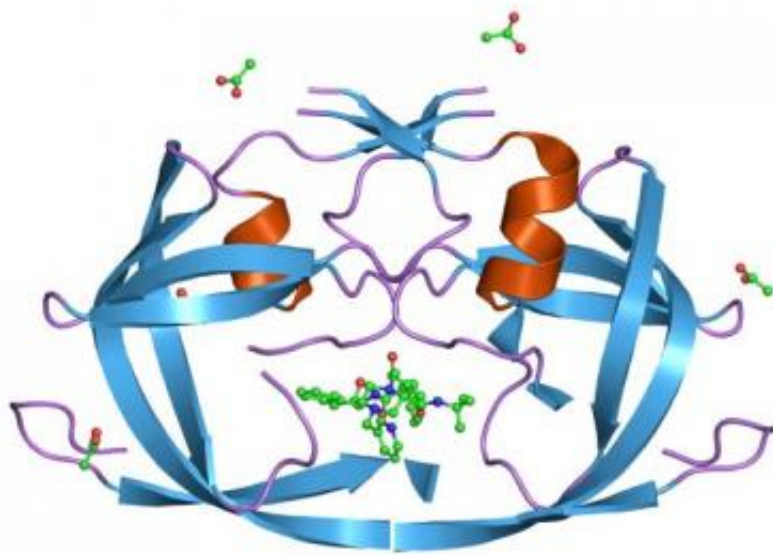


Gene discovery in monkeys could shed light on defense against HIV

November 1 2017, by Laura Otto



Diagrammatical representation of the molecular structure of HIV-1 protease complexed with the inhibitor indinavir. Credit: Public Domain

Vervet monkeys found in Africa and the Caribbean are the most abundant natural hosts of the simian immunodeficiency virus (SIV), a close relative of HIV/AIDS.

Unlike humans with HIV however, vervets infected with SIV rarely get sick with simian AIDS.

In a new study, a research group that includes UWM anthropologist Trudy Turner found that the animals' curious tolerance comes from evolution that has favored certain genes in their DNA.

This discovery of a possible defense mechanism against the [virus](#), described in a paper published Oct. 30 in *Nature Genetics*, could offer clues for developing new treatments for HIV/AIDS and could inform the future evolution of genes involved in HIV.

Vervet [monkeys](#), also called African green monkeys, have long been used as biomedical models for a variety of human diseases.

While it's a long way from monkeys to humans, Turner said, this work has comparative value. For example, it could help scientists determine whether HIV and SIV share a common ancestry, as some research has suggested.

Among the researchers' findings is that selection occurred in the animals' genes that regulate viruses, rather than fight them. And genes that govern progression of immunodeficiency were highly selected.

The team's sampling strategy was designed to investigate the selection of genes across a diverse group of the monkeys from sub-Saharan Africa to the Caribbean.

"The diversity of the sample was important so that we could determine how widespread the evolutionary selection is from a large geographical area," said Turner, professor of biological anthropology.

The animals' tolerance for SIV was the main reason an international

consortium of scientists who study the monkeys formed in 2009. The consortium had the whole genome sequenced in 2015, opening up new lines of inquiry.

"We trapped and released 2,000 animals and looked at all sorts of biological questions from that," said Turner, who directed the African fieldwork operations. "But the hot part of it was the relationship that showed up between some of the [genes](#) and SIV."

The group's discovery of genomic adaptation to viruses agrees with recent evidence that viruses played a major role in the evolution of proteins, which carry out all the functions of life, in mammals.

More information: Ancient hybridization and strong adaptation to viruses across African vervet monkey populations, *Nature Genetics* (2017). [nature.com/articles/doi:10.1038/ng.3980](https://doi.org/10.1038/ng.3980)

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