THE LINK BETWEEN GUT BACTERIA AND P-GP

New research may explain the role that gut bacteria play in inflammatory bowel disease. Researchers found that people with a healthy balance of gut bacteria also have healthy levels of a protein called P-gp. But people who have inflammatory bowel diseases have lower levels of this important protein. That matters because P-gp cuts inflammation in the gut and protects the walls of the intestines. Without enough P-gp, gut inflammation can get out of control. Understanding the connection between gut bacteria and this important protein can lead to new treatments for IBD or dietary recommendations to improve the makeup of the gut bacteria.

SOURCE: Microbiome

6.8 MILLION
Estimated number of IBD cases worldwide.

SOURCE: The Lancet

COULD A NEW PILL PREVENT INFLAMMATION?

Researchers at Harvard’s Brigham and Women’s Hospital have souped up a healthy microbe in the hope that it will treat inflammatory bowel disease. They took the same type of yeast that’s in beer and genetically reengineered it to detect inflammation in the gut. Aptly named a “Yeast Bot,” the idea is that when the pill reaches the gut, it will find inflammation, and emit an anti-inflammatory substance to stop the process and repair the damage. But before you can get a prescription for this newfangled drug, the researchers must put it through clinical trials. Stay tuned.

SOURCE: Nature Medicine

GET STRESS IN CHECK

Does stress trigger your Crohn’s disease flares? Researchers may have discovered why. In a small laboratory study, they found that hormones the body releases when under stress weakened the part of the immune system whose job is to protect the gut from harmful bacteria. This can make the gut more prone to infection with Enterobacteriaceae, a group of bacteria that include E. coli and has links to Crohn’s flares. If unchecked stress is causing you frequent flares, talk to your doctor about how to manage it. Exercise, proper sleep, meditation, and “me” time may help.

SOURCE: Nature Communications

2X
Smokers’ risk of developing Crohn’s disease compared with nonsmokers’ risk.

SOURCE: Crohn’s & Colitis Foundation of America
STATS & FACTS

By Sonya Collins
Reviewed by Neha Pathak, MD, WebMD Lead Medical Editor

3 million
Estimated number of Americans living with IBD.

70,000
Estimated number of new cases of IBD in the U.S. every year.

80,000
Estimated number of children with IBD in the U.S.

36%
Amount of people who develop IBD when both their parents have it.

15 TO 35
Ages at which most people with IBD get a diagnosis.

1 in 20
Number of IBD cases that occur in children.

2X
How much more often Crohn’s disease appears in children compared with ulcerative colitis.

1 IN 5
Number of people with IBD who have a next of kin with the condition.

UP TO 1 IN 5

SOURCES: CDC, Crohn’s & Colitis Foundation of America

BY THE NUMBERS
YOU WEREN’T MADE FOR MODERATE TO SEVERE ULCERATIVE COLITIS OR CROHN’S DISEASE.

BUT ENTYVIO IS.

ENTYVIO has helped many people achieve long-term relief and remission. It may help you too. At your next doctor visit, make sure to:

• Bring up ALL symptoms that are bothering you (don’t worry, they’ve heard it all!)
• Ask how ENTYVIO works
• Ask if ENTYVIO is right for you

Important Safety Information for ENTYVIO® (vedolizumab)

Do not receive ENTYVIO if you have had an allergic reaction to ENTYVIO or any of its ingredients.

• ENTYVIO may cause serious side effects, including:
  o Infusion-related and serious allergic reactions can happen while you are receiving ENTYVIO or several hours after treatment. You may need treatment if you have an allergic reaction. Tell your healthcare provider or get immediate medical help if you get any of these symptoms during or after an infusion of ENTYVIO: rash, itching, swelling of your lips, tongue, throat or face, shortness of breath or trouble breathing, wheezing, dizziness, feeling hot, or palpitations (feel like your heart is racing).
  o ENTYVIO may increase your risk of getting a serious infection. Before receiving and during treatment with ENTYVIO, tell your healthcare provider if you think you have an infection or symptoms of an infection, such as fever, chills, muscle aches, cough, shortness of breath, runny nose, sore throat, red or painful skin or sores on your body, tiredness, or pain during urination.
  o People with weakened immune systems can get progressive multifocal leukoencephalopathy (PML) (a rare, serious brain infection caused by a virus). Although unlikely while receiving ENTYVIO, a risk of PML cannot be ruled out. PML can result in death or severe disability. There is no known treatment, prevention, or cure for PML. Tell your healthcare provider right away if you have any of the following symptoms: confusion or problems thinking, loss of balance, change in the way you walk or talk, decreased strength or weakness on one side of the body, blurred vision, or loss of vision.
  o Liver problems can happen in people who receive ENTYVIO. Tell your healthcare provider right away if you have any of the following symptoms: tiredness, loss of appetite, pain on the right side of your abdomen, dark urine, or yellowing of the skin and eyes (jaundice).
  o The most common side effects of ENTYVIO include common cold, headache, joint pain, nausea, fever, infections of the nose and throat, tiredness, cough, bronchitis, flu, back pain, rash, itching, sinus infection, throat pain, and pain in extremities. These are not all the possible side effects of ENTYVIO.

• Before receiving ENTYVIO, tell your healthcare provider about all of your medical conditions, including if you have or think you may have an infection or have infections that keep coming back; have liver problems; have tuberculosis (TB) or have been in close contact with someone with TB; have been in close contact with someone with HIV; or have liver problems; or have certain blood cell count problems. You may need to get extra or extra tests before starting ENTYVIO.

• Before receiving ENTYVIO, tell your healthcare provider if you are pregnant, breastfeeding, plan to become pregnant, or plan to breastfeed. You may need to discontinue ENTYVIO before pregnancy.

• Before receiving ENTYVIO, tell your healthcare provider if you have had any vaccinations recently received or are scheduled to receive a vaccine; or if you are pregnant, breastfeeding, plan to become pregnant, or plan to breastfeed.

Please see the Important Facts for ENTYVIO on the following page and talk with your healthcare provider.

You are encouraged to report negative side effects of prescription drugs to the FDA. Visit www.fda.gov/medwatch or call 1-800-FDA-1088.

Uses of ENTYVIO® (vedolizumab) ENTYVIO is a prescription medicine used in adults for the treatment of:

• moderately to severely active ulcerative colitis
• moderately to severely active Crohn’s disease

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ENTYVIO may cause serious side effects, including:

- **Infusion-related and serious allergic reactions.** These reactions can happen while you are receiving ENTYVIO or several hours after treatment. You may need treatment if you have an allergic reaction. Tell your healthcare provider or get medical help right away if you get any of these symptoms during or after an infusion of ENTYVIO: rash, itching, swelling of your lips, tongue, throat or face, shortness of breath or trouble breathing, wheezing, dizziness, feeling hot, or palpitations (feel like your heart is racing).

- **Infections.** ENTYVIO may increase your risk of getting a serious infection. Before receiving ENTYVIO and during treatment with ENTYVIO, tell your healthcare provider if you think you have an infection or have symptoms of an infection such as fever, chills, muscle aches, cough, shortness of breath, runny nose, sore throat, red or painful skin or sores on your body, tiredness, or pain during urination.

- **Progressive Multifocal Leukoencephalopathy (PML).** People with weakened immune systems can get progressive multifocal leukoencephalopathy (PML) (a rare, serious brain infection caused by a virus). Although unlikely while receiving ENTYVIO, a risk of PML cannot be ruled out. PML can result in death or severe disability. There is no known treatment, prevention, or cure for PML. Tell your healthcare provider right away if you have any of the following symptoms: confusion or problems thinking, loss of balance, change in the way you walk or talk, decreased strength or weakness on one side of the body, blurred vision, or loss of vision.

- **Liver Problems.** Liver problems can happen in people who receive ENTYVIO. Tell your healthcare provider right away if you have any of the following symptoms: tiredness, loss of appetite, pain on the right side of your stomach (abdomen), dark urine, or yellowing of the skin and eyes (jaundice). See “What are the possible side effects of ENTYVIO?” for more information about side effects.

What is ENTYVIO?

ENTYVIO is a prescription medicine used in adults for the treatment of:

- moderately to severely active ulcerative colitis.
- moderately to severely active Crohn’s disease.

It is not known if ENTYVIO is safe and effective in children under 18 years of age.

Who should not receive ENTYVIO?

Do not receive ENTYVIO if you have had an allergic reaction to ENTYVIO or any of the ingredients in ENTYVIO. See the end of this Medication Guide for a complete list of ingredients in ENTYVIO.

Before receiving ENTYVIO, tell your healthcare provider about all of your medical conditions, including if you:

- have an infection, think you may have an infection or have infections that keep coming back (see “What is the most important information I should know about ENTYVIO?”).
- have liver problems.
- have tuberculosis (TB) or have been in close contact with someone with TB.
- have recently received or are scheduled to receive a vaccine. Talk to your healthcare provider about bringing your vaccines up-to-date before starting treatment with ENTYVIO.
- are pregnant or plan to become pregnant. It is not known if ENTYVIO will harm your unborn baby. Tell your healthcare provider right away if you become pregnant while receiving ENTYVIO.
- are breastfeeding or plan to breastfeed. ENTYVIO passes into your breast milk. Talk to your healthcare provider about the best way to feed your baby if you take ENTYVIO.

Tell your healthcare provider about all the medicines you take, including prescription and over-the-counter medicines, vitamins and herbal supplements. Especially tell your healthcare provider if you take or have recently taken Tysabri (natalizumab), a Tumor Necrosis Factor (TNF) blocker medicine, a medicine that weakens your immune system (immunosuppressant), or corticosteroid medicine.

How will I receive ENTYVIO?

- ENTYVIO is given through a needle placed in a vein (intravenous infusion) in your arm.
- ENTYVIO is given to you over a period of about 30 minutes.
- Your healthcare provider will monitor you during and after the ENTYVIO infusion for side effects to see if you have a reaction to the treatment.

What are the possible side effects of ENTYVIO?

ENTYVIO may cause serious side effects, see “What is the most important information I should know about ENTYVIO?”

The most common side effects of ENTYVIO include: common cold, headache, joint pain, nausea, fever, infections of the nose and throat, tiredness, cough, bronchitis, flu, back pain, rash, itching, sinus infection, throat pain, and pain in extremities. These are not all of the possible side effects of ENTYVIO. Call your doctor for medical advice about side effects. You may report side effects to FDA at 1-800-FDA-1088.

General information about ENTYVIO

Medicines are sometimes prescribed for purposes other than those listed in a Medication Guide. You can ask your pharmacist or healthcare provider for information about ENTYVIO that is written for health professionals.

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For more information, go to www.ENTYVIO.com or call 1-877-TAKEDA-7 (1-877-825-3327).
When you have inflammatory bowel disease (IBD), it’s natural to struggle with mental health and tough emotions. Fatigue, pain, and frequent trips to the bathroom can get in the way of doing what you love. But, there’s more to the link between IBD and mental health than this, according to Charles Bernstein, MD, a gastroenterologist and director of the IBD Clinical and Research Centre at the University of Manitoba in Winnipeg, Canada.

“People with IBD are more likely to have mood disorders or depression or generalized anxiety disorders for years prior to the IBD,” Bernstein says.

In addition, the symptoms of IBD and daily challenges of having any chronic disease add stress to one’s life. You also may take steroids to help with the inflammation that comes with IBD.

“Steroids are one medication we know can have an adverse affect on emotional state,” Bernstein says.

**MANAGE YOUR MENTAL HEALTH**

Bernstein says it’s not clear that stress actually makes your gut inflamed. But he has noticed that people feeling under stress more often do have symptoms flare up. Mental health challenges may make it harder for you to stick to your medications, too.

He says that your IBD doctor should ask you not just about IBD symptoms, but about how you are feeling in other ways, too. Whether your doctor asks you or not, let them know if you are having trouble, even if you don’t think it’s related to your IBD. Ask if they have ways to help or can refer you to someone who can.

Online resources might help you, too. In fact, Bernstein is studying whether an online program that teaches how to manage stress can help in dealing with IBD symptoms.

**TRY SOME ALTERNATIVES**

“There are many things people can do,” Bernstein says. “In general, mindfulness techniques, yoga, meditation, physical exercise—these are all things I would encourage.”

You might not feel up to boot camp or some other intensive exercise regimen and that’s OK. Try long walks, cycling, or tai chi—anything that works for you to relieve stress and get active, Bernstein says. Think about what your stressors are and who can help. Those stressors “may be work or financial or may be related to other health problems,” he adds.

Bernstein says he has at times referred his IBD patients to a social worker. To manage your IBD, it’s important to take an open and holistic approach. Find a doctor who is willing to consider everything that’s going on in your life. The first step for you, Bernstein says, is having some awareness that your “mental health may affect how you feel from a GI perspective.”
I was 37 when I started having digestive pain that just kept getting worse. I was having mucus in my stool, and lost so much blood that I developed anemia. For a cyclist, anemia isn’t a good thing; you want your blood count to be high so your body has the oxygen it needs. But I was constantly fatigued.

Still, I avoided the doctor because I dreaded procedures like colonoscopies. I’m also a little stubborn and tend to wait until things are really bad before addressing them. Finally, when I couldn’t ignore things anymore, I told my wife, who’s a medical student, what was going on. She immediately told me to get my butt to the doctor. (No pun intended.)

THE WINDING ROAD TO DIAGNOSIS AND TREATMENT

But even once I was under medical care, it took a while to get a diagnosis. My general practitioner ran tests, but couldn’t pinpoint what was going on. I just thought this was part of getting older. My best guess was that I had hemorrhoids. Finally, I saw a gastrointestinal therapist, and a year after my symptoms began, I got my ulcerative colitis diagnosis, and a whole new chapter of my life started.

The doctors tried steroids and other first-line treatments, but ultimately, biologics are what I needed to get my UC under control. It took about 6 months, but we found a biologic that got me into remission, which is incredible. I still have symptoms from time to time, so I try to stay aware of what might trigger a flare. Usually for me it’s stress.

DAILY LIFE WITH UC

Even through my worst symptoms, I tried to keep training and racing and parenting just like I always had. I didn’t want to let UC disrupt the enjoyment of my life. But there were times I had to stop—literally—and deal with the reality of my disease. Road trips required a lot more bathroom breaks, for example. It’s kind of a joke in the IBD community that we know where all the bathrooms are.

As an athlete, “pushing through” is what I know to do, but the physical toll of UC was really wearing me down and making that tactic unsustainable. Mentally, I was suffering, too. I rely on the physicality of training as an outlet for my mental health. So to lose that was really hard. I was fortunate to find a lifeline in Ride4IBD—a movement that combines my love of cycling and a passion for IBD awareness.

Being diagnosed with UC brought out the fighter spirit in me and gave me a deeper appreciation of life and health. I no longer take these things for granted. In that respect, it really has changed me for the better.
TEST YOURSELF

How much do you know about IBD?

By Kendall K. Morgan
Reviewed by Brunilda Nazario, MD, WebMD Chief Physician Editor, Medical Affairs

Looking for some new ways to take control of your IBD? Take this quiz to find out how much you already know about IBD and ways you might manage it even better.

1. Your diet might have caused your IBD and the foods you eat now can make it go away.
   - True
   - False

2. Diarrhea, pain, and fatigue are the hardest parts of having IBD.
   - True
   - False

3. You should watch for trouble in your joints, skin, eyes, and many internal organs.
   - True
   - False

4. Therapy can help you with your belly pain.
   - True
   - False

5. You can put your smartphone to work for your IBD.
   - True
   - False

1. **False.** It’s tempting to think that your diet somehow caused your IBD and that eating “better” can make it go away. Watching what you eat can help with symptoms, but the reason you have IBD has nothing to do with your eating habits, now or in the past. You can’t cure IBD with your diet either. Even so, it’s a good idea to pay attention to how different foods make you feel. You might find that bananas, bread, rice, cheese, crackers, and other simple or bland foods help. You might not do as well with food or drinks with lots of caffeine, fiber, or spices. But people are different and there aren’t any set rules. Keeping a journal that tracks what you eat and how you feel is a good way to get a handle on eating habits that work for you.

2. **False.** Of course, what you find hardest about IBD might differ from someone else. But, for many, the toughest parts aren’t the physical aspects so much as the mental ones. A flare may make you feel that you’ve lost control of your body and your life. Some say it’s like living in a different world from other people who don’t have IBD. When you’re flaring, you might have trouble seeing yourself as the person you really are. People with IBD are more likely than the average person to have depression and anxiety. To help you through the hard times, take control of the things that you can: what you eat, when you sleep, and how you’re managing stress. Try taking a walk or meditating. Ask your friends, family, and health care team for the support you need and remember that your IBD doesn’t define you.

3. **True.** IBD primarily describes conditions with chronic inflammation in your intestines or gut. But the inflammation that comes with IBD can affect many other parts of your body, too. These symptoms of IBD are called “extraintestinal manifestations,” or EIMs. Actually, many people with IBD notice symptoms in other parts of the body even before the gut problems. If you have IBD and pain or other signs of inflammation somewhere else, ask your doctor if it’s related and make sure you’re getting the help you need for those EIMs, too.

4. **True.** Many people with IBD have belly pain, even when the IBD is under good control or in remission. You might think the way to help with pain is medicine or your diet. While those certainly might help and you should let your doctor know how you’re feeling, it turns out that therapy often helps, too. Multiple studies looking at one-on-one or group therapy interventions found that they helped with belly pain in people with IBD. It goes to show again just how important the mental aspects of IBD can be. If you want to try it out, ask your doctor if they can recommend a therapist, counselor, or stress management program.

5. **True.** When you have IBD, it’s a good idea to track your symptoms and help catch a flare quickly, before it gets worse. You can do this with pencil and paper, of course. The Crohn’s and Colitis Foundation has an easy-to-use form for tracking the many ways that IBD may be affecting you. But if you use a smartphone, search “IBD” in the app store. You’ll find multiple apps you can use to track your mood, food, sleep, stress, symptoms, and more. All that data can help you make note of any changes over time and work with your doctor to take control of your IBD.
BEYOND THE GI SYSTEM

SYMPTOMS OF IBD THAT AFFECT OTHER PARTS OF THE BODY

By Kendall K. Morgan
Reviewed by Neha Pathak, MD, WebMD Lead Medical Editor

The inflammation caused by IBD comes with plenty of gastrointestinal (GI) trouble. But at least 1 in 4 people with IBD have signs in other parts of the body. Doctors sometimes call these “extraintestinal manifestations” of IBD.

ARTHRITIS
“The predominant one is joint pain,” says Charles Bernstein, MD, gastroenterologist at the University of Manitoba in Canada. “About a quarter of people with IBD also have redness and other signs of arthritis.”

People with IBD may have arthritis symptoms at a young age in large joints of the arms and legs. They also may have symptoms in various parts of the spine, although this is less common. Often, the joint pain gets better as GI symptoms improve with treatment.

SKIN CONDITIONS
About 1 in 5 people with IBD have skin conditions, either related to the IBD or its treatment. Some common ones include erythema nodosum, in which tender red bumps appear on the shins, and pyoderma gangrenosum, in which lesions or ulcers form often where there’s an injury.

Bernstein says that the red bumps typically show up when IBD is most active. The lesions of pyoderma gangrenosum can arise separately from IBD flares. They’re often related to ostomy surgery, which allows waste to leave your body through a pouch, or other trauma. IBD also can come with painful tears around the anus and small ulcers called canker sores in your mouth.

EYES, BONES, AND LIVER
Approximately 1 in 10 people with IBD have problems with their eyes. These may involve redness and pain in different parts of the eye. While it can change your vision, these eye problems in IBD are treatable with eye drops or other medicines.

Up to 60% of people with IBD have thin and weakened bones, too. As a result, Bernstein says they are more at risk for osteoporosis. This can happen from using steroids for a long time or from the inflammation itself. While relatively rare, about 5% of people with IBD have serious liver disease.

WHICH COMES FIRST?
Bernstein says that these other conditions may come up before, after, or around the same time as a diagnosis of IBD. While it’s good to know these can happen, he says he doesn’t want folks with IBD to worry about every minor discomfort.

“Aches and pains can be normal,” he says, and all people feel them. “It’s good to be aware of it, but just because you have red eyes one day doesn’t mean you’ve got iritis [inflammation of the eye’s iris]. I try not to make my patients too vigilant about short-lived symptoms.”

QUESTIONS TO ASK YOUR DOCTOR

+ Could other symptoms I have been linked to my IBD?
+ Are there other medicines to take for them?
+ What can I do to prevent bone loss?
+ I feel tired a lot. Could it be low iron (anemia)?
OPTIONS TO TREAT CROHN’S DISEASE

TALK TO YOUR DOCTOR ABOUT WHICH ONE IS BEST FOR YOU

By Kendall K. Morgan
Reviewed by Neha Pathak, MD,
WebMD Lead Medical Editor

There’s no cure for Crohn’s, but there are ways to control it. The goal is to get your Crohn’s disease into remission and keep it there.

You and your doctor will have many medicines to choose from. They fall into different categories based on how they work to control the disease and its symptoms. Najwa El-Nachef, MD, a gastroenterologist at the University of California at San Francisco, says that the best place for you and your doctor to start will depend on how severe your disease and inflammation is.

AMINOSALICYLATES
Aminosalicylates, which lower inflammation, can be an early step in treating Crohn’s disease. There are several to choose from. They can be applied to the surface of your intestine with an enema or suppository or you can take them by mouth.

“If someone has very mild disease, you may start with these more topical therapies,” El-Nachef says. But, she adds, they less often work well for Crohn’s compared with other types of inflammatory bowel disease. Doctors used them more in the past.

STEROIDS
Corticosteroids can fight inflammation in your gut. But steroids aren’t a long-term solution. That’s both because they aren’t the best way to keep Crohn’s in remission and they have side effects if taken too long.

“They’re not considered good long-term agents.”

IMMUNE SUPPRESSORS
You might also hear these drugs, which are taken by mouth, called “immune modulators.” There are several of them, all of which work to lower the activity in your immune system that’s causing the problems in your gut. But, El-Nachef says, these treatments also are falling out of favor for treating Crohn’s as safer and more effective treatment options have grown.

“There’s an idea out there that a pill is safer than an infusion or injection, but that’s not always the case,” El-Nachef says.

BIOLOGICS
You’ll get biologic therapies as an infusion or injection once every several weeks. They’re antibodies that target the immune system in one of a few different ways. Your doctor will help you decide which to try first.

El-Nachef says any of today’s biologics “are reasonable first-line treatments” for Crohn’s. They work better than earlier medicines and come with fewer side effects.

DIET AND SURGERY
In addition to biologics or other medicines, your doctor may suggest changes to your diet or letting your bowel rest. Sometimes people with Crohn’s also need surgery, but this is becoming less common thanks to treatment advances.

“[Biologic] medications have changed the whole landscape of what it means to have Crohn’s,” El-Nachef says. “People have less surgery now and, if they do, they are less likely to have another surgery [later].”
Are you one of them?
The Crohn’s & Colitis Foundation is here for you.

Whether you were diagnosed while in service or after you were discharged, you may have questions about your disease, need resources to navigate your healthcare options, or want to connect with others who understand what you are experiencing.

Visit www.crohnscolitisfoundation.org/veterans to find resources just for you or contact our IBD Help Center at info@crohnscolitisfoundation.org or 888-MY-GUT-PAIN to get answers to your questions.

Over 66,000 veterans in the United States live with inflammatory bowel disease (IBD).