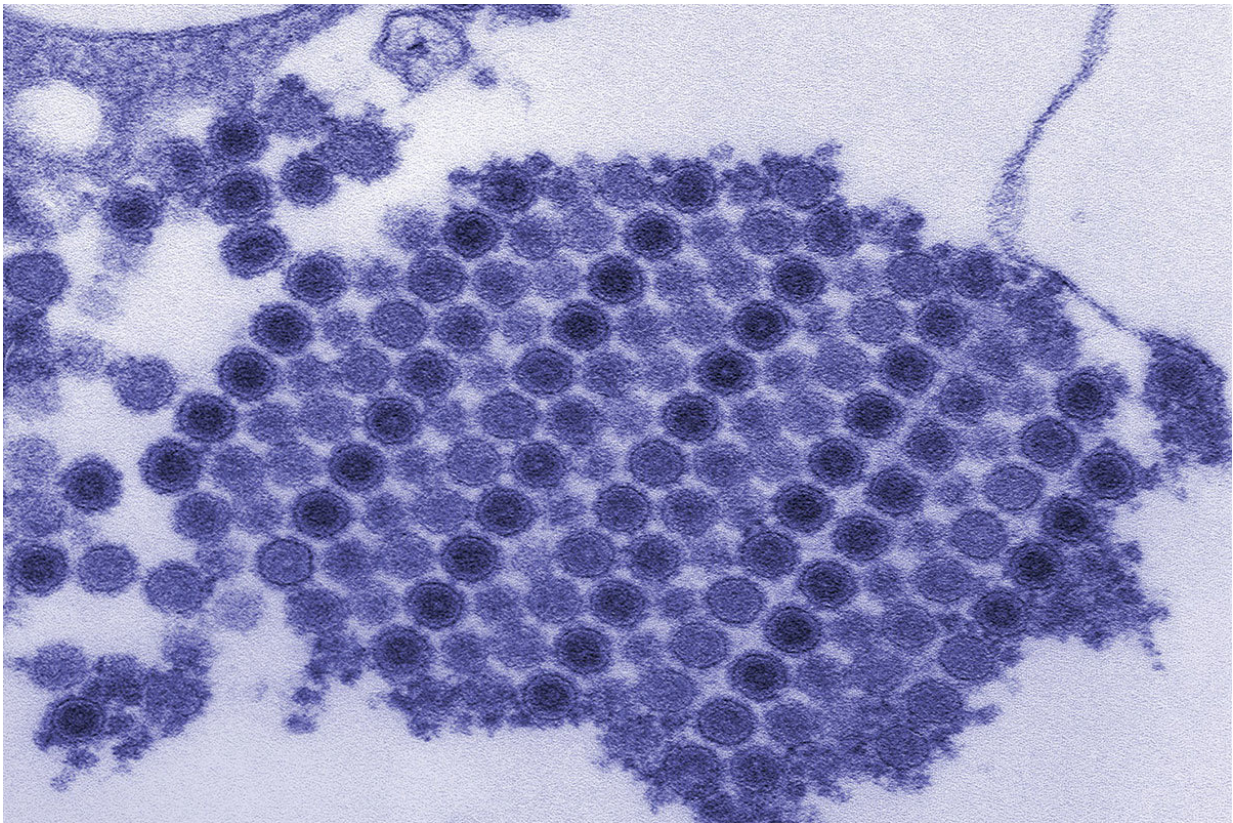


Why chikungunya, other arthritis-causing viruses target joints

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Shown are chikungunya virus particles crystallized into a neat grid. A new study has found the molecular 'handle' the virus uses to enter cells and cause arthritis. Blocking the virus from latching on to the handle could prevent or treat disease. Credit: CDC/CYNTHIA GOLDSMITH

Chikungunya virus is a growing threat to the United States and other

regions of the world as the mosquito that carries the virus expands its reach. Telltale symptoms of chikungunya infection are fever and joint pain that last about a week. But in up to half of patients, the virus can cause a debilitating form of arthritis that persists for months or even years.

Scientists have understood little about how chikungunya and related viruses cause arthritis. Now, researchers at Washington University School of Medicine in St. Louis have identified the molecular handle that chikungunya grabs to get inside cells. The findings, published May 16 in the journal *Nature*, could lead to ways to prevent or treat disease caused by chikungunya and related viruses.

The handle, or receptor, is located on cells that build cartilage, muscle and bone. Joints are filled with such cells, which helps explain patients' painful symptoms. Further, by creating decoy handles, the researchers showed that they could reduce chikungunya infection and signs of arthritis.

"The name chikungunya comes from the Makonde language of Tanzania, and it means 'to walk bent over.' That's how painful the arthritis can be," said senior author Michael S. Diamond, MD, Ph.D., the Herbert S. Gasser Professor of Medicine at the School of Medicine. "We now know how chikungunya gets into cells, and we may have found a way to block the infection. If the [virus](#) cannot get into the cell, it is unable to replicate and cause infection and disease.

There are no specific treatments or vaccines for chikungunya and related viruses, known as arthritogenic alphaviruses. Doctors simply recommend rest, fluids and over-the-counter pain relievers such as acetaminophen or ibuprofen.

With the aid of a warming planet and modern means of transportation,

mosquitoes that carry chikungunya and related viruses are spreading. Once limited to Asia and Africa, chikungunya virus has infected more than a million people in the Caribbean and South America in an outbreak that began in 2013 and continues to this day.

Figuring out how the virus gets inside cells is considered a step toward slowing its spread. Diamond, first author and postdoctoral researcher Rong Zhang, Ph.D., and colleagues identified the [protein](#) on cells that [chikungunya virus](#) latches onto.

The protein is called Mxra8, and it is needed for chikungunya to invade both human and mouse cells, the researchers found. Additional experiments showed that not just chikungunya but its arthritis-causing relatives—Mayaro, Ross River, O'nyongnyong and Barmah Forest viruses—require the protein to get into cells.

Since chikungunya uses Mxra8 protein as a handle to open a door into cells, the researchers tested whether preventing the virus from grabbing that handle could reduce infection. They deluged the virus with decoy handles, reasoning that [chikungunya](#) would grab the decoy and be locked out of cells. Only the few individual viruses that latched onto a true handle could infect cells, so the overall infection rate—and signs of arthritis—would fall.

And that's just what they found. A day after infection, the level of virus in the mice's ankles and calf muscles was between tenfold and a hundredfold lower in the animals that had been treated with Mxra8 proteins or blocking antibodies than those that received placebo, and the numbers remained lower over the next two days. In addition, three days after treatment, the mice that had received the protein exhibited much less swelling in their ankles than those that received the placebo.

The results suggest that a compound that blocks the virus from attaching

to Mxra8 on the surface of [cells](#) could prevent or reduce arthritis.

"Not much is known about what Mxra8 does in the human body, so we need more information before developing a drug that targets Mxra8," said Diamond, who also is a professor of molecular microbiology, and of pathology and immunology. "But we could more immediately develop a drug that targets the virus and prevent it from attaching to this protein."

The researchers are working on mapping the structure of the protein and locating the exact spot to which the virus attaches. Such information could help researchers design a compound to interfere with the virus's ability to hold onto the protein, or to design vaccines to prevent [infection](#).

More information: Mxra8 is a receptor for multiple arthritogenic alphaviruses, *Nature* (2018).

[nature.com/articles/doi:10.1038/s41586-018-0121-3](https://doi.org/10.1038/s41586-018-0121-3)

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