Study offers new insights into immune mechanisms of inflammatory disease
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Preclinical model of intestinal parasitic infection. Digitally colored: Red (intestine), green (parasitic worms), large blue dots within intestine (specialized cells called goblet cells, a signature of type 2 inflammation). Credit: Hiroshi Yano and David Artis/Provided

The study, published Nov. 2 in Nature, shows that although ILC2s have many functional similarities to immune cells called T helper type 2 cells (Th2 cells), the latter cell type cannot adequately compensate for loss of the protective response of ILC2s against parasitic worm infection in the gut as well as gut inflammation. Underscoring the clinical relevance of the study, the researchers found evidence that ILC2s in humans respond in a manner similar to mouse ILC2s.

"This advances our understanding of the complexity of the immune system, and gives us a potential new set of targets for future therapies," said study senior author David Artis, director of the Jill Roberts Institute for Research in Inflammatory Bowel Disease, director of the Friedman Center for Nutrition and Inflammation and the Michael Kors Professor of Immunology at Weill Cornell Medicine.

ILC2s are part of a family of cells, innate lymphoid cells, that were discovered by multiple groups only about 12 years ago. With their strong presence in barrier tissues, innate lymphoid cells are generally considered to serve as sentinels and first responders against various types of infection. But scientists also recognize that ILCs may hold the keys to understanding common inflammatory and autoimmune conditions such as asthma and IBD.

It is thought that both ILC2s and Th2 cells evolved at least in part to defend the body from parasitic worm infections, biting insects and other environmental triggers. When triggered by such challenges, both help marshal what is called a type 2 immune response. These similarities have led researchers to suggest that they are functionally almost the same but ILC2s specialize in earlier, more localized responses, whereas T cells are more blood-borne and mobile, concentrating in multiple tissues where needed.

However, in the new study, the researchers found that ILC2s have an essential immune role rather
than being redundant as type 2 immune responders.

When ILC2s and Th2 cells are activated by a worm infection, they both produce an anti-worm, tissue-protecting protein called amphiregulin (AREG). To determine if Th2 cells can compensate for loss of this protein from ILC2s, the researchers engineered mice in which AREG production is selectively deleted in ILC2s, but not in Th2 cells.

They found that these mice were more susceptible to parasitic worm infection in the gut due to reduced capacity to mount an anti-parasitic immune response, compared with mice with normal ILC2s. The mice lacking ILC2 AREG were also much more susceptible to gut damage from inflammation.

"This finding clarifies that ILC2s are playing the major role in this tissue protective response—without them the response is inadequate," said study co-first author Hiroshi Yano, a postdoctoral research associate in the Artis laboratory.

Clarifying the functional importance of a major immune cell type is a significant achievement in basic immunology and the results of the study also suggest clinical applications. The researchers showed that the ILC2 immune response, either to worm infection or inflammatory gut damage, is selectively controlled by a signaling molecule produced by neurons in the gut.

Giving the molecule to mice with experimental gut inflammation boosted AREG production in ILC2s and protected the animals from gut damage. Preliminary experiments with gut ILC2s taken from patients with inflammatory bowel disease showed that the molecule could boost the protective response in the human cells as well. These findings suggest that neurons in the gut communicate with ILC2s to generate a protective response that can't be replaced by other immune cells, thus offering new therapeutic opportunities, Artis said.
