

Gene identified for immune system reset after infection

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Duke researchers have uncovered the genes that are normally activated during recovery from bacterial infection in the *C. elegans* worm. The finding could be key to new antibiotics and countering auto-immune disorders. Credit: Duke University, Alejandro Aballay

When pathogenic bacteria like *Salmonella* or *Staphylococcus* invade a host, the host organism should respond by going into a state of high alert,



altering its metabolism to defend against the attack.

But if the host doesn't reverse course once the battle is won, its efforts will be wasted on defense rather than on repairing the damage done by bacterial invaders.

Duke University researchers have uncovered the genes that are normally activated during recovery from bacterial infection. The finding could lead to ways to jumpstart this recovery process and possibly fend off autoimmune diseases and chronic inflammatory disorders that can result from the body staying in attack mode for too long.

The study appears Oct. 23, 2014 in PLOS Genetics.

"While the steps involved in recognizing microbial pathogens and inducing the immune response have been extensively studied, the pathways involved in host recovery after an infection are not well understood," said Alejandro Aballay, Ph.D., an associate professor of molecular genetics and microbiology at Duke University School of Medicine.

To examine the biological changes associated with this transition from wartime to peacetime, Aballay and lead study author Brian Heard, PhD, a postdoctoral fellow in his lab, infected the worm *C. elegans* with the common bacterial pathogen *Salmonella enterica*.

The researchers used gene chip technology to identify genes in the worm that were "turned off" or "turned on" during that initial infection. Then they treated the worm with the antibiotic tetracycline and looked to see what genes were dampened or activated as the organism recovered from infection.

They discovered that as the infection resolves, genes involved with



stimulating the immune response were turned off, while genes involved in returning to business as usual—a process known as "cellular homeostasis" – were turned on. These recovery genes play important roles in repairing tissue damage, eliminating bacterial toxins and clearing the cellular fighters that could cause excessive inflammation and further harm tissues.

When the researchers looked closer at the genes involved in recovery, they discovered many were controlled by the ELT-2 transcription factor, a type of <u>master switch</u> that had already been shown to regulate when and where <u>immune response genes</u> were expressed. The researchers showed that shutting down this master switch prevented infected animals from recovering with antibiotic treatment, showing that ELT-2 functions not only in defense but also in recovery.

"Our studies indicate that host or bacterial signals during the initial decline in infection may shift ELT-2's duties from priming the innate immune to protecting against chemical insults," Aballay said. "This opens a number of possibilities for further research."

Aballay says a number of studies in *C. elegans* have shown that the worm's nervous system, which can respond rapidly to all sorts of environmental stimuli, is ultimately responsible for activating the immune system in response to <u>bacterial infection</u>. So now he plans to conduct experiments in the worm to determine whether the nervous system also controls the recovery after infections have been cleared.

More information: "Recovery from an Acute Infection in C. elegans Requires the GATA Transcription Factor ELT-2," Brian Head and Alejandro Aballay. *PLOS Genetics*, Oct. 23, 2014. <u>DOI:</u> <u>10.1371/journal.pgen.1004609</u>



Provided by Duke University

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