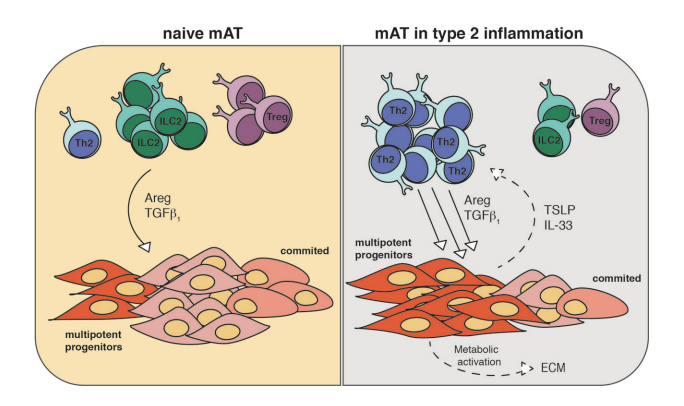


Fatty tissue surrounding intestinal tracts of mice found to help eject gut-infesting worms

October 17 2022, by Bob Yirka



The graphic is a schematic summary of the main findings of the paper, where changes in mesenteric adipose tissue during infection are highlighted. Credit: Agnieszka Kabat and Edward Pearce

A team of researchers at the Max Planck Institute for Immunobiology and Epigenetics, working with colleagues from the Washington University School of Medicine, the Medical University of Vienna, the



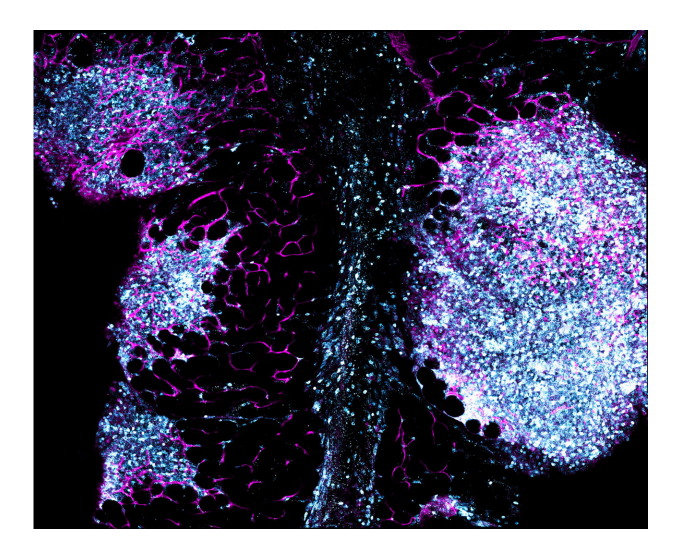
Van Andel Research Institute and the USDA has found that fatty tissue surrounding the intestinal tracts of mice help to eject gut-infesting worms. Their paper is published in the journal *Science Immunology*.

Prior research has shown that mesenteric adipose tissue aids the <u>immune system</u> in responding to pathogens and also certain types of cancer. In this new effort, the researchers found that it also assists with combating <u>parasitic infections</u>.

The work began when the researchers inadvertently noticed that mesenteric adipose tissue (<u>fatty tissue</u> that surrounds the <u>intestinal tract</u>) stiffened when a mouse was infected with helminths, a type of worm. Such stiffening, the team noted, helped to eject the worms.

In taking a closer look at how fatty tissue might recognize and respond to a parasitic infection, the researchers found that a type of T cell found in such tissue, called Th2, communicated with <u>stromal cells</u>, which play a role in differentiating cells that grow into different types of structural support cells. They then conducted <u>flow cytometry</u>, histology, cell culture and single-cell RNA sequencing on the fatty tissue to learn more about how it stiffened in response to an infection.





An immunofluorescence image of T cells (white) in the mesenteric adipose tissue accumulating in lymphoid structures and within interstitial spaces during infection. The vascularisation of these lymphoid structures is also highlighted (purple). Credit: Agnieszka Kabat and Edward Pearce

They discovered that Th2 cells infiltrated fat tissue even though the parasite did not infect such tissue. They also found that the Th2 cells released both Amphiregulin and cytokine TGF β . Additionally, the stromal cells became more active when in the presence of Amphiregulin and cytokine TGF β and thus produced more cytokines. And finally, they



found that the stromal and Th2 cells were combining in interstitial spaces in the fatty tissue, forcing them to become enlarged and to secrete collagen, which resulted in tissue stiffening.

The researchers also found that the stiffening lasted up to a year before tissue returned to normal, rather than progressing to fibrosis. This allowed the mice to respond more quickly to a secondary infection.

More information: Agnieszka M. Kabat et al, Resident T H 2 cells orchestrate adipose tissue remodeling at a site adjacent to infection, *Science Immunology* (2022). DOI: 10.1126/sciimmunol.add3263

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