BioFire® Blood Culture Identification 2 Panel (BCID2) ®

What is it?

A multiplex PCR test that detects 33 pathogens and 10 genetic resistance markers directly from positive blood culture bottles. Results are available about an hour after the test is initiated. This technology informs decision making when moving from empiric to targeted antimicrobial therapy earlier in treatment.

What does it identify?

Gram Positive (+) Bacteria	Gram Negative (-) Bacteria		
Enterococcus faecalis Enterococcus faecium Listeria monocytogenes Staphylococcus species • Staphylococcus aureus • Staphylococcus epidermidis • Staphylococcus lugdunensis Streptococcus species • Streptococcus agalactiae • Streptococcus pyogenes • Streptococcus pneumoniae	 Enterobacterales Enterobacter cloacae complex Escherichia coli Klebsiella aerogenes Klebsiella oxytoca Klebsiella pneumoniae group Proteus species Salmonella species Serratia marcescens 	Acinetobacter calcoaceticus- baumannii complex Bacteroides fragilis Haemophilus influenzae Neisseria meningitidis Pseudomonas aeruginosa Stenotrophomonas maltophilia	
Gram Positive (+) Resistance Genes	Gram Negative (-) Resistance Genes		
Methicillin Resistance: mecA/C mecA/C and MREJ (MRSA) <u>Vancomycin Resistance:</u> vanA/B	<u>Carbapenemases:</u> IMP KPC OXA-48-like NDM VIM		
Yeast Candida albicans Candida auris Candida glabrata Candida krusei Candida parapsilosis Candida tropicalis Cryptococcus neoformans/gattii	Colistin Resistance: mcr-1 <u>Extended Spectrum Beta-Lactamases (ESBL):</u> CTX-M		

Table 1. Note: The BCID2 panel is **ONLY** capable of detecting organisms and resistance genes included in the list above. Absence of a detected organism does **NOT** indicate a false-positive culture or reporting error.

How is it performed?

The test is not ordered. It is performed <u>automatically</u> on **positive** blood samples meeting one of the following criteria:

• All positive blood cultures with a gram-positive cocci, gram-negative, or fungal organism seen on gram stain

<u>Note</u>: The BCID2 panel will be performed on the first blood sample meeting testing criteria and will not be repeated on samples collected in the subsequent 72 hours. <u>All samples will receive standard culture testing as well.</u>

How are results reported?

Results will appear within the blood culture report or as **Rapid Blood Culture Identification** under the micro tab, indicating the genus or species identified by multiplex PCR. When the organism identification and susceptibilities from standard testing are available, the culture report will be updated with this new information below the BCID2 result as shown below:

Example:

Blood Culture

Escherichia coli ESBL 🚦

ESBL Extended Spectrum Beta-lactamase Producer.

	Component	Ref Range & Units	
24	Acinetobacter calcoaceticus -baumannii complex PCR	Not Detected	Not Detected
<u>~</u>	Bacteroides fragilis PCR	Not Detected	Not Detected
∽:	Enterobacterales PCR	Not Detected	Detected !

	Enterobacter cloacae complex PCR	Not Detected	Not Detecte
]	Escherichia coli PCR	Not Detected	Detected !
	Klebsiella aerogenes PCR	Not Detected	Not Detect
	Klebsiella oxytoca PCR	Not Detected	Not Detect
	Klebsiella pneumoniae Group PCR	Not Detected	Not Detect
	Proteus species PCR	Not Detected	Not Detect
	Salmonella species PCR	Not Detected	Not Detect
	Serratia marcescens PCR	Not Detected	Not Detect
	Haemophilus influenzae PCR	Not Detected	Not Detect
]	Neisseria meningitidis PCR	Not Detected	Not Detect
	Pseudomonas aeruginosa PCR	Not Detected	Not Detect
	Stenotrophomonas maltophilia PCR	Not Detected	Not Detect
	Enterococcus faecalis PCR	Not Detected	Not Detect
	Enterococcus faecium PCR	Not Detected	Not Detect
	Staphylococcus PCR	Not Detected	Not Detect
	Staphylococcus aureus PCR	Not Detected	Not Detect
	Staphylococcus epidermidis PCR	Not Detected	Not Detect
	Staphylococcus lugdunensis PCR	Not Detected	Not Detect
	Streptococcus PCR	Not Detected	Not Detect
	Streptococcus agalactiae PCR	Not Detected	Not Detect
]	Streptococcus pneumoniae PCR	Not Detected	Not Detect
]	Streptococcus pyogenes PCR	Not Detected	Not Detect
	Candida albicans PCR	Not Detected	Not Detect
	Candida auris PCR	Not Detected	Not Detect
	Candida glabrata PCR	Not Detected	Not Detect
	Candida krusei PCR	Not Detected	Not Detect
	Candida parapsilosis PCR	Not Detected	Not Detect
	Candida tropicalis PCR	Not Detected	Not Detect
	Cryptococcus neoformans/gatti PCR	Not Detected	Not Detect
	Kpc Carbapenem Resistant Gene	Not Detected, Not Applicable	Not Detect
	MCR-1 Resistant Gene	Not Detected, Not Applicable	Not Detect
	Imp Carbapenem Resistant Gene	Not Detected, Not Applicable	Not Detect
	Oxa-48-Like Carbapenem Resistant Gene	Not Detected, Not Applicable	Not Detect
	Ndm Carbapenem Resistant Gene	Not Detected, Not Applicable	Not Detect
	Vim Carbapenem Resistant Gene	Not Detected, Not Applicable	Not Detect
1	CTX-M Cephalosporin Resistance Gene	Not Detected, Not Applicable	Detected !

What decisions can be made based on the BCID2 panel results?

As with all diagnostic tests, results from the BCID2 panel need to be evaluated <u>in view of the patient's unique</u> <u>clinical scenario</u>, <u>severity of illness</u>, <u>and source of infection</u>. BCID2 results help to optimize antimicrobial therapy.</u> However, treatment should be re-evaluated when standard organism identification and susceptibility testing results are reported.

<u>Common Scenarios</u>: In certain situations, BCID2 can be used to de-escalate antimicrobials. Examples of common situations where de-escalation may be reasonable include but are not limited to:

- If *Staphylococcus aureus* is identified in the absence of methicillin resistance genes (MecA/C, MREJ): MRSA-active empiric therapy (i.e. vancomycin) should be discontinued and therapy can be changed to cefazolin or oxacillin.
- If *Escherichia coli* or *Klebsiella pneumoniae* is identified in the absence of the CTX-M gene in patients on antipseudomonal coverage (i.e. cefepime or piperacillin/tazobactam): de-escalate to ceftriaxone
 - Internal microbiology data for *Escherichia coli* and *Klebsiella spp.* isolates:
 - When 60 *E. coli* isolates and 45 *Klebsiella spp.* isolates at Atrium Health Wake Forest Baptist were reviewed for detection of CTX-M by BCID2 and correlating ceftriaxone susceptibility by standard culture and susceptibility report, the positive predictive value (PPV) was 100% for *E. coli* and *Klebsiella spp.* and the negative predictive value (NPV) was 94% for *E. coli* and 100% for *Klebsiella spp.*
 - Patients in whom de-escalation may not be appropriate:
 - Lack of source control, suspected polymicrobial infection, ANC ≤500, other infection caused by resistant organism / recent culture history with known resistance, hemodynamic instability or metastatic infection
- If only gram-negative organism(s) identified: discontinue empiric anti-MRSA agent (i.e vancomycin, daptomycin, linezolid)
- Refer to Treatment Decision Algorithms (Table 2-5) below for more information

In certain situations, BCID2 can be used to escalate antimicrobial therapy. If an organism or resistance gene is identified that is not covered by current therapy, modify therapy to treat the pathogen(s). An example of a common situation where escalation is advised may include but is not limited to:

- o If gram-negative with CTX-M gene is identified: escalation to a carbapenem may be appropriate
- Refer to Treatment Decision Algorithms (Table 2-5) below for more information

Other considerations:

- BCID2 is reported from blood cultures and will <u>only</u> detect what is present in the blood; therefore, source of infection should <u>always</u> be considered.
- Data suggest that BCID2 is highly sensitive (99%) and specific (99.8%) for monomicrobial bloodstream infections; however, in polymicrobial infections the capacity to correctly identify all present organisms is decreased.
- BCID2 can report to certain family (*Enterobacterales*) or genus (*Staphylococcus, Streptococcus*) levels, with or without identification of a species. These flags should not be confused with identification of a species.
 - When BCID2 is only able to identify *Enterobacterales* to the family level, the species is not one of the included *Enterobacterales* species on the BCID2 panel (i.e. *E. cloacae, E. coli, K. aerogenes, K. oxytoca, K. pneumoniae, Proteus* spp, *Salmonella* spp, *S. marcescens*). If any of the listed *Enterobacterales* organisms are detected, *Enterobacterales* will also report as detected.
 - When BCID2 only identifies *Staphylococcus* species to the genus level, (i.e. NOT *aureus*, *lugdunensis*, *epidermidis*), the species is likely to be a coagulase negative *Staphylococcus* other than *epidermidis*. If any of the *Staphylococcus* species included on the panel are detected, the *Staphylococcus* genus will also report as detected.

- When BCID2 is only able to identify *Streptococcus* species to the genus level, the species is not one of the included *Streptococcus* species on the BCID2 panel (i.e. *agalactiae*, *pneumoniae*, *pyogenes*). If any of the *Streptococcus* species included on the panel are detected, the *Streptococcus* genus will also report as detected.
- *S. epidermidis* and other coagulase negative *Staphylococcus* species (CoNS) are normal skin flora and may contaminate cultures. In situations where a single blood culture is positive for a CoNS and there is an alternative explanation for the patient's symptoms or no concern for active infection, the sample may be considered contaminated. In cases where multiple sets of blood cultures or repeat cultures are positive with the same CoNS, true infection from CoNS should be considered.
- *S. aureus* and *S. lugdunensis* should not be considered contaminants. ID consultation is recommended for cases of bacteremia caused by these organisms.

Whom do I contact for questions about the BCID2 panel?

- Microbiology lab (6-2658) or CAUSE (secure chat group: WFMC CAUSE Antimicrobial Stewardship Approval)
- Rarely, discrepancies between the BCID2 and the antimicrobial susceptibility testing results occur (e.g. mecA/C is identified but oxacillin is susceptible by MIC). The microbiology lab or ID/CAUSE may be contacted for questions about these cases.

The following treatment decision algorithms outline potential options for revised empiric antimicrobial therapy in response to BCID2 panel results that would be reasonable in most situations. Response to the BCID2 results should consider the entire clinical picture and these recommendations should not supersede clinical judgement. If choosing an additional or alternative empiric antimicrobial than what is listed in the algorithms below, please utilize institutional antibiograms that provide local resistance data for decision making (found on the Atrium Health Wake Forest Baptist Intranet). If uncertain of optimal therapy, consult Infectious Diseases or message CAUSE.

Table 2. Gram-Positive Organism BCID2 Treatment Decision Algorithm DCID2 D I				
BCID2 Result	Revised Empiric Therapy	Comments		
Staphylococcus species				
Staphylococcus aureus Negative mecA/C and MREJ	cefazolin or oxacillin	 MecA/C and MREJ detection indicates methicillin resistant <i>Staphylococcus aureus</i> (MRSA) MREJ is specific for methicillin resistant 		
Staphylococcus aureus Positive mecA/C and MREJ	vancomycin	Staphylococcus aureus (MRSA) • Recommend ID consult		
Staphylococcus lugdunensis Negative mecA/C	cefazolin or oxacillin	MecA/C detection indicates methicillin resistant Staphylococcus lugdunensis		
Staphylococcus lugdunensis Positive mecA/C	vancomycin	• Recommend ID consult		
Staphylococcus epidermidis Negative mecA/C	cefazolin or oxacillin	• MecA/C detection indicates methicillin resistant <i>Staphylococcus epidermidis</i> (MRSE)		
<i>Staphylococcus epidermidis</i> Positive mecA/C	vancomycin	• <i>S. epidermidis</i> and other coagulase negative <i>Staphylococcus</i> species (CoNS) are normal skin flora and may contaminate cultures. In situations where a single blood culture is positive for a CoNS and there is an alternative explanation for the patient's symptoms or no concern for active infection, the sample may be considered contaminated		
Staphylococcus PCR only	vancomycin	 When BCID2 only identifies <i>Staphylococcus</i> species to the genus level, the species is NOT one of the included <i>Staphylococcus</i> species on the BCID2 panel MecA/C detection is not assessed when a species is no identified 		
Enterococcus species				
Enterococcus faecalis Negative vanA/B	ampicillin	•CAUSE/ID approval required for daptomycin and IV linezolid		
Enterococcus faecalis Positive vanA/B	ampicillin alternative: linezolid or daptomycin	• Even when E. faecalis is vancomycin resistant, 90-92% or isolates remain susceptible to ampicillin		
Enterococcus faecium Negative vanA/B	vancomycin	• CAUSE/ID approval required for daptomycin and IV		
Enterococcus faecium Positive vanA/B	linezolid or daptomycin	linezolid		
Streptococcus species	1			
Streptococcus pyogenes (Group A)	penicillin	• <i>Streptococcus pyogenes</i> (Group A) is 100% susceptible to penicillin		
Streptococcus agalactiae (Group B)	penicillin	• <i>Streptococcus agalactiae</i> (Group B) is 100% susceptible to penicillin		
Streptococcus pneumoniae	ceftriaxone	• If meningitis is suspected, add vancomycin		
Streptococcus PCR only	ceftriaxone	• When BCID2 is only able to identify <i>Streptococcus</i> species to the genus level, the species is not one of the included <i>Streptococcus</i> species on the BCID2 panel		
Other Gram Positives				
Listeria monocytogenes	ampicillin	Recommend ID consult		

Table 3. Gram-Negative Organism BCID2 Treatment Decision Algorithm			
BCID2 Result	Revised Empiric Therapy	Comments	
Enterobacterales with <u>NO resistance g</u>	enes detected		
Escherichia coli	ceftriaxone		
Klebsiella pneumoniae group	ceftriaxone		
Klebsiella oxytoca	ceftriaxone		
Klebsiella aerogenes (formerly known as Enterobacter aerogenes)	cefepime	• High rates of AmpC production – cefepime preferred	
Proteus species	ceftriaxone		
Enterobacter cloacae complex	cefepime	• High rates of AmpC production – cefepime preferred	
Serratia marcescens	ceftriaxone		
Salmonella species	ceftriaxone		
Enterobacterales (WITHOUT species identification)	cefepime or piperacillin/tazobactam	 Enterobacterales is the name of a family of gram-negative bacteria, it is not the <i>Enterobacter</i> genus When BCID2 is only able to identify Enterobacterales to the family level, the species is not one of the included Enterobacterales species on the BCID2 panel Consider source when choosing between cefepime or piperacillin-tazobactam (e.g. piperacillin-tazobactam for intra-abdominal infection due to additional anaerobic coverage) 	
Other Gram-Negative Organisms with	NO resistance genes detected	eororago)	
Acinetobacter calcoaceticus-baumannii complex	ampicillin/sulbactam	• Refer to CAUSE antimicrobial dosing guide for high dose ampicillin-sulbactam dosing	
Pseudomonas aeruginosa	cefepime or piperacillin/tazobactam		
Stenotrophomonas maltophilia	sulfamethoxazole/ trimethoprim	Refer to CAUSE antimicrobial dosing guide for optimal dosing	
Haemophilus influenzae	ampicillin/sulbactam or ceftriaxone		
Neisseria meningitidis	ceftriaxone	Recommend ID consult	
Bacteroides fragilis	metronidazole	• For polymicrobial infections requiring additional coverage, piperacillin/tazobactam will treat <i>B. fragilis</i> (additional anaerobic coverage (e.g. metronidazole) is not needed)	
Resistance Genes Detected			
Positive CTX-M (ESBL)	carbapenem*	 Indicates an extended-spectrum beta-lactamase (ESBL) producing organism In clinically stable patients with a suspected or confirmed urinary source, piperacillin/tazobactam is usually an appropriate alternative* CAUSE/ID approval required for carbapenems 	
Positive KPC or OXA48-like	Consult ID/CAUSE to discuss preferred therapy ceftazidime/avibactam	 Indicates a carbapenem resistant organism CAUSE/ID approval required for ceftazidime/avibactam 	
Positive IMP, NDM, VIM	Consult ID/CAUSE to discuss preferred therapy cefiderocol	 Indicates a carbapenem resistant organism producing a metallo-beta-lactamase CAUSE/ID approval required for cefiderocol 	
Positive mcr-1	Treat as indicated above	Indicates a colistin reistant organism	

Table 4. Fungal Organism BCID2 Treatment Decision Algorithm			
BCID2 Result	Revised Empiric Therapy	Comments	
Candida species			
Candida albicans	micafungin	Recommend ID consult	
Candida auris	Consult ID/CAUSE to	Recommend ID consult	
	discuss preferred empiric therapy	• High rates of resistance possible	
	micafungin		
Candida glabrata	micafungin	Recommend ID consult	
Candida krusei	micafungin	Recommend ID consult	
		Intrinsically resistant to fluconazole	
Candida parapsilosis	micafungin	• Recommend ID consult	
		• Isolates can have higher MICs to echinocandins, consider fluconazole as alternative if concerned for echinocandin resistance or non-response	
Candida tropicalis	micafungin	Recommend ID consult	
Other Fungal Organisms			
Cryptococcus (C. neoformans/C.	Consult ID/CAUSE to		
gattii)	discuss preferred empiric therapy	• CAUSE/ID approval required for liposomal amphotericin B	
	liposomal amphotericin B + flucytosine		

Table 5. What to do when no organisms are detected on BCID2?

The BCID2 panel is **ONLY** capable of detecting organisms and resistance genes included in table 1. Absence of a detected organism does **NOT** indicate a false-positive culture or reporting error.

When no organisms are detected on BCID2 consider the blood culture gram stain results (e.g. gram positive or negative), growth environment (e.g. aerobic, anaerobic, or both), gaps in antimicrobial coverage, past culture history, suspected source of bacteremia and clinical stability of patient when deciding on an appropriate empiric regimen.