Clinical Evaluation of a New PCR Assay for Detection of
Coxiella burnetii in Human Serum Samples

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A nested PCR method was developed for the detection of Coxilclaimburnetii in human serum samples. Two pairs of oligonucleotide primers were designed to amplify a 438-bp fragment of the com1 gene encoding a 27-kDa outer membrane protein of C. burnetii. The primers amplified the predicted fragments of 21 various strains of C. burnetii but did not react with DNA samples from other microorganisms. The 438-bp amplification products could be digested with restriction enzymes SspI and SalI. The utility of the nested PCR was evaluated by testing human serum samples. The com1 gene fragment was amplified from 135 (87%) of 155 indirect immunofluorescence test (IF)-positive serum samples and from 11 (11%) of 100 IF-negative serum samples. The nested PCR with primers targeted to the com1 gene appeared to be a sensitive, specific, and useful method for the detection of C. burnetii in serum samples.

Coxiella burnetii is the causative agent of acute Q fever and chronic endocarditis in humans (1). Acute Q fever is a flu-like illness which is self-limiting and which is easily treated with antibiotics when an appropriate diagnosis is made. Chronic Q fever is a severe disease that requires prolonged antibiotic therapy, because the infection can result in endocarditis (14, 18, 24) or granulomatous hepatitis (28). Rapid diagnosis of the disease is very important, because appropriate antibiotic treatment may lead to a better prognosis for individuals suffering from Q fever.

Routine diagnosis of Q fever is usually established by serological tests, since isolation of C. burnetii from patients is time-consuming, difficult, and hazardous. Serological methods, including the indirect immunofluorescence test (IF) (5, 6, 16), complement fixation test (5, 22, 23), enzyme-linked immunosorbent assay (21, 25, 26), and high-density particle agglutination test (17), can be used to detect antibodies to C. burnetii antigens. However, these serological tests have some limitations. Antibodies cannot be detected during the early stage of the infection, and it is difficult to discriminate between current and past infection by a test with a single serum sample, because antibodies often persist after the organisms disappear from the blood. Thus, serological tests offer only a retrospective diagnosis and are useless for the treatment of the afflicted patients.

Recently, PCR has become a useful tool for the detection of C. burnetii in clinical samples (10, 27, 29, 30, 31). The PCR appeared to be a very sensitive method for the laboratory diagnosis of Coxilclaimburnetii infection, able to detect DNA sequences in very small samples. Most recently, we demonstrated that the com1 gene encoding a 27-kDa outer membrane protein (OMP) was highly conserved among 21 strains of C. burnetii from a variety of clinical and geographical sources (32). The com1 gene is the genetic target for the detection of C. burnetii in clinical samples.

In the present study, we have developed a useful nested PCR assay based on the com1 gene sequence for the detection of C. burnetii in human serum samples.

**MATERIALS AND METHODS**

Microorganisms. The microorganisms used in the study included 21 isolates of C. burnetii (strains Nine Mile VR 615, California 76 VR 614, Bangui VR 730, Ohio 314 VR 542, Henzerling VR 145, Priscilla, MAN, ME, GQ212, SQ217, and KoQ229 and 10 Japanese isolates) and 14 other bacterial isolates (Bordetella bronchiseptica GIFU 1127, Chlamydia pneumoniae TW185, Chlamydia psittaci GCP-1, Chlamydia trachomatis E, Escherichia coli C600, Haemophilus influenzae GIFU 3191, Klebsiella pneumoniae GIFU 2926, Legionella pneumophila SL94-1, L. pneumophila SL94-2, Mycoplasma pneumoniae, Orientia tsutsugamushi Karp, O. tsutsugamushi Kato, O. tsutsugamushi Gilliam, and Streptococcus pneumoniae GIFU 8766). The isolates of C. burnetii were propagated in Buffalo green monkey (BGM) cell cultures as described elsewhere (9).

Serum. A total of 255 human serum samples were used in this study (155 IF-positive serum samples and 100 IF-negative serum samples) were selected from among 3,000 samples collected from 1,740 patients between September and December 1995. The patients were from the Gifu University Medical Faculty Hospital, where sera are randomly tested for antibodies to C. burnetii by IF (17). In addition, 50 serum samples from patients with pneumonia of viral or bacterial origin (influenza virus, parainfluenza virus, respiratory syncytial virus, C. psittaci, L. pneumophila, or M. pneumoniae) served as negative controls for the PCR.

DNA extraction. DNA was extracted from the C. burnetii isolates as described previously (8). Briefly, the purified organisms from BGM cell cultures were suspended in TNE buffer (10 mM Tris-HCl [pH 8.0], 300 mM NaCl, 1 mM EDTA) and digested with proteinase K in the presence of 0.1% sodium dodecyl sulfate at 35°C for 60 min. DNA was extracted with phenol, phenol-chloroform, and chloroform; this was followed by ethanol precipitation. Dried under vacuum, the DNA was resuspended in TE buffer (10 mM Tris-HCl [pH 8.0], 1 mM EDTA). The DNA concentration and purity were determined by measuring the optical density at both 260 and 280 nm with a DNA calculator (GeneQuant II; Pharmacia Biotech), and the DNA was kept at −20°C.

Preparation of samples for PCR. The serum samples used for PCR were prepared as described previously (10). Ten microliters of each serum sample was mixed with 40 μl of sample buffer (1% Nonidet P-40, 1% Tween 20, 10 mM Tris-HCl [pH 8.0]), the mixture was boiled for 10 min and then centrifuged at 12,000 × g for 5 min, and the supernatant was directly used for the PCR analysis.

Nucleotide primers. All oligonucleotide primers were obtained from a commercial source (Rikenak Co., Ltd., Nagoya, Japan). The first primer system included primers Q3-Q5 and Q4-Q6, which were designed from the nucleotide sequence of the htpB gene encoding a 62-kDa protein and which were used to specifically amplify 501- and 325-bp fragments (10). The second primer system, including primers OMP1 (5'-AGT AGA AGC ATC ACA GAC ATT G-3'), OMP2 (5'-TGC CTC GTA CTT GAC ACC ATT G-3'), OMP3 (5'-GAA GCC CAA CAA GAA GAA CCC-3'), and OMP4 (5'-TAT GAA GTT ATC ACC CAG TTG-3'), was designed from the nucleotide sequence of the com1 gene encoding a 27-kDa OMP and was used to specifically amplify 501- and 438-bp fragments (7). These primers were designed from a conserved region of the com1 gene of C. burnetii on the basis of the gene sequences of 21 strains (32). The
sequence specificities of these primers were checked by using the sequences in the GenBank database, and no homology with the sequences of other viral or bacterial organisms was detected by a search with the BLAST program.

The nested PCR was performed with serial 10-fold dilutions of total DNA (from 500 ng to 0.5 fg) extracted from the Nine Mile strain of C. burnetii to determine the minimum level of DNA detectable by the assay.

**DNA amplification.** Amplification programs for the primers Q3-Q5 and Q4-Q6 were described previously (10). For the nested PCR with primers OMP1-OMP2 and OMP3-OMP4, the first amplification was performed in a total volume of 50 μl containing 5 μl of DNA sample, 50 mM KCl, 10 mM Tris-HCl (pH 8.3), 1.5 mM MgCl₂, 200 μM (each) dATP, dCTP, dGTP, and dTTP, 0.5 μM primer OMP1, 0.5 μM primer OMP2, and 2 U of Taq DNA polymerase (Takara Shuzo, Co., Ltd., Shiga, Japan). The mixtures were overlaid with 2 drops of mineral oil. PCR was performed at 94°C for 3 min and then for 36 cycles of 94°C for 1 min, 54°C for 1 min, and 72°C for 1 min in a DNA thermal cycler (Perkin-Elmer GeneAmp PCR system 9600; Takara Biomedicals, Kyoto, Japan). In the second amplification, the reaction mixture and conditions were the same as those in the first amplification except for the primers and DNA templates. Primers OMP3 and OMP4 were used at 0.5 μM each, and 1 μl of the first amplification product was used as the DNA template. A positive control with 5 μg of C. burnetii DNA as the template and a negative control without DNA template were included in each PCR run.

**Detection of PCR products.** The PCR-amplified products were examined by electrophoresis in a 1.5% agarose gel, stained with ethidium bromide (0.5 μg/ml), visualized under UV illumination (TM-20; UVP, Inc.) at 320 nm, and photographed.

**Restriction endonuclease digestion.** The products were digested with restriction enzymes known to cut within the target sequence to confirm the identities of the amplified products. The 438-bp amplification products were digested with the restriction enzymes SspI and SalI. One SspI site and one SalI site were present in the amplified region of the com1 gene sequence of C. burnetii. These were compared with the amplification products of reference strains and human serum samples digested with SspI and SalI. The restriction products were also examined by electrophoresis and UV illumination for photography as described above.

**RESULTS**

**Specificity of the nested PCR.** The primers OMP1-OMP2 and OMP3-OMP4 amplified the predicted products of the 501-bp DNA in the first amplification and the 438-bp DNA in the second amplification of PCR with DNA templates from all 21 of the isolates of C. burnetii used. No products were amplified when the DNAs from the 14 other microorganisms and negative controls were used. The specificities of the newly synthesized primers OMP1-OMP2 and OMP3-OMP4 were further demonstrated by digesting the amplified products from a reference strain of C. burnetii with the restriction enzymes SspI and SalI. Digestion of the first PCR products of 501 bp of DNA with SspI and SalI yielded 318- and 183-bp and 348- and 153-bp fragments, respectively. Digestion of the second PCR products of 438 bp of DNA with SspI and SalI yielded 283- and 155-bp and 313- and 125-bp fragments, respectively (Fig. 1).

**Sensitivity of the nested PCR.** The primers OMP1-OMP2 and OMP3-OMP4 amplified the predicted products in reactions with about 5 fg of total DNA (corresponding to 1 organism) (Fig. 2A and B). The primers Q3-Q5 and Q4-Q6 amplified the predicted products in reactions with about 500 fg of total DNA (corresponding to 100 organisms) (Fig. 2C and D).

**Detection of C. burnetii DNA sequences in human serum samples.** The efficacies of the two primer systems for the detection of C. burnetii DNA in human serum samples were compared (Table 1). The primers Q3-Q5 and Q4-Q6 amplified the predicted products with DNA templates from 86 of 255 serum samples. The primers OMP1-OMP2 and OMP3-OMP4 amplified the predicted products with DNA templates from 146 of 255 serum samples (Fig. 3). Among sera positive by IF, 55.5% (86 of 155) were positive with primers Q3-Q5 and Q4-Q6, 32.3% (153 of 475) were positive with primers OMP1-OMP2 and OMP3-OMP4, while 31.6% (49 of 155) were positive with primers OMP1-OMP2 and OMP3-OMP4 but negative with primers Q3-Q5 and Q4-Q6. Among the IF-negative sera, none were positive with primers Q3-Q5 and Q4-Q6, but 11% (11 of 100) were positive with primers OMP1-OMP2 and OMP3-OMP4.

![FIG. 1. Analysis of the restriction endonuclease profile of the 438-bp amplification products of 10 reference strains of C. burnetii. (A) The amplification products were digested with SspI, electrophoresed on agarose gels, and stained with ethidium bromide. Lane 1, molecular size markers (100-bp DNA ladder); lanes 2 to 8, seven reference strains (Nine Mile VR 615, Priscilla, MAN, ME, GQ212, SQ217, and KoQ229, respectively); lanes 9 to 11, three Japanese isolates (307, 605, and TK-1, respectively); lane 12, negative control. (B) The amplification products were digested with SalI. The samples in lanes 2 to 12 are the same as those in panel A. The numbers on the right are in base pairs.](http://jcm.asm.org)
primers Q3-Q5 and Q4-Q6 for the detection of *C. burnetii* in serum samples.

Comparison of the nested PCR results with those of IF indicated agreement for most of the serum samples, but discrepant results were found for some serum samples. We found that 11 of 100 IF-negative serum samples were PCR positive, probably resulting from the failure of IF to detect antibodies in serum samples collected during the early stage of infection. This possibility was further supported by the occurrence of *C. burnetii* in the serum samples obtained during the acute phase (2), during which antibodies were sometimes undetectable by IF (12, 22). We have also found that some serum samples from patients with the acute phase of Q fever were PCR positive but IF negative (10). These results suggest that the nested PCR is more sensitive than IF for the primary diagnosis of acute Q fever.

Antibodies against *C. burnetii* often persist for long periods after the organisms disappear from the blood of Q fever patients who are convalescing or receiving antibiotic therapy (4, 6). Musso and Raoult (15) also indicated that *C. burnetii* could not be isolated from the blood of similar patients by using cell culture. Hence, in our present study, the occurrence of 20 PCR-negative samples among 155 IF-positive serum samples may be explained by antibody persistence in convalescing patients who are devoid of *C. burnetii* at levels above the detection limits of the PCR assay. Therefore, the PCR results appear to indicate the presence or absence of the *com1* gene fragment of *C. burnetii* in the blood, so PCR may be used to evaluate the efficacy of antibiotic therapy or optimize the antibiotic regimen for Q fever patients.

Our results also indicated that a high level of antibody detected by IF was correlated with the presence of *C. burnetii* DNA sequences in human serum samples. This observation is not surprising, since antibodies apparently do not play a direct role in resistance to *C. burnetii* infections (11) or prevent the occurrence of chronic disease (19, 20). Blood culture and serology can be positive at the same time in patients with acute Q fever, and *C. burnetii* can persist in patients for long periods, despite the presence of high levels of antibodies (15). Kazar et al. (13) also demonstrated that immune sera containing either phase II antibody or both phase I and phase II antibodies did not neutralize the organisms.

The results of this study suggest that the nested PCR with primers targeted to the *com1* gene is highly specific and sen-

TABLE 1. Comparison of IF and nested PCR assay results with two sets of primers for detection of *C. burnetii* in human serum samples

<table>
<thead>
<tr>
<th>IF result</th>
<th>No. of serum samples with the indicated result by PCR with the following:</th>
<th><em>htpB</em> gene&lt;sup&gt;a&lt;/sup&gt;</th>
<th><em>com1</em> gene&lt;sup&gt;b&lt;/sup&gt;</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Total</td>
<td>Positive</td>
<td>Negative</td>
</tr>
<tr>
<td>Positive</td>
<td>155</td>
<td>86</td>
<td>69</td>
</tr>
<tr>
<td>Negative</td>
<td>100</td>
<td>0</td>
<td>100</td>
</tr>
<tr>
<td>Total</td>
<td>255</td>
<td>86</td>
<td>169</td>
</tr>
</tbody>
</table>

<sup>a</sup> Detected by nested PCR with primers Q3-Q5 and Q4-Q6.<br><sup>b</sup> Detected by nested PCR with primers OMP1-OMP2 and OMP3-OMP4.
sitive for the detection of C. burnetii, and it may be useful for laboratory diagnosis and assessment of the efficacy of antibiotic therapy for Q fever. Particularly in combination with IF, it may provide a more reliable yet quick means of diagnosing Q fever. We suggest that the clinical diagnosis of Q fever could be made on the basis of both the results of IF and the results of PCR for the detection of C. burnetii in serum samples.

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