

## Imaging study of key viral structure shows how HIV drugs work at atomic level

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This illustration depicts the molecular structure of an HIV drug known as an INSTI binding to key sites on the intasome (yellow), the viral machine that allows HIV to invade cells. The red spheres represent DNA. Credit: Salk Institute

Salk scientists have discovered how a powerful class of HIV drugs binds to a key piece of HIV machinery. By solving, for the first time, three-



dimensional structures of this complex while different drugs were attached, the researchers showed what makes the therapy so potent. The work, which appeared in *Science* on January 30, 2020, provides insights that could help design or improve new treatments for HIV.

"The drugs we studied are the latest compounds available in the clinic today, as well as several important pre-clinical molecules. Until now, no one knew exactly how they bound to this HIV complex," says the study's senior author Dmitry Lyumkis, an assistant professor in Salk's Laboratory of Genetics. "A better understanding of how the drugs work will help us improve them and design new therapeutic compounds."

The intasome is a crucial structure of the virus that enables infection, composed of the HIV protein integrase and strands of viral DNA that form when the virus enters <u>human cells</u>. The intasome moves into each human cell and then carries out the chemical reactions necessary to integrate the virus' genetic material into human DNA.

Some drugs, called integrase strand transfer inhibitors (INSTIs), have managed to block the intasome; HIV can't infect human cells when the complex can't integrate viral DNA into the human genome. There are currently four INSTIs approved by the US Food and Drug Administration, as well as others under development.

Despite the success of these molecules, researchers have struggled to study how they inhibit the HIV intasome, largely due to difficulty in isolating intasomes for structural studies. In the past, most research on the intasome and INSTIs was carried out on another retrovirus called prototype foamy virus, or PFV. In 2017, Lyumkis and his colleagues were the first to determine the structure of purified HIV intasomes.

In the new work, Lyumkis' team went a step further: they obtained the structure of HIV intasomes while they were being actively blocked by



one of four INSTIs—the commercially available <u>drug</u> bictegravir or three experimental compounds known as 4f, 4d and 4c. The team used tilted single-particle cryo-electron microscopy (cryo-EM), an imaging technique they've helped optimize, to reveal the structure of each intasome-drug complex.

The first observation that Lyumkis made was just how differently the drugs attached to the HIV intasome than what had been seen with the PFV intasome. The compound known as 4f, for instance, loops backwards onto itself as it binds to the PFV intasome but remains relatively flat as it attaches to the HIV version of the complex, details which can help researchers improve the binding properties of potential future molecules.

"To this day, everyone is still using the PFV intasome structure to rationalize and understand the mechanism of action of these drugs," says Dario Passos, the study's co-first author and a staff scientist in Lyumkis' laboratory. "But we've shown that the field really needs to move and study the HIV structure if we want to make further progress."

"We and many others have been working towards this goal for several decades and it is exciting that at long last we can now understand how HIV inhibitors work in detail and aid the development of new drugs." says Min Li, co-first author and a staff scientist at the National Institute of Diabetes and Digestive and Kidney Diseases.

The structures also revealed why the drugs are so potent and what makes them so good at avoiding drug resistance. The INSTIS, Lyumkis and his colleagues found, fill the entire space that's normally occupied by DNA. That means if the HIV intasome develops a mutation that blocks the INSTI drugs from binding, it also blocks the DNA from attaching, rendering the complex useless for invading human cells.



Finally, the extremely high resolution of the structure obtained by the Salk researchers lets them see details on how the drugs chemically interacted with this binding pocket, and how INSTIs displaced water molecules to do so, which gave the team even more information on what makes INSTIs so successful in the clinic.

"In previous structures, we learned about intasome biology," says Lyumkis. "But here, we've really started to gain insight into the therapeutic angle of how drugs can target these important viral assemblies."

The researchers are planning additional work on the experimental drugs—focusing on the compound known as 4d, which, based on both preclinical testing and the new structural insight, shows more promise against HIV than other compounds. They also want to better understand what happens to the structure of the intasome in cases where it develops resistance to INSTIs. This could help them design more efficient drugs in the future, says Lyumkis.

**More information:** "Structural basis for strand transfer inhibitor binding to HIV intasomes" *Science* (2020). <u>science.sciencemag.org/lookup/ ... 1126/science.aay8015</u>

Provided by Salk Institute

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