Phylogenetic analysis of the Australasian paralysis ticks and their relatives (Ixodidae: Ixodes: Sternalixodes)

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Abstract

Background: The Australasian paralysis ticks and their relatives, Ixodes Latrielle, subgenus Sternalixodes Schulze, are some of the most important ticks in the region. However, very little is known about their phylogenetic relationships. The aim of this study was to elucidate the evolutionary relationships of members of the subgenus Sternalixodes by undertaking phylogenetic analyses of morphological and molecular datasets.

Methods: Adult females (n = 64) of Sternalixodes, including Ixodes anatis Chilton, 1904, Ixodes confusus Roberts, 1960, Ixodes cornuatus Roberts, 1960, Ixodes cordifer Neumann, 1908, Ixodes dendrolagi Wilson, 1967, Ixodes hirsti Hassall, 1931, Ixodes holocyclus Neumann, 1899, Ixodes myrmecobii Roberts, 1962 and Ixodes trichosuri Roberts, 1960, were examined morphologically. Subsequently, these Ixodes spp. were genetically characterised using cytochrome c oxidase subunit 1 (cox1) gene and the internal transcribed spacer 2 (ITS-2) of the rRNA. Both morphological and molecular datasets were analysed using various phylogenetic methods to assess the evolutionary relationship of various members of the subgenus Sternalixodes.

Results: Phylogenetic analyses of the cox1 sequences and morphological characters datasets revealed that the Australian and Papuan Sternalixodes formed a distinct clade with the New Zealand member of the group I. anatis positioned basally, in a separate clade. Ixodes holocyclus, I. cornuatus and I. myrmecobii formed a distinctive clade in both the cox1 and morphological phylogenies. However, based on phylogenetic analysis of the ITS-2 data, I. holocyclus formed a separate clade whereas I. cornuatus and I. myrmecobii grouped in a different clade.

Conclusions: The cox1 and morphological data suggest that the subgenus Sternalixodes is paraphyletic, and I. anatis is not a sternalixodid tick; hence, it should not be included in the subgenus. Based on the phylogenetic analyses of cox1 and ITS-2 sequences, it appears that I. myrmecobii and I. cornuatus are not subspecies of I. holocyclus. Although this study provided better insights into the taxonomic status of the subgenus Sternalixodes, a complete morphological and molecular (using multiple markers) phylogenetic analysis including all members of the subgenus would be required to more accurately elucidate the evolutionary relationships within the subgenus.

Keywords: Ixodes, Sternalixodes, Phylogeny, Molecular, Morphological, Tick

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Background
Ticks (Arachnida: Ixodidae) are important ectoparasites of humans and animals and can cause direct (e.g., paralysis, anaemia) as well as indirect (e.g., transmission of pathogens) effects on their hosts [1–3]. Ticks can be found on all continents and are known to feed on all types of terrestrial vertebrates, including mammals, birds, reptiles and amphibians [1, 3]. The life-cycle of ixodid ticks consists of four developmental stages, the egg, and three active parasitic stages, larva, nymph, and adult (male and female). Depending on the type (hard, Ixodidae or soft, Argasidae) and species of ticks, their life-cycle can vary significantly.

To date, 70 species (56 and 14 members of families Ixodidae and Argasidae, respectively) of ticks have been recorded from a variety of hosts (humans and domestic animals = 16; mammals, reptiles and birds = 54) from Australia [4]. Among different genera of hard ticks prevalent in Australia, *Ixodes* is arguably the most important and its members are known to transmit and/or harbour pathogens, including *Rickettsia australis* (the causative agent of Queensland tick typhus) in humans [5], and flaviviruses, bunyaviruses and *Cercopithifilaria johnstoni* (Nematoda: Filarioidea) in wildlife [6–8]. Some *Ixodes* species such as *I. holocyclus* and *I. cornuatus* can also cause paralysis in humans, domestic animals, and wildlife [9]. *Ixodes holocyclus* is known to have an immunoeffectory action on humans, causing tick bite anaphylaxis [10].

Members of the genus *Ixodes* have not been intensively examined (e.g., phylogeny using combined morphological and molecular datasets, biology, life-cycle) within Australia. Few life-cycles have been elucidated and the bionomics of very few species are understood. Of the subgenera of *Ixodes*, *Sternalixodes* has received most attention. The subgenus comprises nine members, including *Ixodes anatis*, *I. confusus*, *I. cornuatus*, *I. cordifer*, *I. dendrolagi*, *I. hirsti*, *I. holocyclus*, *I. myrmecobii* and *I. trichosuri* [11]. However, the majority of studies have focussed on *I. holocyclus* and *I. cornuatus*, a species morphologically similar to *I. holocyclus*, aiming to determine their distribution [1, 12], morphological and molecular identification [1, 13] and phylogenetic relationships [14]. A number of questions therefore remain to be answered regarding the members of the subgenus *Sternalixodes*. For instance, the status of *I. myrmecobii* as subspecies of *I. holocyclus* as proposed by Roberts [1] needs to be tested. To date, Australian paralysis ticks and their relatives have not been analysed using morphological as well as molecular phylogenetics. Being an important subgenus, *Sternalixodes* requires systematic investigations to address a number of taxonomic questions regarding the validity of its members. Therefore, this study was designed to elucidate the evolutionary relationships of members of the subgenus *Sternalixodes* by undertaking phylogenetic analyses of morphological and molecular datasets.

Methods
 Tick collection and morphological identification
Female ticks (*n* = 74) used in this study were either available from The University of Melbourne (Ian Beveridge and Abdul Jabbar) or museums in Australia (South Australian Museum, Western Australian Museum, and Australian National Insect Collection), New Zealand (A. Heath, AgResearch, New Zealand), Papua New Guinea (Ifor L. Owen, National Veterinary Laboratory, Papua New Guinea) and South America (A. Guglielmone, Instituto Nacional de Tecnologia Agropecuaria, Argentina) (Fig. 1; Table 1). Following collection, each tick specimen was stored in 70% ethanol until used. Developmental stages of all species of *Sternalixodes* could not be examined as many are not yet described.

For morphological identification, each tick was examined using a dissecting microscope (Olympus, Japan). In

![Fig. 1 Collection sites for *Ixodes* species from Argentina, Australia, New Zealand and Uruguay, used in this study. Information linked to each unique number on the map is provided in Table 1](image-url)
addition, electron micrographs were taken using a Hitachi TM3030 Tabletop Scanning Electron Microscope, Germany. All Australian and Papuan tick species were identified using keys by Roberts [1]; whereas *I. anatis* specimens were identified following Hardwick [15] and *I. auritulus* Neumann, 1904 specimens were identified by A. Heath and A. Guglielmone.

One or two legs were removed from each specimen using flame sterilized forceps and stored in 70% ethanol for molecular work.

**DNA extraction, PCR amplification and DNA sequencing**

Prior to DNA extraction, ethanol was removed and leg(s) of individual ticks were washed three times (30 min) in distilled H$_2$O, and then ground using a plastic mortar. DNA was extracted using a DNeasy Blood and Tissue Kit (Qiagen, Hilden, Germany) following the protocol provided by the manufacturer.

Two loci (one mitochondrial and one nuclear ribosomal DNA) were PCR-amplified separately from each individual genomic DNA sample. The first locus (partial *cox*1 gene, ~850 bp) was amplified using the primers HCO2064 (5′-GGT GGG CTC ATA CAA TAA ATC C-3′) and HCOX1215 (5′-GCC ATT TTA CCG CGA TGA-3′); the second locus (partial second internal transcribed spacer, ITS-2; ~760 bp) was amplified employing primers ITS865 (5′-GTC TCC TGA TCG TGA GGT CG-3′) and ITS105 (5′-GGT CGA ATT GCC CCT CGT CC-3′) [14]. All PCRs were performed in a final volume of 50 μl, containing 10 mM Tris–HCl (pH 8.4), 50 mM KCl, 3.5 mM of MgCl$_2$, 200 μM of deoxynucleotide triphosphate, 100 pmol of each primer and 1 U of GoTaq polymerase (Promega, Madison, WI, USA) under the following cycling conditions: 94 °C for 5 min (initial denaturation); 35 cycles of 94 °C for 30 s (extension), 48 °C (cox1) or 50 °C (ITS-2) for 30 s (annealing) and 72 °C for 50 s (extension), followed by final extension.
Table 2  List of morphological characters (character numbers, name of character, character states)

| No. | Feature or structure | Character states |
|-----|----------------------|------------------|
|     |                      | 0    | 1      | 2      | 3      |
| 1   | Hypostome - 1        | lanceolate   | spatulate   |        |        |
| 2   | Hypostome - 2        | blunt       | bluntly-pointed | acutely pointed |        |
| 3   | Hypostome - 3        | Not bilobed | bilobed     |        |        |
| 4   | Dentition - 1        | 3/3 apically | 4/4 apically | 5/5 apically |        |
| 5   | Dentition - 2        | only 3/3 mid-hypostome | 4/4 and 3/3 mid-hypostome |        |        |
| 6   | Dentition - 3        | 2/2 basally | 3/3 basally (1) |        |        |
| 7   | Palpal article 1-1   | does not ensheath mouthparts | ensheathes basal portion of mouthparts |        |        |
| 8   | Palpal article 1-2   | no internal horn-like projection | internal horn-like projection |        |        |
| 9   | Palpal article 1-3   | rounded dorsally | rectangular dorsally | sub-rectangular dorsally | triangular dorsally |
| 10  | Palpal articles 2 and 3-1 | separate | faint suture present | between articles | fused |
| 11  | Palpal articles 2 and 3-2 | short and broad | long and slender |        |        |
| 12  | Palpal article 2 and 3-3 | distal spur absent | distal spur present |        |        |
| 13  | Auriculae            | absent      | present     |        |        |
| 14  | Cornua               | absent      | present     |        |        |
| 15  | Ventral posterior lobe on basis capituli | absent | present |        |        |
| 16  | Porose areas         | separated by equal to or less than half their width | separated by more than half their width |        |        |
| 17  | Median depression between porose areas | present | absent |        |        |
| 18  | Dorsal lateral carina(e) on basis capituli - 2 | absent | not extending to base of hypostome | extending to base of hypostome |        |
| 19  | Dorsal carinae on basis capituli | median carina present | median carina absent |        |        |
| 20  | Ventral lateral carinae on basis capituli - 2 | absent | not extending to base of hypostome | extending to base of hypostome |        |
| 21  | Ventral carinae on basis capituli - 3 | no carinae | two carinae | three carinae |        |
| 22  | Scutum - 1           | longer than wide | wider than long | as long as wide |        |
| 23  | Scutum - 2           | lateral carinae absent | lateral carinae present |        |        |
| 24  | Scutum - 3           | cervical grooves extending less than halfway down scutum | cervical grooves extending halfway or more down scutum |        |        |
| 25  | Scutum - 4           | emarginations absent | emarginations present |        |        |
| 26  | Scapulae             | absent      | present     |        |        |
| 27  | Sternal plate - (0), (1) | absent | present |        |        |
| 28  | Genital aperture     | level with third intercoxal space | level with mid-fourth intercoxal space |        |        |
| 29  | Coxae I              | external spur present | external spur absent |        |        |
| 30  | Coxae II             | external spur present | external spur absent |        |        |
| 31  | Coxae III            | external spur present | external spur absent |        |        |
| 32  | Coxae IV             | external spur present | external spur absent |        |        |
| 33  | Syncoxae             | Present     | Absent     |        |        |
| 34  | Ridges/rugosities - 1 | absent on coxa I | present on coxa I |        |        |
Table 2 List of morphological characters (character numbers, name of character, character states) (Continued)

| Character number | Character | Character state | Character state |
|-----------------|-----------|-----------------|-----------------|
| 35.             | Ridges/rugosities - 2 | absent on coxa II | present on coxa II |
| 36.             | Ridges/rugosities - 3 | absent on coxa III | present on coxa III |
| 37.             | Ridges/rugosities - 4 | absent on coxa IV | present on coxa IV |
| 38.             | Anal groove | does not meet posteriorly | meets posteriorly |

at 72 °C for 5 min. For each set of PCRs, negative (no-DNA) and positive (I. holocyclus DNA) controls were included. No amplification was detected in any of the negative control reactions at any time during the study. Amplicons (5 µl) were examined on 1.5% agarose gels stained with ethidium bromide. Gels were examined using transillumination and were photographed using a GelDoc system (BioRad, Hercules, CA, USA). If amplicons were not detected on agarose gel, then semi-nested PCRs were used as follows: HCOX1240 (5′-CCA CAA ATC ATA AAG ACA TTG G-3′) was used in conjunction with HCO2064 to amplify cox1 and ITS130 (5′-AGT TGT ACA TTG G-3′) in conjunction with ITS865 was used to amplify ITS-2. PCR cycling conditions for semi-nested PCRs were same as used above.

For each locus, amplicon(s) representing each *Ixodes* species were purified using shrimp alkaline phosphatase and exonuclease I [16] prior to automated DNA sequencing (ABI3730XL automatic sequencer at Macrogen Cooperation, South Korea). Sequencing of the *I. cornuatus* and *I. myrmecobii* (ABI3730XL automatic sequencer at Macrogen Cooperation, South Korea). Sequencing of the ITS-2 region was conducted using the primers HCO2064 and HCOX1215 or HCO2064 and HCOX1240 (cox1) and ITS865 and ITS105 or ITS865 and ITS130 (ITS-2), in separate reactions. The quality of each sequence obtained was appraised using the program Geneious Pro 6.5 (Biomatters Ltd., Auckland, New Zealand) [17]. Partial cox1 sequences were identified by local alignment comparison (set reading frames) using amino acid sequences conceptually translated using an online tool http://www.ebi.ac.uk/Tools/st/emboss_transeq/ from the respective loci of the reference sequence of *I. holocyclus* are available from GenBank.

**Phylogenetic analyses**

For morphological phylogenetics, the character matrix was based on adult female specimens. All characters are morphological, collected by examining specimens using light and/or scanning electron microscopy. Characters that could not be scored with complete accuracy in some taxa were excluded from the analysis. Morphological data were analysed employing Maximum Parsimony (MP) in TnT [18], gaps were treated as missing characters, and bootstrap replicates and maximum trees were set at 10,000. In addition, data were analysed using Bayesian Inference (BI) by employing the Markov K model in MrBayes 3.2.6 [19–21]. Lset rates were set to gamma and coding was set to variable. Four simultaneous tree-building chains were used to calculate posterior probabilities (pp) for 2,000,000 generations, saving every 100th tree produced. Based on the final 75% of trees generated, a consensus tree was constructed. *Ixodes tasmani* Neumann, 1899 was used as the outgroup.

For molecular phylogenetics, nucleotide sequences were aligned using the MUSCLE V 3.8.31 program [22] and adjusted manually employing the program Mesquite V 3.03 [23]. Based on pairwise comparisons, sequence differences were calculated using the program MEGA 6.0. [24]. Two separate datasets representing cox1 and ITS-2 were compiled, together with reference sequences from GenBank [14, 25, 26]; *I. tasmani* and *I. uriae* White, 1852 were used as the outgroups, respectively. Both cox1 and ITS-2 sequences were aligned over 519 and 610 bp, respectively, and adjusted manually as described above. Phylogenetic analyses were performed on individual cox1 and ITS-2 datasets using Maximum Likelihood (ML), Neighbour-Joining (NJ) and BI methods. The ML and NJ analyses were performed using MEGA 6.0, and the nodes were tested for robustness with 10,000 bootstrap replicates. The data format was set to DNA and gaps were treated as missing data (10,000 bootstrap replicates, Max. trees was set at 10,000). The likelihood parameters for the BI (TIM2+1 +G for pCXO1 and TVM+G for ITS-2) and ML (Tamura 3-parameter model for both cox1 and ITS-2) analyses were selected based on the Akaike Information Criterion (AIC) test in jModeltest v2.1.5 [27]. The BI was conducted, using Monte Carlo Markov Chain (MCMC) analysis in MrBayes 3.1.2. Four simultaneous tree-building chains were used to calculate posterior probabilities (pp) for 2,000,000 generations, saving every 100th tree produced. Based on the final 75% of trees generated, a consensus tree was constructed.

The phylogenetic trees produced for both morphological and molecular datasets were visually compared separately for concordance in their topologies.

**Results**

**Morphological characterisation**

Out of 74 individual specimens of female *Ixodes* examined, 64 belonged to the subgenus *Stenolixodes*, including *I. anatis* (*n* = 7), *I. dendrolagi* (*n* = 3), *I. cordifer* (*n* = 5), *I. cornutus* (*n* = 5), *I. hirsti* (*n* = 10), *I. holocyclus* (*n* = 13), *I. myrmecobii* (*n* = 17) and *I. trichosuri* (*n* = 4);
whereas, remaining 10 belonged to two subgenera Endopalpiger Schulze (I. tasmani; n = 6) and Multidentatus Neumann (I. auritulus; n = 4).

Character states are presented in Table 2, and the morphological data matrix is provided in Table 3. In addition, principal features of the capitulum used as characters are shown in Fig. 2.

Molecular characterisation

PCR amplification was successful for 27 (out of 64) genomic DNA samples extracted from individual tick specimens (Table 1). Considerable variation in the size (~650 to 750 bp) of amplicons (n = 15) for the ITS-2 was detected on agarose gel, whereas the amplicon size (~700 bp) for cox1 (n = 27) did not differ. DNA sequencing of amplicons for both loci revealed 27 and 15 unique sequences for cox1 and ITS-2, respectively. Sequence length, G+C content, pairwise differences and GenBank accession numbers for cox1 (KY213767–KY213793) and ITS-2 (KY213752–KY213766) sequences are given in Table 4. The length of cox1 sequences for each tick species was 674 bp, whereas that of ITS-2 ranged from 630 to 704 bp. Among various members of Sternalixodes, the highest genetic variation was detected

**Table 3** Morphological character matrix of character states for each taxon used to construct morphological phylogeny

| Species/State | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 2 | 2 | 2 | 2 | 2 | 2 | 2 | 2 | 3 | 3 | 3 | 3 | 3 | 3 | 3 |
|---------------|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|---|
| L. auritulus  | 0 | 0 | 0 | 2 | 1 | 1 | 0 | 1 | 3 | 1 | 1 | 0 | 1 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 |
| L. anatis     | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 |
| L. tasmani    | 1 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 | 0 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | 0 |
| L. holocyclus | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 1 | 1 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 0 |
| L. cornutus   | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 1 | 0 |
| L. myrmecobii | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 1 | 0 | 0 | 0 | 1 | 1 | 1 | 0 | 1 | 1 | 1 | 1 | 1 | 0 |
| L. cordifer   | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 0 | 1 | 1 | 1 | 1 |
| L. dendrolagi | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 1 | 1 | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| L. confusus   | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 1 | 1 | 1 | 1 | 1 | 1 | 1 |
| L. hirsti    | 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 1 | 2 | 0 | 2 | 1 | 0 | 1 | 1 | 1 | 1 | 1 |
| L. trichosuri| 0 | 2 | 0 | 0 | 0 | 0 | 0 | 0 | 0 | 1 | 0 | 1 | 0 | 0 | 0 | 1 | 2 | 0 | 2 | 1 | 0 | 1 | 1 | 1 | 1 | 1 |

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in *I. cornuatus* (number of sequences = 4; pairwise differences 0.2–12%) followed by *I. anatis* (n = 2; 1.2%), *I. myrmecobii* (n = 6; 0.2–1.1%), *I. auritulus* (n = 2; 1.1%), *I. holocyclus* (n = 5; 0.2–0.9%), *I. hirsti* (n = 3; 0.2–0.3%) and *I. trichosuri* (n = 2; 0.3%) (Table 4). Based on ITS-2 sequences, multiple sequences for individual ticks were obtained only for *I. holocyclus* and *I. myrmecobii* and their pairwise differences were 1.4–8.6% and 0.8–3.6%, respectively (Table 4).

**Phylogenetic analyses**

The topology of the phylogenetic trees generated for morphological data employing BI and MP methods were similar (data not shown); hence, the MP tree is presented here, with nodal support values given for both methods (Fig. 3). The morphological phylogram showed six main clades, clade numbers including taxa of the preceding clade. *Ixodes confusus* and *I. dendrolagi* grouped together in clade 1, with moderate statistical support (posterior probability for BI: 0.99; bootstrap value for MP: 87%) (Fig. 3). *Ixodes hirsti*, *I. trichosuri*, *I. auritulus* and *I. anatis* each formed a clade (2, 3, 5 and 6, respectively), with no to high statistical support (Fig. 3). The common Australian paralysis tick, *I. holocyclus*, and *I. cordifer*, *I. cornuatus* and *I. myrmecobii* formed clade 4, with low to moderate statistical support (0.92, 79%).

Molecular phylogenetic analyses revealed that the topology of trees generated from the *cox1* (aligned over 519 positions) and ITS-2 (608 positions) sequence data were similar using BI, NJ and ML (data not shown); hence, only the NJ trees for both loci are presented here (Figs. 4

| Table 4 | Characteristics of sequences of *Ixodes* species determined in this study. GenBank accession numbers, G+C content and length of each sequence, and pairwise differences for each species with more than one specimens are provided | Species | Specimen voucher | GenBank accession no. | Length (bp) | G+C content (%) | Pairwise difference (%) | GenBank accession no. | Length (bp) | G+C content (%) | Pairwise difference (%) |
|---------|-------------------------------------------------------------------------------------------------|--------|------------------|---------------|-----------------|-------------------------|------------------|---------------|-----------------|-------------------------|
| *I. auritulus* | S63 | KY213767 | 674 | 31.75 | 1.1 | – | – | – | – |
| | S64 | KY213768 | 674 | 31.75 | – | – | – | – | – |
| *I. anatis* | S28 | KY213769 | 674 | 31.90 | 1.2 | KY213757 | 703 | 54.62 | – | – |
| | S29 | KY213770 | 674 | 31.90 | – | – | – | – | – |
| *I. holocyclus* | S1 | KY213782 | 674 | 32.20 | 0.2–0.9 | KY213766 | 679 | 55.38 | 1.4–8.6 | – |
| | S4 | KY213783 | 674 | 32.05 | – | KY213765 | 630 | 55.70 | – | – |
| | S37 | KY213781 | 674 | 32.34 | – | KY213756 | 684 | 55.40 | – | – |
| | S17 | KY213779 | 674 | 32.49 | – | KY213762 | 638 | 55.80 | – | – |
| | S39 | KY213780 | 674 | 31.90 | – | KY213755 | 676 | 55.47 | – | – |
| *I. myrmecobii* | S26 | KY213784 | 674 | 30.86 | 0.2–1.1 | KY213758 | 649 | 53.80 | 0.8–3.6 | – |
| | S46 | KY213785 | 674 | 31.16 | – | KY213753 | 656 | 53.70 | – | – |
| | S56 | KY213786 | 674 | 31.00 | – | KY213752 | 657 | 53.60 | – | – |
| | S44 | KY213787 | 674 | 31.00 | – | – | – | – | – |
| | S25 | KY213788 | 674 | 30.70 | – | KY213759 | 668 | 53.30 | – | – |
| | S42 | KY213789 | 674 | 30.86 | – | KY213754 | 647 | 53.80 | – | – |
| *I. cornuatus* | S19 | KY213792 | 674 | 30.42 | 0.2–1.2 | – | – | – | – |
| | S20 | KY213793 | 674 | 30.12 | – | – | – | – | – |
| | S18 | KY213790 | 674 | 30.12 | – | KY213761 | 654 | 53.36 | – | – |
| | S41 | KY213791 | 674 | 30.27 | – | – | – | – | – |
| *I. dendrolagi* | S14 | KY213776 | 674 | 30.70 | – | KY213763 | 672 | 55.20 | – | – |
| *I. trichosuri* | S23 | KY213777 | 674 | 31.90 | 0.3 | – | – | – | – |
| | S21 | KY213778 | 674 | 31.90 | – | KY213760 | 704 | 56.39 | – | – |
| *I. hirsti* | S10 | KY213773 | 674 | 33.10 | 0.2–0.3 | – | – | – | – |
| | S12 | KY213774 | 674 | 33.38 | – | – | – | – | – |
| | S9 | KY213775 | 674 | 33.23 | – | KY213764 | 667 | 56.97 | – | – |
| *I. tasmani* | S68 | KY213771 | 674 | 32.20 | 10.9 | – | – | – | – |
| | S69 | KY213772 | 674 | 32.05 | – | – | – | – | – |

**Notes**

*cox1*: cytochrome *c* oxidase subunit 1  *ITS-2*: second internal transcribed spacer
The cox1 tree had three major clades (Fig. 4) in which *I. cornuatus*, *I. holocyclus* and *I. myrmecobii* formed Clade 1, with mixed statistical support (posterior probability for BI: 0.90; bootstrap value for NJ and ML: 97 and 86%). Individually, five cox1 sequences of *I. holocyclus* determined herein (GenBank accession nos. KY213779–KY213782) grouped together with those previously published from Australia, with strong statistical support (0.99, 100, 99%) (Fig. 4). All six cox1 sequences of *I. myrmecobii* grouped together with strong statistical support (0.99, 100, 96%), whereas four cox1 sequences of *I. cornuatus* found in this study formed two sub-clades with strong statistical support (1.0, 100, 99%) in which two sequences from Tasmania (KY213792 and KY213793) grouped outside the other two sequences from this study (KY213790 and KY213791) as well as previously published sequences (Fig. 4). However, individual sequences of *I. hirsti* determined here (KY213773–KY213777) formed a separate sub-clade compared with previously published sequences of this species. *Ixodes auritulus* and *I. anatis* formed Clade 3 with weak to moderate statistical support (0.98, 67, 70%) (Fig. 4).

The ITS-2 tree contained four major clades (Fig. 5). The composition of some clades was different from that found in the cox1 tree. For example, in the ITS-2 tree, Clade 1 contained only *I. holocyclus* with strong statistical support (1.0, 100, 99%) whereas *I. cornuatus* and *I. myrmecobii* formed a separate clade (Clade 3) with strong statistical support (1.0, 100, 99%) (Fig. 5). Similarly in the cox1 tree, *I. dendrolagi*, *I. hirsti* and *I. trichosuri* formed a separate clade (Clade 2) with strong statistical support (1.0, 99, 95%) whereas *I. anatis* formed a separate clade outside Australian and Papuan *Ixodes* spp. (Fig. 5).

**Discussion**

This study addressed the evolutionary relationships amongst the species of *Sternalixodes*, using both morphological and molecular phylogenetic methods. The cox1 and morphological data suggest that the subgenus is paraphyletic with *I. anatis* which is congruent with a previous suggestion by Heath [28].

The topology was similar in the cox1 and morphological trees, with the Australian and Papuan *Sternalixodes* forming a distinct clade and the New Zealand member of the group *I. anatis* positioned basally, in a separate clade. *Ixodes holocyclus*, *I. cornuatus* and *I. myrmecobii* formed a distinctive clade in both the cox1 and morphological phylogenies. This pattern supports comments made by Roberts [1], who suggested that these three species were closely related and that *I. myrmecobii* and *I. cornuatus* may be subspecies of *I. holocyclus*. *Ixodes hirsti*, *I. trichosuri* and *I. dendrolagi* formed a separate clade distinct from the *I. holocyclus* species group. It is possible that the resulting tree would show *I. dendrolagi* forming a distinct clade with these northern species rather than grouping with *I. hirsti*. The cox1 phylogeny suggests that the outgroup *I. tasmani* contains a cryptic species based on the long branch lengths of the two *I. tasmani* samples (see Fig. 4) as previously proposed by Roberts [1].

While the morphological and cox1 trees showed similar topologies, there were some differences. The position of *I. auritulus* was different between the cox1 tree and the morphological tree, while *I. auritulus* and *I. anatis*
formed a distinct clade in the \( \text{cox}1 \) phylogram. However, in the morphological tree \textit{I. auritulus} formed a clade with the Australian/Papuan \textit{Sternalixodes}. It is likely that the groups are only distantly related as \textit{I. anatis}, \textit{I. auritulus} and the Australian \textit{Sternalixodes} are all morphologically and molecularly distinct. More extensive morphological and molecular phylogenetic analyses are required to adequately illuminate the evolutionary relationships between the three groups. This would require more extensive morphological character sets as well as the use of other molecular markers such as 16S, 28S or complete mitochondrial genomes.

Topological differences were also present between the two molecularly derived trees (see Figs. 4 and 5). The
members of the *I. holocyclus* species group did not form a distinct clade in the ITS-2 tree as was seen in the *cox1* tree, but instead formed two separate clades comprising *I. cornutus* and *I. myrmecobii* in one and *I. holocyclus* in the other. The morphologically similar species, *I. holocyclus* and *I. cornutus* grouped in highly divergent clades contrary to morphological evidence, as well as inferences from previous study of the interspecific relationship of these species by Song et al. [14]. This was likely due to the conserved nature of this marker in ticks. Song et al. [14] commented on the conserved nature of ITS-2 in *Sternalixodes* noting that intraspecific variation between *I. holocyclus* and *I. cornutus* was as low as 0.19%. Despite suggestions that ITS-2 is suitable for inferring evolutionary relationships in ticks [14], it appears that it may not be suited for revealing the relationships between more distantly related species within subgenera.

Within the *cox1* phylogeny, *I. hirsti* was divided into two distinct subclades (see Fig. 4). The GenBank sequences utilised were from ticks collected in South Australia [29], while the sequences obtained during this study were from Victorian specimens. This pattern in the *cox1* sequences coupled with the geographic difference between the two groups suggests the group may be undergoing genetic differentiation.

The results of this study inform a number of historical questions and uncertainties concerning the subgenus *Sternalixodes*. Previously, Song et al. [14] used molecular techniques to assess the validity of the *I. holocyclus* species group. In the present study, based on the *cox1* and ITS-2 trees, it appears that *I. myrmecobii* and *I. cornutus* are not subspecies of *I. holocyclus*, each being a valid species. This contradicts the suggestion made by Roberts [1] that these two species may be subspecies of the widespread *I. holocyclus*. The results of this study are congruent with the results of Song et al. [14] and Jackson et al. [13]. However neither of these studies included the Western Australian species, *I. myrmecobii*.

The findings of this study also provided insights into the validity of *Sternalixodes* as a subgenus. The apparent paraphyletic status of the subgenus based on the position of *I. anatis* in the *cox1* and morphological phylograms validates the suggestion by Heath [28] that the species should not be included in *Sternalixodes*. Heath [28] made the suggestion, citing the morphology of *I. anatis* as being incongruent with the morphological definition of *Sternalixodes*. Based on both morphological and *cox1* data, it appears that *I. anatis* is not a sternalixodid tick and should not be included in the subgenus. This species does not meet the criteria of any of the subgenera of *Ixodes* defined by Clifford et al. [11]. However, Clifford et al. [11] noted that the classification of the subgenera of *Ixodes* was inaccurate in some situations,
especially with regard to the subgenus *Ixodes*. An extensive examination of all existing subgenera using molecular and morphological data should be made in future to provide a more accurate hypothesis of the evolutionary relationships between the subgenera and the validity of the species within them. Based on the distinctive morphology of *I. anatis* and the fact that it does not meet the diagnostic requirements of any of the known subgenera of *Ixodes*, it may require the erection of a new subgenus. However, more extensive molecular data should be accumulated and examined before this can occur.

Although seven of the nine species of *Sternalixodes* were examined in this study, molecular sequences and morphologically complete specimens were not located for *I. confusus* and *I. cordifer*. Although these species are most likely members of *Sternalixodes*, a complete molecular phylogenetic analysis including these species would be desirable to more accurately illuminate the evolutionary relationships within the subgenus.

As *I. myrmecobii* clusters within the *I. holocyclus* species group, a set of ticks known to cause paralysis, the question of its ability to also induce paralysis is raised. Tick-induced paralysis has been extensively studied on the east coast of Australia; however, little information exists concerning ticks in Western Australia, let alone tick paralysis in Western Australia [9]. Studies of *I. myrmecobii* should be undertaken to determine if this species can induce paralysis. Roberts [30] and Kemp [31] noted that *I. hirsti* has been recorded to cause paralysis. Kemp [31] also proposed that all sternalixodid ticks may be capable of causing paralysis. As *I. hirsti* clustered with *I. trichosuri* and *I. dendrolagi* within the morphological and cox1 phylogeny, it is possible that these species may also be capable of inducing paralysis, however, this should be investigated.

**Conclusion**

In conclusion, the cox1 and morphological data suggest that the subgenus *Sternalixodes* is paraphyletic, and *I. anatis* should not be included in this subgenus. Based on the phylogenetic analyses of cox1 and ITS-2 sequences, it appears that *I. myrmecobii* and *I. cornuatus* are not subspecies of *I. holocyclus*, each being a valid species. Although this study has improved insights into the taxonomic status of the subgenus *Sternalixodes*, a complete morphological and molecular (using multiple markers) phylogenetic analysis including all nine species of the subgenus would be desirable to more accurately illuminate the evolutionary relationships within the subgenus.

**Abbreviations**

AIC: Akaike information criterion; BI: Bayesian inference; cox1: Cytochrome c oxidase subunit 1 gene; ITS-2: Internal transcribed spacer 2; MCMC: Monte Carlo Markov Chain; ML: Maximum likelihood; Nt: Neighbour-joining

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**Availability of data and materials**

The data supporting the conclusions of this article are included within the article. The nucleotide sequences generated during this study are deposited in GenBank under the accession numbers KY213752–KY213793.

**Authors’ contributions**

MK, IB and AJ conceived the project and participated in the study design. MK carried out the laboratory work, data analyses, interpretation of data, and also drafted the manuscript, with guidance from co-authors. MK, AVK and AJ undertook phylogenetic analyses. IB and AJ participated in data interpretation and provided critical inputs on the draft manuscript. All authors read and approved the final manuscript.

**Competing interests**

The authors declare that they have no competing interests.

**Consent for publication**

Not applicable.

**Ethics approval and consent to participate**

Not applicable.

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