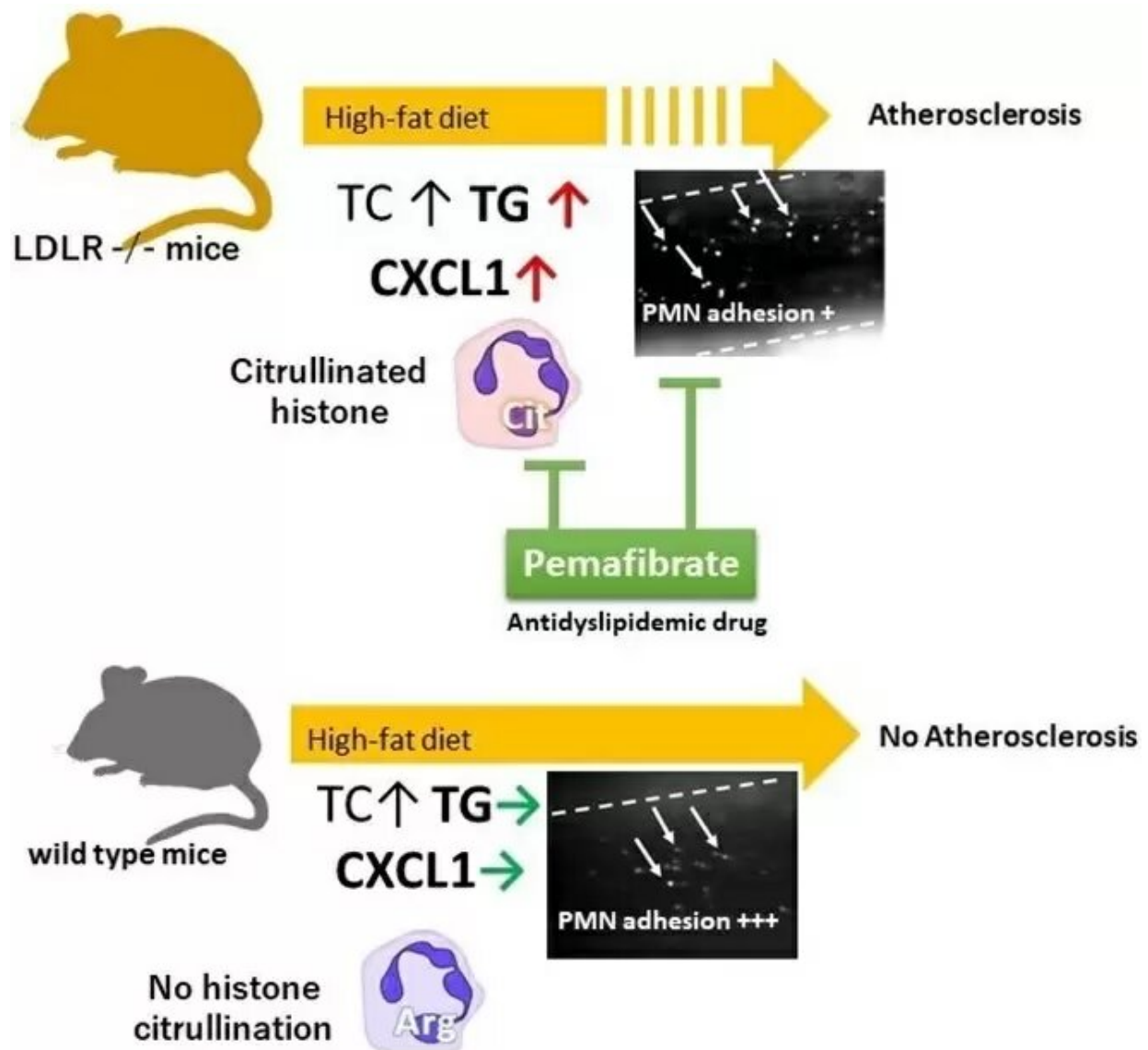


# Using mice to explore the prevention of blocked arteries

July 9 2021



Proposed mechanisms of vascular inflammation in LDLR -/- mice fed High Fat

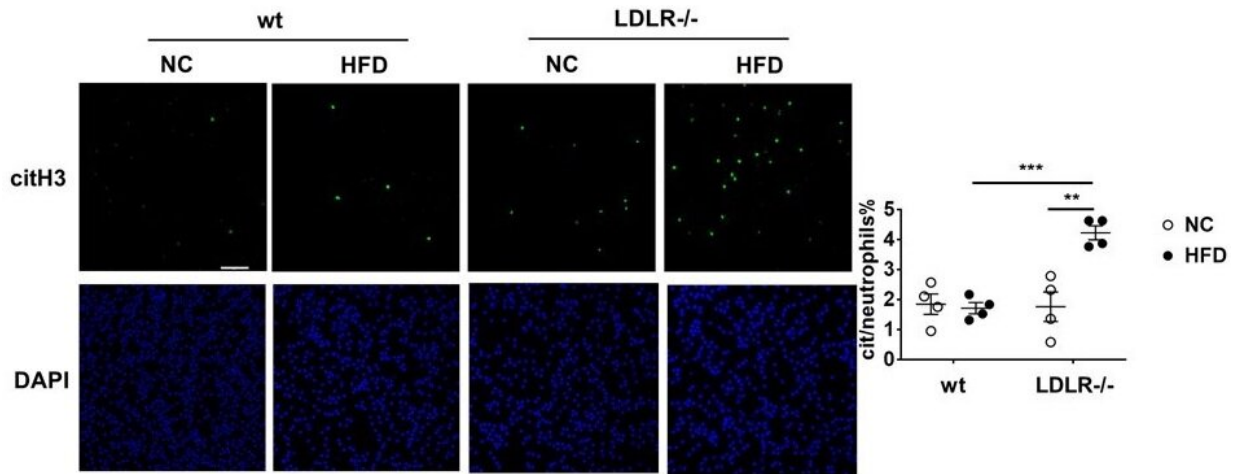
Diet. Credit: Tokyo Medical and Dental University

It's long been known that a high-fat diet can lead to clogged arteries, but we have only recently begun to learn in detail how the process works. A new study in experimental mice could go a long way to finding treatments to keep arteries open and flowing.

In a study published this month in *JACC Basic To Translational Science*, researchers from Tokyo Medical and Dental University (TMDU) have continued their research into how a [high-fat diet](#) causes atherosclerosis, the fatty buildup of plaques on the walls of blood vessels that can lead to heart attacks and other vascular disease. Atherosclerosis is not as simple as fats getting stuck in your arteries—it's actually an inflammatory disease driven by the body's own immune response, particularly neutrophils, the [white blood cells](#) that attack infections and respond to injuries.

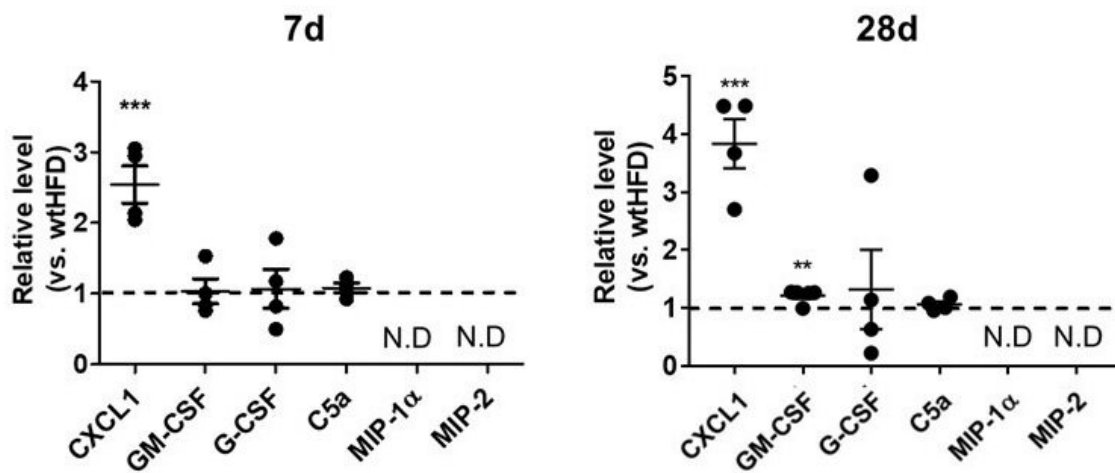
Normally, neutrophils act to regulate the immune response, so that inflammation can resolve the problem allowing the body to return to its normal state. But when a stimulus is persistent, the immune response can shift strategies, becoming a long-term chronic condition.

"We applied this theory to what happens in atherosclerosis—in this case, the persistent stimulus is dyslipidemia (unhealthy levels of fats and [cholesterol](#) in the blood)," says lead author of the study Mizuko Osaka. "So detailing the role of neutrophils in acute inflammation can help us understand how it becomes chronic inflammation."



Citrullination of histoneH3 enhanced in peripheral neutrophils of LDLR<sup>-/-</sup> mice fed HFD. Green: citrullinated histoneH3, Blue: nuclei of neutrophils. \*\*\*P<0.001, \*\*P<0.01.

Peripheral neutrophils are collected after 4 weeks of high fat diet (HFD) to LDLR<sup>-/-</sup> mice or wild type control mice. Significant citrullinated HistoneH3 signals are only recognized in LDLR<sup>-/-</sup> mice with HFD. Credit: Department of Life Science and Bioethics, TMDU



Cytokines and chemokines in blood of LDLR<sup>-/-</sup> mice fed HFD relative to wild-type mice fed HFD. \*\*\*P<0.001, \*\*P<0.01

Extensive serum analysis from LDLR<sup>-/-</sup> mice after High Fat Diet for 7 days and

28 days revealed that elevation of CXCL1 as early as 7 days after HFD suggesting its pivotal role in observed citrullination of neutrophil histone H3. Credit: Department of Life Sciences and Bioethics, TMDU

The group compared a group of regular wild-type mice against mice specially bred to genetically lack [low-density lipoprotein](#) (LDL) receptors on their cell membranes. This is because mice carry most plasma cholesterol in [high-density lipoprotein](#) and overall have far lower cholesterol levels; lacking these receptors means these lab mice metabolize LDL (the "bad" cholesterol) more like humans do. When fed a "humanized" high-fat diet, these mice developed high triglycerides and cholesterol, showed significantly increased levels of neutrophil adhesion to their blood vessel walls, and experienced much more inflammation.

"On the basis of certain biomarkers in the LDLR-null mice, we suspected that neutrophil extracellular traps (NETs, which help trap and eliminate pathogens from the body) could be involved in activating neutrophils," explains Masayuki Yoshida, senior author of the study. "And indeed, we found enhanced citrullination of histone H3, a known marker of NETs, in these mice. Going further, we specifically identified plasma levels of CXCL1 (a peptide that acts to attract immune cells to the site of an injury or infection) to be significantly increased in the LDLR-null mice after 7 days and 28 days of the high-fat diet."

This suggests that CXCL1 is the link between the high-fat diet and citrullination, part of the process of making NETs—and which leads to neutrophils forming plaques on the blood vessel walls when overstimulated in the long term. In fact, blocking the citrullination process led to a reversal of the increased neutrophil adhesion from the high-fat [diet](#).

Once we fully understand how a disease happens, it's easier to develop strategies to prevent it. In fact, although [mice](#) are metabolically different from people, it's possible that certain medications already being used to control cholesterol could be used to decrease neutrophil adhesion by affecting the pathways identified in this study.

**More information:** Mizuko Osaka et al, High-Fat Diet Enhances Neutrophil Adhesion in LDLR-Null Mice Via Hypercitrullination of Histone H3, *JACC: Basic to Translational Science* (2021). [DOI: 10.1016/j.jacbts.2021.04.002](#)

Provided by Tokyo Medical and Dental University

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