Evaluation of the FilmArray Blood Culture Identification Panel: Results of a Multicenter Controlled Trial

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Sepsis is a major cause of morbidity, mortality, and increased medical expense. Rapid diagnosis improves outcomes and reduces costs. The FilmArray blood culture identification panel (BioFire Diagnostics LLC, Salt Lake City, UT), a highly multiplexed PCR assay, can identify 24 etiologic agents of sepsis (8 Gram-positive, 11 Gram-negative, and 5 yeast species) and three antimicrobial resistance genes (mecA, vanA/B, and blakpc) from positive blood culture bottles. It provides results in about 1 h with 2 min for assay setup. We present the results of an eight-center trial comparing the sensitivity and specificity of the panel with those of the laboratories’ standard phenotypic identification techniques, as well as with molecular methods used to distinguish Acinetobacter baumannii from other members of the A. calcoaceticus-A. baumannii complex and to detect antimicrobial resistance genes. Testing included 2,207 positive aerobic blood culture samples, 1,568 clinical and 639 seeded. Samples were tested fresh or were frozen for later testing within 8 h after the bottles were flagged as positive by an automated blood culture system. At least one organism was detected by the panel in 1,382 (88.1%) of the positive clinical specimens. The others contained primarily off-panel organisms. The panel reported multiple organisms in 81 (5.86%) positive clinical specimens. The unresolved blood culture identification sensitivity for all target detections exceeded 96%, except for Klebsiella oxytoca (92.2%), which achieved 98.3% sensitivity after resolution of an unavoidable phenotypic error. The sensitivity and specificity for vanA/B and blakpc were 100%; those for mecA were 98.4 and 98.3%, respectively.

Bacteremia and sepsis constitute major health and financial burdens in the United States and internationally (1). In the United States, rates for hospitalization that included the diagnostic codes for sepsis or severe sepsis doubled between 2000 and 2008, partly because of the aging of the population (2, 3). In 2008, hospital costs for the treatment of sepsis were $14.6 billion, and the inflation-adjusted cost is increasing by almost 12% annually. Treatment of sepsis is complicated by the continuing increase in antibiotic resistance (4).

A definitive diagnosis of a bacteremic/septic episode is made from blood cultures. They consist of an aerobic and an anaerobic blood culture bottle inoculated with blood from the patient and incubated in an automated blood culture instrument. When the bottles are flagged as positive, identification (ID) and antibiotic susceptibility testing (AST) of the etiologic agents generally require 2 or more additional days. Patients who are treated rapidly with appropriate antibiotics have better outcomes, with decreased morbidity and mortality rates, hospital stay lengths, and hospital costs (5). Excess mortality attributable to inadequate antimicrobial therapy ranges from 10 to almost 40% (6, 7). Since decreasing the time to administration of appropriate antibiotics improves survival and decreases costs (6, 8–10), a goal of the clinical microbiology laboratory is to expedite organism ID and AST results. Molecular techniques play an increasing role in speeding these determinations (9, 11–13).

Between July 2012 and February 2013, eight clinical microbiology laboratories participated in a study of the BioFire FilmArray blood culture ID (BCID) system. BCID is a two-stage, highly multiplexed, nested PCR test that is carried out in a closed, disposable, single-use pouch. It requires about 2 min for assay setup and provides results in approximately 1 h. It is designed to identify simultaneously 24 etiologic agents of sepsis (eight Gram-positive, 11 Gram-negative, and five Candida species) as well as three antimicrobial resistance genes (mecA, vanA/B, and blakpc) which encode Klebsiella pneumoniae carbapenemase. In general, the comparator assays were the standard phenotypic organism ID and antibiotic susceptibility testing methods employed by each participating laboratory. The clinical specimens consisted of 2,207 positive aerobic blood culture samples tested within 8 h of bottle inoculation. After bacterial growth was confirmed by the BCID system, the sample was inoculated into an automated blood culture instrument. Identification of the etiologic agent and antibiotic resistance profiles were determined in parallel by the BCID system and comparator methods. The study was conducted by the Microbiology Assessment Network (MANET) of the American Society for Microbiology. The study protocol was approved by the institutional review boards of all participating sites. A quality assurance program was used to ensure the accuracy and reliability of the data. The study was performed in compliance with federal regulations and HIPAA guidelines for the protection of patient privacy.

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methods utilized by the eight participating laboratories. However, molecular techniques were used for confirmation of *Acinetobacter baumannii* to the species level and detection of the antibiotic resistance genes. The results of this study were submitted to the U.S. Food and Drug Administration as part of the information required for the 2013 clearance of the system.

(These results were presented in part in 2013 at the 113th General Meeting of the American Society for Microbiology and at the 23rd European Congress of Clinical Microbiology and Infectious Diseases.)

**Materials and Methods**

### Study Sites

The eight clinical microbiology laboratories that participated in this study were in the following U.S. cities: New York, NY; Baltimore, MD (two sites); Galveston, TX; Chicago, IL; Detroit, MI; Salt Lake City, UT; and Los Angeles, CA.

### Overall Design

The comparison had two arms. The prospective clinical arm tested the residual specimens from diagnostic patient blood cultures and compared them with the results of phenotypic analysis of the same specimens. The seeded arm tested specimens from similar bottles that had been injected with human blood (screened for pathogens such as HIV and the hepatitis B and C viruses; Bioreclamation, Westbury, NY) and a pure culture of a known microorganism. One laboratory tested only seeded cultures, and three analyzed only clinical specimens. The other four performed both types of testing. This study was approved by the Institutional Review Board at each site, and a waiver of informed consent was obtained.

All laboratories used the BD Bactec Plus Aerobic/F blood culture bottle (BD, Sparks, MD). For both the prospective and seeded arms, a positive Gram stain from the flagged blood culture bottle was required before the specimen was included in the study. BCID testing and the freezing of aliquots had to be initiated within 8 h after the bottle was flagged as positive by the automated blood culture system. This time interval was chosen for reproducibility on the basis of preliminary experiments that showed that BCID positivity was stable for at least 12 h. Specimens from both arms of the study were deidentified, coded with unique study numbers, and treated identically once they were flagged as positive by the blood culture instrument. Organisms from all positive blood culture bottles, both clinical and seeded, were subjected to standard blood culture ID and AST. The laboratory ID was generally regarded as the gold standard and as the basis for determination of results as true or false positive (TP or FP) or true or false negative (TN or FN), respectively. However, in the results reported in 1 h, BCID results for a specimen were displayed only if the two internal pouch controls for the run were valid. For general descriptions of the FilmArray system and of the methodology used to develop the primers and probes for BCID and other FilmArray pouches, see references 14 and 15. All target sequences are longer than 200 bp and are proprietary, developed by BioFire for the FilmArray system. RNA was not used as a target in the BCID assay.

BCID targets included the following genera or families of bacteria: *Enterococcus*, *Staphylococcus* (two targets), *Streptococcus*, and members of the family *Enterobacteriaceae*. There were also assays for organisms included in the above groups. Among the staphylococci was *Staphylococcus aureus*; among the streptococci were *Streptococcus* agalactiae, *S. pneumoniae*, and *S. pyogenes*, and the members of the family *Enterobacteriaceae* included the *Enterobacter cloacae* complex, *Escherichia coli*, *Klebsiella oxytoca*, *K. pneumoniae*, *Proteus* species, and *Serratia* species. Other targets for bacteria not included in the above groups included *A. baumannii*, *Haeomophilus influenzae* (two targets) and *Pseudomonas aeruginosa*. Assays for five different *Candida* spp. (*Candida albicans*, *C. glabrata*, *C. krusei*, *C. parapsilosis*, and *C. tropicalis*) were also included.

If any of the three *Staphylococcus* assays (two genus assays or the *S. aureus* assay) was positive, the culture was identified as containing a member of the genus *Staphylococcus*. If the *S. aureus* assay was positive, *S. aureus* was also reported as detected, and the presence/absence of other staphylococci could not be determined. If *S. aureus* was reported as not detected by BCID, a *Staphylococcus* species other than *S. aureus* was assumed. Similarly, if BCID reported that either a *Streptococcus* species or a member of the family *Enterobacteriaceae* was detected and the sample was also reported as positive for a specific organism in that genus or family, the specific organism was presumed to be the one responsible for the positive genus/family BCID result. If a positive result for a specific member of the genus *Streptococcus* or the family *Enterobacteriaceae* was not reported, the analyte was presumed to be a *Streptococcus* species or a members of the family *Enterobacteriaceae* for which a specific assay was not included in the panel. Therefore, as with the staphylococci, the presence of a *Streptococcus* species or a member of the family *Enterobacteriaceae* that did not have a specific assay in the BCID panel could not be ruled either in or out if a related organism with a specific assay was present.

**Bacterial and Fungal Comparator Methods.** For organism ID, with the exception of *A. baumannii*, the reference methods used to evaluate BCID performance in both the prospective and seeded arms of the study were the standard phenotypic ID procedures used in each laboratory. Automated methods included MicroScan Walk-Away (Siemens Medical Solutions, Deerfield, IL), Vitek 2 (bioMérieux, Durham, NC), and Phoenix (BD Diagnostics, Sparks, MD). Testing was performed according to the manufacturers’ instructions. Manual techniques were performed according to the individual laboratory protocols, which conformed to the standards of the Clinical and Laboratory Standards Institute (16).
Phenotypic methods do not separate *A. baumannii* from other members of the *A. calcoaceticus-A. baumannii* complex (17), while BCID specifically targets *A. baumannii*. Therefore, the gold standard for the comparator identification of *A. baumannii* was a two-step process in which the phenotypic ID of a complex member was followed by bidirectional 16S rRNA gene testing to confirm its identity as *A. baumannii sensu stricto*. The 16S rRNA gene PCR was performed by research associates at BioFire who were blind to the original BCID and AST results. Amplicon cleanup and bidirectional sequencing were performed by Macrogen USA, Rockville, MD.

**Antibiotic resistance gene detection methods.** The BCID resistance gene targets are within the sequences of the resistance genes, at least 200 bp in length, and are proprietary. Antibiotic resistance gene results are reported as detected by BCID only if an organism known to carry that gene is also detected: *Staphylococcus* spp. for *mecA*; *Enterococcus* spp. for *vanA/B*; and *A. baumannii*, *P. aeruginosa*, or a member of the family *Enterobacteriaceae* for *blaKPC*. The comparator methods for these resistance genes were molecular rather than phenotypic and consisted of PCR assays performed with aliquots from the frozen blood culture bottles and targeting a sequence of the resistance gene distinct from that of the BCID target. The comparator PCR targets were developed by BioFire in conjunction with the design of their BCID targets; they were different from and did not overlap those of the BCID assays. PCR was performed by research associates at BioFire who were blind to the original BCID and AST results. Amplicon cleanup and bidirectional sequencing were performed by Macrogen USA, Rockville, MD.

Comparison of BCID resistance gene determinations with the results of AST of the isolated organisms by the phenotypic techniques is presented elsewhere (18).

**Investigation of discrepancies.** Each discrepant result occurring between BCID and the comparator methods was investigated at BioFire by technologists who were blind to the results of the original assays. They applied one or more of the following strategies to frozen aliquots of the original blood culture specimens: direct molecular testing of the aliquot or subculturing and molecular testing of an organism grown from the aliquot. The molecular methods used included repeat BCID and/or sequencing of the 16S rRNA gene for bacteria or of the ITS1 region for yeast. Resolution of discrepant determinations from polymicrobial cultures by molecular techniques required isolation of the organisms prior to 16S rRNA gene or ITS1 analysis. Human error leading to a possible sample mix-up was also considered in the investigation of discrepant results. A sample mix-up involving two bottles, each containing one organism, would lead to two FP and two FN results.

**Statistics.** Sensitivity and specificity were determined from standard two-by-two performance tables. Briefly, sensitivity (or positive percent agreement [PPA]) = 100TP/(TP + FN), while specificity (or negative percent agreement [NPA]) = 100TN/(TN + FP). Sensitivity and specificity are applicable to the prospective clinical cultures, and PPA and NPA are applicable to the seeded cultures. Both terms are included in the tables. However, for simplicity, only sensitivity and specificity are used in the text. The binomial two-sided 95% confidence intervals were calculated for both sensitivity and specificity by using the exact method of Clopper and Pearson (19). The number of patients enrolled in the prospective clinical study ensured that organisms with a prevalence rate of >2% could achieve statistical significance. For rare analytes, the seeded specimens ensured an adequate number of samples for analysis. The statistical analysis was conducted separately for each subgroup of each organism or resistance gene (clinical fresh, clinical frozen, overall clinical, seed fresh, seeded frozen, and overall frozen) and for the combined total. Since no significant difference between the categories of each analyte was noted, they were combined in the performance summary tables (see Tables 3, 4, and 6).

**RESULTS**

**Overview.** This study included 2,207 samples, 1,568 clinical and 639 seeded. Specimen testing by BCID was either initiated within 8 h after the bottles turned positive (821 or 52% of the patient specimens and 419 or 66% of the seeded cultures) or performed with aliquots of the fluid from the bottles that had been frozen within the 8-h time window (the remainder). Of the total number of samples tested by BCID, 41 (1.9%) needed repeat testing: 1 was aborted by the operator, 5 had instrument or software errors, internal control failure occurred in 16, and pouch hydration failures occurred in 19. All were successfully retested within the 8-h limit.

**Table 1** shows all of the BCID targets and the number and percentage of each detected by BCID in the prospective clinical specimens from pediatric and adult patients. More than one organism was reported by BCID in 81 (5.2%) of the cultures, including two organisms in 74 cultures, three organisms in five cultures, and four organisms in two cultures.

The data from both prospective clinical samples and seeded specimens are combined (see Tables 3, 4, and 6). The columns marked clinical arm and seeded arm give the numbers of positive specimens determined by BCID over those of the comparator as-

| TABLE 1 FilmArray BCID panel targets and prevalence of positive results by age group for the prospective clinical study arm |
|---------------------------------------------------------------|
| **FilmArray BCID result** | **No. (%) of patients** |
| **Gram-positive bacteria** | **Total** | **Pediatric** | **Adult** |
| Enterococcus | 102 (7) | 5 (3) | 97 (7) |
| Listeria monocytogenes | 0 (0) | 0 (0) | 0 (0) |
| Staphylococcus | 780 (50) | 74 (50) | 706 (50) |
| S. aureus | 257 (16) | 25 (17) | 232 (16) |
| Streptococcus | 140 (9) | 21 (14) | 119 (8) |
| *S. pneumoniae* | 26 (2) | 3 (2) | 23 (2) |
| *S. pyogenes* (GAS) | 8 (1) | 2 (1) | 6 (<1) |
| **Gram-negative bacteria** | | | |
| *A. baumannii* | 16 (1) | 1 (1) | 15 (1) |
| *Enterobacteriaceae* | 307 (20) | 27 (18) | 280 (20) |
| *Enterobacter cloacae* | 24 (2) | 4 (3) | 20 (1) |
| *Escherichia coli* | 149 (10) | 16 (11) | 133 (9) |
| *Klebsiella oxytoca* | 6 (<1) | 1 (1) | 5 (<1) |
| *Klebsiella pneumoniae* | 74 (5) | 5 (3) | 69 (5) |
| Proteus | 22 (1) | 0 (0) | 22 (2) |
| *Serratia marcescens* | 22 (1) | 1 (1) | 21 (1) |
| Haemophilus influenzae | 8 (1) | 3 (2) | 5 (<1) |
| Neisseria meningitidis | 1 (<1) | 0 (0) | 1 (0) |
| *P. aeruginosa* | 52 (3) | 4 (3) | 48 (3) |
| **Yeast** | | | |
| *Candida albicans* | 20 (1) | 1 (1) | 19 (1) |
| *Candida glabrata* | 14 (1) | 0 (0) | 14 (1) |
| *Candida krusei* | 4 (<1) | 1 (1) | 3 (<1) |
| *Candida parapsilosis* | 8 (1) | 0 (0) | 8 (1) |
| *Candida tropicalis* | 3 (<1) | 0 (0) | 3 (<1) |
| **Antimicrobial resistance genes** | | | |
| *mecA* | 491 (31) | 46 (31) | 445 (31) |
| *vanA/B* | 36 (2) | 0 (0) | 36 (3) |
| *blaKPC* | 6 (<1) | 0 (0) | 6 (<1) |

*a Regardless of results of comparator assays.

*b E. cloacae complex. Individual members are not separated.

*c n = 1,568 (100%).

*d Those <18 years old; n = 149 (10% of the total).

*e Those ≥18 years old; n = 1,419 (90% of the total).
**Summary of results for organisms not detected by the BCID panel.** Organisms detectable by the comparator assays for which the BCID pouch did not contain primers are called off-panel or OPOs. Other organisms identified during the resolution of discrepancies between BCID and the phenotypic comparator assays included one species of each of seven genera: Abiotrophia, Flavobacterium, Chryseomonas, Chryseobacterium, Capnocytophaga, Brevundimonas, and Propionibacterium. Other organisms identified included one of seven species of Flavobacterium, one of seven species of Capnocytophaga, and 10 of 14 species of Bacillus. Of the 23 isolates, 17 were Acinetobacter spp. not A. baumannii and 10 were phenotypically A. baumannii. The latter were revealed to be A. pittii, A. nosocomialis, A. bereziniae, and A. radioreistantes by 16S rRNA gene sequencing.

**Gram-positive bacteria.** Table 3 presents a summary of BCID results for Gram-positive and Gram-negative bacteria from the clinical and seeded arms compared to phenotypic determinations. These are the original data and are not altered by the resolution of the discrepancies. These data are presented in the context of the resolution of discordant results (see Table 5).

**Enterococcus spp.** Relative to the comparator assay, BCID exhibited four FP and three FN results (sensitivity and specificity, 97.7% and 99.8%). The species detected in prospective specimens by phenotypic techniques included 56 of Enterococcus faecalis (55.4%; 1 missed by BCID), 37 of Enterococcus faecium (36.6%; 1 missed by BCID), 2 of Enterococcus avium, 2 of Enterococcus casseliflavus (1 missed by BCID), and 1 each of Enterococcus durans and Enterococcus gallinarum. Of the four specimens with BCID FP results for Enterococcus, one was also FN for E. coli. The frozen aliquot of the latter specimen was Gram stained again and recultured, and the fluid showed only Gram-negative rods and grew E. coli. On retesting by BCID, the results were concordant with the phenotypic results: negative for Enterococcus and positive for E. coli. This was presumed to be a bottle selection error at the time of original testing. The remaining three FP results came in bottles that also contained a Staphylococcus sp. target assay. Several less common Staphylococcus spp. are not detected or are detected with lower sensitivity (18).
### TABLE 3 Performance summary of the BCID panel versus the comparator assays for bacteria in both clinical and seeded positive blood cultures combined

| Organism | Isolates detected\(^b\): BCID/comparator | No. of results: BCID/comparator | Sensitivity or PPA\(^b\): TP/(TP + FN) (%) | Specificity or NPA\(^b\): TN/(TN + FP) (%) | 95% CI |
|----------|-----------------------------------------|----------------------------------|----------------------------------------|----------------------------------------|------|
| **Gram-positive bacteria** | | | | | |
| Enterococcus | 102/101 | 29/29 | 127 4 3 2,073 | 127/130 (97.7) | 93.4–99.5 | 2,073/2,077 (99.8) | 99.5–99.9 |
| L. monocytogenes | 0/0 | 36/36 | 36 0 0 2,171 | 36/36 (100) | 90.3–100 | 2,171/2,171 (100) | 99.8–100 |
| Staphylococcus | 780/797 | 2/1 | 770 12 28 1,397 | 770/798 (96.5) | 95.0–97.7 | 1,397/1,409 (99.1) | 98.5–99.6 |
| S. aureus | 257/257 | 0/0 | 253 4 4 1,946 | 253/257 (98.4) | 96.1–99.6 | 1,946/1,950 (99.8) | 99.3–99.9 |
| Streptococcus | 140/141 | 63/62 | 198 5 5 1,999 | 198/203 (97.5) | 94.3–99.2 | 1,999/2,004 (99.4) | 99.4–99.9 |
| S. pneumoniae | 26/25 | 12/12 | 36 2 1 2,168 | 36/37 (97.3) | 85.8–99.9 | 2,168/2,170 (99.9) | 99.7–100 |
| S. pyogenes | 8/7 | 31/31 | 38 1 0 2,168 | 38/38 (100) | 90.7–100 | 2,168/2,169 (99.9) | 99.7–100 |
| Total | 1,331/1,346 | 191/189 | 1,494 28 41 | 16,093 | 1,494/1,535 (97.3) | 96.4–98.1 | 16,093/16,121 (99.8) | 99.7–99.9 |
| **Gram-negative bacteria** | | | | | |
| A. baumannii | 16/14 | 40/37 | 51 5 0 2,151 | 51/51 (100) | 93.0–100 | 2,151/2,156 (99.8) | 99.7–100 |
| Enterobacteriaceae | 307/310 | 187/188 | 490 4 8 | 1,705 | 490/498 (98.4) | 96.9–99.3 | 1,705/1,709 (99.8) | 99.4–99.9 |
| E. cloacae complex | 24/22 | 17/17 | 38 3 1 | 2,165 | 38/39 (97.4) | 85.6–99.9 | 2,165/2,168 (99.9) | 99.6–100 |
| E. coli | 149/148 | 6/5 | 150 5 3 | 2,049 | 150/153 (98.0) | 94.4–99.6 | 2,049/2,054 (99.8) | 99.4–99.9 |
| K. oxytoca | 6/6 | 54/58 | 59 1 5 | 2,142 | 59/64 (92.2) | 82.7–97.4 | 2,142/2,143 (99.9) | 99.7–100 |
| K. pneumoniae | 74/71 | 37/34 | 102 9 3 | 2,093 | 102/105 (97.1) | 91.9–99.4 | 2,093/2,102 (99.6) | 99.2–99.8 |
| Proteus | 22/22 | 17/17 | 39 0 0 | 2,168 | 39/39 (100) | 91.0–100 | 2,168/2,168 (100) | 99.8–100 |
| S. marcescens | 22/22 | 55/55 | 76 1 1 | 2,129 | 76/77 (98.7) | 93.0–100 | 2,129/2,130 (99.9) | 99.7–100 |
| H. influenzae | 8/8 | 35/35 | 43 0 0 | 2,164 | 43/43 (100) | 91.8–100 | 2,164/2,164 (100) | 99.8–100 |
| N. meningitidis | 1/1 | 35/35 | 36 0 0 | 2,171 | 36/36 (100) | 90.3–100 | 2,171/2,171 (100) | 99.8–100 |
| P. aeruginosa | 52/52 | 0/0 | 51 1 1 | 2,154 | 51/52 (98.1) | 89.7–100 | 2,154/2,155 (99.9) | 99.7–100 |
| Total | 681/676 | 483/481 | 1135 29 22 | 23,091 | 1,135/1,157 (98.1) | 97.1–98.8 | 23,091/23,120 (99.9) | 99.8–99.9 |

\(^a\) Culture or culture and 16S rRNA gene sequencing for A. baumannii.

\(^b\) The data in these columns are based on the combined results from the clinical and seeded specimens. Sensitivity refers to performance with clinical specimens, while PPA refers to performance with seeded specimens. Despite the differences in names and usage, they were calculated identically. Likewise, specificity refers to performance with prospective samples, while NPA is used for seeded specimens. They were also calculated identically. These data were obtained in the initial analyses and have not been changed to reflect subsequent investigations of discordant results. Investigations of discordant results are described in Results and summarized in Table 5.

\(^c\) K. oxytoca is the only organism that apparently failed to meet the prespecified criterion of a sensitivity of >95%. However, this was due to an unavoidable error in phenotypic identification. When this was resolved, the sensitivity for this organism exceeded 98.3%.
Gram-negative bacteria and also targets members of the family Enterobacteriaceae that do not have a specific assay. The unresolved results for Gram-negative bacteria detected during this evaluation are presented in Table 3. Discrepancies are discussed below (for a summary, see Table 5).

A. baumannii. A. baumannii was identified by phenotypic identification of an A. calcoaceticus-A. baumannii complex member, followed by bidirectional 16S rRNA gene PCR sequencing. After utilization of both comparator assays, BCID detected 51/51 cultures that were found to be positive for A. baumannii (sensitivity, 100%). Ten isolates phenotypically identified as members of the A. calcoaceticus-A. baumannii complex were not confirmed by 16S rRNA gene sequencing to be A. baumannii but rather proved to be A. pittii (four isolates), A. nosocomialis (four isolates), A. bereziniae (one isolate), and A. radioresistens (one isolate). All except the four A. pittii isolates are included among the Acinetobacter isolates not A. baumannii in Table 2. In addition, BCID falsely identified the four A. pittii isolates and one A. junii isolate as A. baumannii (a total of five FP results, specificity of 99.8%; see also Table 5).

Enterobacteriaceae. Members of the family Enterobacteriaceae are targeted by a family level assay that detects most of the common genera/species present in human infections. In addition, the E. cloacae complex, E. coli, K. oxytoca, K. pneumoniae, Proteus spp., and S. marcescens all have their own specific assays. If the family level assay and/or one of the genus/species-specific assays was positive, BCID reported “Enterobacteriaceae detected.” As detailed in Table 3, the reference phenotypic methods detected 310 members of the family Enterobacteriaceae in the prospective cultures with four BCID FP and seven BCID FN results. In addition, 188 cultures were seeded with members of the family Enterobacteriaceae. One gave a BCID FN result, bringing the total number of FN results to eight. Two discrepant results were from sequential specimens and were resolved as specimen mix-ups. Another specimen was both FN for Enterobacteriaceae and FP for Enterococcus by BCID (discussed above as a bottle selection error). A third FN BCID Enterobacteriaceae determination came from a specimen that was reported to contain both an E. coli and a Pasteurella species isolate by phenotypic methods, but the E. coli isolate could not be detected on resubculture. The remaining FN specimens were analyzed by 16S rRNA gene sequencing directly from the bottle. Among the four definite BCID FN results were two Pantoaea sp. isolates, one S. marcescens isolate, and one Citrobacter koseri isolate. Sequencing of the eighth FN isolate failed, and it remains unresolved. Of the four FP specimens, one was resolved by attribution to a specimen mix-up described above, while three came from mixed cultures, failed to grow on resubculture, and remain unresolved.

E. cloacae complex. Individual members of the E. cloacae complex were not differentiated either by BCID or by phenotypic assays. Phenotypic assays detected the E. cloacae complex in 22 prospective clinical specimens and 17 cultures seeded with members of the E. cloacae complex, including 14 clinical isolates and one reference strain (ATCC 13407; three cultures). Relative to the phenotypic comparator methods, BCID detected 21 (95.5%) of the 22 positive clinical isolates and all 17 of the seeded cultures. The sensitivity and specificity were 97.4 and 99.9%, respectively. On resolution, the single FN culture grew an organism identified by bidirectional 16S rRNA gene sequencing as E. cloacae, but the presence of E. cloacae in the three FP samples was not confirmed.
Instead, three other members of the family Enterobacteriaceae were detected: one Cedecia daviesiae isolate and two K. pneumoniae isolates.

**E. coli.** Phenotypic assays detected *E. coli* in 148 of the prospective clinical specimens of which BCID detected 145. Five specimens were seeded with distinct clinical isolates that were included to increase the number of *blaKPC*-containing *E. coli* isolates, and both culture and BCID detected all five. Five BCID FP results were obtained, four among the 1,420 prospective clinical cultures that were negative for *E. coli* by phenotypic methods and one among the 634 cultures seeded with other organisms. Three FN results were also obtained. One FN result and one FP result were obtained with sequential specimens and were interpreted as specimen mix-ups. Among the four remaining FP cultures, one was confirmed to contain *E. coli* on subculture. Three contained other members of the family Enterobacteriaceae; two contained *K. pneumoniae*, and one contained *E. cloacae*. These are not known to cross-react in the BCID *E. coli* assay. Of the two remaining BCID FN samples, one was already discussed as a possible sample selection error (see the *Enterococcus* and Enterobacteriaceae sections). One *E. coli* BCID FN specimen is also discussed in the Enterobacteriaceae section.

**K. oxytoca.** Phenotypic analysis and BCID detected *K. oxytoca* in six prospective clinical specimens. However, there was one FP BCID result and one FN BCID result. BCID also detected 54/55 cultures seeded with *K. oxytoca*. The overall unresolved sensitivity of the *K. oxytoca* assay was therefore 92.2% (Table 3). The five FN results and one FP result were investigated. No *K. oxytoca* could be grown from the frozen aliquot of the single FP specimen. One of the five apparent BCID FN results was confirmed to be *K. oxytoca* by bidirectional 16S rRNA gene sequencing. The other four were identified by 16S rRNA gene sequencing to be *K. ornithinolytica*, which cannot be distinguished from *K. oxytoca* by phenotypic methods (20, 21). After resolution of this unavoidable phenotypic error, the resolved sensitivity of the BCID assay for *K. oxytoca* was 98.3%.

**K. pneumoniae.** The phenotypic assays detected *K. pneumoniae* in 71 prospective clinical specimens, of which BCID detected 68. BCID also detected 34/34 cultures seeded with individual clinical isolates. There were nine apparent BCID FP results and three FN results. Of the three FN BCID results, one was confirmed to be *K. pneumoniae*. One was found by sequencing to be *Raoultella planticola*, and one isolate did not grow for further analysis. The nine BCID FP results were investigated by attempting to isolate *K. pneumoniae* from the blood culture specimens, but no *K. pneumoniae* could be grown. Two isolates were identified as *Enterobacter aerogenes*, and two were identified as *E. coli*. Four BCID FP results, including two of seven that were also BCID positive for other members of the family Enterobacteriaceae, were not resolved.

**Proteus.** In prospective clinical cultures, both the comparator phenotypic methods and BCID identified *Proteus* in 22/22 clinical cultures. BCID also detected *Proteus* in 22/22 seeded cultures. There were no FN or FP BCID results. Thus, the sensitivity and specificity were 100%.

**S. marcescens.** Both the phenotypic methods and BCID identified *S. marcescens* in the same 22 prospective cultures, with one FP determination. BCID detected 54/55 *S. marcescens* isolates in seeded cultures with one FN result. The unresolved sensitivity and specificity were 98.7 and 99.9%, respectively. Both the FN and FP results were investigated and found to be due to organisms other than *S. marcescens*. The apparent FN specimen contained a member of the *Serratia proteamaculans-S. grimesii* complex that had been misidentified by the phenotypic methods as *S. marcescens*, while the apparent FP specimen contained an *R. ornithinolytica* isolate that was identified phenotypically as *K. oxytoca* but also cross-reacted with the *S. marcescens* assay (data not shown).

**H. influenzae.** In prospective clinical cultures, both the phenotypic methods and BCID identified *H. influenzae* in the same 8 prospective clinical cultures and in 35/35 (100%) cultures seeded with 22 distinct clinical isolates. There were no FN or FP determinations. The sensitivity and specificity of BCID for *H. influenzae* were both 100%.

**Neisseria meningitidis.** Both the phenotypic methods and BCID identified *N. meningitidis* in the same clinical culture and in 35/35 (100%) cultures seeded with 22 distinct clinical isolates. There were no FN or FP determinations. The sensitivity and specificity of BCID for *N. meningitidis* were both 100%.

**P. aeruginosa.** In prospective clinical cultures, both the phenotypic assays and BCID identified *P. aeruginosa* in 52 cultures. However, BCID reported one FN result and one FP result (sensitivity, 98.1%; specificity, 99.7%). One seeded culture were included. The apparent BCID FN specimen was found to contain *P. putida*, not *P. aeruginosa*. The BCID FP specimen did not contain detectable *Pseudomonas*, but rather *S. aureus* was detected.

**Candida.** The unresolved results of BCID detection of the *Candida* species included in the panel are shown in Table 4. Res-
olution of discordant results is discussed in each organism section and summarized in Table 5.

**C. albicans.** The phenotypic assays and BCID detected *C. albicans* in the same 16 clinical specimens. In addition, both the phenotypic and BCID assays detected 48/48 of cultures seeded with *C. albicans*. BCID also reported four FP *C. albicans* detections. *C. albicans* could not be grown from frozen aliquots of these specimens, although *C. glabrata* was grown from one. The sensitivity and specificity of BCID for *C. albicans* were 100 and 99.8%, respectively.

**C. glabrata.** The phenotypic assays and BCID detected *C. glabrata* in the same 12 prospective clinical specimens and 37/37 cultures seeded with *C. glabrata* (27 clinical isolates, ATCC MYA-2950 [nine cultures], and ATCC 15126 [one culture]). BCID produced two FP results among the prospective clinical cultures (sensitivity, 100%; specificity, 99.9%). *C. glabrata* could not be grown from frozen aliquots of the two FP specimens, although *C. albicans* was grown from one.

**C. krusei.** The phenotypic method and BCID detected *C. krusei* in the same 4 prospective clinical specimens and in all 33 seeded specimens (15 distinct clinical isolates, ATCC 14343 [15 cultures], and ATCC 6248 [3 cultures]). BCID did not report any FP or FN

| Result and analyte | No. of results | Comparator result confirmed | FilmArray result confirmed | Both results incorrect | Inconclusive |
|--------------------|----------------|-----------------------------|---------------------------|------------------------|-------------|
| **BCID FP results** |                |                             |                           |                        |             |
| Enterococcus       | 4              | 1                           | 1                         | 3                      |             |
| Staphylococcus     | 12             | 11                          | 1                         | 1                      |             |
| *S. aureus*        | 4              | 0                           | 1                         | 3                      |             |
| Streptococcus      | 5              | 5                           | 1                         | 3                      |             |
| *S. pneumoniae*    | 2              | 2                           |                           | 1                      |             |
| *S. pyogenes*      | 1              | 1                           |                           | 1                      |             |
| *A. baumannii*     | 5              | 5                           |                           | 1                      |             |
| Enterobacteriaceae | 4              | 1                           |                           | 3                      |             |
| *E. cloacae complex* | 3              |                             |                           |                        |             |
| *E. coli*          | 5              | 1                           | 2                         |                        |             |
| *K. pneumoniae*    | 9              | 4                           | 3                         |                        |             |
| *K. oxytoca*       | 1              | 1                           |                           |                        |             |
| *S. marcescens*    | 1              | 1                           |                           |                        |             |
| *P. aeruginosa*    | 1              |                             |                           |                        |             |
| *C. albicans*      | 4              |                             |                           |                        |             |
| *C. glabrata*      | 2              |                             |                           |                        |             |
| *C. parapsilosis*  | 2              |                             |                           |                        |             |
| **Total**          | 65             | 12                          | 18                        | 5                      | 30          |
| % of total FP results | 18            | 28                          | 8                         | 46                     |             |

| **BCID FN results** |                |                             |                           |                        |             |
| Enterococcus        | 3              | 3                           |                           |                        |             |
| Staphylococcus      | 28             | 20                          | 1                         | 7                      |             |
| *S. aureus*         | 4              | 1                           | 2                         |                        |             |
| Streptococcus       | 5              | 3                           |                           |                        |             |
| *S. pneumoniae*     | 1              | 1                           |                           |                        |             |
| Enterobacteriaceae  | 8              | 4                           | 3                         |                        |             |
| *E. cloacae complex* | 1              |                             |                           |                        |             |
| *E. coli*           | 3              | 3                           |                           |                        |             |
| *K. pneumoniae*     | 3              | 1                           |                           |                        |             |
| *K. oxytoca*        | 5              | 1                           | 4                         |                        |             |
| *S. marcescens*     | 1              | 1                           |                           |                        |             |
| *P. aeruginosa*     | 1              | 1                           |                           |                        |             |
| *C. parapsilosis*   | 2              |                             |                           |                        |             |
| **Total**           | 65             | 34                          | 18                        | 1                      | 12          |
| % of total FN results | 52            | 28                          | 2                         | 18                     |             |

The method used was investigation of possible sample mix-ups, reculturing from frozen aliquots of the blood culture bottle fluid, sequencing directly from the bottle if isolated organisms were expected, or sequencing from resubcultured isolated organisms if more than one isolate per bottle was present. For bacterial resolution, 16S rRNA gene sequences were analyzed and for yeast, ITS1 was analyzed. See Materials and Methods for further information. Three apparent blood culture bottle mix-ups and one apparent bottle selection error were resolved in favor of BCID.

a See Results for further information regarding the indicated organism.
detections of *C. krusei*. Thus, the unresolved sensitivity and specificity were both 100%.

**C. parapsilosis.** The phenotypic method and BCID both detected *C. parapsilosis* in the same seven clinical specimens and 52/54 (96.3%) seeded cultures (two apparent FN results). BCID also produced two FP results (sensitivity, 96.7%; specificity, 99.9%). The two FN results and the two FP results were investigated. Sequencing determined that the two apparent BCID FN samples both contained *Candida metapsilosis*, which is identified by phenotypic methods as *C. parapsilosis*. *C. parapsilosis* could not be grown from either of the two cultures with FP BCID results, although *C. albicans*, also detected by BCID, grew from one.

**C. tropicalis.** Both the phenotypic methods and BCID detected *C. tropicalis* in the same three prospective clinical specimens and 36/36 cultures seeded with *C. tropicalis* (35 unique clinical isolates and 1 reference strain [CAP proficiency sample]), with no FP or FN results. Thus, the unresolved sensitivity and specificity were both 100%.

**Resolution of discrepant organism identification.** The resolution of BCID FP and FN organism detection is discussed in the section relevant to each organism (*Tables 3 and 4* contain unresolved data) and summarized in Table 5. There were 130 apparently discrepant BCID organism detections, 65 FP and 65 FN results. Bottle selection errors apparently occurred with seven (0.3%) of the bottles analyzed by BCID. One that occurred at the time of specimen selection is discussed in the sections on *Enterococcus*, *Enterobacteriaceae*, and *E. coli*. Sixteen additional discrepancies were attributed to three bottle mix-ups (two bottles each). The bottles involved in the mix-ups turned positive at approximately the same time and were erroneously picked by testing personnel. After investigation, discrepancies attributed to bottle selection errors were resolved in favor of BCID.

Of 130 discordant results, 36 were resolved in favor of BCID and 46 were resolved in favor of the comparator assay. In an additional six discordant results, both methods were incorrect because of cross-reactivity with closely related organisms or misidentification by phenotypic methods, leaving 42 discordant results unresolved. Of the BCID FP detections, 25/65 (38.5%) occurred in the 81 prospective clinical cultures that were polymicrobial, as did 16/65 (24.6%) of the BCID FN results.

**Antibiotic resistance gene detection.** The results of the BCID antibiotic resistance gene assays are compared to data derived from gene sequencing directly from frozen aliquots from the blood culture bottles (Table 6). A comparison of the BCID resistance gene detection with the phenotypic assays utilized by the participating laboratories is presented elsewhere (18).

**Methicillin resistance (meca) gene detection in *Staphylococcus spp.*** The meca PCR comparator assay detected *mecA* in 496 cultures containing staphylococci (494 clinical and 2 seeded) and BCID detected 488 (98.4%) with eight FN results. Of 286 *Staphylococcus*-containing clinical cultures in which the comparator assay did not detect *mecA*, BCID was concordant in 281 (98.3%) with five FP results. The sensitivity of *mecA* detection in all *Staphylococcus* spp. was 98.4%. The specificity was 98.3%. The BCID assay is also capable of detecting meCC, formerly known as *mecA*LGAA251. No isolates with *mecC* were detected in this study.

**meca detection in *S. aureus*.** Of 139 prospective samples that were positive for *mecA* by the comparator molecular method, BCID detected *mecA* in 137 (sensitivity, 98.6%; specificity, 100%). The two isolates with apparent BCID FN results were phenotypically methicillin susceptible.

**Vancomycin resistance in *Enterococcus spp.* (vanA/B).** Among the clinical specimens, BCID vanA/B results were concordant with the molecular comparator method for 36/36 positive and 67/67 negative cultures. In all 29 seeded cultures, vanA/B detection by comparator and BCID agreed (28 positive, 1 negative).

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**TABLE 6 Comparison of FilmArray BCID resistance gene results to the prespecified comparator assay (PCR/sequencing directly from blood culture bottle)**

| Antimicrobial resistance gene(s) | Isolates detected: BCID/comparator | No. of results: BCID/comparator | Sensitivity or PPA*: | Specificity or NPA*: |
|----------------------------------|------------------------------------|-------------------------------|---------------------|---------------------|
|                                  | Clinical arm                      | Seeded arm                    |                     |                     |
|                                  | TP(+/-) | FP (+/-) | FN (-/-) | TN (-/-) | Sensitivity or PPA* (%) | 95% CI | Specificity or NPA* (%) | 95% CI |
| meca in association with:        |                                |                                |                     |                     |
| All *Staphylococcus* isolates    | 491/494 | 2/2     | 488    | 5     | 281 | 488/496 (98.4) | 96.8–99.3 | 281/286 (98.3) | 96.0–99.4 |
| detected*                        |                                |                                |                     |                     |
| *Staphylococcus* and *S. aureus* | 137/139 | 0/0     | 137    | 0     | 2   | 118/139 (98.6) | 94.9–99.8 | 118/118 (100) | 96.9–100  |
| isolates detected                |                                |                                |                     |                     |
| vanA/B in association with       | 36/36    | 28/28   | 64     | 0     | 67  | 64/64 (100)* | 94.4–100 | 67/67 (100) | 94.6–100  |
| *Enterococcus* isolates detected |                                |                                |                     |                     |
| bla<sub>KPC</sub> in association with | 6/6* | 33/33   | 39     | 0     | 0   | 39/39 (100)* | 91.0–100 | 558/558 (100) | 99.3–100  |

* Sensitivity and specificity refer to performance with the prospective specimens only. PPA and NPA refer to performance with the seeded specimens. These are unresolved data.

* Either *Staphylococcus* or *S. aureus* isolates detected or both.

* Of the 64 *Enterococcus* isolates, 11 (17.2%) carried the van<sub>B</sub> gene as determined by bidirectional sequence analysis.

* All six isolates were identified as *K. pneumoniae*.

* These isolates included 30 of *K. pneumoniae*, 2 of *E. cloacae*, and 1 of *E. coli*. 

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The sensitivity and specificity of the vanA/B detection assay were both 100%.

**bla\textsubscript{KPC}, in Gram-negative bacteria.** BCID reported a total of 597 specimens to be positive for members of the family Enterobacteriaceae, A. baumannii, or P. aeruginosa. BCID and the comparator method were concordant for 6/6 positive specimens and 364/364 negative specimens. In cultures seeded with members of the family Enterobacteriaceae, A. baumannii, or P. aeruginosa, BCID and the comparator methods were concordant in 33/33 positive detections and in 194/194 negative determinations. The sensitivity and specificity of the BCID **bla\textsubscript{KPC}, detection assay are both 100%.

**DISCUSSION**

Rapid identification of the etiologic agents of bloodstream infection/sepsis and their resistance genes by molecular methods allows early targeted antibiotic therapy, which may improve patient outcomes, reduce mortality, and impact the length of stay of septic patients (5–8, 22, 23). This paper presents the results of a large (1,568 clinical and 639 seeded specimens), multicenter trial of the FilmArray BCID panel. Of the 2,207 BCID analyses performed, 98.1% were completed on the initial run. Repeat analyses of specimens with failed controls were successful in all cases, and no specimens were lost because the repeat test was not initiated within the 8-h time limit. The comparator assays for organism ID were the phenotypic assays used in the participating laboratories. For A. baumannii, 16S rRNA gene sequencing was prespecified to be the final arbiter. This strategy eliminated several organisms phenotypically misidentified as A. baumannii, but it also revealed cross-reactivity of BCID with A. pittii and A. junii.

We did not change the results presented in Tables 3, 4, and 6 after the resolution of discrepancies. The unresolved sensitivities of all BCID assays of organisms other than K. oxytoca were \( \geq 96\% \) (Tables 3, 4, and 6). After four apparent K. oxytoca BCID misses were determined to be R. ornithinolytica by 16S rRNA gene sequencing, the resolved sensitivity for K. oxytoca exceeded 98%. Phenotypic methods incorrectly identify R. ornithinolytica as K. oxytoca (20, 21).

Discordant results, both FP and FN, are a problem when the new assay may be more sensitive and specific than the supposed gold standard. The detection of discrepant results among the seeded bottles was not unexpected since bottles were seeded with archived clinical isolates that were identified phenotypically. These are subject to the same limitations of phenotypic testing as the prospective clinical specimens. Organisms phenotypically identified as K. oxytoca and K. pneumoniae and reported to be BCID misses proved to be R. ornithinolytica and R. planticola, as discussed above. In addition, it was determined by BioFire, subsequent to the research presented here, that 16/28 FN Staphylococcus sp. assay results were due to the failure of the BCID assay to detect S. pettenkoferi. For detailed information on other organisms that may cause cross-reactivity or organisms that may be undetectable or detected with reduced efficiency by BCID, see reference 18.

FilmArray BCID is capable of detecting organisms in mixed cultures. Of prospective clinical specimens, 81 (5.2%) contained multiple (two to four) BCID-detectable organisms. However, other BCID studies have shown that the presence of multiple organisms favors discordant BCID results (15, 24). We obtained BCID FP results with 37 (46%) of the specimens containing multiple organisms. In addition, although polymicrobial cultures represented approximately 10% of those in which Staphylococcus was detected, they were associated with 29% (8/28) of the FN results and 58% (7/12) of the FP results for Staphylococcus. A similar trend was seen for other target organisms.

Investigation of the 130 discrepancies between BCID and the comparator phenotypic assays was approached by investigating the possibility of human error, by growing the organisms from subcultures of frozen aliquots of the specimens, and/or by identification by sequencing methods either directly from aliquots of the frozen specimens or from colonies grown from these aliquots. These investigations revealed one bottle selection error and three possible sample mix-ups (six bottles). The supposed mix-ups led to seven FP and nine FN determinations that were resolved in favor of BCID (Table 5). Overall, 36 (27.6%) of the discrepant results were resolved in favor of BCID and 46 (35.3%) were resolved in favor of the comparator. In 6 (4.6%) cases, both identifications were incorrect, while 42 (32.3%) remain unresolved. The sensitivity and specificity of the BCID detections were not altered in Tables 3 and 4 regardless of the results of these resolution investigations.

There are several possible sources of FP results in nucleic acid-based assays, as molecular methods may detect the genetic material present in nonviable organisms. One cause of molecular FP results could be the ingredients in the blood culture bottles. Package inserts for blood culture bottles state that although the fluid in the bottles is sterile, it may contain nonviable, Gram-stainable organisms. These could theoretically also be detectable by molecular techniques. A preliminary study showed that uninoculated, charcoal-containing medium (BacT/ALERT FA FAN, aerobic; bioMérieux) gave multiple but infrequent FP BCID results (18). Any blood culture bottle type could theoretically contain nucleic acid sequences from nonviable organisms. BCID results that do not agree with the bottle Gram strains should be evaluated carefully. The resin-containing bottles used in this study (BD Bactec Plus Aerobic/F), gave no false-positive results when 92 bottles from 16 lots were screened (data not shown; 95% confidence interval, 0.00 to 0.04). However, some of the bottles seeded with one organism were BCID FP for an additional organism that might have come from the bottle medium.

Another source of these FP results could be rare microbial nucleic acids in the blood injected into the seeded bottles to create the simulated samples. In addition, BCID FP results with clinical samples could be caused by nonviable organisms present in the patient’s blood, especially since we do not know whether the patients were on antibiotic therapy at the time of specimen collection. The patient might have originally experienced polymicrobial bacteremia, and one organism might still be able to grow in the blood culture bottles while a different, but no longer viable, organism could remain detectable by BCID. Alternatively, an organism that was no longer viable at the time of culture collection could have been detected by BCID in a patient specimen also containing a contaminating skin organism; these may be expected in about 3% of positive blood cultures (16). Thus, an overrepresentation of “biologic” BCID FP results would be expected in polymicrobial cultures, both in those in which BCID detected other panel organisms and in those containing only OPOs detected by culture.

The unresolved sensitivities and specificities for detection of mecA in all members of the genus Staphylococcus and for S. aureus alone exceeded 98%. The unresolved sensitivities and specificities...
for vanA/B and blaKPC were 100%. No blaKPC was detected in A. baumannii or P. aeruginosa, as blaKPC in these organisms is currently rare or absent in the United States but has been detected in Puerto Rico (25, 26).

Several previous studies of the BCID have been published (15, 24, 27-30). Most of these analyzed specimens from adults, but one (24) analyzed only pediatric specimens. The majority included 100 to 169 specimens; but one included 204 prospective specimens (27) and another included 111 prospective and 102 archived samples (15). All generally found excellent sensitivity and specificity, although the number of pathogens was limited. Altun et al. (27) showed the reproducibility and stability of the BCID results by retesting five positive bottles for up to 4 weeks. Two studies (15, 27) analyzed all cultures that were flagged as positive by the blood culture instruments, regardless of Gram stain results, and found that some of the bottles grew no organisms and were BCID negative as well. We did not analyze culture-negative bottles since our protocol required a positive Gram stain on bottles flagged as positive prior to BCID testing.

Although our investigation is the largest clinical study of BCID, it is limited by a low number of clinical isolates of certain organisms. No L. monocytogenes isolates were detected and only one N. meningitidis isolate was detected in the prospective clinical specimens. Nonetheless, the presence of these rare and extremely pathogenic organisms is highly significant clinically, and the inclusion of targets for them in this assay seems warranted. For some rare organisms and also for the resistance genes, many of the analyses were performed with seeded specimens. Some of these were archived clinical isolates. In many cases, this did not fulfill the desired numbers, and so reference strains were included, many of which were tested repeatedly. This is not ideal, but a ready alternative to attain statistically significant results does not seem available.

Contamination of blood cultures with skin organisms is unavoidable (16). While Staphylococcus spp. other than S. aureus are the most prevalent contaminating organisms, they may also be the etiologic agents of disease (16) and thus worthy of reporting. However, in this study, the numbers of other OPOs assumed to be skin contaminants (49 Corynebacterium spp., 33 Bacillus spp.; not B. anthracis) and 27 Micrococcus spp.; 16) exceeded many of the pathogens detected by BCID. Identification of a common blood culture contaminant can be useful clinically, as it may facilitate the discontinuation of antibiotics (5, 31). Thus, inclusion of such organisms in the BCID panel might be valuable.

Another limitation is that BCID did not report the resistance genes as detected unless an organism generally recognized as containing that gene was also detected. This seems sensible but may cause occasional problems. For example, it would have prevented the detection of the world’s first vancomycin-resistant S. aureus strain. This was isolated in one of the participating laboratories and was found to have acquired vanA (32).

An additional limitation is that the study sites, although they are distributed widely throughout the United States, did not include other countries. Resistance mechanisms and organism strains are known to vary geographically, and information about the detectability of such variations could prove useful, as such organisms and resistance genes often disseminate widely over time (25, 26).

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