Blood culture-negative endocarditis Improving the diagnostic yield using new diagnostic tools

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Improving the diagnostic yield using new diagnostic tools

Blood culture-negative endocarditis (BCNE) may represent up to 70% of all endocarditis cases, depending on series. From 2001 to 2009, we implemented in our laboratory a multimodal diagnostic strategy for BCNE that included systematized testing of blood, and when available, valvular biopsy specimens using serological, broad range molecular, and histopathological assays. A causative microorganism was identified in 62.7% of patients.

In this study from January 2010 to December 2015, in an effort to increase the number of identified causative microorganisms, we prospectively added to our diagnostic protocol specific real-time (RT) polymerase chain reaction (PCR) assays targeting various endocarditis agents, and applied them to all patients with BCNE admitted to the 4 public hospitals in Marseille, France.

A total of 283 patients with BCNE were included in the study. Of these, 177 were classified as having definite endocarditis. Using our new multimodal diagnostic strategy, we identified an etiology in 138 patients (78.0% of cases). Of these, 3 were not infective (2.2%) and 1 was diagnosed as having Mycobacterium bovis BCG endocarditis. By adding specific PCR assays from blood and valvular biopsies, which exhibited a significantly greater sensitivity $(P < 10^{-5})$ than other methods, causative agents, mostly enterococci, streptococci, and zoonotic microorganisms, were identified in an additional 27 patients (14 from valves only, 11 from blood only, and 2 from both). Finally, in another 107 patients, a pathogen was detected using serology in 37, valve culture in 8, broad spectrum PCR from valvular biopsies and blood in 19 and 2, respectively, immunohistochemistry from valves in 3, and a combination of several assays in 38.

By adding specific RT-PCR assays to our systematic PCR testing of patients with BCNE, we increased the diagnostic efficiency by 24.3%, mostly by detecting enterococci and streptococci that had not been detected by other diagnostic methods, but also agents requiring specific management such as Mycoplasma hominis and Tropheryma whippelii.

**Abbreviations:** BCNE = blood culture-negative endocarditis, CIED = cardiovascular implantable electronic device, LCSF = LightCycler SeptiFast, PCR = polymerase chain reaction, RT-PCR = real time polymerase chain reaction.

**Keywords:** blood, diagnosis, endocarditis, serology, specific PCR, valve

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**1. Introduction**

Blood culture-negative endocarditis (BCNE), that is, endocarditis in which blood cultures using usual laboratory methods remain sterile, may account for 2.5% to 70% of all cases of endocarditis, depending on countries. This geographical variation in incidence may be explained by several factors including: differences in the diagnostic criteria used; specific epidemiological factors, as is the case for fastidious zoonotic agents; variations in early use of antibiotics prior to blood sampling; differences in sampling and testing strategies; and involvement of unknown pathogens or noninfective etiologies.

Our center serves as a reference center for the diagnosis of BCNE. From 2010 to 2015, we received specimens from more than 1500 patients worldwide for the diagnosis of BCNE. In an effort to reduce the proportion of endocarditis with no identified etiology, we have continuously diversified the diagnostic tests used. In addition to systematic serological testing for the detection of fastidious agents, especially Coxiella burnetii and Bartonella spp., we have demonstrated that, when available, valvular biopsies are the most useful specimens with regard to diagnostic performance, notably thanks to histological examination and broad range polymerase chain reaction (PCR). In addition, we have demonstrated that syndrome-based sampling and testing was particularly suited to the diagnosis of BCNE.
In a previous study of 759 BCNE cases, we identified an infectious and a noninfective etiology in 62.7% and 2.5% of cases, respectively, using systematic serological testing as well as broad range PCR from blood and/or cardiac valves. However, in that study, patient recruitment was biased as patients’ specimens were referred to our laboratory from around the world, many for confirmation of a specific diagnosis.

Since 2010, in an effort to reduce the proportion of BCNE with no identified etiology, we added to our diagnostic scheme specific real-time (RT)-PCR assays for common agents of this disease such as Bartonella species, C. burnetti, Enterococcus faecalis, E faecium, Escherichia coli, Mycoplasma hominis, Staphylococcus aureus, streptococci from the gallolyticus and oralis groups, and Tropheryma whippelii. In this study, we report the results of prospective testing of all patients admitted to Marseille University Hospitals with a suspected diagnosis of BCNE from 2010 to 2015 using systematic specific RT-PCR assays from valves or blood.

2. Methods

2.1. Patients

From January 1st, 2010 to December 31st, 2015, we prospectively included all patients admitted with suspected BCNE to the 4 university hospitals in Marseille, France. Blood cultures were considered negative when no microorganism grew after 5 days of incubation. For each studied patient, a questionnaire was completed by the physician in charge. Requested information included age, sex, the involved cardiac valve and its type (native or prosthesis, and the type of preexisting valvular defect, if any), contact with cats or body lice, drug abuse, immunodeficiency (and its type), antibiotic uptake prior to blood cultures, clinical and echocardiographic data used in the Duke score, clinical and biological data used in the Marseille score, antibiotic treatment, and outcome (valvular surgery, recovery, and death). Signed informed consent was obtained from all patients. Systematic testing was performed for all BCNE patients using a diagnostic kit and, when available, the patients’ valvular biopsies. The study was approved by the ethics committee of the Institut Federatif de Recherche 48 under reference 07-015.

2.2. Diagnostic procedures

2.2.1. Serology. Indirect immunofluorescence assays to detect significant levels of antibodies to C burnetti (phase I IgG titer ≥1:800), Bartonella quintana, B henselae (IgG ≥ 1:800), and Legionella pneumophila (total antibody titer ≥ 1:256) were performed as previously described. Specific antibodies to Brucella melitensis and Mycoplasma pneumoniae were detected with an immunoenzymatic antibody test (titer ≥ 1:200) and the Platellia M pneumoniae IgM kit (BioRad, Marnes-la-Coquette, France), respectively. When 1st rank tests were negative, we systematically performed a Western blot using Bartonella spp antigens as previously described.

2.2.2. Detection of auto-antibodies. The presence of rheumatoid factor, antinuclear antibodies, and anti-DNA antibodies was determined using the Rheumatoid Factor IgM kit (Orientec, Trappes, France), ANA Hep2 kit (BMD, Marne-la-Vallee, France), and MuST Connective kit (Inodiag, Signes, France), respectively.

Patients with porcine bioprostheses were systematically tested for total (TlgE) and specific immunoglobulin E to pork (SlgEp) using the FEIA ImmunoCAP kit (Phadia, Sweden).

2.2.3. Molecular detection methods. Bacterial DNA was extracted from surgically excised valves or EDTA blood, when no valve was available, using the QIAmp Tissue kit (QIAGEN, Hilden, Germany) as described by the manufacturer. Prosthetic valvular material for which no tissue was available for direct DNA extraction or histological examination (eg, some of the cardiovascular implantable electronic device [CIED] leads) were vortexed in 10 mL of sterile tryptase soy broth, and the DNA was extracted from this suspension. PCR and RT-PCR primers and targets are detailed in Table 1.

2.2.4. Histopathology. Paraffin-embedded heart valves were examined with hematoxylin-eosin for histopathologic features.

To detect microorganisms within tissues, the Giemsa, Gram (Brown-Brenn and Brown-Hopps), periodic-acid Schiff, Grocott-Gomori, Warthin-Starry, Gimenez, and Ziehl-Nielsen stains were systematically performed as described elsewhere. Bartonella henselae and quintana, and C burnetti and T whippelii were tested for in valvular specimens with immunohistochemistry using specific polyclonal antibodies as previously described for patients for whom all other techniques remained negative, we performed auto-immunohistochemistry as previously described.

2.2.5. Statistical methods. All comparisons were performed using the Mantel-Haenszel chi-square test and the EPI info software (version 3.3.2) (http://www.cdc.gov/epiinfo/index.htm). Observed differences were considered significant when P was <.05 for 2-tailed tests.

3. Results

3.1. Patients

Over the study period, 918 patients admitted to Marseille public hospitals were diagnosed as having definite or possible endocarditis, according to the modified Duke criteria. Of these, blood cultures were positive in 635 patients (69.2%) (Table 2). Of the remaining 283 patients, 177 (62.5%) were classified as having definite endocarditis, including 138 in whom our diagnostic strategy (Fig. 1) allowed the identification of an etiology (Table 2, Figs. 1 and 2, Supplemental digital content, http://links.lww.com/MD/B950). This classification was based on pathological criteria in 114 patients, an association of 2 major Duke criteria in 30 patients, 1 major and 3 minor criteria in 26 patients, and 5 minor criteria in 7 patients. The remaining 106 patients (37.5%) were classified as having possible endocarditis. The mean age (±standard deviation) of these 177 patients was 65.3 ± 15.3 years old (range 1–100 years old). Their sex ratio (M/F) was 131/46. Seven patients were admitted to the Nord hospital and the remaining 170 to the Timone hospital (Supplemental digital content, http://links.lww.com/MD/B950). A total of 113 patients had received antibiotics prior to blood culture collection (Supplemental digital content, http://links.lww.com/MD/B950). Valvular specimens were available for 119/177 patients (67.2%), including native valves in 93 patients, bioprosthetic valves in 8, mechanic prostheses in 7, and CIED leads in 11. Both EDTA blood and serum samples were available for all 177 patients. The involved valves were the aortic, mitral, tricuspid, both the aortic and mitral valves, and a CIED in 83, 70, 4, 4, and 16 cases, respectively.

3.2. Diagnostic procedures

The overall sensitivity of our diagnostic strategy (138 etiologies identified/177 patients with definite BCNE, Fig. 2) was 78.0%.
### Table 1

Primer name | Nucleotide sequence (5'-3') | Microorganisms detected | Molecular target | Reference |
---|---|---|---|---|
536F | CAGCAGCCCGCGGTAATAC | All bacteria | 16S rRNA | [11] |
RP2F | AGTGCTCTCTGCAAGCTTCT | | | |
CUEF | TCCCTAGGGTAGAACCTCGG | | | |
CUEF | GCTGCGTTCTTCTGAAGTC | | | |
SI111F | CAGGACGGATCTCTGTCG | C. burnetii | htpAB-associated element | [13] |
SI111R | CAGGAGACGACCACTGATG | | | |
ITSF | GGGCCCGTACCGCAACTG | Bartonella sp | 16S-23S rRNA spacer | [14] |
ITSR | TTAATATATCTTCTCTCATCAGTTC | | | |
ITSprobe | CCAGACATGCGCAAAAT | | | |
199F | GCCACAGGGGCACTGG | | | |
492R | AACCCCTGCTGACCC | | | |
TWprobe | CTTTGTTATGAGATATCTTCTTCATCTTCC | | | |
Etacaeal_F | GTATCGCGGACTCTGAGCC | Enterococcus faecalis | recN | Present study |
Etacael_R | CATGTCGATTCTTGGCGGA | | | |
Etacael_P | AGTCAGAAGGAGAACAA | | | |
Etacalium_F | GGAGAATCTGCGCAACAAAT | Enterococcus faecium | sodA | Present study |
Etacalium_R | CCCATCGAAGAACATCTG | | | |
Etacalium_P | GCTGCGTGGAAGCTACAGAGAA | Staphylococcus aureus | nucA | Present study |
NucAMGBf | TGATCCTCTTTGCAAGATG | Streptococcus oralis-group | Gdh | Present study |
NucAMGBd | CGAGTTCGAAACAATGAGGGCTG | | | |
Efaecalis_F | GTATCGCGGACTCTGAGCC | Enterococcus faecalis | recN | Present study |
Efaecalis_R | CATGTCGATTCTTGGCGGA | | | |
Efaecalis_P | AGTCAGAAGGAGAACAA | | | |
Efaecium_F | GGAGAATCTGCGCAACAAAT | Enterococcus faecium | sodA | Present study |
Efaecium_R | CCCATCGAAGAACATCTG | | | |
Efaecium_P | GCTGCGTGGAAGCTACAGAGAA | Staphylococcus aureus | nucA | Present study |
S_MOS_gdh_F | AYAGCATGATGHTCCCTG | Streptococcus oralis-group | Gdh | Present study |
S_MOS_gdh_R | GASTCCATYTGGTTTAAACG | | | |
S_MOS_gdh_P | TTCTTGGGGTCTGTAAGCTCTVA | | | |
bovis_16S_F | TTTAACMCATTGATAGTCCTGGAAG | Streptococcus gallolyticus-group | 16S rRNA | Present study |
bovis_16S_R | GTAGGAGTCTGGGCCGTGGTC | | | |
bovis_16S_P | GCCCATCGCTTTCTGACAACAC | | | |
ECOmpGMGBAluf | GCTGCGTGGAAGCTACAGAGAA | Escherichia coli | ompG | Present study |
ECOmpGMGBAlur | CATGTCGATTCTTGGCGGA | | | |
ECOmpGMGB | CATGTCGATTCTTGGCGGA | | | |
MHMG16Sd | GTGATATGCGGAGAACCATTTCAAT | Mycoplasma hominis | 16S rRNA | Present study |
MHMG16Sf | GCCATCGGCTTCTGCAAGGG | | | |
MHMG16Sf | AATGATTTGACGATGAC | | | |

| Primer name | Nucleotide sequence (5'-3') | Microorganisms detected | Molecular target | Reference |
---|---|---|---|---|
NucAMGBf | TGATCCTCTTTGGA | Streptococcus oralis-group | Gdh | Present study |
NucAMGBd | CGAGTTCGAAACAATGAGGGCTG | | | |
Efaecalis_F | GTATCGCGGACTCTGAGCC | Enterococcus faecalis | recN | Present study |
Efaecalis_R | CATGTCGATTCTTGGCGGA | | | |
Efaecalis_P | AGTCAGAAGGAGAACAA | | | |
Efaecium_F | GGAGAATCTGCGCAACAAAT | Enterococcus faecium | sodA | Present study |
Efaecium_R | CCCATCGAAGAACATCTG | | | |
Efaecium_P | GCTGCGTGGAAGCTACAGAGAA | Staphylococcus aureus | nucA | Present study |
S_MOS_gdh_F | AYAGCATGATGHTCCCTG | Streptococcus oralis-group | Gdh | Present study |
S_MOS_gdh_R | GASTCCATYTGGTTTAAACG | | | |
S_MOS_gdh_P | TTCTTGGGGTCTGTAAGCTCTVA | | | |
bovis_16S_F | TTTAACMCATTGATAGTCCTGGAAG | Streptococcus gallolyticus-group | 16S rRNA | Present study |
bovis_16S_R | GTAGGAGTCTGGGCCGTGGTC | | | |
bovis_16S_P | GCCCATCGCTTTCTGACAACAC | | | |
ECOmpGMGBAluf | GCTGCGTGGAAGCTACAGAGAA | Escherichia coli | ompG | Present study |
ECOmpGMGBAlur | CATGTCGATTCTTGGCGGA | | | |
ECOmpGMGB | CATGTCGATTCTTGGCGGA | | | |
MHMG16Sd | GTGATATGCGGAGAACCATTTCAAT | Mycoplasma hominis | 16S rRNA | Present study |
MHMG16Sf | GCCATCGGCTTCTGCAAGGG | | | |
MHMG16Sf | AATGATTTGACGATGAC | | | |

1 Forward primer.
2 Reverse primer.
3 5'-FAM-3'TAMRA probe.
4 5'-FAM-3'Mgb probe.

### Table 2

Comparison of microorganisms identified in patients with positive (n = 635) or negative (n = 283) blood cultures.

| Microorganism | Positive blood culture, % | Negative blood culture, % | $P^*$ | Odds ratio (95% CI) |
---|---|---|---|---|
**Intracellular bacteria**
Bartonella sp | 0 | 19 (6.7) | <.01 | Undefined |
Coxiella burnetii | 0 | 23 (8.1) | <.01 | Undefined |
T whipplei | 0 | 3 (1.1) | .03 | Undefined |
**Gram-positive bacteria**
Enterococcus sp | 90 (14.2) | 15 (5.3) | <.01 | 0.34 (0.18–0.42) |
Streptococcus sp | 206 (32.4) | 24 (8.9) | <.01 | 0.19 (0.12–0.31) |
Staphylococcus sp | 266 (41.9) | 31 (10.9) | <.01 | 0.17 (0.11–0.26) |
Other gram-positive bacilli | 9 (1.4) | 15 (5.3) | <.01 | 5.05 (1.9–13.85) |
**Gram-negative bacteria**
HACEK bacteria | 6 (0.9) | 1 (0.3) | .67 | 0.38 (0.02–3.36) |
Other gram-negative bacteria | 47 (7.4) | 1 (0.3) | <.01 | 0.04 (0.00–0.30) |
**Other microorganisms**
Other bacteria | 4 (0.6) | 2 (0.7) | 1.0 | 1.28 (0.15–9.45) |
Fungi | 7 (1.1) | 1 (0.3) | .43 | 0.32 (0.01–2.66) |
Total | 635 | 135 |

*Confidence interval.
*Statistically significant $P$ values are indicated in bold characters.
*One patient was diagnosed as having an allergy to pork, 1 as suffering from marantic endocarditis, and 1 as having Loeffler endocarditis.

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**Table 2**

Comparison of microorganisms identified in patients with positive (n = 635) or negative (n = 283) blood cultures.

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T whipplei | 0 | 3 (1.1) | .03 | Undefined |
**Gram-positive bacteria**
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Staphylococcus sp | 266 (41.9) | 31 (10.9) | <.01 | 0.17 (0.11–0.26) |
Other gram-positive bacilli | 9 (1.4) | 15 (5.3) | <.01 | 5.05 (1.9–13.85) |
**Gram-negative bacteria**
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Other gram-negative bacteria | 47 (7.4) | 1 (0.3) | <.01 | 0.04 (0.00–0.30) |
**Other microorganisms**
Other bacteria | 4 (0.6) | 2 (0.7) | 1.0 | 1.28 (0.15–9.45) |
Fungi | 7 (1.1) | 1 (0.3) | .43 | 0.32 (0.01–2.66) |
Total | 635 | 135 |
Serology using immunofluorescence assay (Fig. 1) provided a diagnosis in 41/177 patients (23.2%) (Fig. 2). Chronic Q fever (IgG titer to phase I C. burnetii ≥1:800) was diagnosed in 23 patients. Seventeen patients had an IgG titer to B. quintana and/or B. henselae ≥1:800. Of these, cross-adsorption was able to identify 13 B. quintana and 4 B. henselae infections. Lastly, 1 patient had positive L. pneumophila serology, with an Ig titer of 1:512.

Broad spectrum and specific PCR assays from EDTA blood detected pathogens in 3/177 (1.7%) and 24/177 (13.5%) patients, respectively, for a total of 27 diagnoses (Fig. 1, Supplemental digital content, http://links.lww.com/MD/B950). When applied to valvular specimens, broad spectrum and specific PCR assays were positive in 52/119 (43.7%) and 45/119 (37.8%) patients, respectively, for a total of 85 diagnoses (14% overlap). Negative controls were negative in all assays. Overall, broad range PCR identified a causative agent in 51 patients and specific PCR in 56 (12.1% overlap, for a total number of PCR-positive patients of 99). Culture from valvular biopsies or CIED leads was positive in 41 patients (34.4%), but did not detect any microorganism that had not been identified by PCR (Supplemental digital content, http://links.lww.com/MD/B950). Immunohistochemistry

Figure 1. Proposed diagnostic strategy for patients with blood culture-negative endocarditis (BCNE).

Figure 2. Distribution of identified etiological agents according to the diagnostic method used. The percentages of positive specimens per diagnostic method are indicated in parentheses. Etiological agents identified using newly added specific polymerase chain reaction (PCR) assays are indicated in red.
the positivity of valves from patients infected with *B. henselae*, *B. quintana*, and *C. burnetii*, and with *T. whipplei*. Although useful in previous studies, auto-immunohistochemistry did not provide any additional diagnosis in this series.

The microorganisms detected using culture and/or PCR included * Bartonella* species in 4 patients, * C. burnetii* in 4, * T. whipplei* in 3 (Fig. 1, Supplemental digital content, http://links.lww.com/MD/B950), * Enterococcus* spp in 13, * S. aureus* in 12, other staphylococci in 19, streptococci from the *gallolyticus* and *oralis* groups in 12 and 4, respectively, other streptococci in 7, other gram-positive germs in 15 (*Actinobacillus defectiva, Corynebacterium jeikeium, Gemella morbillorum, Granulicatella adiacens, and Propionibacterium spp*), gram-negative bacteria in 1 (*Haemophilus parainfluenzae*), other bacteria in 1 (*M. hominis*), and fungi in 1 (*Candida albicans*). In addition, in an 86-year-old male who developed aortic BCNE following intrabladder BCG instillations for bladder cancer, specific blood cultures for mycobacteria in Middlebrook liquid medium grew *M. bovis* BCG. With the exception of this latter case, cultures did not provide any diagnosis that was not made by another method. A 53-year-old male with recurrent mitral BCNE episodes and who had undergone 4 valve replacements with porcine bioprostheses was diagnosed as having allergy to pork[20] by the absence of anti-porcine IgG. A 53-year-old male with a history of chronic pulmonary BCNE was also diagnosed as suffering from eosinophilic leukemia complicated with Loeffler endocarditis as confirmed by histopathological examination of the removed valve.

Although broad spectrum and specific PCR assays from blood and valves were complementary, each being the only one to identify an etiological agent in 2, 12, 11, and 14 patients, respectively (Supplemental digital content, http://links.lww.com/MD/B950), broad range PCR from blood was statistically less sensitive than all other diagnostic methods used (Table 3). Similarly, specific PCR from blood provided significantly fewer diagnoses than serology and culture and PCR from valves (Table 3). In contrast, culture, broad range, and specific PCR from valve samples did not significantly differ (Table 3). *Bartonella* spp and *C. burnetii* were only diagnosed by serology and/or specific PCR and/or immunohistochemistry, and *T. whipplei* by specific PCR and/or immunohistochemistry. Among the 177 patients with definite BCNE (Supplemental digital content, http://links.lww.com/MD/B950), enterococci were more likely to be detected by specific PCR than culture (15 vs 2, *P* < 10^-2, OR 8.44, range 1.88–37.77) or broad range PCR (15 vs 3, *P* < 10^-2, OR 5.57, range 1.57–19.81). Streptococci were detected significantly more in valves using specific PCR assays as compared to culture (18 vs 4, *P* < 10^-2, OR 5.12, range 1.67–15.64), but not more than broad spectrum PCR (18 vs 12, *P* = .32, OR 1.58, range 0.73–3.46), although the detected species differed. Streptococci from the *gallolyticus* and *oralis* groups were detected more by specific than broad range PCR (18 vs 6, *P* = .01, OR 3.35, range 1.28–8.78). The detection of staphylococci using broad spectrum PCR or culture from valvular specimens was not statistically different (23 vs 24, *P* = .88, OR 0.95, range 0.5–1.79). However, *S. aureus* was statistically more identified by specific than broad spectrum PCR (12 vs 3, *P* = .02, OR 4.35, range 1.19–15.78). Thus, overall, specific RT-PCR was significantly more sensitive than broad range PCR (58 vs 12, *P* < 10^-2, OR 8.47, range 4.22–17.0) and valve culture (58 vs 13, *P* < 10^-2, OR 7.75, range 3.93–15.28).

All patients in this series received an empirical antibiotic treatment as per the guidelines of the European Society for Cardiology.[21] In 27 patients, the antibiotic regimen was adapted to the identified etiological agents. These included all patients infected with *C. albicans, C. burnetii, L. pneumophila, M. bovis* BCG, and *M. hominis*. Five patients died within 6 to 30 days following admission. Of these, 4 were male. An etiological agent was identified by PCR in 2 patients, including *E. faecalis* from a valvular biopsy in 1 and *P. acnes* from a CIED lead in the other. Four died from heart failure and 1 from a cerebral embolism despite antibiotic treatment.

### 4. Discussion

We report here the results of the diagnostic strategy that we systematically applied to patients with BCNE from 2010 to 2015. To evaluate the effectiveness of our syndrome-based diagnostic strategy for BCNE without being biased by heterogeneous patient populations from areas with distinct epidemiologies, we only included patients admitted to Marseille public hospitals. Specific PCR assays were designed to target BCNE agents that had either been found to be common in Marseille[3] and/or that were fastidious, such as * Bartonella* spp, *C. burnetii, M. hominis*, and *T. whipplei*.

Of 283 patients with BCNE, 177 patients were diagnosed as having definite endocarditis. An etiological diagnosis was made in 138 patients (78.0%), including 135 in whom a microorganism was identified. Of these, *C. burnetii, Bartonella* spp, and *T. whipplei* accounted for 32.8% of diagnoses and 15.9% of all BCNE cases (Table 2, Fig. 1). Such a high prevalence of these agents when compared to other series (4.9% of all IE cases from 2010 to 2015) may be explained by the endemicity of Q fever in the Marseille area and by the Marseille University hospitals’ role as reference center for these diseases. This may also, at least partially, explain the high rate of BCNE (30.8%) in our IE series during this period. Thus, we acknowledge the fact that the
obtained in one of our previous series (65.2%[3]), in which we 177 patients with de
dsensitivity of our diagnostic strategy (138 etiologies identi
dated, several authors have reported using the commercially
fi
re
half of BCNE patients in this study, is needed to improve
which enabled the identi
methodology. In addition, unlike our specific PCR panel, LCSF does not take into account the specific epidemiology of IE, particularly in our region. When considering the 135 causative microorganisms identified in our series, 91 were in the spectrum of our specific PCR panel versus only 71 for LCSF (P = .01, OR = 0.33, range 0.33–0.88).

Regarding the impact of our diagnostic approach to patient management, the results prompted a specific adaptation of the antibiotic treatment in 27 patients, all of whom recovered from their infections, and a continuation of the empirical treatment in the other 256. Among the latter group, 5 patients died within 30 days of admission, including 2 in whom a microorganism was identified. However, in these 2 patients, the administered empirical antibiotic treatment was active on the detected agents.

5. Conclusion

In summary, we demonstrated that using specific PCR assays targeting the most common pathogens in a given area has the potential to significantly increase the diagnostic yield of BCNE, but that these assays should be included in a global diagnostic strategy involving other methods such as serology, broad range PCR, and valve culture.

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