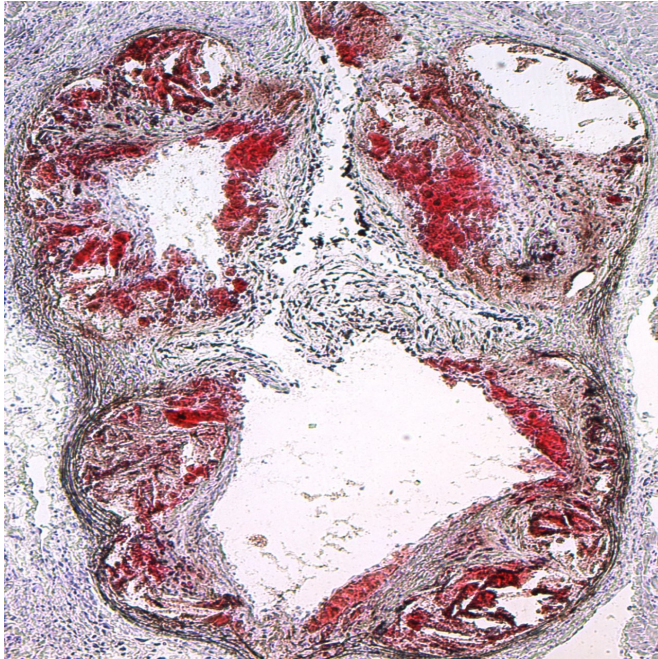


New cellular pathway helps explain how inflammation leads to artery disease

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Atherosclerotic lesions appear in red in this image of a mouse aorta. Credit: Cedars-Sinai

Investigators have identified a new cellular pathway that may help explain how arterial inflammation develops into atherosclerosis—deposits of cholesterol, fats and other substances that create plaque, clog arteries and promote heart attacks and stroke. The findings could lead to improved therapies for atherosclerosis, a leading cause of death worldwide.

"We have known for decades that [atherosclerosis](#) is a disease of chronic inflammation that ultimately results in the scarring of arteries and tissue damage," said Moshe Arditi, MD, director of the Infectious and Immunologic Disorders Translational Research Center in the Department of Biomedical Sciences at Cedars-Sinai. "But the ongoing stimulus for this inflammation has been

unclear."

A study published today in the journal *Cell Metabolism* sheds light on this mystery by using a [bacterial infection](#) to reveal a cascade of cellular events that can lead to inflammation and atherosclerosis. Arditi is the co-senior author and the lead author of the study, which was led by investigators at Cedars-Sinai.

Investigators focused on interleukin-1 beta, a type of protein that is assembled and released by immune system cells in response to infection and injury, including [tissue damage](#) caused by atherosclerosis. While interleukin-1 beta helps rally the immune system against these threats, it also can cause [chronic inflammation](#). The study team wanted to understand how the interleukin-1 beta pathway might promote atherosclerosis.

Using laboratory mice bearing a bacterial infection, along with human cells cultured in a petri dish, the team discovered that several harmful processes related to interleukin-1 beta can lead to buildup of [cholesterol](#) in the arteries:

- To make its way out of the immune system cell, interleukin-1 beta can also use the same chemical channels that are used by cholesterol to exit the cell. The result is a "traffic rush" on those channels that blocks the exit of artery-damaging cholesterol and causes it to accumulate in the cell.
- Once it is released by the cell into the body, interleukin-1 beta suppresses a chemical receptor that enables niacin, or Vitamin B3, to be used in the body. This action is harmful because niacin works by removing cholesterol from [cells](#) in the artery walls. When niacin is blocked, cholesterol can accumulate in the walls.
- The suppression of the niacin receptor has another negative effect: It reduces the number of chemical channels that

cholesterol uses to exit the immune system cell, causing more cholesterol to be trapped inside. That is because the niacin receptor, besides enabling niacin, also increases these channels as part of its normal function.

Chlamydia pneumoniae Hijacks a Host Autoregulatory IL-1? Loop to Drive Foam Cell Formation and Accelerate Atherosclerosis, *Cell Metabolism* (2018). DOI: [10.1016/j.cmet.2018.05.027](https://doi.org/10.1016/j.cmet.2018.05.027)

Arditi said these discoveries are especially significant because drugs that inhibit interleukin-1 beta have shown promise in combatting atherosclerosis and heart disease. A major clinical trial, led by another research institution and published last year, reported that administering one such drug to patients who had a prior heart attack reduced inflammation and lowered the risk of another cardiovascular event.

Provided by Cedars-Sinai Medical Center

The Cedars-Sinai study raises the possibility that by using drugs to block the initial production of interleukin-1 beta, rather than just neutralizing it, a stronger positive effect could be obtained for these patients, said Ardit, professor of Pediatrics and Biomedical Sciences.

Prediman K. Shah, MD, director of the Atherosclerosis Prevention and Management Center at Cedars-Sinai, noted that a drug, colchicine, already exists that blocks interleukin-1 beta production, but it is FDA-approved only to treat gout and Mediterranean fever. Two clinical trials are underway elsewhere to evaluate the drug's potential for treating atherosclerosis and preventing heart attacks.

In addition, Shah, a professor of Medicine who was not involved in the Cedars-Sinai study, said, "A very intriguing aspect of these findings is that they could prompt a re-examination of niacin therapy for atherosclerosis and heart disease." He explained that physicians long used niacin to treat atherosclerosis until the 1980s, when statin drugs were shown to be more effective at reducing cholesterol and cardiovascular risk.

The newly released study suggests that combining niacin with an interleukin-1 beta inhibitor might enhance niacin therapy by making [niacin](#) more available to the body, Ardit said.

More information: Gantsetseg Tumurkhuu et al,

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