Romero-Viva, Claudia M.; Thiry, Dorothy; Rodríguez, Virginia; Calderón, Alfonso; Arrieta, Germán; Mátar, Salim; Cuello, Margarett; Levett, Paul N.; Falconar, Andrew K.
Molecular serovar characterization of Leptospira isolates from animals and water in Colombia
Biomédica, vol. 33, núm. 1, 2013, pp. 179-184
Instituto Nacional de Salud
Bogotá, Colombia

Available in: http://www.redalyc.org/articulo.oa?id=84328376019
**Molecular serovar characterization of *Leptospira* isolates from animals and water in Colombia**

Claudia M. Romero-Vivas¹, Dorothy Thiry², Virginia Rodríguez¹, Alfonso Calderón³, Germán Arrieta³, Salim Máttar³, Margarett Cuello¹, Paul N. Levett², Andrew K. Falconar¹

¹ Grupo de Investigaciones en Enfermedades Tropicales, Departamento de Medicina, Universidad del Norte, Barranquilla, Colombia
² Saskatchewan Disease Control Laboratory, Regina, Saskatchewan, Canada
³ Instituto de Investigaciones Biológicas del Trópico, Universidad de Córdoba, Montería, Colombia

**Introduction:** Leptospirosis is a bacterial disease transmitted directly or indirectly from animals to humans that may result in severe hemorrhagic, hepatic/renal and pulmonary disease. There are 20 known *Leptospira* species and hundreds of serovars, some of which belong to different species. It is essential to identify pathogenic *Leptospira* serovars and their potential reservoirs to prepare adequate control strategies.

**Objective:** To characterize the *Leptospira* serovars isolated from rodents, dogs, pigs and water samples in Colombia.

**Materials and methods:** *Leptospira* organisms were isolated and cultured, and pathogenic strains were identified using a polymerase chain-reaction (PCR). *Leptospira* DNA and *Salmonella* Braenderup H9812 (molecular weight standard) DNA were cleaved using *NotI* and subjected to pulsed-field gel electrophoresis (PFGE). The PFGE patterns were analyzed based on bacterial strain-typing criteria and Dice coefficients (DCs) between these isolates and over 200 *Leptospira* organisms isolated from other parts of the world.

**Results:** All of the isolates were pathogenic strains, and five were genetically characterized. The P275 (84% DC) and P282 (95% DC) pig isolates were related to the *Leptospira interrogans* Pomona serovar; the I15 (DC: 100%) rat isolate was identical to the *Leptospira interrogans* Icterohameorrhagiae or Copenhageni serovars, while the C67 (64% DC) dog and A42 (60% DC) water isolates were not related (< 73.7% DC) to any of the 200 reference serovars; the closest serovars were the *Leptospira noguchii* Nicaragua and Orleans serovars, respectively.

**Conclusion:** This was the first molecular characterization of Colombian *Leptospira* spp isolates; these isolates will be used to develop a Colombian diagnostic panel.

**Key words:** *Leptospira*; electrophoresis, gel, pulsed-field; Colombia

doi: http://dx.doi.org/10.7705/biomedica.v33i0.731

**Caracterización molecular de serovariedades de *Leptospira* spp. aisladas de muestras de animales y agua en Colombia**

**Introducción.** La leptospirosis es una infección bacteriana transmitida directa o indirectamente de animales a humanos, la cual puede resultar en una enfermedad hemorrágica grave, hepática o renal y pulmonar. Hay 20 especies de *Leptospira* conocidas y cientos de serovariedades, algunas de las cuales pertenecen a diferentes especies. Es esencial identificar las serovariedades patógenas y sus reservorios potenciales para enfocar estrategias de control.

**Objetivo.** Caracterizar las serovariedades de *Leptospira* aisladas de muestras de roedores, perros, cerdos y agua en Colombia.

**Materiales y métodos.** Las cepas de leptospiras aisladas fueron identificadas como patógenas usando la reacción en cadena de la polimerasa (PRC). Sus ADN y el ADN de *Salmonella* Braenderup H9812 (marcador de peso molecular) fueron cortados con *NotI* y corridos en electroforesis de campo pulsado. Los patrones de la ECP se analizaron con base en los criterios de tipificación para cepas bacterianas y el coeficiente de Dice, cuando se compararon con 200 cepas aisladas en otras partes del mundo.

**Author contributions:**
Andrew K. Falconar designed the study
Margarett Cuello, Virginia Rodríguez, Alfonso Calderón, Germán Arrieta and Salim Mattar performed the *Leptospira* spp. isolations and the PCRs.
Dorothy Thiry and Paul N. Levett provided training in the PFGE and result analysis and prepared the figures.
Andrew K. Falconar wrote the article.
Claudia M. Romero-Vivas participated in all activities.
Los perfiles de ADN con un coeficiente de Dice entre 73,7 % y 100 % se consideraron pertenecientes a la misma especie.

Resultados. Todos los aislamientos fueron cepas patógenas y cinco se caracterizaron genéticamente. El aislamiento P275 (coeficiente de Dice: 84 %) y el P282 (coeficiente de Dice: 95 %) de cerdos, se relacionaron con Leptospira interrogans de serovariety Pomona; el aislamiento de rata (I15) fue indistinguible de Leptospira interrogans de serovarieties Icterohaemorrhagiae o Copenhageni (coeficiente de Dice: 100 %), mientras que los aislamientos de perro (C67) y agua (A42) no se relacionaron (coeficiente de Dice <73,7 %) con ninguna de las 200 cepas de referencia; las más cercanas fueron Leptospira noguchii de serovarieties Nicaragua (coeficiente de Dice: 63 %) y Orleans (coeficiente de Dice: 60 %).

Conclusiones. Esta fue la primera caracterización molecular de serotipos de aislamientos colombianos, los cuales serían los primeros miembros de un panel diagnóstico colombiano.

Palabras clave. Leptospira, electroforesis en gel de campo pulsado, Colombia
doi: http://dx.doi.org/10.7705/biomedica.v33i0.731

Pathogenic Leptospira spp. are chronically maintained in the renal tubules of a wide range of wild and domestic mammals, and enzootic cycles are maintained by direct contact with infected urine or indirect contact with contaminated soil or water (1). More than 260 pathogenic serovars have been serologically identified and grouped into 24 serogroups adapted to different animal species (2). Although some animal species may act as maintenance hosts for some serovars, they can also be incidental hosts for other serovars, which may result in a range of clinical symptoms depending on the infecting strain, the geographical location and the host immune response (3). Leptospirosis is a rural and occupational disease (4) considered to be an emerging zoonotic disease. In tropical and subtropical areas, the transmission of leptospirures is increased during heavy rainfall, flooding, in conditions of poor sanitation and in areas of high host biodiversities; leptospirosis has therefore become a major public health problem in these areas (5).

The microscopic agglutination test (MAT) is the gold standard diagnostic assay for identifying leptospirosis but requires a large panel of live Leptospira serogroups/serovars to be maintained and used. Because it is difficult to isolate Leptospira serovars, the sensitivity of the MAT may be reduced when local serogroups/serovars are not included in these panels, thus limiting the possibility of characterizing other serovars (6). Knowledge of the prevalent serovars and their maintenance hosts and the monitoring of the emergence of new serovars is essential for understanding the epidemiology of the disease in a region and for developing effective control strategies (7).

Leptospira spp. isolates were identified as belonging to the pathogenic L. interrogans and the saprophytic L. biflexa species and were typed to the serovar level based on the expression of surface-exposed lipopolysaccharide (LPS) antigens using the cross agglutinin absorption test (CAAT) (8). Subsequently, molecular techniques such as DNA-DNA hybridization, restriction fragment length polymorphism (RFLP), and 16S rRNA sequence analysis (9) allowed the Leptospira genus to be classified into 14 pathogenic and intermediately pathogenic and 6 saprophytic species (10); however, a particular serovar can also belong to different species (11). Because of the epidemiological importance of precisely typing Leptospira organisms, molecular techniques such as multi-locus sequencing typing (MLST) (12) and pulsed-field gel electrophoresis (PFGE) (13) are useful tools for genetically typing isolates at the serovar level. However, MLST cannot always be used for all Leptospira spp. as some serovars share the same sequence type or a particular serovar has multiple sequence types. In contrast, PFGE is generally applicable to all pathogenic species as this tool can identify strains of serovars that belong to different species and new serovars (10). In this study, we isolated and PFGE-characterized pathogenic Leptospira spp. isolates from animals (dogs, pigs and rats) and water samples collected in Colombia.

Materials and methods

Origin of Leptospira spp. isolates

The Leptospira spp. isolates were obtained from urine samples from two dogs (C67 and C76) (3.7%)
of the 54 dogs sampled and from three pigs (P237, P275 and P282) (0.8%) of the 383 pigs sampled. Two other isolates (A42 and A44) (3.7%) were obtained from two of the 54 water samples. These isolates were obtained in 2008 from six pig farms in Monteria (Cordoba). One other isolate (115) (6.3%) was obtained from a homogenized kidney sample from one of the 16 black rats (Rattus rattus) sampled from Barranquilla (Atlántico) in 2009. All of these isolates were maintained in three 5 ml culture tubes containing Ellinghausen-McCullough-Johnson-Harris (EMJH) liquid medium (Difco, USA) at 28°C.

Detection of pathogenic Leptospira spp isolates using a conventional polymerase chain reaction (PCR)

After five days of incubation, one 5-ml culture from each sample was centrifuged at 5000 x g for 5 min; the supernatants were then discarded and the pellets were re-suspended in 1 ml of media. DNA extraction was performed on 100 μl samples using Qiaamp DNA mini kits (Qiagen, Hilden, Germany). The PCRs were performed in total volumes of 25 μl that contained 2-5 μl of DNA, 0.25 mM dNTPs (Bioline, USA), 1.0 IU of Taq DNA polymerase (Fermentas, Thermo Scientific, USA), 3.0 mM MgCl₂, and 0.1 μM of LipL32/270F and LipL32/692R primers. The PCR program consisted of an initial cycle at 95°C for 5 minutes followed by 35 cycles at 94°C for 1 min and 55°C for 1 min, and a final extension step at 72°C for 5 min (14).

Leptospira isolate typing

The Leptospira isolates were grown in EMJH medium until a turbidity of 0.40 to 0.42 was reached (Dade Behring turbidity meter, Germany). Pulsed-field gel electrophoresis (PFGE) was performed as previously described (10). For this method, Leptospira DNA samples and Salmonella Braenderup H9812 strain DNA (used as standard size markers) were digested with Not I (Roche, Mannheim, Germany). PFGE was performed on a 1% agarose gel using a CHEF DR III (Bio-Rad) for 18 hr at 14°C, with 0.5 times Tris-Borate/EDTA (TBE) buffer recirculation, 2.16 and 35.07 second switching times at 120°, a gradient of 6 V/cm, and a linear ramping factor. The resultant gel was stained with GelRed (Biotium, USA) for 30 minutes and analyses were performed using BioNumerics version 4.0 software (Applied Maths, Inc., Austin, TX, USA). The PFGE profiles were then compared with a library of over 200 Leptospira spp. reference serovars. In the PFGE gel analysis, genetic relatedness was based on the numbers of bands; the Dice coefficients were determined using BioNumerics version 4.0. Genetic relatedness was based on bacterial strain typing criteria and classified as indistinguishable (same number of bands with the same apparent size), closely related (two- or three-band differences), possibly related (four- to six-band differences) or unrelated (seven or more band differences) as previously described (15), and serovars with Dice coefficients between 73.7-100% were considered to be the same serovar.

Results

The 423 bp lipL32 gene fragment, specific to pathogenic Leptospira, was amplified by PCR in all of the pig (P237, P275 and P282), dog (C67, C76), and water samples (A42, A44) from the same farm in Monteria (Cordoba), and from the rat (115) collected in Barranquilla (figure 1).

The PFGE analyses were only performed on isolates that attained turbidities of at least 0.40, and the numbers of PFGE bands obtained for each of the five isolates were as follows: P275 (n = 13), P282 (n = 10), C67 (n =14), A42 (n = 9) and I15 (n = 9) (figure 2). When the DNA profiles of the two pig isolates (P275 and P282) were analyzed, the most similar restriction band patterns were observed for the 2006006986 pathogenic Leptospira strain that belonged to the Leptospira interrogans Pomona serovar. The P275 isolate had 4 additional bands.
and an 84% DC and was therefore related to the Pomona serovar (figure 3). The P282 isolate, however, had only one additional band, resulting in a 95% DC and was therefore classified as more closely related to the Pomona serovar. In contrast, an indistinguishable restriction band pattern (100% DC) was obtained between the I15 rat isolate and the 200600691 strain, which belonged to the *Leptospira interrogans* Icterohaemorrhagiae serovar (figure 3). Although the C67 dog isolate had the same number of bands (n= 14) as lepto0282 *Leptospira noguchii* Nicaragua serovar, it had different restriction band patterns, resulting in a low DC of 64%. The A42 water isolate had two fewer bands than the nearest lepto0258 *Leptospira noguchii* Orleans serovar, which also resulted in a low DC of 60% (figure 3). Thus, because these two isolates had DCs below the threshold range (73.7-100%), they were classified as new serovars.

**Discussion**

To identify and understand alterations in leptospirosis epidemiology, it is essential to characterize the serovars of the circulating *Leptospira* strains in each endemic area so that appropriate control strategies can be implemented. In most endemic areas, this characterization has not been performed due to the slow growth of *Leptospira* organisms, requirements for special nutrients in the culture medium, frequent contamination with other micro-organisms, and the high cost of maintenance and genetic analyses (16). Although we obtained eight *Leptospira* isolates, PFGE could only be performed on five (5/8: 62.5%) of them. More efficient isolation and maintenance methods therefore need to be developed.

While the *L. interrogans* Icterohaemorrhagiae and Copenhageni serovars cause no disease in their rat reservoir hosts, they cause severe icteric Weil's disease in humans, which is particularly common in urban slum areas (17). This was consistent with the fact that the *L. interrogans* Icterohaemorrhagiae/Copenhageni serovars that cannot be differentiated by PFGE analyses (11,21), the *L. interrogans* Pomona serovar, and possibly the *L. noguchii* Nicaragua and Orleans serovars are circulating in the Caribbean coast of Colombia. Despite the fact that a relatively low number of isolates were studied due to the well-documented difficulties in isolating and maintaining *Leptospira* spp., the isolate from one pig (P275) was found to have a different band pattern (84% Dice coefficient (DC)) than the reference *Leptospira interrogans* Pomona serovar, while the isolates from a dog (C67) (64% DC) and water (A42) (60% DC) were uniquely different from the closest reference *Leptospira interrogans* Pomona serovar and the isolates from a dog (C67) (64% DC) and water (A42) (60% DC) were uniquely different from the closest reference *Leptospira noguchii* Nicaragua and Orleans serovars. These important isolates are the first Colombian isolates to be PFGE characterized; they will be further genetically analyzed using assays such as the gold standard cross agglutinin absorption test (CAAT) (9). Because it is advisable to use locally isolated *Leptospira* strains to obtain optimal sensitivities in serological assays (such as the MAT) (1), these uniquely different isolates will be used to establish a Colombian panel of genetically typed strains to improve *Leptospira* diagnosis in Colombia.

**Acknowledgements**

We thank the Dirección de Investigaciones, Desarrollo e Innovación de la Universidad del Norte, and the Centro de Investigaciones Universidad de Córdoba, código del proyecto FMV-2006.

**Conflict of interest**

The authors declare that no conflicts of interest exist.
Figure 3. PFGE DNA fragment patterns of the Colombian isolates from pigs (P275, P282), a rat (I15), a dog (C67) and a water sample (A42), control strains with species and serogroup information of Pomona, Icterohaemorrhagiae, Orleans, and Nicaragua serovars from the PFGE database (10) with the percentage similarities (Dice coefficients) shown on the left.

Financial support
This study was supported by the Departamento Administrativo de Ciencia Tecnología e Innovación, Colciencias ID Project Number 1215-408-20551, Universidad del Norte, internal project 2011 Leptospirosis and the Universidad de Córdoba Project Number FMV-2006.

References


