Molecular identification of *Rickettsia felis* in ticks and fleas from an endemic area for Brazilian Spotted Fever

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Rickettsioses are arthropod-borne diseases caused by parasites from the Order Rickettsiales. The most prevalent rickettsial disease in Brazil is Brazilian Spotted Fever (BSF). This work intends the molecular detection of those agents in ectoparasites from an endemic area of BSF in the state of Espírito Santo. A total of 502 ectoparasites, among them Amblyomma cajennense, Amblyomma dubitatum (A. cooperi), Riphicephalus sanguineus, Anocentor nitens and Ctenocephalides felis, was collected from domestic animals and the environment and separated in 152 lots according to the origin. Rickettsia sp. was detected in pools of all collected species by amplification of 17kDa protein-encoding gene fragments. The products of PCR amplification of three samples were sequenced, and Rickettsia felis was identified in R. sanguineus and C. felis. These results confirm the presence of Rickettsia felis in areas previously known as endemic for BSF, disease caused by Rickettsia rickettsii. Moreover, they show the needing of further studies for deeper knowledge of R. felis-spotted fever epidemiology and differentiation of these diseases in Brazil.

Key words: Rickettsia felis - Rickettsia rickettsii - Ctenocephalides felis - Brazilian Spotted Fever - R. felis-spotted fever - tick

Brazilian Spotted Fever (BSF) is the most prevalent rickettsial disease in Brazil (Lemos et al. 2001). The illness is caused by *Rickettsia rickettsii*, a bacterial organism belonging to the family Rickettsiaceae and to the Order Rickettsiales, whose members are intracellular mandatory microorganisms, many of them causing infections worldwide-distributed in humans and other vertebrate and invertebrate hosts. These organisms are classically transmitted to humans via arthropod vector bites. In Brazil, the tick *Amblyomma cajennense* is the main vector of *Rickettsia rickettsii* (Labruna et al. 2004) and reservoir of BSF (Aragão & Fonseca 1961, Figueiredo et al. 1999).

Rickettsia felis-spotted fever is caused by R. felis, whose main vector is the cat flea, Ctenocephalides felis. Once in this host, the pathogen is maintained by transovarial transmission with no lethal effect, which represents a factor of great importance for the epidemiology of this emerging rickettsiosis (Azad et al. 1992, Rolain et al. 2003).

Studies have demonstrated the presence of *R. felis* in Brazil by serology performed in human cases (Raoult et al. 2001) and *C. felis* infection confirmed by polymerase chain reaction (PCR) (Horta et al. 2006, Oliveira et al. 2002).

The objective of this work was the molecular detection of rickettsiae circulation in ectoparasites collected from domestic animals and from domestic environments, following a BSF outbreak, in an endemic area of BSF. This study sought to contribute to a wider understanding and knowledge of the rickettsial diseases related to these agents in Brazil.

MATERIALS AND METHODS

Study area - The study began in November 2003, in the municipal district of Nova Venécia (18°43'38"S, 40°24'02"W), Northwestern of state of Espírito Santo, Brazil, which is considered an endemic area for BSF. Nova Venécia has an estimated population of 44,380 inhabitants (http://www.ibge.gov.br). From August 1st to October 31st 2003, 16 suspected cases of rickettsiosis and two confirmed cases by indirect immunofluorescence assay (IFA) for *R. rickettsii* (data supplied for Vigilance of Health Secretary, Espírito Santo, Brazil) were identified, as well as the occurrence of one death, in this municipal district. The majority of cases were concentrated in the rural community named Patrimônio do XV, with about 1,200 inhabitants.

Collection and preparation of material for analysis - A total of 502 ectoparasites (ticks and fleas) were collected from domestic animals (24 dogs and 5 horses) and from the natural environment, in the nymph and adult stages, as well as eggs laid in a controlled chamber. The ectoparasites were hand-picked from the animals, using tweezers when necessary. CO₂ traps were used to capture free-living ticks in sites detected as being the probable origin of infection, according to the epidemiologic inquiry, considering possible environments for contact between human and ticks and evidence for the presence of capybaras. The ectoparasites were separated in 152 lots according to the origin, and adult stages identified through a stereoscopic microscope using Aragão and

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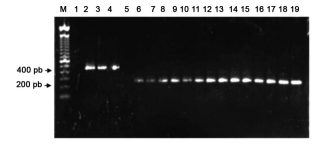
+ Corresponding author: mafra@ufv.br Received 5 October 2007 Accepted 20 March 2008 Fonseca's (1961) taxonomic keys for ticks and Linardi and Guimarães' keys (2000) for fleas. At the laboratory, the ectoparasites were kept in a BOD incubator, at 25°C for 2-5 days, for reactivation and multiplication of the rickettsial agents according to Hayes and Burgdorfer (1982), and stored at -20°C until nucleic acid extraction.

DNA extraction and PCR - After the surface sterilization by immersion in absolute ethanol for 10 min, the ectoparasites were washed in PBS, crushed in 200 ul of the same buffer, and the DNA extraction was performed using the method described by Billings et al. (1998). For the amplification of 434 bp-portion of the gene encoding Rickettsia genus-specific 17-kDa protein, the primers 17kD1 (5'- GCTCTTGCAACTTCTATGTT-3') and 17kD2 (5'-CATTGTTCGTCAGGTTGGCG-3') described by Webb et al. (1990) were used, under the following conditions: an initial stage at 94°C for 3 min, 30 cycles at 95°C for 1 min, 50°C for 1 min, and 72°C for 2 min, followed by an step at 72°C for 5 min. The reactions were performed in 25 µl containing PCR 1X (InvitrogenTM) buffer, 0.4 mM of each primer, 1 mM of MgCl₂, 0.05 mM of each dNTP, 1U of recombinant Tag DNA Polymerase (InvitrogenTM), and 2 ul of DNA from each sample or 0.5 ul of purified Rickettsia prowazekii DNA for positive control. The amplifications were initially carried out in 30 pools, each containing about 5 lots of ectoparasites. After detecting the positive pools, they were separated and each lot was subjected to a new PCR to identify the infected lots. Nested-PCR was used for the negative samples in the first amplification, using 1 µl of the first reaction as template and the pair of genus-specific primers 17kN1 (5'- CATTACTTGGTTCTCAATTCGGT-3') and 17kN2 (5'-GTTTTATTAGTGGTTACGTAA-3') described by Schriefer et al. (1994), to amplify an inner 232-pb region of the gene encoding the 17 kDa protein. The reagent concentrations and amplification conditions were the same as described on the first amplification reaction. The reaction products of PCR were visualized in 1% agarose gel.

Sequencing of PCR products - The PCR amplification products were cloned into the vector pCR® 2.1-TOPO® (InvitrogenTM TOPO® TA Cloning® Kit), following the manufacturer's protocol. Plasmid DNA extraction was performed using the High Pure Plasmid Isolation Kit (Roche Diagnostic), following the manufacturer's instructions. The generated nucleotide sequences were edited with CHROMAS (http://www.mb.mahidol.ac.th/pub/chromas/chromas.htm) and compared with the corresponding homologous sequences available through GenBank, using Discontiguous Mega Blast (http://www.ncbi.nlm.nih.gov).

RESULTS

Ticks and fleas were identified as *Amblyomma dubitatum* (*A. cooperi*), *A. cajennense*, *Riphicephalus sanguineus*, *Anocentor nitens* and *C. felis*. In this study, 28 lots of ectoparasites contained *Rickettsia* DNA amplified by nested-PCR, from these lots two were positive in the first reaction and 26 were positive in the second reaction, using genus-specific primers (Table I). Figure shows the generated band patterns.



Result of PCR to detect *Rickettsia*, in 1% agarose gel. Lanes - M: 100 bp DNA ladder size marker; 1 to 4: products of first reaction; 1: negative control, 2: positive control, 3 and 4: lots 45 and 71, respectively; 5 to 19: products of second reaction, 5: negative control, 6: positive control, 7 to 19: lots 4, 10, 26, 31, 34, 35, 36, 38, 42, 44, 59, 63, and 65, respectively.

The sequencing of PCR amplification products of ectoparasite lots 45, 71 and 88 (Table I) was successful. The generated nucleotide sequences had greater similarity with *R. felis* sequences deposited in GenBank, and the sequence obtained from *R. sanguineus* lot (lot 88) showed 96% similarity to *R. felis*, whereas the sequences obtained from lots 45 and 71, both *C. felis*, showed 98% similarity to *R. felis* (Table II).

DISCUSSION

These results present a contribution to the initial understanding of rickettsial ecology in the studied area. The amplification of gene fragments encoding the 17kDa protein showed that *Rickettsia* spp. circulates among all the ectoparasite species collected in the studied area, confirming the role of different ectoparasites in the maintenance of these organisms.

The detection of *R. felis* in *R. sanguineus* and *C. felis* is corroborated by other studies carried out in Brazil (Galvão et al. 2003, 2006). Knowing that by IFA with specific antigens, the sera of patients with rickettsiosis during the outbreak showed a positive title for *R. rickettsii*, the presence of *R. felis* in ectoparasites from the studied area indicates the possibility of occurrence of *R. felis*-spotted fever, concomitant or not with BSF, in the studied area.

Some studies (Phillip et al. 1978, Oliveira et al. 2002, Horta et al. 2007) have indicated that at least two species of pathogenic rickettsia can be circulating in the endemic foci of BSF. Horta et al. (2007) found evidences of at least four Rickettsia species (R. rickettsii, R. parkeri, R. felis and R. bellii) in the studied areas (4 endemic and 1 non-endemic for BSF), however serological evidences of rickettsial infection in humans and/or animals were found for only two species: R. rickettsii and R. parkeri. These results suggest that some BSF cases occurred in endemic areas may have been caused by other rickettsiae rather than R. rickettsii, and being R. rickettsii the sole antigen regularly used in BSF serological diagnosis, some human cases of BSF due to other rickettsiae of this group have been wrongly identified as BSF in Brazil (Horta et al. 2007).

TABLE I

Ectoparasite lots that were positive for *Rickettsia* spp., using polymerase chain reaction, in Nova Venécia, state of the Espírito Santo, Brazil

Lot	Origin of ectoparasite	Species	Stages	Number of ectoparasites/lo
4	Dog	-	Eggs	ND
10	Dog	Ctenocephalides felis	Adult	7
26	Environment	Amblyomma dubitatum (A. cooperi)	Adult/male	4
31	Environment	<u>-</u>	Nymphs	5
34	Horses	-	Nymphs	4
35	Horses	Anocentor nitens	Adult/female	8
36	Horses	Anocentor nitens	Adult/female	4
38	Horses	Amblyomma cajennense	Adult/female	3
42	Environment	Amblyomma dubitatum (A. cooperi)	Adult/female	3
44	Environment	-	Nymphs	4
45	Dog	Ctenocephalides felis	Adult	8
59	Dog	Riphicephalus sanguineus	Adult/female	4
63	Dog	Riphicephalus sanguineus	Adult/male	10
65	Dog	Riphicephalus sanguineus	Adult/female	2
69	Dog	Ctenocephalides felis	Adult	8
70	Dog	Ctenocephalides felis	Adult	8
71	Dog	Ctenocephalides felis	Adult	1
74	Horses	-	Nymphs	3
74	Horses	Anocentor nitens	Adult/male	2
85	Dog	Riphicephalus sanguineus	Adult/male	3
87	Environment	Amblyomma dubitatum (A. cooperi)	Adult/male	3
88	Environment	Riphicephalus sanguineus	Adult/female	3
88	Environment	<u> </u>	Nymphs	4
98	Horses	Amblyomma cajennense	Adult/male	5
99	Horses	-	Nymphs	3
104	Environment	-	Nymphs	1
131	Dog	Riphicephalus sanguineus	Adult/male	4
136	Dog		Eggs	ND
137	Dog	Ctenocephalides felis	Adult	1
152	Dog		Eggs	ND

ND: not determined.

TABLE II
Similarity analysis of nucleotide sequences from lots 45, 71 and 88 with *R. felis* sequences from GenBank, according to Discontiguous MegaBlast

Accession number at GenBank	Species	Similarity ^a	Vector (lot) ^b
CP 000053.1	R. felis URRWX Cal 2	98% (410/415)	C. felis (45)
DQ 102709.1	R. felis ssc50	98% (379/383)	C. felis (71)
DQ 102709.1	R. felis ssc50	96% (225/232)	R. sanguineus (88)

a: percentage of similarity calculated as identical bases/total bases; b: lot number according to Table I.

It is also known that the occurrence of cross-reaction between *R. rickettsii* and *R. felis* serology is common, as reported by Raoult et al. (2001). This fact could also explain the detection of *R. felis* in the studied area in this present work, where cases of rickettsiosis were diagnosed as BSF due to *R. rickettsii*, not excluding, however, the potential presence of this and/or other *Rickettsia* species in the analysed ectoparasites. It is important to point out that the patients' clinical diagnosis was not completely compatible with the BSF symptoms, therefore confirming the possibility of facing, in this area, a pathology of emerging character in Brazil - *R. felis*-spotted fever.

These results emphasize the need for more specific techniques to the diagnosis of rickettsiosis that will allow a better characterization of these diseases in Brazil, as well as the development of strategies more compatible with the real situation of the country.

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