



Monthly Injectable Treats HIV as Well as Daily Meds

Given every four weeks, combo injections of cabotegravir and rilpivirine will likely face an FDA decision by early 2020.

March 8, 2019 By [Benjamin Ryan](#)

Treating HIV with monthly combination injections of long-acting formulations of ViiV Healthcare's cabotegravir and Janssen's rilpivirine (sold in daily pill form as Edurant) suppresses the virus as effectively as a standard daily oral antiretroviral (ARV) regimen. Given the success of the experimental regimen in a pair of advanced trials, the long-acting combination of cabotegravir and rilpivirine is poised to achieve approval from health regulators, including the Food and Drug Administration (FDA).

This would mark the first time an approved regimen does not require taking daily oral medications.

People with HIV who receive a prescription for an approved long-acting injectable regimen would need to receive their injections intramuscularly in a clinic, most ; they would not be able to inject themselves at home.

Susan Swindells, MBBS, of the University of Nebraska Medical Center in Omaha, presented findings from the Phase III open-label, multicenter ATLAS study at the 2019 Conference on Retroviruses and Opportunistic Infections (CROI) in Seattle. Participants started the trial with HIV fully suppressed by daily oral ARVs and were randomized to stay on their initial regimen or switch to monthly long-acting cabotegravir and rilpivirine.

Susan Swindells of the University of Nebraska Medical Center at CROI 2019 Benjamin Ryan

Findings from the Phase III [FLAIR](#) study of people starting their first ARV regimen with either the long-acting injectable regimen or a daily oral regimen were also presented at CROI. That trial reached findings comparable to the ATLAS study.

These studies carried the torch from the Phase IIb [LATTE-2](#) study, which established that a four-week dosing interval, as opposed to the eight-week interval that was also tested, was preferred for long-acting injectable cabotegravir and Edurant.

All 616 HIV-positive participants enrolled in the ATLAS study had had a fully suppressed viral load for at least six months prior to entering the study and had no history of virologic failure. They were all initially taking daily oral ARV regimens of two nucleoside/nucleotide reverse transcriptase inhibitors (NRTIs) plus one integrase inhibitor, non-nucleoside reverse transcriptase inhibitor (NNRTI) or protease inhibitor.

The median age of participants was 42 years old. Twenty-six percent were 50 years old or older, 33 percent were female, 68 percent were white and 23 percent were Black.

At the study's baseline, in addition to taking two NRTIs, 50 percent of the participants took an NNRTI, 33 percent took an integrase inhibitor and 17 percent took a protease inhibitor. They had been on ARVs for an average of about four years.

The participants were randomly assigned to two even groups. The members of one group stayed on their daily oral ARV regimen. The other group received a daily oral formulation of 30 milligrams of cabotegravir and 25 mg of Edurant for four weeks, during which time they were monitored to ensure that the regimen was safe.

Then the participants in the injectable regimen group received a loading dose of 600 mg of long-acting cabotegravir and 900 mg of long-acting rilpivirine through intramuscular injection. Four weeks later they began a regimen of injections of 400 mg of long-acting cabotegravir and 600 mg of long-acting rilpivirine given every four weeks. (They could receive the injections up to one week before or after the four-week point, a protocol that Swindells anticipated would likely make its way into the drug label of an approved regimen.)

At week 48 of the study, five participants (1.6 percent) in the long-acting treatment arm of the study and three people (1 percent) in the daily oral ARV arm had a viral load of 50 or greater. Consequently, the study authors concluded that the long-acting regimen was noninferior to, or as effective as, standard daily or ARV regimens.

The long-acting regimen was also deemed noninferior to the daily regimen as a result of the fact that a respective 92.5 percent and 95.5 percent of each group had a viral load below 50 at the 48-week point of the study.

The FLAIR study also found that the long-acting regimen was noninferior to daily oral ARVs.

In ATLAS, three people in the long-acting arm and four in the daily-drug arm experienced virologic failure, defined as a viral load of 200 or greater according to two consecutive tests. All were found to have ARV drug resistance that predated their entry into the study.

Among those who received the injectable treatment, 231 participants (75 percent) experienced pain at the injection site. Four participants (1 percent) withdrew from the study because of such pain.

In the long-acting and daily-drug arms, a respective 35 (11 percent) and 23 (7 percent) of participants experienced grade 3 or 4 adverse health events and a respective 13 (4 percent) and 14 (5 percent) experienced serious adverse events.

An analysis of the metabolism of the injectable treatment indicated it led to drug levels in the body comparable to those seen with standard daily oral therapy.

One person died in the daily-drug arm.

Of the 273 people in the long-acting treatment arm who completed a questionnaire at week 48, 266 (97 percent) preferred their new regimen over their old daily regimen.

"It has surprised me how much people do like it," Swindells said of the long-acting regimen. "They just love not having to think about it," she continued, referring to participants' freedom from

adhering to a daily oral regimen.

The FDA has already given the long-acting regimen a fast-track designation. According to a Janssen representative, the company plans to apply for approval of the regimen in the United States and the European Union later in the year. Consequently, the FDA will likely issue a decision on the regimen by early 2020.

To read the ATLAS abstracts, [click here](#).

To read a press release about the ATLAS and FLAIR studies, [click here](#).

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