‘Candidatus Rickettsia asemboensis’ and Wolbachia spp. in Ctenocephalides felis and Pulex irritans fleas removed from dogs in Ecuador

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Abstract

Background: Flea-borne infections are distributed worldwide. Up to date there are no reports about microorganisms associated to fleas in Ecuador.

Methods: Seventy-one Pulex irritans and 8 Ctenocephalides felis fleas were removed from dogs in two Ecuadorian areas (Pastaza and Chimborazo Provinces) in December 2012. DNA extracts were tested by polymerase chain reaction (PCR) assays targeting universal 16S rRNA, as well as screened for the presence of Rickettsia spp. (gltA, htrA, ompB, sca4 and ompA genes) and Bartonella spp. (rpoB, gltA and ITS genes).

Results: Our results showed the presence of ‘Candidatus Rickettsia asemboensis’ (highly similar to R. felis) in C. felis and Wolbachia spp. endosymbionts in P. irritans collected from animals in Ecuador. No fleas were found to be positive for any Bartonella species or Yersinia pestis.

Conclusions: Clinicians should be aware of the potential risk of this new Candidatus Rickettsia sp. and keep in mind other flea-borne infections since these flea species frequently bite humans.

Keywords: Fleas, Pulex irritans, Ctenocephalides felis, Ecuador, ‘Candidatus Rickettsia asemboensis’, Wolbachia spp., Bartonella spp., Yersinia pestis, Plague

Background

Flea-borne diseases are worldwide-distributed emerging and re-emerging infections. Among them, plague, which is caused by Yersinia pestis, is the most severe human infection transmitted by fleas [1]. In South America, permanent plague foci exist among native rodent and flea populations in Bolivia, Brazil, Ecuador and Peru [2]. Rats have been the responsible hosts and from them, the disease has spread to other rodents. Ecuador is considered a plague ‘hot-spot’ since its introduction in 1908, and has experienced important outbreaks. Chimborazo Province has historically been a highly endemic area and the last fatal Ecuadorian cases of plague were reported there in 2004 [3,4].

In addition, fleas are vectors of murine typhus (caused by Rickettsia typhi), flea-borne spotted fever (caused by Rickettsia felis) and harbour Bartonella spp. [1,5-7]. Recent evidence of murine typhus in Ecuador is lacking, but the disease may be endemic in localities where commensal rodents (Rattus spp.) are abundant. To the best of our knowledge, data about distribution of R. felis in this country are unknown and there are no reports describing Bartonella spp. in fleas from Ecuador. For these reasons, our interest was focused on the study of flea-borne agents in two Ecuadorian areas (one of them where the last plague outbreak occurred) using molecular biological methods [polymerase chain reaction (PCR) and DNA sequencing].

Methods

In December 2012, a total of 79 fleas were removed from dogs by members of Red Iberoamericana para la Investigación y Control de las Enfermedades Rickettsiales, Programa Iberoamericano de Ciencia y Tecnologías para el Desarrollo (RIICER, CYTED; no. 210RT0403). Fifty-two specimens (44 Pulex irritans and 8 Ctenocephalides felis)
were collected in the Estación Experimental Fátima (ESPOCH), Cantón Puyo, in Pastaza Province (01°24’34.6”S; 77°59’57.5”W), and 27 specimens (all P. irritans) were collected in Cantón Guamote, in Chimborazo Province (02°00’25.0”S; 78°47’09.3”W). The study areas had altitudes of 1,034 m and 3,657 m, respectively. After identification at the species level, samples were kept in 70% ethanol at room temperature before being tested. DNA of each arthropod was extracted by lysis with 0.7 M ammonium hydroxide and tested by PCR with the universal primers fD1 and rp2 [8]. This primer pair amplifies the main part of the 16S rRNA gene and has been used for the identification of Y. pestis as the causative agent of plague in India [9]. Samples were also screened for the presence of Rickettsia spp. with PCR assays targeting rickettsial citrate synthase (gltA) and 17 kDa antigen (htrA) genes [10,11]. In accordance with the taxonomic scheme [12], additional rickettsial genes (ompB, sca4 and ompA) were tested to properly identify Rickettsia-positive specimens [13-17]. Moreover, Bartonella spp. was tested using RNA polymerase β-subunit–encoding gene (rpoB), gltA and intergenic spacer region gene (ITS) PCR primers, which amplify fragments of Bartonella genes [18-20].

Each PCR included positive controls consisting of Bartonella henselae DNA extracted from a cat flea (C. felis) from La Rioja - Spain, or Rickettsia slovaca strain S14ab DNA (obtained from Vero cells inoculated in our facility with a Dermacentor marginatus tick from La Rioja - Spain, and known to be infected with R. slovaca). Negative controls (DNA-free water) were included in all assays. Sequences generated by each pair of primers were then compared with those in GenBank using BLAST (www.ncbi.nlm.nih.gov/blast/Blast.cgi).

Results and discussion

PCR assays using universal eubacterial primers for 16S rRNA gene yielded amplicons of different intensity for 69 out of 79 fleas (8/8 C. felis and 35/44 P. irritans from Pastaza and 26/27 P. irritans from Chimborazo). All C. felis (n = 8) were also found to be infected with Rickettsia species using gltA and htrA as rickettsial PCR targets, whereas no evidence of Rickettsia spp. was found in P. irritans. Moreover, no sample was positive for Bartonella species as determined either by rpoB, gltA or ITS PCR assays. Positive and negative controls worked as expected in all cases.

Sequences of rickettsial 16S rRNA gene obtained from 7/8 C. felis (1303–1373 bp) showed the closest identity (99.6-99.9%) with ‘Candidatus Rickettsia asemboensis’, a potentially new Rickettsia species according to the established criteria [12,21] (Table 1). In these samples, the percentage of identity with 16S rRNA gene of a validly published Rickettsia species reached 99.4% for R. felis (accession no. NR074483). The 1387 bp-long sequence of 16S rRNA gene obtained from the remaining C. felis had the highest identity (97.1%) with a sequence from uncleaved flea-associated bacterium [22], and showed 93.6% identity with Snodgrassella alvi, betaproteobacteria classified in the family Neisseriaceae and previously isolated from the bee gut [23] (Table 1).

Sequencing of gltA fragments (350 bp) identified R. felis-like rickettsiaeae in C. felis (100% identity with Rickettsia sp. genotype RF2125) [24] and showed 99.7% identity (349/350 bp) with ‘Candidatus R. asemboensis’ (Table 1). The sequences of the 17-kDa amplicons (394 bp) were homologous (100% identical) to each other and to ‘Ca. R. asemboensis’ as well as to other molecular isolates included in the R. felis-like genotype group (Rickettsia sp. SE313 detected in Echidnophaga gallinacea from Egypt and Rickettsia sp. cf1and5 detected in C. felis from USA) (Table 1).

The sequences of ompB and sca4 amplicons (464 and 352 bp, respectively) were also 100% identical to ‘Ca. R. asemboensis’ (Table 1). Unfortunately, ompA PCR primers did not yield amplicons of the expected size, and inconclusive sequences were obtained for this target gene.

Percentages of identity with validated species R. felis (accession no. CP000053) were 98.6, 96.2, 97.9 and 96.6% for gltA, htrA, ompB and sca4 genes, respectively.

In addition, 59 out of 61 sequences of 16S rRNA gene obtained from 61 P. irritans specimens provided evidence of the presence of probable endosymbionts similar to those found within other arthropods and belonging to the genus Wolbachia (Table 2). Unfortunately, in two cases (corresponding to two P. irritans samples from Chimborazo) it was not possible to get a good-quality sequence to identify the bacteria.

In our study ‘Candidatus R. asemboensis’ has been found in fleas that bite humans (C. felis) removed from dogs in Ecuador. This potential new species was previously detected in Ctenocephalides canis from Kenya and whether it is a human pathogen remains unknown [21]. ‘Ca. R. asemboensis’ is highly similar to Rickettsia RF2125, a member of the R. felis-like genotype group that circulates in fleas from Brazil and other South American countries [26]. Nevertheless, in the Pastaza province (one of our sampling areas) cases of acute undifferentiated febrile illness compatible with rickettsioses have been reported [27].

Up to the present study, the presence of R. felis and/or Bartonella spp. has not been demonstrated in fleas from Ecuador. However, R. felis has been found in South American fleas from Brazil, Peru, Uruguay, Chile, Argentina and Colombia [11,25,28-31]. In addition, human
infection with *R. felis* in South America has been confirmed in Brazil by molecular methods [32], and human serological evidence of *R. felis* infection has been recently reported in Colombia [33]. Moreover, there are limited reports describing *Bartonella* spp. in fleas from South America. A molecular study conducted in a *Pulex* specimen found on a Peruvian person evidenced the presence of a potential new *Bartonella* species [5]. Years later, our research group detected *B. rochalimae, B. clarridgeiae,* and *B. henselae* in *P. irritans* and *C. felis* collected from cats and dogs in Chile, suggesting the role of fleas as possible vectors of *Bartonella* spp. [7].

Lastly, *Wolbachia* spp. are alphaproteobacteria included in the family *Anaplasmataceae* that were first detected in fleas in 2000 [34]. In this study, the detection rate of *Wolbachia* spp. in *P. irritans* was 83% (59/71). On the contrary, no *C. felis* analysed (0/8) showed evidence of carriage of *Wolbachia* endosymbionts. Previous studies had identified *Wolbachia* in around 20% of cat fleas [35,36]. It has been suggested that *R. felis* infection in fleas might diminish the richness of flea microbiota [37].

According to our data, there is a strong association of *C. felis* with ‘Ca. *R. asemboensis*’ whereas *P. irritans* is associated with *Wolbachia* spp. Nevertheless, the interaction of *Wolbachia* with *R. felis* or other related species, such as ‘Ca. *R. asemboensis*’ in fleas needs further investigation.

Based on 16S rDNA analysis, the presence of *Y. pestis* DNA has not been demonstrated in our fleas despite *P. irritans* having been previously described as vectors of *Y. pestis* in Ecuador [2] and plague outbreaks have been repeatedly reported in the Chimborazo region [3,4], where some flea specimens were collected. The rodent flea *Xenopsilla cheopis*, which is the main vector, does not exist in the inter-Andean region of Ecuador (2,500-4,000 m above sea level) possibly, due to very sudden changes in the climatic conditions [38].

**Conclusions**

In summary, our result confirms the presence of ‘*Candidatus* *R. asemboensis*’ and *Wolbachia* spp. in fleas removed from dogs in Ecuador. Clinicians should be aware of the potential risk of this new *Candidatus Rickettsia* sp. and keep in mind other flea-borne infections in areas where humans are exposed to fleas.

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### Table 1 Results of nucleotide sequence analysis corresponding to PCR products amplified from the 8 *Ctenocephalides felis* specimens of this study

| Gene* | Flea gender | Location | Length of sequence (bp) | % identity | Bacteria with closest identity in the BLAST search | GenBank accession no. |
|-------|-------------|----------|-------------------------|------------|--------------------------------------------------|-----------------------|
| 16S rRNA | 1/2 M; 6/6 F | Pastaza | 1303-1373 | 99.6-99.9 | ‘Candidatus Rickettsia asemboensis’ (Rickettsia F30) | JN315967 |
|         | 1/2 M; 0/6 F | Pastaza | 1387 | 97.1 | Uncultured flea-associated bacterium | EU137419 |
|         |             |         |             | 93.6 | *Snodgrassella alvi* | JQ746645 |
| gilA   | 2/2 M; 6/6 F | Pastaza | 350 | 100 | *Rickettsia felis*-like (Rickettsia RF2125) | AF516333 |
|         |             |         |             | 99.7 | ‘Candidatus Rickettsia asemboensis’ (Rickettsia F30) | JN315968 |
| htrA   | 2/2 M; 6/6 F | Pastaza | 394 | 100 | ‘Candidatus Rickettsia asemboensis’ (Rickettsia F30) | JN315969/DQ166937/AY953286 |
| ompB   | 2/2 M; 6/6 F | Pastaza | 464 | 100 | ‘Candidatus Rickettsia asemboensis’ (Rickettsia F30) | JN315972 |
| sco4   | 2/2 M; 6/6 F | Pastaza | 352 | 100 | ‘Candidatus Rickettsia asemboensis’ (Rickettsia F30) | JN315970 |

*Inconclusive sequences were obtained for ompA gene.

1 Number of positive specimens / Total number of specimens from each flea gender; M: Male; F: Female.

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### Table 2 Results of nucleotide sequence analysis corresponding to 59 PCR products amplified from the 71 *Pulex irritans* of this study (44 specimens from Pastaza and 27 from Chimborazo)

| Gene | Flea gender | Location | Length of sequence (bp) | % identity | Bacteria with closest identity in the BLAST search | GenBank accession no. |
|------|-------------|----------|-------------------------|------------|--------------------------------------------------|-----------------------|
| 16S rRNA | 9/16 M; 10/28 F | Pastaza | 1287-1386 | 98.3-100 | *Wolbachia* sp. wRI endosymbiont of *Drosophila simulans* | NR074437 |
|         | 11/13 M; 13/14 F | Chimborazo |             |         |                                                   |                       |
|         | 4/16 M; 2/28 F | Pastaza | 1338-1383 | 98.4-98.7 | *Wolbachia* sp. endosymbiont of *Pseudopolyphaga cananiensis* | DQ115538 |
|         | 0/16 M; 2/28 F | Pastaza | 1300-1372 | 98.3-99.3 | *Wolbachia* sp. endosymbiont of *Gryllus crickets* | U83094 |
|         | 0/16 M; 1/28 F | Pastaza | 1305 | 99.3 | *Wolbachia* sp. endosymbiont of *Gryllus ovipops* | U83093 |
|         | 0/16 M; 1/28 F | Pastaza | 1374 | 98.3 | *Wolbachia* sp. endosymbiont of *Curculio hachijoensis* | AB746399 |
|         | 0/16 M; 6/28 F | Pastaza | 1291-1368 | 98.3-99 | *Wolbachia* sp. endosymbiont of *Kleidocerys resedae* | JQ726770 |

*In two cases we did not obtain enough good-quality sequences to identify the bacteria, and we did not obtain amplicons for ten specimens.

1 Number of positive specimens / Total number of specimens from each flea gender; M: Male; F: Female.
Competing interests
The authors declare they have no competing interests.

Authors’ contribution
Designed the study: JMV, MBL, JAO, JZC. Collected and identified fleas: JMV, MBL, JAO, FP, JZC. Processed samples and analyzed sequences: AP. Analyzed the data: AP, JAO. Wrote the paper: AP, JAO. All authors read and approved the final version of the manuscript.

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