

## GENOTYPIC IDENTIFICATION OF AN UNDESCRIBED SPOTTED FEVER GROUP *RICKETTSIA* IN *IXODES RICINUS* FROM SOUTHWESTERN SPAIN

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**Abstract.** An undescribed rickettsia was directly analyzed with specific rickettsial molecular biology tools on *Ixodes ricinus* L. collected in different localities of the province of Cadiz (southwestern Spain). On the basis of the results of the citrate synthase (*gltA*) gene, 190 kD-outer membrane protein (*rOmpA*) gene, and 16S ribosomal RNA (16S rRNA) gene partial sequence data, it was found that this rickettsia is sufficiently genetically distinct from other *Rickettsia* to be considered a distinct taxonomic entity. The isolation and culture of this organism, as well as comparative antigenic analysis, are required to ensure its conclusive taxonomic placement among spotted fever rickettsiae. The epidemiologic role of this new rickettsial agent and its possible pathogenicity to wild and domestic animals or humans is still unknown and needs to be investigated.

Rickettsiae are gram-negative bacteria that multiply only inside host cells and require arthropods either as reservoirs or as vectors.<sup>1</sup> The bacterial genus *Rickettsia* is traditionally divided into three biotypes: the spotted fever group (SFG), the typhus group, and the scrub typhus group, based on vector host and antigenic cross-reactivity.

Some *Rickettsia* infect lice, fleas, and mites, but most are associated with various species of Ixodid ticks. As a first approach, field studies oriented to the detection of rickettsial and rickettsial-like organisms in ticks and other arthropods, and the subsequent precise identification of these bacteria are of critical importance in understanding the epidemiology, natural history, and potential threat to human health, along with a reconsideration of established components of the vector-reservoir cycle.<sup>2,3</sup>

Until the mid-1980s, the identification of new species of rickettsia was a tedious and time-consuming process that required successful propagation of the organism in a cell culture system to characterize such parameters as growth and cytopathology.<sup>4</sup> In the first trials, the identification of *Rickettsia* harbored by individual ticks was made using indirect microimmunofluorescent serologic typing<sup>5,6</sup> and microagglutination<sup>7</sup> after isolation of the strain, and comparing the isolate with prototype strains belonging to known rickettsia species. Recent advances in molecular biology, and the availability of amplification, restriction, and sequencing techniques have determined the development of new detection and characterization tests concerning rickettsiae. In this regard, the analysis of restriction endonuclease digest of DNA from purified rickettsiae has been used primarily for its genotypic identification and estimation of intraspecies divergence,<sup>8–11</sup> and secondarily for the detection of such rickettsiae in different vectors (ticks,<sup>12–14</sup> fleas,<sup>15,16</sup> or mites<sup>17</sup>) and humans.<sup>3,18,19</sup>

Analysis of ribosomal gene sequences has proven to be useful for identifying genotypic relationships between major groups of rickettsia-like organisms,<sup>20</sup> differentiating at a specific level between recognized and unrecognized species of the genus *Rickettsia*,<sup>21–23</sup> despite the extreme sequence conservation observed among these microorganisms. In this way, a consensus sequence polymerase chain reaction (PCR) relies on the use of highly conserved and specific DNA sequences, such as ribosomal 16S rRNA gene sequences, and

has been demonstrably useful in amplifying DNA from as yet undiscovered though related organisms.<sup>24,25</sup>

Rickettsiosis are zoonoses limited geographically by the distribution area of their infected vector. *Ixodes ricinus* is an exophilic, three-host tick widely distributed in European countries. To date, only one species of *Rickettsia* (*R. helvetica*, formerly called Swiss agent), has been isolated from *I. ricinus* ticks in Switzerland and Sweden.<sup>26–31</sup> Recently, the presence of *Ehrlichia phagocytophila* was reported in *I. ricinus* in Switzerland,<sup>32</sup> and *Coxiella burnetii* and an undefined rickettsial organism were isolated from *I. ricinus* in Austria.<sup>33</sup>

### MATERIALS AND METHODS

**Collection of ticks.** Adult *I. ricinus* were obtained from red deer (*Cervus elaphus*) hunted in woodland areas of Ubrique, Alcalá de los Gazules, Jimena de la Frontera and Los Barrios (Cadiz Province, Spain) (Figure 1) during the winters of 1995–1996 and 1996–1997, and classified by an entomologist (FJM).

**Extraction of DNA.** Individual *I. ricinus* adults were crushed in 340  $\mu$ l of lysis buffer (100 mM Tris-HCl, pH 8.0, 100 mM NaCl, 10 mM EDTA), added to 40  $\mu$ l of 10% sodium dodecyl sulfate and 20  $\mu$ l of proteinase K (Promega, Madison, WI) (50 mg/ml), and maintained in a bath preset at 55°C for 1 hr. After incubation, samples were extracted with phenol-chloroform as described elsewhere.<sup>34</sup> Nucleic acids were precipitated adding 0.1 volumes of potassium acetate solution (5 M based on acetate), 2.5 volumes of absolute ethanol, and chilling to -20°C.

**Polymerase chain reaction–restriction fragment length polymorphism (RFLP) method.** Amplification was assayed using three sets of primers: RpCS primer set (RpCS.877p and RpCS.1258n) and Rr190 primer set (Rr190.70p and Rr190.602n) derived from the sequences of *R. prowazekii* and *R. rickettsii* respectively,<sup>8</sup> and the primer pair derived from the 120-kD outer membrane protein (*rOmpB*) gene of *R. rickettsii* (BG 1-21 and BG 2-20).<sup>10</sup> The sequences and orientations of these primers (Boehringer, Mannheim, Germany) are described in Table 1.

A 100- $\mu$ l reaction mixture, which contained 1  $\mu$ l of sample, 56  $\mu$ l of distilled water, 10  $\mu$ l of 10 $\times$  Taq polymerase



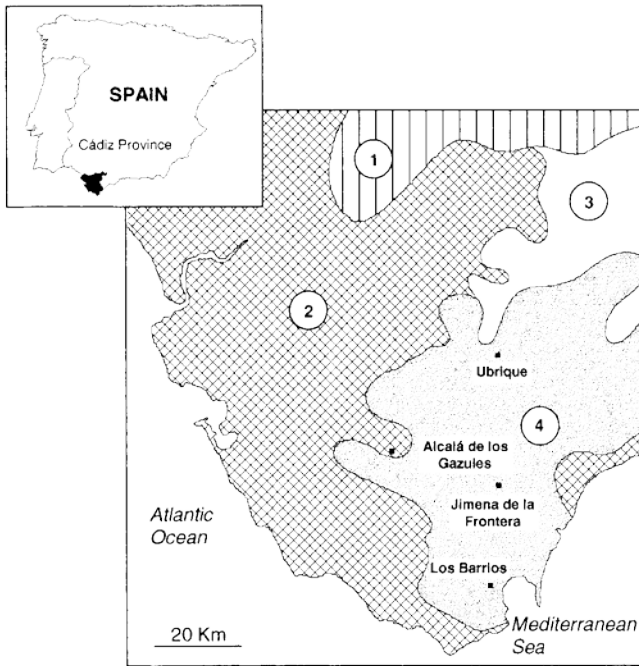


FIGURE 1. Geographic localization of the sites in the province of Cadiz, Spain where *Ixodes ricinus* were collected. Numbered areas correspond to very dry (1), dry (2), subhumid (3), and humid (4) climate areas.

buffer (Promega), 8 µl of 25 mM MgCl<sub>2</sub>, 20 µl of 1 mM dNTPs, and 0.5 U of *Taq* DNA polymerase (Promega), was prepared. Each of the 35 cycles of amplification consisted of denaturation at 95°C for 20 sec, annealing at 48°C for 30 sec, and sequence extension at 72°C for 2 min.<sup>8,10</sup> All amplification reactions were conducted in a DNA Thermocycler 9600 (Perkin Elmer, Norwalk, CT), using thin-wall microamp® tubes (Perkin Elmer). After amplification, 7 µl of the PCR mixture was electrophoresed for 1 hr at 7 V/cm in a 4% Nusieve® GTG agarose (FMC Bioproducts, Rockland, ME) gel in 1 × TAE buffer (0.04 M Tris-acetate, 0.001 M EDTA), using the molecular weight marker VI (MWM VI) (Boehringer Mannheim) as a control, stained with ethidium bromide, and observed under ultraviolet light illumination.

Samples for RFLP analysis were prepared by digesting 15 µl of the PCR mixture containing the amplified fragment with the appropriate restriction endonuclease (Promega)<sup>8</sup> for the citrate synthase (*gltA*) and the outer membrane protein (*rOmpA*) genes.

**Sequencing.** Amplification of 5' region of the 16S rRNA gene was made using the primers fD1<sup>20</sup> and Rc16S.452n. To ensure the uniqueness of the Rc16S.452 primer, the 16S rRNA sequence of *R. canada* (GenBank accession numbers L36104 and U15162) was compared by homology with other GenBank related sequences using the FASTA algorithm of the Genetic Computer Group (GCG) (University of Wisconsin, Madison, WI) program, allowing for three or less mismatches. The *Rickettsia*-exclusive sequences were selected from the FASTA alignments file and the candidate primers were tested for exclusivity to SFG rickettsiae using the FINDPATTERNS algorithm from GCG. The PCR was per-

TABLE 1  
Oligonucleotide primers used for genotypic identification of rickettsial species\*

Species	Gene	Primer	Nucleotide sequence (5' 3')	Amplified fragment size (bp)	Authors and reference
<i>R. prowazekii</i>	CS	RpCS.877p	GGGGCCTGCTCAGCGGG	381	Regnery and others <sup>8</sup>
<i>R. rickettsii</i>	120-kD genus common antigen (rompB)	RpCS1258n	ATTGCCAAAAGTACAGTGAACA	650	Anderson and others <sup>39</sup>
		RrBG.1-21	GGCAATTAATATCGCTGACGG		
	190-kD protein antigen (rompA)	RrBG.2-20	GCATCTGCACTAGCACATTC	532	Regnery and others <sup>8</sup>
Protobacteria <i>R. canada</i>	16S rRNA	Rr190.70p	ATGGCGAATATTTCTCCAAA	426	Weisburg and others <sup>20</sup>
		Rr190.602n	AGTGCAGCATTGGCTCCCCCT		
		fD1	AGAGTTTGATCTGGCTCAG		This report
		Rc16S.452n	AACGTCATTATCTTCTTGC		

\* bp = basepairs; CS = citrate synthase (*gltA*).

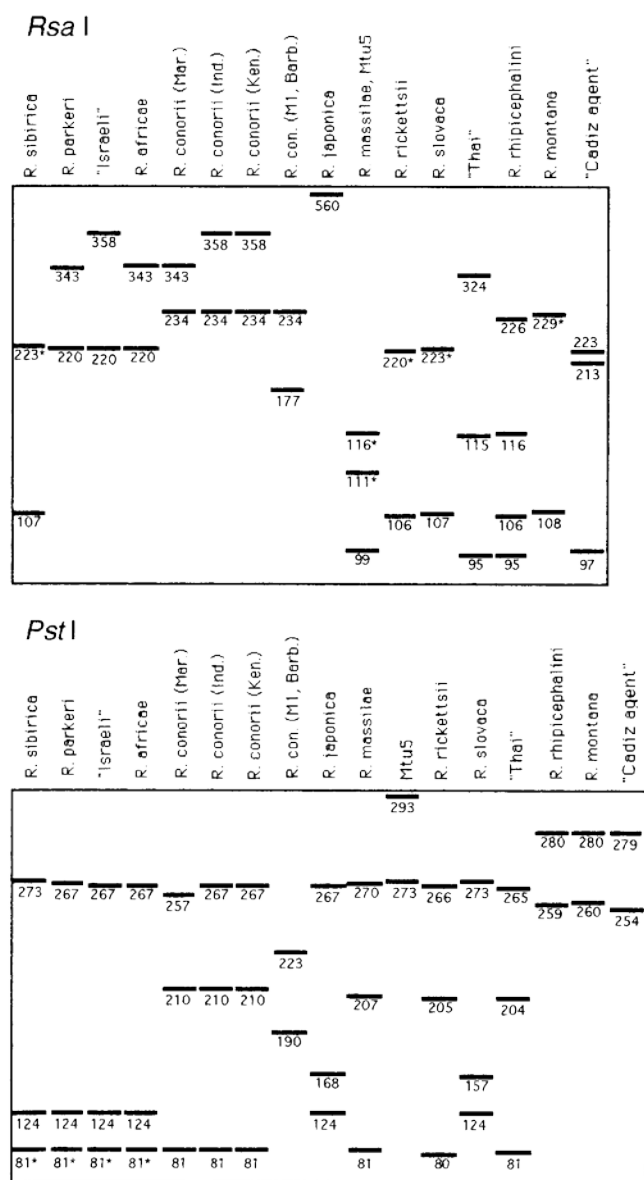


FIGURE 2. Electrophoretic migration patterns of polymerase chain reaction-amplified rickettsial DNA with the Rr 190.70p and Rr 190.602n primer pair digested with the restriction endonucleases *Rsa* I and *Pst* I. The schematic profiles of the other species of *Rickettsia* were adopted from Regnery and others<sup>8</sup> and Eremeeva and others.<sup>10</sup> Mar. = Morocco strain; Ind. = India strain; Ken. = Kenya strain; con = *conorii*; Barb. = Barbados strain. \* = double band.

formed for 30 cycles in a reaction mixture at 95°C for 20 sec, 59°C for 30 sec, and 72°C for 45 sec.

The specific primers assayed were used to sequence the amplified fragments of *gltA*, *rOmpA*, and a 5' portion of the 16S rRNA genes of the rickettsiae detected in the infected ticks. Thermal cycling conditions and product observation were as described in the above protocols.

After amplification, primers and nucleotides were removed from 300  $\mu$ l of PCR products by purification on the Wizard<sup>TM</sup> PCR preps DNA purification system (Promega), according to the manufacturer's protocol. The remaining DNA was eluted in 50  $\mu$ l of TE buffer (10 mM Tris-HCl, 1

mM EDTA, pH 7.6). In this case, the amount of DNA obtained was quantified in a 4% Nusieve<sup>®</sup> GTG agarose gel in 1  $\times$  TAE buffer by comparing fluorescence emission with 1  $\mu$ g of MWM VI. Approximately 100 fmol of the purified PCR product (4–5  $\mu$ l) was used directly in the sequencing reaction.

The PCR cycle sequencing was performed (Silver sequence<sup>TM</sup> DNA Sequencing System; Promega) for each amplicon using the correct forward or reverse primers. Four reactions of 9  $\mu$ l containing 3  $\mu$ l of the appropriate termination mixture and 6  $\mu$ l of the sample reaction mixture were made (100 fmol of purified product of the PCR, 1.8  $\mu$ l of 5 $\times$  DNA sequencing buffer [75 mM Tris-HCl, pH 9.0, 3 mM MgCl<sub>2</sub>], 6.75 pmol of respective primer and 0.25 U of *Taq* DNA polymerase sequencing grade). Tubes were preheated to 95°C for 2 min and subjected to linear amplification for 55 cycles at 95°C for 30 sec, 42°C or 59°C depending on the calculated melting temperature of the primer concerned for 30 sec, and 72°C for 1 min. A volume of 4.5  $\mu$ l of sequencing stop solution (10 mM NaOH, 95% formamide, 0.05% bromophenol blue, 0.05% xylene cyanol) was added to each sequencing tube and heated at 65°C for 2 min.

Sequencing reaction products were loaded twice on 40-cm, 6% polyacrylamide, 7 M urea gels by electrophoresis in the Sequi-Gen Nucleic Acid Sequencing System (Bio-Rad, Hercules, CA) at 55 W of constant electrophoresis (55°C) and separated for 4.5 and 2.5 hr, respectively. Gels were fixed in 2 liters of fix/stop solution (10% glacial acetic acid) for 20 min (until diffusion of the tracking marks in the gel occurred), rinsed three times with ultrapure water for 2 min, stained with 2 liters of 0.1% AgNO<sub>3</sub> and 0.055% formaldehyde for 30 min, rinsed for 5 sec with ultrapure water, developed with 2 liters of 0.14 M sodium carbonate, 0.055% formaldehyde, and 4 mg of sodium thiosulfate, and chilled to 8–10°C (8–12 min) until the bands appeared. To terminate the developing reaction, an equal volume of fix/stop solution was added. Finally, the gel was rinsed twice with ultrapure water and completely dried in an oven at 45°C. A permanent record was made using APC film (Promega) exposed to overhead fluorescent lighting. Development of APC film was carried out according to the manufacturer's protocol.

For electron microscope studies, infected *I. ricinus* females were dissected in a mixture of 2% paraformaldehyde and 2% glutaraldehyde in 0.1 M cacodylate buffer, postfixed in a 2% osmium tetroxide solution, dehydrated in increasing concentrations of ethanol, and embedded in the resin Epon 812. Ultrathin sections were cut by ultramicrotome (model Ultracut S; Reichert-Jung, Vienna, Austria), mounted on grids, stained with uranyl acetate and lead citrate, and examined by transmission electron microscope (model EM10C; Carl Zeiss, Oberkochen, Germany) at 80 kV.

## RESULTS

Preliminary detection and identification of SFG rickettsia in *I. ricinus* adults was performed using PCR/RFLP with specific rickettsial molecular biology tools. We attempted to amplify the *gltA* gene with rickettsial genus-specific primers, and the *rOmpA* and *rOmpB* genes with SFG-specific primers.

The RpCS.877p-RpCs1258n (CS) and Rr190.70p-





"Cadiz agent"	GIKTDLFTTSTAALMLSSSGALGVAA-GDLFTINDAAFSDLAAAGNWNKITAGGVANGTSDVDPQDNKAFYTGGA
<i>R. montana</i>	.L.AA.....I.....F..V..VIS-N.....N..VN...E..V...T...PA...R..V.....D
Astrakhan	.L.AA.....I.....VIA.N.N.....NVGN...E...A..T...PAGS..N.W.....D
Israeli	.L.AA.....I.....VIA.N.N.....NVGN...E...A...VAG...N.W.....D
<i>R. africae</i>	.L.AA.....I.....VIA.N.N.....NVGN...E...A...NPAGS..N.W.....D
BAR-29	.L.AA.....I.....V..V..VIS.N.N.....V.N...D...Q...P.....NGM.....D
<i>R. conorii</i> #1	.L.AA.....I.....I.VS.VIA.N.N.....NVGN...E...A...PAH...N.W.....D
<i>R. conorii</i> #2	.L.AA.....I.....I.VS.VIA.N.N.....NVGN...E...A...PAR...N.W.....D
<i>R. conorii</i> #3	.L.AA.....I.....I.VS.VIA.N.N.....NVGN...E...A...PAR...N.W.....D
HA-91	.L.AA.....I.....VIA.N.N.T...NVGN...E...A...APAG...N.W.....D
<i>R. japonica</i>	.L.AA.....I.....VIA.N.....ND...N..Q...R.E.D.NPAG...N.W.....D
<i>R. massiliae</i>	.L.AA.....I.....V..V..VIS.N.N.....V.N...D...Q...P.....NGM.....D
MC16	.L.AA.....I.....VVSNN.N.....N..D...E..I...P.....NGM.....D
<i>R. parkeri</i>	.L.AA.....I.....VIA.N.N.....DVNN...SE...A...VPAGS..N.W.....D
<i>R. rhipiceph.</i>	.L.AA.....I.....V..VIS.N.N.....V.N...D...K...P.....NGM.....D
<i>R. rickettsii</i>	.L.AA.....I.....T.VIA.N.N.....NVGN...E...A...PAG...N.W.....D
Rick. S	.L.AA.....I.....VIA.N.N.T...NVGN...E...A...NPAG...N.W.....D
<i>R. sibirica</i>	.L.AA.....I.....VIA.N.N.I...NVGN...E...A...PAG...N.W.....D
<i>R. slovacica</i>	.L.AA.....I.....VIA.N.N.....NVGN...E...A...PAGS..N.W.....D
Thai	.L.AA.....I.....T.VIA.N.N.....NVGN...E...A...NPAGS..N.W.....D
"Cadiz agent"	HIITADEVGRITAINVAATNPIGIKIAGNTSVGSIVTDRNLLPVNITAGKSLTLTGTAAFVPRHGVGAFADTY
<i>R. montana</i>	.T...K.....G.T.V.LD.TQ..V.....GG.....T.I.....N.AN.I.AN..FD.P..N.
Astrakhan	YT...AAD.....D.T.V.LN..Q..V.....RG.....T.....N.NN.VAAN..F..P..N.
Israeli	YT...VAD.....D.T.V.LN.VQ..V.....RG.....T.....N.NN.VAAN..F..P..N.
<i>R. africae</i>	YT...AAD.....G.T.V.LN..Q..V.....I.GG.....T.....N.NN.VAAN..FD.P..N.
BAR-29	.T...A.....G.T.V.LN.TQ..V.....GG.....T.....N.TN.VAAN..FD.P..N.
<i>R. conorii</i> #1	YT...VADH.....D.T...LN..Q..V.....GG.....T.....N.NN.DAAN..F..P..N.
<i>R. conorii</i> #2	YT...VADH.....D.T...LN..Q..V.....GG.....T.....N.NN.DAAN..F..P..N.
<i>R. conorii</i> #3	YT...VADH.....D.T...LN..Q..V.....GG.....T.....N.NN.DAAN..F..P..N.
HA-91	YT...AAD.....G.T.V.LD..Q..V.....I.GG.....T.....N.NN.VAAN..FD.P..N.
<i>R. japonica</i>	YT...V.NC..K...N...V.LN..Q..V.....I.GG.....T.....N.NN.VAAN..FD.P..N.
<i>R. massiliae</i>	.T...A.....G.T.V.LN.TQ..V.....GG.....T.....N.TN.VAAN..FD.P..N.
MC16	.T...A.....G.T.V.LN.TQ..V.....GG.....T.....N.TN.VAAN..FD.P..N.
<i>R. parkeri</i>	YT...AAD.....G.T.V.LN..Q..V.....I.GG.....T.....N.NN.VAAN..FD.P..N.
<i>R. rhipiceph.</i>	.T...A.C.....G.T.V.LN.TQ..V.....GG.....T.....N.TN.VAAN..FD.P..N.
<i>R. rickettsii</i>	YTV...AADR..K...G.T.V.LN.TQ..V.....I.RG.....TLN.....N.NN.VAAN..FD.P..N.
Rick. S	YT...AAD.....G.T.V.LN..Q..V.....I.GG.....T.....N.NN.VAAN..FD.P..N.
<i>R. sibirica</i>	YT...AAD.....G.T.V.LN..Q..V.....I.GG.....T..V.....N.NN.VAAN..FD.P..N.
<i>R. slovacica</i>	YT...AAD.....G.T.V.LN..Q..V.....I.RG.....T.N.....N.NN.VAAN..FDSP..N.
Thai	YT...AAD.....G.T.V.LNV.Q..V.....I.RG.....T.....N.TN.VAAN..FDAP.NS.

FIGURE 5. Alignment of the deduced amino acid sequence of the 190-kD outer membrane protein (*rOmpA*) gene fragment between positions 25 and 468 (*Rickettsia rickettsii* numbering in GenBank accession number U43804). Amino acids are conserved between sequences (.) except where indicated. Amino acid deletions are indicated (-). Accession numbers of GenBank deposited sequences are U43801 (*R. montana*), U43791 (Astrakhan rickettsiae), U43797 (Israeli tick typhus), U43790 (*R. africae*), U43792 (BAR-29), U43794 (*R. conorii* #1), U43798 (*R. conorii* #2), U45244 (*R. conorii* #3), U43796 (*Rickettsia* HA-91), U43795 (*R. japonica*), U43793 (*R. massiliae*), U43800 (*Rickettsia* MC16), U43802 (*R. parkeri*), U43803 (*R. rhipicephali* [*rhipiceph.*]), U59722 (*Rickettsia* sp. #2), U43794 (*R. conorii*), U59732 (*R. parkeri*), U43804 (*R. rickettsii*), U43805 (*Rickettsia* S), U43807 (*R. sibirica*), U43808 (*R. slovacica*), and U43809 (Thai tick typhus).

gene in *I. ricinus* infected with *R. helvetica* collected in Switzerland.<sup>30</sup>

Ereemeva and others<sup>10</sup> pointed out that different methods of identification of rickettsiae (analysis of antigenic diversity, restriction profiles in pulsed-field electrophoresis, or PCR/RFLP) gave the same results, mainly because these identification methods are based on the analysis of the major surface proteins of the rickettsiae, *rOmpA* and *rOmpB*, which are extensively implicated in the antigenic properties of different species and strains, and thus in the protective immune response of the vertebrate host.<sup>21, 39, 40</sup>

Throughout the study, this new *Rickettsia* has differed considerably from other known rickettsiae. We believe that it represents a new genotype, with notable differences in the 16S rRNA sequence,<sup>21-23</sup> in the *rOmpA* PCR/RFLP pattern, in partial sequence data, and in the deduced protein sequences. Nonetheless, it must be included in the SFG by virtue of similarities inferred from the partial sequence of 16S rRNA and *gltA* genes and the electrophoretic band pattern obtained from the restriction of the PCR-amplified *gltA* fragment, according to criteria previously proposed.<sup>8, 10</sup> Extending the discussion initiated by Beati and others,<sup>30</sup> the failure to amplify the 190-kD (*R. akari*, *R. australis*, *R. bellii*, and *R.*

*helvetica*) and 120-kD (*R. akari*, *R. bellii*, *R. helvetica*, and *R. massiliae*) outer membrane proteins may imply that the major differences among the SFG *Rickettsia* are located in the outer membrane surface proteins. The *rOmpA* sequences indicate that this new species is different from other known SFG rickettsiae.

The Cadiz mountains are the wettest areas of the Iberian Peninsula, with annual rainfall reaching 1,200 mm per year at the Ubrique Meteorological Station, and they have a temperate climate (with an average maximum temperature of 15.7°C and an average minimum temperature of 4.3°C) that is very propitious to the successful growth of an isolated *I. ricinus* population.

The epidemiologic role of the Cadiz agent and its possible pathogenicity requires confirmation. As for *R. helvetica*,<sup>29</sup> low heterologous cross-reactive titers to sera derived from natural and experimental infection caused by this new *Rickettsia* could be expected. Responses to these rickettsiae may confuse the interpretation of serologic tests in patients from the province of Cadiz and contiguous areas who are suspected of having Mediterranean spotted fever, particularly if the high number of bites due to *I. ricinus* in humans and the main role played by this tick as the vector of Lyme borre-

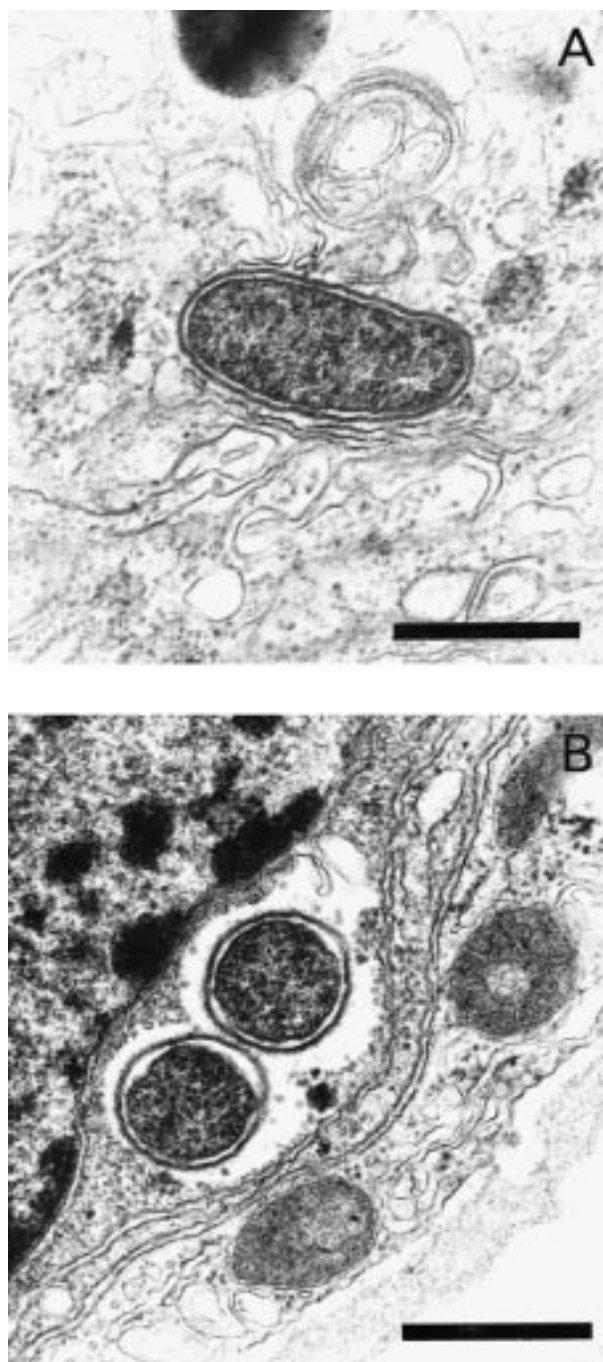


FIGURE 6. Electron micrographs of rickettsiae-infected *Ixodes ricinus* tissues. **A**, longitudinal section of a rickettsiae living free in the cytoplasm of a *Ixodes ricinus* cell. **B**, transversal sections of two rickettsiae associated with the endoplasmic reticulum. Bars = 0.5  $\mu$ m.

liosis in Europe are considered. In addition, immunologic characterization must be carried to conclude that this rickettsia is a new species.

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#### REFERENCES

1. Winkler HH, 1990. *Rickettsiae* species (as organisms). *Annu Rev Microbiol* 44: 131–153.
2. Drancourt M, Kelly PJ, Regnery R, Raoult D, 1992. Identification of spotted fever group rickettsiae using polymerase chain reaction and restriction-endonuclease length polymorphism analysis. *Acta Virol* 36: 1–6.
3. Schriefer ME, Sacci JB Jr, Taylor JP, Higgins JA, Azad AF, 1994. Murine typhus: updated roles of multiple urban components and a second typhus like rickettsia. *J Med Entomol* 31: 681–685.
4. Higgins JA, Radulovic S, Schieffer ME, Azad AF, 1996. *Rickettsia felis*: a new species of pathogenic rickettsia isolated from cat fleas. *J Clin Microbiol* 34: 671–674.
5. Philip RN, Casper EA, Burgdorfer W, Gerloff RK, Hughes LE, Bell EJ, 1978. Serological typing of rickettsiae of the spotted fever group by microimmunofluorescence. *J Immunol* 121: 1961–1968.
6. Philip RN, Casper EA, Ormsbee RA, Peacock MG, Burgdorfer W, 1976. Microimmunofluorescence test for the serological study of Rocky Mountain spotted fever and typhus. *J Clin Microbiol* 3: 51–61.
7. Fiset P, Ormsbee RA, Silbermann R, Peacock M, Spielman SH, 1969. A microagglutination technique for detection and measurement of rickettsial antibodies. *Acta Virol* 13: 60–63.
8. Regnery RL, Spruill CL, Plikaytis BD, 1991. Genotypic identification of rickettsiae and estimation of intraspecies sequence divergence for portions of two rickettsial genes. *J Bacteriol* 173: 1576–1589.
9. Yu X, Jin Y, Fan M, Xu G, Liu Q, Raoult D, 1993. Genotypic and antigenic identification of two new strains of spotted fever group rickettsiae isolated from China. *J Clin Microbiol* 31: 83–88.
10. Eremeeva M, Yu X, Raoult D, 1994. Differentiation among spotted fever group rickettsiae species by analysis of restriction fragment length polymorphism of PCR-amplified DNA. *J Clin Microbiol* 32: 803–810.
11. Furuya Y, Katayama T, Yoshida Y, Kaiho I, 1995. Specific amplification of *Rickettsia japonica* DNA from clinical specimens by PCR. *J Clin Microbiol* 33: 487–489.
12. Gage KL, Gilmore RD, Karstens RH, Burgdorfer W, Schwan TG, 1994. DNA typing of rickettsiae in naturally infected ticks using a polymerase chain reaction/restriction fragment length polymorphism system. *Am J Trop Med Hyg* 50: 247–260.
13. Dupont HT, Cornet JP, Raoult D, 1994. Identification of rickettsiae from ticks collected in the Central African Republic using the polymerase chain reaction. *Am J Trop Med Hyg* 50: 373–380.
14. Uchida T, Yan Y, Kitaoka S, 1995. Detection of *Rickettsia japonica* in *Haemaphysalis longicornis* ticks by restriction fragment length polymorphism of PCR product. *J Clin Microbiol* 33: 824–828.
15. Webb L, Carl M, Malloy DC, Dasch GA, Azad AF, 1990. Detection of murine typhus infection in fleas by using the polymerase chain reaction. *J Clin Microbiol* 28: 530–534.
16. Azad AF, Webb L, Carl M, Dasch GA, 1990. Detection of rickettsiae in arthropod vectors by DNA amplification using the polymerase chain reaction. *Ann NY Acad Sci* 590: 557–563.
17. Kelly DJ, Dasch GA, Chan TC, Ho TM, 1994. Detection and



- characterization of *Rickettsia tsutsugamushi* (Rickettsiales: Rickettsiaceae) in infected *Leptotrombidium* (*Leptotrombidium flecheri* chiggers (Acari: Trombiculidae) with the polymerase chain reaction. *J Med Entomol* 31: 691–699.
18. Sexton DJ, Kanj SS, Wilson K, Corey GR, Hegarty BC, Levy MG, Breitschwerdt EB, 1994. The use of a polymerase chain reaction as a diagnostic test for Rocky Mountain spotted fever. *Am J Trop Med Hyg* 50: 59–63.
  19. Williams WJ, Radulovic S, Dasch GA, Lindstrom J, Kelly DJ, Oster CN, Walker DH, 1994. Identification of *Rickettsia conorii* infection by polymerase chain reaction in a soldier returning from Somalia. *Clin Infect Dis* 19: 93–99.
  20. Weisburg WG, Dobson ME, Samuel JE, Dasch GA, Mallavia L, Baca O, Mandelco L, Sechrest JE, Weiss E, Woese CR, 1989. Phylogenetic diversity of the rickettsiae. *J Bacteriol* 171: 4302–4306.
  21. Stothard DR, Clark JB, Fuerst PA, 1994. Ancestral divergence of *Rickettsia bellii* from the spotted fever and typhus groups of *Rickettsia* and antiquity of the genus *Rickettsia*. *Int J Syst Bacteriol* 44: 798–804.
  22. Roux V, Raoult D, 1995. Phylogenetic analysis of the genus *Rickettsia* by 16S rDNA sequencing. *Res Microbiol* 146: 385–396.
  23. Stothard DR, Fuerst PA, 1995. Evolutionary analysis of the spotted fever and typhus group *Rickettsia* using 16S rRNA gene sequences. *Syst Appl Microbiol* 18: 52–61.
  24. Relman DA, 1993. The identification of uncultured microbial pathogens. *J Infect Dis* 168: 1–8.
  25. Amann RI, Ludwig W, Schleifer KH, 1995. Phylogenetic identification and in situ detection of individual microbial cells without cultivation. *Microbiol Rev* 59: 143–169.
  26. Aeschlimann A, Burgdorfer W, Matile H, Péter O, Wyler R, 1979. Aspects nouveaux du rôle de vecteur joué par *Ixodes ricinus* L. en Suisse. *Acta Trop* 36: 181–191.
  27. Burgdorfer W, Aeschlimann A, Peter O, Hayes SF, Philip RN, 1979. *Ixodes ricinus*: vector of a hitherto undescribed spotted fever group agent in Switzerland. *Acta Trop* 36: 357–367.
  28. Péter O, Burgdorfer W, Aeschlimann A, 1981. Enquête épidémiologique dans un foyer naturel de rickettsies a *Ixodes ricinus* du plateau suisse. *Ann Parasitol Hum Comp* 56: 1–8.
  29. Beati L, Péter O, Burgdorfer W, Aeschlimann A, Raoult D, 1993. Confirmation that *Rickettsia helvetica* sp. nov. is a distinct species of the spotted fever group of rickettsiae. *Int J Syst Bacteriol* 43: 521–526.
  30. Beati L, Humair PF, Aeschlimann A, Raoult D, 1994. Identification of spotted fever group rickettsiae isolated from *Dermacentor marginatus* and *Ixodes ricinus* ticks collected in Switzerland. *Am J Trop Med Hyg* 51: 138–148.
  31. Nilsson K, Jaenson TGT, Uhnoo I, Lindquist O, Pettersson B, Uhlén M, Friman G, Pahlson C, 1997. Characterization of a spotted fever group rickettsia from *Ixodes ricinus* ticks in Sweden. *J Clin Microbiol* 35: 243–247.
  32. Zhu Z, Aeschlimann A, Gern L, 1992. Rickettsia-like organisms in the primordia of molting *Ixodes ricinus* (Acari: Ixodidae) larvae and nymphs. *Ann Parasitol Hum Comp* 67: 99–110.
  33. Rěháček J, Kaaserer B, Urvö Igyi J, Lukáčová M, Kováčová E, Kocianova E, 1994. Isolation of *Coxiella burnetii* and of an unknown rickettsial organism from *Ixodes ricinus* ticks collected in Austria. *Eur J Epidemiol* 10: 719–723.
  34. Sambrook J, Fritsch EF, Maniatis T, 1989. *Molecular Cloning*. Second edition. Cold Spring Harbor, NY: Cold Spring Harbor Press.
  35. Roux V, Rydkina E, Eremeeva M, Raoult D, 1997. Citrate synthase gene comparison, a new tool for phylogenetic analysis, and its application for the rickettsiae. *Int J Syst Bacteriol* 47: 252–261.
  36. Roux V, Fournier PE, Raoult D, 1996. Differentiation of the spotted fever group rickettsiae by sequencing and analysis of restriction fragment length polymorphism of PCR-amplified DNA of the gene encoding the protein *rOmpA*. *J Clin Microbiol* 34: 2058–2065.
  37. Gage KL, Schrupf ME, Karstens RH, Schwan TG, 1992. Detection of *Rickettsia rickettsii* in saliva, hemolymph and triturated tissues of infected *Dermacentor andersoni* ticks by polymerase chain reaction. *Mol Cell Probes* 6: 333–341.
  38. Lange JV, El Dessouky AG, Manor E, Merdan I, Azad AF, 1992. Spotted fever rickettsiae in ticks from the northern Sinai Governorate, Egypt. *Am J Trop Med Hyg* 46: 546–551.
  39. Anderson BE, McDonald GA, Jones DC, Regnery RL, 1990. A protective protein antigen of *Rickettsia rickettsii* has tandemly repeated, near-identical sequences. *Infect Immun* 58: 2760–2769.
  40. Yan Y, Uchiyama T, Uchida T, 1994. Nucleotide sequence of polymerase chain reaction product amplified from *Rickettsia japonica* DNA using *Rickettsia rickettsii* 190-kilodalton surface antigen gene primers. *Microbiol Immunol* 38: 865–869.