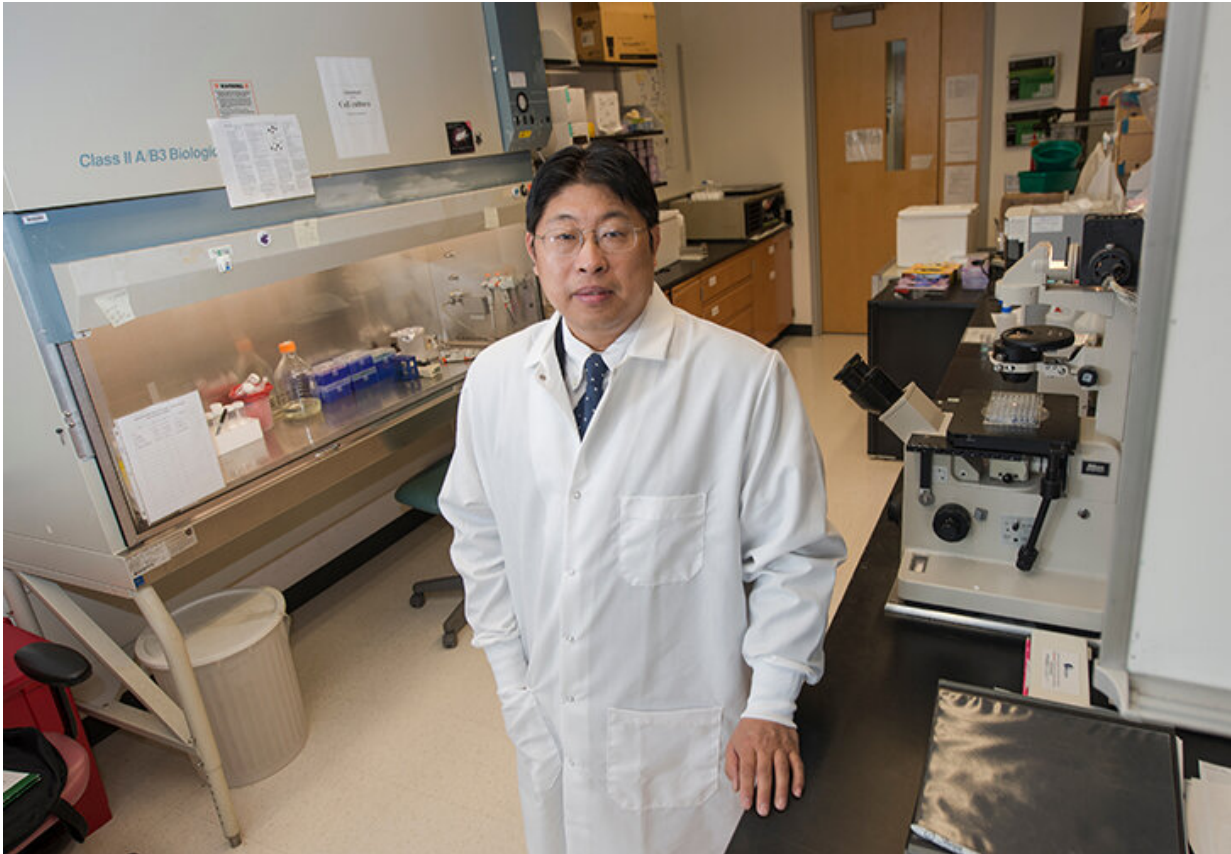


Finding clues to a functional HIV cure

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George Mason University College of Science Professor Yuntao Wu is part of a team researching a protein that mobilizes cells to fight against infection.
#MasonSolveX Credit: Evan Cantwell

George Mason University's Yuntao Wu is the lead scientist on a research team that has identified a measurable indicator that could prove

instrumental in the fight against HIV.

The research focuses on cofilin, a key protein that regulates [cells](#) to mobilize and fight against infection.

In an HIV-infected patient, cofilin dysfunction is a key factor in helper T cell defects, according to the research recently published in the journal *Science Advances*. Helper T cells augment the body's immune response by recognizing the presence of a foreign antigen and then helping the immune system mount a response.

"When you have an infection, you need to mobilize the T cells," said Wu, a College of Science professor of Molecular and Microbiology within Mason's School of Systems Biology and National Center for Biodefense and Infectious Diseases. "In HIV infection, there is a profound depletion of helper T cells in lymphoid tissues, such as those in the gut."

Antiretroviral therapy has significantly increased the lifespan of HIV-infected people, although it offers neither a cure nor a full restoration of the body's [immune system](#), he said. The natural course of the HIV infection leads to multiple immune defects, including the impairment of T cell migration, according to the research team.

Wu and his team found that patients with HIV have "significantly lower" levels of cofilin phosphorylation—which provides a control of cofilin's activity with the addition of a phosphate—than healthy patients. Cofilin is a key protein that helps cells generate the driving force for migration. Proper cofilin phosphorylation is needed for cells to move in and out of tissues.

Their findings suggest that a lasting immune control to HIV isn't likely to come from [antiretroviral therapy](#) alone because it is not sufficient to

repair the cofilin damage caused by HIV and to restore normal T cell migration in and out of tissues.

But the researchers found that by stimulating the T cells with additional therapeutics, such as the $\alpha 4\beta 7$ integrin antibody, they could modulate the levels of [cofilin](#) activity needed to restore T cell mobility. The remedy has shown lasting effects in immune control of simian immunodeficiency virus (SIV), the simian form of the AIDS virus, in a monkey trial, but it has not showed the same results in HIV-infected human patients.

"Now we have a marker, and at least one target that we can focus on to discover new therapies to repair the immune damages for a functional cure," Wu said.

More information: Sijia He et al, Cofilin hyperactivation in HIV infection and targeting the cofilin pathway using an anti- $\alpha 4\beta 7$ integrin antibody, *Science Advances* (2019). [DOI: 10.1126/sciadv.aat7911](https://doi.org/10.1126/sciadv.aat7911)

Provided by George Mason University

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