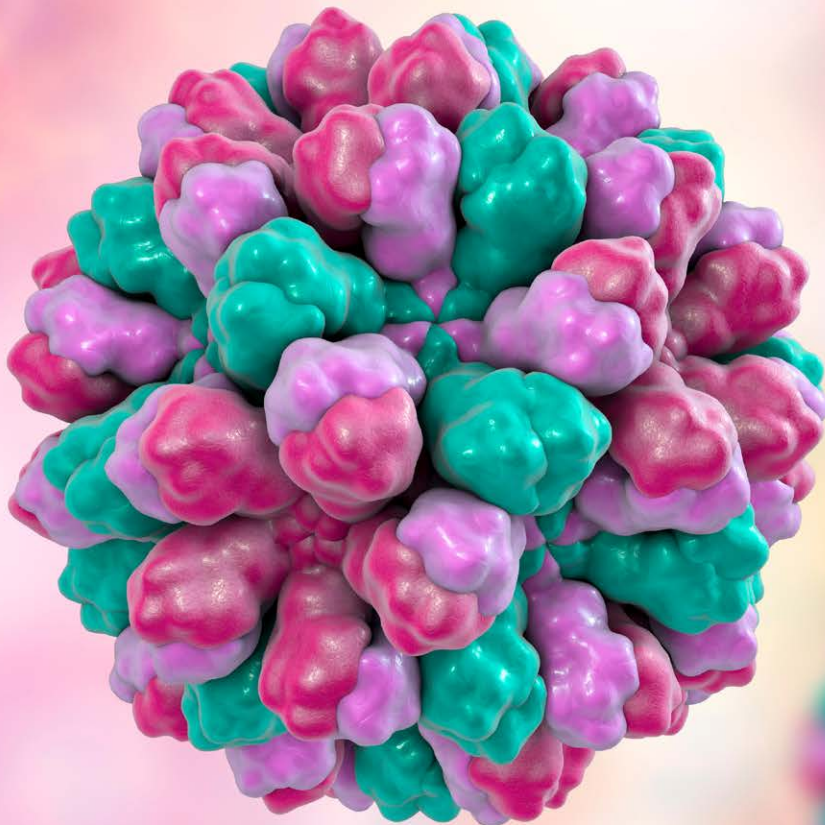


Faecal Multiplex PCR

*For accurate and timely diagnosis
of gastroenteritis*



Gastroenteritis is a common presentation in both adults and children. Most acute cases are due to infection, with chronic cases more likely to be due to non-infectious causes such as inflammatory bowel disease or malabsorption syndrome. When infectious diarrhoea is suspected, two decisions need to be made: firstly, when to perform stool testing, and secondly, whether antibiotic therapy is required.

Most infectious diarrhoea is mild and self-limiting. In such instances, supportive therapy, such as rehydration, is sufficient, and microbiological testing is not required¹. However, in patients with severe illness and/or high-risk comorbidities, a diagnosis will help guide further management.

Who to test

Patients with severe illness

- Dehydration/hypovolaemia
- Hospitalisation
- Fever > 38°C
- Bloody diarrhoea/dysentery

Patients with co-morbidities

- Age > 70
- Malignancy
- Immunosuppressed
- Inflammatory bowel disease
- Pregnancy

Additional reasons to test

- Prolonged symptoms > 1 week
- Recent antibiotic exposures (*C.difficile* only)
- If directed by public health/outbreak investigations

Causes of infectious gastroenteritis and respective testing

Infectious diarrhoea can broadly be categorised according to its aetiology: bacterial, viral or parasitic. Viral causes are the most common, while bacterial causes are more likely to cause severe illness². Identifying the underlying aetiology assists with ongoing management (see Table 1 below).

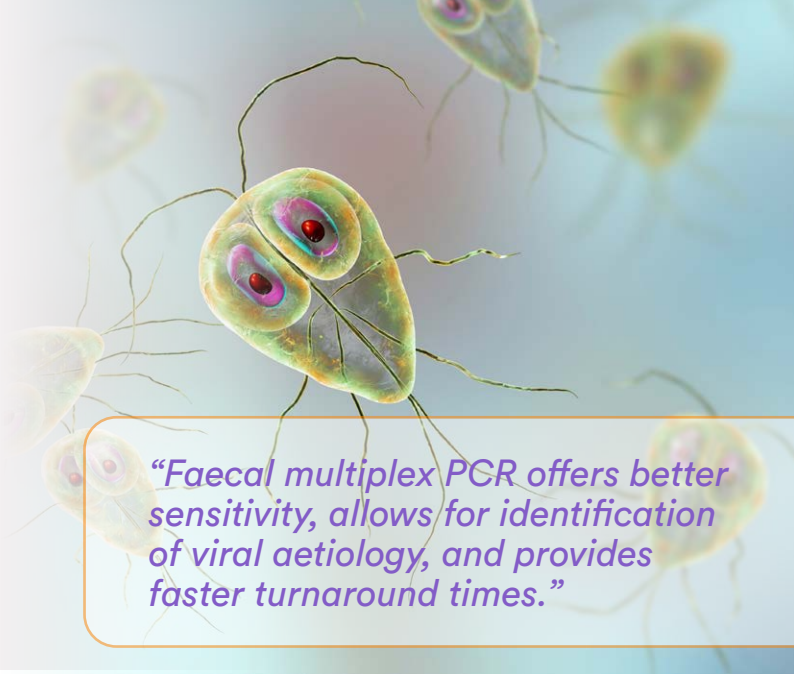
Table 1. Pathogens and diagnostic testing available at Clinical Labs WA

Bacteria	Testing	Comments
<i>Campylobacter</i>	Culture + PCR	Accounted for 94% of national notifiable enteric diseases in 2017 ³ . PCR cannot differentiate between typhoid/non-typhoid strains.
<i>Salmonella</i>	Culture + PCR. Culture is required for serotyping of Typhi/non-Typhi strains. Blood cultures in returned travellers suspicious of typhoid fever.	
<i>Shigella</i>	Culture + PCR	Can cause dysentery.
<i>C.difficile</i>	PCR	May be bowel commensals (especially in children <2 years old). Test only in symptomatic patients with recent antibiotic exposure.
Viruses	Testing	Comments
Rotavirus	Multiplex viral PCR	Can be vaccinated. Common cause of childhood diarrhoea.
Norovirus	Multiplex viral PCR	Common cause of outbreaks in nursing homes/schools.
Adenovirus	Multiplex viral PCR	Most adenoviruses can cause gastroenteritis. Adenovirus F40/F41 common cause of gastroenteritis outbreaks in children.
Enterovirus, astrovirus, sapovirus, bocavirus	Multiplex viral PCR	
Parasites	Testing	Comments
<i>Giardia spp.</i>	OCP microscopy/PCR	Cause of dysentery and liver abscess in returned travellers.
<i>Cryptosporidium spp.</i>	OCP microscopy/PCR	
<i>Entamoeba histolytica</i>	OCP microscopy/PCR	
<i>Dientamoeba fragilis, Blastocystis hominis</i>	OCP microscopy/PCR	Pathogenicity has not been established. Treatment is not routinely recommended. Exclude other causes in the first instance.
<i>Helminths – e.g. Enterobius, Strongyloides, Taenia, Schistosomiasis</i>	OCP microscopy only. Serology available for certain helminths.	Seen mainly in returned travellers. Travel history important. Collect 3 x specimens to improve sensitivity.

Benefits of faecal PCR testing

Multiplex PCR has become more readily available and commercially affordable, offering many advantages over traditional culture testing. PCR offers better sensitivity, allows for identification of viral aetiology, and provides faster turnaround times.

Although faecal multiplex PCR offers many benefits, referrers need to be aware of the following: Identification of the pathogen genome does not necessarily indicate disease. This is most classically seen with *C.difficile*, which is a bowel commensal and may not cause disease in healthy individuals. Similarly, in immunosuppressed patients, persistent viral shedding can often be found and does not represent active infection.



“Faecal multiplex PCR offers better sensitivity, allows for identification of viral aetiology, and provides faster turnaround times.”

Are faecal culture and faecal microscopy still needed?

Faecal microscopy and culture have remained the gold standard for many years and are still commonly requested. PCR will only identify the specific pathogens on the testing panel, potentially missing other causes of infection. Also, PCR does not allow for antimicrobial susceptibility testings for bacterial pathogens. Therefore, stool cultures remain an important part of microbiological workup.

Faecal culture

Faecal culture continues to be routinely performed and will identify many bacterial pathogens. However, one of its weaknesses is the failure to identify viral pathogens, which account for a significant number of infectious diarrhoea, particularly in children.

Faecal microscopy

Faecal microscopy is another important diagnostic tool, particularly when a parasitic cause is suspected, such as in returned travellers or those with agricultural exposure. In these instances, patient history should be included on the request, and specific ova, cyst, parasite (OCP) microscopy should be requested as these samples require special processing in the laboratory. Sensitivity of microscopy is time-dependent and can vary significantly depending on the stage of illness and severity. Three specimens are recommended for increased sensitivity.

How to order Faecal Multiplex PCR Testing

Request ‘Faecal Multiplex PCR’ using the standard Clinical Labs request form. This will test for the bacterial and parasitic pathogens as listed in Table 1. Faecal M/C/S will also be completed by the lab. If Multiplex Viral PCR testing is required, please add this separately to the request form.

Additional tests:

- In patients with gastrointestinal symptoms suggestive of inflammatory or functional bowel disease of more than 6 weeks’ duration a Faecal Calprotectin test may be ordered.
- Faecal occult bloods can also be requested.
- If helminth parasites (worms) are suspected, then add OCP.
- *C. difficile* needs to be specified as an additional test on the request form.

- If *Strongyloides* is suspected, please also request *Strongyloides* serology (serum sample).
- *Dientamoeba fragilis* and *Blastocystitis hominis* must be specifically requested.

Specimens required:

A fresh faecal sample in brown top container. Frozen faecal samples are also accepted; however, culture cannot be performed on these.

Test cost:

Bulk-billing is available through Medicare.

References

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Dr Eric Chu graduated from The University of Western Australia with a Bachelor of Medicine/Bachelor of Surgery in 2007. He undertook his post-graduate training primarily at Sir Charles Gairdner Hospital, and completed his clinical microbiology and physicians training in infectious diseases. He obtained dual fellowship in 2020 after additional training at Fiona Stanley Hospital, PathWest and Princess Margaret Hospital. Eric currently works as an Infectious Diseases Physician at Sir Charles Gairdner Hospital and joined Australian Clinical Labs as a clinical microbiologist in 2020.

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Dr Sudha Pottumarthy-Boddu comes to us from Houston, Texas, where she was Assistant Professor in the Department of Pathology and Laboratory Medicine at the University of Texas, School of Medicine. She was also the Technical Director of the Clinical Laboratory Services at the Houston Department of Health and Human Services. After graduating from medical school in India, Dr Pottumarthy-Boddu migrated to New Zealand and completed her Pathology/Microbiology Fellowship training with the Royal College of Pathologists of Australasia. She is a recipient of various awards and scholarships, including the Neil Prentice Memorial Prize of RCPA. She is also a Diplomate of the American Board of Medical Microbiology. Over the last 10 years she gained experience in various hospital, research, and public health laboratories in the US, publishing over 30 articles in peer-reviewed journals and presenting at various national and international conferences. Detection of the first USA isolate of *Enterobacter* spp., with NmcAcarbapenem hydrolyzing enzyme and establishing clinical significance of *Nocardia verterana* are noteworthy. Dr Pottumarthy-Boddu's main research interests are antimicrobial susceptibility trends and molecular methods in the diagnosis of infectious diseases.

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