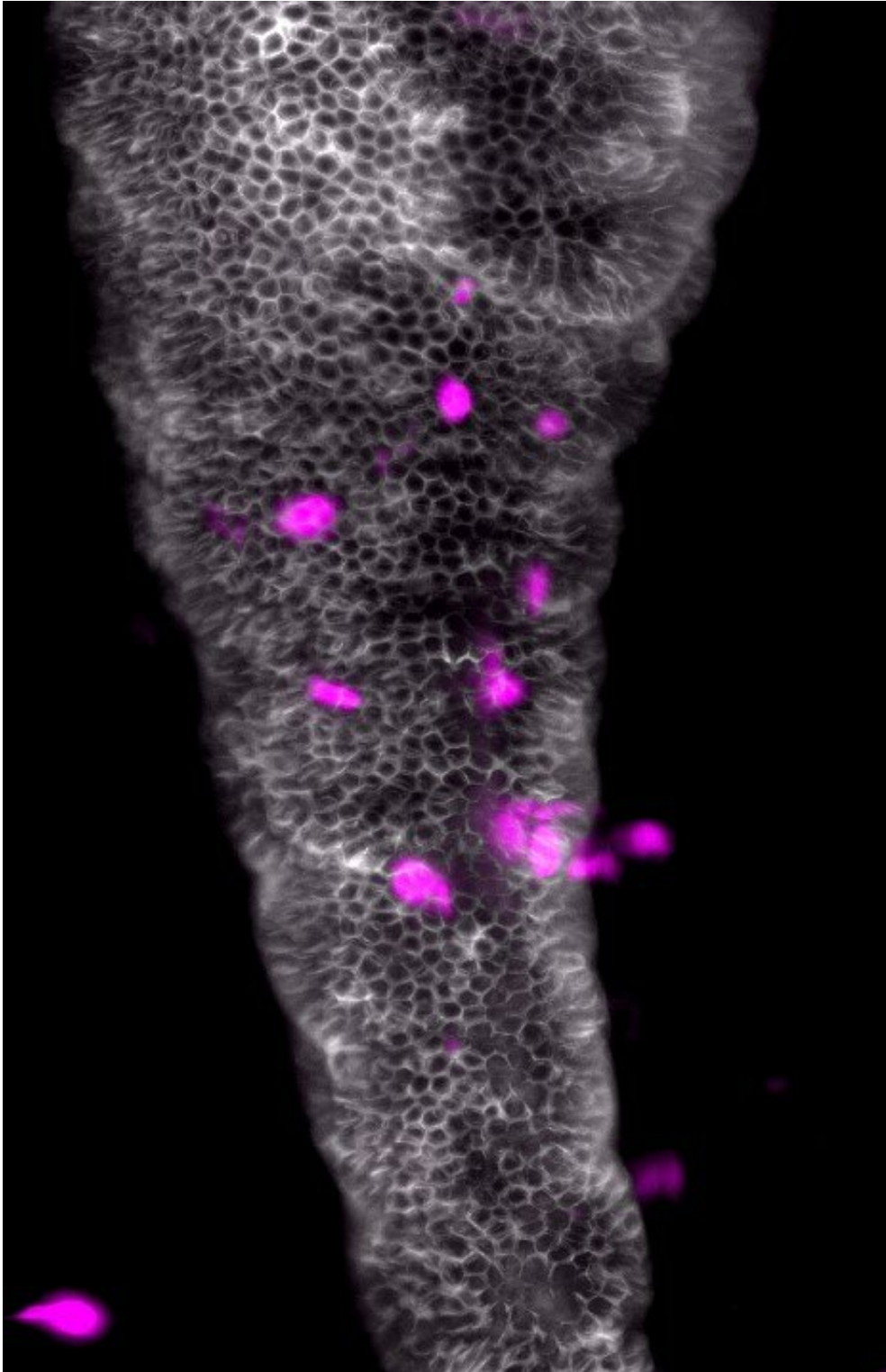


How the microbiota controls neutrophil activity

March 7 2019



Neutrophils take their cue: A protein produced in the intestinal epithelium in response to the microbiome educates innate immune cells. High resolution in vivo imaging of a zebrafish larva reveals the intimate association between

neutrophils (magenta) and the intestine (gray). Credit: Caitlin Murdoch (2019)

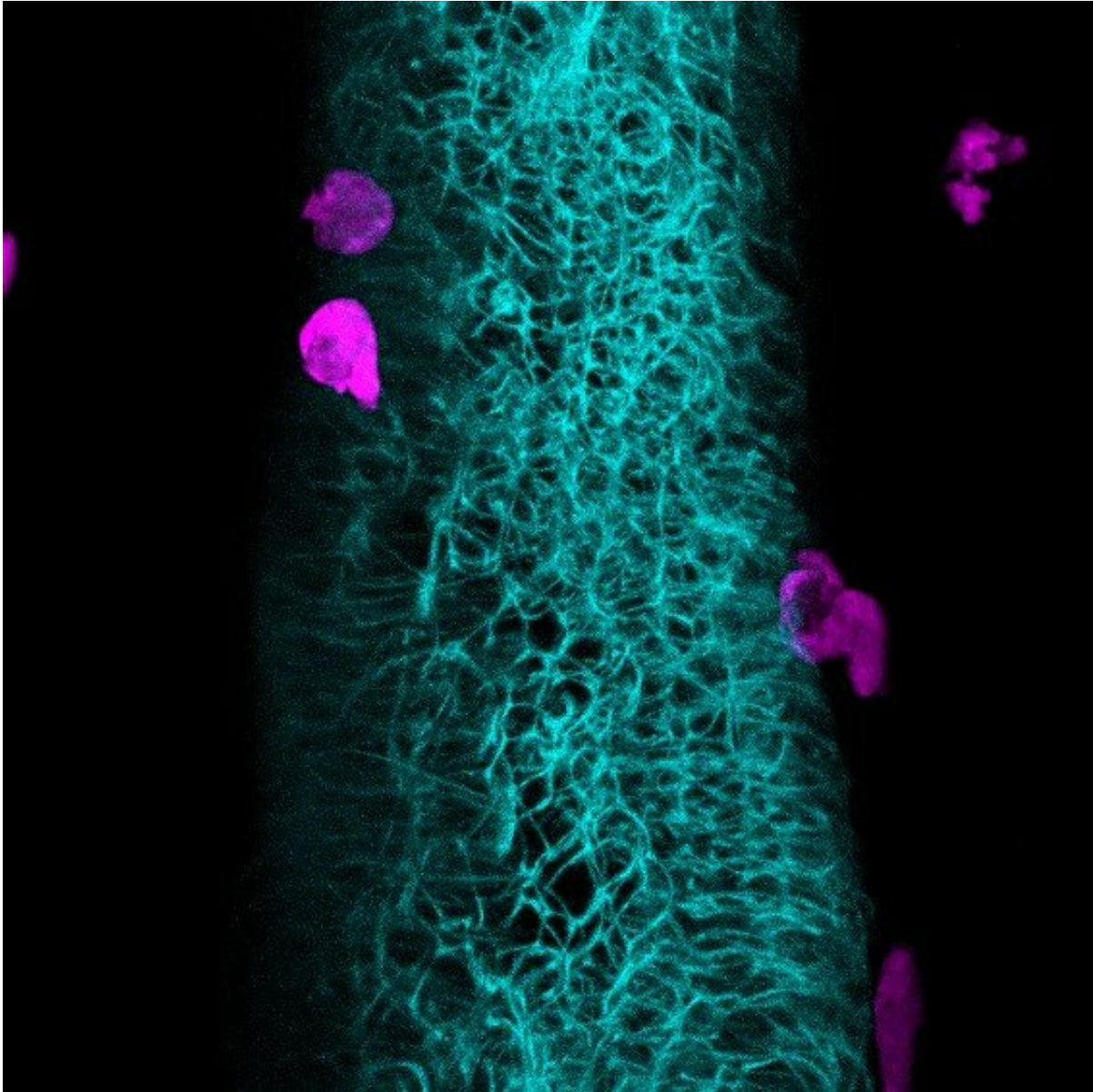
A host protein called Serum Amyloid A (Saa) is a major factor mediating the effects of the microbiota on the function of immune cells called neutrophils, according to a study published March 7 in the open-access journal *PLOS Pathogens* by John Rawls of Duke University School of Medicine, and colleagues.

Experiments in zebrafish showed that Saa produced by the intestine in response to the microbiota serves as a signal to [neutrophils](#) to restrict aberrant activation, decreasing inflammation and bacterial killing potential while simultaneously enhancing their ability to migrate to wounds.

The intestine is colonized by complex microbial communities called the microbiota, which impact diverse aspects of host physiology, including the development and [function](#) of the immune system. The microbiota strongly impacts the activity of innate immune cells called neutrophils, which are essential for host defense against infection, but the underlying mechanisms remain poorly understood. Caitlin Murdoch, a [graduate student](#) in Dr. Rawls' lab suspected that the effects of the microbiota on immunity could be mediated by Saa. This [host protein](#) is produced by the intestine and liver in response to the microbiota, but until now, its functions in living organisms were elusive.

Using normal and mutant zebrafish, the researchers demonstrate that Saa promotes the recruitment of neutrophils to sites of injury, yet restricts the clearance of pathogenic bacterial infection. Analysis of isolated neutrophils revealed that Saa reduces the bactericidal activity of these cells and their expression of pro-inflammatory genes. These effects depend on microbiota colonization, suggesting that this protein mediates

the microbiota's effects on host immunity. Collectively, these data establish that Saa induced by the microbiota in the intestine signals to neutrophils, tuning the extent to which they may be activated by other microbes or respond to injury. Because antibiotic treatment results in reduced intestinal Saa in mice, the authors suggest that antibiotic treatment could be associated with Saa-mediated aberrations in neutrophil function. They speculate that secondary infections that can occur following antibiotic use could be due in part to concomitant alterations in the production of this protein.



Neutrophils take their cue: A protein produced in the intestinal epithelium in response to the microbiome educates innate immune cells. High resolution in vivo imaging of a zebrafish larva reveals the intimate association between neutrophils (magenta) and the intestine (cyan). Credit: Caitlin Murdoch (2019)

The authors add, "Our study demonstrates that a host protein made in the

intestine following exposure to the [microbiota](#) affects innate immune cell populations, both in the gut and other tissues. Microbially responsive secretion of host factors from the [intestine](#) can therefore serve as a molecular rheostat which controls the development and function of the [host](#) innate immune system."

More information: Murdoch CC, Espenschied ST, Matty MA, Mueller O, Tobin DM, Rawls JF (2019) Intestinal serum amyloid a suppresses systemic neutrophil activation and bactericidal activity in response to microbiota colonization. *PLoS Pathog* 15(3): e1007381. doi.org/10.1371/journal.ppat.1007381

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