BIOFIRE[®] FILMARRAY[®] Gastrointestinal (GI) Panel Mid

Instructions for Use	https://www.biofiredx.com/e-labeling/ITI0203					
Quick Guide	https://www.biofiredx.com/e-labeling/ITI0211					
Safety Data Sheet (SDS)	https://www.biofiredx.com/e-labeling/ITI0227					
Pouch Module	https://www.biofiredx.com/e-labeling/ITIFA20GIMid21					
Customer and Technical	Phone: 1-800-735-6544 (toll free)					
Customer and Technical Support Information	Phone: 1-800-735-6544 (toll free) Email: <u>BioFireSupport@biomerieux.com</u>					



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Rx Only

INTENDED PURPOSE

Intended Use

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The BIOFIRE® FILMARRAY® Gastrointestinal (GI) Panel Mid is an automated qualitative multiplexed nucleic acid-based *in vitro* diagnostic test intended for use with BIOFIRE® FILMARRAY® Systems. The BIOFIRE FILMARRAY GI Panel Mid is capable of the simultaneous detection and identification of nucleic acids from multiple bacteria, viruses, and parasites directly from stool samples in Cary Blair transport media obtained from individuals with signs and/or symptoms of gastrointestinal infection. The following bacteria, parasites, and viruses are identified using the BIOFIRE FILMARRAY GI Panel Mid:

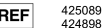
Bacteria	Viruses
Campylobacter (C. jejuni/C. coli/ C. upsaliensis)	Norovirus GI/GII
Clostridioides (Clostridium) difficile (toxin A/B)	
Salmonella	Parasites
Shiga-like toxin-producing E. coli (STEC) stx1/stx2	Cryptosporidium
Shigella/Enteroinvasive E. coli (EIEC)	Cyclospora cayetanensis
Vibrio (V. parahaemolyticus/V. vulnificus/ V. cholerae)	Giardia lamblia
Yersinia enterocolitica	

The BIOFIRE FILMARRAY GI Panel Mid is indicated as an aid in the diagnosis of specific agents of gastrointestinal illness and results are meant to be used in conjunction with other clinical, laboratory, and epidemiological data. Positive results do not rule out co-infection with organisms not included in the BIOFIRE FILMARRAY GI Panel Mid. The agent detected may not be the definite cause of the disease.

Concomitant culture is necessary for organism recovery and further typing of bacterial agents.

This device is not intended to monitor or guide treatment for C. difficile infection.

Due to the small number of positive specimens collected for certain organisms during the prospective clinical study, performance characteristics for *Yersinia enterocolitica* were established primarily with retrospective clinical specimens.



Performance characteristics for *Vibrio* (*V. parahaemolyticus, V. vulnificus,* and *Vibrio cholerae*) were established primarily using contrived clinical specimens.

Negative BIOFIRE FILMARRAY GI Panel Mid results in the setting of clinical illness compatible with gastroenteritis may be due to infection by pathogens that are not detected by this test or non-infectious causes such as ulcerative colitis, irritable bowel syndrome, or Crohn's disease.

A gastrointestinal microorganism multiplex nucleic acid-based assay also aids in the detection and identification of acute gastroenteritis in the context of outbreaks.

Intended User and Use Environment

The BIOFIRE FILMARRAY GI Panel Mid is intended for use by trained medical and laboratory professionals in a laboratory setting or under the supervision of a trained laboratory professional.

SUMMARY AND EXPLANATION OF THE TEST

Despite advances in food safety, sanitation, and medical treatment, infectious gastroenteritis remains a significant problem in industrialized countries among all age groups. In the United States, around 76 million cases of foodborne disease, resulting in 325,000 hospitalizations and 5,000 deaths, are estimated to occur each year.^{1–3} Additionally, there are over 300,000 *C. difficile* diagnoses per year in the US⁴ resulting in estimated costs of at least \$1 billion.⁵ Globally, infectious diarrheal illness is a significant cause of mortality in young children resulting in an estimated 800,000 deaths per year in children under the age of 5.⁶ In addition to this significant morbidity and mortality, diarrhea in children contributes to malnutrition, increased susceptibility to other infections, and may lead to delays in growth and intellectual development.^{7,8} The BIOFIRE FILMARRAY GI Panel Mid simultaneously tests for 11 pathogens (Table 1) from stool specimens collected in Cary Blair transport medium. Results from the BIOFIRE FILMARRAY GI Panel Mid test are available within about one hour.

Bacteria	Viruses
Campylobacter (C. jejuni/C. coli/ C. upsaliensis)	Norovirus GI/GII
Clostridioides (Clostridium) difficile (toxin A/B)	
Salmonella	Parasites
Shiga-like toxin-producing E. coli (STEC) stx1/stx2	Cryptosporidium
Shigella/Enteroinvasive E. coli (EIEC)	Cyclospora cayetanensis
Vibrio (V. parahaemolyticus/V. vulnificus/ V. cholerae)	Giardia lamblia
Yersinia enterocolitica	

Table 1. Bacteria, Viruses, and Parasites Detected by the BIOFIRE FILMARRAY GI Panel Mid

Summary of Detected Organisms

Bacteria

Campylobacter (C. jejuni/C. coli/C. upsaliensis). Campylobacter are gram-negative, non-spore forming, s-shaped or spiral bacteria that are usually motile. Most sporadic infections are acquired through ingestion of undercooked poultry or from cross-contamination of other foods. Outbreaks have been associated with unpasteurized dairy, contaminated water, poultry, and produce. Transmission from the stool of household pets has also been documented.⁹ *C. jejuni* and *C. coli* are the species most commonly associated with diarrheal illness, followed distantly by *C. upsaliensis*. Other species such as *C. lari* and *C. fetus* are more uncommon.¹⁰ Infection with *Campylobacter* species is common throughout the world, representing a large and perhaps under-recognized health burden.¹¹ *Campylobacter* are a leading cause of bacterial enteritis in the US (est. 845,000 infections annually with almost 8,500 hospitalizations¹²) and the most common cause of foodborne illness in the EU (over 220,000 confirmed cases reported by EU member states in 2011¹³). Enteric *Campylobacter* infections range from asymptomatic to severe infections characterized by bloody or non-bloody diarrhea, fever, and abdominal cramping. Infections may also lead to long-term health issues such as Guillain-Barré syndrome (GBS) and reactive arthritis.¹¹ *Campylobacter* infections are a notifiable disease in the US and are tracked by the European Surveillance System (TESSy).

Clostridium difficile (re-classified as Clostridioides difficile) are obligately anaerobic, gram-positive rods capable of forming hardy spores and are widespread in nature. These bacteria are acquired from the environment or transmitted via the fecal-oral route. Some *C. difficile* strains produce two enterotoxins, toxin A and toxin B, that damage the large intestine of the infected individual. *C. difficile* infection (CDI) is the major cause of hospital-acquired diarrhea and is responsible for more than 300,000 cases of diarrheal disease and 14,000 deaths annually in the US, resulting in over a billion dollars in health care costs.¹⁴ CDI presents a similar healthcare burden in the EU.¹⁵ Antibiotic treatment, which severely disrupts the normal gastrointestinal flora, is a major risk factor for the development of CDI. Community-acquired CDI, which has a

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somewhat lower association with antibiotic exposure, has also been emerging in the last few years.¹⁶ Clinical manifestations of *C. difficile* infection range from asymptomatic carriage (estimated to occur in 3-5% of healthy adults and up to 30% of healthy neonates¹⁷) to pseudomembranous colitis, involving bloody diarrhea, severe abdominal pain, and fever. Due to the high asymptomatic carriage rates, especially in young children, the clinical relevance of the detection of toxigenic *C. difficile* from stool should be considered in the context of other clinical findings, patient age, and risk factors including hospitalization and antibiotic exposure.^{18,19}

Salmonella. Salmonella enterica and *S. bongori* are the sole members of the Salmonella genus. Greater than 2,500 different serotypes of Salmonella have been recognized, with the majority of pathogenic serotypes being within the *S. enterica* species.²² These motile, rod-shaped, gram-negative, facultative bacteria are commonly recognized as a food contaminant associated with meat, poultry, produce, and manufactured products. Salmonella may be classified as typhoidal and non-typhoidal based on the disease that they cause. The non-typhoidal Salmonella are associated with intestinal illness resulting in acute, watery diarrhea, often with fever, and are a common cause of foodborne illness. Though rare in developed countries, it is common in the developing world (>70% of US cases are related to foreign travel).² In contrast, infection with non-typhoidal Salmonella is one of the most common causes of foodborne illness in the US and EU mon-typhoidal Salmonella is one of the most common causes of foodborne illness in the US and EU with greater than one million cases per year.^{12,13} While large outbreaks do occur, the majority of cases are sporadic with peak in incidence in late summer/early fall. The highest incidence is seen in children aged <5 years.²³ In general Salmonella-related gastroenteritis is self-limiting, except in cases of severe or typhoidal illness. Salmonellosis is a notifiable disease in the US and is tracked by TESSy in the EU.

Vibrio (V. parahaemolyticus/V. vulnificus/V. cholerae). Vibrio are motile, gram-negative, comma-shaped bacteria typically found in marine environments. Several species are capable of causing illness in humans, both extraintestinal (soft tissue infection, septicemia, eye and ear infections) and intestinal. Gastrointestinal illness is most commonly associated with *V. cholerae, V. parahaemolyticus, V. vulnificus, V. fluvialis, V.mimicus* or *V. alginolyticus* and infections are associated with consumption of contaminated food, particularly in coastal regions.²⁴

V. cholerae is the only *Vibrio* species that causes endemic, epidemic, and pandemic cholera. There are three major subgroups of *V. cholerae*: *V. cholerae* O1, *V. cholerae* O139, and *V. cholerae* non-O1/non-O139. Classic cholera is characterized by passing copious amounts of watery diarrhea leading to extreme dehydration and death. Severe disease is mediated by the presence of cholera toxin (CTX). Cholera is endemic in many parts of the world and new outbreaks often follow natural disasters or social upheaval. As such, cholera remains a significant cause of morbidity and mortality in much of the world. In the US and EU, occasional cases of cholera are seen in travelers returning from overseas.

Vibriosis and cholera are notifiable diseases in the US and are tracked by the Cholera and Other *Vibrio* Illness Surveillance Network (COVIS). While *V. cholerae* infections are exceedingly rare in the US, other *Vibrio* species are estimated to cause approximately 50,000 food-borne infections per year^{12,13} though only ~400 isolates recovered from stool were reported to COVIS in 2009 (the majority of which were *V. parahaemolyticus*).²⁴ This discrepancy between estimated prevalence and actual detections is due to specialized testing required to recover *Vibrio* organisms from stool, leaving most cases undiagnosed. The risk of *Vibrio* infection in Europe is thought to be very low and is not tracked by TESSy.²⁵

Yersinia enterocolitica are small, gram-negative bacilli, which generally appear as single cells or short chains. *Y. enterocolitica* is transmitted through ingestion of contaminated food or water, often raw undercooked meats (especially pork), and is estimated to cause almost 100,000 foodborne illnesses in the US annually (though only about 1,000 cases are laboratory-confirmed; possibly because *Y. enterocolitica* are not identified by routine enteric pathogen testing).¹² A higher incidence of Yersiniosis is observed in European countries, particularly in continental Europe²⁶ with nearly 7,000 confirmed cases reported in 2011.¹³ The severity of the illness is based on the serotype of the infecting strain and ranges from self-limiting gastroenteritis to terminal ileitis and mesenteric lymphadenitis. Symptoms of illness mimic appendicitis and may lead to unnecessary surgery, highlighting the importance of properly identifying this organism when it is present in stool specimens. Yersiniosis is a notifiable disease in the US and is tracked by TESSy in the EU.

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Shiga-like toxin-producing *E. coli* (STEC). There are two main types of Shiga-like toxins, Shiga-like toxin 1 (Stx1) and Shiga-like toxin 2 (Stx2) (also known as verotoxins). Shiga-like toxin-producing *E. coli* (STEC) may contain either one or both *stx* genes. STEC are a primary cause of bloody diarrhea^{10,36} and can progress to a potentially fatal condition known as hemolytic uremic syndrome (HUS; caused by Shiga-like toxin destruction of red blood cells that leads to renal failure), especially in the very young and very old. STEC are important foodborne pathogens. Infections may also be waterborne, transmitted person-to-person, or via contact with animals (especially cattle, which are a reservoir for STEC). Antimicrobial therapy for STEC may lead to an increased risk for HUS, especially in antibiotic-resistant strains, potentially by up-regulating production and thus increasing the amount of Shiga-like toxin available for absorption. Therefore, identification of Shiga-like toxin genes in a patient with gastrointestinal illness can aid in the decision of whether or not to prescribe antibiotics for patient care.

*Shigella/*Enteroinvasive *E. coli* (EIEC). There are four subgroups of *Shigella* species: subgroup A (*S. dysenteriae*), subgroup B (*S. flexneri*), subgroup C (*S. boydii*), and subgroup D (*S. sonnei*). All *Shigella* are non-motile, gram-negative rods which are typically transferred through person-to-person contact or ingestion of contaminated food or water (humans and other primates are the only known animal reservoirs). Infections are most common where hygiene is compromised, for example institutional settings (day care, nursing homes) and may become endemic in developing societies without running water and indoor plumbing.¹⁰ *Shigella* are responsible for multiple illnesses including shigellosis and bacillary dysentery which can result in bloody or non-bloody diarrhea.

Enteroinvasive *E. coli* (EIEC), unlike most *E. coli* do not decarboxylate lysine, and do not ferment lactose. EIEC strains contain a plasmid encoding virulence factors (such as invasion plasmid antigen *ipaH*) that allow the bacteria to invade the colon and produce a watery diarrhea syndrome that is identical to that caused by *Shigella*. *Shigella* and EIEC infections are generally treated in the same manner.

Multiple copies of the *ipaH* gene are present in all four *Shigella* species (*S. dysenteriae*, *S. flexneri*, *S. boydii*, and *S. sonnei*) as well as in the virulence plasmid of Enteroinvasive *E. coli* (EIEC).^{38,39} *IpaH*, along with other factors encoded by the invasion plasmid, mediate entry of *Shigella* and EIEC into host cells. This is a common target for laboratory developed molecular tests.^{38,39}

There are an estimated 130,000 *Shigella* infections associated with foodborne illness in the US each year¹² however no data exists for EIEC. Shigellosis is a notifiable disease in the US and is tracked by TESSy in the EU.

Parasites

Cryptosporidium is a genus of protozoa capable of causing infections of the human stomach, intestine, and biliary ducts following ingestion of chlorine-tolerant oocysts which are shed in fecal material and can contaminate drinking water, recreational water, or food. *Cryptosporidium* are among the most common parasitic causes of diarrhea in developed nations.⁴⁰ There are an estimated 60,000 illnesses every year in the US due to *Cryptosporidium* infection¹² with rates being highest in summer months.²³ At least 10 species infect humans though *C. hominis* and *C. parvum* are the most common.¹⁰ Illness is generally characterized by short-term gastroenteritis that resolves without treatment. However, severe illness is possible in immunocompromised individuals, particularly those with AIDS, where illness resolves slowly or not at all and can be fatal. Cryptosporidiosis is a notifiable disease in the US and is tracked by TESSy in the EU.

Cyclospora cayetanensis are parasitic protozoa that cause gastroenteritis in humans, which are the only known hosts. Unsporulated oocysts are disseminated in feces. After a period of maturation (days to weeks), the oocysts become infectious and can cause illness if ingested through contaminated food or water.¹⁰ Infections are most common in tropical, subtropical, or warm temperate regions. In the US and EU, infections are associated with travelers' diarrhea in persons returning from endemic areas. Additionally, outbreaks have been associated with consumption of contaminated food from other countries.^{10,41} There are an estimated 11,000 foodborne illnesses due to *C. cayetanensis* infections annually in the US¹² but the true incidence may be underestimated due to the difficulty of diagnosing infection.⁴² Illness presents as non-bloody



diarrhea that may be up to several months in duration. Cyclosporiasis is a notifiable disease in the US but is not tracked by TESSy in the EU.

Giardia lamblia (also referred to as *G. duodenalis* and *G. intestinalis*) are intestinal flagellate parasites found world-wide. *Giardia* are the most common intestinal parasites isolated in the US and EU and are a leading cause of parasitosis worldwide.^{12,13,40} Populations with the highest risk of *G. lamblia* infection include children in day care centers, hikers, and the immunocompromised. *G. lamblia* prevalence is about 1-7% in developed countries and as high as 50% in developing countries.¹⁰ Transmission occurs through ingestion of contaminated food or water, with approximately 77,000 foodborne illnesses in the US annually.¹² Infection rates are highest during summer months.²³ The majority of *G. lamblia* infections are generally self-limiting; though symptoms are long-lasting, and some patients go on to develop chronic illness, which can lead to complications. Giardiasis is a notifiable disease in the US and is tracked by TESSy in the EU.

Viruses

Norovirus GI/GII. Noroviruses are highly contagious members of the *Caliciviridae* family of RNA viruses and can be divided into at least ten genogroups (GI – GX) and many genotypes and variants within each genogroup based on VP1 sequences.⁴⁸ Genogroups GI, GII, and GIV have been found most commonly in humans (though GIV is very rare) where they cause moderate to severe gastroenteritis consisting primarily of nausea, vomiting, and diarrhea accompanied by fever. Transmission occurs via the fecal-oral route or through aerosolized vomitus and the infectious dose may be as low as 18 particles.⁴⁹ Symptoms of infection generally last 24-48 hours⁵⁰ and the illness is self-limiting; though immunocompromised persons may suffer chronic diarrhea and some children have been reported to develop necrotizing colitis. Outbreaks are common in closed communities such as cruise ships, hospitals, nursing homes, schools, and military installations. Norovirus infections are the leading cause of foodborne gastroenteritis in the US, causing nearly 5.4 million illnesses (and over 14,000 hospitalizations) annually¹² and are also a significant source of illness in the EU.¹³ Peak infection rates occur during winter months.⁵¹ Immunity following Norovirus illness is short lived as re-infection is possible within 6 months, even in the presence of high serum antibody titers.⁵²

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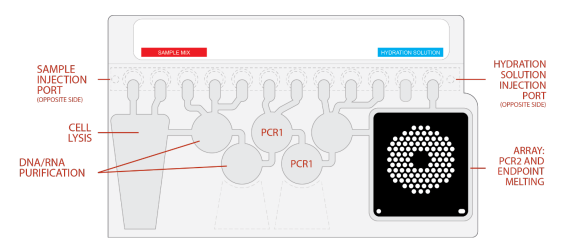
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Principle of the Procedure

The BIOFIRE FILMARRAY GI Panel Mid pouch is a closed system disposable that stores all the necessary reagents for sample preparation, reverse transcription, polymerase chain reaction (PCR), and detection in order to isolate, amplify, and detect nucleic acid from multiple gastrointestinal pathogens from a single stool specimen. After sample collection, the user injects Hydration Solution and sample combined with Sample Buffer into the pouch, places the pouch into a BIOFIRE[®] FILMARRAY[®] System module, and starts a run. The entire run process takes about one hour. Additional detail can be found in the appropriate BIOFIRE System Operator's Manual.

During a run, the BIOFIRE System:

- Lyses the sample by agitation (bead beating) in addition to chemical lysis mediated by the Sample Buffer
- Extracts and purifies all nucleic acids from the sample using magnetic bead technology.
- Performs nested multiplex PCR by:
 - First performing a single, large volume, massively multiplexed reaction (PCR1).
 - Then performing multiple simultaneous second-stage PCR reactions (PCR2) in the array to amplify sequences within the PCR1 products.
- Uses endpoint melting curve data to detect and generate a result for each target on the BIOFIRE FILMARRAY GI Panel Mid array.



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MATERIALS PROVIDED

The BIOFIRE FILMARRAY GI Panel Mid contains materials consisting of primers, buffers, dNTPs, polymerase, molecular grade water, guanidinium chloride (50 - < 60%), Triton-X 100 (10 - < 20%), and LCGreen® Plus.

Each kit contains sufficient reagents to test 6 or 30 samples (6 test kit- REF #425089 or 30 test kit- REF #424898):

- Individually packaged BIOFIRE FILMARRAY GI Panel Mid pouches
- Sample Preparation Reagent Kits (SPRKs)
 - Single-use Sample Buffer ampoules
 - Single-use pre-filled Hydration Injection Vials (blue)
 - Single-use Sample Injection Vials (red)
 - Individually packaged Transfer Pipettes
- BIOFIRE FILMARRAY GI Panel Mid Pouch Module Software This software is required to run the BIOFIRE FILMARRAY GI Panel Mid and can be downloaded at

https://www.biofiredx.com/e-labeling/ITIFA20GIMid21.

NOTE: The BIOFIRE FILMARRAY GI Panel Mid Pouch Module software and the BIOFIRE System Software provides customers with the option for selective reporting of *Clostridioides* (*Clostridium*) difficile (toxin A/B). This is an optional feature. Please refer to the appropriate BIOFIRE System Operator's Manual for more information.

MATERIALS REQUIRED BUT NOT PROVIDED

- BIOFIRE System including:
 - BIOFIRE 2.0 or BIOFIRE TORCH Systems
 - Including accompanying system-specific core software and panel-specific software
 - o BIOFIRE® Pouch Loading Station
- 10% bleach solution or similar disinfectant

WARNINGS AND PRECAUTIONS

General Precautions

- 1. For *in vitro* diagnostic use only.
- 2. A trained healthcare professional should carefully interpret the results from the BIOFIRE FILMARRAY GI Panel Mid in conjunction with a patient's signs and symptoms, results from other diagnostic tests, and relevant epidemiological information.
- 3. BIOFIRE FILMARRAY GI Panel Mid pouches are only for use with BIOFIRE 2.0 and BIOFIRE TORCH Systems.
- 4. Performance characteristics of the BIOFIRE FILMARRAY GI Panel Mid have only been determined with stool specimens in Cary Blair transport media.

- 5. Always check the expiration date on the kit. Do not use kit components or a pouch after its expiration date.
- 6. BIOFIRE FILMARRAY GI Panel Mid pouches are stored under vacuum in individually wrapped canisters. To preserve the integrity of the pouch vacuum for proper operation, be sure that an instrument module will be available and operational before unwrapping any pouches for loading.
- 7. Contamination can cause unexpected positive results in negative or positive external controls. If unexpected positive results are observed, thoroughly clean and decontaminate the workspace and contact customer support if the unexpected results persist.

Safety Precautions

- Wear appropriate Personal Protective Equipment (PPE), including (but not limited to) disposable powder-free gloves and lab coats. Protect skin, eyes and mucus membranes. Change gloves often when handling reagents or samples.
- 2. Handle all samples and waste materials as if they are capable of transmitting infectious agents. Observe safety guidelines such as those outlined in:
 - CDC/NIH Biosafety in Microbiological and Biomedical Laboratories,⁶³
 - CLSI Document M29 Protection of Laboratory Workers from Occupationally Acquired Infections,⁶⁴
- 3. Follow your institution's safety procedures for handling biological samples.
- 4. Dispose of materials used in this assay, including reagents, samples, and used buffer ampoules, according to federal, state, regional, and local regulations.
- 5. Sample Buffer contains Guanidinium chloride and Triton X100.

The following statements apply:

- Health Hazards
 - Acute Toxicity, oral (Category 4)
 - H302 Harmful if swallowed.
 - Skin corrosion/irritation (Category 2)
 - H315 Causes skin irritation.
 - Serious eye damage/eye irritation (Category 1)
 - H318 Causes serious eye damage.
- Environment Hazards
 - Hazardous to the aquatic environment, acute aquatic hazard (Category 1)
 - H400 Very toxic to aquatic life.
 - Hazardous to the aquatic environment, long-term aquatic hazard (Category 1)
 - H410 Very toxic to aquatic life with long lasting effects.
- Precautionary Statements
 - Prevention
 - P273 Avoid release to the environment.
 - P280 Wear protective gloves/protective clothing/eye protections/face protection.
- Response
- P332 + P313 If skin irritation occurs: Get medical advice/attention.

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P305 + P351 + P338 - IF IN EYES: Rinse cautiously with water for several minutes. Remove contact lenses, if present and easy to do. Continue rinsing.

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- P301 + P312 IF SWALLOWED: Call a POISON CENTRE/doctor if you feel unwell.
- P337 + P313 If eye irritation persists: Get medical advice/attention.

Please refer to the BIOFIRE FILMARRAY GI Panel Mid Safety Data Sheet (SDS) for more information: <u>https://www.biofiredx.com/e-labeling/ITI0227.</u>

Sample Buffer will form hazardous compounds and fumes when mixed with bleach or other disinfectants.

WARNING: Never add Bleach to Sample Buffer or sample waste.

- 6. Bleach, a recommended disinfectant, is corrosive and may cause severe irritation or damage to eyes and skin. Vapor or mist may irritate the respiratory tract. Bleach is harmful if swallowed or inhaled.
 - Eye contact: Hold eye open and rinse with water for 15-20 minutes. Remove contact lenses after the first 5 minutes and continue rinsing eye. Seek medical attention.
 - Skin contact: Immediately flush skin with plenty of water for at least 15 minutes. If irritation develops, seek medical attention.
 - Ingestion: Do not induce vomiting. Drink a glassful of water. If irritation develops, seek medical attention.
 - Please refer to the appropriate Safety Data Sheet (SDS) for more information.

Laboratory Precautions

1. Preventing organism contamination

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Due to the sensitive nature of the BIOFIRE FILMARRAY GI Panel Mid, it is important to guard against contamination of the work area by carefully following the testing process outlined in this booklet, including these guidelines:

- Laboratory personnel may carry or shed common gastrointestinal pathogens asymptomatically and can
 inadvertently contaminate the specimen while it is being processed. Stool samples may also contain a
 high concentration of organisms. Careful adherence to the sample processing steps described in this
 document is recommended to avoid possible contamination. Samples should be processed in a clean
 biosafety cabinet if available, or according to local laboratory guidelines. If a biosafety cabinet is not used,
 a dead air box (e.g., AirClean PCR workstation), a splash shield (e.g., Bel-Art Scienceware Splash
 Shields), or a face shield can be used when preparing samples instead.
- It is recommended to avoid handling specimens or pouches in an area used to routinely process stool pathogen testing (e.g., culture, EIA).
- Prior to processing specimens, thoroughly clean both the work area and the Pouch Loading Station using a suitable cleaner such as freshly prepared 10% bleach or a similar disinfectant. To avoid residue buildup and potential damage to the specimen or interference from disinfectants, wipe disinfected surfaces with water.
- Specimens and pouches should be handled and/or tested one-at-a-time. Always change gloves and clean the work area between each pouch and specimen.
- Use clean gloves when removing Sample Buffer ampoules and Sample/Hydration Injection Vials from bulk packaging bags and reseal packaging bags when not in use.

2. Preventing amplicon contamination

A common concern with PCR-based assays is false positive results caused by contamination of the work area with PCR amplicon. Because the BIOFIRE FILMARRAY GI Panel Mid pouch is a closed system, the risk of amplicon contamination is low provided that pouches remain intact after the test is completed. Adhere to the following guidelines, in addition to those above, to prevent amplicon contamination:

- Discard used pouches in an appropriate biohazard container immediately after the run has completed.
- Avoid excessive handling of pouches after test runs.
- Change gloves after handling a used pouch.
- Avoid exposing pouches to sharp edges or anything that might cause a puncture.

WARNING: If liquid is observed on the exterior of a pouch, the liquid and pouch should be immediately contained and discarded in a biohazard container. The module and workspace must be decontaminated as described in the appropriate BIOFIRE Operator's Manual.

DO NOT PERFORM ADDITIONAL TESTING UNTIL THE AREA HAS BEEN DECONTAMINATED.

3. Cary Blair media may contain non-viable organisms and/or nucleic acids at levels that can be detected by the BIOFIRE FILMARRAY GI Panel Mid.

The presence of non-viable organisms and/or nucleic acids in Cary Blair may lead to false positive test results.

Precaution Related to Public Health Reporting in the United States

Local, state, federal, and country regulations for notification of reportable disease are continually updated and include a number of organisms for surveillance and outbreak investigations^{65,66}.

The U.S. Centers for Disease Control (CDC) recommends that when pathogens from reportable diseases are detected by a culture independent diagnostic test (CIDT), the laboratory should facilitate obtaining the isolate or clinical materials for submission to the appropriate public health laboratory to aid in outbreak detection and epidemiological investigations.

Laboratories are responsible for following their state, local, and/or country regulations and should consult their local and/or state public health laboratories for isolate and/or clinical sample submission guidelines.

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REAGENT STORAGE, HANDLING AND STABILITY

- 1. Store the test kit, including reagent pouches and buffers, at room temperature (15-25 °C).
- 2. Avoid storage of any materials near heating or cooling vents or in direct sunlight.
- 3. Always check the expiration date on the kit. Do not use reagents beyond the expiration date printed on the pouch or kit.
- 4. All kit components should be stored and used together. Do not use components from one kit with those of another kit. Discard any extra components from the kit after all pouches have been consumed.
- 5. Do not remove pouches from their packaging until a sample is ready to be tested. Once the pouch packaging has been opened, the pouch should be loaded as soon as possible (within approximately 30 minutes).
- 6. Once a pouch has been loaded, the test run should be started as soon as possible (within approximately 60 minutes). Do not expose a loaded pouch to temperatures above 40°C (104°F) prior to testing.

SAMPLE REQUIREMENTS

The following table describes the requirements for specimen collection, preparation, and handling that will help ensure accurate test results.

Stool Specimen Collection	Stool specimens should be collected in Cary Blair transport media according to manufacturer's instructions.
Minimum Sample Volume	0.2 mL (200 μ L) of sample is required for testing
	Specimens should be tested with the BIOFIRE FILMARRAY GI Panel Mid as soon as possible. ^a
Transport and Storage	 If storage is required, specimens can be held: At room temperature for up to 4 days Refrigerated for up to 4 days Note: refer to Cary Blair transport media manufacturer's instructions for any additional transport and storage recommendations.

Table 2. Sample requirements for the BIOFIRE FILMARRAY GI Panel Mid

^a The performance validation included the evaluation of clinical specimens that were frozen at -70°C or lower for up to 90 days. However, longer frozen storage may be acceptable. Please follow your institutions rules and protocols regarding sample storage validation.

PROCEDURE

Refer to the BIOFIRE FILMARRAY GI Panel Mid Quick Guide or the appropriate BIOFIRE System Operator's Manual for more detail and pictorial representations of these instructions.

Use clean gloves and other Personal Protective Equipment (PPE) when handling pouches and samples. Only prepare one BIOFIRE FILMARRAY GI Panel Mid pouch at a time. Once sample is added to the pouch, promptly transfer to the instrument to start the run. After the run is complete, discard the pouch in a biohazard container.

Step 1: Prepare Pouch

1. Thoroughly clean the work area and the Pouch Loading Station with freshly prepared 10% bleach (or suitable disinfectant) followed by a water rinse.

2. Remove the pouch from its vacuum-sealed package by tearing or cutting the notched outer packaging and opening the protective canister. Use caution when opening the vacuum-sealed package as the edges may be sharp.

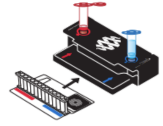
NOTE: The pouch may still be used even if the vacuum seal of the pouch is not intact. Attempt to hydrate the pouch using the steps in the Hydrate Pouch Section. If hydration is successful, continue with the run. If hydration fails, discard the pouch and use a new pouch to test the sample.

- 3. Check the expiration date on the kit. Do not use expired kit components or pouches.
- 4. Insert the pouch into the Pouch Loading Station, aligning the red and blue labels on the pouch with the red and blue arrows on the Pouch Loading Station.
- 5. Place a red-capped Sample Injection Vial into the red well of the Pouch Loading Station.
- 6. Place a blue-capped Hydration Injection Vial into the blue well of the Pouch Loading Station.

.....

Step 2: Hydrate Pouch

- 1. Unscrew the Hydration Injection Vial from the blue cap.
- 2. Remove the Hydration Injection vial, leaving the blue cap in Pouch Loading Station.
- 3. Insert the cannula tip of the Hydration Injection Vial into the pouch hydration port located directly below the blue arrow of the Pouch Loading Station.
- 4. Forcefully push down in a firm and quick motion to puncture seal until a faint "pop" is heard and there is an ease in resistance. Wait as the correct volume of Hydration Solution is pulled into the pouch by vacuum.
 - If the Hydration Solution is not automatically drawn into the pouch, repeat
 Step 2 to verify that the seal of the pouch hydration port was broken. If Hydration Solution is again not drawn into the pouch, discard the current pouch, and retrieve a new pouch, and repeat from *Step 1: Prepare Pouch.*
- 5. Verify that the pouch has been hydrated.





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BIOFIRE® FILMARRAY® Gastrointestinal Panel Mid

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- Flip the barcode label down and check to see that fluid has entered the reagent wells (located at the base of the rigid plastic part of the pouch). Small air bubbles may be seen.
- If the pouch fails to hydrate (dry reagents appear as white pellets), repeat Step 2 to verify that the seal of the pouch hydration port was broken. If Hydration Solution is still not drawn into the pouch, discard the current pouch, retrieve a new pouch, and repeat from *Step 1: Prepare Pouch*.

Step 3: Prepare Sample Mix

- 1. Add Sample Buffer to the Sample Injection Vial.
 - Hold the Sample Buffer ampoule so that the tip is facing up.

NOTE: Use care to avoid touching the tip during handling, as this may introduce contamination.

- To open the Sample Buffer ampoule:
 - Gently twist and remove tab at the tip of the Sample Buffer ampoule.
- Invert the ampoule over the red-capped Sample Injection Vial and dispense Sample Buffer using a slow, forceful squeeze followed by a second squeeze.
- NOTE: Avoid squeezing the ampoule additional times. This will generate foam, which should be avoided.

WARNING: The Sample Buffer is harmful if swallowed and can cause serious eye damage and skin irritation.

- 2. Mix the patient specimen by inversion.
- 3. Use the Transfer Pipette provided in the test kit to draw specimen to the second line (approximately 0.2 mL) of the Transfer Pipette.
- 4. Add the specimen to the Sample Buffer in the Sample Injection Vial.
- 5. Tightly close the lid of the Sample Injection Vial and discard the Transfer Pipette in a biohazard waste container.

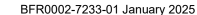
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NOTE: DO NOT use the transfer pipette to mix the sample once it is loaded into the Sample Injection Vial.

- 6. Remove the Sample Injection Vial from the Pouch Loading Station and invert the vial at least 3 times to mix.
- 7. Return the Sample Injection Vial to the red well of the Pouch Loading Station.

Step 4: Load Sample Mix

1. Slowly twist to unscrew the Sample Injection Vial from the red cap and wait for 5 seconds with the vial resting in the cap.

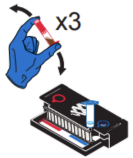




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NOTE: Waiting 5 seconds decreases the risk of dripping and contamination from the sample.

- 2. Lift the Sample Injection Vial, leaving the red cap in the well of the Pouch Loading Station, and insert the Sample Injection Vial cannula tip into the pouch sample port located directly below the red arrow of the Pouch Loading Station.
- 3. Forcefully Push down in a firm and quick motion to puncture seal (a faint "pop" is heard) and sample is pulled into the pouch by vacuum.
- 4. Verify that the sample has been loaded.
 - Flip the barcode label down and check to see that fluid has entered the reagent well next to the sample loading port.
 - If the pouch fails to pull sample from the Sample Injection Vial, the pouch should be discarded. Retrieve a new pouch and repeat from *Step 1: Prepare Pouch*.

- 5. Discard the Hydration Injection Vial and Sample Injection Vial in an appropriate biohazard sharps container.
- 6. Record the Sample ID in the provided area on the pouch label (or affix a barcoded Sample ID) and remove the pouch from the Pouch Loading Station.

Step 5: Run Pouch

The BIOFIRE® FILMARRAY® Software includes step-by-step on-screen instructions that guide the operator through performing a run. Brief instructions for BIOFIRE 2.0 and BIOFIRE TORCH Systems are given below. Refer to the appropriate BIOFIRE System Operator's Manual for more detailed instructions.

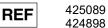
BIOFIRE 2.0

- 1. Ensure that the System (instrument and computer) is powered on and the software is launched.
- 2. Follow on-screen instructions and procedures described in the Operator's Manual to place the pouch in a module, enter pouch, sample, and operator information.
- 3. Pouch identification (Lot Number and Serial Number), Pouch Type and Protocol information will be automatically entered when the barcode is scanned. If it is not possible to scan the barcode, the pouch Lot Number, Serial Number, Pouch Type, and Protocol can be manually entered from the information provided on the pouch label into the appropriate fields. To reduce data entry errors, it is strongly recommended that the pouch information be entered by scanning the barcode

NOTE: When selecting a Pouch Type manually, ensure that the Pouch Type matches the label on the BIOFIRE FILMARRAY GI Panel Mid pouch.

- 4. Enter the Sample ID. The Sample ID can be entered manually or scanned in by using the barcode scanner when a barcoded Sample ID is used.
- 5. If necessary, select and/or confirm the appropriate protocol for your sample type from the Protocol drop down list. The BIOFIRE FILMARRAY GI Panel Mid has a single protocol available in the drop down list.





6. Enter a username and password in the Name and Password fields.

NOTE: The font color of the username is red until the username is recognized by the software.

7. Review the entered run information on the screen. If correct, select Start Run.

Once the run has started, the screen displays a list of the steps being performed by the instrument and the number of minutes remaining in the run.

NOTE: The bead-beater apparatus makes an audible, high-pitched noise during the first minute of operation.

8. When the run is finished, follow the on-screen instructions to remove the pouch, then immediately discard it in a biohazard waste container.

The run file is automatically saved in the BIOFIRE Software database, and the test report can be viewed, printed, and/or saved as a PDF file.

BIOFIRE TORCH

- 1. Ensure that the System is on.
- 2. Select an available Module on the touch screen or scan the barcode on the pouch using the barcode scanner.
- 3. Pouch identification (Lot Number and Serial Number), Pouch Type and Protocol information will be automatically entered when the barcode is scanned. If it is not possible to scan the barcode, the pouch Lot Number, Serial Number, Pouch Type, and Protocol can be manually entered from the information provided on the pouch label into the appropriate fields. To reduce data entry errors, it is strongly recommended that the pouch information be entered by scanning the barcode.

NOTE: When selecting a Pouch Type manually, ensure that the Pouch Type matches the label on the BIOFIRE FILMARRAY GI Panel Mid pouch.

- 4. Enter the Sample ID. The Sample ID can be entered manually or scanned in by using the barcode scanner when a barcoded Sample ID is used.
- 5. Insert the pouch into the available Module.
 - Ensure that the pouch fitment label is lying flat on top of pouch and not folded over. As the pouch is inserted, the Module will grab onto the pouch and pull it into the chamber.
- 6. If necessary, select and/or confirm the appropriate protocol for your sample type from the Protocol drop down list. The BIOFIRE FILMARRAY GI Panel Mid has a single protocol available in the drop down list.
- 7. Enter a username and password, then select Next.

NOTE: The font color of the username is red until the username is recognized by the software.

8. Review the entered run information on the screen. If correct, select Start Run.

Once the run has started, the screen displays a list of the steps being performed by the instrument and the number of minutes remaining in the run.

NOTE: The bead-beater apparatus can be heard as a high-pitched noise during the first minute of operation.

9. At the end of the run, remove the partially ejected pouch, then immediately discard it in a biohazard waste container.



10. The run file is automatically saved in the BIOFIRE Software database, and the test report can be viewed, printed, and/or saved as a PDF file.

QUALITY CONTROL

Process Controls

Two process controls are included in each pouch:

1. RNA Process Control

The RNA Process Control assay targets an RNA sequence from the yeast *Schizosaccharomyces pombe*. The yeast is present in the pouch in a freeze-dried form and becomes rehydrated when sample is loaded. The control material is carried through all stages of the test process, including lysis, nucleic acid purification, reverse transcription, 1st stage PCR, dilution, 2nd stage PCR and DNA melting. A positive control result indicates that all steps carried out in the pouch were successful.

2. PCR2 Control

The PCR2 Control assay detects synthetic DNA that is dried into wells of the array along with the corresponding primers. A positive result indicates that 2nd stage PCR was successful.

Both control assays must be positive for the test run to pass. If either control fails, the Controls field of the test report (upper right-hand corner) will display Failed and all results will be listed as Invalid. If the controls fail, the sample should be retested using a new pouch.

Monitoring Test System Performance

The BIOFIRE Software will automatically fail the run if the melting temperature (Tm) for either the RNA Process Control or the PCR2 Control is outside an acceptable range (80.2-84.2 for the RNA Process Control and 74.1-78.1 for the PCR2 Control). If required by local, state, or accrediting organization quality control requirements, users can monitor the system by trending Tm values for the control assays and maintain records according to standard laboratory quality control practices.^{67,68} Refer to the appropriate BIOFIRE FILMARRAY System Operator's Manual for instructions on control assay Tm values. The PCR2 Control is used in several BIOFIRE pouch types and can, therefore, be used to monitor the system when multiple pouch types are used on the same BIOFIRE System.

External Controls

External controls should be used in accordance with laboratory protocols and the appropriate accrediting organization requirements, as applicable. Cary Blair can be used as an external negative control. Previously characterized positive stool samples or negative samples spiked with well characterized organisms can be used as external positive controls. Commercial external control materials may be available from other manufacturers; these should be used in accordance with the manufacturers' instructions and appropriate accrediting organization requirements, as applicable.

INTERPRETATION OF RESULTS

The BIOFIRE Software automatically analyzes and interprets assay results and displays the results in a test report (see the BIOFIRE FILMARRAY GI Panel Mid Quick Guide to view an example of a test report). The analyses performed by the BIOFIRE Software and details of the test report are described below.

Assay Interpretation

When PCR2 is complete, the BIOFIRE instrument performs a high-resolution DNA melting analysis on the PCR products and records the change in fluorescence signal generated in each well (for more information see appropriate BIOFIRE System Operator's Manual). The BIOFIRE Software then performs several analyses and assigns a final assay result. The steps in the analysis are described below.

Analysis of melt curves. The BIOFIRE Software evaluates the DNA melt curve for each well of the PCR2 array to determine if a PCR product was present in that well. If the melt profile indicates the presence of a PCR product, then the analysis software calculates the melting temperature (Tm) of the curve and compares it against the expected Tm range for the assay. If the software determines that the Tm value falls inside the assay-specific Tm range, the melt curve is called positive. If the software determines that the melt curve is not in the appropriate Tm range, the melt curve is called negative.

Analysis of replicates. Once positive melt curves have been identified, the software evaluates the three replicates for each assay to determine the assay result. For an assay to be called positive, two of the associated melt curves must be called positive, and both Tm values must be similar. Assays that do not meet these criteria are called negative.

Organism Interpretation

For many organisms detected by the BIOFIRE FILMARRAY GI Panel Mid, the organism is reported as Detected if a single corresponding assay is positive. For example, *Salmonella* will have a result of *Salmonella* Detected if at least two of the three replicates of the Salm assay have similar positive melt peaks with Tm values that are within the assay-specific Tm range.

The following organisms are detected using a single assay: *Clostridioides* (*Clostridium*) *difficile* (toxin A/B), *Salmonella*, *Yersinia enterocolitica, Shigella//*Enteroinvasive *E. coli* (EIEC), *Cyclospora cayetanensis,* and *Giardia lamblia*.

Test results for several other organisms rely on the combination of multiple assays. These include *Campylobacter (C. jejuni/C. coli/C. upsaliensis)*, Shiga-like toxin-producing *E. coli* (STEC) *stx1/stx2*, *Vibrio* (*V. parahaemolyticus/ V. vulnificus/V. cholerae)*, *Cryptosporidium*, and Norovirus GI/GII. Interpretation rules for these assays are described below. Also included are summary descriptions of the assays' expected reactivity; see also the Analytical Reactivity (Inclusivity) section.

NOTE: If four or more distinct organisms are detected in a specimen, retesting is recommended to confirm the polymicrobial result.

Bacteria

Campylobacter (C. jejuni/C. coli/C. upsaliensis)

The BIOFIRE FILMARRAY GI Panel Mid contains two assays (Campy 1 and Campy 2) designed to together detect, but not differentiate, the most common *Campylobacter* species associated with human gastrointestinal illness: *C. jejuni, C.*

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coli, and *C. upsaliensis*. These are the same three species that are identified using standard clinical laboratory practices. Other *Campylobacter* species will not be identified by the BIOFIRE FILMARRAY GI Panel Mid. Empirical testing and *in silico* sequence analysis indicates reduced sensitivity for a less common subspecies of *C. jejuni* (*C. jejuni* subsp. *doylei*). A positive result for one or both assays will give a *Campylobacter* (*C. Jejuni/C. Coli/C. Upsaliensis*) Detected test result.

Clostridioides (Clostridium) difficile (toxin A/B)

The BIOFIRE FILMARRAY GI Panel Mid contains a single multiplexed assay (Cdiff) for the identification of toxigenic *C*. *difficile* which targets both the toxin A gene (*tcdA*) and the toxin B gene (*tcdB*). Typical toxigenic strains produce both toxins, but the presence of either is indicative of a pathogenic strain. Empirical testing and *in silico* sequence analysis support that the assay will detect all toxinotypes and the epidemic BI/NAP1/027 hypervirulent strain, although these will not be specifically differentiated by the assay. Detection of either or both toxin genes by this assay gives a test result for *Clostridioides* (*Clostridium*) *difficile* (toxin A/B) Detected. As rates of asymptomatic carriage of *C. difficile* can be high in very young children and hospitalized patients, the detection of toxigenic *C. difficile* should be interpreted within the context of guidelines developed by the testing facility or other experts (e.g., guidelines/policy statements published by The American Academy of Pediatrics¹⁸ or the Society for Healthcare Epidemiology of America and the Infectious Disease Society of America).¹⁹

Salmonella

The BIOFIRE FILMARRAY GI Panel Mid contains a single assay (Salm) designed to detect both species of *Salmonella*; *S. enterica* and *S. bongori*. Empirical testing and *in silico* sequence analysis support detection of all subspecies and serovars of *Salmonella*. Cross-reactivity may occur with certain *E. coli* strains containing variants of the cryptic ETT2 type-III secretion system (see Analytical Reactivity (Inclusivity) for additional information).

Vibrio (V. parahaemolyticus/V. vulnificus/V. cholerae)

The BIOFIRE FILMARRAY GI Panel Mid contains two assays (Vibrio) for detection of *Vibrio* species most commonly implicated in gastroenteritis (*V. parahaemolyticus*, *V. vulnificus*, and *V. cholerae*). Empirical testing and in silico sequence analysis indicate that the assay may also react with some less common *Vibrio* species (i.e., *V. alginolyticus*, *V. fluvialis*, and *V. mimicus*) and with *Grimontia* (formerly *Vibrio*) *hollisae* (see Analytical Specificity (Cross-Reactivity and Exclusivity)). The Vibrio assay does not indicate which species has been detected and the Vibrio assay is not expected to detect the more rare species *V. cincinnatiensis*, *V. furnissii* and *V. metschnikovii*.

Yersinia enterocolitica

The BIOFIRE FILMARRAY GI Panel Mid contains a single assay (Yent) designed to detect all known serotypes/biotypes of *Y. enterocolitica*. Empirical testing and *in silico* sequence analysis indicate a potential for cross-reactivity with *Y. kristensenii* and *Y. frederiksenii* when present at high levels (>10⁸ CFU/mL). These two species are in the *Y. enterocolitica* group and are difficult to differentiate from *Y. enterocolitica* by culture methods; both are suspected human pathogens.

Shiga-like toxin-producing *E. coli* (STEC) Shiga-like toxin genes 1 and 2 (*stx1/stx2*)

The BIOFIRE FILMARRAY GI Panel Mid contains two assays (STEC 1 and STEC 2) for the detection of Shiga-like toxin 1 (*stx1*) and Shiga-like toxin 2 (*stx2*) sequences. The reported results do not indicate which of these toxin(s) have been detected. A positive result for either or both of these assays will give a Shiga-like toxin-producing *E. coli* (STEC) *stx1/stx2* Detected test result.

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Note: Shiga toxin (*stx*; identical to *stx1* of STEC) is found in *Shigella dysenteriae*; therefore, a BIOFIRE FILMARRAY GI Panel Mid report with positive test results for Shiga-like toxin-producing *E. coli* (STEC) *stx1/stx2* and *Shigella*/Enteroinvasive *E. coli* (EIEC) in the same sample may indicate the presence of *S. dysenteriae*.

Shigella/Enteroinvasive E. coli (EIEC)

The BIOFIRE FILMARRAY GI Panel Mid contains a single assay (Shig) for the detection of *ipaH*, a gene specifically found in all *Shigella* species as well as Enteroinvasive *E. coli* (EIEC). It is not possible to differentiate *Shigella* from EIEC using this method, and detection of *ipaH* will result in a *Shigella*/Enteroinvasive *E. coli* (EIEC) Detected test result.

Note: Shiga toxin (*stx*; identical to *stx1* of STEC) is found in *Shigella dysenteriae*, therefore a BIOFIRE FILMARRAY GI Panel Mid report with positive test results for Shiga-like toxin-producing *E. coli* (STEC) *stx1/stx2* with *Shigella*/Enteroinvasive *E. coli* (EIEC) in the same sample may indicate the presence of *S. dysenteriae*.

Parasites

Cryptosporidium

The BIOFIRE FILMARRAY GI Panel Mid contains two assays (Crypt 1 and Crypt 2) for detection of *Cryptosporidium* species. Empirical testing and/or *in silico* sequence analysis support detection of approximately 23 different *Cryptosporidium* species, including the most common species of human clinical relevance (i.e., *C. hominis* and *C. parvum*), as well as several less common species (e.g., *C. meleagridis*, *C. felis*, *C. canis*, *C. cuniculus*, *C. muris*, and *C. suis*). The assays do not differentiate between species and the very rare species *C. bovis*, *C. ryanae* and *C. xiaoi* may not be detected. A positive result for either or both assays will give a *Cryptosporidium* Detected test result.

Cyclospora cayetanensis

The BIOFIRE FILMARRAY GI Panel Mid contains a single assay (Ccayet) for the detection of *C. cayetanensis*, the only *Cyclospora* species implicated in human disease.

Giardia lamblia

The BIOFIRE FILMARRAY GI Panel Mid contains a single assay (Glam) designed to detect *G. lamblia* (aka *G. intestinalis*, *G. duodenalis*), the only *Giardia* species infectious to humans. A very low frequency of cross-reactivity with commensal microorganisms (i.e., *Bifidobacterium* and *Ruminococcus*) was observed in the clinical evaluation.

Viruses

Norovirus GI/GII

The BIOFIRE FILMARRAY GI Panel Mid contains two assays (Noro 1 and Noro 2) that together target the Norovirus genogroups most commonly associated with human infections (GI and GII). The reported results do not indicate which genogroup(s) (GI and/or GII) have been detected. A positive result for either or both assays will produce a test result of Norovirus GI/GII Detected. The assay(s) will also detect Norovirus GIX.1, which was formerly classified as GII.15.⁴⁸ Cross-reactivity of the Norovirus assay(s) with sequences linked to a variety of commensal microorganisms (e.g. *Mediterraneibacter gnavus, Prevotella sp.*) has been identified by investigation of specimens in clinical studies and via post-market monitoring (see Analytical Specificity (Cross-Reactivity and Exclusivity)).



BIOFIRE FILMARRAY GI Panel Mid Test Report

The BIOFIRE FILMARRAY GI Panel Mid test report is automatically displayed upon completion of a run and can be printed or saved as a PDF file. Each report contains a Run Summary, a Result Summary, and a Run Details Section.

	Array® Panel Mid			В	10 Ş f i r i				
				B	Y BIOMERIEU:				
					www.BioFireDx.com				
Run Summary	,								
Sample I				Run Date:	10 Sep 2024 3:05 PM				
Detected	I: Cyclospora cayetanensis			Controls:					
	Norovirus GI/GII								
Result Summa	ary								
		Bacteria							
Not Detected		C. jejuni/C. coli/C. upsalien							
Not Detected		Clostridioides (Clostridium) difficile (toxin A/B)							
Not Detected	Salmonella								
Not Detected		Shiga-like toxin-producing E. coli (STEC) stx1/stx2							
Not Detected	Shigella/Enteroinvasive Ē. coli (EIEC)								
Not Detected		Vibrio (V. parahaemolyticus/V. vulnificus/V. cholerae)							
Not Detected	d Yersinia enteroce								
		Parasites							
Not Detected									
 Detected 	Cyclospora caye	tanensis							
Not Detected	d Giardia lamblia								
		Viruses							
 Detected 	Norovirus GI/GII								
Change Sumn	nary								
Field	Changed To	Changed From	Operat	or	Date				
¹ Sample ID	New Sample ID	Cyclospora and Noro	Mandy	Mieu (mm)	10 Sep 2024				
Run Details									
Pouch	: GI Panel Mid v2.1		Protocol	: Stool FA	v3.4				
	Completed		Operator		Vieu (mm)				
Serial No			Instrument		FAULT Sim 1				
L at No	.: SIM000								

Figure 1. BIOFIRE FILMARRAY GI Panel Mid Example Test Report

Run Summary

The Run Summary section of the test report provides the Sample ID, time and date of the run, internal process control results and an overall summary of the test results. Any organism with a Detected result will be listed in the Detected field of the summary. If all of the organisms were Not Detected, then None will be displayed in the Detected field. Internal process controls are listed as Passed, Failed or Invalid. See the Controls Field section below for detailed information about the interpretation of internal process controls and appropriate follow-up in the case of internal process control failures.

Result Summary

The Result Summary section of the test report lists the result for each organism tested by the panel. Possible results for each organism are Detected, Not Detected, or Invalid. See Results Summary section below for detailed information about interpretation of test results and appropriate follow-up for Invalid results.

Run Details

The Run Details section provides additional information about the run including: pouch information (type, lot number, and serial number), Run Status (Completed, Incomplete, Aborted, Instrument Error, or Software Error), the protocol that was used to perform the test, the identity of the operator that performed the test, and the instrument used to perform the test.

Change Summary

Once a run has completed, it is possible to edit the Sample ID. If this information has been changed, an additional section called Change Summary will be added to the test report. This Change Summary section lists the field that was changed,

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the original entry, the revised entry, the operator that made the change, and the date that the change was made. Sample ID is the only field of the report that can be changed.

Controls Field

The Controls field on the test report will display Passed, Failed, or Invalid. The Controls field will display Passed only if the run completed successfully (no errors) and both of the pouch control assays (RNA Process Control and PCR2 Control) were successful. The Controls field will display Failed if the run was completed successfully (no errors) but one or both of the pouch control assays failed. If the control result is Failed, then the result for all of the tests on the panel are displayed as Invalid and the sample will need to be retested with a new pouch.

Table 3 provides a summary and explanation of the possible control results and follow-up actions.

Control Result	Explanation	Action Required	Outcome
Passed	The run was successfully completed AND Both pouch controls were successful.	None	Report the results provided on the test report.
Failed	The run was successfully completed BUT At least one of the pouch controls (RNA Process Control and/or PCR2 Control) failed.	Repeat the test using a new pouch.	Accept the results of the repeat testing. If the error persists, contact Customer Technical Support for further instruction.
Invalid	The controls are invalid because the run did not complete (Aborted, Incomplete, Instrument Error, or Software Error). (Typically this indicates a software or hardware error)	Note any error codes displayed during the run and the Run Status field in the Run Details section of the report. Refer to the appropriate BIOFIRE Operator's Manual or contact Technical Support for further instruction. Once the error is resolved, repeat the test or repeat the test using another module.	Accept the valid results of the repeat testing. If the error persists, contact Customer Technical Support for further instruction.

Table 3. Interpretation of Controls Field on the BIOFIRE FILMARRAY GI Panel Mid Test Report

Results Summary – Interpretations

The Results Summary – Interpretations section provides a complete list of the test results. Possible results for each organism include Detected, Not Detected, and Invalid. Table 4 provides an explanation for each interpretation and any follow-up necessary to obtain a final result.

Result	Explanation	Action
Detected	The run was successfully completed AND	None. Report results.
	The pouch controls were successful (Passed), AND	

Table 4. Reporting of Results and Required Actions



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Result	Explanation	Action
	The assay(s) associated with the interpretation were positive based on the following requirements for at least 2 of the 3 assay replicates:	
	-a positive melt curve, and	
	-the Tm for the melt data were within the assay specific limits, and	
	-the Tm for the melt data were within 1°C of each other.	
Not Detected	The run was successfully completed	None. Report results.
	AND	
	The pouch controls were successful (Passed)	
	AND	
	The assay(s) associated with the interpretation were negative (did not meet the requirements for a positive assay described in Detected).	
Invalid	The run did not complete successfully (Aborted, Incomplete, Instrument Error, or Software Error) OR	See Table 3, Interpretation of Controls Field on BIOFIRE Report, for instruction.
	The pouch controls were not successful (Failed)	

LIMITATIONS OF THE PROCEDURE

- 1. For prescription use only.
- 2. The BIOFIRE FILMARRAY GI Panel Mid is intended for use only on the BIOFIRE 2.0 and BIOFIRE TORCH Systems. Performance of the panel was established on the BIOFIRE FILMARRAY System (no longer manufactured or distributed), BIOFIRE 2.0 System, and BIOFIRE TORCH System. This test is a qualitative test and does not provide a quantitative value for the organism(s) in the sample.
- 3. The performance of this test has only been validated with human stool collected in Cary Blair transport medium, according to the media manufacturers' instructions. It has not been validated for use with other stool transport media, raw stool, rectal swabs, endoscopy stool aspirates, or vomitus.
- 4. This product should not be used to test stool samples in fixative (e.g., formalin or polyvinyl alcohol; PVA).
- 5. The performance of this product has not been established for the screening of stool for stool transplants.
- 6. The performance of this test has not been established for patients without signs and symptoms of gastrointestinal illness.
- 7. False positive and false negative results can be the result of a variety of sources and causes, it is important that these results be used in conjunction with other clinical, epidemiological, or laboratory information.
- 8. Virus, bacteria, and parasite nucleic acid may persist *in vivo* independently of organism viability. Additionally, some organisms may be carried asymptomatically. Detection of organism targets does not imply that the corresponding organisms are infectious or are the causative agents for clinical symptoms.
- 9. Results from this test must be correlated with the clinical history, epidemiological data, and other data available to the clinician evaluating the patient. Due to high rates of asymptomatic carriage of *C. difficile*, especially in very young children and hospitalized patients, the detection of toxigenic *C. difficile* should be interpreted within the context of guidelines developed by the testing facility or other experts (e.g., guidelines/policy statements published by The American Academy of Pediatrics or the Society for Healthcare Epidemiology of America and the Infectious Disease Society of America).^{18,19}
- 10. The performance of this test has not been established for monitoring treatment of infection with any of the panel organisms.
- 11. Discrepancies between the BIOFIRE FILMARRAY GI Panel Mid and other microbial identification methods may be caused by the inability to reliably differentiate species based on standard phenotypic microbial identification methods. Examples include differentiation of *Yersinia enterocolitica* from other *Y. enterocolitica* group members such as *Y. kristensenii* or *Y. frederiksenii* and differentiation of *Helicobacter pullorum* from *Campylobacter*. Also refer to the Organism Interpretation section of this document.
- 12. There is a risk of false negative results due to the presence of sequence variants in the gene targets of the assay, procedural errors, amplification inhibitors in specimens, or inadequate numbers of organisms for amplification.
- 13. The identification of several diarrheagenic *E. coli* pathotypes has historically relied upon phenotypic characteristics, such as adherence patterns or toxigenicity in certain tissue culture cell lines. The BIOFIRE FILMARRAY GI Panel Mid targets genetic determinants characteristic of most pathogenic strains of these organisms but may not detect all strains having phenotypic characteristics of a pathotype.
- 14. Target genes associated with the diarrheagenic *E. coli/Shigella* pathotypes are capable of horizontal transfer between strains, thus Detected results for multiple diarrheagenic *E. coli/Shigella* may be due to co-infection with multiple pathotypes or, less frequently, may be due to the presence of a single organism containing genes characteristic of multiple pathotypes.
- 15. Shigella dysenteriae possess a shiga toxin gene (stx) that is identical to the stx1 gene of STEC. The detection of both Shigella/Enteroinvasive E. coli (EIEC) and STEC stx1/stx2 analytes in the same specimen may indicate the presence of S. dysenteriae. Rare instances of the detection of shiga-like toxin genes in other genera/species have been reported; e.g., Aeromonas caviae, Acinetobacter haemolyticus, Shigella sonnei, Enterobacter cloacae, Citrobacter freundii, and Klebsiella pneumoniae.



- 16. This test only detects Campylobacter jejuni, C. coli and C. upsaliensis and does not differentiate between these three species of Campylobacter. Additional testing is required to differentiate between these species and to detect other Campylobacter species that may be present in stool specimens.
- 17. The LoD value for *Giardia intestinalis* was determined in units of cells/mL based on microscopic examination. However, parasites have a variable number of nuclei per cell in different life cycle stages (e.g. 1-2 nuclei per trophozoite and 4 (or more) nuclei per mature cyst). Because there is not a constant ratio of nucleic acid copy number per cell, detection by the BIOFIRE FILMARRAY GI Panel Mid may be variable when testing at an LoD measured in cells/mL.
- 18. The detection of organism nucleic acid is dependent upon proper sample collection, handling, transportation, storage, and preparation. Failure to observe proper procedures in any one of these steps can lead to incorrect results or no result. There is a risk of false positive and false negative results caused by improperly collected, transported, or handled specimens. The RNA process control and the PCR 2 control will not indicate whether or not nucleic acid has been lost due to inadequate collection, transport or storage of specimens.
- 19. A negative BIOFIRE FILMARRAY GI Panel Mid result does not exclude the possibility of gastrointestinal infection. Negative test results may occur from sequence variants in the region targeted by the assay, the presence of inhibitors, technical error, sample mix-up, or an infection caused by an organism not detected by the panel. Test results may also be affected by concurrent antimicrobial therapy or levels of organism in the sample that are below the limit of detection for the test. Negative results should not be used as the sole basis for diagnosis, treatment, or other management decisions.
- 20. Cary Blair transport medium may contain non-viable organisms and/or nucleic acid at levels that can be detected by the BIOFIRE FILMARRAY GI Panel Mid.
- 21. Due to the complex and highly variable nature of stool specimens, freezing may affect analyte integrity and subsequent test results for some specimens.
- 22. Organism, nucleic acid, and amplicon contamination may produce erroneous results for this test. Particular attention should be given to the Laboratory Precautions noted under the Warnings and Precautions section.
- 23. If four or more distinct organisms are detected in a specimen, retesting is recommended to confirm the polymicrobial result.
- 24. The effect of interfering substances has only been evaluated for those listed in the labeling. Interference by substances other than those described in the Interference section below could lead to erroneous results.
- 25. Several organisms were shown to have the potential to cross-react with BIOFIRE FILMARRAY GI Panel Mid assays. These include *Bifidobacterium* spp. and *Ruminococcus* spp. (*G. lamblia* assay); atypical sequences of unculturable/ uncharacterized *Prevotella* species, some sequences of *Anaerostipes hadrus*, *Enterobacter hormaechei* and *Parabacteroides* species, and *Mediterraneibacter* (*Ruminococcus*) gnavus (Noro 1 assay), *E. coli* containing a variant type III secretion protein (*Salmonella* assay), *Grimontia hollisae* which was formerly classified as a *Vibrio* sp. (*Vibrio* assay), *Yersinia frederiksenii* and *Yersinia kristensenii*, which are members of the Y. *enterocolitica* group (Y. *enterocolitica* assay). Please refer to the Organism Interpretation and Analytical Specificity (Cross-Reactivity and Exclusivity) sections of this document.
- 26. Cross-reactivity with organisms other than those listed above or in the Organism Interpretation or Analytical Specificity sections may lead to erroneous results.
- 27. Campylobacter inclusivity testing and in silico analyses demonstrated that the BIOFIRE FILMARRAY GI Panel Mid may have variable detection or reduced sensitivity for some organisms detected by the Campylobacter assays (Note: the Campylobacter assays only detect C. jejuni, C. coli, and C. upsaliensis). Campylobacter upsaliensis strain ATCC 43954 and Campylobacter jejuni subsp. doylei may not be detected and in silico analysis indicates primer mismatches that might lead to reduced assay sensitivity or lack of reactivity with 11/138 C. coli sequences evaluated from the NCBI database.
- 28. Empirical testing and *in silico* sequence analysis indicate that the *Vibrio* assay (*V. parahaemolyticus*/*V. vulnificus*/*V. cholerae*) may react with some less common *Vibrio* species (i.e., *V. alginolyticus*, *V. fluvialis*, and *V. mimicus*) but it is not expected to detect the rarer *Vibrio cincinnatiensis*, *Vibrio furnissii*, and *Vibrio metschnikovii* (Note: *Vibrio* spp. not associated with human disease were not evaluated).



- 29. Rare isolates of *V. harveyi*, *V. mimicus*, and *V. vulnificus* that have acquired a homolog of the *toxR* gene have been reported and may show cross-reactivity with the Vchol assay.
- 30. Based on the available sequences, a few *Cryptosporidium* species, or certain variants of species, including *C. bovis*, *C. ryanae*, and *C. xiaoi*, may not be efficiently detected by the Cryptosporidium assays. These species are rarely detected in human samples.
- 31. There is a risk of false positive values resulting from cross-contamination by target organisms, their nucleic acids or amplified product, or from non-specific signals in the assay.
- 32. There is a risk of false negative results due to the presence of strains with sequence variability or genetic rearrangements in the target regions of the assays. Refer to the inclusivity testing section of this document for additional information.
- 33. Unexpected results obtained from testing isolates from culture collections (e.g., during quality control testing) may occur due to mislabeling or miscategorization of the isolate, contamination of the stock, or genetic rearrangements (including loss of virulence plasmids) during repeated passaging.
- 34. Not all *Salmonella* serotypes were tested in validation studies; however, representatives of the 20 most prevalent serotypes recently circulating in the US (CDC National *Salmonella* Surveillance Annual Summary 2009) were evaluated. *In silico* sequence analysis supports detection of all subspecies and serotypes of *Salmonella*.
- 35. Cross-reactivity with the *Salmonella* assay may occur with certain *E. coli* strains containing variants of the cryptic ETT2 type-III secretion system (see Analytical Reactivity (Inclusivity) for additional information).
- 36. Positive and negative predictive values are highly dependent on prevalence. False negative results are more likely during peak activity when prevalence of disease is high. False positive results are more likely during periods when prevalence is moderate to low.
- 37. The performance of this test has not been evaluated for immunocompromised individuals.
- 38. State and local public health authorities have published guidelines for notification of reportable diseases in their jurisdictions including *Salmonella*, *Shigella*, and Shiga-like toxin-producing *E. coli* (STEC) *stx1/stx2* to determine necessary measures for verification of results to identify and trace outbreaks. Laboratories are responsible for following their state or local regulations for submission of clinical material or isolates on positive specimens to their state public health laboratories.

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EXPECTED VALUES

NOTE: The BIOFIRE FILMARRAY GI Panel Mid has an abbreviated panel menu compared to the original BIOFIRE FILMARRAY GI Panel, reporting only 11 of the original 22 analytes. The performance presented here was established during the original clinical and analytical evaluations for the BIOFIRE FILMARRAY GI Panel.

In the prospective clinical evaluation of the BIOFIRE FILMARRAY GI Panel Mid, 1556 eligible specimens (stool in Cary Blair transport medium) were collected and tested at four study sites across the United States (Pacific, North Central, Great Lakes, and Northeast regions) over approximately five months (May–September 2013). The number and percentage of positive results as determined by the BIOFIRE FILMARRAY GI Panel Mid, stratified by age group, are presented in the following table. Overall, the BIOFIRE FILMARRAY GI Panel Mid detected at least one organism in 30.2% (470/1556) of the prospective specimens.

BIOFIRE FILMARRAY GI Panel Mid Result	Overall (n=1556)	<1 year (n=121)	1-5 years (n=418)	6-12 years (n=193)	13-21 years (n=240)	22-64 years (n=411)	65+ years (n=173)			
Bacteria										
Campylobacter (C. jejuni/C. coli/C. upsaliensis)	58 (3.7%)	1 (0.8%)	11 (2.6%)	12 (6.2%)	6 (2.5%)	19 (4.6%)	9 (5.2%)			
Clostridioides (Clostridium) difficile toxin A/B	204 (13.1%)	49 (40.5%)	66 (15.8%)	18 (9.3%)	33 (13.8%)	29 (7.1%)	9 (5.2%)			
Salmonella	37 (2.4%)	5 (4.1%)	7 (1.7%)	5 (2.6%)	5 (2.1%)	11 (2.7%)	4 (2.3%)			
Shiga-like toxin-producing <i>E. coli</i> (STEC) stx1/stx2	38 (2.4%)	1 (0.8%)	24 (5.7%)	2 (1.0%)	4 (1.7%)	5 (1.2%)	2 (1.2%)			
Shigella/Enteroinvasive E. coli (EIEC)ª	49 (3.1%)	0 (0%)	31 (7.4%)	7 (3.6%)	5 (2.1%)	6 (1.5%)	0 (0%)			
Vibrio (V. parahaemolyticus/V. vulnificus/V. cholerae)	2 (0.1%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)	2 (0.5%)	0 (0%)			
Yersinia enterocolitica	1 (0.1%)	1 (0.8%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)			
		Parasites								
Cryptosporidium	24 (1.5%)	0 (0%)	9 (2.2%)	3 (1.6%)	6 (2.5%)	5 (1.2%)	1 (0.6%)			
Cyclospora cayetanensis ^b	19 (1.2%)	0 (0%)	0 (0%)	0 (0%)	0 (0%)	13 (3.2%)	6 (3.5%)			
Giardia lamblia	27 (1.7%)	1 (0.8%)	6 (1.4%)	5 (2.6%)	2 (0.8%)	13 (3.2%)	0 (0%)			
Viruses										
Norovirus GI/GII°	70 (4.5%)	15 (12.4%)	31 (7.4%)	5 (2.6%)	7 (2.9%)	9 (2.2%)	3 (1.7%)			

Table 5. Expected Values (as determined by the BIOFIRE FILMARRAY GI Panel Mid) Summary by Age Group for the
Prospective Clinical Evaluation (May through September 2013)

^a 10 of 49 Shigella/EIEC were detected at a study site in Providence, RI, in July, 2013, during a regional Shigella outbreak.

^b All 19 C. cayetanensis were detected at a study site in Omaha, NE, between June and July, 2013, during a multi-state Cyclospora outbreak.

° Refer to Table 7 below for additional Norovirus GI/GII Expected Values results.

In the prospective clinical evaluation, the BIOFIRE FILMARRAY GI Panel Mid reported a total of 53 specimens with multiple organism detections (i.e., mixed infections). This represents 11.3% (53/470) of positive specimens and 3.4% of all specimens tested (53/1556). The expected values for each BIOFIRE FILMARRAY GI Panel Mid organism result in mixed infections are presented in the following table.

Table 6. Expected Values for Analytes in Mixed Infections (as determined by the BIOFIRE FILMARRAY GI Panel Mid) in the Prospective Clinical Evaluation (May through September 2013)

Analyte	Number of Specimens Containing Analyte in Mixed Infections	Prevalence of Analyte in Mixed Infections (<i>N</i> = 53)								
Bacteria										
Campylobacter (C. jejuni/C. coli/C. upsaliensis)	14	26.4%								
Clostridioides (Clostridium) difficile toxin A/B	37	69.8%								
Salmonella	8	15.1%								
Shiga-like toxin-producing E. coli (STEC) stx1/stx2	11	20.8%								
Shigella/Enteroinvasive E. coli (EIEC)	8	15.1%								
Vibrio (V. parahaemolyticus/V. vulnificus/V. cholerae)	0	0%								
Yersinia enterocolitica	1	1.9%								
Parasites										
Cryptosporidium	4	7.5%								
Cyclospora cayetanensis	0	0%								
Giardia lamblia	7	13.2%								
Viruses										
Norovirus GI/GII	21	39.6%								

In 2023, bioMérieux conducted a new prospective clinical evaluation to assess norovirus performance using the most recent version of the U.S. Centers for Disease Control and Prevention (CDC) CaliciNet Norovirus GI/GII assay. This assay was updated with new probe designs and quantification cycle cutoff thresholds compared to the version that was used in the original clinical evaluation (2013, above).

The prospective clinical evaluation for norovirus detection on the BIOFIRE FILMARRAY GI Panel Mid included 872 eligible specimens (stool in Cary Blair transport medium) that were collected and tested at three study sites across the United States over approximately four months (April – July 2023). The number and percentage of positive norovirus results as determined by the BIOFIRE FILMARRAY GI Panel Mid, stratified by age group, are presented in the following table.

Table 7. Expected Values (EV) (as determined by the BIOFIRE FILMARRAY GI Panel Mid) by Age Group for the Prospective Clinical Evaluation (April through July 2023)

BIOFIRE FILMARRAY GI	Overall (n=872)			/ear 70)	,	/ears 120)	6-12 (n=	,		years 127)	22-64 (n=2	,	65+ y (n=2	/ears 208)
Panel Mid Result	#	EV	#	EV	#	EV	#	EV	#	EV	#	EV	#	EV
Norovirus GI/GII	63	7.2%	13	18.6%	16	13.3%	7	10.9%	11	8.7%	12	4.2%	4	1.9%

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CLINICAL PERFORMANCE CHARACTERISTICS

NOTE: The BIOFIRE FILMARRAY GI Panel Mid has an abbreviated panel menu compared to the original BIOFIRE FILMARRAY GI Panel, reporting only 11 of the original 22 analytes. The performance presented here was established during the original clinical and analytical evaluations for the BIOFIRE FILMARRAY GI Panel.

NOTE: BIOFIRE FILMARRAY GI Panel Mid performance was initially established on the first generation BIOFIRE FILMARRAY System (REF: FLM1-ASY-0001). The BIOFIRE FILMARRAY System is no longer being manufactured or distributed, but the performance characteristics established on that system are relevant to the BIOFIRE FILMARRAY GI Panel Mid and remain in this Instructions for Use. Comparison studies have established that BIOFIRE FILMARRAY GI Panel Mid performance characteristics are equivalent between BIOFIRE FILMARRAY System, BIOFIRE TORCH Systems.

Clinical Performance (2013)

The clinical performance of the BIOFIRE FILMARRAY GI Panel Mid was established during a multi-center study conducted at four geographically distinct U.S. study sites between May and September 2013. A total of 1578 prospective residual stool specimens in Cary Blair transport medium were acquired for the clinical study; 22 of these were excluded. The most common reasons for exclusion were that a valid external control was not completed on the day of testing, that the specimen was not plated to all of the appropriate bacterial culture media required for the reference method, or that the specimen was beyond four days from the date of collection. The final data set consisted of 1556 specimens. Table 8 provides a summary of demographic information for the 1556 specimens included in the prospective study.

Prospective Study Specimens			
Total Specimens	1556		
Sex	Number of Specimens (%)		
Male	718 (46%)		
Female	838 (54%)		
Age Group	Number of Specimens (%)		
<1 year	121 (8%)		
1-5 years	418 (27%)		
6-12 years	193 (12%)		
13-21 years	240 (15%)		
22-64 years	411 (26%)		
65+ years	173 (11%)		
Status	Number of Specimens (%)		
Outpatient	1350 (87%)		
Hospitalized	164 (11%)		
Emergency	42 (3%)		

Table 8. Demographic Summary for Prospective BIOFIRE FILMARRAY GI Panel Mid Clinical Evaluation	on

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The performance of the BIOFIRE FILMARRAY GI Panel Mid was evaluated by comparing the BIOFIRE FILMARRAY GI Panel Mid test result for each member of the panel with the appropriate comparator/reference methods shown in the table below.

BIOFIRE FILMARRAY System Test Results	Reference/Comparator Method			
Campylobacter (C. jejuni/ C. coli/C. upsaliensis)	Stool culture ^a (Blood agar, Blood agar with Ampicillin, MacConkey agar,			
Salmonella	Sorbitol-MacConkey agar, GN broth + Hektoen enteric agar,			
Vibrio (V. parahaemolyticus/V. vulnificus/V. cholerae)	Campylobacter agar, Cefsulodin-Irgasan™-Novobiocin agar, and Thiosulfate Citrate Bile Salts agar) with standard manual and automated microbiological/biochemical			
Yersinia enterocolitica	identification methods			
Clostridioides (Clostridium) difficile toxin A/B				
Shiga-like toxin-producing <i>E. coli</i> (STEC) <i>stx1/stx2</i>				
Shigella/Enteroinvasive E. coli (EIEC) ^b	PCR with Bi-directional Sequencing ^e			
Cryptosporidium				
Cyclospora cayetanensis	1			
Giardia lamblia ^c				
Norovirus GI/GII ^d				

Table 9. Comparator Methods for BIOFIRE FILMARRAY GI Panel Mid Clinical Evaluation

^a Any bacteria isolated from stool culture that could not be identified to the species level by laboratory methods were sequenced using an assay capable of providing species information (e.g., 16S).

^b Shigella may be identified by routine culture methods; however, culture detection will be reported for informational purposes only.

^c G. lamblia comparator assays consisted of one well-validated, sequenceable assay and one published assay that was not sequenceable.

^d CDC Calicinet assays (non-sequenceable) were used for the comparator method for Norovirus.

^e PCR assays were designed to amplify different sequences than those targeted by BIOFIRE FILMARRAY GI Panel Mid. Positive results for sequenceable assays required a sequence of adequate quality that matched a sequence of the expected organism/gene from the National Center for Biotechnology Information (NCBI) GenBank database (<u>www.ncbi.nlm.nih.gov</u>), with an acceptable E-value.

A total of 1556 specimens were evaluated in this study. Clinical sensitivity or positive percent agreement (PPA) was calculated as $100\% \times (TP / (TP + FN))$. True positive (TP) indicates that both the BIOFIRE FILMARRAY GI Panel Mid and reference/comparator method had a positive result for the specific analyte, and false negative (FN) indicates that the BIOFIRE FILMARRAY GI Panel Mid result was negative while the comparator result was positive. Specificity or negative percent agreement (NPA) was calculated as $100\% \times (TN / (TN + FP))$. True negative (TN) indicates that both the BIOFIRE FILMARRAY GI Panel Mid and the reference/comparator method had negative results, and a false positive (FP) indicates that the BIOFIRE FILMARRAY GI Panel Mid result was positive, but the comparator result was negative. The exact binomial two-sided 95% confidence interval was calculated. The results are summarized in Table 10.

Table 10. BIOFIRE FILMARRAY GI Panel Mid Performance in the Prospective Clinical Evaluation (May through September

	2013)					
	Sensitivity/PPA ^a			Specificity/NPA ^a		
BIOFIRE FILMARRAY GI Panel Mid Analyte	TP/(TP + FN)	%	95% CI	TN/(TN + FP)	%	95% CI
Bacteria						
Campylobacter (C. jejuni/C. coli/C. upsaliensis)	34/35 ^b	97.1	85.1-99.9%	1497/1521 ^b	98.4	97.7-99.0%
Clostridioides (Clostridium) difficile toxin A/B ^a	163/165°	98.8	95.7-99.9%	1350/1391°	97.1	96.0-97.9%
Salmonella	31/31	100	88.8-100%	1519/1525 ^d	99.6	99.1-99.9%
Shiga-like toxin-producing E. coli (STEC) stx1/stx2	33/33	100	89.4-100%	1518/1523°	99.7	99.2-99.9%
Shigella/Enteroinvasive E. coli (EIEC)	47/49	95.9	86.0-99.5%	1505/1507	99.9	99.5-100%

	Sensitivity/PPA ^a			Specificity/NPA ^a		
BIOFIRE FILMARRAY GI Panel Mid Analyte	TP/(TP + FN)	%	95% CI	TN/(TN + FP)	%	95% CI
Vibrio (V. parahaemolyticus/V. vulnificus/V. cholerae)	0/0	-	-	1554/1556 ^f	99.9	99.5-100%
Yersinia enterocolitica	1/1	100	-	1555/1555	100	99.8-100%
Parasites				•		
Cryptosporidium	18/18	100	81.5-100%	1532/1538 ^g	99.6	99.2-99.9%
Cyclospora cayetanensis	19/19	100	82.4-100%	1537/1537	100	99.8-100%
Giardia lamblia	20/20	100	83.2-100%	1529/1536 ^h	99.5	99.1-99.8%
Viruses	·					
Norovirus GI/GII	52/55 ^q	94.5	84.9-98.9%	1483/1501 ⁱ	98.8	98.1-99.3%

^a C. difficile performance is reported as positive percent agreement/negative percent agreement in contrast to the table headings. The performance measures of sensitivity and specificity only refer to those analytes for which the gold-standard bacterial culture was used as the reference method; *Campylobacter, Salmonella, Vibrio,* and *Yersinia enterocolitica.* Performance measures of positive percent agreement (PPA) and negative percent agreement (NPA) refer to all other analytes, for which PCR/sequencing assays were used as comparator methods.

^b Campylobacter jejuni subsp. doylei was identified in the single false negative specimen using bi-directional sequence analysis. Campylobacter was detected in 19/24 false positive specimens using bi-directional sequence analysis.

° C. difficile was detected in 1/2 false negative specimens and 41/41 false positive specimens using bi-directional sequence analysis.

^d Salmonella was detected in 6/6 false positive specimens using bi-directional sequence analysis.

^e STEC was detected in 5/5 false positive specimens using bi-directional sequence analysis.

^f Vibrio was detected in 2/2 false positive specimens using bi-directional sequence analysis.

⁹ Cryptosporidium was detected in 6/6 false positive specimens using bi-directional sequence analysis.

^h *G. lamblia* was detected in 4/7 false positive specimens using bi-directional sequence analysis. Two false positive results appear to be caused by cross-reactivity with *Bifidobacterium longum* and *Ruminococcus callidus*.

¹ The BIOFIRE FILMARRAY GI Panel Mid detected Norovirus in 1/3 false negative specimens when retested. Norovirus was detected in 1/2 remaining false negative specimens and 8/18 false positive specimens using bi-directional sequence analysis. Refer to "Clinical Performance (2023)" section below for additional Norovirus GI/GII performance results.

BIOFIRE FILMARRAY GI Panel Mid reports genus level (or multiple species group) results for three bacterial analytes; i.e., *Campylobacter (C. jejuni/C. coli/C. upsaliensis), Salmonella*, and *Vibrio (V. parahaemolyticus/V. vulnificus/V. cholerae)*. Standard laboratory methods identified various species/serovars within each of these groups during the clinical evaluation. Where standard methods did not provide a species identification, bi-directional sequencing was used to identify the species of the isolate. Stratification of performance by species/serovar is presented below. For *Vibrio*, no organisms were isolated by the culture methods; however, bi-directional sequencing from the original specimens identified one *V. parahaemolyticus* and one *V. cholerae*.

Table 11. Campylobacter Clinical Performance Stratified by Species

Campylobacter species ^a	Sensitivity
C. jejuni ^b	31/31 (100%)
C. coli	2/2 (100%)
C. jejuni subsp. doylei	0/1 (0%)
C. upsaliensis	1/1 (100%)
Overall Campylobacter	34/35 (97.1%) 95%Cl = 81.3-99.3%

^a Fifteen (15) Campylobacter were not speciated by the source laboratory and were subject to sequencing of the cadF gene. This method identified 11 *C. jejuni*, two *C. coli*, one *C. jejuni* subsp. doy/ei, and one *C. upsaliensis*.

^b Two *C. jejuni* were originally identified by the source lab as "*Campylobacter* species". Sequencing of the isolates provided by the laboratory identified them as *C. jejuni*. However, molecular testing of the specimen from which the isolates were obtained also detected the presence of *C. upsaliensis*, representing co-infection by these two species.

Table 12. Salmonella Clinical Performance Stratified by Species/Serovar

Salmonella species/serovar	Sensitivity
S. enterica ser. Enteritidis	7/7 (100%)
S. enterica ser. Typhimurium (i:-)	7/7 (100%)

Salmonella species/serovar

S. enterica ser. Typhimurium

S. enterica ser. Javiana

S. enterica ser. Newport

S. enterica ser. Agbeni

S. enterica ser. Berta

S. enterica ser. Ealing

S. enterica ser. Gaminara

S. enterica ser. Mbandaka

S. enterica ser. Muenchen

S. enterica ser. Thompson

Overall Salmonella

S. enterica ser. Paratyphi B var L-Tartrate

S. enterica ser. Infantis

S. enterica ser. Miami

95%01 = 88.8-100%
The BIOFIRE FILMARRAY GI Panel Mid reported multiple organism detections (i.e., mixed infections) for a total of 53
specimens. This represents 11.3% of positive specimens (53/471) and 3.4% of all specimens (53/1556). The majority of
multiple detections (49/53; 92.5%) contained two organisms, while 5.7% (3/53) contained three organisms, and 1.9% (1/53)
contained four organisms. The three organisms that were most prevalent in co-infections were C. difficile, Campylobacter,
and Norovirus). Out of the 53 specimens with multiple detections, 24 specimens (45.3%; 24/53) were concordant with the
reference methods. Twenty-nine (29) specimens (54.7%; 29/53) contained one or more organisms that had not been
detected by the reference/comparator methods (i.e., 32 false positive results); however, bi-directional sequence analysis
confirmed the presence of the analyte for 78.1% (25/32) of the discrepant results.

The most prevalent mixed infection was *C. difficile* with Norovirus (0.9% of all specimens; 14/1556) followed by *C. difficile* with STEC (0.5% of all specimens; 8/1556. Mixed infections were observed for all combinations of analyte classes (e.g., bacteria with viruses and/or parasites) and co-infections were observed within classes.

Multiple Detection Combination	Number of Specimens
C. difficile toxin A/B + Norovirus GI/GII	14
C. difficile toxin A/B + STEC stx1/stx2	8

Table 13. Most Prevalent Multi	nle Detection	Combinations	(>5 instances)	
	pie Detection	Compinations	(25 mstances)	1

The overall success rate for initial specimen tests in the prospective study was 99.2% (1544/1557). Four tests were incomplete due to software errors (3) or a user aborted run (1), and nine tests were invalid due to pouch control failures. All specimens but one were retested within four days of specimen collection and were successful after a single retest, for a final success rate of 99.9% (1556/1557).

Clinical Performance (2023)

The clinical performance of norovirus detection on the BIOFIRE FILMARRAY GI Panel Mid was assessed using the most recent version of the CDC CaliciNet Norovirus GI/GII assay with specimens from a multi-center study conducted at three geographically distinct U.S. study sites between April and July 2023. A total of 872 prospective residual stool specimens in

Sensitivity

3/3 (100%)

2/2 (100%)

2/2 (100%)

1/1 (100%)

1/1 (100%)

1/1 (100%)

1/1 (100%)

1/1 (100%)

1/1 (100%)

1/1 (100%)

1/1 (100%)

1/1 (100%)

1/1 (100%) **31/31 (100%)** 425089 424898

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Cary Blair transport medium were eligible for this evaluation; none of these were excluded. Table 14 provides a summary of the demographic information for the 872 specimens included in this prospective clinical evaluation.

Table 14. Demographic Summary for the BIOFIRE FILMARRAY GI Panel Mid Prospective Clinical Evaluation (April through
July 2023)

Stu	dy Specimens
Total Specimens	872
Sex	Number of Specimens (%)
Male	394 (45.2%)
Female	478 (54.8%)
Age Group	Number of Specimens (%)
< 1 year	70 (8.0%)
1-5 years	120 (13.8%)
6-12 years	64 (7.3%)
13-21 years	127 (14.6%)
22-64 years	283 (32.5%)
65+ years	208 (23.9%)
Status	Number of Specimens (%)
Outpatient	368 (42.2%)
Hospitalized	204 (23.4%)
Emergency	84 (9.6%)
Unknown	216 (24.8%)

The performance of the BIOFIRE FILMARRAY GI Panel Mid Norovirus GI/GII assay was evaluated by comparing the BIOFIRE FILMARRAY GI Panel Mid test result with an updated version of the comparator method used in the original (2013) prospective clinical evaluation: the U.S. Centers for Disease Control and Prevention (CDC) CaliciNet Norovirus GI/GII assay (updated in 2017).

A total of 872 specimens were evaluated. Positive percent agreement (PPA) was calculated as $100\% \times (TP / (TP + FN))$. True positive (TP) indicates that both the BIOFIRE FILMARRAY GI Panel Mid and the comparator method had positive results, and a false negative (FN) indicates that the BIOFIRE FILMARRAY GI Panel Mid result was negative while the comparator result was positive. Negative percent agreement (NPA) was calculated as $100\% \times (TN / (TN + FP))$. True negative (TN) indicates that both the BIOFIRE FILMARRAY GI Panel Mid and the comparator method had negative results, and a false positive (FP) indicates that the BIOFIRE FILMARRAY GI Panel Mid and the comparator method had negative results, and a false positive (FP) indicates that the BIOFIRE FILMARRAY GI Panel Mid result was positive, but the comparator result was negative. The exact binomial two-sided 95% confidence interval was calculated. The results are shown in Table 15.

Table 15. BIOFIRE FILMARRAY GI Panel Mid Norovirus GI/GII Performance in the Prospective Clinical Evaluation (April through July 2023)

BIOFIRE FILMARRAY GI	Positive Perce	ent Agr	eement (PPA)	Negative Percent Agreement (NPA)				
Panel Mid Result	TP/(TP + FN)	%	95% CI	TN/(TN + FP)	%	95% CI		
Norovirus GI/GII	34/35ª	97.1	85.1-99.9%	808/837ª	96.5	95.1-97.7%		

^a Norovirus was detected in the single FN specimen using bi-directional sequencing analysis. Norovirus was detected in 3/29 false positive specimens using bi-directional sequencing analysis. Twenty (20) of the remaining false positive results appear to have been caused by cross-reactivity; refer to the Analytical Specificity (Cross-Reactivity and Exclusivity) section for cross-reactive organisms.

The overall success rate for initial specimen tests in the prospective clinical evaluation was 99.0% (863/872). Nine specimens required repeat testing. Eight tests were incomplete due to instrument errors and one test was invalid due to a

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pouch control failure. All specimens were retested and were successful after a single retest (eight specimens) or two retests (one specimen), for a final success rate of 100% (872/872).

Clinical Comparison on the BIOFIRE 2.0

Clinical and non-clinical studies have established that the performance characteristics of the BIOFIRE FILMARRAY GI Panel Mid, including LoD (see Limit of Detection section below), positive percent agreement and negative percent agreement, and reproducibility are equivalent on the previously sold BIOFIRE FILMARRAY System and current BIOFIRE 2.0 systems. Non-clinical studies also demonstrate similar performance characteristics on BIOFIRE TORCH Systems.

NOTE: BIOFIRE TORCH Modules are BIOFIRE 2.0 modules that have been re-configured into a stacked system for higher throughput in a smaller workspace.

A clinical comparison study of the BIOFIRE FILMARRAY System and BIOFIRE 2.0 was performed using specimens previously obtained during the BIOFIRE FILMARRAY GI Panel Mid prospective clinical evaluation and supplemented with other archived specimens collected from external medical facilities and reference laboratories to increase the number of specimens being tested for low prevalence analytes. Contrived clinical specimens were also used for GI analytes which are extremely rare and for which no clinical specimens were available (*Vibrio* spp.). A total of 104 specimens were selected such that each analyte was represented 3-5 times. Each specimen was thawed or contrived and tested using the BIOFIRE FILMARRAY System and BIOFIRE 2.0 systems. Overall positive percent agreement (PPA) between systems was 98.3% with the lower bound of the two-sided 95% confidence interval (95% CI) at 90.8%. Overall negative percent agreement (NPA) was 99.4% with the lower bound of the two-sided 95% CI at 98.8%.

	BIOFIRE 2.0 / BIOFIRE FILMARRAY System						
Analyte	PPA	%	95% CI	NPA	%	95% CI	
Bacteria							
Campylobacter (C. jejuni/C. coli/C. upsaliensis)	5/5	100	47.8-100%	96/97	99.0	94.4-100%	
Clostridioides (Clostridium) difficile toxin A/B	5/5	100	47.8-100%	95/97	97.9	92.7-99.7%	
Salmonella	5/5	100	47.8-100%	97/97	100	96.3-100%	
Shiga-like toxin-producing E. coli (STEC) stx1/stx2	6/6	100	54.1-100%	96/96	100	96.2-100%	
Shigella/Enteroinvasive E. coli (EIEC)	6/6	100	54.1-100%	96/96	100	96.2-100%	
Vibrio (V. parahaemolyticus/V. vulnificus/V. cholerae)	6/7	85.7	42.1-99.6%	94/95	98.9	94.3-100%	
Yersinia enterocolitica	4/4	100	39.8-100%	98/98	100	96.3-100%	
Parasites							
Cryptosporidium	6/6	100	54.1-100%	96/96	100	96.2-100%	
Cyclospora cayetanensis	4/4	100	39.8-100%	98/98	100	96.3-100%	
Giardia lamblia	6/6	100	54.1-100%	96/96	100	96.2-100%	
Viruses							
Norovirus GI/GII	4/4	100	39.8-100%	96/98	98.0	92.8-99.8%	
Overall agreement	57/58	98.3	90.8-100%	1058/1064	99.4	98.8-99.8%	

Table 16. Analyte Results from BIOFIRE FILMARRAY System Clinical Comparison Study

System performance for testing of 104 specimens on each platform was calculated. For the BIOFIRE FILMARRAY System, a total of 105 runs were attempted, 104 of which were completed (99.0%; 104/105). One run was aborted by the user

(0.9%). There were no control failures. For the BIOFIRE 2.0, a total of 104 runs were attempted, all of which completed (100%; 104/104). There was one control failure.

Testing of Preselected Archived Specimens

Several analytes were either not encountered or had a low prevalence in the clinical study. To supplement the results of the prospective clinical study, an evaluation of 85 preselected archived specimens was performed. These specimens were archived clinical specimens that were selected because they had previously tested positive for one of the following analytes: *Vibrio, Y. enterocolitica, Cryptosporidium, G. lamblia,* or had been negative in previous laboratory testing. Prior to testing with the BIOFIRE FILMARRAY GI Panel Mid, the presence (or absence for negative specimens) of the expected analytes was verified in each specimen using analyte-specific PCR followed by bi-directional sequencing.

The specimens were organized into "test panels" and randomized such that the users performing the BIOFIRE FILMARRAY GI Panel Mid testing were blinded as to the expected test result. A summary of the available demographic information of the tested samples is provided in Table 17 and the results of the BIOFIRE FILMARRAY GI Panel Mid testing are presented in

Table 18.

Preselected Archived Specimens				
Total Specimens	85			
Sex	Number of Specimens (%)			
Male	14 (16.5%)			
Female	16 (18.8%)			
Unknown	55 (64.7%)			
Age Group	Number of Specimens (%)			
<1 year	2 (2.4%)			
1-5 years	8 (9.4%)			
6-12 years	11 (12.9%)			
13-21 years	4 (4.7%)			
22-64 years	5 (5.9%)			
65+ years	0 (0%)			
Unknown	55 (64.7%)			

Table 17. Demographic Summary for Preselected Archived Specimens

Table 18. BIOFIRE FILMARRAY GI Panel Mid Archived Specimen Performance Data Summary

Analista	Positive Perc	ent Agree	ment (PPA)	Negative Percent Agreement (NPA)			
Analyte	TP/(TP + FN)	%	95% CI	TN/(TN + FP)	%	95% CI	
Bacteria							
Vibrio (V. parahaemolyticus/ V. vulnificus/V. cholerae)	1/1	100	-	18/18	100	81.5-100%	
Yersinia enterocolitica	8/8	100	63.1-100%	9/9	100	66.4-100%	
Parasites							
Cryptosporidium	29/30	96.7	82.8-99.9%	36/36	100	90.3-100%	
Giardia lamblia	26/26	100	86.8-100%	36/36	100	90.3-100%	

Testing of Contrived Specimens

Some analytes are so rare that both prospective and archived testing efforts were insufficient to demonstrate system performance. To supplement the prospective and archived data, an evaluation of contrived specimens was performed. Surrogate specimens were prepared using residual specimens from the prospective clinical study that had previously tested negative for all BIOFIRE FILMARRAY GI Panel Mid analytes. Specimens were spiked at clinically relevant levels using five different quantified strains for each organism (or unspiked; 50 of each). The analyte status of each contrived specimen was blinded to the users analyzing the specimens. The results of the BIOFIRE FILMARRAY GI Panel Mid testing are presented in Table 19.

Table 19. BIOFIRE FILMARRAY GI Panel Mid Performance using Contrived Specimens
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Analyta	Positive Perce	ent Agree	ement (PPA)	Negative Percent Agreement (NPA)		
Analyte	TP/(TP + FN)	%	95% CI	TN/(TN + FP)	%	95% CI
Vibrio (V. parahaemolyticus/ V. vulnificus/V. cholerae)ª	112/115	97.4	92.6-99.5%	60/60	100	94.0-100%
Yersinia enterocolitica	65/65	100	94.5-100%	110/110	100	96.7-100%

^a Includes 64/65 *V. cholerae* (five different strains were used in spiking; one specimen spiked near the assay limit of detection was not detected) and 48/50 non-*V. cholerae* (four *V. parahaemolyticus* strains and one *V. vulnificus* strain were used in spiking; two specimens spiked with *V. parahaemolyticus* near the assay limit of detection were not detected).

ANALYTICAL PERFORMANCE CHARACTERISTICS

Limit of Detection

The Limit of Detection (LoD) for BIOFIRE FILMARRAY GI Panel Mid analytes was estimated with dilutions of single-spiked and multi-spiked samples (up to four organisms per sample). Detection was equivalent between single-spiked and multi-spiked samples, and LoD confirmation testing was performed by spiking one or more organism(s) into stool samples at the estimated LoD concentration and testing 20 replicates per sample. The LoD concentrations listed in Table 20 were confirmed on the BIOFIRE 2.0 and BIOFIRE TORCH Systems with analyte detection in at least 19/20 replicates (≥95%).

GI Panel Mid Test Result	Species/Isolate Tested	LoD Concentration				
BACTERIA						
Campylobacter	Campylobacter coli ATCC 33559	4 x 10 ⁴ cells/mL				
	Campylobacter jejuni ATCC BAA-1234					
	Campylobacter upsaliensis ATCC BAA-1059					
Clostridium difficile (toxin A/B)	Clostridium difficile Toxinotype 0 A+B+ ATCC 9689	4 x 10 ⁵ cells/mL				
	Clostridium difficile (NAP1) Toxinotype III A+B+ Zeptometrix #801619	4 x 10 ⁴ cells/mL				
Salmonella	Salmonella bongori O66:H1z41:H2- SGSC RKS#3041 SarC11	1 x 10 ⁴ CFU/mL				
	Salmonella enterica ssp. enterica Serovar Typhimurium O1,4,[5],12:H1i:H21,2 SGSC RKS#4194 SarC1	5 x 10 ³ CFU/mL				

Table 20. Limit of Detection (LoD) for BIOFIRE GI Panel Analytes

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GI Panel Mid Test Result	Species/Isolate Tested	LoD Concentration		
Vibrio (V. parahaemolyticus/V. vulnificus/V.	Vibrio cholerae Ogawa serotype O:1 ATCC 14035	8 x 10 ³ cells/mL		
cholerae)	Vibrio parahaemolyticus ATCC 17802	8 x 10 ⁴ cells/mL		
Yersinia enterocolitica	Yersinia enterocolitica Biovar1 serogroup O:8 ATCC 9610	5 x 10 ⁴ CFU/mL		
Shiga-like toxin-producing E. coli (STEC) stx1/stx2	<i>Escherichia coli</i> O26:H11 ATCC BAA-2196	1 x 10 ³ CFU/mL		
Obice//Subcesius 5 acti/SUSO	Escherichia <i>coli</i> O29:NM ATCC 43892	5 x 10 ³ CFU/mL		
Shigella/Enteroinvasive E. coli (EIEC)	Shigella sonnei ATCC 29930	100 CFU/mL		
	PARASITES			
Cryptosporidium ª	Cryptosporidium parvum Iowa isolate (Harley Moon) Waterborne, Inc. P102C Cryptosporidium hominis Clinical Specimen	5 x 10 ³ oocysts/mL ^{a,b}		
Cyclospora cayetanensis	Cyclospora cayetanensis Clinical Specimen	180 genome equivalents (GE)/mL		
Giardia lamblia	Giardia intestinalis (aka G. lamblia) ATCC 30957	50 cells/mL ^b		
VIRUSES				
Norovirus GI/GII	Norovirus GI Clinical Specimen Norovirus GII Clinical Specimen	− 1 x 10 ⁴ RNA copies/mL		

^a Limited testing with a clinical specimen containing *Cryptosporidium meleagridis* indicates that the LoD for *C. meleagridis* is similar to that of *C. parvum* and *C. hominis*. ^b Note that parasites will have variable numbers of nuclei per cell (up to 16) in different stages of reproduction and maturation (e.g. cyst versus trophozoite). Therefore, LoD established in units of cells/mL or oocysts/mL may not be reproducible, due to variable DNA concentration in each cell/cyst/trophozoite in the sample or culture tested.

Analytical Reactivity (Inclusivity)

The analytical reactivity (inclusivity) of the BIOFIRE FILMARRAY GI Panel Mid was evaluated with a collection of isolates that represent the diversity of the BIOFIRE FILMARRAY GI Panel Mid analytes. Isolates were selected to represent relevant subspecies or serotypes and selection was biased toward more common species and known human pathogens. When possible, *in silico* analysis of sequence data was used to make predictions of assay reactivity for less common species, strains, serovars or serotypes that were not tested but that may be detected by the BIOFIRE FILMARRAY GI Panel Mid.

Organisms were tested at concentrations near the limit of detection (LoD). If a sample containing a particular strain was positive (detected) at the initial test level, no further testing was required. If a strain was not detected, the strain was retested at the same level (up to five additional times) and if necessary, additional testing was performed at 10- and 100-fold higher concentrations to determine if the strain can be detected by the BIOFIRE FILMARRAY GI Panel Mid . Based upon predicted assay reactivity, a few select isolates were initially tested at a high concentration, followed by evaluation at lower concentrations if detection was observed. Results are provided below for each BIOFIRE FILMARRAY GI Panel Mid test result.

Organism	Isolate ID	Concentration Detected (cells/mL)	Multiple of LoD Detected
Campylobacter coli ^a	ATCC BAA-1061	1.2 x 10⁵	3×LoD

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Organism	Isolate ID	Concentration Detected (cells/mL)	Multiple of LoD Detected
	BEI HM-296	1.2 x 10 ⁵	3×LoD
	ATCC 43485	1.2 x 10⁵	3×LoD
	ATCC 43478	1.2 x 10⁵	3×LoD
	ATCC 33559 ^b	4.0 x 10 ⁴	1×LoD
	ATCC 49349	4.0 x 10 ⁶	Not Detected ^c
<i>Campylobacter jejuni</i> subsp. <i>doylei</i> ⁰	ATCC 49351	4.0 x 10 ⁶	100×LoD ^c
	ATCC 49350	4.0 x 10 ⁶	Not Detected ^c
	ATCC 43430	1.2 x 10⁵	3×LoD
Commutato a factoria in terrativati	ATCC BAA-1062	1.2 x 10⁵	3×LoD
Campylobacter jejuni subsp. jejuni	ATCC BAA-1234 ^b	4.0 x 10 ⁴	1×LoD
	BEI NR-128	1.2 x 10⁵	3×LoD
	ATCC BAA-1059	4.0 x 10 ⁴	1×LoD
Campylobacter upsaliensis	CCUG 24191	1.2 x 10⁵	3×LoD
	ATCC 43953	1.2 x 10⁵	3×LoD
	ATCC 43954 ^d	4.0 x 10 ⁶	Not Detected ^d
	ATCC 49815	1.2 x 10⁵	3×LoD
	BEI HM-297	1.2 x 10⁵	3×LoD

^a In silico analysis indicates primer mismatches that might lead to reduced assay sensitivity or lack of reactivity with 11/138 *C. coli* sequences.
 ^b Isolate was used to establish the LoD for this assay.
 ^c In silico analysis indicates primer mismatches that might lead to reduced assay sensitivity for this subspecies.
 ^d Sequencing under the primers identified an insertion/deletion in the primer binding region of the target gene.

Table 22. Clostridium difficile toxin A/B Inclusivity Results

Organism	Toxinotype	Isolate ID	Concentration Detected (cells/mL)	Multiple of LoD Detected
		ATCC 9689ª	4.0 x 10⁵	1xLoD
		ATCC BAA-1382	1.2 x 10 ⁶	3×LoD
		ATCC 17857	1.2 x 10 ⁶	3×LoD
		ATCC 17858	1.2 x 10 ⁶	3×LoD
		ATCC 43255	1.2 x 10 ⁶	3×LoD
	0 A+B+	ATCC 43594	1.2 x 10 ⁶	3×LoD
		ATCC 43596	1.2 x 10 ⁶	3×LoD
		ATCC 43599	1.2 x 10 ⁶	3×LoD
Clostridium difficile		ATCC 43600	1.2 x 10 ⁶	3×LoD
		ATCC 51695	1.2 x 10 ⁶	3×LoD
		ATCC 700792	1.2 x 10 ⁶	3×LoD
		ATCC BAA-1805 (NAP1)	1.2 x 10 ⁶	3×LoD
	III A+B+	Zeptometrix #0801619 (NAP1)ª	4.0 x 10 ⁴	1×LoD
	V A+B+	ATCC BAA-1875	1.2 x 10 ⁶	3×LoD
	VIII A-B+	ATCC 43598	1.2 x 10 ⁶	3×LoD
	X A-B+	CCUG 8864	1.2 x 10 ⁶	3×LoD

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Organism	Toxinotype	Isolate ID	Concentration Detected (cells/mL)	Multiple of LoD Detected
	XII A+B+	ATCC BAA-1812	1.2 x 10 ⁶	3×LoD
	XXII A+B (unknown)	ATCC BAA-1814	1.2 x 10 ⁶	3×LoD

^a This isolate was used to establish the LoD for this assay.

			-	
Organism (species, subspecies	and serovar)	Isolate ID	Concentration Detected (cells/mL)	Multiple of LoD Detected
		SGSC RKS 3041ª	1.0 x 10⁴	1xLoD
Salmonella bongori		NCTC 10946	3.0 x 10 ⁴	3×LoD
		SGSC RKS 3044	3.0 x 10⁴	3×LoD
Salmonella enterica su	bsp. <i>salamae II</i>	SGSC RKS 2985	1.5 x 10⁴	3×LoD
Salmonella enterica su	bsp. <i>arizonae IIIa</i>	SGSC RKS 2980	1.5 x 10⁴	3×LoD
Salmonella enterica su	bsp. <i>diarizonae IIIb</i>	SGSC RKS 2978	1.5 x 10⁴	3×LoD
Salmonella enterica su	bsp. <i>houtenae IV</i>	SGSC RKS 3027	1.5 x 10 ⁴	3×LoD
Salmonella enterica su	bsp. <i>indica VI</i>	SGSC RKS 2995	1.5 x 10⁴	3×LoD
	Typhimurium	SGSC RKS 4194ª	5.0 x 10 ³	1xLoD
	Enteritidis	ATCC BAA-708	1.5 x 10 ⁴	3×LoD
	Newport	ATCC 27869	1.5 x 10 ⁴	3×LoD
	Javiana	ATCC 10721	1.5 x 10 ⁴	3×LoD
	Heidelberg	ATCC 8326	1.5 x 10 ⁴	3×LoD
	Montevideo	ATCC BAA-710	1.5 x 10 ⁴	3×LoD
	l 4,[5],12:i:-	Cornell CU0580	1.5 x 10 ⁴	3×LoD
	Oranienburg	ATCC 9239	1.5 x 10 ⁴	3×LoD
	Saintpaul	ATCC 9712	1.5 x 10 ⁴	3×LoD
	Muenchen	ATCC 8388	1.5 x 10 ⁴	3×LoD
Salmonella enterica subsp. enterica	Braenderup	ATCC 700136	1.5 x 10 ⁴	3×LoD
Subop. entendu	Infantis	ATCC BAA-1675	1.5 x 10⁴	3×LoD
	Thompson	ATCC 8391	1.5 x 10 ⁴	3×LoD
	Mississippi	Cornell CU0633	1.5 x 10 ⁴	3×LoD
	Paratyphi B var. L(+) tartrate+ (formerly java)	CCUG 9561	1.5 x 104	3×LoD
	Typhi (Purified DNA)⁵	ATCC 700931D-5	1.5 x 10 ⁴	3×LoD
	Agona	ATCC 51957	1.5 x 10 ⁴	3×LoD
	Schwarzengrund	CCUG 21280	1.5 x 10 ⁴	3×LoD
	Bareilly	ATCC 9115	1.5 x 10 ⁴	3×LoD
	Hadar	ATCC 51956	1.5 x 10 ⁴	3×LoD

^a This isolate was used to establish the LoD for this assay. The organism was quantified in CFU/mL by plate enumeration. ^b Purified DNA was quantified in GE/mL by spectrophotometer (GE = genomic equivalents).



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Note: In addition to those evaluated in this study, *in silico* sequence analysis indicates the Salmonella assay should react with all species and subspecies of *Salmonella*, including all serovars of *S. enterica* subsp. *enterica*.

Organism (species, b	biotype and serotype)	Isolate ID	Concentration Detected (cells/mL)	Multiple of LoD Detected
	O:1 Ogawa	ATCC 14035 ^a	8.0 x 10 ³	1xLoD
	O:1 Inaba, Biotype El Tor	BEI NR-147	2.4 x 10 ⁴	3xLoD
Vibrio	O:1 Ogawa, Biotype El Tor	BEI NR-148	2.4 x 10 ⁴	3xLoD
cholerae	non-O:1,non-O:139 (O:2)	BEI NR-149	2.4 x 10 ⁴	3xLoD
non-O:1,non-O:139 (O:7)		BEI NR-152	2.4 x 10 ⁴	3xLoD
	O:1 Inaba, Biotype El Tor	ATCC 25870	2.4 x 10 ⁴	3xLoD
		ATCC 17802ª	8.0 x 10 ⁴	1xLoD
		ATCC BAA-242	2.4 x 10⁵	3xLoD
Vibrio para	haemolyticus	ATCC 27969	2.4 x 10⁵	3xLoD
		ATCC 33845	2.4 x 10⁵	3xLoD
		BEI NR-21990	2.4 x 10⁵	3xLoD
		BEI NR-21992	2.4 x 10⁵	3xLoD
Vibrio vulnificus		ATCC 29306	2.4 x 10⁵	3xLoD
		ATCC 33817	2.4 x 10⁵	3xLoD
		ATCC BAA-88	2.4 x 10⁵	3xLoD
		ATCC 27562	2.4 x 10 ⁴	0.3xLoD
		ATCC BAA-86	2.4 x 10 ⁴	0.3xLoD

Table 24. Vibrio (V. parahaemolyticus/V. vulnificus/V. cholerae) Inclusivity Results

^a Isolate was used to establish the LoD for this assay.

Organism	Serotype	Isolate ID	Concentration Detected (cells/mL)	Multiple of LoD Detected
	O:8	ATCC 9610ª	5.0 x 10 ⁴	1xLoD
Yersinia enterocolitica		ATCC 23715	1.5 x 10⁵	3xLoD
		BEI NR-207	1.5 x 10⁵	3xLoD
	O:5, 27	NCTC 10463	1.5 x 10⁵	3xLoD
	O:3	ATCC 700822	1.5 x 10⁵	3xLoD
		BEI NR-212	1.5 x 10⁵	3xLoD
	O:9	ATCC 55075	1.5 x 10⁵	3xLoD

Table 25. Yersinia enterocolitica Inclusivity Results

^a Isolate was used to establish the LoD for this assay. The organism was quantified in CFU/mL by plate enumeration.

Note: In addition to those evaluated in this study, *in silico* sequence analysis indicates the BIOFIRE FILMARRAY GI Panel Mid should react with all strains/serotypes of *Y. enterocolitica*.

	s. Shiga-like toxin			Concentration			
				Detected	Multiple of LoD		
Organism	Serotype	stx1/stx2	Isolate ID	(cells/mL)	Detected STEC		
	STEC (non-O157)						
	O26:H11	+/+	ATCC BAA-2196ª	1.0 x 10 ³	1×LoD		
	O113:H21	+/+	ATCC BAA-177	3.0 x 10 ³	3×LoD		
	O45:H2	Unknown	STEC Center DEC11C	3.0 x 10 ³	3×LoD		
	O103:H2	+/Unknown	STEC Center 107-226	3.0 x 10 ³	3×LoD		
	O104:H21	-/+	STEC Center G5506	3.0 x 10 ³	3×LoD		
	O111:NM	+/+	STEC Center 95-3208	3.0 x 10 ³	3×LoD		
	O111:H2	-/+	STEC Center RD8	3.0 x 10 ³	3×LoD		
	O111:H8	+/+	STEC Center DEC8B	3.0 x 10 ³	3×LoD		
Shiga-like toxin producing	O121:H19	Unknown	STEC Center F6173	3.0 x 10 ³	3×LoD		
E. coli (STEC)	O26:NM	+/-	STEC Center DA-22	3.0 x 10 ³	3×LoD		
	O26:H11	+/-	STEC Center H19	3.0 x 10 ³	3×LoD		
	O145:NM	+/-	STEC Center GS G5578620	3.0 x 10 ³	3×LoD		
	0104:H4 [♭] (Purified DNA) [°]	-/+	ATCC BAA-2326D-5⁵	3.0 x 10 ^{3c}	3×LoD		
			STEC O157				
	O157:NM	+/+	STEC Center DA-26	3.0 x 10 ³	3×LoD		
	O157:H7	-/+	STEC Center E32511	3.0 x 10 ³	3×LoD		
	O157:HNT	+/+	STEC Center DA-74	3.0 x 10 ³	3×LoD		
	O157:H7	+/+	ATCC 43895	1.0 x 10 ⁴	10xLoD		
	O157:H7	+/+	STEC Center A8993-CS2	3.0 x 10 ⁴	30×LoD		

Table 26. Shiga-like toxin producing E. coli (STEC) stx1/stx2 Inclusivity Results

^a Isolate was used to establish the LoD. The organism was quantified in CFU/mL by plate enumeration.

^b2011 European Outbreak Strain..

^c Purified DNA was quantified in GE/mL by spectrophotometer.

Note: Based on *in silico* analysis, *stx2* subtypes e and f are predicted to be detected with reduced sensitivity or not detected by the BIOFIRE FILMARRAY GI Panel Mid STEC assays.

Table 27. Shigella/Enteroinvasive E. coli (EIEC) Inclusivity Results

Organism	Serotype (Year/Location)	Isolate ID	Concentration Detected (cells/mL)	Multiple of LoD Detected
	O29:NM	ATCC 43892ª	5.0 x 10 ³	1×LoD
	O29:HNM (1977)	STEC Center 1885-77	3.0 x 10 ³	0.6×LoD
Enteroinvasive <i>E.</i> <i>coli</i> (EIEC)	O124:HNM (1978)	STEC Center 929-78	3.0 x 10 ³	0.6×LoD
	O29:H27 (1979; VA, USA)	STEC Center 1827-79	3.0 x 10 ³	Not Detected ^b
	O28:H- (1983, Brazil)	STEC Center LT-15	3.0 x 10 ³	0.6×LoD

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Organism	Serotype (Year/Location)	Isolate ID	Concentration Detected (cells/mL)	Multiple of LoD Detected
	O136:H- (1983, Bangladesh)	STEC Center LT-41 Strain 1111-55	3.0 x 10 ³	0.6×LoD
	Туре 2	ATCC 8700	3.0 x 10 ²	3×LoD
Chinalla havdii	Type 4	CDPH HUM-2010029296	3.0 x 10 ²	3×LoD
Shigella boydii (Serogroup C)	Type 1	ATCC 9207	3.0 x 10 ²	3×LoD
	Туре 20	ATCC BAA-1247	3.0 x 10 ²	3×LoD
	Туре 10	ATCC 12030	3.0 x 10 ²	3×LoD
	Type 1	BEI NR-520	3.0 x 10 ²	3×LoD°
Shigella	Type 2	CDPH PHM-2004008089	3.0 x 10 ²	3×LoD
dysenteriae	Type 13	ATCC 49555	3.0 x 10 ²	3×LoD
(Serogroup A)	Туре 3	ATCC 29028	3.0 x 10 ²	3×LoD
	Type 12	ATCC 49551	3.0 x 10 ²	3×LoD
	Type 2a	ATCC 700930	3.0 x 10 ²	3×LoD
	Type 1a	ATCC 9199	3.0 x 10 ²	3×LoD
Shigella flexneri	Туре 6	CDPH PHM-2006004043	3.0 x 10 ²	3×LoD
(Serogroup B)	Type 2b	ATCC 12022	3.0 x 10 ²	3×LoD
	Туре 2а	ATCC 29903	3.0 x 10 ²	3×LoD
	Unknown	STEC Center VA-6	3.0 x 10 ²	3×LoD
	N/A	ATCC 29930 ^a	1.0 x 10 ²	1×LoD
	N/A	ATCC 11060	3.0 x 10 ²	3×LoD
Shigella sonnei	N/A	CDPH HUM-2010027998	3.0 x 10 ²	3×LoD
(Serogroup D)	N/A	ATCC 29031	3.0 x 10 ²	3×LoD
	N/A	ATCC 25931	3.0 x 10 ²	3×LoD
	N/A	ATCC 9290	3.0 x 10 ²	3×LoD

^a Isolate was used to establish the LoD for this assay. The organism was quantified in CFU/mL by plate enumeration. ^b Secondary PCR assay could not confirm the presence of the target gene(s); plasmid/gene loss suspected.

° This isolate gave the expected STEC Detected and Shigella/EIEC Detected results due to the presence of stx in Shigella dysenteriae.

Organism	Location/Source of Isolate or Sample	Concentration Detected (copies/mL)	Multiple of LoD Detected	
	Scotland Clinical Sample ^ь	2.1 x 10 ^{3 b}	1×LoD	
Cryptosporidium hominis	Scotland Clinical Sample	6.4 x 10 ³	3×LoD	
	Scotland Clinical Sample	6.4 x 10 ³	3×LoD	
	BEI NR-2520 (Purified DNA Isolate TU502)	6.4 x 10 ³	3×LoD	
Cryptosporidium meleagridis	BEI NR-2521 (Purified DNA Isolate TU1867)	1.8 x 10 ³	3×LoD	
Cryptosporidium muris	Waterborne, Inc.P104	1.5×10 ⁴ oocycts/mL	3×LoD	

Table 28. Cryptosporidium Inclusivity Results^a

Organism	Location/Source of Isolate or Sample	Concentration Detected (copies/mL)	Multiple of LoD Detected
	Waterborne, Inc. P102C ^c	6.0 x 10 ^{2 c}	1×LoD
Cryptosporidium parvum	Scotland Clinical Sample	1.8 x 10 ³	3×LoD
	Scotland Clinical Sample	1.8 x 10 ³	3×LoD
	BEI NR-2519 (Purified DNA Isolate Iowa)	1.8 x 10 ³	3×LoD
Cryptosporidium ubiquitum	Scotland Purified DNA from Clinical Sample	Unknown	≤LoD ^d
	Scotland Purified DNA from Clinical Sample	Unknown	≤LoD ^d

^a Testing also included a clinical sample containing *Cryptosporidium canis* at a very low or non-amplifiable concentration (based on evaluation with a commercial quantitative real-time PCR (qPCR) assay). *Cryptosporidium* was not detected in the sample, but sequence analysis predicts that the species *C. canis* will be reliably detected when present in a sample at a concentration equal to or greater than the LoD of the Cryptosporidium assay(s) (i.e., no limitation on reactivity with *C. canis*).

with *C. canis*). ^b This *C. hominis* sample was used to establish the LoD for *C. hominis* (LoD of 5.0×10³ oocysts/mL was determined to be equivalent to 2.1×10³ copies/mL based on qPCR).

^c This *C. parvum* isolate was used to establish the LoD for *C. parvum* (LoD of 5.0×10³ oocysts/mL was determined to be equivalent to 6.0×10² copies/mL based on qPCR).

^d C. ubiquitum DNA purified from clinical samples was detected by the BIOFIRE FILMARRAY GI Panel Mid Cryptosporidium assay(s), though the DNA concentration tested was estimated (by qPCR) to be at or below the 95% LoD.

Note: In silico sequence analysis indicates the *Cryptosporidium* assay(s) should react with approximately 23 different *Cryptosporidium* species (including those evaluated in this study) as well as sequences not assigned to specific species (see also the Organism Interpretation section above). In silico analysis predicts that the *Cryptosporidium* assay(s) may not react with the rare or non-human species *C. bovis*, *C. ryanae* and *C. xiaoi*.

Table 29. Cyclospora cayetanensis Inclusivity Results

Organism		_ocation/Sample	Concentration Detected (GE/mL)	Multiple of LoD Detected
		Clinical Specimen ^a	180	1×LoD
	Nebraska	Clinical Specimen	540	3×LoD
		Clinical Specimen	540	3×LoD
Cyclospora cayetanensis		Clinical Specimen	540	3×LoD
		Clinical Specimen	540	3×LoD
	Peru	Clinical Specimen	540	3×LoD
		Clinical Specimen	540	3×LoD

^a Specimen was used to establish the LoD for this assay.

Table 30. Giardia lamblia Inclusivity Results

Organism	Location/Year of Isolation	Isolate ID	Concentration Detected (cells/mL)	Multiple of LoD Detected
	New Orleans, LA 1985	ATCC 50137	150	3×LoD
	Portland, OR 1971	ATCC 30888	150	3×LoD
<i>Giardia lamblia</i> (aka <i>G. intestinalis</i> or	Bethesda, MD 1979	ATCC 30957ª	50	1×LoD
G. duodenalis)	Unknown	Waterborne P101	150	3×LoD
	Egypt	ATCC PRA-243	150	3×LoD
	United States	ATCC PRA-247	150	3×LoD

^a Isolate was used to establish the LoD for this assay.

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Table 31. Norovirus GI/GII ^a Inclusivity Results				
S		Isolate ID	Concentration Dete	
notype		(Clinical Samples)	(copies/mL)	

Norovirus Genogroup/Genotype		Isolate ID (Clinical Samples)	Concentration Detected (copies/mL)	Multiple of LoD Detected
	3	Noro1_036 ^b	1.0 x 10 ⁴	1×LoD
	2	Noro1_002	6.0 x 10 ³	0.6×LoD
		Noro1_003	6.0 x 10 ³	0.6×LoD
	3	Noro1_012	6.0 x 10 ³	0.6×LoD
		Noro1_030	6.0 x 10 ³	0.6×LoD
Norovirus Gl	4	Noro1_031	6.0 x 10 ³	0.6×LoD
0.	6	Noro1_021	1.0 x 10⁵	10×LoD
		Noro1_009	2.0 x 10⁵	20×LoD ^d
	7	Noro1_029	6.0 x 10 ³	0.6×LoD
		Noro1_034	6.0 x 10 ³	0.6×LoD
	8	Noro G1.8 ^c	6.0 x 10 ⁴	6×LoD
	Unknown	Noro2_013 ^b	1.0 x 10 ^{4a}	1×LoD
	2	Noroll.2 ^b	6.0 x 10 ³	0.6×LoD
	3	China-5	6.0 x 10 ³	0.6×LoD
		SGB_038	6.0 x 10 ³	0.6×LoD
	4	GI-PILOT-SPDRL-077	2.0 x 10⁵	20×LoD ^d
		Noro2_004	2.0 x 10⁵	20×LoD ^d
		Noro2_032	2.0 x 10⁵	20×LoD ^d
Norovirus		PCMC_025 (Sydney)	6.0 x 10 ³	0.6×LoD
GII		PCMC_031 (Sydney)	6.0 x 10 ³	0.6×LoD
	6	NYH-A	6.0 x 10 ³	0.6×LoD
	7	Noroll.7°	6.0 x 10 ³	0.6×LoD
	8	Noroll.8°	6.0 x 10 ³	0.6×LoD
	12	Noroll.12 ^c	6.0 x 10 ³	0.6×LoD
	16	Noroll.16 ^c	6.0 x 10 ³	0.6×LoD
	20	Noroll.20c ^c	2.0 x 10 ⁵	20×LoD ^d
	20	Noroll.20°	6.0 x 10 ³	0.6×LoD

^a Sequence analysis indicates the Norovirus assays(s) will also detect Norovirus GIX.1 (formerly classified as genogroup GII.15).

^b Isolate was used to establish the LoD for this assay.

^c Isolate obtained as RNA extract from a clinical sample. Genotype provided by the source laboratory.

^d Noroviruses are genetically diverse. *In silico* analysis predicts that most strains of all genotypes will be detected, though some variant strains may be detected with reduced sensitivity or may not be detected due to inefficient amplification or exclusion by melt analysis.

Analytical Specificity (Cross-Reactivity and Exclusivity)

The potential for cross-reactivity between assays contained in the BIOFIRE FILMARRAY GI Panel Mid was evaluated by testing high concentrations of analyte. Both on-panel (identified by the GI Panel assays) and off-panel (not identified by the BIOFIRE FILMARRAY GI Panel Mid assays) organisms/viruses were tested.

On-panel organisms were tested to verify that they only react with the appropriate assay(s) on the panel. All on-panel organisms gave only the expected positive results; no false positive results were reported.

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Over 175 off-panel organisms were selected for specificity testing based on a combination of several factors including (1) relatedness to specific species detected by the BIOFIRE FILMARRAY GI Panel Mid (near-neighbors), (2) clinical relevance, (3) likelihood of being present in stool specimens, and (4) genetic similarity to BIOFIRE FILMARRAY GI Panel Mid assay primers, as determined by *in silico* analyses during assay design. When an organism of interest could not be obtained for testing, a separate organism-specific *in silico* analysis of whole genome sequence(s) directed against all BIOFIRE FILMARRAY GI Panel Mid primers was attempted for reactivity predictions. Several of the off-panel organisms were selected and tested to evaluate the specificity of particular assays, while many others were tested because they are commensal or pathogenic organisms with the potential to be found at high levels in stool. All organisms were tested at a high concentration (typically $\geq 1.0 \times 10^8$ CFU/mL for bacteria and fungi, $\geq 1.0 \times 10^4$ cells/mL for protozoa/parasites, and $\geq 1.0 \times 10^5$ units/mL for viruses).

Table 32 lists the organisms for which cross-reactivity was identified (either observed in testing or predicted by *in silico* analyses). With the exception of *Vibrio fluvialis* and *Vibrio mimicus* detection by the *Vibrio* assay, cross-reactivity has only been observed when the cross-reactive organism was known or suspected to be present in the sample at a high level.

Table 33 contains a complete list of the off-panel bacteria, fungi, protozoa/parasites, and viruses that were tested or for which *in silico* analysis was performed. Species with cross-reactivity observed in analytical testing are in bold font. Species without bold font received the expected BIOFIRE FILMARRAY GI Panel Mid test result (negative for all assays; no cross-reactivity) or for which *in silico* analysis does not predict cross-reactivity.

BIOFIRE FILMARRAY GI Panel Mid Test Result	Cross-Reactive Organism(s)
Giardia lamblia	Bifidobacterium spp °
Glaidia lambia	Ruminococcus spp ^a
Norovirus GI/GII [Noro 1 assay] ^b	Prevotella spp. (sequences from unculturable/uncharacterized species) ^c Mediterraneibacter (Ruminococcus) gnavus Parabacteroides spp. (P. merdae, P. acidifaciens ^d , P. distasonis ^e) Anaerostipes hadrus (select sequences) ^f Enterobacter hormaechei (select sequences) ^g
Salmonella	E. coli with variant type III secretion protein h
Vibrio (V. parahaemolyticus/V. vulnificus/ V. cholerae)	Vibrio alginolyticus Vibrio fluvialis ⁱ Vibrio mimicus ⁱ Grimontia (formerly Vibrio) hollisae
Yersinia enterocolitica	Yersinia frederiksenij ^{a.j} Yersinia kristensenii ^j

Table 32. Observed or Predicted Cross-Reactivity with Off-Panel Organisms

^a Cross-reactivity was not observed when tested at high concentration (1.5×10⁹ cells/mL). However, cross-reactivity was suspected or confirmed in clinical specimens and/or the potential for cross-reactivity is supported by *in silico* predictions.

^b Cross-reactivity was identified by post-market investigation of suspected false positive Norovirus GI/GII results in clinical specimens. Cross-reactivity with the species listed was confirmed by analytical testing at high concentration (>2.4x10⁸ cells/mL) and/or is supported by sequence analysis.

^c Cross-reactive sequences are inconsistent with other *Prevotella* sequence data, suggesting non-specific interaction with atypical or uncharacterized species and/or sequences.

^d P. acidifaciens was not tested but was determined by sequence analysis to have a similar risk of cross-reactivity as P. merdae.

^e Norovirus GI/GII Not Detected was reported when *P. distasonis* was tested at high concentration (3.1x10⁹ cells/mL). However, non-specific amplification products with Tm values close to the assay specific Tm range have been observed and the potential for false positive Norovirus GI/GII test results exists.

^f The risk of false positive Norovirus GI/GII results due to cross-reactivity with *A. hadrus* is associated with only a subset of *A. hadrus* RefSeq genome sequences (<35% as of June 2024).

⁹ The risk of false positive Norovirus GI/GII results due to cross-reactivity with *E. hormaechei* is associated with only a subset of *E. hormaechei* RefSeq genome sequences (<25% as of June 2024).

^h Cross-reactivity resulting in false positive *Salmonella* results has not been observed in analytical or clinical testing. However, non-specific amplification products with Tm values close to the assay specific Tm range have been observed and the potential for false positive *Salmonella* test results exists.

ⁱ Detected at concentrations near the Vibrio assay LoD.

¹ Y. kristensenii and Y. fredericksenii are difficult to distinguish from Y. enterocolitica by standard laboratory methods.



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 Table 33. Off-Panel Organisms Tested or Evaluated by in silico Analysis for BIOFIRE FILMARRAY GI Panel Mid Analytical Specificity

 Species with cross-reactivity observed in analytical testing are in bold font. On-panel species were also tested at high concentration (not shown).

		BACTERIA		
		Tested		
Abiotrophia defectivia	Campylobacter mucosalis	Enteropathogenic <i>E. coli</i> (EPEC)	Lactobacillus reuteri	Ruminococcus flavefaciens ^b
Acinetobacter baumannii	Campylobacter rectus	Enterotoxigenic <i>E. coli</i> (ETEC)	Lactococcus lactis	Selenomonas ruminantium
Acinetobacter Iwoffii	Campylobacter showae	Escherichia blattae	Leminorella grimontii	Serratia liquefaciens
eromonas hydrophila	Campylobacter sputorum	Escherichia fergusonii	Listeria monocytogenes	Serratia marcescens
keromonas nyuropinia	Campyiobacter spatoram	Eschenchia lergusolili	Mediterraneibacter	Serralia marcescens
Alcaligenes faecalis	Campylobacter ureolyticus	Escherichia hermannii	(Ruminococcus) gnavus	Shewanella algae
naerococcus tetradius	Cedecea davisae	Escherichia vulneris	Megamonas hypermegale	Staphylococcus aureus
Anaerostipes hadrus ^{a,c}	Chlamydia trachomatis	Edwardsiella tarda	Megasphaeara elsdenii	Staphylococcus epidermidis
rcobacter butzleri	Citrobacter amalonaticus	Egglerthella lenta	Methanobrevibacter smithii	Stenotrophomonas maltophilia
rcobacter cryaerophilus	Citrobacter freundii	Enterobacter cloacae	Morganella morganii	Streptococcus agalactiae
acillus cereus	Citrobacter koseri °	Enterobacter hormaechei ^{c,d}	Parabacteroides distasonis ^e	Streptococcus intermedius
acteroides fragilis	Citrobacter sedlakii	Enterococcus faecalis	Parabacteroides merdae ^f	Streptococcus pyogenes
Bacteroides thetaiotaomicron	Clostridium acetobutylicum	Enterococcus faecium	Peptoniphilus asaccharolyticus	Streptococcus salivarius
acteroides vulgatus	Clostridium botulinum	Eubacterium cylindroides	Peptostreptococcus anaerobius	Trabulsiella guamensis
Bifidobacterium adolescentis ^b	<i>Clostridium difficile</i> non-toxigenic [°]	Eubacterium rectale	Photobacterium damselae	Veillonella parvula
Bifidobacterium bifidum ^b	Clostridium histolyticum	Faecalibacterium prausnitzii	Plesiomonas shigelloides	Vibrio alginolyticus
Bifidobacterium longum ^b	Clostridium methylpentosum	Fusobacterium varium	Porphyromonas asaccharolytica	Vibrio fluvialis
Bifidobacterium pseudocatenulatum	Clostridium novyi	Gardnerella vaginalis	Prevotella bivia ^g	Vibrio mimicus
lautia (Ruminococcus) obeum	Clostridium perfringens	Gemella morbillorum	Prevotella copri ^g	Yersinia bercovieri
lautia wexlerae	Clostridium ramosum	Grimontia (Vibrio) hollisae	Prevotella intermedia 9	Yersinia frederiksenii ^h
ampylobacter concisus	Clostridium septicum	Haemophilus influénzae	Prevotella histicola ^g	Yersinia intermedia
Campylobacter curvus	Clostridium sordellii	Hafnia alvei	Prevotella melaninogenica ^g	Yersinia kristensenii
Campvlobacter fetus	Clostridium tetani	Helicobacter fennelliae	Proteus mirabilis	Yersinia mollaretii
Campylobacter gracilis	Collinsella aerofaciens	Helicobacter pylori	Proteus penneri	Yersinia pseudotuberculosis
Campylobacter helveticus	Corynebacterium genitalium	Klebsiella (Enterobacter) aerogenes	Proteus vulgaris	Yersinia rohdei
Campylobacter hominis	Desulfovibrio piger	Klebsiella oxytoca	Providencia alcalifaciens	
Campylobacter hyointestinalis	Diffusely adherent <i>E.coli</i>	Klebsiella pneumoniae	Pseudomonas aeruginosa	
Campylobacter Iari	Enteroaggregative <i>E. coli</i> (EAEC)	Lactobacillus acidophilus	Ruminococcus bromii ^b	
	PROTOZOA/F			FUNGI
Tes	ted	In silico Anal	ysis Only	Tested
abesia microti	Entamoeba histolytica	Ancylostoma duodenale	Entamoeba hartmanni	Aspergillus fumigatus
lastocystis hominis	Entamoeba moshkovskii	Ascaris lumbricoides	Entamoeba polecki	Candida albicans
onidiobolus lachnodes	Giardia muris	Balantidium coli	Enterobius vermicularis	Candida catenulate
onidiobolus lobatus	Pentatrichomonas hominis	Chilomastix mesnili	Enteromonas hominis	Penicillium marneffei
ncephalitozoon hellem	Schistosoma mansoni	Dientamoeba fragilis	Isospora belli	Saccharomyces boulardi
ncephalitozoon intestinalis	Toxoplasma gondii	Endolimax nana	Necator americanus	Saccharomyces cerevisiae
ntamoeba dispar	Trichomonas tenax	Entamoeba coli		•
Entamoeba gingivalis				
		VIRUSES		
	Test			In silico Analysis Only
denovirus A:31	Adenovirus F41	Coxsackievirus B3	Rhinovirus 1A	Adenovirus G52
Adenovirus B:34	Astrovirus type 8	Cytomegalovirus (CMV)	Rotavirus A	Norovirus GIV
Adenovirus C:2	Astrovirus variant VA1	Echovirus 6	Sapovirus	Rotavirus B
Adenovirus D:37	Astrovirus variant MLB	Enterovirus 68		Rotavirus C

Adenovirus E:4a Adenovirus F40	Bocavirus Type 1 Coronavirus 229E	Hepatitis A Herpes Simplex Type 2				
^a Anerostipes hadrus isolates (DSM sequence.	23942 and ATCC 29173) were tested at >2.4×10 ⁸	cells/mL. Norovirus GI/GII Detected results were only observed	rved with DSM 23942. The ATCC 29173 isolate does not carry the cross-reactive			
•	testing, cross-reactivity of the Giardia lamblia assa	y with one or more <i>Bifidobacterium</i> and <i>Ruminococcus</i> spec	cies was observed in the clinical evaluation (see Table 43).			
° Two isolates of this species were t	ested for analytical specificity.					
^d Enterobacter hormaechei isolates be cross-reactive by in silico analy		at >5.0×10 ⁸ cells/mL. Norovirus GI/GII Detected results were	e only observed with ATCC 49162. The ATCC BAA-2082 sequence is not predicted to			
e Though not observed in analytical testing, cross-reactivity of the Noro 1 assay with a sequence identified in roughly half (~50%) of the P. distasonis genomes evaluated could occur at high concentration.						
^f A similar risk of cross-reactivity wa	s identified with sequences annotated as Parabact	eroides sp. and P. acidifaciens.				
^g No cross-reactivity with high conce	entrations of various Prevotella species (commensa	al and pathogenic) was observed in analytical testing, but the	e potential for weak cross-reactivity between the Noro 1 assay and unique variant			

sequences annotated as unculturable Prevotella sp. has been identified via investigation of discrepant results in clinical specimens.

^{hi} Though not observed in analytical testing, in silico analysis indicates that, similar to *Y. kristensenii*, cross-reactivity between the Yersinia entercolitica assay and Yersinia fredericksenii is possible at high concentrations (see Table 43).

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Cross-Contamination and Carryover

The potential for sample-to-sample carryover was evaluated by alternately testing samples containing a high concentration or organism (10⁷ - 10⁹ organism/mL) with samples containing no organism. No false positive results were observed during testing of five sets of a high positive sample followed directly by a negative sample; demonstrating that the system design and recommended sample handling and testing practices are effective in preventing false positive results due to carryover or cross-contamination between samples.

Reproducibility

A multicenter reproducibility study was performed to determine between-site and overall reproducibility of the BIOFIRE FILMARRAY GI Panel Mid. Reproducibility testing occurred at three test sites using a panel of contrived stool samples, each spiked with various combinations of four different BIOFIRE FILMARRAY GI Panel Mid analytes. Each analyte was evaluated at three different concentrations (Negative, Low Positive and Moderate Positive).

The study incorporated a range of potential variation introduced by 13 different operators, 4 different pouch lots, and 16 different BIOFIRE modules. Samples were stored refrigerated (4°C) or frozen (\leq -70°C) prior to testing. Frozen samples were tested on five different days at three testing sites for 90 data points per sample and refrigerated samples were tested on four different days at three testing sites for 108 data points per sample. A summary of results (percent (%) agreement with the expected result) for each analyte (by site and overall) is provided in Table 34. The BIOFIRE FILMARRAY GI Panel Mid provided highly accurate and reproducible test results for all analytes (15,891/15,912 = 99.87% overall agreement with a 95% confidence interval of 99.81% - 99.92%).

				% Agreemen	t with Expecte	ed Result ^a
Organism Tested	Concentration Tested	Expected Result	Site A	Site B	Site C	All Sites (95% Confidence Interval)
	Moderate Positive 3xLoD 1.2x10 ⁵ cells/mL	Detected	30/30 100%	30/30 100%	30/30 100%	90/90 100% (96.0 - 100%)
Campylobacter jejuni ATCC BAA-1234	Low Positive 1xLoD 4x10 ⁴ cells/mL	Detected	30/30 100%	30/30 100%	30/30 100%	90/90 100% (96.0 - 100%)
	None	Not Detected	192/192 100%	192/192 100%	192/192 100%	576/576 100% (99.4 - 100%)
	Moderate Positive 3xLoD 1.2x10 ⁶ cells/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6 - 100%)
Clostridium difficile ª ATCC 9689	Low Positive 1xLoD 4x10 ⁵ cells/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6 - 100%)
	None	Not Detected	120/120 100%	120/120 100%	120/120 100%	360/360 100% (96.6 - 100%)
	Moderate Positive 3xLoD 1.5x10 ⁴ CFU/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6 - 100%)
Salmonella enterica ª SarC1 (SGSC)	Low Positive 1xLoD 5x10 ³ CFU/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6 - 100%)
	None	Not Detected	120/120 100%	120/120 100%	120/120 100%	360/360 100% (96.6 - 100%)

Table 34. Reproducibility of the BIOFIRE FILMARRAY GI Panel Mid Test Results on the BIOFIRE FILMARRAY System



	-			% Agreement	t with Expecte	ed Resultª
Organism Tested	Concentration Tested	Expected Result	Site A	Site B	Site C	All Sites (95% Confidence Interval)
	Moderate Positive 3xLoD 3x10 ² CFU/mL	Detected	30/30 100%	30/30 100%	30/30 100%	90/90 100% (96.0 - 100%)
Shigella sonnei ATCC 29930	Low Positive 1xLoD 1x10 ² CFU/mL	Detected	30/30 100%	30/30 100%	30/30 100%	90/90 100% (96.0 - 100%)
	None	Not Detected	192/192 100%	192/192 100%	192/192 100%	576/576 100% (99.4 - 100%)
	Moderate Positive 3xLoD 2.4x10 ⁵ cells/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6 - 100%)
Vibrio parahaemolyticusª ATCC 17802	Low Positive 1xLoD 8x10 ⁴ cells/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6 - 100%)
	None	Not Detected	120/120 100%	120/120 100%	120/120 100%	360/360 100% (96.6 - 100%)
	Moderate Positive 3xLoD 1.5x10 ⁴ oocysts/mL	Detected	30/30 100%	30/30 100%	30/30 100%	90/90 100% (96.0 - 100%)
Cryptosporidium parvum Waterborne, Inc. P102C	Low Positive 1xLoD 5x10 ³ oocysts/mL	Detected	30/30 100%	30/30 100%	30/30 100%	90/90 100% (96.0 - 100%)
1 1020	None	Not Detected	192/192 100%	192/192 100%	192/192 100%	576/576 100% (99.4 - 100%)
Olevalia	Moderate Positive 3x LoD 150 cells/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6 - 100%)
Giardia intestinalisª (syn. Giardia lamblia) ATCC 30957	Low Positive 1xLoD 50 cells/mL	Detected	30/36 83.3%	30/36 83.3%	31/36 86.1%	91/108 84.3% (77.0 - 91.0%)
A100 30837	None	Not Detected	120/120 100%	120/120 100%	120/120 100%	360/360 100% (96.6 - 100%)
	Moderate Positive 3xLoD 3x10 ⁴ copies/mL	Detected	29/30 96.7%	30/30 100%	30/30 100%	89/90 98.9% (96.0 - 100%)
Norovirus GI Clinical Specimen	Low Positive 1xLoD 1x10 ⁴ copies/mL	Detected	28/30 93.3%	29/30 96.7%	30/30 100%	87/90 96.7% (96.0 - 100%)
	None	Not Detected	192/192 100%	192/192 100%	192/192 100%	576/576 100% (99.4 - 100%)

^a Reproducible, but suboptimal (<95%) detection was observed at one or both concentrations in frozen contrived samples. Data presented are from samples stored at ~4°C for up to 4 days prior to testing.

Similar reproducibility studies were also performed to determine between-site/system and overall reproducibility of the BIOFIRE FILMARRAY GI Panel Mid on BIOFIRE 2.0 and BIOFIRE TORCH Systems. For the BIOFIRE 2.0 study, testing occurred at three test sites using contrived stool samples, each spiked with various combinations of three different BIOFIRE FILMARRAY GI Panel Mid analytes representing the types of organisms detected by the panel (bacteri, parasites, DNA viruses, and RNA viruses). Each analyte was evaluated at three different concentrations (Negative, Low Positive and Moderate Positive). Negative results for each assay were obtained from samples that were not spiked with a corresponding organism (analyte not in the sample).

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The data include 108 replicates per analyte and incorporate a range of potential variation introduced by 9 different operators, 3 different pouch lots, and 9 different BIOFIRE 2.0 modules configured on 3 different multi-instrument systems. Similar to the reproducibility of the BIOFIRE FILMARRAY GI Panel Mid on the BIOFIRE FILMARRAY System (Table 34 above), percent (%) agreement with the expected Detected, Not Detected or N/A result was 99.1% or better (Table 35).

The reproducibility of the BIOFIRE FILMARRAY GI Panel Mid results on the BIOFIRE TORCH System showed agreement with the expected Detected results in 97.0% of replicates for various representative analytes at 1×LoD and agreement with the expected Not Detected or N/A results was \geq 99.0% (not shown).

Note: Giardia lamblia was tested in the BIOFIRE FILMARRAY System and BIOFIRE TORCH Reproducibility studies and was not detected in ≥95% of replicates tested at the 1× LoD concentration in either study (84.3% on BIOFIRE FILMARRAY System and 91.7% on BIOFIRE TORCH). It was subsequently determined that the integrity of the stock culture used to prepare the contrived samples had been compromised (requires storage in liquid nitrogen). Retesting of 20 replicates of a sample prepared from a new stock culture resulted in the expected ≥95% detection at the 1× LoD concentration when tested on the on the BIOFIRE FILMARRAY System, BIOFIRE 2.0, and BIOFIRE TORCH Systems.

	Concentration Expected Tested Result	Function	1	% Agreement with Expected Result			
Organism Tested		Result	Site/System			Total	
	resteu	Result	Α	В	С	(95% Confidence Interval)	
	Moderate Positive 3× LoD 1.2x10 ⁶ CFU/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6-100%)	
Clostridium difficile (toxinotype 0 A+B+) ATCC 9689	Low Positive 1× LoD 4.0x10 ⁵ CFU/mL	Detected	35/36 97.2%	36/36 100%	36/36 100%	107/108 99.1% (95.0-100%)	
	Negative	Not Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6-100%)	
Shine toyin producing	Moderate Positive 3× LoD 3.0x10 ⁴ CFU/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6-100%)	
Shiga-toxin producing Escherichia coli O157 (STEC O157) ATCC 43895	Low Positive 1× LoD 1.0x10 ⁴ CFU/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6-100%)	
A100 43033	Negative	Not Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6-100%)	
	Moderate Positive 3× LoD 1.5x10 ⁴ oocysts/mL	Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6-100%)	
Cryptosporidium parvum Waterborne P102C	Low Positive 1× LoD 5.0x10 ³ oocysts/mL	Detected	35/36 97.2%	36/36 100%	36/36 100%	107/108 99.1% (95.0-100%)	
	Negative	Not Detected	36/36 100%	36/36 100%	36/36 100%	108/108 100% (96.6-100%)	

Table 35. Reproducibility of the BIOFIRE FILMARRAY GI Panel Mid Test Results on the BIOFIRE 2.0 System

Interference

Substances that could be present in stool samples (preserved in Cary Blair medium) or introduced during sample handling were evaluated for their potential to interfere with assay performance. A potentially interfering substance was added to a contrived stool sample containing representative BIOFIRE FILMARRAY GI Panel Mid organisms. Each contrived sample contained a mix of four different organisms, each present at approximately three times (3×) the limit of detection (LoD). Unspiked samples (no test substance) served as positive controls (no interference) for comparison. Spiked samples

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(containing the test substance) were reviewed for performance of controls and accuracy of the test results for each sample. Reproducible control failures or unexpected test results (false positive or false negative) were a sign of interference.

No inhibition or unexpected test result were obtained in the presence of the endogenous and exogenous substances tested (Table 36).

	-	
Endogenous Substances	Exogenous Substances	S
Human Whole Blood	Bacitracin	Glycerin
Triglycerides	Doxycycline	Hydrocortisone
Cholesterol	Nystatin	Loperamide hydrochloride
Fatty acids (palmitic acid)	Metronidazole	Magnesium hydroxide
Fatty acids (stearic acid)	Naproxen sodium	Mineral oil
Bovine Mucin ^a	Bisacodyl	Phenylephrine hydrochloride
Human Bile	Bismuth subsalicylate	Sodium phosphate
Human Urine	Calcium carbonate	Nonoxynol-9
Human stool (overfill of Cary Blair vial)	Docusate sodium	Bleach
		Ethanol

 Table 36. Endogenous and Exogenous Substances Tested – No Interference

^a Unexpected EPEC results reported due to contamination of the mucin with EPEC nucleic acid.

No inhibition or unexpected test results were obtained in the presence of high concentrations of potentially competing microorganisms (Table 37).

Table 37. Potentially Competing Microorganisms Tested – No Interference

Off-Panel Organisms	
Adenovirus F41	Enterotoxigenic <i>E. coli</i> (ETEC)
Aeromonas hydrophila	Non-pathogenic <i>E. coli</i>
Bacteroides vulgatus	Helicobacter pylori
Bifidobacterium bifidum	Saccharomyces boulardii
Human Rhinovirus 87	

Contrived stool samples prepared in various transport media, including Cary Blair (see Table 38), were evaluated for the potential of different media to interfere with the accuracy of BIOFIRE FILMARRAY GI Panel Mid test results. No interference was observed for samples collected in Protocol Cary Blair or other brands of enteric transport media (Para-Pak Enteric Plus and Para-Pak C&S media); performance has not been established in these media. However, accurate detection of analytes was impaired (false negative results) for samples prepared in media containing fixatives, particularly those containing formalin.

Table 38. Transport Media Tested

Enteric Transport Media – No Interference Observed				
PROTOCOL™ Cary Blair	Para-Pak Enteric Plus ^a	Para-Pak C&S ^a		
Fixative-containing Transport Media - Interference Observed ^a				
Modified (Cu) PVA Fixative	Para-Pak 10% Formalin Fixative⁵	Para-Pak SAF Fixative ^a		
Para-Pak ECOFIX Fixative	Para-Pak LV-PVA Fixative	Para-Pak Zn-PVA Fixative		

^a Performance has not been established in these media.

^b Impaired detection of analytes (false negative results) in formalin containing media.

APPENDIX A

Symbols Glossary

ISO 15223-1 Medical devices - Symbols to be used with medical devices labels, labeling and information to be supplied					
5.1.1	Manufacturer	5.1.4	Use-By date (YYYY-MM-DD	5.1.5 LOT	Batch Code (Lot Number)
5.1.6 REF	Catalog Number	5.1.7 SN	Serial Number	5.2.8	Do Not Use if Package Is Damaged
5.3.2	Keep Away from Sunlight	5.3.7	Temperature Limit	5.4.2	Do Not Reuse
5.4.3	Consult Instructions for Use	5.5.1	<i>In vitro</i> Diagnostic Medical Device	5.5.5 S	Contains Sufficient For <n> Tests</n>
5.7.10 UDI	Unique Device Identifier				
	Use of Symbol	s in Labeling – 81 FR 3	8911, Docket No. (FD	A-2013-N-0125)	
Rx Only			Prescription Use Only		
United Nation	ons Globally Harmoniz	ed System of Classific	ation and Labeling of	chemicals (GHS) (ST/	SG/AC.10/30)
THE STREET	Serious eye damage, Category 1	$\langle \mathbf{\hat{l}} \rangle$	Acute toxicity, oral, Category 4 & Skin corrosion, irritation, Category 2	×	Acute aquatic hazard, Category 1 & Long- term aquatic hazard, Category 1
	Ма	nufacture Symbols (BI	OFIRE Diagnostics, L	LC)	
G	BIOFIRE FILMAR	RAY GI Panel Mid	Ş	A product in the BIO	FIRE GI Panel family

REF

BioFire Diagnostics, LLC 515 Colorow Drive Salt Lake City, UT 84108

USA

APPENDIX B

Contact and Legal Information

Customer and T	echnical Support
Reach Us on the Web http://www.BioFireDX.com	Reach Us by Phone 1-800-682-2666 – Toll Free
Reach Us by Email BioFireSupport@biomerieux.com	Reach Us by Fax
Reach Us by Mail 515 Colorow Drive Salt Lake City, UT 84108 USA	(801) 588-0507

Note: A paper copy of this Instructions for Use is available upon request by contacting Customer Support.

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Warranty Information

Product warranty information is available online at:

http://www.biofiredx.com/support/documents/

For warranty information for customers outside the United States, contact the local bioMérieux sales representative or an authorized distributor.

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REVISION HISTORY

Version	Revision Date	Description of Revision(s)
01	January 2025	Initial Release

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