Evidence of *Ehrlichia chaffeensis* in Argentina through molecular detection in marsh deer (*Blastocerus dichotomus*)

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**A B S T R A C T**

Vector-borne pathogens are responsible for serious emerging diseases and have been widely described in wildlife. *Ehrlichia chaffeensis* causes the zoonotic “monocytic ehrlichiosis” in humans, is transmitted by the tick *Amblyomma americanum* and its reservoir host is the white-tailed deer (*Odocoileus virginianus*) in North America. Little is known about the native reservoir and the tick vectors involved in the transmission cycle in South America. We report here the detection of *E. chaffeensis* in a study on marsh deer (*Blastocerus dichotomus*) mortality in Argentina, in different time periods between 2007 and 2016. Four deer, from two distinct populations, were positive for *E. chaffeensis* through molecular methods. Additionally, the variable-length PCR target (VLPT) region of positive samples was genotyped. Our results provide the first evidence of *E. chaffeensis* in autochthonous *Cervidae* from Argentina, contributing to uncover the distribution of this tick-borne infection in South America.

1. Introduction

Within the emerging tick-borne infections, ehrlichioses are potentially zoonotic diseases that stand out because of the multiple vertebrate hosts and hard tick vectors that are involved in their transmission cycle. Like other tick-borne diseases, ehrlichioses are strongly influenced by climate change because it may alter the density, activity patterns and geographic distribution of arthropod and hosts involved (Randolph, 2010). The genus *Ehrlichia* comprises Gram-negative intracellular bacteria from the *Anaplasmataceae* family, which are transmitted by hard ticks (Dumler et al., 2001). This genus currently includes recognized species (*E. canis*, *E. chaffeensis*, *E. ewingii*, *E. muris*, *E. ovis*, *E. ruminantium*) and potentially new species under characterization (Tate et al., 2013; Pritt et al., 2017; Yang et al., 2017).

*Ehrlichia chaffeensis* is responsible for the symptomatic monocytic ehrlichiosis in humans (Paddock and Childs, 2003). In the United States *Amblyomma americanum* ticks are the transmitting vectors of *E. chaffeensis* and the white-tailed deer (*Odocoileus virginianus*) is the main reservoir host (Ewing et al., 1995), while humans and dogs are incidental hosts. Bovines are susceptible to experimental infection with *E. chaffeensis* (Delos Santos et al., 2008).

In Argentina Tomassone et al. (2008) detected *E. chaffeensis* in three developmental stages of *Amblyomma parvum* tick through molecular methods and Ripoll et al. (1999) reported serological evidence of *E. chaffeensis* in humans in the north-west region of the country. The lack of simultaneous evidence of *E. chaffeensis* in mammalian host and arthropods has prevented the recognition of the local species capable of maintaining and transmitting the pathogen respectively. In this study, we analyzed the occurrence of *E. chaffeensis* in marsh deer (*Blastocerus dichotomus*) that inhabit the Paraná River Delta and Iberá Wetlands in Argentina.

2. Materials and methods

Samples were collected during mortality events of marsh deer in 2007, 2015 and 2016 in two different populations located along the alluvial plain of the Paraná River, in Argentina: Paraná River Delta and Iberá Wetlands. The Paraná River Delta region, located between 60° 39′ S and 58° 30′ W, 32° 30′ S and 58° 30′ W, 34° 30′ S, is a complex floodplain covered by herbaceous vegetation on frequently flooded area, shrub savannas.
and forests highly disturbed by afforestation, cattle rearing and fishing (Malvárez, 1999, Iriondo, 2004) (Fig. 1, Parana River Delta). The Ibera Wetlands, which lay on the paleolithic-river beds of the Parana River between 56° 25′ W, 27° 30′ S and 58° W 29° S in Corrientes province, is about 800 km North to Parana River Delta and rather pristine. (Fig. 1, Ibera Wetlands). It is a temporarily and permanently waterlogged area, combined with forests, scrublands, grasslands, pastures, lakes, wetlands and peat lands. Remarkably, both areas have no hydrographic connection between each other.

Complete necropsies of succumbed marsh deer were performed. All deer showed cachexia and signs of anemia and were clearly weakened (Orozco et al., 2013). Full blood samples from the heart of recently succumbed animals (within the hour) were collected and stored with EDTA: a 2-ml aliquot was refrigerated at 4 °C and 1 ml was frozen at −20 °C. Right afterwards, the succumbed deer was examined for ticks, which were removed and stored in 70% ethanol. Tick examination was conducted by visual inspection on predilection sites (face, ears, brisket, withers, knees, perineum region and tail). The tick burden was roughly estimated only in recently dead marsh deer. Biosafety procedures for animal management were performed according to protocols approved by the Argentinian CICUAL (Institutional Committee for the Care and Use of Experimental Animals; Protocol N° 2014-40, issued by the Facultad de Ciencias Veterinarias, Universidad de Buenos Aires). Wildlife permits (including transit permits for biological samples) were obtained from the provincial government through “Natural Resources Agency of Corrientes” (Disposition N° 845, Proceeding N° 193-15-09-410/2014) and the General Directorate for Natural Resources of Entre Ríos (Authorization N° 007/16/Resolution 1721/14 D.G.R., Proceeding N°, 1845661).

The ticks were taxonomically identified (Guglielmone and Viñabal, 1994; Nava et al., 2014) under a stereoscopic magnifier (10X-40X) before processing. DNA was extracted by the phenol/chloroform method followed by a standard ethanol precipitation whether from crushed ticks using liquid nitrogen or 400 μl of whole blood (Halos et al., 2004). DNA quality and concentration were determined using a micro-volume spectrophotometer (NanoDrop ND-1000. ThermoFisher Scientific).

Detection of *E. chaffeensis* DNA from the blood samples and ticks was conducted with genus- and species-specific primers. Initially, all samples were screened using a PCR protocol targeting the 16S rRNA gene common to *Ehrlichia* spp. and *Anaplasma* spp. (Bekker et al., 2002). Subsequently, positive samples were also tested employing a specific 16S rRNA *E. chaffeensis* Forward primer (Anderson et al., 1992) together with the Reverse primer employed previously (Bekker et al., 2002). Lastly, the variable-length PCR target (VLPT) region of positive samples was genotyped (Sumner et al., 1999).

The PCRs were performed in a 50 μl reaction mixture (0.4 μmol of each primer, 0.2 mM of each deoxyribonucleotide triphosphate, 1.25 U of GoTaq DNA polymerase - Promega Madison, Wi. USA, 10 μl of 5x PCR buffer and purified water for 50 μl of final volume) using 200 ng of
genomic DNA (both for blood and tick samples). Amplification was carried out in a thermocycler (Bio-Rad MyCycler Thermal Cycler) under specific cycling conditions (Anderson et al., 1992; Sumner et al., 1999; Bekker et al., 2002). For each amplification reaction, positive (DNA from cultured *E. chaffeensis* Arkansas strain) and negative (pure water) controls were included. An aliquot of 5 μl of each amplified product was analyzed by electrophoresis in 1% agarose gel stained with ethidium bromide. A molecular size marker (1 Kb Plus DNA Ladder, Invitrogen) was used to determine PCR product size.

Both strands from 16S rRNA *Ehrlichia/Anaplasma* and VLPT fragments were sequenced with a Big Dye Terminator v3.1 kit from Applied Biosystems and analyzed on an ABI 3130XL genetic analyzer from the same supplier (Genomic Unit, Consorcio Argentino de Tecnología Genómica (CATG), Instituto de Biotecnología, CICVyA, INTA).

Raw files from target regions were processed using the Vector NTI Advanced 10 program (Invitrogen). For each fragment, forward and reverse chromatograms (ab1 files) were used for assembling and the outcome contig (FASTA file) was used for further sequence analysis.

### 3. Results

In this study, we assessed the *E. chaffeensis* presence in blood samples from *B. dichotomus* (n = 38) as well as in ticks *Amblyomma triste* (n = 7) and *Rhipicephalus microplus* (n = 173) from 7 succumbed deer (Table 1).

From the 38 tested deer blood samples, 4 were positive to the generic 16S rRNA (KY644143-KY644146) and the *E. chaffeensis*-specific PCR reaction. Five adult *R. microplus* from a positive deer (CP_C7) tested positive for *E. chaffeensis* 16SrRNA. The remaining 168 *R. microplus* and *A. triste* (n = 7) tested negative for generic 16SrRNA (Fig. 2). Although the PCR reaction rendered the expected size for the 16SrRNA target region, only one of the positive tick samples contained enough DNA for sequencing (KY644147). The obtained 16S rRNA 410 bp fragment, which hold the hypervariable V1 region, showed high identity values with *E. chaffeensis* strains deposited in GenBank (Table 2). The V1 region alignment strengthened species identification (Fig. 2, bottom).

To further characterize the positive samples, we then amplified the variable-length PCR target (VLPT). This assay yielded an amplicon for two samples (blood from CP_C2 and CP_C7 from Ibera) (Fig. 2). We subsequently compared the protein sequences retrieved for the VLPT gene (KY652924 - KY652925) against two previously reported polymorphic regions (EU826517 and EU826518) amplified from ticks collected in Argentina (Tomassone et al., 2008) and the *E. chaffeensis* strain Arkansas (WP011452439.1). The new target sequences revealed a novel genotype that was identical for both samples but different from the formerly reported sequences (Table 3).

The *E. chaffeensis* positive blood samples were collected in 2007, 2015 and 2016 from three adult deer (CP_C2, CP_C7 and CP_C12) and one newborn deer (CP_11). The adult deer died during a mortality event in Ibera Wetlands and had high *R. microplus* tick infestation (more than 50 ticks grouped in one or more predilection sites), with cutaneous lesions and exudation of tissue fluid, especially on external ears and face (Fig. 3). The deer CP_D21 was a newborn deer dead by starvation after illegal hunters killed his mother during the floods of 2016 in the Paraná River Delta. No ticks were detected on CP_D11 (Fig. 2).

### 4. Discussion

In our study, *E. chaffeensis* DNA was detected in blood samples from two geographically distinct marsh deer population during mortality events at different time points in 2007, 2015 and 2016, providing the first evidence of *E. chaffeensis* presence in a mammalian host in Argentina. *E. chaffeensis* has also been reported in some Brazilian Paraná river marsh deer population (Fig. 1) (Machado et al., 2006; Sacchi et al., 2012), but we could not use the sequences for comparison (DQ345720.1 and JQ085940.1) as the authors targeted a different 16S rRNA fragment that does not contain the V1species specific region (Dumler et al., 2001).

Although in South America *E. chaffeensis* can be present in ticks from domestic and wild mammals (Tomassone et al., 2008), little is known about the native mammalian hosts and vectors involved in its transmission cycle. White-tailed deer are the most common host for the three mobile stages of *A. americanum* (Yabsley, 2010) in USA. The exposure to the agent appears to occur early in the life in this host species (Paddock and Yabsley, 2007) and ticks can become infected by feeding on animals with a persistent bacteremia without apparent clinical signs of disease (Davison et al., 2001). In North America, *E. chaffeensis* has been also found in *A. maculatum*, *Dermacentor variabilis* and *Ixodes pacificus* among others (Paddock and Yabsley, 2007).

We found *E. chaffeensis* DNA in the main *B. dichotomus* populations in Argentina, where *A. triste* is endemic. Even though *E. chaffeensis* was apparently absent from the collected *A. triste* (n = 7), we expect that the agent will be detected if a large sample is analyzed. Ticks feeding on infected deer could be acquiring the organism in a blood meal but not necessarily transmitting or maintaining the infection. Whereas whole full engorged *R. microplus* from infected marsh deer were positive for *E. chaffeensis* DNA, further studies should be performed so as to appraise for its vector competence.

From an epidemiological point of view, correlation between samples and geographic origin together with variations in gene sequences under immunological selection are used as markers for tracking infection cycles (Sumner et al., 1999). For revealing the intraspecific variation of circulating isolates, we sequenced the VLPT gene fragment, a size variation gene resulting from loss or gain of long of direct repeats (Yabsley, 2010) in USA. The ex-posure to the agent appears to occur early in the life in this host species (Paddock and Yabsley, 2007) and ticks can become infected by feeding on animals with a persistent bacteremia without apparent clinical signs of disease (Davison et al., 2001). In North America, *E. chaffeensis* has been also found in *A. maculatum*, *Dermacentor variabilis* and *Ixodes pacificus* among others (Paddock and Yabsley, 2007).

The findings of *E. chaffeensis* in *B. dichotomus*, during independent mortality events over time and in two geographic areas, suggest that *E. chaffeensis* is circulating in marsh deer populations from Argentina. Previous reports of *E. chaffeensis* in *A. parvum* ticks, serological evidence in humans and our present findings provide valuable information to understand *E. chaffeensis* epidemiology in Argentina. Forthcoming

### 5. Conclusions

The findings of *E. chaffeensis* in *B. dichotomus*, during independent mortality events over time and in two geographic areas, suggest that *E. chaffeensis* is circulating in marsh deer populations from Argentina.

### Table 1

| Samples obtained          | Number of marsh deer | Total |
|---------------------------|----------------------|-------|
|                           | Ibera Wetlands CP_C  |       |
| Deer blood                | 17                   | 21    |
| *Rhipicephalus microplus* | 5 (n = 173)          | 0     |
| *Amblyomma triste* ticks  | 1 (n = 2)            | 2 (n = 5) |
|                           | 21                   | 5 (n = 173) |
|                           | 2 (n = 5)            |       |
|                           | 3 (n = 7)            |       |
studies should comprise larger sampling, including a wide range of hosts and vectors in order to determine the extent of the *E. chaffeensis* in the region.

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**Table 2**

Results of the global nucleotide alignment of deer sample sequences in the present study.

| Deer Sample | Percent identity to closest match (E. chaffeensis str. West Paces) |
|-------------|-------------------------------------------------------------------|
| CP_C2 Blood (KY644143) | 97% |
| CP_C7 Blood (KY644144) | 99% |
| R. microplus tick (KY644147) | 99% |
| CP_D11 Blood (KY644145) | 97% |
| CP_C12 Blood (KY644146) | 99% |

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**Table 3**

Aminoacid sequence of VLPT tandem repeats and profiles for previously reported *E. chaffeensis* in Argentina (EU826517 and EU826518), CP_2 and CP_7 deer samples and the *E. chaffeensis* positive control Arkansas strain (Reference).

| Repeat sequence | Sample Name |
|-----------------|-------------|
| SDSHESSHTVPNLSEEVQVESLQQS | EU826517 EU826518 CP_C2 CP_C7 Arkansas strain |
| SDFDQGFSVELDPFKEAVQLGNDLQQHS | ✓ ✓ ✓ ✓ |
| SNSDYRSSVPVELPGSKEEVQVESAVQP | ✓ ✓ ✓ ✓ |
| SDSLHGPSPHELPSAEVEVMQELDDLQQQ | ✓ ✓ ✓ ✓ |
| SDFDQGFSVELPSKESQEVQLESATKN Vy | ✓ ✓ ✓ ✓ |
| SDSLHGPSPHELPSAEVEVMQELDDLQQQ | ✓ ✓ ✓ ✓ |
| SFGLHSSSVELPSKESQEVQLESATKN Vy | ✓ ✓ ✓ ✓ |
| SDFDQGFSVELDPFKEAVQLGNDLQQQS | ✓ ✓ ✓ ✓ |
| SDSLHGSVELPQEEQVESLQLQGQSN | ✓ ✓ ✓ ✓ |
| SDSLHGSVELPQEEQVESLQLQGQSN | ✓ ✓ ✓ ✓ |
| SDSLHGSVELPQEEQVESLQLQGQSN | ✓ ✓ ✓ ✓ |

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