Molecular detection of *Bartonella*, *Ehrlichia* and *Mycoplasma* in feral dogs of El Pedregal de San Angel Ecological Reserve in Mexico City

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Abstract

Free-ranging and feral dogs represent a group of unattended companion animals. They impact wild animal populations by predating native species, displacing predators and introducing exotic pathogens. The aim of this work was to describe the molecular occurrence of *Rickettsia*, *Ehrlichia*, *Anaplasma*, *Mycoplasma* and *Bartonella* in feral dogs. The study was carried out in the last relict of a protected area in Mexico City. Blood clots samples from 19 dogs were obtained and analyzed for detection of specific fragments of the 16S-rRNA gene for *Anaplasma*, *Ehrlichia* and *Mycoplasma* and citrate synthase (*gltA*) for *Bartonella* and *Rickettsia*. Our results showed that DNA from three bacteria species (*Bartonella vinsonii* subsp. *berkhoffii*, *Ehrlichia canis* and *Mycoplasma haemocanis*) was present with frequencies ranging from 5.3 to 15.8%. This is the first record of *B. vinsonii* subsp. *berkhoffii* and *M. haemocanis* in dogs from México, and also the first finding of *Ehrlichia canis* in Mexico City. It is important to perform surveillance of feral dog populations in order to identify the impact of these pathogens on wild animal populations and Public Health in order to establish prevention and protection programs.

Keywords: Canis familiaris, flea-borne pathogens, Mexico, tick-borne pathogens.

Resumo

Cães errantes e selvagens representam um grupo de animais de companhia livres. Eles impactam as populações de animais selvagens pela predação de espécies nativas, deslocando predadores e introduzindo patógenos exóticos. O objetivo deste trabalho foi descrever a ocorrência molecular de *Rickettsia*, *Ehrlichia*, *Anaplasma*, *Mycoplasma* e *Bartonella* em cães selvagens. O estudo foi realizado no último ecossistema de uma área protegida na Cidade do México. Amostras de coágulos sanguíneos de 19 cães foram obtidas e analisadas para detecção de segmentos específicos do gene 16S-rRNA para *Anaplasma*, *Ehrlichia* e *Mycoplasma* e síntese de citrato (*gltA*) para *Bartonella* e *Rickettsia*. Nosso resultado mostrou que DNA de três espécies de bactérias (*Bartonella vinsonii* subsp. *berkhoffii*, *Ehrlichia canis* e *Mycoplasma haemocanis*) estava presente com frequências variando de 5,3 a 15,8%. Isso é o primeiro registro de *B. vinsonii* subsp. *berkhoffii* e *M. haemocanis* em cães de México, e também a primeira descrição de *Ehrlichia canis* na Cidade do México. É importante realizar a vigilância das populações de cães selvagens para identificar o impacto desses patógenos nas populações de animais silvestres e na Saúde Pública, a fim de estabelecer programas de prevenção e proteção.

Palavras-chave: Canis familiaris, patógenos transmitidos por pulgas, México, patógenos transmitidos por carrapatos.

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Introduction

Invasive mammalian predators, such as feral dogs, are arguably the most damaging group of exotic species for global biodiversity (DOHERTY et al., 2016). They can live in extreme conditions in human-dominated ecosystems, restructuring food webs and endangering native fauna (SUZÁN & CEBALLOS, 2005). Dogs display a wide range of social organization, from solitary living in human homes to living in packs as free-ranging and feral dogs (PAUL et al., 2014). Free-roaming dogs (those that are not permanently under human control) are thought to account for about 75% of the global dog population. Dogs requiring no human contact (often described as feral) (HUGHES & MACDONALD, 2013) have been considered as potential reservoirs for zoonotic pathogens, a significant public health concern worldwide (CRUZ-REYES, 2009; DEPLAZES et al., 2011; CHEN et al., 2012; FANG et al., 2015).

El Pedregal de San Angel Ecological Reserve [REPSA after its Spanish acronym] produced by the Xitle Volcano eruption approximately 1670 years ago, is located inside the Universidad Nacional Autónoma de México [UNAM]. This reserve covers 237.3 ha, and is inhabited by least 1,821 species of different tax, including 130 birds, 20 reptiles and 33 mammals among others (CASTILLO-ARGUERO et al., 2007; ESTAÑOL-TECUATL & CANO-SANTANA, 2017). Buildings and wide avenues are surrounding this reserve and it has been constantly affected by irregular garbage deposition and invasion of feral fauna, among other problems (MARTÍNEZ-OREA et al., 2012). With respect to the feral fauna, it has been particularly worrisome with the presence of antibody titers in dogs against rabies, Toxoplasma gondii, and parvovirus (SUZÁN & CEBALLOS, 2005), suggesting an important risk for public and ecosystem health. However, no studies have been conducted to evaluate the presence of tick-borne and flea-borne pathogens in feral dogs, although previous studies have shown the presence of Bartonella henselae in feral dogs from UK using serological techniques (BARNES et al., 2000), and Ehrlichia canis from Costa Rica using serological and molecular techniques (BARRANTES-GONZÁLEZ et al., 2016).

For this reason, the aims of this study were to perform the molecular detection and characterization of five bacterial pathogens (Anaplasma, Bartonella, Ehrlichia, Mycoplasma and Rickettsia) in feral dogs of El Pedregal de San Angel Ecological Reserve in Mexico City.

Materials and Methods

Study site

The study was conducted in El Pedregal de San Angel Ecological Reserve [REPSA] in Mexico City (N19°18’31”–19°19’17”, W99°10’20”–99°11’52”), located in southern Mexico City, within the Universidad Nacional Autónoma de México [UNAM]. This reserve consists of 237.3 ha divided into three core zones and 13 buffer zones (171 ha correspond to the core zones and 66 ha to buffer zones) (Figure 1A) that are under long-term disturbance (fragmentation and isolation) (SUZÁN & CEBALLOS, 2005). The average annual temperature is 15°C and 700-900 mm of rainfall.

Figure 1. (A) Study area (southern portion of Mexico City) showing the sighting of feral dogs in several parts of El Pedregal de San Angel Ecological Reserve, Mexico; (B) Photographic records of feral dogs in which these animals are observed moving in different parts of El Pedregal de San Angel Ecological Reserve, Mexico.
Capture of dogs

The study was carried out in a period of three years, spanning from September 2015 to July 2018. Dogs were captured either inside the reserve or in the UNAM campus. For activity patterns, camera traps (Bushnell Trophy Cam HD) were used. Tomahawk traps (Tomahawk Live Trap Co., Tomahawk, Wisconsin, USA) and collar traps (Collarum, Wildlife Control Supplies, East Granby, CT) baited with meat, chicken or dog food, were activated late in the afternoon between 17:00-19:00 h, and checked every two hours during the night until dawn. Dogs that were not captured with the traps were sedated using a dart gun. A fetch pole was used for restraining dogs prior to sedation. The capture methods used tried to avoid unnecessary stress of the animals. Chemical immobilization was performed with a combination of ketamine hydrochloride (10 mg/kg) and xylazine/HCl (2 mg/kg) (Wildlife Pharmaceutics Mexico SA de CV 04930, Mexico). Dog captured was subject to a clinical evaluation where the age (juveniles or adults) and sex were also estimated. Age was determined based on tooth wear. Dogs were euthanized with an overdose of sodium pentobarbital (60 mg/kg) (PLUMB, 2011) (Salud y Bienestar Animal, 03310, Mexico) and referred to the invasive species control area of the Faculty of Veterinary-UNAM. The procedures were conducted after approval from Ministry of Environment and Natural Resources [SEMARNAT] permit: SGPA/DGVS/03670/2015 and SGPA/DGVS/005615/18: Authorization for management and control of exotic species (dogs and cats).

Sample collection

As a part of an ongoing project to identify the exposure of feral dogs to viral agents and *Leptospira* sp., serum samples of the animals were recovered and were used to screen for the presence of several flea-borne and tick-borne bacteria (MYLONAKIS et al., 2009). Blood samples were collected after immobilization and sedation by jugular or cephalic venipuncture in BD Vacutainer blood collection tubes (10-15 ml). Samples were preserved at 4°C until needed. The serum was extracted at 3,500 rpm for 15 minutes and the blood clots were recovered and stored frozen at -20°C before testing.

DNA extraction

Five hundred microliters of a 10% Cheelex-100 solution and 20 μL of proteinase K were added to each blood clot sample, incubated at 56°C in a dry bath for one hour. The temperature was increased to 100°C for 10 minutes to make the protein denaturation more efficient. The samples were then centrifuged at 14,000 rpm for 15 min, and the supernatant was collected in new tubes and storage at -20°C until further use (BALLADOS-GONZÁLEZ et al., 2018). The amount (in ng/μL) and quality (260/280 ratio) of DNA was measure using a Nanodrop Spectrophotometer (Thermo Scientific™ NanoDrop™ 2000/2000c).

Molecular detection

As an internal control of extraction, we amplified a fragment of 620 bp of cytochrome oxidase subunit I gene (COI) of the dogs, using primers and conditions of Marsico et al. (2010). For bacterial pathogens detection, we amplified several fragments of the 16S-rRNA and citrate synthase genes using primers and conditions reported previously (Table 1). The reaction mixture was done in a final volume of 25 μL, using 12.5 μL of GoTaq® Green Master Mix, 2 μL of Promega Corporation (Madison, WI, USA), 2 μL of primers (2 μM, 1 μL each), 8.5 μL nuclease-free water, and 500 ng DNA (2 μL). A negative (a reaction mix with-out DNA template) and positive (*Anaplasma ovis* [MG733099], *Bartonella vinsonii vinsonii* [KT326174], *Mycoplasma ovis* [MG733068] and *Rickettsia amblyommatis* [KX363842] DNA) controls were included. PCR products were resolved into 2% agarose gels, using a 100 bp molecular weight marker (nucleic acid markers, Axygen) in 1×TAE buffer.

Sequencing procedures

All positive products were purified using the Agencourt AMPure PCR purification kit. Samples were amplified and sequenced using BigDye Terminator v3.1 Cycle Sequencing Kit (Thermo Fisher, Cat.4337455). Sequencing products were purified by BigDye XTerminator (Thermo Fisher, Cat. 4376486) prior to loading on the ABI 3730xL DNA Analyzer at Sequencing Unit of the National Institute of Genomic Medicine, Ministry of Health, Mexico.

<table>
<thead>
<tr>
<th>Pathogen</th>
<th>Gene</th>
<th>Primers</th>
<th>Size (bp)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Ehrlichia</em> spp./<em>Anaplasma</em> spp.</td>
<td>16S rDNA (RNA ribosomal 16S)</td>
<td>EHR01F, EHR02R</td>
<td>495</td>
<td>Murphy et al. (2017)</td>
</tr>
<tr>
<td><em>Mycoplasma</em> spp.</td>
<td>16S rDNA (RNA ribosomal 16S)</td>
<td>HemMycop16S-322s, HemMycop16S-1420as</td>
<td>1000</td>
<td>Maggi et al. (2013)</td>
</tr>
</tbody>
</table>
**Phylogenetic analysis**

Consensus sequences were constructed using the software Genious, and were aligned with those of reference of each bacterial group deposited in GenBank using Bioedit. Phylogenetic analysis was done using the maximum likelihood method and the best substitution model was selected for each alignment based on the lowest Bayesian Information Criterion (BIC) in Mega 6.0. Sequences generated in this study were deposited in GenBank under accession numbers MH917710- MH917715, and MN294708.

**Results**

Activity patterns were observed during day and night. A pack of feral dogs was detected moving across different parts of REPSA (core and buffer zones). Nineteen dogs were captured, 9 males (3 juvenile and 6 adults) and 10 females (4 young and 6 adults) (Figure 1B, Table 1). Thirteen dogs were captured inside the core zones, two in buffers zones and four dogs were captured outside the reserve. All samples were positive for the amplification of the COI gene (internal extraction control). From DNA extraction, we obtained a median concentration of DNA of 173.75 ng/μl, with a minimum of 22.90 ng/μl and a maximum of 730.60 ng/μl, with a 260/280 coefficient between 1.8 and 2.0. Six dogs (three females and three males) were positive for at least one vector-borne pathogen (Table 2). None of the juvenile dogs tested positive for any pathogen analyzed. Two females and one male were positive for *Bartonella* DNA; the three sequences exhibited an identity of 99% (343/344 bp) with sequences of *Bartonella vinsonii* subsp. *berkhoffii* detected in dogs and wild carnivores in US and northern Mexico (Accession numbers CP003124.1 and KT807806.1) (Figure 2A). Two more samples were positive for *Ehrlichia* DNA; one female and a male, whose sequences were identical to each other and exhibited a identity of 100% (341/341 bp) with sequences of *Ehrlichia canis* from dogs, also from the US and South of Mexico (Accession numbers NR_118741.1 and MH487666.1) (Figure 2B). A single sample tested positive for the presence of a hemotrophic *Mycoplasma*, and exhibit an identity of 100% (882/882 bp) with *Mycoplasma haemocanis* detected in dogs from Chile (Accession number KY117659.1) (Figure 2C). We did not detect the presence of *Anaplasma* or *Rickettsia* in any of the analyzed samples. The presence of coinfections was not detected in any of the blood samples (Table 1). The Maximum Likelihood analysis confirms the identity of the three pathogens, which form monophyletic groups with support values that go between 98-100 with sequences of *B. vinsonii* subsp. *berkhoffii*, *E. canis* and *M. haemocanis*, respectively (Figure 2).

<table>
<thead>
<tr>
<th>Code</th>
<th>Collection place</th>
<th>Sex</th>
<th>Age</th>
<th>Bartonella spp.</th>
<th>Ehrlichia spp.</th>
<th>Mycoplasma spp.</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>ZNO</td>
<td>Female</td>
<td>Adult</td>
<td>(-)</td>
<td>Ehrlichia canis</td>
<td>(-)</td>
</tr>
<tr>
<td>2</td>
<td>ZNO</td>
<td>Male</td>
<td>Adult</td>
<td>(-)</td>
<td>Ehrlichia canis</td>
<td>(-)</td>
</tr>
<tr>
<td>3</td>
<td>ZNSO</td>
<td>Female</td>
<td>Adult</td>
<td>(-)</td>
<td>Bartonella vinsonii berkhoffii 99% (343/344) CP003124.1</td>
<td>(-)</td>
</tr>
<tr>
<td>4</td>
<td>ZNSO</td>
<td>Male</td>
<td>Adult</td>
<td>Bartonella vinsonii berkhoffii 99% (346/349) CP003124.1</td>
<td>(-)</td>
<td></td>
</tr>
<tr>
<td>5</td>
<td>ZNSO</td>
<td>Female</td>
<td>Adult</td>
<td>Bartonella vinsonii berkhoffii 99% (346/349) CP003124.1</td>
<td>(-)</td>
<td></td>
</tr>
<tr>
<td>6</td>
<td>ZNO</td>
<td>Female</td>
<td>Adult</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>7</td>
<td>ZNP</td>
<td>Female</td>
<td>Adult</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>8</td>
<td>ZNO</td>
<td>Male</td>
<td>Juvenile</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>9</td>
<td>DGAE</td>
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<td>Adult</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>10</td>
<td>ZNO</td>
<td>Female</td>
<td>Juvenile</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>11</td>
<td>ZNO</td>
<td>Male</td>
<td>Juvenile</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>12</td>
<td>ZNO</td>
<td>Female</td>
<td>Juvenile</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>13</td>
<td>CU</td>
<td>Male</td>
<td>Adult</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>14</td>
<td>CU</td>
<td>Male</td>
<td>Adult</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>15</td>
<td>MUAC</td>
<td>Female</td>
<td>Juvenile</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>16</td>
<td>CU</td>
<td>Male</td>
<td>Adult</td>
<td>(-)</td>
<td>(-)</td>
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</tr>
<tr>
<td>17</td>
<td>ZNO</td>
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<td>Juvenile</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>18</td>
<td>ZNO</td>
<td>Female</td>
<td>Juvenile</td>
<td>(-)</td>
<td>(-)</td>
<td>(-)</td>
</tr>
<tr>
<td>19</td>
<td>A3CO</td>
<td>Male</td>
<td>Adult</td>
<td>(-)</td>
<td>(-)</td>
<td>Mycoplasma haemocanis 100% (882/882) KY117659.1</td>
</tr>
</tbody>
</table>

A3CO: Cantera Oriente; CU: Circuito Universitario; DGAE: Dirección General Administración Escolar; MUAC: Museo Universitario de Arte Contemporáneo; ZNO: Zona Núcleo Oriente; ZNSO: Zona Núcleo Suroriente; ZNP: Zona Núcleo Poniente DNA; (-): Not detected.
To the best of our knowledge, this is the first study to detect the presence of tick and flea-borne pathogens in feral dogs from Mexico, and it is also the first report of *B. vinsonii* subsp. *berkhoffii* and *E. canis* in Mexico City. Besides, our study confirmed, through molecular methods, the presence of *B. vinsonii* subsp. *berkhoffii* and *M. haemocanis* in Mexican dogs. These results contribute to determining the negative relationship between feral dogs and wild animals of REPSA regarding disease transmission. *Ehrlichia canis* (the causative agent of canine monocytic ehrlichiosis) is the main rickettsial agent registered in dogs of the Neotropical region, with prevalence ranging between 15 and 70% (BARRANTES-GONZÁLEZ et al., 2016; MONTENEGRO et al., 2017; ROTONDANO et al., 2017; GEIGER et al., 2018; PAULINO et al., 2018). Clinical signs of an *E. canis* infection can be variable, depending on the strain, the immune response of the dog, and the presence of concomitant infections with other tick or flea-borne pathogens (SAINZ et al., 2015), however, no clinical signs were observed in the animals captured. In Mexico, it has been reported in several states, mainly in the coastal regions of the Gulf of Mexico and also in the states of the northern border close to the United States of America (RODRÍGUEZ-VIVAS et al., 2005; JIMÉNEZ-COELLO et al., 2009; SOSA-GUTIÉRREZ et al., 2013; PAT-NAH et al., 2015; ALMAZÁN et al., 2016). *Ehrlichia canis* is not common in humans; however it has been reported causing human disease in Venezuela (PEREZ et al., 2006) and recently in Panama (DAZA et al., 2018).

*Bartonella vinsonii* subsp. *berkhoffii* is an emerging pathogen that represents a growing public health problem (ROUX et al., 2000; LANTOS et al., 2014). Initially isolated in 1990, in blood samples of a dog and its owner who had an endocarditis, it has been shown to be a widespread pathogen, which has been detected mainly in the United States and recently in several other countries of South America (KORDICK et al., 1996; FENIMORE et al., 2011; PÉREZ et al., 2011; DINIZ et al., 2013). This is a common pathogen of pets, which is transmitted by contact with infected fleas (MÜLLER et al., 2018). In our study, all positive dogs were captured in the same area of the reserve during a season of the same year. In the North of Mexico, it was reported the presence of this *Bartonella* species in two wild carnivores, coyotes (*Canis latrans*) and striped skunks (*Mephitis mephitis*) (LÓPEZ-PÉREZ et al., 2017), however, its effect on these species is unknown because it...
is the cause of endocarditis in domestic dogs (BILLETTER et al., 2012; LÓPEZ-PÉREZ et al., 2017; ROURA et al., 2018).

*Mycoplasma haemocanis* causes hemolytic anemia in immunocompromised dogs (KAEWMONGKOL et al., 2017). The first record of this pathogen in Mexico was in 1960, and detected in splenectomized dogs (OSORNO & RISTIC, 1972). Recently it has been demonstrated that the tick *Rhipicephalus sanguineus* s.l. is capable of transmitting this pathogen (AKTAS & OZUBEK, 2017). It is possible that the frequency of the three microorganisms is low compared with other studies due to the use of clots instead of whole blood; however it is shown that the sera can be useful for the surveillance of these microorganisms (MYLONAKIS et al., 2009).

Furthermore, because feral dogs have a high vagility and are able to establish close interactions with other groups of wild mammals, the exchange of ectoparasites carrying bacterial agents is highly possible. For this reason, it is essential to determine if any of these agents have been established in wild mammal populations because other animals could be endangered. In REPSA there are records of carnivore species that are classified in risk categories for conservation, for example, the recently rediscovered gray fox (*Urocyon cinereoargenteus*), and several species of skunks (*Spilogale gracilis*) and ringtail cats (*Bassariscus astutus*). Also, because these bacterial species can infect other wild carnivores and some of them have been detected causing disease in humans, it is important to continue with active surveillance in feral dog populations in order to identify the impact of these pathogens in the wild and to establish prevention and protection measures in this group of unattended animals.

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**References**


