MOLECULAR

# Gastrointestinal Pathogen Simplicity Pane $I^{\mathrm{TM}}$ 384 Well 

## Product Specification Sheet

RUO Product*

## Product Description:

The Gastrointestinal (GI) Pathogen Simplicity Pane ${ }^{T W M}$ is a multiplex polymerase chain reaction (PCR) assay for the identification of nucleic acids in 16 organisms frequently associated with GI tract infections. This method is highly accurate and analytically sensitive. It is used to identify organisms by amplifying and detecting genetic material of pathogens in samples. The panel will aid in the research of causative agents of Gl tract infections and their prevalence.
The target organisms included in the panel are as follows:
Bacteria: C. difficile ( $t c d A, t c d B$ ), Enterotoxigenic E. coli (ETEC), Shiga-like toxin-producing E. coli (STEC), Enteroinvasive E. coli (EIEC)/Shigella, Campylobacter (C. coli, C. jejuni, and C. Iari), Salmonella spp., Yersinia enterocolitica, and Vibrio spp.
Parasites: Cryptosporidium spp., Entamoeba histolytica, and Giardia lamblia
Viruses: Adenovirus F40/F41, Norovirus GI/GII, Rotavirus A, and Astrovirus

| Product Information |  | Product Specifications |  |
| :---: | :---: | :---: | :---: |
| Gastrointestinal Pathogen Simplicity Panel ${ }^{T m}$ (384 Well Plate) |  | QC Test | qPCR Cycle Threshold Percent CV |
| Part Number | $\begin{aligned} & \text { P-GI384-001-A } \\ & \text { P-GI384-002-A } \end{aligned}$ | Specification | $\leq 2.5$ |
| Number of Panels | 48 | QC Results |  |
|  |  | Positive | meets specification |
| Positive Control | Refer Page 3, step 6 | Negative | meets specification |
| Storage Temperature | $-25^{\circ} \mathrm{C}$ to $-15^{\circ} \mathrm{C}$ | Targets | meets specification |
| Disclaimer - Use of PCR and Patent <br> This product is for basic PCR and is outside of any valid US patents assigned to Hoffman La-Roche. | ISO Certification <br> This product was manufactured in a facility whose Quality Management System is certified as being in conformity with ISO 13485:2016 by Intertek. <br> * Limitations of Use <br> For Research Use Only. Not for use in diagnostic procedures. |  | oduct Guarantee <br> kit has been shown to generate able, repeatable and high-performance ults. <br> se contact Molecular Designs for nical assistance. If not completely sfied, our team will help you identify address the issue and replace the ays as needed. |

## Reagent Storage and Use Guidelines

1. Store all reagents at $-25^{\circ} \mathrm{C}$ to $-15^{\circ} \mathrm{C}$.
2. Do not freeze-thaw plates more than 3 times.

## The Following is Included in the Kit:

384 well PCR plate pre-loaded with the Gastrointestinal Pathogen Simplicity Panel ${ }^{\text {TM }}$ assays and positive controls. Negative control assay are plated but negative control is user supplied.

## The Following is Supplied by the User: Materials

1. Extracted Sample(s)
2. PCR optical film
3. Sealer for optical film
4. Negative Control

## Equipment

1. Manual defrost $-20^{\circ} \mathrm{C}$ freezer
2. Laminar Flow or PCR Dead Air Box for general plate setup. Do not use Laminar Flow for infectious samples
3. Pipette and appropriate filtered pipette tips
4. Plate Vortex [recommend Vortex Genie 2 (Model G560) with the 3-inch platform and rubber cover]
5. Plate centrifuge

## Instrumentation

1. CFX384 Touch Real-Time PCR Detection System (or equivalent)

## General Guidelines and Safety Precautions

1. As with any test procedure, good laboratory practice is essential to the proper performance of this assay. Due to the high sensitivity of this test, care should be taken to keep reagents and amplification mixtures free of contamination.
a) Do not pipette by mouth.
b) Do not eat, drink, or smoke in designated work areas.
c) Wear laboratory gloves, laboratory coats, and eye protection when handling samples and reagents. Gloves must be changed between handling samples to prevent contamination. Avoid contaminating gloves when handling samples and controls.
d) Wash hands thoroughly after handling samples and kit reagents, and after removing the gloves.
e) Thoroughly clean and disinfect all laboratory work surfaces.
NOTE: Do not use sodium hypochlorite solution (bleach) to clean up a spill or to disinfect a plate before disposal as it can react with the common extraction reagents and generate toxic byproducts.
If spills occur, follow internal procedures to immediately clean and decontaminate the surface of instrument.
2. A laminar flow or PCR Dead Air Box is recommended to reduce contamination probability.
3. The use of filtered, sterile and nuclease-free pipette tips is recommended.
4. False positive results may occur if carryover of samples is not adequately controlled during sample handling and processing.

## Usage Information

## Reaction Plate Setup

1. Remove a reaction plate from the $-20^{\circ} \mathrm{C}$ manual defrost freezer.
2. Use the plate within 1 hour of thawing, keep sealed and store refrigerated at $4^{\circ} \mathrm{C}$, if not using immediately.
3. Spin down the plate for 30 seconds in a plate centrifuge.
4. Carefully remove the foil seal from the plate.
5. Add $4.0 \mu \mathrm{~L}$ of the sample being tested to each of the target wells.
6. Do not add any additional liquid to the Positive Control wells. All components have been added to these wells. Add $4.0 \mu \mathrm{~L}$ of Negative Control to each Negative Control well.

The following wells have the controls:

- Negative well M19/ Positive well 019:
C. difficile tcdA / C. difficile tcdB (CFO)/ Endogenous Control (CFR)
- Negative well M20/ Positive well O20:
E. coli: Enterotoxin E. coli / Shiga-like toxin-producing E. coli (CFO)/ Shigella/Enteroinvasive E. coli (STEC) (CFR)
- Negative well M21/ Positive well 021: Parasites: Cryptosporidium spp. (CFO)/ E. histolytica / G. Lamblia (CFR)
- Negative well M22/ Positive well 022: : Bacteria: Campylobacter (CFO) /Salmonella spp. (CFR) / Y. enterocolitica (Q670) / Vibrio spp.
- Negative well M23/ Positive well 023:

Virus 1: Adenovirus (Q670) /Rotavirus (CFR)/ Astrovirus

- Negative well M24/ Positive well 024: :

Virus 2: Norovirus (CFO)
7. Seal the PCR plate using optical qPCR film. Note: If using a partial plate, remove the excess optical seal using the utility knife and ensure the plate is sealed.

1. Optional: vortex the plate, at least 5 seconds per plate quadrant.
2. Optional: spin down the plate in a plate centrifuge

## Procedural Notes

1. Do not reuse consumables. They are for one-time use only.
2. Always use caution when transferring specimens from a primary collection tube to a secondary tube.
3. Use pipettes with aerosol-barrier or positivedisplacement tips to handle specimens.
4. Always use a new pipette tip for each specimen.
5. For testing of previously frozen sample, ensure samples are equilibrated to room temperature and well mixed prior to use.

Target Layout per panel (384 well plate, 48 panels per plate). See page 5 for the complete layout of a 384 well plate.

| C. difficile |
| :---: |
| C. difficile tcdA / C. difficile tcdB (CFO) / |
| Endogenous Control (CFR) |
| E. coli |
| Enterotoxigenic E. coli (ETEC) / Shiga-like |
| toxin-producing E. coli (STEC) (CFO) / |
| Enteroinvasive E. coli (EIEC)/Shigella (CFR) |
| Parasites |
| E. Histolytica / Cryptosporidium spp. (CFO) |
| $/$ |
| G. lamblia (CFR) |
| Bacteria |
| Vibrio spp. / Campylobacter (CFO) / |
| Salmonella spp. (CFR) / |
| Y. enterocolitica (Q670) |
| Virus 1 |
| Astrovirus / Rotavirus (CFR) / |
| Adenovirus (Q670) |
| Virus 2 |
| Norovirus (CFO) |

If not noted, the fluorophore is FAM and additional channels are noted in parentheses signifying multiple targets in the primer and probe mixture which will be detected on the qPCR instrument. CFO is CAL Fluor Orange 560, CFR is CAL Fluor Red 610 and Q670 is equivalent to Quasar 670.

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## Usage Information

## Real-Time PCR Detection System PCR Run Setup

1. Open the specified run template and fill in the sample name fields with unique sample IDs corresponding to the samples being processed.
2. NOTE: This step can also be done prior to reaction plate setup if sample IDs have already been specified.
3. Place the reaction plate into the instrument in the appropriate orientation (A1 in the upper left corner), close the instrument lid and initiate the run.
4. NOTE: When running a partial plate, a balance is required at the other side of the instrument to ensure that the lid is sealed properly and doesn't break the instrument.

## Thermocycling Protocol

1. Reverse Transcription
a) 15 minutes at $55^{\circ} \mathrm{C}$
2. Denaturation
a) 3 minutes at $95^{\circ} \mathrm{C}$
3. Annealing and Extension 40 cycles consisting of:
a) 5 seconds at $95^{\circ} \mathrm{C}$
b) 30 seconds at $60^{\circ} \mathrm{C}$, with fluorescence acquisition during this step

## Amplification Interpretation and Troubleshooting

1. The laboratory should establish cycle threshold (CT) cutoffs as appropriate for their sample workflow and procedures. It is recommended that CT cutoffs are determined during the validation of the test.
2. The laboratory should evaluate the curve shape when considering whether a sample with a given CT should be considered positive:
a) Plate sealing issues can lead to jagged curve shapes or rising/decreasing baselines that lead to inaccurate data (erroneous CT value).
b) Inappropriate mixing or centrifuging can lead to inaccurate data.
3. If user suspects contamination, it is recommended to clean and disinfect the laboratory area and re-test to ensure proper results.
4. Any failure of the positive or negative control should require a repeat run. If the control failure continues, it is recommended to have the qPCR instrument and the sample extraction workflow evaluated to ensure they are functioning properly.


