Borrelia sp. phylogenetically different from Lyme disease- and relapsing fever-related Borrelia spp. in Amblyomma varanense from Python reticulatus

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Abstract

Background: Species of the genus Borrelia are causative agents of Lyme disease and relapsing fever. Lyme disease is the most commonly reported vector-borne disease in the northern hemisphere. However, in some parts of the world Lyme borreliosis and relapsing fever may be caused by novel Borrelia genotypes. Herein, we report the presence of a Borrelia sp. in an Amblyomma varanense collected from Python reticulatus.

Methods: Ticks were collected from snakes, identified to species level and examined by PCR for the presence of Borrelia spp. flaB and 16S rRNA genes. Phylogenetic trees were constructed using the neighbour-joining method.

Results: Three A. varanense ticks collected from P. reticulatus were positive for a unique Borrelia sp., which was phylogenetically divergent from both Lyme disease- and relapsing fever-associated Borrelia spp.

Conclusion: The results of this study suggest for the first time that there is a Borrelia sp. in A. varanense tick in the snake P. reticulatus that might be novel.

Keywords: Borrelia, Amblyomma varanense, Amblyomma pattoni, Tick, Thailand

Background

Tick infestation in snakes occurs worldwide and involves the following species: Amblyomma gervaisi in the northern region of western Ghats in India [1], Rhipicephalus sanguineus (sensu lato) in Malaysia [2], Amblyomma varanense and Amblyomma helvolum in Thailand [3, 4] and Amblyomma hydrosauri in Australia [5]. Hirunkanokpun et al. [6] detected several bacterial species in the national parks of Thailand, but no Borrelia spp. were found. The aim of this study was to determine the presence of Borrelia spp. within Amblyomma spp. ticks collected from five snake species. In addition, phylogenetic analyses of Borrelia spp. are also presented.

Methods

Tick collection and identification

Tick collection from snakes was performed in February 2014 in Lopburi Province, Thailand (14°48′1.61″N, 100°38′10.75″E). We observed snake scales to identify partial protrusions of tick bodies outside of the scales. Ticks were collected from the skin beneath the scales using forceps. Ticks were identified according to their morphology using standard taxonomic keys [7–11].

DNA extraction and amplification

The ticks were washed individually and homogenised in 200 μl of 10× PBS solution. DNA extraction was performed using the QIAamp DNA Blood Mini Kit (Qiagen, Hilden, Germany). Oligonucleotide primer pairs FLA1-FLA2 (BflaPAD 5′-GAT CA(G/A) GC(T/A) A TA CCA(A/T) ATG CA-3′; BflaPBU, nest- 5′-GCT GAA GAG CTT GGA ATG CAA CC-3′) were used.
3′; BflaPCR, nest-5′-TGA TCA GTT ATC ATT CTA ATA GCA-3′; BflaPDU 5′-AGA TTC AAG TCT GTT TTG GAA AGC-3′) and 16S rDNA (16SF1 5′-ATA ACG AAG AGT TTG ATC CTG GC-3′; 16SR 5′-CAG CCG CAC TTT CCA GTA CG/3′) were used in this study to identify target *Borrelia* DNA in the ticks [7, 8]. The positive PCR products from tick samples were purified using a High Pure PCR product purification kit (Roche, Basel, Switzerland). Sequencing reactions were performed with BigDye Terminator v3.1 Cycle Sequencing Kits (Applied BioSystems, Waltham, Massachusetts, USA) based on the fluorescent-label terminator method. Sequencing products were analyzed using a Genetic Analyzer 3730XL automated DNA sequencing system (Applied BioSystems, Waltham, Massachusetts, USA).

**Phylogenetic analysis**

Phylogenetic trees were constructed using the neighbour-joining method (PAUP 4.0b1) [12]. DNA gaps or missing data were excluded from the analyses. Confidence values for individual branches of the resulting tree were determined by bootstrap analysis with 1000 replicates.

**Results and discussion**

Ticks were collected from the following five snake species: *Python reticulatus*, *Ophiophagus hannah*, *Ptyas korros*, *Naja kaouthia* and *Elaphe radiata*. Four *Amblyomma varanense* ticks (three males and one female) were collected from one *P. reticulatus*. One *A. varanense* tick (one female) and two *Amblyomma pattoni* (males) were collected from *O. hannah*. Two *A. pattoni* (males) were collected from *N. kaouthia*. In addition, two *A. pattoni* males

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**Fig. 1** Pictures of ticks identified in this study. a *Amblyomma pattoni*, male, dorsal view. b *Amblyomma pattoni*, male ventral view. c *Amblyomma varanense*, male, dorsal view. d *Amblyomma varanense*, male, ventral view. e *Amblyomma varanense*, female, dorsal view. f *Amblyomma varanense*, female, ventral view. Scale-bars: 1 mm
were collected from *E. radiata*. Finally, one *A. varanense* tick (male) was collected from *E. radiata*. Tick species reported in this study have hypostomal dentition 3/3. The male of *A. pattoni* has coxa I with an inconspicuous internal spur, which is sometimes fused with the more prominent external spur, and cervical pits are comma-shaped. The male of *A. varanense* has coxa I with the external spur noticeably longer than the internal and the female has coxa I with the internal spur smaller than the external spur, but always separated from the latter (Fig. 1).

A total of 12 ticks was collected from snakes and examined by PCR for the presence of the *Borrelia* spp. genes. Of these, three ticks, all identified as *A. varanense* isolated from *P. reticulatus*, were positive for *Borrelia* spp. No *Borrelia* spp. were detected in *A. pattoni* or in *A. varanense* collected from other snake species.

*Borrelia* sp. DNA sequences were compared with sequences in the NCBI GenBank database by nucleotide BLAST. The 16S rRNA gene sequence of this *Borrelia* sp. is 100 % identical (1449/1449 bp) to *Borrelia* sp. BF16 (GenBank: AB473538) and was submitted to GenBank and assigned as KU497718 (*Borrelia* sp. in *Amblyomma varanense* from *Python reticulatus*). The amplified PCR product of the flaB gene for flagellin was approximately 384 bp. A sequence analysis of the flaB gene for flagellin from the *Borrelia*-positive ticks revealed that this gene sequence is 99 % similar (294/296 bp) to a sequence for *Borrelia* sp. BF16 flaB gene for flagellin (isolate BF16; GenBank: AB473488). The 384 bp *Borrelia* flaB consensus sequence was submitted to GenBank and assigned as *Borrelia* sp. KT758064. By contrast, this sequence was only 87 % similar (337/387 bp) to the following: the *B. turcica* flaB gene for flagellin (strain IST4; GenBank: AB109244); *B. turcica* IST7 flagellin gene (GenBank: KF422815) (336/387 bp); and *Borrelia* sp. tAG66 M flaB gene for flagellin (GenBank: AB529322) (336/387 bp).

Phylogenetic trees were constructed using the neighbour-joining method (PAUP 4.0b1) [12]. Individual branch confidence values were determined by bootstrap analysis with 1000 pseudoreplicates (Figs. 2 and 3). The phylogenetic trees for both genes of *Borrelia* spp. inferred from the complete 16S rRNA gene sequences (Fig. 2) and partial sequences of the flaB genes (Fig. 3) indicated that *Borrelia* sp. from the present study is related to *Borrelia* sp. (GenBank: AB473538) from reptiles and to *B. turcica* IST7 (GenBank: KF422815) flagellin gene, respectively, but belongs in a different group from *Borrelia burgdorferi*. The phylogenetic relationships among relapsing fever-associated *Borrelia* spp. and Lyme disease-associated *Borrelia* spp. were reported previously using rrs and 16S rDNA [13, 14]. The flaB and 16S rRNA gene sequences of *Borrelia* sp. of *A. varanense* from *P. reticulatus* isolated in this study, formed a separate branching root from both Lyme disease-associated *Borrelia* species and relapsing fever-associated *Borrelia* species.

![Fig. 2 Neighbour-joining phylogenetic analysis of Borrelia 16S rRNA gene sequences including the newly-generated KU497718 for Borrelia sp. in Amblyomma varanense from Python reticulatus](image-url)
Conclusion
Our findings suggest that Borrelia sp. in A. varanense from P. reticulatus might be novel and phylogenetically divergent from both Lyme disease- and relapsing fever-associated Borrelia species.

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Availability of data and material
The sequences for Borrelia sp. in Amblyomma varanense from Python reticulatus are submitted to the GenBank database under accession numbers KU497718 (16S rRNA) and KT758064 (flaB gene sequence).

Authors’ contributions
WT and AA planned and designed the study. WL and DB carried out the field collection of samples. WT and AA carried out the majority of the laboratory and tick identification work. SH performed the phylogenetic analysis. RS generated 16S rRNA gene sequences of Borrelia and helped in identification section. WT and AA wrote the manuscript with some advice from VB. All authors read, discussed, and approved the final version of the 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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