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HIV Cure-Related Research Strategies

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Image



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Although a few people appear to have been cured of HIV under special circumstances, there is currently no general cure for this virus. See our fact sheet on [Finding a Cure for HIV](#) for information on general challenges to developing such a cure and what the research means for women. The current fact sheet provides information on what scientists are trying to do to develop a cure.

Different Kinds of Cure

There are several terms currently used in HIV cure-related research. All of them assume that a person no longer needs to take today's HIV drugs, at least for long periods of time:

- Eradication or complete elimination – getting rid of all virus from all locations in the body; sometimes referred to as "complete cure"
- Durable antiretroviral (ART)-free control – HIV may still be in the body, but it is not active; the body is not fully rid of HIV, but the virus cannot affect one's health and cannot be transmitted to others; also sometimes called "functional cure"
- Remission – a term borrowed from the cancer field, it means that HIV is not active in the body; there is no guarantee of lifelong control of the virus, and it suggests the need for continued monitoring (to make sure HIV is still inactive)

Read on for the different strategies that could be used to achieve one of these types of cure. All are still being researched – some are at more advanced research stages than others. Each strategy includes different ways to achieve the same thing – e.g., for "kick and kill," different substances could be used to "awaken" the hidden virus and strengthen the immune system. There are also challenges with each of these strategies, which are explained below.

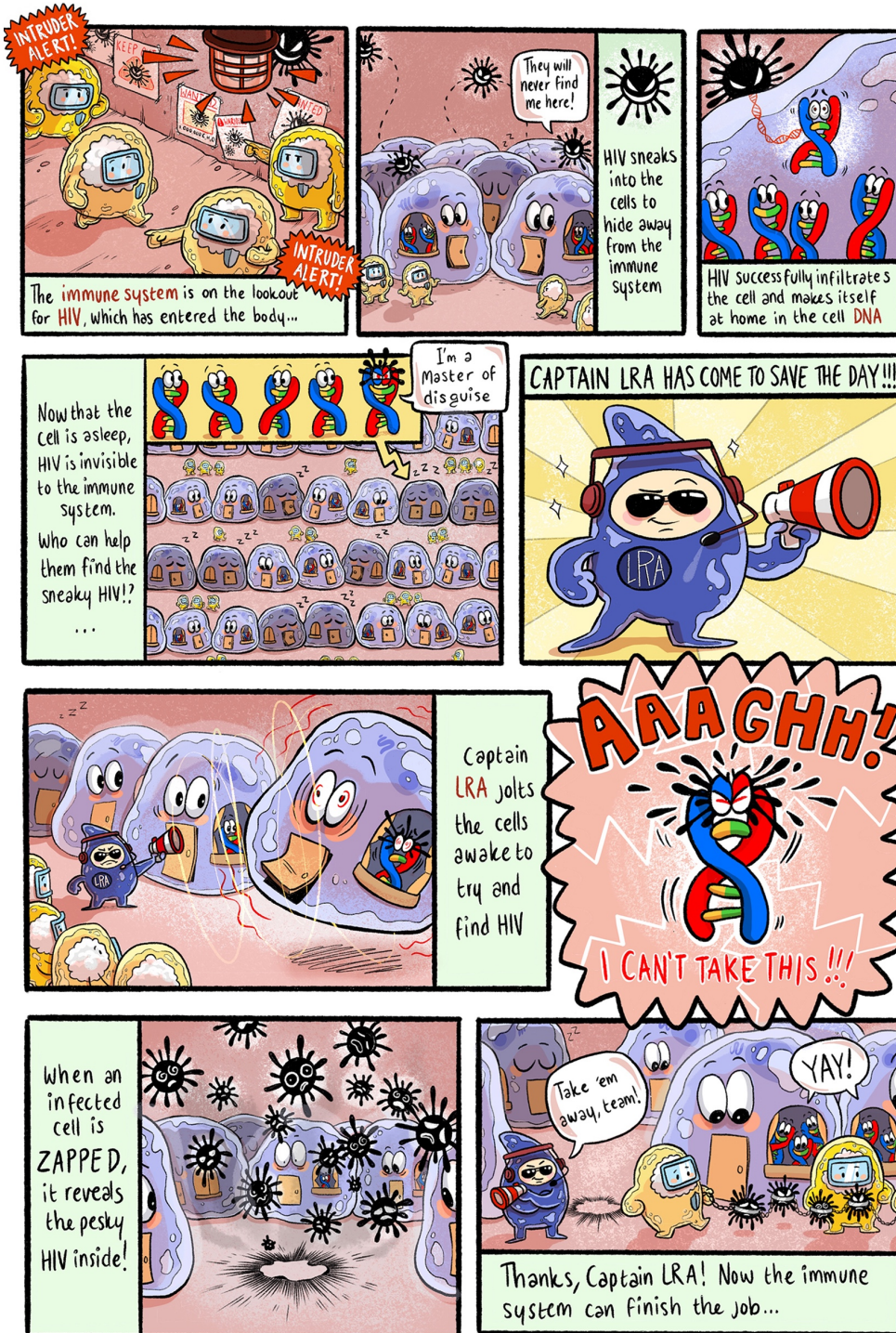
Latency Reversing Agents ("Kick and Kill")

Also called "shock and kill" (or, more gently, "poke and clear"), the game plan here is to "kick" or "poke" the resting cells in the reservoirs into action, then "kill" or "clear" the newly activated cells when HIV returns to the bloodstream. Once the cells become active, they are no longer hidden from the immune system. The substances that provide the kick are called latency reversing agents, as they interrupt HIV's ability to remain inactive within cells.

At the same time, regular antiretroviral therapy would prevent uninfected cells from becoming infected with the newly active virus that has been kicked into action. Ideally, this strategy would empty the reservoirs and thereby rid the body of infection. Latency reversing agents would need to be taken along with strategies that enhance the immune system.

Image

CAPTAIN LRA SAVES THE DAY!



KEY

Immune System

a System of cells, tissues and organs within the body that help fight off infections & diseases.



HIV

(Human Immunodeficiency Virus)
A virus that enters the body and attacks cells that help the body to fight off infections, making the body highly susceptible to diseases and infections



DNA

Genetic material found in all living organisms that contains the main constituent of chromosomes. It is self-multiplying and contains all genetic information.



LRA

Pharmacological approach to eliminating the HIV reservoirs. This strategy attempts to flush the virus out of the resting cells by reawakening the dormant viruses in the latent reservoirs.



Story by: Eric Lee, Matylda Mai & Jazmin Guzman
(Pencils) (Inks & Lettering) (colors)



UCSF Center for AIDS
Prevention Studies
Division of Prevention Sciences

Challenges with this approach:

- Finding substances (latency reversing agents) that will safely and effectively activate, or kick the resting cells into action
- Ensuring that every cell infected with HIV is reactivated when kicked, leaving no infected cell untouched
- Ensuring that infected cells that are kicked into action to start producing virus end up dying (i.e., the "kill" part of kick and kill); approaches to bolstering the immune system's ability to recognize and kill the newly activated cells are listed below.

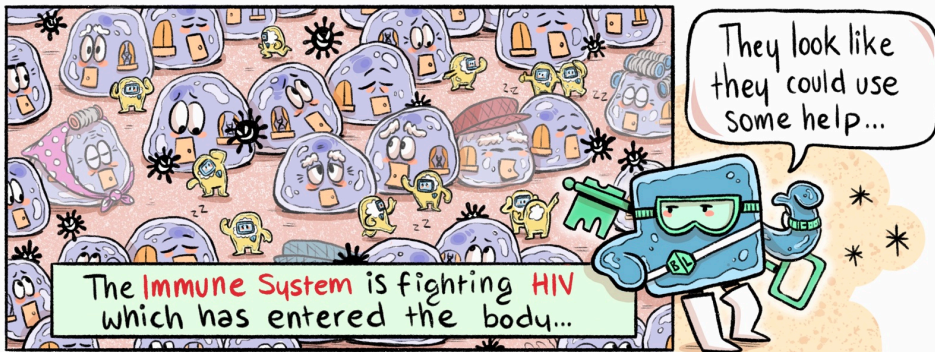
Block and Lock Strategies

Rather than activating and clearing off cells as with latency reversal strategies, the "block and lock" approach aims to permanently silence latent virus using latency promoting agents (LPAs) to "block" the transcription step in the cell's [lifecycle](#). Over time, this would "lock" the part of the cell or the gene responsible for HIV transcription into a deep latent state. Permanent control of that part of the cell means HIV drugs would no longer be necessary. Another key aspect of these "block and lock" approaches is that the agent would be specific to HIV (e.g., Tat gene), so it would not block other processes needed to overcome other infections.

Studies in humans will be needed to test whether this kind of approach could work to achieve treatment-free durable suppression. However, unlike latency reversal approaches that are studied in HIV after being used to manage other health conditions, no block and lock strategy has been approved by the US Food and Drug Administration (FDA). Therefore, clinical trials of these therapies have lagged behind.

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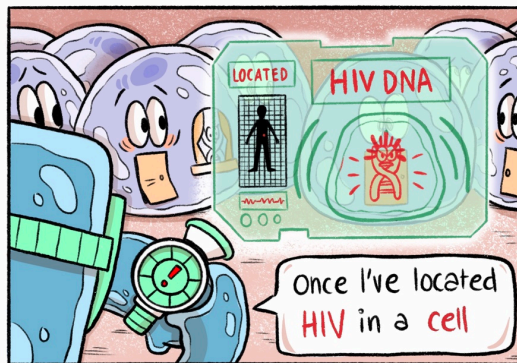
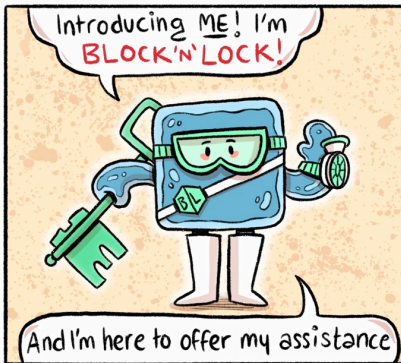
AGENT BLOCK 'N' LOCK



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A system of cells, tissues and organs within the body that help fight off infections and diseases

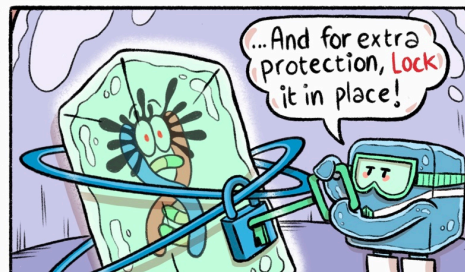
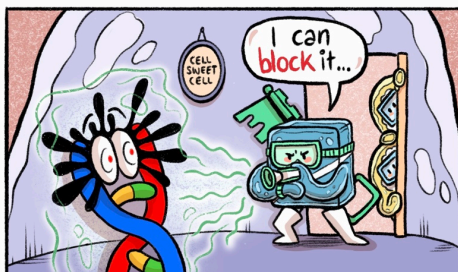


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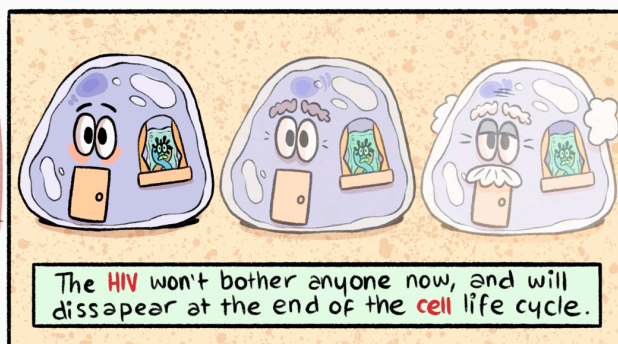


Latently-Infected Cell

A cell that is affected by the HIV but not actively producing the virus. It's hard for the immune system cell to recognize it as an affected cell because of its inactivity.

Block and Lock Strategy

A strategy that targets and silences the HIV virus DNA in the latently-infected cell. The cell can return to its normal activity but the viral DNA stays silent.



Story by: Eric Lee, Matyllda Mai & Jazmin Guzman
(Pencils) (Inks & Lettering) (Colors)

Gene Therapy

There are three broad approaches to gene therapy:

- knocking *out* genes in the virus that allow it to enter and infect immune cells;
- knocking *in* genes to our immune cells that make them resistant to infection; and
- cutting out the genetic pieces of HIV that have become integrated into the DNA of infected immune cells.

The first strategy looks to disable HIV and make it unable to enter cells in your body by editing the genetic code of the virus. The genes that code for or provide the manufacturing instructions for HIV's ability to enter cells would be deleted or "knocked out." Thus, HIV would remain in the body, but it would be unable to infect cells in your or other people's bodies.

The second approach involves adding, or "knocking in" genes to a person's immune cells that would protect them against HIV. We know what those protective genes look like because some people are naturally born with them. These people are protected by their inability to produce a receptor called CCR5 on the outside of their immune cells that HIV needs to enter and infect the cells.

One popular example of this approach is Timothy Brown, once known as the "Berlin patient." He received a stem cell transplant after he was diagnosed with leukemia (a form of blood cancer). Stem cells are cells that have not yet received instructions from the body to make them into a specific type of cell. Stem cells can renew themselves, develop and grow into several different types of cells, including several different kinds of immune cells.

The stem cells Brown received were from a person who was naturally protected from HIV by having genes that lacked the ability to produce a functional CCR5 receptor, one of two T-cell receptors that HIV needs to bind to in order to enter and infect cells. Brown was considered cured of HIV, though he died of cancer in September 2020.

A similar procedure appears to have been successful in two other people. The first one is known as the "London patient" – a man named Adam Castillejo. The second person, called the "Dusseldorf patient," also received a bone marrow transplant to treat leukemia. He was declared cured of HIV in 2023.

In early 2022, researchers announced that a fourth person – and the first woman – has reached ART-free control of HIV following a procedure. The "New York patient," who at this point wishes to remain anonymous, is of mixed race and was also diagnosed with leukemia. She underwent a transplant using stem cells from umbilical cord blood, but from an individual with a CCR5 mutation like in the three previous cases. She remains in remission without HIV medications at the time of this writing.

In mid-2022, a potential cure of an older person, the "City of Hope patient," was announced. He had lived with HIV for a long time, and after a stem cell transplant has been free of HIV without treatment since 2021.

Two other people living with HIV and cancer who received stem cell transplants to treat their cancer are commonly referred to as the "Boston patients." These people stopped taking HIV drugs for a couple of years after their transplants and were able to go for several weeks or months without the virus reappearing. Unlike Timothy Brown, however, they did not receive donor stem cells from a person resistant to HIV. As a result, although they appeared to have no evidence of HIV after their transplants,

the virus ultimately returned in both of them after they stopped taking their HIV drugs.

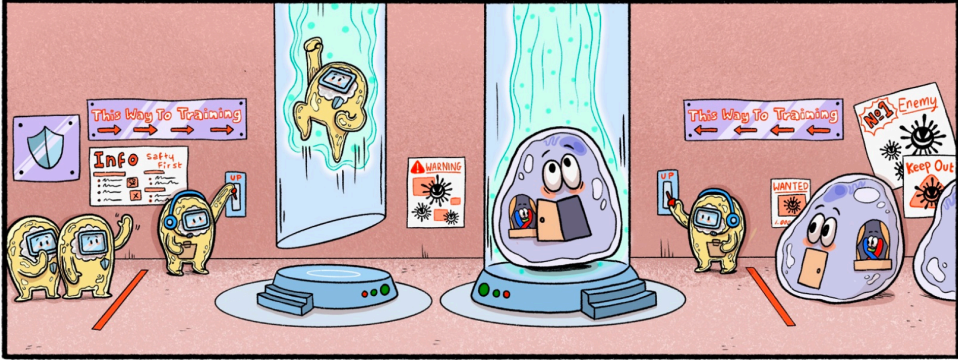
Another person with cancer who had a stem cell transplant, the "Geneva patient," has remained without the virus after stopping ART. He had started HIV treatment early and had an undetectable viral load for a long time before receiving a transplant from a donor who did not have the HIV-protective CCR5 mutation. His viral load remains undetectable, but very small amounts of defective virus remain. Researchers therefore call this a case of HIV remission, not cure. A similar approach was used with another person, the "second Berlin patient" whose HIV also remains in remission.

The third gene therapy approach uses relatively new technologies (e.g., CRISPR) that are able to latch precisely onto the HIV genes that have integrated into human DNA and cut them out without harming the cell or causing it to malfunction.

Image

IMMUNOTEAM: DEFEND 'N' ASSIST

The Immune System and cells are being taken to a training school to learn how to defend the body against HIV...



=KEY=

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Gene Editing

There are two main forms:

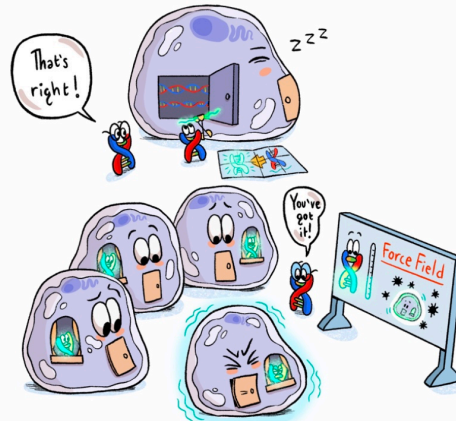
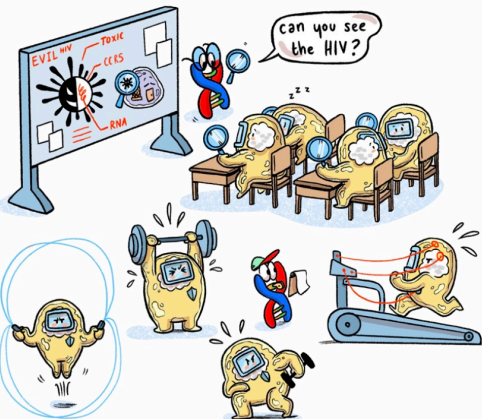
1. Cells are taken out of the body to have some of their genetic characteristics modified.
2. Genes in the cells are modified while they are still inside the body.

Goal: To make specific cells resistant to or better at fighting HIV, or to change the HIV itself so it becomes ineffective.

Gene Direct Approach

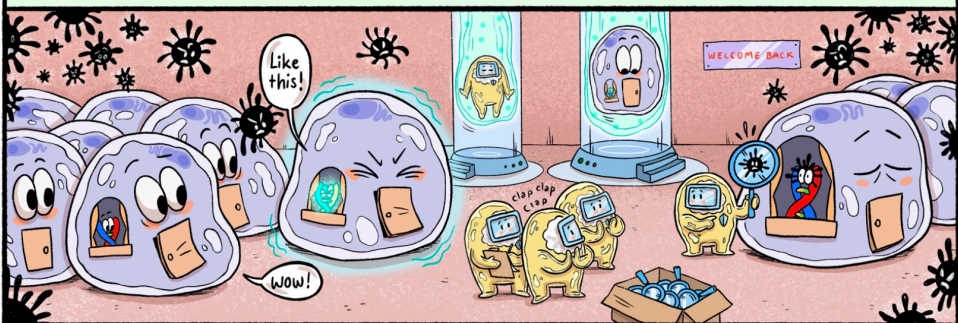
- (Immune system cell)
To make the immune system better at locating and fighting HIV
- (non-immune system cell)
To make non-immune system cells resistant to HIV entry

The Immune system learns how to locate and protect against HIV...



The other cells learn how to shield themselves against HIV

After their training, they are taken back to teach and assist the other cells how to find, protect and defend against HIV.



Story by: Eric Lee, Matyllda Mai & Jazmin Guzman
(Pencils) (Inks & Lettering) (colors)

Challenges with these approaches:

- Changing the genetic sequence or code can produce unexpected results, including unintended side effects
- Stem cell therapies require wiping out a person's existing immune cells. This process can involve several types of drugs as well as radiation to create a 'clean slate' for the newly transplanted stem cells to flourish and grow. It is a lengthy, uncomfortable, and dangerous process. In addition, there are few individuals naturally immune to HIV (few potential stem cell donors), and the process is very costly.
- Virus-removal techniques would need to be highly specific (removing only HIV genetic material and not human genetic material) and highly sensitive (able to find almost all the infected cells)

Immune-Based Approaches

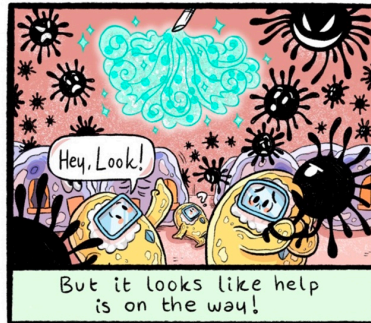
Immune-based approaches work by enhancing the immune system's ability to clear infected cells and achieve a cure. Scientists are looking at three main approaches:

- Broadly neutralizing antibodies (bNAbs): When our immune cells attack and destroy invaders like HIV, they display pieces of the virus — known as antigens (from antibody-generating) — on their surfaces. An antibody is a protein that attaches to an antigen like a key fits a lock. When an antibody has matched up with an antigen, it has marked the intruder for destruction by immune cells. Broadly neutralizing HIV antibodies can recognize and target for destruction several different strains of HIV, unlike standard antibodies, which are usually only able to latch onto antigens from a single strain of the virus.
- Natural killer (NK) cells and interferon-gamma: NK cells destroy infected cells and are an important part of the body's early response to viral infections. They clear infected cells while the body is prodding killer T cells into action. Unlike some immune cells, NK cells do not need to be infected by HIV in order to effectively identify or clear HIV. Scientists are hoping that this new understanding of how NK cells work can lead to a therapeutic vaccine or durable ART-free control of HIV.
- Chimeric antigen receptor (CAR) T-cell therapy: In this approach, the T cells of a person living with HIV would be engineered to have receptors that reprogram those cells to identify and eliminate cells infected with HIV. The FDA has approved this therapy in a certain type of leukemia, and it is being studied in other kinds of cancers. There have been promising studies in animal models. Clinical trials exploring CAR T cells as an approach to curing HIV have begun in humans.

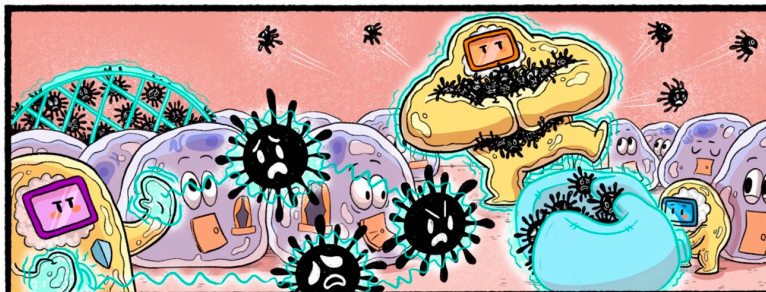
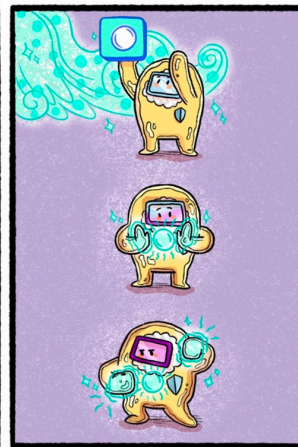
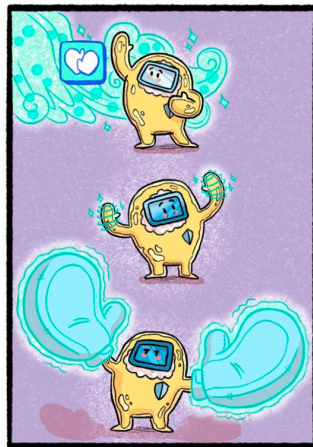
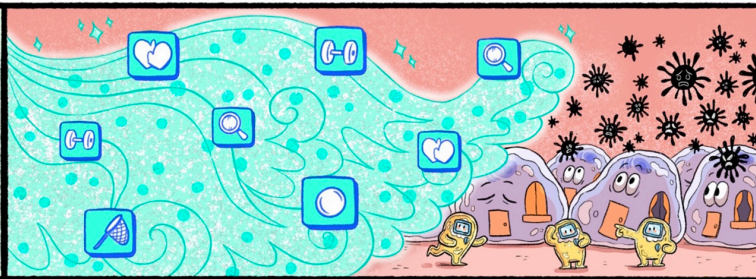
About 0.5 percent of people living with HIV around the world who acquired HIV continue to have low viral loads and high CD4 counts without ever taking HIV drugs. They are called "elite controllers." One of these is Loreen Willenberg. After acquiring HIV in 1992, but not progressing to worse HIV disease or AIDS, she participated in research studies for many years, hoping that a path to an HIV cure might include what scientists could learn from her unique cells. In 2011, researchers realized they could not find any virus inside Willenberg's body that could make copies of (replicate) itself – including in reservoirs. The virus had entirely cleared from her body by itself. In 2021, this was found to have occurred with a second woman in Argentina, known as the "Esperanza patient." Now, the job of scientists is to figure out how to repeat the process in other people. These cases are inspiring research efforts towards "block and lock" approaches (see above) to keep HIV from replicating inside the body.

Image

IMMUNOTEAM: POWER UP!



The body has been injected in an effort to help the immune system control the spread of HIV more effectively.



Thanks to the injection, the immune system is now stronger and more able to protect the body against HIV.

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Immune Base Strategy

Designed to boost an immune response against HIV in someone who already has the virus.



Killer T-Cell

An immune system cell that can clear certain cells. The injection gives the killer T-Cell more strength to fight off the HIV

Helper T-Cell

An immune system cell that activates the majority of the immune system. The injection gives the helper T-cell better tools to fight the virus.

B-Cell

An immune system cell that makes antibodies and develops from the stem cells in the bone marrow. The injection gives the B-Cell a better way to create the antibodies to help the immune system find HIV

Story by: Eric Lee, Matyllda Mai & Jazmin Guzman
(Pencils) (Inks & Lettering) (Colors)

Challenges with immune-based approaches:

- Both broadly neutralizing antibodies and "post-treatment controller" NK cells are hard to find. They occur in only a small minority of people and may not be able to get into all parts of the body where HIV hides.
- Stimulating the immune system can increase the number of cells that HIV can target for infection
- There are many different strains of HIV, and HIV mutates, or changes, very rapidly. This may make even broadly neutralizing antibodies ineffective over time. Experts believe people may need to receive a combination of broadly neutralizing antibodies just as today's effective HIV treatment regimens include multiple drugs.

Combination Approaches

Most likely, we will need a combination of approaches to achieve ongoing control of HIV without HIV drugs. Scientists are investigating safe ways to combine different HIV cure research approaches.

The Well Project would like to thank Karine Dubé, DrPH (UNC-Chapel Hill) and Danielle M. Campbell, MPH (Charles R. Drew University) for thorough reviews and updates of the fact sheet content in 2022; as well as Rowena Johnston, PhD (amfAR) and David Evans (Project Inform) for their assistance authoring the original edition of this fact sheet.

Additional Resources

Select the links below for additional material related to HIV Cure-Related Research Strategies.

- [Are Scientists Getting Closer to an HIV Cure? \(The Body.com\)](#)
- [Research Toward a Cure Trials \(Treatment Action Group\)](#)
- [HIV Cure Research Illustrations \(University of California-San Francisco\)](#)
- [#Cure \(POZ\)](#)
- [5 Strategies to Cure HIV \(Desmond Tutu Health Foundation\)](#)
- [Meet the London Patient: Adam Castillejo \(amfAR\)](#)
- [A Third Person Living With HIV Has Been Cured by Transplant \(Medscape\)](#)
- [The Often-Overlooked Düsseldorf Patient \(CATIE\)](#)
- [An Exception to the Rule \(POZ\)](#)
- [Research Toward HIV Cure \(US National Institutes of Health\)](#)
- [HIV Cure Strategies \(Weill Cornell Medicine\)](#)
- [HIV Cure Research Lacks Diversity From the Earliest Stages, Scientists Warn \(Th...](#)
- [HIV Cure Research Information Sheet \(Treatment Action Group\)](#)
- [We Still Have No HIV Cure. But We Have a Better Strategy for Finding One \(aidsm...](#)
- [New HIV Remission Case: 'Geneva Patient' Is a Potential Leap for Stem Cell Tran...](#)
- [A seventh case of HIV cure reported at AIDS 2024 \(World Health Organization\)](#)



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