Molecular detection of *Rickettsia* spp. in ticks collected from dogs from the Department of Piura, Peru

Detecção molecular de *Rickettsia* spp. em carrapatos coletados de cães provenientes do Departamento de Piura-Peru

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Abstract

The aim of this study was to determine the frequency of ticks positive for genus *Rickettsia* bacteria among ticks collected from domestic dogs in the Department of Piura, Peru, using polymerase chain reaction (PCR) analysis. Ticks were collected from dogs in urban areas of the metropolitan region of Piura, Peru. Only three species of ticks were identified; 977 *Rhipicephalus sanguineus* (180 nymphs, 417 females, and 380 males), Six *Amblyomma triste* females, and one *Amblyomma tigrinum* male. After classifying the specimens morphologically by stage, species, and sex, their total DNA was tested by PCR using primers that amplify fragments of the *gltA*, *ompA*, *ompB*, and *htrA* genes. The resulting positive sample was sequenced, compared to the GenBank database, and analyzed phylogenetically. The *Rickettsia* spp. infection rate in the tick pools was 0.2% (1/484); the positive specimen was an *R. sanguineus* tick. GenBank analysis of the positive sequence revealed 100% identify with *Rickettsia felis*; however, no products of the *htrA*, *ompA* and *ompB* genes were amplified from this sample. To the best of our knowledge, this is the first report of *R. felis* in *R. sanguineus* in Peru.

Key words: Acari. Ixodidae. Molecular analysis. Rickettsiosis. South America.

Resumo

O objetivo do presente estudo foi determinar a frequência de carrapatos coletados de caninos domésticos positivos para bactérias do gênero *Rickettsia* por meio da Reação em cadeia da polimerase (PCR) no Departamento de Piura, Peru. Desta forma, carrapatos foram coletados de caninos de zonas urbanas da região Metropolitana de Piura, Peru. Apenas três espécies de carrapatos foram identificadas, sendo 977 *R. sanguineus* (180 ninhãs, 417 fêmeas e 380 machos), 6 fêmeas da espécie *Amblyomma triste* e um macho *Amblyomma tigrinum*. Após classificados morfologicamente por estádio, espécie e sexo, o DNA total dos espécimes foi testado pela PCR utilizando iniciadores que amplificam fragmentos dos genes *gltA*, *ompA*, *ompB* e *htrA*. A amostra positiva foi sequenciada, comparada com o banco de dados depositados no GenBank e analisada filogeneticamente. Verificou-se, mediante PCR, uma taxa de...
infecção de 0,2% (1/484) “pools” de carrapatos positivos para *Rickettsia* spp. O espécime positivo foi um carrapato *Rhipicephalus sanguineus*. A análise no GenBank da sequência positiva apresentou uma similaridade de 100% com a espécie *R. felis*. Essa amostra não amplificou produtos para os genes htrA, ompA e ompB. Este é o primeiro relato de *R. felis* em *R. sanguineus* no Peru.


Genus *Rickettsia* bacteria belong to the alpha subdivision of the class Proteobacteria. This genus is subdivided into four groups: 1) the typhus group (TG); 2) the spotted fever group (SFG), containing >20 species; 3) the transitional group, represented by species such as *R. akari*, *R. australis*, and *R. felis*; and 4) the basal group (LABRUNA et al., 2011).

The geographic reach of spotted fever (SF) is entirely determined by vector (tick) distribution and activity. In Mexico, *Rhipicephalus sanguineus* has been implicated as a vector of *R. rickettsii*. This intradomiciliary tick mainly parasitizes domestic dogs, but can also parasitize humans; thus, it could play an important role as a vector in some endemic areas with specific ecological characteristics (EREMEEVA et al., 2011).

A number of studies have reported the presence of several genera of ticks in different regions of Peru, such as Piura; the genera *Amblyomma*, *Dermacentor*, *Ixodes*, and *Rhipicephalus* have been detected on dogs and other hosts. Furthermore, there is also molecular evidence of the presence of rickettsial agents in ticks in these regions (FORSHEY et al., 2010).

Advances in laboratory techniques for the study of different agents, especially molecular biology methods, have enabled the identification of new species around the world, as well as pathogenic species in both dogs and tick triturates in areas or continents without previous reports (EREMEEVA et al., 2011; FORSHEY et al., 2010).

The clinical importance of these agents, the large population of hemoparasite vectors, and the fact that many of these parasites have zoonotic potential, emphasize the need to characterize the agents present in the northern region of Peru in order to delineate control strategies and diagnostic methods and to further elucidate the epidemiological characteristics of dog infections caused by these agents.

Thus, the aim of this study was to determine the frequency of genus *Rickettsia* in domestic canine ticks from the metropolitan region of the Piura Province, Department of Piura, Peru, using polymerase chain reaction (PCR) analysis and to examine the frequency of *Rickettsia felis* in *Rhipicephalus sanguineus* ticks.

This study was approved by the PRÓ-REITORIA DE PESQUISA E PÓS-GRADUAÇÃO (Office of the Vice-President for Research and Graduate Studies) Animal Research Ethics Committee (CEPA), registration number 759.

This was a cross-sectional study with convenience sampling. Samples were obtained from dogs in urban areas of the Piura metropolitan region, located in the central zone of the Department of Piura, Peru (Figure 1) at an altitude of 29 meters above sea level, latitude 4°59’24.00”S, and longitude 80°24’36.00”W (Greenwich). The region is characterized by maximum and minimum annual average temperatures of 31.2 and 17.7 °C, respectively, and an average relative air humidity of 83% in the morning and 51% in the evening.

Once the dog owners agreed with the proposed procedures and signed the informed consent form, the ticks were removed from the dogs using hemostatic forceps with a vertical rotational movement, taking care to avoid separating the body from the mouthpiece. The greatest possible number of ticks was collected within a period of physical containment that prevented excessive stress to the animal. Following collection, the ticks were stored and transported in flasks containing absolute isopropanol.
The species were identified morphologically using a binocular stereoscope. The ticks were separated into groups according to species, sex, and host. Females were stored individually, males in pools of up to seven individuals, and nymphs in pools of up to ten specimens; each species pool from the same animal was placed in a 1.5 mL microtube containing absolute isopropanol. Females, especially engorged ones, were processed individually because they exceeded the ideal weight for DNA extraction.

Figure 1. (A) Geographical location of the Department of Piura, Peru, South America. (B) Geographical location of the districts (names in green, inside of red circle) within the eight provinces (names in red) of the Department of Piura.

DNA was extracted from each tick sample using the tick DNA extraction method with phenol/phenol-chloroform (SANTOLIN et al., 2013).

PCR amplification was conducted with primers Rr19O.70p/Rr19O.602n (EREEMEEVA et al., 1994), BG1-21/BG2-20 (REGNERY et al., 1991), 17k-5/17k-3 (LABRUNA et al., 2004), and RpCS.877p/RpCS.1258n (REGNERY et al., 1991), which amplify fragments of approximately 532 bp, 650 bp, 549 bp, and 381 bp of the ompA (membrane protein) ompB (membrane protein), htrA (a 17 kDa protein), and gltA (citrate synthase) genes, respectively.

The final concentrations of the reagents used were: 0.2 mM of each dNTP, 10 μM of each primer, 3.0 mM MgCl₂, 1.25 U GoTaq Flexi DNA Polymerase (Promega, Madison, WI, USA), 1× Green GoTaq Reaction Buffer, and 5 μL of the DNA
sample in a total reaction volume of 25 μL. The PCR reaction conditions included an initial DNA denaturation step at 95 °C for 5 minutes; followed by 40 consecutive cycles of 95 °C for 40 seconds (denaturation), 55 °C for 60 seconds (annealing), and 72 °C for 70 seconds (extension); and a final extension at 72 °C for 7 minutes (REGNERY et al., 1991; EREMEEVA et al., 1994). For each reaction, a positive sample of *R. parkeri* strain At24 and a negative control (Ultra Pure TMD DNase/RNase-Free Distilled Water) were included.

The resulting products were visualized by electrophoresis on a 1.5% agarose gel stained with GelRed that was exposed to ultraviolet light to reveal the bands. A 100 base pair molecular weight marker (100 bp DNA Ladder, Invitrogen, Carlsbad, CA, USA) was used to determine the size of the amplified products.

The amplified PCR products from the positive samples were purified using the Illustra GFX PCR DNA and Gel Band Purification Kit commercial kit (GE Healthcare Life Sciences) and sequenced with a capillary-type Sanger platform on an ABI 3730 DNA Analyzer (Applied Biosystems, Life Technologies). The nucleotide sequences were compared to the corresponding homologs available in the National Center for Biotechnology Information (www.ncbi.nlm.nih.gov) GenBank database using the Basic Local Alignment Search Tool (BLAST; www.ncbi.nlm.nih.gov/blast.cgi).

The sequences were aligned using the Clustal W program version 2.1 and the alignment gaps were removed. Next, phylogenetic analysis was carried out using PHYLIP (Phylogeny Inference Package) with maximum stringency analysis and a bootstrap value of 1000 and a consensus tree was generated (using Seqboot, Dnapars, and Consense software). Graphical processing of the tree was performed with Mega version 6.06 software. The sequences used in the comparison were chosen because they cluster with *R. felis* according to the phylogeny proposed by Tsui et al. (2007); additional sequences of other Rickettsia species were also used for comparison.

A total of 984 dog ticks were collected from 216 dogs. Only three species of ticks were identified: 977 *R. sanguineus* (180 nymphs, 410 females, and 380 males), six *Amblyomma triste* females, and one *Amblyomma tigrinum* male. It is important to mention that ticks were grouped into 484 pools.

In a total of 484 tick pools, an infection rate of 0.2% (1/484) was determined by PCR. The positive tick was morphologically identified as *R. sanguineus* (adult female); analysis of the *gltA* gene sequence obtained demonstrated 100% (387 bp) identity with *R. felis* (KY887024). Although positive results were obtained for the *gltA* gene, no PCR product amplification was achieved for the *htrA, ompA*, and *ompB* genes.

The vertebrate host of the tick was a 2-year-old dog from the Tacalá neighborhood in the Castilla district; the dog did not exhibit any clinical signs of disease.

The positive sample sequence (KY887024) was 97% and 98% identical to *R. felis* sequences isolated in Taiwan (AF210692) and Spain (AF540555), respectively. Phylogenetic analysis showed that the sequence grouped in the *R. felis* clade, as shown in Figure 2. Moreover, the isolate identified in this study (KY887024) demonstrated 92% and 94% identify with isolates obtained in Panama (HM582437) and Peru (GU117906), respectively. In addition, compared to other *Rickettsia* species, the sequence isolated in this study exhibited 93%, 94%, and 91% identity with *R. rickettsii* (U59729), *R. akari* (U59717), and *R. typhi* (U59714) isolated in the USA, respectively.

Several surveys of *R. felis* in different countries have demonstrated that this agent has a worldwide distribution. *R. felis* has been identified in *Ctenocephalides* spp. fleas in Mexico, USA, Brazil, Peru, Ethiopia, Spain, and Germany (LABRUNA et al., 2011). However, it has also been reported in other species of fleas, mosquitoes, and ticks (CARDOSO et al., 2006, REIF; MACALUSO, 2009).
This study is the first to confirm the presence of *R. felis* in *R. sanguineus* in Peru. Previous studies in Brazil have detected *R. felis* in *Amblyomma sculptum* (*A. cajennense sensu lato*) and in *R. sanguineus* using nested PCR and conventional PCR directed at the rickettsial *htrA* gene. In addition, Tanikawa et al. (2013), were able to detect *R. felis* in *R. sanguineus* by conventional PCR and real-time PCR using the rickettsial *gltA* gene. Similar to the results of Cardoso et al. (2006), Oliveira et al. (2008), and Tanikawa et al. (2013), who failed to amplify other *R. felis* gene fragments from their tick specimens, no other rickettsial genes (*ompA* and *htrA*) could be amplified by standard PCR in this study. These negative PCR results could be due to the low rickettsial load in *R. felis*-infected *R. sanguineus* when using PCR protocols that amplify larger fragments, which are therefore less sensitive.

![Figure 2](image)

**Figure 2.** Phylogenetic tree based on the partial *gltA* gene sequence of the *R. felis*-positive sample from the metropolitan region of the province of Piura, Peru and similar sequences obtained from the GenBank database. Bootstrap values obtained after 1000 re-samplings are shown. The scale bar represents the difference (number of nucleotides) of each *Rickettsia* species within each branch. The sequence marked with (*) is the sample determined as *Rickettsia* spp.-positive by PCR and confirmed by sequencing as *R. felis*.

The role of this tick in the ecology and horizontal transmission of *R. felis* remains unclear (REIF; MACALUSO, 2009). It is likely that the source of *R. sanguineus* infection by *R. felis* is blood feeding from a host parasitized by *R. felis*-infected fleas. However, many rickettsial species exhibit robust growth in cell lines derived from ticks and other arthropods. Recently, the ISE6 tick cell line was used to isolate previously uncultured *R. felis* from cat fleas (PORNWIROON et al., 2006). If future studies confirm the ability of *R. sanguineus* to transmit *R. felis*, the results of the current study would support the need for tick and flea control in the prevention of spotted fever.
Using molecular detection, Blair et al. (2004) identified the presence of SFG agents in three locations in rural areas of northwestern Peru (department of Piura); SFG was found in four of 16 ticks of the species Amblyomma maculatum, Dermacentor (Anocentor) nitens, and Ixodes boliviensis. In this study, the rate of infection by genus Rickettsia agents in dog ticks from the metropolitan region of Piura Province in the Department of Piura, Peru was 0.2% (1/484).

The low prevalence (<2%) of rickettsia in epidemiological studies, even in regions considered endemic (SANGIONI et al., 2005), can be explained by the high pathogenicity of some species of Rickettsia in the vector, resulting in a low survival rate. In addition, the presence of vertebrate hosts with a degree of bacteremia sufficient to infect new susceptible ticks must be considered (SOARES et al., 2012).

Previous studies have demonstrated the efficacy of both dogs and other species as indicators of spotted fever agents (SANGIONI et al., 2005). Based on the results observed in the metropolitan region of Piura, it can be concluded that dogs may constitute an important indicator of the spotted fever agent; thus, future research should screen dogs for ticks.

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References


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