

## A new culprit for multiple sclerosis relapses

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Demyelination by MS. The CD68 colored tissue shows several macrophages in the area of the lesion. Original scale 1:100. Credit: <u>CC BY-SA 3.0</u> Marvin 101/Wikipedia

A molecule that helps blood clot may also play a role in multiple sclerosis relapses, researchers report in the May 6 issue of *PNAS*. The new research may help answer the mystery of why remissions happen, as well as find early markers of the disease.



The research also shows a new way to study multiple sclerosis (MS) in  $\underline{\text{mice}}$  that is closer to the human form of the disease.

MS affects about a million people in the United States and many more globally. It damages the brain's ability to communicate with the rest of the body, making it hard to walk, write, or hold a fork and knife. This happens because of damage to the insulation around the nerves. Just like a frayed wire, a nerve with damaged insulation can short out or send bad signals.

But the damage isn't permanent, at least not at first. Most people with multiple sclerosis have recurring episodes of disability, followed by remissions when their symptoms lessen or disappear. Why these relapses and remissions happen is a great mystery. We know that the damage to the nerves is caused by the immune system, the army of cells in our body that is supposed to protect us from disease-causing invaders. For some reason, in MS, the immune system turns on cells in the brain and spinal cord. In MS patients, a particular type of immune cell—CD8+ cells, a part of the immune system that normally kills cells that are cancerous or infected—seem to be the ones doing the damage.

Although researchers have been able to develop drugs to help fight MS using a mouse version of MS, these experimental mice develop a slightly different immune system response than what happens in MS in humans. Different cells do the damage in MS mice: CD4+ cells. The mice have CD8+ cells, but those CD8+ cells are generally quiescent. This has been a big stumbling block to understanding how the <u>immune system</u> develops in MS.

But a team of researchers from UConn Health, the University of Illinois at Chicago (UIC), and the Gladstone Institutes have figured out how CD8+ cells are activated in MS mice, and the result seems very close to what happens in humans. The new findings hinge on how cells



talk to each other. Cells will often secrete little bubbles containing proteins and genetic signals. These bubbles are called extracellular vesicles, or EVs. EVs are made by most cells in the body, and float in the blood stream like a message in a bottle.

So the team injected EVs from normal, healthy mice into mice that had that experimental MS-like disease. When they did this, the mice acquired a relapsing-remitting disease and active CD8+ cells, like human MS patients. The researchers examined the EVs in mice and patients with MS and found they contained <u>fibrinogen</u>, a protein that neuroscientist Katerina Akassoglou's lab at Gladstone had been studying in MS. Fibrinogen normally helps blood clot and seal up wounds. But in these MS mice, the EVs with fibrinogen seemed to activate the CD8+ immune cells. When they injected the MS mice with EVs that did not have fibrinogen, they could not cause the relapsing-remitting illness.

"These findings expand our understanding of how fibrinogen contributes to the progression of MS pathology" says Akassoglou, senior investigator at Gladstone and professor of neurology at UC San Francisco. "Fibrinogen in exosomes may have far-reaching implications for therapies and as a biomarker for disease progression in MS and potentially, other neurological diseases," she says.

"We now have a robust model of relapsing/remitting disease driven by CD8+ cells," says UConn neuroscientist Stephen Crocker, who directed the study. "There's all these clinically important questions we can now ask." Crocker and his colleagues want to study this model further to understand how and why the remissions of disease happen.

"Understanding the causes of relapses is a key step on the path to a cure for MS," says study co-author Ernesto Bongarzone, an anatomy and cell biology neuroscientist and professor at UIC. "The results of this study and the identification of fibrinogen as a key molecule contributing to



relapses are exciting steps forward."

The researchers would like to understand how the fibrinogen stimulates the CD8+ <u>cells</u> that cause the relapsing and remitting disease activity. They would also like to test whether fibrinogen and related proteins found in the EVs also play a role in humans with MS, and test if these molecular signals in EVs might be early warnings of relapses or <u>disease</u> progression.

**More information:** Cory M. Willis el al., "Extracellular vesicle fibrinogen induces encephalitogenic CD8+ T cells in a mouse model of multiple sclerosis," *PNAS* (2019).

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