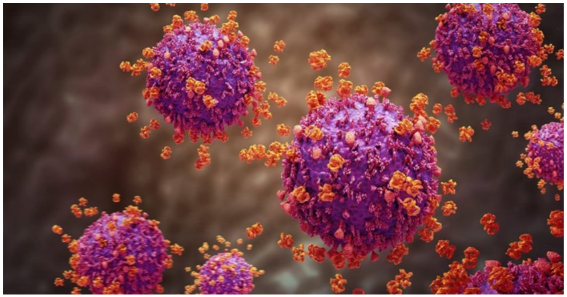
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Genome sequencing reveals new HIV strain

Next-generation genome sequencing has played a significant role in identifying a new subtype of the human immunodeficiency virus (HIV), HIV-1 Group M, subtype L, a new <u>study</u> shows.



ΗV

Since the beginning of the <u>global AIDS pandemic</u>, 75-million people have been infected with HIV and 37.9-million people today are living with the virus.

Thanks to the work done by the global health community over the past few decades, the goal of ending the HIV pandemic is becoming feasible. Yet researchers must remain vigilant to monitor for new strains to make sure testing and treatments continue to work.

"In an increasingly connected world, we can no longer think of viruses being contained to one location," said Carole McArthur, professor in the departments of oral and craniofacial sciences, University of Missouri — Kansas City, and one of the study authors.

"This discovery reminds us that to end the HIV pandemic, we must continue to outthink this continuously changing virus and use the latest advancements in technology and resources to monitor its evolution."

This research marks the first time a new subtype of "Group M" HIV virus has been identified since guidelines for classifying

new strains of HIV were established in 2000. Group M viruses are responsible for the global pandemic, which can be traced back to the Democratic Republic of Congo (DRC) in sub-Saharan Africa.

The science behind genetic sequencing to discover new viruses

To determine whether an unusual virus is in fact a new HIV subtype, three cases must be discovered independently.

The first two samples of this subtype were discovered in DRC in the 1980s and the 1990s. The third, collected in 2001, was difficult to sequence at that time because of the amount of virus in the sample and the existing technology. Today, next-generation sequencing technology allows researchers to build an entire genome at higher speeds and lower costs. In order to utilise this technology, scientists had to develop and apply new techniques to help narrow in on the virus portion of the sample to fully sequence and complete the genome.

"Identifying new viruses such as this one is like searching for a needle in a haystack," said Mary Rodgers, a principal scientist and head of the Global Viral Surveillance Program, Diagnostics, Abbott, and one of the study authors.

"By advancing our techniques and using next generation sequencing technology, we are pulling the needle out with a magnet. This scientific discovery can help us ensure we are stopping new pandemics in their tracks."

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