Probiotics and their Role in GI Diseases

S PERVEENA, MA AHMEDB

Summary:
Objective: To evaluate role of probiotics in human physiology, metabolism, health, immunity and GI disorders. It is important for gastroenterologists to improve their understanding of the mechanisms of probiotics and the evidence that support their use in clinical practice.

Data Sources: A medline search (1948-December 2014) was conducted using GI diseases and probiotics as terms for identifying pertinent studies. Search limits included English language. Additional information was obtained from bibliographies.

Data Selection And Data Extraction: The information provided is based on review of primary literature from randomized controlled trials (RCTs), meta-analyses, expert consensus panel recommendations and society-based practice recommendations. References are provided for more reading and figure summarizes key information about their mechanism of action.

Data Synthesis: The need for objective, evidence-based guidance on the role of probiotics is becoming increasingly important as public awareness grows. This consensus is intended as a practical reference to help physicians make appropriate, evidence-based recommendations to patients who might benefit from probiotic treatment. Overall, the randomised, placebo-controlled trials included in this article support, with a high evidence level, the therapeutic effects of probiotic agents for several disorders including antibiotic or Clostridium difficile-associated diarrhea, irritable bowel syndrome, and the inflammatory bowel diseases. Although probiotic research is a rapidly evolving field, there are sufficient data to justify a trial of probiotics for treatment or prevention of some of these conditions. However, the capacity of probiotics to modify disease symptoms is likely to be modest and varies among probiotic strains and not all probiotics are right for all diseases. The goal of this review is to provide clinicians with an overview of the rationale and data which support or refute the role of probiotics for treating commonly encountered gastrointestinal disorders.

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Introduction:

Our gut is home to an estimated 100 trillion microorganisms representing more than 1000 different bacterial species, 400 of which are estimated to be of probiotic species1. There is growing recognition of the role of diet and other environmental factors in modulating the composition and metabolic activity of the human gut microbiota, which in turn can impact health. Up to 10% of an individual’s daily energy needs can be derived from the by-products of bacterial fermentation. Gastrointestinal microbiota are also critical for normal immune system development.2 Intestinal bacteria weigh up to 1 kg and bacterial cells outnumber human cells by 10:1. The bacterial genome or DNA may outnumber the human genome by 100:1. The physiologic impact mediated by these resident microbes is important enough to be labeled as “other organ” or “hidden metabolic organ”

Probiotics are defined as ‘live microorganisms’ which when administered in adequate amounts confer a health benefit on the host’.3 There is plentiful evidence that probiotics( pro and biota meaning ‘for life’) impact the microbiome and benefit human health4. Probiotics are distinct from prebiotics (dietary substances such as indigestible oligosaccharides including bananas, whole grains, honey, garlic and onions) that provide a health benefit by selectively promoting the growth of beneficial bacteria in the gut and synbiotics (products containing a synergistic combination of prebiotics and probiotics).

Although research on the microbiome is emerging, scientists have already made tremendous progress in
understanding the microbial makeup and associating microbiome diversity with human physiology, health and disease. It is likely that much of this impact is mediated through diet. What we eat and drink influences the microbiome and the microbiome in turn converts dietary content into biological signals providing enzymatic machinery thus influencing what the human host is able to extract from its diet including energy.5

**Probiotics and the Immune System**

Humans and their trillions of intestinal microorganisms coevolve to form a largely beneficial symbiosis. Human gut microbiome is acquired during the last trimester of pregnancy6. Unlike the human genome, the human microbiome is acquired. Vaginally born babies acquire different microbiomes than babies born by cesarean section7. Gastrointestinal microbiota are critical for normal immune system development8. The intestinal mucosa comprising the largest surface area of the body, is constantly exposed to a vast array of microbes, food antigens, and toxins. The intestinal epithelium must ‘tolerate’ the commensal flora that maintain mucosal homeostasis by controlling inflammatory responses as well as sensing danger signals of potentially harmful pathogens.

Probiotics can down regulate the effects of luminal bacteria in initiating and sustaining an intestinal inflammatory response particularly important in ulcerative colitis. It is becoming increasingly clear that the intestinal microbiota composition, the intestinal barrier and the mucosal immune system plays pivotal roles in the development of a variety of allergic and autoimmune diseases.

**Types of Probiotics**

Most of the probiotics are bacteria, some are fungus (yeast) and some contain mixture. *Lactobacillus acidophilus* is the “friendly” bacteria and most commonly used probiotic. Such healthy bacteria inhabit the intestines and protect against the entrance and proliferation of pathogens. Since the mid 1990s, clinical studies have established that probiotic therapy can help treat several gastrointestinal illnesses.9

*Lactobacillus acidophilus, Lactobacillus casei* both convert lactose into lactic acid helping in lactose intolerance. *Lactobacillus bulgaricus* (discovered by Bulgarian doctor) can be found in many yogurts and soft cheese. It is helpful for those who are lactose intolerant. *L. Acidophilus* may also be helpful in reducing cholesterol levels. *Lactobacillus rhamnosus* GG means the genus is *Lactobacillus*, the species is *rhamnosus* and the strain is GG.10

*Bifidobacteria* is a family of bacteria that has been studied for its ability to prevent and treat various gastrointestinal disorders, including infections, irritable bowel syndrome and constipation. In addition to making lactic acid, it also makes some important short-chain fatty acids that are then absorbed and metabolized by the body. Certain bifidobacteria may actually protect the host from carcinogenic activity of other intestinal flora.

*Saccharomyces boulardii* is the only yeast probiotic. Some studies have shown that it is effective in preventing and treating diarrhea associated with the use of antibiotics and to reduce side effects of *H. pylori* therapy, traveler’s diarrhea, Crohn’s disease.

**Probiotic mixture (VSL#3)** consisting of 8 strains of live freeze-dried lactic acid bacteria. Each sachet contains 450 billion live probiotic bacteria. *Streptococcus thermophilus, Bifidobacterium breve, B longum, B infantis, Lactobacillus acidophilus, L plantarum, L paracasei, L bulgaricus.*

Probiotic therapy may also help people with Crohn’s disease and irritable bowel syndrome. Clinical trial results are mixed, but several small studies suggest that certain probiotics may help maintain remission of ulcerative colitis and prevent relapse of Crohn’s disease and the recurrence of pouchitis (a complication after surgery for ulcerative colitis). Because these disorders are so frustrating to treat, many people are giving probiotics a try even before all the evidence is in for the particular strains. More research is needed to find out which strains work best for what conditions.

**Mechanism of Action of Probiotics**

Probiotics work by several different mechanisms. These commensal microorganisms contribute energy and cellular precursors in the form of short-chain fatty acids, prevent infections and modulate and train the host immune system.12 They act as a barrier by lining the intestinal tract close to the brush border. Through competitive inhibition, they prevent other luminal bacteria stimulating the mucosal immune system. They enhance mucus production so that patients will have a thicker mucus layer, which protects against invasive bacteria. Probiotics influence the mucosal immune system to secrete protective immunoglobulins (Ig) such as secretory IgA and protective defensins and bacteriocins into the lumen (Fig I). Finally, probiotics alter the function of the mucosal immune system to make...
dendritic cells more anti-inflammatory and less pro-inflammatory so that they are slightly less responsive and less reactive to luminal bacteria in initiating and sustaining an intestinal inflammatory response which is particularly important in ulcerative colitis.\textsuperscript{13}

\textbf{Indications of Probiotics}

Although still being studied, probiotics may help several specific illnesses. In 2011, experts at Yale University reviewed the research studies. They concluded that probiotics are best case in the treatment of diarrhea. Controlled trials have shown that \textit{Lactobacillus GG} can shorten the course of infectious diarrhea in infants and children (but not adults). Although studies are limited and data are inconsistent, two large reviews, taken together, suggest that probiotics reduce antibiotic-associated diarrhea by 60\% when compared with a placebo. Probiotics can improve intestinal function and maintain the integrity of the intestinal lining and help fight bacteria that cause diarrhea. In childhood diarrhea in the pediatric population, rotavirus has been the most common cause of infectious diarrhea. Data suggests that the benefit of probiotics in preventing acute infectious diarrhea is modest.\textsuperscript{14}

In treating necrotizing enterocolitis, a type of infection and inflammation mostly seen in infants \textit{Lactobacillus rhamnosus} GG (LGG) is the most effective probiotic reported to date, reducing both severity and duration of diarrhea by \~1 day.\textsuperscript{15} The American Academy of Pediatrics supports the recommendation of LGG early in the course of acute infectious diarrhea to reduce symptom duration.

\begin{figure}[h]
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\caption{Diagrammatic representation of mechanism of action of Probiotics}
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For antibiotic associated diarrhea including those receiving *Helicobacter pylori* eradication therapy and viral gastroenteritis supporting data are strong for probiotics (*L. rhamnosus*GG, *Saccharomyces boulardii*) are among the few treatment modalities available. However, the duration of symptoms in these conditions is typically short regardless of probiotic use. Patients given the higher dose of probiotic concurrent with antibiotics (for 5 days afterward) had fewer occurrences of AAD (15.5 vs. 44.1%). As a secondary endpoint these studies also showed a reduction in development of *C. difficile*-associated diarrhea.

Diarrhea caused by *C. difficile* bacteria is a common nosocomial and community-based medical condition. *C. difficile* sickens half a million Americans and kills more than approximately 14,000 people a year in the United States. Antibiotic therapy with metronidazole, oral vancomycin and now fidaxomicin makeup the current treatment regimen but recurrence remains a clinical problem.

Recently, in patients with recurrent CDI the fecal communities were highly variable in bacterial composition and were characterized by markedly decreased diversity. Preservation and restoration of the microbial diversity could represent novel strategies for prevention and treatment of recurrent CDAD.

Fecal microbiota transplants (FMTs), also known as stool transplants from healthy donors have emerged as an effective means of stopping infections and succeeded in 91% of patients with recurrent CDI. Patients who underwent colonoscopic FMT found it so effective that more than half (53%) said that a fecal microbiota transplant would be their top choice if they contracted CDI again.

Typically, patients receive the fecal transplants by enema, colonoscopy or a nasal tube. As a potentially less costly and less invasive alternative, researchers made fecal pill. Dr. Louie first created a fecal transplant pill when a *C. difficile* infection persisted in one patient who could not tolerate a nasal tube. Currently, it is approved only for recurrent *C. difficile* infections.

After the patients ingested the capsules, the researchers found significantly increased numbers of *Bacteroides*, *C. coccoides*, *C. leptum*, *Prevotella*, *Bifidobacteria*, and *Desulfovibrio* and significantly decreased numbers of *Enterobacteriaceae* and *Veillonella*. They found no *C. difficile* in the intestines.

Symptoms of irritable bowel syndrome (IBS) and other functional GI disorders (FGID) frustrate people with overlapping symptoms and exhaust them with prolonged conventional medications having considerable adverse effects. They can be treated with foods and supplements containing probiotics.

The first practical consensus on the role of probiotics in the management of GI symptoms in adults recommend specific probiotics in the management of some IBS symptoms and can also be used as an adjunct to conventional treatment. For overall symptoms and abdominal pain in IBS, probiotics (*Bifidobacterium bifidum* MIMBb75, *Escherichia coli* DSM17252) has strong supportive evidence for benefit and should be tried. In overall symptoms in IBS-D (*B. longum* subsp. *infantis* 35624), improving the frequency and/or consistency of bowel movements and bloating/distension (*B. animalis* subsp. *lactis* DN-173 010 - *Activia*) probiotics could be tried. In overall symptoms in IBS-C (*B. animalis* - *Activia*) probiotics may be considered. Flatus in IBS has currently no evidence to support use of probiotics. No probiotic alleviates the full range of symptoms in IBS. Though it has shown improvement in some aspects of health-related quality of life there is need for more research. This consensus publication with new research and input from patient groups is intended to be updated in 3 years.

The need for objective, evidence-based guidance on the role of probiotics is becoming increasingly important as public awareness for probiotics grows. This consensus is intended as a practical reference to help physicians make appropriate, evidence-based recommendations to patients who might benefit from probiotic treatment. Overall, the randomised, placebo-controlled trials included with a high evidence level, a role for specific probiotics in the management of overall symptoms and abdominal pain in patients with IBS. Preventing or reducing diarrhoea in patients receiving antibiotics or *H. pylori* eradication triple therapy. The trials support, with a moderate evidence level, a role for specific probiotics in managing overall symptoms in patients with IBS-D; improving bowel movements and bloating/distension in patients with IBS; and improving some aspects of health-related quality of life.

The lack of consensus on the role of probiotics in the management of constipation is consistent with the World Gastroenterology Organisation guideline on
prebiotics and probiotics, which recommends certain prebiotics but not probiotics, for the treatment of constipation.26

Inflammatory bowel disease (IBD) e.g Crohn’s disease, ulcerative colitis and pouchitis (after ileal pouch anal anastomosis in ulcerative colitis patients) are often refractory to standard therapy. The rational to use probiotics and its beneficial efficacy in the treatment of chronic inflammatory bowel disease (IBD) is increasingly scrutinized. The role of probiotics for inducing and maintaining remission especially regarding induction of remission for ulcerative colitis has shown breakthroughs.27

A relationship between immune response and gastrointestinal microbials appears to be involved in the mechanism of ulcerative colitis (UC). Alternative IBD treatment approaches aimed at modifying the composition of the intestinal microbiota in order to overcome gut dysbiosis have become a subject of major interest in recent years. There is also some evidence that probiotics may facilitate and stabilize clinical remission.28

Changes in the relative abundance of the families Enterobacteriaceae and Lachnospiraceae in UC patients provide useful diagnostic indications of clinical response after FMT.29 Randomized, placebo-controlled trials with larger cohorts will be necessary to establish cause–effect relationships for the successfully transmitted donor phylotypes such as F. prausnitzii, B. ovatus, and R. faecis.

Trial results comparing the probiotic (Escherichia coli Nissle1917) to mesalazine have reported equivalent rates of UC relapse. Treatment with Lactobacillus rhamnosus GG strain alone or in combination with mesalazine resulted in a nonsignificant odds ratio decrease for relapse and a significant increase in time to relapse compared to treatment with mesalazine alone. Additionally, bifidobacteria fermented milk supplemented patients had significant reductions in UC exacerbations when compared to nonsupplemented patients. Probiotics were well tolerated, with adverse event rates similar between treatments. One study of 90 volunteers found significantly higher remission rates in people with ulcerative colitis who were given the beneficial bacteria E. coli Nissle. The higher the dose, the longer their remission.30

It is suggested that patients with ulcerative colitis benefit by using bacterial therapies and patients with Crohn’s disease from S. boulardii, an yeast.31

Chronic or recurrent pouchitis is an important complication occurring in ~10–20% of UC patients after ileal anal pouch formation surgery. VSL#3 was shown beneficial in prophylaxis against pouchitis onset after surgical take-down and in maintaining clinical remission after antibiotic induction. Clinical expert guidelines concur that probiotics (VSL#3) can be effective for preventing recurrence of pouchitis.32,33

Diverticulosis is present in approximately two thirds of the elderly population and a large majority of those affected remain entirely asymptomatic. However, an estimated 20% of patients may develop ‘diverticular disease’ with time. Change in the colonic microflora, resulting in a decrease in healthy flora and an increase in pathogenic bacteria, may be detected in patients with diverticular disease. This may allow chronic inflammation and epithelial cell proliferation in and around the diverticula.34 Probiotics, restoring the colonic microenvironment, have been proposed to treat those patients.35

The most recently published RCT in Diverticulosis compared mesalazine or mesalazine plus Bifidobacterium infantis 35624 for 12 weeks and followed up for nine additional months. Global symptom scores were assessed over the 52 weeks of follow-up. Mesalamine demonstrated a consistent trend in reducing symptoms. Addition of probiotic did not increase mesalamine efficacy.36

Another Multicentre Double-blind, Randomised, Placebo-Controlled Study showed both cyclic mesalazine and L. casei subsp. DG 24 billion/day appear to be better than placebo for maintaining remission in symptomatic uncomplicated diverticular disease (SUDD) especially when used in combination. Moreover, both treatments alone or in combination are significantly better than placebo in preventing occurrence of acute diverticulitis in SUDD patients.37

Alcoholic liver disease is characterized by fatty liver (steatosis), which may progress to alcoholic hepatitis, fibrosis, and cirrhosis.38 Intestinal bacterial overgrowth is common in patients with alcoholic liver disease. In alcoholics, translocation of bacteria and bacterial products into the circulation contributes to liver disease.
Treatment with prebiotics partially restored gut microbiome, reduced bacterial overgrowth and lessened alcoholic steatohepatitis. Disbiosis (disruption or dysregulation of intestinal antimicrobial molecules) contributed to changes in the enteric microbiome leading to alcoholic steatohepatitis in mice.39

Role in Metabolic disease or obesity shows comparisons of the distal gut microbiota of genetically obese and lean human volunteers revealing that obesity is associated with changes in the relative abundance of the two dominant bacterial divisions, the Bacteroidetes and the Firmicutes. Obese microbiome has an increased capacity to harvest energy from the diet.40 Gut microbiota is an additional contributing factor to the pathophysiology of obesity.41 Studies in mice have associated the phylum Firmicutes with obesity and the phylum Bacteroidetes with weight loss.42

There is clear evidence that the intestinal microbiota influences the host through its effect on body weight, bile acid metabolism, proinflammatory activity and insulin resistance, and modulation of gut hormones.43 A synergism between role of microflora, aberrant intestinal microbiota (either quantitatively or qualitatively), a “leaky” gut mucosal barrier and altered mucosal immunity contributing to type 1 diabetes has begun to evolve. Changes in gut microbiota and thus cell wall components are involved in the epigenetic regulation of inflammatory reactions. An improved diet targeted to induce gut microbial balance and epigenetic changes of pro-inflammatory genes may be effective in the prevention of metabolic syndrome.44

**Precautions about Probiotics**

Probiotics preparations are generally considered as food not drug. They are already present in a normal digestive system and therefore safe and may cause few side effects. In the USA, formulations of probiotics are medical foods and must be used under medical supervision. In a systematic review of studies, rate of side effects was same in probiotics users as in those taking mesalazine. No serious side effects have been associated with prolonged use of probiotics in ulcerative colitis.45

Probiotics have enjoyed an impeccable safety reputation but immune compromise (including a debilitated state or malignancy) has been identified as a risk factor for rare cases of bacteremia or fungaemia in patients taking certain probiotics (most commonly *Saccharomyces boulardii*).46 No serious adverse events attributed to FMT were observed.47

One high profile multicenter placebo controlled Dutch RCT examining probiotic supplementation in severe acute pancreatitis found a higher incidence of mesenteric ischemia and death in the treatment group.48 This is the only trial supporting the concept that probiotics should be avoided in critically ill patients.

Probiotic ingredients should be clearly marked on the label. There’s no way to judge the safety of unidentified mixtures. Probiotics are food particles and they are rapidly washed away within days, although strain-specific differences occur.49 For a chronic GI problem, it is critically important that the product is taken in adequate doses on a regular basis (e.g. just before a meal) for a reasonable period of time, which should be at least a month, unless it cannot be tolerated for any reason. In order to get the full benefits of probiotics, dose selection should be based on available evidence and manufacturers’ recommendations. Some yogurts contain the friendly bacteria but as they are sensitive to oxygen, light and dramatic temperature changes, when heat-treated or pasteurized, they lose these valuable “live and active cultures.”

Despite their long history, wide availability and substantial publication record, the clinical role of probiotics is in general, inadequately characterised and remains ill-defined. Probiotic research are complicated by the wide variety of probiotic strains obtained with one strain not applicable to others. According to age and health status of the target group effects of probiotics may vary.

The variable range of formulations (capsules, sachets, yogurts and fermented milks or fruit drinks), differ in dose and the presence of supporting substrates add further sources of variation.50

**Conclusion:**

Despite widespread use of natural therapies by patients allured by advertisements, health care providers may be unfamiliar with probiotics as a treatment modality. Many patients consume probiotic products in attempts to manipulate the intestinal microbiota for health benefit. Questions regarding optimal probiotic, dosing, specific patient populations and placement in therapy are to be answered by large, randomized,
controlled trials conducted before probiotics can be routinely recommended. Evidence supports a role for considering the recommendation of conventional probiotics for some clinical conditions. Probiotic strain selection should focus on quality tested products with clinically demonstrated benefit for the given disorder. Patients and physicians should expect modest effects and consider using probiotics as a supplement rather than a replacement for conventional therapy.\textsuperscript{[51]}

Probiotic research is evolving rapidly and this article reflect physicians' rather than patients' perspectives. Future clinicians will have the opportunity to use directed selection of a probiotic or probiotic derived product to specifically address a unique disease causing physiologic or genetic defect. Though challenges exist, ongoing investigations offer great promise for the future.

Some patients have interest in probiotics and their potential and they may take probiotics (or products incorrectly identified as probiotics) to reduce their symptoms before consulting their physician. Therefore, educational materials for the general public are also needed to improve their understanding and to ensure appropriate use of probiotics.

References:
19. Dr. Mellow, Dr. Schiller, and Dr. Kenneley. American College of Gastroenterology (ACG) 2011 Annual Scientific Meeting and Postgraduate Course; President’s Plenary Session II. Presented 2011.
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S Perveen & MA Ahmed


