Molecular Method for *Bartonella* Species Identification in Clinical and Environmental Samples

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A new, efficient molecular method for detection of *Bartonella*, based on the 16S-23S rRNA intergenic spacer and 16S rRNA amplification by multiplex PCR combined with reverse line blotting, was designed. This assay could simultaneously detect 20 different known species and other *Bartonella* species not described previously.

*Bartonella* species are gram-negative facultative intracellular bacteria that can infect erythrocytes and endothelial cells (7). These zoonotic pathogens have complex cycles in the nature, which include different reservoir hosts and arthropod vectors. *Bartonella* species have been classified as reemerging bacteria with a worldwide distribution, except for *B. bacilliformis*, which is located in northwestern South America.

More than 20 different *Bartonella* species have been described so far, and 8 of them have been associated with human disease (14). However, new species have been identified recently as human pathogens, such as *B. alsatica* (13), *B. koehleae* (3), and *B. rochalimae* (5), as a result of the implementation of more efficient molecular tools for diagnosis. *Bartonella* spp. can produce a wide variety of clinical manifestations, including severe ones such as encephalitis or endocarditis (14). In fact, the species of this genus are a major cause of culture-negative endocarditis, representing in some studies up to 10% of all the cases (12, 16).

The goal in clinical and environmental studies is not only the identification of positive samples but also the identification of the species involved. Therefore, we describe a new molecular method, based on multiplex PCR combined with reverse line blotting (RLB), for the simultaneous detection of 20 different *Bartonella* species.

The targets selected for the method were the conservative 16S rRNA, as a generic target for detecting any *Bartonella* species, and the hypervariable intergenic transcribed spacer 16S-23S rRNA (ITS), which allows distinguishing among the different *Bartonella* species (4, 9). Also, an internal amplification control (IAC), based on delta-9-tetrahydrocannabinolic acid synthase of *Cannabis sativa*, was added to evaluate the presence of PCR inhibitors (8).

To design primers and probes (Table 1), available sequences were retrieved from GenBank and were aligned by using ClustalX (6). The ITSs of *B. chomelii* and *B. capreoli* were sequenced from strains of the collection of the Institute Pasteur, i.e., *B. chomelii* A828 (GenBank accession no. EU098133) and *B. capreoli* IBS 193 (GenBank accession no. EU098130 and EU098131). Interestingly, the latter species had two different ITS sequences, which differed in a 12-bp repetition and were determined after cloning the obtained amplicons from a single colony of *B. capreoli*. Regions of interest, between 18 and 24 bp long and with melting temperatures above 60°C, were identified by visual analysis. Their feasibility for use as primers and probes was checked with Oligo6 software (Molecular Biology Insights, Inc., West Cascade, CO). The Basic Local Alignment Search Tool (BLASTn) (1) was used for a preliminary assessment of the oligonucleotide specificity. A generic probe for all species was designed based on 16S rRNA, and 17 specific probes were selected from ITSs (Table 1). Given the high homology of the ITS sequences between *B. chomelii*, *B. schoenbuchensis*, *B. capreoli*, and *B. birtlesii*, a common probe for the four species was designed (S-CHOSCA [Table 1]). The probe and primers for the IAC used in this study have been described previously (8).

The primers were combined in a multiplex PCR, which was performed in a 50-μl reaction volume with 10 mM Tris-HCl, 50 mM KCl, 2 mM MgCl2, 200 μM of each deoxynucleoside triphosphate (Promega, Madison, WI), 1.5 U of Taq Gold DNA polymerase (Applied Biosystems, Branchburg, NJ), and 0.8 μg/μl of DNase-free bovine serum albumin (Amershams Biosciences, Barcelona, Spain). Primers were used at final concentrations of 0.4 μM for the ITS (Bart/16-23F and Bart/16-23R) and 1 μM for the other two sets of oligonucleotides (16S rRNA and IAC). PCR cycling included an initial denaturing step of 9 min at 94°C, followed by 40 cycles of 30 s at 94°C, 1 min at 64.3°C, and 1 min 30 s at 72°C and a final elongation step of 7 min at 72°C. The amplification was performed in an MJ Research PCT-200 (Ecogen, Barcelona, Spain). The obtained amplicons (438 bp for 16S rRNA, 371 for the IAC, and 176 to 452 bp for the ITS) were further analyzed by RLB. The combination of PCR amplification and hybridization with specific probes allows us to avoid the interference of nonspecific amplifications due to close ubiquitous organisms, such as *Mesorhizobium*, which was recently described as a cause of difficulty in the detection of *Bartonella* by PCR (10).

The RLB was performed as previously described (8) with minor modifications as follows: 3.2 or 0.4 pmol/μl of each
probe (Table 1) was attached to the membrane, the hybridization was performed at 50°C for 1 h, and the washing steps were done at 44°C. The overall time required for the RLB was 3.5 h.

The specificity of the probes was tested with genomic DNAs from different *Bartonella* species (Fig. 1), obtained after purification with the QIAamp DNA minikit (IZASA S.A., Barcelona, Spain). One hundred genomic equivalents (GE) of each species and 10² copies of the cloned IAC were amplified by PCR, and amplicons were analyzed by RLB. Positive hybridization signals were obtained with the IAC and the generic probe for all the samples, as well as for each specific probe (Table 1). In clinical samples, we detected 10 GE of *B. burgdorferi* subsp. *berkhoffii* (Fig. 1A, lanes 16 to 18). It has been suggested that the simultaneous presence of different *Bartonella* species in the same sample could be underestimated using the current diagnostic methods (11). Cases of mixed infections with *B. vinsonii* subsp. *berkhoffii* and *B. henselae* or *B. quintana* have been described. Using our method, we have observed a preferential amplification of *B. henselae* or *B. quintana* over *B. vinsonii* subsp. *berkhoffii* (data not shown).

Finally, the sensitivity of the technique was determined with 10² copies of IAC plus 10³, 10², 10, and 1 GE of *B. schoenbuchensis* (Fig. 1B, lane 1 to 4). The test was repeated in the presence of foreign DNA free of pathogens (300 ng of human DNA [Fig. 1B, lanes 5 to 8] or 300 ng of DNA from an *Ixodes ricinus* specimen [Fig. 1B, lanes 9 to 12]). As expected, in all the samples the IAC was amplified, and the sensitivity for *B. schoenbuchensis* was 10 GE with the 16S rRNA probe and 1 GE with the ITS probe. In the presence of human DNA there was no loss of sensitivity, while with arthropod DNA we lost a logarithmic unit with the ITS probe (Fig. 1B, lane 9 to 12) but still detected until 10 GE with the 16S rRNA probe.

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The method was later validated with clinical and environmental samples (Fig. 1). In clinical samples, we detected *B. henselae* and *B. quintana* in 7% (15/210) and 10% (21/210) of the samples, respectively. In environmental samples, we detected *B. henselae* in 4% (4/100) of the samples. The specificity of the method was tested with 10² copies of IAC plus 10³, 10², 10, and 1 GE of *B. schoenbuchensis* (Fig. 1B, lane 1 to 4). The test was repeated in the presence of foreign DNA free of pathogens (300 ng of human DNA [Fig. 1B, lanes 5 to 8] or 300 ng of DNA from an *Ixodes ricinus* specimen [Fig. 1B, lanes 9 to 12]). As expected, in all the samples the IAC was amplified, and the sensitivity for *B. schoenbuchensis* was 10 GE with the 16S rRNA probe and 1 GE with the ITS probe. In the presence of human DNA there was no loss of sensitivity, while with arthropod DNA we lost a logarithmic unit with the ITS probe (Fig. 1B, lane 9 to 12) but still detected until 10 GE with the 16S rRNA probe.

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quintana and B. henselae from patients with endocarditis and lymphadenopathies (Fig. 1B, lanes 13 to 17). In environmental samples, we detected B. henselae and B. clarridgeiae in cat fleas (Ctenocephalides felis), cats (Felis silvestris), and a wild cat (F. felis), including a mixed infection of both species in a cat (Fig. 1B, lanes 18 to 24 and 33). In small mammals, B. taylorii was detected (Fig. 1B, lanes 25 to 30). However, when the ITS amplicons were sequenced, an undescribed 422-bp ITS fragment (GenBank accession no. EU098135) different from the sequence of the ITS of B. taylorii was detected in two moles (Talpa europaea). In the BLAST server from the NCBI, this fragment showed an 81% homology with an uncultured Bartonella (accession no. AJ269794) which had been detected in a wood mouse (4). The 16S rRNA amplicons from these samples were also sequenced (GenBank accession no. EU098129) and grouped in the dendrogram in the same clade as B. capreoli, B. taylorii, and B. doshiæ (Fig. 2). As it was detected in a mole, it could correspond to B. talpae, although there are currently no available strains or sequences of this species. Consequently, we designed a new probe for this agent, and we cloned the corresponding ITS sequence from a badger (B. taylorii subsp. birtlesii) (GenBank accession no. EU098132) with 88% homology with B. clarridgeiae (GenBank accession no. EU98131) showing a previously undescribed 427-bp sequence (GenBank accession no. EU98130) with 99% homology with B. clarridgeiae, a new species which has recently been detected in a tourist visiting Peru (5). Also, the ITS sequence from a badger (Meles meles) (Fig. 1B, lane 31) had 99% homology with B. rochalimae, a new species which was detected in two moles (B. taylorii subsp. birtlesii) (GenBank accession no. U98127) and 98% homology with B. rochalimae. In the dendrogram (Fig. 2), both 16S rRNA sequences (GenBank accession no. EU98127 and EU98128) from these samples grouped in the same clade as B. clarridgeiae and B. rochalimae.

The molecular method described, which is being patented (2; P. Anda, 29 June 2007, Oficina Española de Patentes y Marcas, pending patent P200701830), has shown an excellent specificity and sensitivity and is a powerful tool for the study of environmental samples and of the etiology of human bartonellosis. Moreover, the technique allows the identification of new
Bartonella species and could be updated progressively with the design of probes for new species, as has been done in the case of the Bartonella detected in small mammals. New probes are being tested for new Bartonella species detected in wild carnivores as well as for other species, such as B. rattimassiliensis or B. phoceensis.

**Nucleotide sequence accession numbers.** The sequences obtained in this study has been submitted to GenBank under the following accession numbers: EU098127 and EU98132 (16S rRNA and ITS of a Bartonella sp. detected in M. meles), EU098128 and EU98134 (16S rRNA and ITS of a Bartonella sp. detected in V. vulpes), EU098129 and EU98135 (16S rRNA and ITS of a Bartonella sp. detected in T. europaea), EU098130 and EU98131 (ITSs from B. capreoli), and EU098133 (ITS from B. chomelii).

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