Sick from Your Stomach: Gut Bacteria May Trigger Diseases such as Rheumatoid Arthritis

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ROCHESTER, Minn. — The billions of bugs in our guts have a newfound role: regulating the immune system and related autoimmune diseases such as rheumatoid arthritis, according to researchers at Mayo Clinic and the University of Illinois at Urbana-Champaign.

Larger-than-normal populations of specific gut bacteria may trigger the development of diseases like rheumatoid arthritis and possibly fuel disease progression in people genetically predisposed to this crippling and confounding condition, say the researchers, who are participating in the Mayo Illinois Alliance for Technology Based Healthcare.

The study is published in the April 2012 issue of PloS ONE.

"A lot of people suspected that gut flora played a role in rheumatoid arthritis, but no one had been able to prove it because they couldn't say which came first — the bacteria or the genes," says senior author Veena Taneja, Ph.D., a Mayo Clinic immunologist. "Using genomic sequencing technologies, we have been able to show the gut microbiome may be used as a biomarker for predisposition."

The roughly 10 trillion cells that make up the human body have neighbors: mostly bacteria that often help, training the immune system and aiding in digestion, for example. The bacteria in the intestines, in addition to a relatively small number of other microorganisms (the gut microbiome), outnumber human cells 10-to-1.

Researchers found that hormones and changes related to aging may further modulate the gut immune system and exacerbate inflammatory conditions in genetically susceptible individuals.

Nearly 1 percent of the world's population has rheumatoid arthritis, a disease in which the immune system attacks tissues, inflaming joints and sometimes leading to deadly complications such as heart disease. Other diseases with suspected gut bacterial ties include type 1 diabetes and multiple sclerosis.
Researchers with the Mayo Illinois Alliance for Technology Based Healthcare say that identifying new biomarkers in intestinal microbial populations and maintaining a balance in gut bacteria could help physicians stop rheumatoid arthritis before it starts.

"This study is an important advance in our understanding of the immune system disturbances associated with rheumatoid arthritis. While we do not yet know what the causes of this disease are, this study provides important insights into the immune system and its relationship to bacteria of the gut, and how these factors may affect people with genetic susceptibilities to disease," says Eric Matteson, M.D., chairman of rheumatology at Mayo Clinic, who was not a study author.

Dr. Taneja and her team genetically engineered mice with the human gene HLA-DRB1*0401, a strong indicator of predisposition to rheumatoid arthritis. A set of control mice were engineered with a different variant of the DRB1 gene, known to promote resistance to rheumatoid arthritis. Researchers used these mice to compare their immune responses to different bacteria and the effect on rheumatoid arthritis.

"The gut is the largest immune organ in the body," says co-author Bryan White, Ph.D., director of the University of Illinois' Microbiome Program in the Division of Biomedical Sciences and a member of the Institute for Genomic Biology. "Because it's presented with multiple insults daily through the introduction of new bacteria, food sources and foreign antigens, the gut is continually teasing out what's good and bad."

The gut has several ways to do this, including the mucosal barrier that prevents organisms — even commensal or "good" bacteria — from crossing the lumen of the gut into the human body. However, when commensal bacteria breach this barrier, they can trigger autoimmune responses. The body recognizes them as out of place, and in some way this triggers the body to attack itself, he says.

These mice mimic human gender trends in rheumatoid arthritis, in that females were about three times as likely to generate autoimmune responses and contract the disease. Researchers believe these "humanized" mice could shed light on why women and other demographic groups are more vulnerable to autoimmune disorders and help guide development of new future therapies.

"The next step for us is to show if bugs in the gut can be manipulated to change the course of disease," Dr. Taneja says.

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