

ID: P0000000000 SEX: Male

DOB: AGE: 37

CLIENT #: 12345 DOCTOR:

Doctor's Data, Inc. 3755 Illinois Ave.

St. Charles, IL 60174 U.S.A.

**Dysbiotic flora** 

# Comprehensive Stool Analysis / Parasitology x3

## **BACTERIOLOGY CULTURE**

## Expected/Beneficial flora Commensal (Imbalanced) flora

2+ Alpha hemolytic strep

1+ Beta strep, not group A or B

2+ Hemolytic Escherichia coli

NG Bifidobacterium spp.

4+ Bacteroides fragilis group

4+ Escherichia coli3+ Lactobacillus spp.

1+ Enterococcus spp.

4+ Clostridium spp.

NG = No Growth

#### **BACTERIA INFORMATION**

**Expected /Beneficial bacteria** make up a significant portion of the total microflora in a healthy & balanced GI tract. These beneficial bacteria have many health-protecting effects in the GI tract including manufacturing vitamins, fermenting fibers, digesting proteins and carbohydrates, and propagating anti-tumor and anti-inflammatory factors.

Clostridia are prevalent flora in a healthy intestine. Clostridium spp. should be considered in the context of balance with other expected/beneficial flora. Absence of clostridia or over abundance relative to other expected/beneficial flora indicates bacterial imbalance. If C. difficile associated disease is suspected, a Comprehensive Clostridium culture or toxigenic C. difficile DNA test is recommended.

Commensal (Imbalanced) bacteria are usually neither pathogenic nor beneficial to the host GI tract. Imbalances can occur when there are insufficient levels of beneficial bacteria and increased levels of commensal bacteria. Certain commensal bacteria are reported as dysbiotic at higher levels.

**Dysbiotic bacteria** consist of known pathogenic bacteria and those that have the potential to cause disease in the GI tract. They can be present due to a number of factors including: consumption of contaminated water or food, exposure to chemicals that are toxic to beneficial bacteria; the use of antibiotics, oral contraceptives or other medications; poor fiber intake and high stress levels.

# YEAST CULTURE

#### **Normal flora**

+ Rhodotorula mucilaginosa

### **Dysbiotic flora**

## **MICROSCOPIC YEAST**

Result:

**Expected:** 

Many

None - Rare

Yeast in stool is expected at a level of nonerare. A microscopic finding of yeast in stool of few, moderate, or many may be helpful in identifying potential yeast overgrowth, or nonviable or dietary yeast.

## YEAST INFORMATION

Yeast may normally be present in small quantities in the skin, mouth, and intestine. When investigating the presence of yeast, disparity may exist between culturing and microscopic examination. Yeast are not uniformly dispersed throughout the stool and this may lead to undetectable or low levels of yeast identified by microscopy, despite culture and identified yeast species. Conversely, microscopic examination may reveal a significant amount of yeast present but no viable yeast cultured. Yeast may not always survive transit through the intestines. Nonviable diet-derived yeast may also be detected microscopically. Consideration of clinical intervention for yeast detected microscopically should be made in the context of other findings and presentation of symptoms.

# Comments:

Date Collected: 06/30/2017 Date Received: 07/03/2017 Date Reported: 07/17/2017 \* Aeromonas, Campylobacter, Plesiomonas, Salmonella, Shigella, Vibrio, Yersinia, & Edwardsiella tarda have been specifically tested for and found absent unless

reported.





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#### PARASITOLOGY/MICROSCOPY

# Sample 1

None Ova or Parasites

Rare Yeast

# Sample 2

None Ova or Parasites

Few Yeast

## Sample 3

None Ova or Parasites

Many Yeast

#### PARASITOLOGY INFORMATION

Intestinal parasites are abnormal inhabitants of the gastrointestinal tract that have the potential to cause damage to their host. The presence of any parasite within the intestine generally confirms that the patient has acquired the organism through fecal-oral contamination. Damage to the host includes parasitic burden, migration, blockage and pressure. Immunologic inflammation, hypersensitivity reactions and cytotoxicity also play a large role in the morbidity of these diseases. The infective dose often relates to severity of the disease and repeat encounters can be additive.

There are two main classes of intestinal parasites, they include protozoa and helminths. The protozoa typically have two stages; the trophozoite stage that is the metabolically active, invasive stage and the cyst stage, which is the vegetative inactive form resistant to unfavorable environmental conditions outside the human host. Helminths are large, multicellular organisms. Like protozoa, helminths can be either free-living or parasitic in nature. In their adult form, helminths cannot multiply in humans.

In general, acute manifestations of parasitic infection may involve diarrhea with or without mucus and or blood, fever, nausea, or abdominal pain. However these symptoms do not always occur. Consequently, parasitic infections may not be diagnosed or eradicated. If left untreated, chronic parasitic infections can cause damage to the intestinal lining and can be an unsuspected cause of illness and fatigue. Chronic parasitic infections can also be associated with increased intestinal permeability, irritable bowel syndrome, irregular bowel movements, malabsorption, gastritis or indigestion, skin disorders, joint pain, allergic reactions, and decreased immune function.

In some instances, parasites may enter the circulation and travel to various organs causing severe organ diseases such as liver abscesses and cysticercosis. In addition, some larval migration can cause pneumonia and in rare cases hyper infection syndrome with large numbers of larvae being produced and found in every tissue of the body.

One negative parasitology x1 specimen does not rule out the possibility of parasitic disease, parasitology x3 is recommended. This test is not designed to detect Cyclospora cayetanensis or Microsproridia spp.

#### GIARDIA/CRYPTOSPORIDIUM IMMUNOASSAY Giardia duodenalis (AKA intestinalis and lamblia) Within Outside Reference Range is a protozoan that infects the small intestine and is passed in stool and spread by the fecal-oral Giardia duodenalis Neg Neg route. Waterborne transmission is the major source of giardiasis. Cryptosporidium is a coccidian protozoa that Cryptosporidium Neg Neg can be spread from direct person-to-person contact or waterborne transmission.

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DIGESTION /ABSORPTION				
	Within	Outside	Reference Range	Elastase findings can be used for the diagnosis or the exclusion of exocrine pancreatic
Elastase		182	] > 200 μg/mL	insufficiency. Correlations between low levels and chronic pancreatitis and cancer have been reported. <b>Fat Stain:</b> Microscopic determination
Fat Stain	None		None - Mod	of fecal fat using Sudan IV staining is a qualitative procedure utilized to assess fat absorption and to detect steatorrhea. <b>Muscle</b>
Muscle fibers	None		None - Rare	<b>fibers</b> in the stool are an indicator of incomplete digestion. Bloating, flatulence, feelings of "fullness" may be associated with increase in
Vegetable fibers	Rare		None - Few	muscle fibers. <b>Vegetable fibers</b> in the stool may be indicative of inadequate chewing, or eating "on the run". <b>Carbohydrates:</b> The presence of
Carbohydrates	Neg		Neg	reducing substances in stool specimens can indicate carbohydrate malabsorption.

			INFLAMMATION	
	Within	Outside	Reference Range	Lactoferrin and Calprotectin are reliable markers for differentiating organic inflammation
Lactoferrin	< 0.5		< 7.3 μg/mL	(IBD) from function symptoms (IBS) and for management of IBD. Monitoring levels of fecal lactoferrin and calprotectin can play an essential
Calprotectin*	< 10		= 50 μg/g	role in determining the effectiveness of therapy, are good predictors of IBD remission, and can indicate a low risk of relapse. <b>Lysozyme*</b> is an
Lysozyme*	174		<= 600 ng/mL	enzyme secreted at the site of inflammation in the GI tract and elevated levels have been identified in IBD patients. <b>White Blood Cells</b>
White Blood Cells	None		None - Rare	(WBC) and <b>Mucus</b> in the stool can occur with bacterial and parasitic infections, with mucosal irritation, and inflammatory bowel diseases such
Mucus	Neg		Neg	as Crohn's disease or ulcerative colitis.

			IMMUNOLOGY	
	Within	Outside	Reference Range	Secretory IgA* (sIgA) is secreted by mucosal tissue and represents the first line of defense of
Secretory IgA*		19.6	51 - 204 mg/dL	the GI mucosa and is central to the norn function of the GI tract as an immune barri Elevated levels of slgA have been associat with an upregulated immune response.

Comments:

Date Collected: 06/30/2017 \*For Research Use Only. Not for use in diagnostic procedures.

Date Received: 07/03/2017 Methodology: Elisa, Microscopy, Colormetric,

Date Completed: 07/17/2017 Gas Chromotography, ph Electrode



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			SHORT CHAIN FATTY AC	EIDS
	Within	Outside	Reference Range	Short chain fatty acids (SCFAs): SCFAs are the end product of the bacterial fermentation
% Acetate	57		40 - 75 %	process of dietary fiber by beneficial flora in the gut and play an important role in the health of the GI as well as protecting against intestinal
% Propionate	20		9 - 29 %	dysbiosis. Lactobacilli and bifidobacteria produce large amounts of short chain fatty acids, which decrease the pH of the intestines and therefore
% Butyrate	19		9 - 37 %	make the environment unsuitable for pathogens, including bacteria and yeast. Studies have shown that SCFAs have numerous implications in
% Valerate	4.0		0.5 - 7 %	maintaining gut physiology. SCFAs decrease inflammation, stimulate healing, and contribute to normal cell metabolism and differentiation. Levels
Butyrate	2.3		0.8 - 4.8 mg/mL	of <b>Butyrate</b> and <b>Total SCFA</b> in mg/mL are important for assessing overall SCFA production,
Total SCFA's	12		4 - 18 mg/mL	and are reflective of beneficial flora levels and/or adequate fiber intake.

INTESTINAL HEALTH MARKERS				
	Within	Outside	Reference Range	Red Blood Cells (RBC) in the stool may be associated with a parasitic or bacterial infection,
Red Blood Cells	None		None - Rare	or an inflammatory bowel condition such as ulcerative colitis. Colorectal cancer, anal fistulas, and hemorrhoids should also be ruled out.
рН	6.4		6 - 7.8	<b>pH:</b> Fecal pH is largely dependent on the fermentation of fiber by the beneficial flora of the gut.
Occult Blood	Neg		Neg	<b>Occult blood:</b> A positive occult blood indicates the presence of free hemoglobin found in the stool, which is released when red blood cells are lysed.

MACROSCOPIC APPEARANCE			
	Appearance	Expected	Color: Stool is normally brown because of pigments formed by bacteria acting on bile introduced into the digestive system from the
Color	Brown	Brown	liver. While certain conditions can cause changes in stool color, many changes are harmless and are caused by pigments in foods
Consistency	Soft	Formed/Soft	or dietary supplements. <b>Consistency</b> : Stool normally contains about 75% water and ideally should be formed and soft. Stool consistency can vary based upon transit time and water absorption.

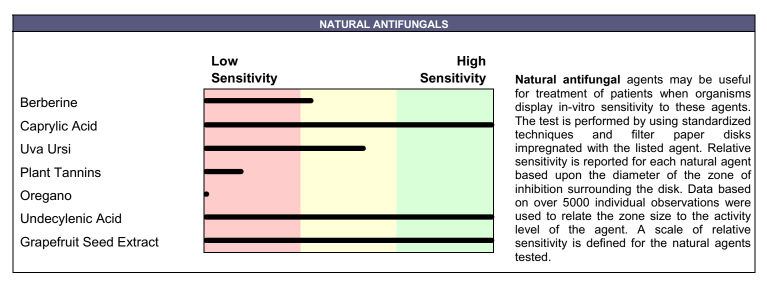


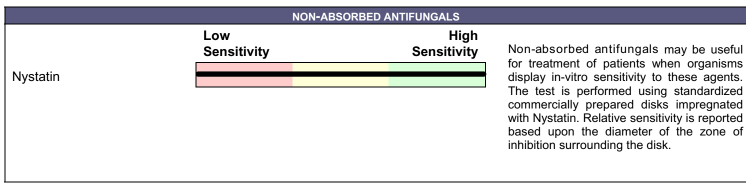
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# Yeast Susceptibilities: Rhodotorula mucilaginosa





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Yeast antifungal susceptibility testing is intended for research use only.

Not for use in diagnostic procedures.

v10.11

Lab number: F000000-0000-0 CSAPx3 Page: 1
Patient: Sample Patient Client: 12345

### INTRODUCTION

This analysis of the stool specimen provides fundamental information about the overall gastrointestinal health of the patient. When abnormal microflora or significant aberrations in intestinal health markers are detected, specific interpretive paragraphs are presented. If no significant abnormalities are found, interpretive paragraphs are not presented.

# Clostridium spp

Clostridia are expected inhabitants of the human intestine. Although most clostridia in the intestine are not virulent, certain species have been associated with disease. Clostridium perfringens is a major cause of food poisoning and is also one cause of antibiotic-associated diarrhea. Clostridium difficile is a causative agent in antibiotic-associated diarrhea and pseudomembranous colitis. Other species reported to be prevalent in high amounts in patients with Autistic Spectrum Disorder include Clostridium histolyticum group, Clostridium cluster I, Clostridium bolteae, and Clostridium tetani.

If these disease associations are a concern further testing may be necessary.

Washington W, Allen S, Janda W, Koneman E, Procop G, Schreckenberger P, Woods, G. Koneman's Color Atlas and Textbook of Diagnostic Microbiology, 6th edition. Lippincott Williams and Wilkins; 2006. pg 931-939

Song Y, Liu C, Finegold SM. Real-Time PCR Quantitation of Clostridia in Feces of Autistic Children. Applied and Environmental Microbiology. Nov. 2004, 6459-6465.

Parracho H, Bingham MO, Gibson GR, McCartney AL. Differences Between the Gut Microflora of Children with Autistic Spectrum Disorders and That of Healthy Children. Journal of Medical Microbiology. 2005;54, 987-991.

#### Imbalanced flora

Imbalanced flora are those bacteria that reside in the host gastrointestinal tract and neither injure nor benefit the host. Certain dysbiotic bacteria may appear under the imbalances category if found at low levels because they are not likely pathogenic at the levels detected. When imbalanced flora appear, it is not uncommon to find inadequate levels of one or more of the beneficial bacteria and/or a fecal pH which is more towards the alkaline end of the reference range (6 - 7.8). It is also not uncommon to find hemolytic or mucoid E. coli with a concomitant deficiency of beneficial E. coli and alkaline pH, secondary to a mutation of beneficial E. coli in alkaline conditions (DDI observations). Treatment with antimicrobial agents is unnecessary unless bacteria appear under the dysbiotic category.

Lab number: F000000-0000-0 CSAPx3 Page: 2
Patient: Sample Patient Client: 12345

Mackowiak PA. The normal microbial flora. N Engl J Med. 1982;307(2):83-93.

## **Cultured Yeast**

Yeast, such as Candida are normally present in the GI tract in very small amounts. Many species of yeast exist and are commensal; however, they are always poised to create opportunistic infections and have detrimental effects throughout the body. Factors that contribute to a proliferation of yeast include frequent use of wide-spread antibiotics/low levels of beneficial flora, oral contraceptives, pregnancy, cortisone and other immunosuppressant drugs, weak immune system/low levels of slgA, high-sugar diet, and high stress levels.

When investigating the presence of yeast, disparity may exist between culturing and microscopic examination. Yeast grows in colonies and is typically not uniformly dispersed throughout the stool. This may lead to undetectable or low levels of yeast identified by microscopy, despite a cultured amount of yeast. Conversely, microscopic examination may reveal a significant amount of yeast present, but no yeast cultured. Yeast does not always survive transit through the intestines rendering it unviable for culturing. Therefore, both microscopic examination and culture are helpful in determining if abnormally high levels of yeast are present.

## Microscopic yeast

Microscopic examination has revealed yeast in this stool sample. The microscopic finding of yeast in the stool is helpful in identifying whether the proliferation of fungi, such as Candida albicans, is present. Yeast is normally found in very small amounts in a healthy intestinal tract. While small quantities of yeast (reported as none or rare) may be normal, yeast observed in higher amounts (few, moderate to many) is considered abnormal.

An overgrowth of intestinal yeast is prohibited by beneficial flora, intestinal immune defense (secretory IgA), and intestinal pH. Beneficial bacteria, such as Lactobacillus colonize in the intestines and create an environment unsuitable for yeast by producing acids, such as lactic acid, which lowers intestinal pH. Also, lactobacillus is capable of releasing antagonistic substances such as hydrogen peroxide, lactocidin, lactobacillin, and acidolin.

Many factors can lead to an overgrowth of yeast including frequent use of antibiotics (leading to insufficient beneficial bacteria), synthetic corticosteroids, oral contraceptives, and diets high in sugar. Although there is a wide range of symptoms which can result from intestinal yeast overgrowth, some of the most common include brain fog, fatigue, reccurring vaginal or bladder infections, sensitivity to smells (perfumes, chemicals, environment), mood swings/depression, sugar and carbohydrate cravings, gas/bloating, and constipation or loose stools.

A positive yeast culture (mycology) and sensitivity to prescriptive and natural agents is helpful in determining which anti-fungal agents to use as part of a therapeutic treatment plan for chronic colonic yeast. However, yeast are colonizers and do not appear to be dispersed uniformly throughout the stool. Yeast may therefore be observed microscopically, but not grow out on culture even when collected from the same bowel movement.

Lab number: F000000-0000-0 CSAPx3 Page: 3
Patient: Sample Patient Client: 12345

## Elastase (low)

The level of Elastase is abnormally low in this specimen. Elastase is a pancreatic enzyme that digests and degrades a number of proteins. A finding of low elastase is an indicator of pancreatic exocrine insufficiency. Moderate pancreatic insufficiency is defined at 100-200 ug/g, and severe pancreatic insufficiency as <100 ug/g [1,2].

Fecal Elastase measured by a sensitive immunoassay is a specific marker for pancreatic function [1,3,4] and maintains a high diagnostic accuracy among patients with small intestinal diseases [5]. This Elastase marker allows for the diagnosis or exclusion of pancreatic exocrine insufficiency and degree of severity, which can be caused by chronic pancreatitis, cystic fibrosis, pancreatic tumor, cholelithiasis or diabetes mellitus [6,7,8]. This test does not differentiate between pancreatic insufficiency due to chronic pancreatitis and that due to pancreatic cancer [9]. Immunoreactive elastase concentrations are similar for children and adults [3].

In cases of severe exocrine pancreatic insufficiencies, triglycerides and/or steatocrit may also be elevated. Supplementation with pancreatic enzymes, minerals, and vitamins may be warranted.

- 1. Garcia-Bueno C, Rossi T, Lee K, Yuwono M, Robinson A, Tjota A. Quantification of fecal elastase-1 using either polyclonal or monoclonal antibodies. AGA Abstracts, Gastroenterology 2002:122(4):A-510.
- Dominguez-Munoz J, Hieronymus C, Sauerbruch T, Malfertheiner P. Fecal Elastase: Evaluation of a new noninvasive pancreatic function test. Am J Gastroenterol 1995;90:1834-7.
- 3. Miendje Y, Maisin D, Sipewa MJ, Deprez P, Buts JP, De Nayer P, Philippe M. Polyclonal versus monoclonal ELISA for the determination of fecal elastase 1: diagnostic value in cystic fibrosis and chronic pancreatic insufficiency. Clin Lab 2004;50(7-8):419-424.
- 4. Gullo L, Ventrucci M, Tomassetti P, Migliroi M, Pezzilli R. Fecal elastase1 determination in chronic pancreatitis. Dig Dis Sci. 2000;45(1):166-167.
- Carroccio A, Verghi F, Santini B, Lucidi V, Iacono G, Cavataio F, Soresi M, Ansaldi N,
  Castro M, Montalto G. Diagnostic accuracy of fecal elastase 1 assay in patients with pancreatic
  maldigestion or intestinal malabsorption. Digestive Diseases and Sciences 2001; 46(6):1335-1342,
  2001.
- 6. Glassbrenner B, Schon A, Klatt S, Beckh K, Adler G. Clinical evaluation of the faecal elastase test in the diagnosis and status of chronic pancreatitis. Eur J Gastroenterol Hepatol 1996;8(11):1117-20.
- 7. Deprez P, Del Natale M, Deji Y, Pauwels S, Philippe M. Comparative evaluation of C-mixed triglyceride breath test and faecal elastase1 tests I the assessment of exocrine pancreatic insufficiency. AGA Abstracts, Gastroenterology 2002;122(4):A-510.
- Nunes AC, Pontes JM, Rosa A, Gomes L, Carvalheiro M, Freitas D. Screening for pancreatic exocrine insufficiency in patients with diabetes mellitus. Am J Gastroenterol 2003;98(12):2672-2675.
- 9. Stein J, Jung M, Sziegoleit A, Zeuzem S, Capary W, Lembcke B. Immunoreactive elastase I: clinical evaluation of a new noninvasive test of pancreatic function. Clinical Chemistry 1996;42 (2):222-226.

Lab number: F000000-0000-0 CSAPx3 Page: 4
Patient: Sample Patient Client: 12345

# Secretory IgA (sIgA)

The concentration of sIgA is abnormally low in this specimen. Immunological activity in the gastrointestinal tract can be assessed using secretory immunoglobulin A (sIgA). Secretory IgA is the predominant antibody, or immune protein the body manufactures and releases in external secretions such as saliva, tears, and milk [1]. It is also transported through the epithelial cells that line the intestines out into the lumen. Secretory IgA represents the first line of defense of the GI mucosa and is central to the normal function of the GI tract as an immune barrier [1]. As the principal immunoglobulin isotype present in mucosal secretions, sIgA plays an important role in controlling intestinal milieu which is constantly presented with potentially harmful antigens such as pathogenic bacteria, parasites, yeast, viruses, abnormal cell antigens, and allergenic proteins [1]. Secretory IgA antibodies exert their function by binding to antigenic epitopes on the invading microorganism, limiting their mobility and adhesion to the epithelium of the mucus membrane [2]. This prevents the antigens from reaching systemic circulation and allowing them to be excreted directly in the feces.

Mental and physical stress as well as inadequate nutrition have been associated with low fecal slgA concentrations. This includes dietary restrictions, excessive alcohol intake, body mass loss, negative moods, and anxiety [3]. One study found depressed levels of slgA in malnourished children, particularly protein malnourishment, that responded well to nutritional rehabilitation with a significant increase in slgA [4]. This may be because the synthesis and expression of slgA requires adequate intake of the amino acid L-glutamine [3]. Animal studies have demonstrated that a glutamine-restricted diet can result in a 50% decrease in slgA levels [5]. An increase of dietary L-glutamine can restore GI immune function by protection of cells that synthesize slgA [6]. Saccharomyces boulardii is a nonpathogenic yeast that has been used for the treatment of acute infectious enteritis and antibiotic-associated diarrhea [7]. Significantly elevated levels of slgA and subsequent enhanced host immune response have been found following S. boulardii administration in mice and rats [8,9].

### References:

- 1. Crago SS, Tomasi TB. Mucosal Antibodies, Food Allergy and Intolerance. Bailliere Tindall/W.B. Saunders 1987;167-89.
- 2. Roberts JA. Factors predisposing to urinary tract infections in children. Ped Neph 1996;10:517-522.
- 3. Carins J, Booth C. Salivary immunoglobulin-A as a marker of stress during strenuous physical training. Aviat Space Environ Med 2002;73(12)1203-7.
- 4. Teodosio MR, Oliveira ECM. Urinary secretory IgA after nutritional rehabilitation. Braz J Med Biolog Res 1999;32-421-426
- Alverdy J. Effects of glutamine-supplemented diets on immunology of the gut. J Parent Enteral Nutr 1990;14(4):1095-1135.
- 6. Burke DJ, et al. Glutamine-supplemented total parenternal nutrition improves gut function. Arch Surg 1989;24:2396-2399.
- 7. Alverdy JA. The effect of total parenternal nutrition on gut lamina propria cells. J Parent. Enteral Nutr 1990;14(suppl).
- 8. Qamar A, Aboudola S, Warny M, et al. Saccharomyces boulardii stimulates intestinal immunoglobulin A immune response to clostridium difficile toxin A in mice. Infect Immun 2001;69(4):2762-5.

Lab number: F000000-0000-0 CSAPx3 Page: 5
Patient: Sample Patient Client: 12345

9. Buts JP, Bernasconi P, Vaerman JP, et al. Stimulation of secretory IgA and secretory component of immunoglobulins in small intestine of rats treated with Saccharomyces boulardii. Dig Dis Sci 1990;35(2):251-6.

#### Beneficial Flora

One or more of the expected or beneficial bacteria are low in this specimen. Normally abundant include lactobacilli, bifidobacteria, clostridia, Bacteroides fragilis group, enterococci, and some strains of Escherichia coli. The beneficial flora have many health-protecting effects in the gut, and as a consequence, are crucial to the health of the whole organism. Some of the roles of the beneficial flora include digestion of proteins and carbohydrates, manufacture of vitamins and essential fatty acids, increase in the number of immune system cells, break down of bacterial toxins and the conversion of flavinoids into anti-tumor and anti-inflammatory factors. Lactobacilli, bifidobacteria, clostridia, and enterococci secrete lactic acid as well as other acids including acetate, propionate, butyrate, and valerate. This secretion causes a subsequent decrease in intestinal pH, which is crucial in preventing an enteric proliferation of microbial pathogens, including bacteria and yeast. Many GI pathogens thrive in alkaline environments. Lactobacilli also secrete the antifungal and antimicrobial agents lactocidin, lactobacillin, acidolin, and hydrogen peroxide. The beneficial flora of the GI have thus been found useful in the inhibition of microbial pathogens, prevention and treatment of antibiotic associated diarrhea, prevention of traveler's diarrhea, enhancement of immune function, and inhibition of the proliferation of yeast.

In a healthy balanced state of intestinal flora, the beneficial flora make up a significant proportion of the total microflora. Healthy levels of each of the beneficial bacteria are indicated by either a 3+ or 4+ (0 to 4 scale). However, some individuals have low levels of beneficial bacteria and an overgrowth of nonbeneficial (imbalances) or even pathogenic microorganisms (dysbiosis). Often attributed to the use of antibiotics, individuals with low beneficial bacteria may present with chronic symptoms such as irregular transit time, irritable bowel syndrome, bloating, gas, chronic fatigue, headaches, autoimmune diseases (e.g., rheumatoid arthritis), and sensitivities to a variety of foods. Treatment may include the use of probiotic supplements containing various strains of lactobacillus and bifidobacterium species and consumption of cultured or fermented foods including yogurt, kefir, miso, tempeh and tamari sauce. Polyphenols in green and ginseng tea have been found to increase the numbers of beneficial bacteria. If dysbiosis is present, treatment may also include the removal of pathogenic bacteria, yeast, or parasites.

Percival M. Intestinal Health. Clin Nutr In. 1997;5(5):1-6.

Fuller R. Probiotics in Human Medicine. Gut. 1991;32: 439-442.

Siitonen S, Vapaatalo H, Salminen S, et al. Effect of Lactobacilli GG Yoghurt in Prevention of Antibiotic Associated Diarrhea. Ann Med. 1990; 22:57-59.

Oksanen P, Salminen S, Saxelin M, et al. Prevention of Travelers' Diarrhea by Lactobacillus GG. Ann Med. 1990; 22:53-56.

Lab number: F000000-0000-0 CSAPx3 Page: 6
Patient: Sample Patient Client: 12345

Perdigon G, Alvarez M, et al. The Oral Administration of Lactic Acid Bacteria Increases the Mucosal Intestinal Immunity in Response to Enteropathogens. J Food Prot. 1990;53:404-410.

Valeur, N, et al. Colonization and Immunomodulation by Lactobacillus reuteri ATCC 55730 in the Human Gastrointestinal Tract. Appl Environ. Microbiol. 2004 Feb; 70(2):1176-81.

Elmer G, Surawicz C, and McFarland L. Biotherapeutic agents - a Neglected Modality for the Treatment and Prevention of Intestinal and Vaginal Infections. JAMA. 1996; 275(11):870-876.

Fitzsimmons N and Berry D. Inhibition of Candida albicans by Lactobacillus acidophilus: Evidence for Involvement of a Peroxidase System. Microbio. 1994; 80:125-133

Weisburger JH. Proc Soc Exp Biol Med 1999;220(4):271-5.