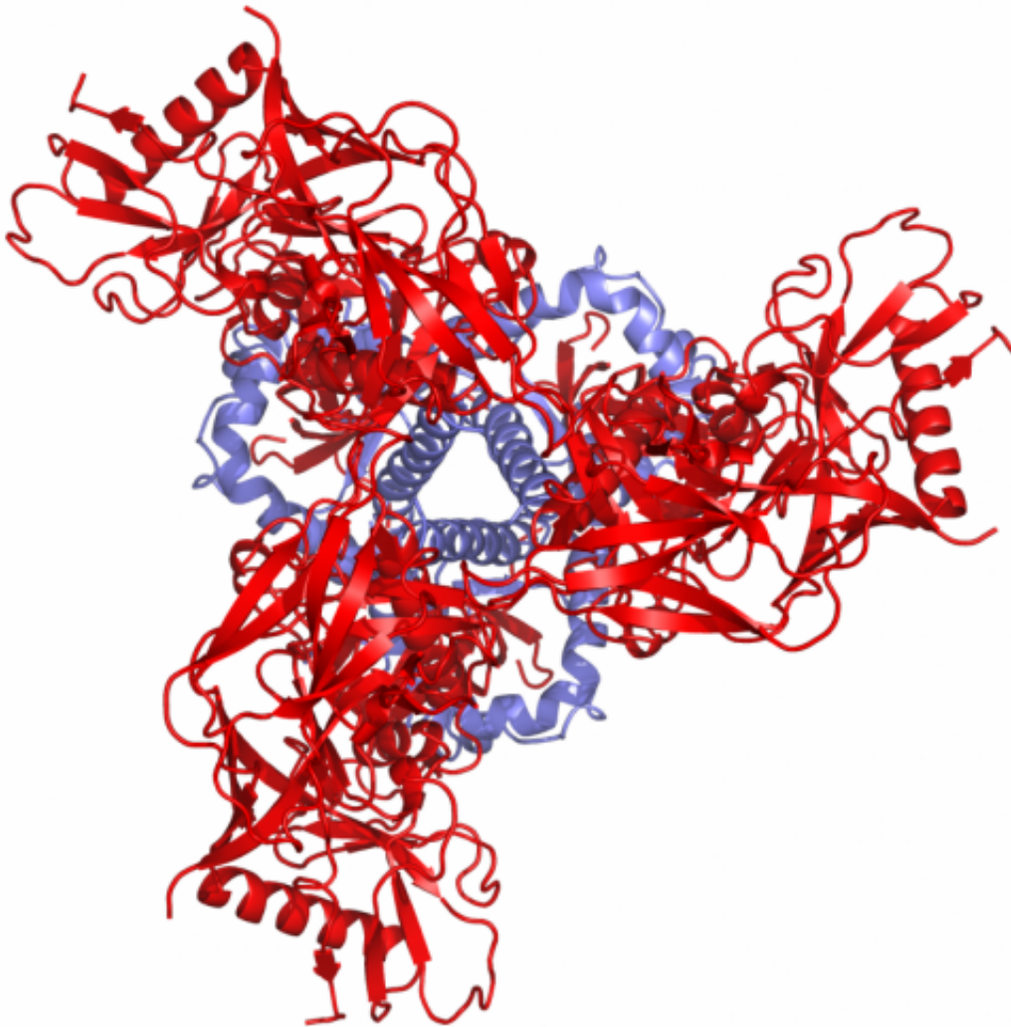


# Researchers watch, in real time, the dynamic motion of HIV as it readies an attack

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This schematic depiction of the protein structure of the pre-fusion HIV spike as viewed from above shows the three gp41 molecules in blue and the three gp120 molecules in red. Credit: NIAID

Researchers at Weill Cornell Medical College have developed technologies that allow investigators, for the first time, to watch what they call the "dance" of HIV proteins on the virus' surface, which may contribute to how it infects human immune cells. Their discovery is described in the Oct. 8 issue of *Science*, and is also a part of a study published the same day in *Nature*.

The new technology platform opens new possibilities for devising an approach to prevent HIV infection, says Dr. Scott Blanchard, an associate professor of physiology and biophysics at Weill Cornell, and one of three co-lead authors on the *Science* study. He is also an author on the *Nature* paper that describes a three-dimensional structure of one of the shapes, or conformations of the HIV [protein](#).

"Making the movements of HIV visible so that we can follow, in real time, how surface proteins on the virus behave will hopefully tell us what we need to know to prevent fusion with human cells—if you can prevent viral entry of HIV into immune cells, you have won," says Dr. Blanchard, who is also associate director of Weill Cornell's chemical biology program.

"What we have shown in the *Science* study is that we now have the means to obtain real-time images of processes happening on the surface of intact HIV particles, which we now plan to use to screen the impact of drugs and antibodies that can shut it down," he says.

"We desperately need solutions to prevent HIV infection, which, to date, has infected or killed more than 70 million people worldwide," Dr. Blanchard says. If this technology proves useful in HIV management, it could potentially be used to decode infection processes for other viruses, he says.

## Using Light to Watch HIV Dance

In the *Science* study, Dr. Blanchard worked with Dr. Walther Mothes, a HIV specialist at the Yale University School of Medicine, and with Dr. James Munro, who was Dr. Blanchard's first graduate student and who is now an assistant professor at Tufts University School of Medicine. Drs. Mothes and Munro are the two other co-lead investigators.

Dr. Blanchard adapted an imaging technique that uses fluorescence to measure distance on molecular scale – single-molecule fluorescence resonance energy transfer (smFRET) imaging – to study viral particles. His group developed fluorescent molecules (fluorophores)—which he dubs "beacons"—and the team inserted them into the virus's outer covering, known as the envelope. With two of these special beacons in place, smFRET imaging can be used to visualize how the molecules move over time, when the virus proteins change conformation. The approach provides a measure of distances, on the order of a billionth of an inch, between two beacons glowing in different colors, and this can be used to detect shape shifting as it occurs and the attached beacons move.

The team used the technology to study motions of proteins on the surface of the HIV virus (called envelope proteins) that are key to the virus's ability to infect [human immune cells](#) carrying CD4 receptor proteins. (CD4 receptor proteins help HIV bind to a cell.) The envelope consists of three gp120 and gp41 proteins positioned close together, and referred to as "trimers," that open up like a flower in the presence of CD4, exposing the gp41 subunit that is essential for subsequent aspects of the mechanism that causes infection.

"There are 10-20 such envelope trimers on the surface of each HIV particle, and they mutate rapidly, thereby evading typical immune responses. This is why it is so difficult for humans to mount an effective immune response and why it is challenging for researchers to develop vaccines targeting the HIV envelope proteins," Dr. Blanchard says. The researchers were able to study proteins from two different strains of

HIV, which contained beacons that did not alter the biology of the particles.

Then they watched.

They saw that the gp120 proteins' virus particles changed shape constantly and that the timing and nature of their movements were both similar and distinct. "This answered the first big question of how opening of the envelope trimer is triggered," Dr. Blanchard says. "Many scientists believe that the particles remain in one conformation until they come across a CD4-positive cell. But we saw that the proteins dance when no CD4 was present—they change shape all the time."

The researchers were then able to watch how the viruses responded when synthetic CD4 was introduced. They also saw that antibodies known to exhibit some effectiveness acted to prevent gp120 from opening, and that these effects correlated with a decrease in the virus' ability to infect cells. Similar things happened when they introduced a small molecule now under development to prevent HIV infection.

"The practical outcome from this technology is that we can begin to understand how the biological system moves. So far we have detected three different conformations of the envelope trimer. We are working now to improve the technology to achieve the imaging precision we need to make broadly effective therapies," Blanchard says.

## **Technologies Work Hand in Hand**

The *Nature* study, led by researchers at the National Institute of Allergy and Infectious Diseases, used X-ray crystallography to capture a three-dimensional structure of one of the conformations revealed in the *Science* paper. The protein constructs used in this investigation were originally developed by a team of researchers headed by Dr. John

Moore, professor of microbiology and immunology at Weill Cornell.

"The antibodies used in the crystallography study are ones that we observed to stop the dance of the HIV envelope proteins, pushing the trimer assembly into a quiescent, ground state," Dr. Blanchard says.

"This concrete, atomic resolution picture of what the pre-fusion machinery looks like and where these antibodies bind provides an important step forward to understanding HIV's biology," he says.

Dr. Blanchard believes both techniques—smFRET and X-ray crystallography—can work hand in hand to help scientists describe the functions of molecules from the perspective motion, including the other two distinct conformations identified in the smFRET study.

"The approach is really a breakthrough for science because most research is done in a test tube where billions of molecules are present, all behaving independently. It is very difficult to extract direct information about these types of movements from indirect observations," such as those that don't use imaging technology, he says. "The single-molecule approach allows practical, interpretable, real-time information to be obtained about molecular processes in complex biological systems."

**More information:** — Structure and immune recognition of trimeric pre-fusion HIV-1 Env, *Nature*, [DOI: 10.1038/nature13808](https://doi.org/10.1038/nature13808) (2014)

— Munro et al. Conformational dynamics of single HIV-1 envelope trimers on the surface of native virions. *Science* [DOI: 10.1126/science.1254426](https://doi.org/10.1126/science.1254426) (2014).

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