pa1 22 (n = 1), length of Pa2 32 (n = 1).

Gnathosoma (Fig. 7B, 7C, 7D) length from palpal apices to cornual apices dorsally 123 (n = 1), width between lateral projection of palpal segments I 155 (n = 1), ratio 0.79 (n = 1). Dorsal basis capituli (Fig. 7B) width 135 (n = 1), subrectangular, posterior margin nearly straight; cornua practically indistinct. Ventral basis capituli (Fig. 7C) subrectangular; lateral margins with slight constrictions at mid-length; auriculae absent. Post-hypostomal setae
pair, length of Ph$_2$ 4 (n = 1); distance between Ph$_1$ 37 (n = 1), distance between Ph$_2$ 27 (n = 1). Palpi (Fig. 7B) short, length dorsally (segments II and III) 77 (n = 1), maximum width (in dorsolateral plane) 42 (n = 1), ratio 1.83 (n = 1), maximum width of palpal segment I ventrally 50 (n = 1); segment I greatly enlarged, greatest dimension in anteromedian-posterolateral direction; dorsally segment I subrectangular with convex medial margin; ventrally segment I subtriangular, posterior margin with long moderately broad spur with narrowly rounded apex; segments II and III fused together with indistinct suture between them, narrower proximally and abruptly widening to broadly rounded apex; segment I lacking setae, segments II and III combined with 9 dorsal and 3 ventral setae. Hypostome (Fig. 7C) length 83 (n = 1), width 34 (n = 1), ratio 2.44 (n = 1); club-shaped, widening to broadly rounded apex with medial indentation; base of hypostome approximately at level of mid-length of palpal segment I; dental formula 2/2, approximately 5 denticles in files; basal half of hypostome without denticles, denticles sharply pointed.

Legs moderately long, slender. Coxae (Fig. 7E): coxae I-III with external spur; spur on coxa I moderately long, on coxa II and III short to very short; spur on coxae I-III with broadly rounded apex; coxae I-III with syncoxae occupying approximately 1/3, 1/4 and 1/5 respectively posteromedian portion of coxal width. Trochanters I-IV without spur ventrally. Tarsus I: length 183 (n = 1); tarsus III length 166 (n = 1); tarsi only slightly humped subapically.

**Remarks**

By having a greatly enlarged palpal segment I that extends inwardly and anteriorly, all active life-stages of *I. barkeri* most closely resembles those of the *Endopalpiger* species of Australasia: *I. acer*, *I. australiensis*, *I. giluwensis*, *I. hydromyidis*, *I. luxuriosus*, *I. mirzai*, *I. planiscutatus*, *I. steini*, *I. stellae*, *I. tasmani*, *I. victoriensis*, *I. woyliei* and *I. zaglossi* (refer to [8, 19, 20]).

The males of *I. australiensis*, *I. tasmani*, *I. victoriensis* and *I. zaglossi* have been described [19, 21, 22, 23, 24]. The male of *I. barkeri* is easily distinguished from the males of the *Endopalpiger* species by the absence of the syncoxal areas on all coxae (vs. well-developed syncoxae on coxae I–IV in all those species).

The female of *I. barkeri* resembles only that of *I. woyliei* by the absence of syncoxal areas on coxae (vs. females of all other *Endopalpiger* with well developed syncoxae). The female of *I. barkeri* can be differentiated from *I. woyliei* by the scutum and basis capituli dorsally and ventrally without lateral carinae and other longitudinal ridges (vs. lateral carinae and longitudinal ridges present in *I. woyliei*), the considerably smaller palpal segment I with a long spur on its posterior margin (vs. greatly enlarged palpal segment I with shorter spur on its posterior margin in *I. woyliei*), 4/4 dental formula on hypostome (vs. 6/6 in *I. woyliei*) and the long spur on trochanter I dorsally (vs. indistinct spur in *I. woyliei*).

The nymph of *I. australiensis*, *I. hydromyidis*, *I. luxuriosus* ([25] wrote that the nymph of *I. luxuriosus* had not been described, although there is a brief description of it in [26]), *I. steini*, *I. tasmani*, *I. victoriensis* and *I. woyliei* have been described [8, 19, 22, 24, 26]. Unfortunately, all of these published descriptions and illustrations are too brief for confident comparison. None the less, we note that the nymph of *I. barkeri* has a scutum without lateral carinae (vs. distinct carinae in *I. victoriensis*), a scutum and basis capituli dorsally and ventrally without distinct longitudinal ridges (vs. with distinct, sharp ridges in *I. woyliei*), a distinct cornua (vs no cornua in *I. australiensis*, *I. hydromyidis*, *I. tasmani* and *I. woyliei*), mostly 3/3 dental formula on the hypostome (vs. 2/2 dental formula in
I. hydromyidis and I. tasmani, 4/4 in I. australiensis), external spurs on coxae I–IV (vs. apparently no spurs on coxae in I. hydromyidis, I. luxuriosus, I. steini and I. tasmani) and tarsi I–IV slightly humped subapically, without a notch (vs. strongly humped tarsi with distinct notch in I. victoriensis).

The larva of I. hydromyidis, I. tasmani and I. victoriensis have been described [22, 24, 27, 28]. Unfortunately, as with the nymphs, all of these published descriptions and illustrations of larvae are too brief for confident comparison. None the less, we note that the larva of I. barkeri has indistinct cornua on the basis capituli dorsally (vs. distinct cornua in I. victoriensis) and has external spurs on coxae I–III (vs. no spurs on coxae in I. hydromyidis and I. tasmani).

Our diagnoses may be broadened and improved once the nymphs and larvae of the other Australasian species of Endopalpiger are redescribed and illustrated accurately.

Mitochondrial Genomes And Phylogeny

Four entire mt genomes are presented here for the first time: I. australiensis, I. barkeri, I. fecialis and I. woyliei (Fig. 9). These mt genomes have the gene-arrangement that is typical of the Australasian Ixodes ([29], Fig. 1) except that in I. fecialis the main cluster of tRNA genes has the arrangement ARN\(\rightarrow\)ESF rather than ARN\(\rightarrow\)SEF: ARN\(\rightarrow\)SEF has been found in all other Ixodidae, Argasidae, Nuttalliellidae and Holothyrida studied so far [3, 4, 5, 6, 7 and Fig. 1 of 29]. Moreover, in I. fecialis we found a 60 bp insertion between tRNA-Asn (N) and tRNA-Glu (E) and a 57 bp insertion between tRNA-Glu (E) and tRNA-Ser (S); neither of these insertions are similar to any other motifs in the mitochondrial genome of I. fecialis.

The phylogeny from these mt genomes indicates that I. barkeri and I. woyliei are best placed in the subgenus Endopalpiger Schulze, 1935 since I. barkeri and I. woyliei were in a lineage with I. (Endopalpiger) australiensis and I. (Endopalpiger) tasmani, to the exclusion of species from the subgenera Ceratixodes, Exopalpiger and Sternalixodes: I. (Ceratixodes) uriae, I. (Exopalpiger) fecialis and I. (Sternalixodes) holocyclus (Fig. 10).

Discussion

The subgeneric classification of Ixodes is complex and sometime ignored, probably because some subgenera are defined ambiguously, making species difficult to place. Yet the names of the subgenera are valid, represent hypotheses of relationships, and deserve closer attention. Previously, neither [8] nor [1] attempted to place I. woylie and I. barkeri, respectively, in a subgenus. We, however, conclude that I. woylie and I. barkeri are best placed in the subgenus Endopalpiger Schulze, 1935 (Fig. 10). Alas, mt genomes from the other species of Endopalpiger were not available to us: (i) I. victoriensis Nuttall, 1916 and I. hydromyidis Swan, 1931 from Australia; and (ii) I. acer Apanaskevich, 2020; I. giluwensis Apanaskevich 2020; I. luxuriosus Schulze, 1935; I. mirzai Apanaskevich, 2020; I. planiscutatus Apanaskevich, 2020; I. steini Schulze, 1935; I. stellae Apanaskevich, 2020; and I. zaglossi Kohls, 1960 from Papua New Guinea.

Paul Schulze was a prolific German taxonomist whose life works were reviewed recently [28]. He described 17 entities that are presently considered as subgenera [30], including Endopalpiger in 1935, with Ixodes luxuriosus Schulze, 1935 as the type species (redescribed by [20]). The subgenus Endopalpiger was based mainly on their prominent and distinctive palps. Later, Schulze [31] gave generic status to Endopalpiger, thus emphasizing the very unusual form of the palps. [32] and [19] considered the subgenus Endopalpiger to be valid, but [33] and [34]
presented the subgenus *Endopalpiger* as a synonym of *Exopalpiger* Schulze, 1935 but without evidence or argument. Here, our phylogenetic trees show that *Endopalpiger* and *Exopalpiger* are not closely related. Rather, *Exopalpiger* is much closer to *Sternalixodes* and *Ceratixodes* than it is to *Endopalpiger* (Fig. 10).

The four species of *Endopalpiger* in our tree formed a monophyletic group (*barkeri, tasmani, woylie, australiensis*); indeed, a monophyletic group with 100% bootstrap support and a posterior probability of 1.0, the highest possible posterior probability (Fig. 10). This is the first phylogenetic tree from entire mt genomes (about 15,000 bps) or any similarly large number of nucleotides. [The only other tree that had more than one species of *Endopalpiger* was by [8] (Fig. 10; ca. 800 bps of *cox1*). Therefore, we found strong support for *Endopalpiger*, albeit with a limited set of taxa. The unique nature of palpal segment (article) I is a morphological synapomorphy of *Endopalpiger*. As described by [19] (p. 13), the female palpal segment I ["I" in Fig. 5C] is greatly enlarged and projects inwardly and forwardly so that it *ensheathes* each side of the base of the mouthparts; and ventrally palpal segment I is strongly salient ("ss" in Fig. 5C). The only similar palp-morphology in adults is that of *Exopalpiger* which, in the words of [19] (p. 13), sounds more like that of *Endopalpiger* than it actually is. According to [19] (p. 13), the female palpal segment 1 of *Exopalpiger* is also greatly enlarged, being attached at right angles to the transverse axis of the basis, but does not project inwardly or forwardly and it does not *ensheathe* any part of the base of the mouthparts; ventrally palpal segment I is salient but not as salient as in *Endopalpiger*.

**Abbreviations**

ANIC: Australian National Insect Collection; mt: mitochondrial.

**Declarations**

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**Availability of data and materials**

The data supporting the conclusions of this article are in the article. The mitochondrial genomes published for the first time in this paper have been submitted to GenBank database; accession numbers KX520004–KX520007.

**Authors’ contributions**
Conception and design of the study that led to the paper, and/or the paper: DB, ODS, MK, SCB, DAA.

Acquisition of research data where the acquisition required significant intellectual judgement, planning, design and/or input: DB, SK, RS, RN, SCB, DAA.

Analysis and interpretation of the data on which the research output is based: DB, ODS, SCB, DAA.

Drafting and/or critical review and re-drafting of the paper so as to contribute to the interpretation: DB, ODS, MK, SCB, DAA.

All authors read and approved the final manuscript.

**Ethics approval and consent to participate**

Not applicable.

**Consent for publication**

Not applicable.

**Competing interests**

The authors declare that they have no competing interests.

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Tables

Due to technical limitations, Table 1 is only available as a download in the Supplemental Files section.

Figures

![Map of Australia with localities marked](image)

**Figure 1**

See image above for figure legend.
Fig. 2. *irodes borkeri* Barker, 2019, scanning electron micrographs of idiosoma of male. A, Dorsal view; B, Dorsolateral view; C, Ventral view. Scale-bars: 0.5 mm.

**Figure 2**

See image above for figure legend.
Fig. 3. *Ixodes barkeri* Barker, 2019, scanning electron micrographs of male. A, Spiracular plate (arrows show orientation of spiracular plate: a, anterior; d, dorsal); B, Gnathosoma, dorsal view; C, Gnathosoma, ventral view; D, Gnathosoma, anterodorsal view; E, Coxae; F, Trochanter I, dorsal view. Scale-bars: A–D, F, 0.1 mm; E, 0.2 mm.

Figure 3

See image above for figure legend.
**Figure 4**

See image above for figure legend.
Fig. 5. *Ixodes barkeri* Barker, 2019, scanning electron micrographs of female. A, Spiracular plate (arrows show orientation of spiracular plate: a, anterior; d, dorsal); B, Gnathosoma, dorsal view; C, Gnathosoma, ventral view (I, palpal article 1; II, palpal article 2; ss the strongly salient part of palpal article 1); D, Gnathosoma, anteroventral view; E, Coxae; F, Trochanter I, dorsal view. Scale-bars: A, F, 0.1 mm; B–E, 0.2 mm.

**Figure 5**

See image above for figure legend.
**Figure 6**

See image above for figure legend.
Figure 7

See image above for figure legend.
**Fig. 8. Ixodes barkeri**
Barker, 2019, light microscopy image of female (Barker & Barker Collection reference # B5321), male (# B4994), nymph (# B5321) and larva (# B5321). Horizontal broken scale-bars: 1 mm; vertical scale-bars also in mm.

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**Figure 8**

See image above for figure legend.
Fig. 9. The mitochondrial genomes of *Ixodes* (*Endopelagifer*) *australiensis*, *I. (Endo.) barkeri*, *I. (Endo.) woylei* and *I. (Exopalpigfer) fecialis* and. Protein-coding genes are shown in green, tRNAs are in yellow, rRNAs are in red whereas the two control regions are in blue. Protein-coding genes are labelled by their four-character abbreviations, tRNAs are labelled by their one-letter amino acid abbreviations whereas the two control regions are labelled as CR1 and CR2. Mitochondrial genome size variation is indicated in parenthesis. The arrangement of genes in these four species is identical except that the main cluster of tRNA genes has the arrangement ARNSEE in the three species of *Endopelagifer* [I. (*Endo.*) *australiensis*, *I. (*Endo.*) barkeri* and *I. (*Endo.*) woylei*] whereas in the one species of *Exopalpigfer* [I. (*Exo.*) *fecialis*] the arrangement is ARNESP. The arrangement in *I. (Exo.) fecialis* is the first known arrangement in an *Ixodidae* tick that is different to ARNSEE. Thus, ARNESP might be a synapomorphy for the subgenus *Exopalpigfer*.

![Diagram of mitochondrial genome](image)

**Figure 9**

See image above for figure legend.

**Figure 10.** Maximum likelihood (ML) phylogenetic tree from entire mt genomes (14,935 bps). The sequence-alignment was put through Gblocks to remove regions with alignment gaps. Tip-labels indicate NCBI accession numbers and (Barker & Barker Collection reference #). Numbers above branches show Maximum Likelihood bootstrap support whereas numbers below branches show the Bayesian Posterior Probability support. *Ixodes pavlovskyi* Pomerantzev, 1946, one of the species “Other *Ixodes*” (entu Barker & Munzell, 2004), for which an entire mitochondrial (mt) genome was available in Genbank, was set as the outgroup. The scale bar indicates 0.06 nucleotide substitutions per nucleotide site for the 14,935 nucleotide sites in our alignment of theses entire mt genomes. So, for example, there were 896 nucleotide substitutions along the branch that leads to *I. (Ceratixodes) uriae* plus *I. (Sternalixodes) holocyclus* plus *I. (Exopalpigfer) fecialis*, which is marked with an asterisk (i.e. 0.06 nucleotide substitutions per nucleotide site x14,935 nucleotide sites (bps) = 896 nucleotide substitutions). Ticks in bold were sequenced in the present study.

![Phylogenetic tree](image)
See image above for figure legend.

**Supplementary Files**

This is a list of supplementary files associated with this preprint. Click to download.

- Table1.png
- BarkerI.barkerigraphicalabstractcopy.pdf