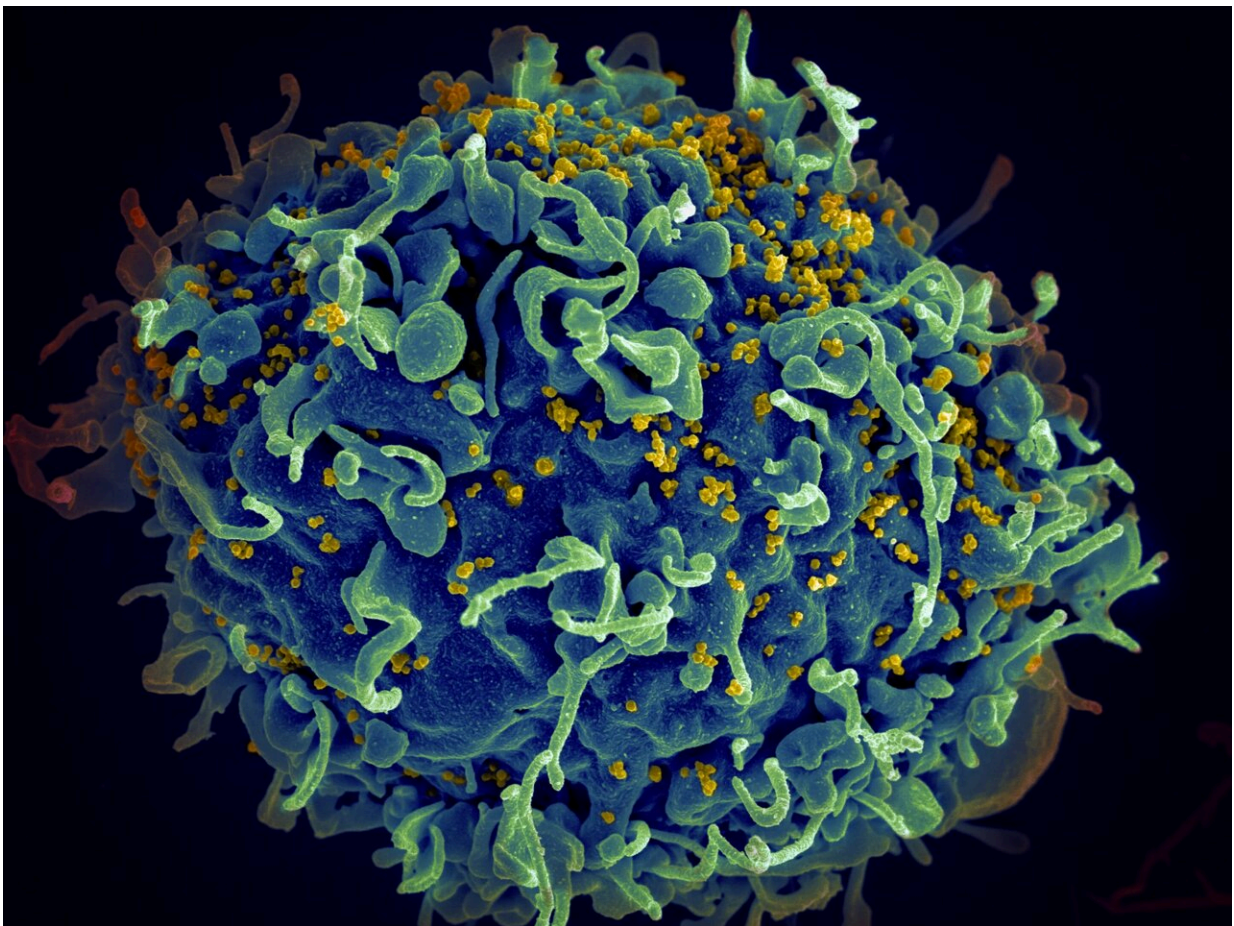


Longer-lasting HIV antibody treatments could offer a one-shot alternative to maintenance therapies

September 26 2022, by Ronald C. Desrosiers



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[Antiretroviral therapy](#) has had an enormous impact on treating HIV infections around the world. The [millions of people](#) currently taking these treatments under medical supervision can reasonably expect to reduce their viral loads to [undetectable levels](#), eliminate the risk of transmission and live a normal life span. However, antiretroviral therapy is not without shortcomings. People need to take these medications regularly for life, and [low compliance](#) can lead to drug resistance.

There is a promising new option on the horizon. I am a [researcher who studies AIDS treatments](#), and I believe that [monoclonal antibodies](#) could become game-changers for the treatment of HIV infections.

HIV presents challenges to antibodies

[Antibodies](#) are proteins that serve as major players in the immune system's response to pathogens, which cause disease, and allergens, which cause allergic reactions. Antibodies recognize specific markers, or antigens, on a potentially harmful substance and help the body eliminate it.

Over the past few decades, researchers have been able to isolate individual antibodies specific to the individual pathogen or allergen they are meant to attack. With this advance, [monoclonal antibodies](#) made in the lab have become a [major segment of the pharmaceutical industry](#). You can see numerous ads on TV or in magazines promoting [monoclonal antibodies](#) to treat osteoporosis, autoimmune disorders and various types of cancers.

Antibodies can also be used to treat [viral infections](#), including [COVID-19](#). But using antibodies gets more complicated with HIV, the virus that causes AIDS in people.

One reason is that HIV has an [enormous number of variants](#) circulating

across the world and even within a single infected individual. In fact, the genetic variation of HIV within a single patient exceeds the genetic variation of all circulating influenza strains worldwide during an entire flu season.

The immune system of an individual infected with HIV creates antibodies to neutralize the virus. However, because these antibodies can usually recognize only one particular strain, they are unable to neutralize other HIV strains circulating in the population. Furthermore, HIV can [mutate within an infected individual](#) and escape antibodies specific to the variant causing the original infection.

This ability to mutate and escape ongoing immune responses is a critical factor in the virus's ability to continuously replicate, a hallmark of AIDS. It also makes it difficult to design an antibody treatment that can account for HIV's enormous genetic variability.

Broadly neutralizing antibodies show promise

The discovery of rare individuals who make anti-HIV antibodies that can be effective against [up to 80% of circulating strains](#), however, has boosted prospects for antibody treatments for HIV.

These [broadly neutralizing antibodies](#), or bnAbs, have seen impressive results. [Monkey studies](#) have found that a single administration of bnAbs can prevent infection from SHIV, the nonhuman primate version of HIV. One study found that [two broadly neutralizing antibodies](#) were able to reduce [viral loads](#) to undetectable levels in infected monkeys.

In people, one study administering [two bnAbs](#) also saw suppression of HIV replication and nearly undetectable viral loads. One [early-phase clinical trial](#) in 2021 showed that one bnAb could potentially offer protection against HIV infection.

Long-term production of antibodies

All the monkey and [human studies](#) mentioned above required re-administering the broadly neutralizing antibodies every three weeks or so to maintain effective concentrations. This runs into the same problem antiretroviral therapies face in terms of requiring the individual to retake the drug frequently for life. But researchers have found a potential solution.

Using a small virus that doesn't cause disease, called an [adeno-associated virus](#), to deliver broadly neutralizing antibodies into the body can stimulate [muscle cells](#) to continually produce these antibodies. Because muscle cells have a [prolonged life span](#) and can last on average 10 to 16 years, they can be turned into factories that produce the antibodies essentially for life.

One study my colleagues and I conducted using adeno-associated virus found that one monkey was able to produce these antibodies for [over six years](#) after a single injection.

Another monkey that researchers dubbed the "[The Miami Monkey](#)" is considered functionally cured, meaning its viral loads have been at undetectable levels for prolonged periods even without continuous antiviral drug therapy. [Two other monkeys](#) have also been cured of their AIDS virus infections with this approach.

Adeno-associated virus vectors for HIV antibody therapies still face one more hurdle: [anti-drug antibodies](#), or antibodies the body produces in response to the antibodies in the treatment. Anti-drug antibodies can result when the body registers an antibody treatment as foreign and mounts an immune response against it, negating the treatment. They have also have caused problems for antibody treatments in [cancer](#) and [autoimmune disorders](#). That may especially be the case for broadly

neutralizing [antibodies](#), which have unusual structures that deviate from what the body normally expects an antibody to look like.

Researchers are working hard to develop simple and accessible approaches to help patients build tolerance to [broadly neutralizing antibodies](#). Some of these approaches include delivering treatments to other areas that have greater immune tolerance than the muscle, such as [to the liver](#) and [through the mouth](#).

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