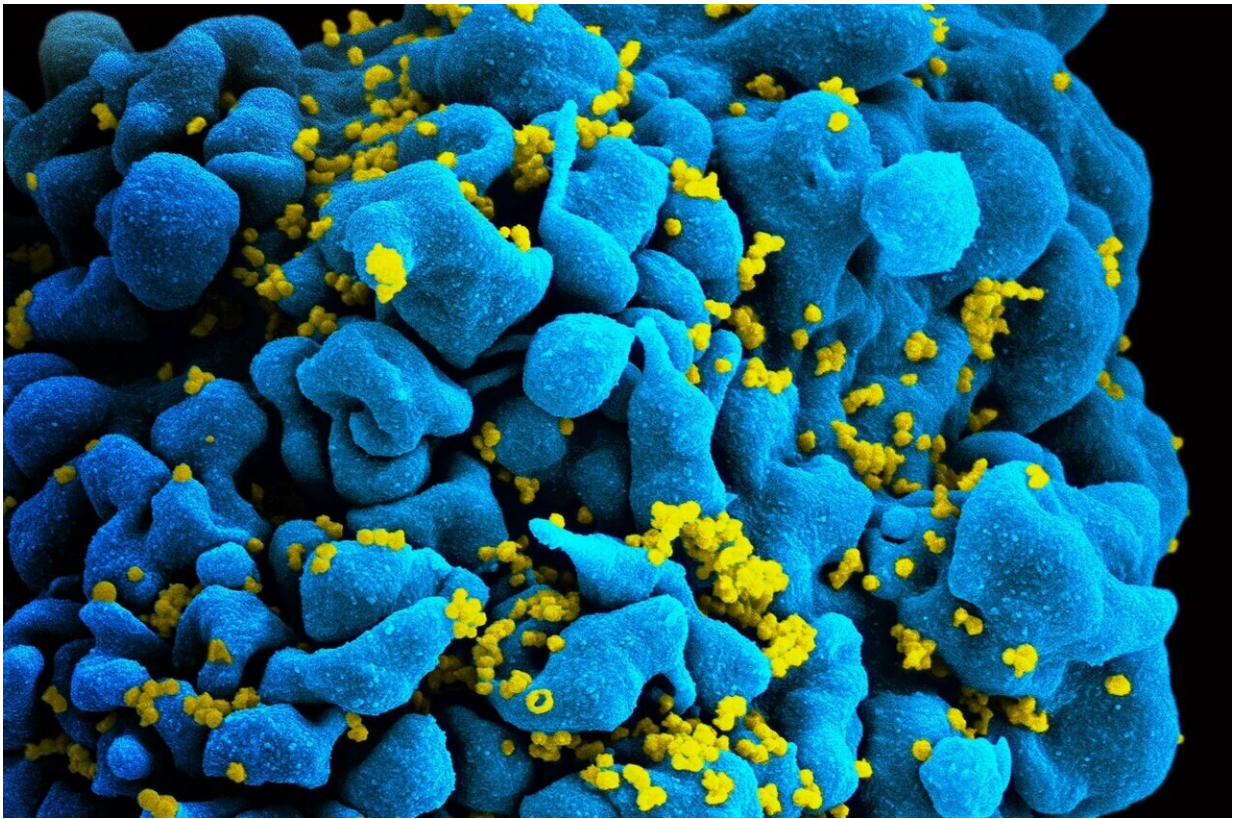


Using anti-cancer immunotherapy to fight HIV

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Scanning electromicrograph of an HIV-infected T cell. Credit: NIAID

Researchers at the University of Montreal Hospital Research Centre (CRCHUM) have shown that immunotherapy treatments against cancer could reduce the amount of virus that persists in people on triple therapy.

In a study published in the journal *Nature Communications*, they show, in the cells of people living with HIV, how these therapies reveal the virus—until now hidden in the hollows of infected cells—to the immune system.

"We identified the mechanism by which anti-cancer immunotherapy 'awakens' the [virus](#) from its hiding places and reduces the size of HIV reservoirs in people on triple therapy. Although most of our experiments have been performed in vitro, our approach could lead to the development of new therapies," stated Nicolas Chomont, a CRCHUM researcher and Université de Montréal professor.

HIV reservoirs are cells and tissue in which the virus persists despite triple therapy. This treatment prevents the infection from developing into acquired immunodeficiency syndrome (AIDS). To survive and replicate, HIV needs to be hosted in a cell. As a rule, it infects CD4+ T lymphocytes, [white blood cells](#) responsible for activating the body's defences against infections.

The virus remains dormant in these cells and builds a reservoir that is controlled, but not eliminated by antiretroviral drugs. The subject of intense study, these reservoir cells are the last hurdle in eradicating the virus and force people living with HIV to take antiretroviral drugs for the rest of their lives.

In 2016, Rémi Fromentin, a research associate in Nicolas Chomont's laboratory, showed that the cells housing the persistent viruses have specific immunological characteristics: three proteins called PD-1, LAG-3 and TIGIT, that are frequently expressed at their surface. Today these molecules are the target of immunotherapies used to treat cancer. The researchers decided to evaluate the effect of these therapies on HIV reservoirs.

A strategy evaluated in a small number of people with HIV and cancer

"Our results prove that immunotherapies targeting molecules such as PD-1 could reduce the amount of virus persisting in people on triple [therapy](#). One of the next steps would be to combine immunotherapy with molecules that, up to now, have been ineffective in eradicating HIV reservoirs. This combination of immunotherapy and chemical molecules could 'awaken' the virus and help remove the [cells](#) infected by HIV," added Chomont.

In this article, Rémi Fromentin and Nicolas Chomont also present data from a patient in Montreal infected with HIV and treated by immunotherapy for a melanoma.

"The size of the patient's HIV reservoirs decreased significantly, which is encouraging. However, we must remain cautious, because this doesn't work with all patients. These treatments also cause considerable side effects," indicated Fromentin. The results of clinical trials currently underway in the United States on patients with cancer and HIV should help guide future research.

Nearly 37 million people around the world live with HIV. Every day, 5,000 cases are reported to global health authorities.

More information: Rémi Fromentin et al, PD-1 blockade potentiates HIV latency reversal ex vivo in CD4+ T cells from ART-suppressed individuals, *Nature Communications* (2019). [DOI: 10.1038/s41467-019-08798-7](https://doi.org/10.1038/s41467-019-08798-7)

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