

Researchers find stomach cancer pathway doesn't work as previously thought

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Scientists had thought that destroying acid-secreting cells in the stomach would lead to a precancerous condition in other stomach cells. In the stomachs of mice, cells on the right (in yellow) are undergoing a process that can lead to cancer. But on the left, where researchers selectively had destroyed acid-secreting cells, the precancerous changes did not happen. Credit: Mills Laboratory



Conventional wisdom holds that the loss of cells that secrete acid in the stomach leads to a condition that eventually can develop into stomach cancer.

But new research at Washington University School of Medicine and Siteman Cancer Center at Barnes-Jewish Hospital and Washington University in St. Louis indicates otherwise. Researchers found that damage to acid-secreting <u>cells</u> alone doesn't jump-start the transformation of <u>healthy cells</u> into precancerous cells—at least in a <u>mouse model</u>.

Their research is published online in the journal Gastroenterology.

"We believe it's easier to stop cancer from starting than to treat it after it occurs, so our goal has been to find ways to stop this cascade before it begins," said first author and doctoral candidate Joseph Burclaff. "But the first steps in the process that leads to cancer, however, are different than what we had assumed, so to prevent cancer, we'll need to identify the real culprit."

Burclaff works in the laboratory of Jason C. Mills, MD, PhD, a professor of medicine in the Division of Gastroenterology. Like other scientists in the field, they had concluded long ago that damage to acid-secreting cells in the stomach leads directly to a precancerous condition called metaplasia. But studying a mouse model of that condition, the researchers learned that the <u>conventional wisdom</u> was wrong.

"We thought we had this process figured out, but when we selectively destroyed only these acid-secreting cells in the stomachs of mice, the animals didn't develop metaplasia," Mills said.

The precancerous process causes cells in adult organs to change in response to inflammation or injury. Those changes may help cells



respond more effectively to particular types of injury. The problem, said Mills, who also is a professor of developmental biology and of pathology and immunology, is that when the process continues over many years, it greatly increases cancer risk.

The two main causes of the precancerous condition are bacterial infections—particularly with H pylori, the microbe associated with stomach ulcers—and inflammation caused by the body's own immune system.

In this study, Burclaff and Mills inserted a human gene into acidsecreting cells in the stomachs of mice. Then, by exposing the animals to a toxin that affects <u>human cells</u> but not mouse cells, they were able to kill the acid-secreting cells without damaging any other cells in the animals. But, surprisingly, the mice did not go on to develop the precancerous condition.

"We had thought the dying cells might signal other cells, sort of like a 911 call," Burclaff said. "If there was such a signal, we could work to block it before it led to cancer. But these experiments demonstrate that if such signals exist, they must be coming from someplace else."

Burclaff and Mills now believe whatever is contributing to the precancerous condition also may be causing the acid-secreting cells to die.

"These experiments suggest the loss of the acid-secreting cells and metaplasia may occur through different mechanisms," Mills said. "Ultimately, the cause may be the same, for example infection with H pylori, but simply damaging or destroying the cells isn't sufficient to launch this cascade. The more we can learn about what actually causes the <u>precancerous condition</u>, the more likely we'll be able to interrupt the cascade and prevent <u>stomach cancer</u>."



More information: Joseph Burclaff et al. Targeted Apoptosis of Parietal Cells is Insufficient to Induce Metaplasia in Stomach, *Gastroenterology* (2016). DOI: 10.1053/j.gastro.2016.12.001

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