

3425 Corporate Way Duluth, GA 30096

Patient: SAMPLE PATIENT

DOB: Sex:

MRN:

2200 GI Effects™ Eundamentals - Stor			
		QUINTILE DISTRIBUTION	
Methodology: GC/MS, Automated Chemistry, ElA	Result		Reference Range
	Diges	tion and Absorption	
Pancreatic Elastase 1 †	158 L	100 200	>200 mcg/g
Products of Protein Breakdown (Total*) (Valerate, Isobutyrate, Isovalerate)	6.0		1.8-9.9 micromol/g
Fecal Fat (Total*)	19.5	+ + + +	3.2-38.6 mg/g
Triglycerides	1.1	+ + + +	0.3-2.8 mg/g
Long-Chain Fatty Acids	12.9	+ + + +	1.2-29.1 mg/g
Cholesterol	0.5		0.4-4.8 mg/g
Phospholipids	5.0		0.2-6.9 mg/g
	Inflamm	ation and Immunology	
Calprotectin †	145 H	50 120	<=50 mcg/g
Eosinophil Protein X (EPX)†	4.9 <b>H</b>	1.1 4.6 ◆	<=4.6 mcg/g
	Gut Mic	crobiome Metabolites	
Metabolic			
Short-Chain Fatty Acids (SCFA) (Total*) (Acetate, n-Butyrate, Propionate)	81.3		>=23.3 micromol/g
n-Butyrate Concentration	18.1		>=3.6 micromol/g
n-Butyrate %	22.3		11.8-33.3 %
Acetate %	63.1		48.1-69.2 %
Propionate %	14.6		<=29.3 %
Beta-glucuronidase	2,297		368-6,266 U/g

\*Total value is equal to the sum of all measurable parts. †These results are not represented by quintile values.

Tests were developed and their performance characteristics determined by Genova Diagnostics. Unless otherwise noted with Ø, the assays have not been cleared by the U.S. Food and Drug Administration.



NG

No Growth

Methodology: Culture/MALDI-TOF MS, Automated and Manual Biochemical Methods, Vitek® 2 System Microbial identification and Antibiotic susceptibility

Ρ

Pathogen

### **Gastrointestinal Microbiome (Culture)**

Human microflora is influenced by environmental factors and the competitive ecosystem of the organisms in the GI tract. Pathogenic significance should be based upon clinical symptoms.

Microbiology Legend

PP

Potential

NP

Non-

#### Additional Bacteria

**Non-Pathogen:** Organisms that fall under this category are those that constitute normal, commensal flora, or have not been recognized as etiological agents of disease.

**Potential Pathogen:** Organisms that fall under this category are considered potential or opportunistic pathogens when present in heavy growth. **Pathogen:** The organisms that fall under this category have a well-recognized mechanism of pathogenicity in clinical literature and are considered significant regardless of the quantity that appears in the culture.



## **KOH Preparation for Yeast**

Methodology: Potassium Hydroxide (KOH) Preparation for Yeast

#### Potassium Hydroxide (KOH) Preparation for Yeast

These yeast usually represent the organisms isolated by culture. In the presence of a negative yeast culture, microscopic yeast may reflect organisms not viable enough to grow in culture. The presence of yeast on KOH prep should be correlated with the patient's symptoms. However, moderate to many yeast suggests yeast overgrowth.

#### Result

KOH Preparation, stool

Few Yeast Present

The result is reported as the amount of yeast seen microscopically: Rare: 1-2 per slide Few: 2-5 per high power field (HPF) Moderate: 5-10 per HPF Many: >10 per HPF

\*\* Indicates testing performed by Genova Diagnostics, Inc. 63 Zi Ilicoa St., Ash eville, NC 28801-0174 A. L. Peace-Brewer, PhD, D(ABM LI), Lab Director - CLI A Lic. #34D0655571 - Medicare Lic. #34-8475

## Parasitology

### **Microscopic O&P Results**

Microscopic O&P is capable of detecting all described gastrointestinal parasites. The organisms listed in the box represent those commonly found in microscopic stool analysis. Should an organism be detected that is not included in the list below, it will be reported in the Additional Results section. For an extensive reference of all potentially detectable organisms, please visit www.gdx.net/product/gi-effects-comprehensive-stool-test

Genus/species	Result
Nematodes - roundworms	
Ancylostoma/Necator (Hookworm)	Not Detected
Ascaris lumbricoides	Not Detected
Capillaria philippinensis	Not Detected
Enterobius vermicularis	Not Detected
Strongyloides stercoralis	Not Detected
Trichuris trichiura	Not Detected
Cestodes - tapeworms	
Diphyllobothrium latum	Not Detected
Dipylidium caninum	Not Detected
Hymenolepis diminuta	Not Detected
Hymenolepis nana	Not Detected
Taenia spp.	Not Detected
Trematodes - flukes	
Clonorchis/Opisthorchis spp.	Not Detected
Fasciola spp./ Fasciolopsis buski	Not Detected
Heterophyes/Metagonimus	Not Detected
Paragonimus spp.	Not Detected
Schistosoma spp.	Not Detected
Protozoa	
Balantidium coli	Not Detected
Blastocystis spp.	Rare Detected
Chilomastix mesnili	Not Detected
Cryptosporidium spp.	Not Detected
Cyclospora cayetanensis	Not Detected
Dientamoeba fragilis	Moderate Detected
Entamoeba coli	Not Detected
Entamoeba histolytica/dispar	Not Detected
Entamoeba hartmanii	Not Detected
Entamoeba polecki	Not Detected
Endolimax nana	Not Detected
Giardia	Not Detected
lodamoeba buetschlii	Not Detected
Cystoisospora spp.	Not Detected
Trichomonads (e.g. Pentatrichomonas)	Not Detected
Additional Findings	
White Blood Cells	Not Detected
Charcot-Leyden Crystals	Not Detected
Other Infectious Findings	

One negative specimen does not rule out the possibility of a parasitic infection.

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Parasitology							
PCR Parasitology - Protozoa**		Add-on testing	Methodologies: DNA by PCR, Next Generation Sequencing				
Organism	Result	Units		Expected Result			
Blastocystis spp.	6.00e2	femtograms/microliter C&S stool	Detected	Not Detected			
Cryptosporidium spp.	<4.87e2	genome copies/microliter C&S stool	Not Detected	Not Detected			
Cyclospora cayetanensis	<2.65e2	genome copies/microliter C&S stool	Not Detected	Not Detected			
Dientamoeba fragilis	6.40e2	genome copies/microliter C&S stool	Detected	Not Detected			
Entamoeba histolytica	<1.14e3	genome copies/microliter C&S stool	Not Detected	Not Detected			
Giardia	<1.57e2	genome copies/microliter C&S stool	Not Detected	Not Detected			

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Additional Results					
Methodology: Fecal Immunochemical Testin	g (FIT)				
	Result	Expected Value			
Fecal Occult Blood	Negative	Negative			
Color††	Green				
Consistency <sup>††</sup>	Formed/Normal				

††Results provided from patient input.

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	Z	onulin Family Peptide	
Methodology: EIA	Result	Reference Range	Zonulin Family Peptide
Zonulin Family Peptide, Stool	100.0	22.3-161.1 ng/mL	This test is for research use only. Genova will not provide support on interpreting the test results. This test does not detect zonulin. <sup>1</sup> The Scheffler paper suggests that the IDK kit may detect a zonulin family peptide, such as properdin. Genova's unpublished data demonstrated that the current IDK kit results were associated with stool inflammation biomarkers and an inflammation-associated dysbiosis profile. The performance characteristics of Zonulin Family Peptide
			have been verified by Genova Diagnostics, Inc. The assay
			has not been cleared by the U.S. Food and Drug

Administration.

#### **Reference:**

1. Scheffler L, et al. Widely Used Commercial ELISA Does Not Detect Precursor of Haptoglobin2, but Recognizes Properdin as a Potential Second Member of the Zonulin Family. *Front Endocrinol.* 2018;9:22.

© Genova Diagnostics · Robert M. David, PhD, Lab Director · CLIA Lic. #11D0255349 · Medicare Lic. #34-8475 · Georgia Lab Lic. Code #067-007 New York Clinical Lab PFI #4578 · Florida Clinical Lab Lic. #800008124 Macroscopic/Direct Exam for Parasites

Methodology: Macroscopic Evaluation

No human parasite detected in sample.

Add-on Testing							
Methodology: EIA	Result			Reference Range			
Fecal secretory IgA	206	- + +	♦	<=885 mcg/g			
Methodology: EIA	Result	Expected Value					
HpSA - <i>H. pylori</i>	Negative	Negative	HpSA ( <i>Helicobacter pylo</i> <i>Helicobacter pylori</i> is a bac	<b>pri stool antigen)</b> terium which causes peptic			
<i>Campylobacter</i> spp.⊠**	Negative	Negative	ulcer disease and plays a r gastric cancer. Direct stool	ole in the development of testing of the antigen (HpSA)			
Clostridium difficile⊠**	Negative	Negative	is highly accurate and is ap follow-up of infection.	propriate for diagnosis and			
Shiga toxin <i>E. coli⊠</i> **	Negative	Negative					

#### Clostridium difficile

*Clostridium difficile* is an anaerobic, spore-forming gram-positive bacterium. After a disturbance of the gut flora (usually with antibiotics), colonization with *Clostridium difficile* can take place. *Clostridium difficile* infection is much more common than once thought.

#### Shiga toxin E. coli

Shiga toxin-producing *Escherichia coli* (STEC) is a group of bacterial strains that have been identified as worldwide causes of serious human gastrointestinal disease. The subgroup enterohemorrhagic *E. coli* includes over 100 different serotypes, with 0157:H7 being the most significant, as it occurs in over 80% of all cases. Contaminated food continues to be the principal vehicle for transmission; foods associated with outbreaks include alfalfa sprouts, fresh produce, beef, and unpasteurized juices.

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Methodology: Vitek 2® System Microbial Antibiotic susceptibility, Manual Minimum Inhibition Concentration

### Mycology Sensitivity

# Azole Antifungals

0							
Candida species	R	I	S-I	DD	S		NI
Fluconazole					0.5		
Voriconazole					<=0.008		
Nystatin	=50						
Natural Agents						•	
Candida species	LOW INHIBITION					ŀ	HIGH INHIBITION
Berberine							
Caprylic Acid							
Garlic							
Undecylenic Acid							
Plant tannins							
Uva-Ursi							

#### **Prescriptive Agents:**

The R (Resistant) category implies isolate is not inhibited by obtainable levels of pharmaceutical agent.

The I (Intermediate) category includes isolates for which the minimum inhibition concentration (MIC) values usually approach obtainable pharmaceutical agent levels and for which response rates may be lower than for susceptible isolates.

The S-DD (Susceptible-Dose Dependent) category implies clinical efficacy when higher than normal dosage of a drug can be used and maximal concentration achieved.

The S (Susceptible) column implies that isolates are inhibited by the usually achievable concentrations of the pharmaceutical agent.

NI (No Interpretive guidelines established) category is used for organisms that currently do not have established guidelines for MIC interpretation.

Refer to published pharmaceutical guidelines for appropriate dosage therapy.

#### Nystatin and Natural Agents:

Results for Nystatin are being reported with natural antifungals in this category in accordance with laboratory guidelines for reporting sensitivities. In this assay, inhibition is defined as the reduction level on organism growth as a direct result of inhibition by a natural substance. The level of inhibition is an indicator of how effective the substance was at limiting the growth of an organism in an in vitro environment. High inhibition indicates a greater ability by the substance to limit growth, while Low Inhibition a lesser ability to limit growth. The designated natural products should be considered investigational in nature and not be viewed as standard clinical treatment substances.

Methodology: Vitek 2® System Microbial Antibiotic susceptibility, Manual Minimum Inhibition Concentration

## **Bacteria Sensitivity**

# **Prescriptive Agents**

Klebsiella pneumoniae	R		L I	S-DD		S		NI
Ampicillin	R							
Amox./Clavulanic Acid						S		
Cephalothin						S		
Ciprofloxacin						S		
Tetracycline						S		
Trimethoprim/Sulfa						S		
Natural Agents		_			_		_	

Klebsiella pneumoniae	LOW INHIBITION	HIGH INHIBITION
Berberine		
Oregano		
Plant Tannins		
Uva-Ursi		

#### **Prescriptive Agents:**

The R (Resistant) category implies isolate is not inhibited by obtainable levels of pharmaceutical agent.

The I (Intermediate) category includes isolates for which the minimum inhibition concentration (MIC) values usually approach obtainable pharmaceutical agent levels and for which response rates may be lower than for susceptible isolates.

The S-DD (Susceptible-Dose Dependent) category implies clinical efficacy when higher than normal dosage of a drug can be used and maximal concentration achieved.

The S (Susceptible) column implies that isolates are inhibited by the usually achievable concentrations of the pharmaceutical agent.

NI (No Interpretive guidelines established) category is used for organisms that currently do not have established guidelines for MIC interpretation.

Refer to published pharmaceutical guidelines for appropriate dosage therapy.

#### Natural Agents:

In this assay, inhibition is defined as the reduction level on organism growth as a direct result of inhibition by a substance. The level of inhibition is an indicator of how effective the substance was at limiting the growth of an organism in an in vitro environment. High inhibition indicates a greater ability by the substance to limit growth, while Low Inhibition a lesser ability to limit growth. The designated natural products should be considered investigational in nature and not be viewed as standard clinical treatment substances.