

Scissor-like enzyme points toward treatment of infectious disease

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Along with Dr. Neal Alto is Nikolay Burnaevskiy, a graduate student in microbiology Credit: UT Southwestern Medical Center

UT Southwestern Medical Center researchers report that a pathogen annually blamed for an estimated 90 million cases of food-borne illness defeats a host's immune response by using a fat-snipping enzyme to cut off cellular communication.

"Our findings provide insight into severe bacterial infectious diseases, as well as some forms of cancer, in which the attachment of fat molecules to proteins is an essential feature of the disease process," said Dr. Neal



Alto, assistant professor of microbiology and senior author of the study in today's print edition of *Nature*. The study's first author is Nikolay Burnaevskiy, a graduate student in microbiology.

The research group discovered a scissor-like enzyme that specifically cuts off functionally-essential fatty acids from proteins. "The one we studied in particular – a 14-carbon saturated fatty acid called myristic acid– has received a lot of attention due to its crucial role in the transformation of normal cells to <u>cancer cells</u> and for promoting <u>cancer cell growth</u>," Dr. Alto said.

Because of the fat's importance in human disease, researchers have tried for years to identify effective methods to remove them from proteins. "To our amazement, bacteria have invented the precise tool for the job," Dr Alto said.

The bacteria used in this study, *Shigella flexneri*, are able to cross the intestinal wall and infect immune cells. Other <u>intestinal bacteria</u>, such as *E. coli*, are unable to do this. Once *Shigella* encounters <u>immune system</u> <u>cells</u>, including <u>white blood cells</u> such as macrophages, the bacteria use a needle-like complex to inject the cells with about 20 <u>bacterial toxins</u>.

The UTSW researchers conducted a series of experiments to characterize one of those toxins, called IpaJ, chosen in part because so little was known about the protein. They not only discovered IpaJ's fatcutting ability, but also determined how the protein disables the immune system's <u>communication infrastructure</u>, which Dr. Alto compared to knocking out a bridge needed to deliver a package.

"Normally, a macrophage will engulf an invading bacteria and send out cytokines, proteins that act as cellular alert signals, which in turn recruit more <u>immune cells</u> to the site of infection," Dr. Alto said. "When the macrophages engulf *Shigella*, however, the bacteria use IpaJ to cut fatty



acids from proteins, which need those fats attached in order to sound the alarm. Doing so buys more time for the bacteria to grow and survive.

"It's very interesting from a disease process point of view, but it's also important because we now have a potential drug target," said Dr. Alto. The next step, he said, will be to identify small molecule inhibitors that are specific to this fat-snipping protease and that might be developed into drugs.

Provided by UT Southwestern Medical Center

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