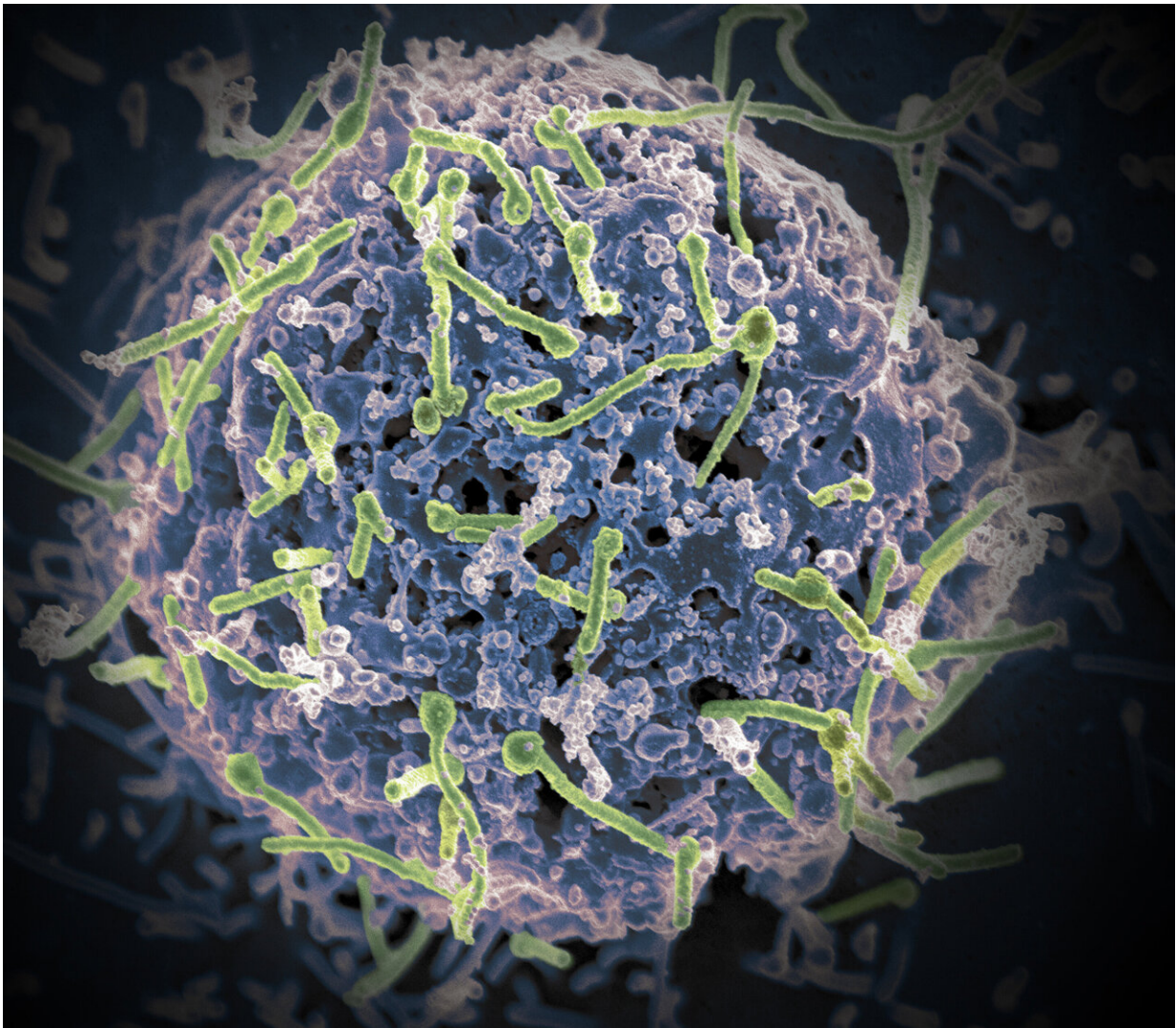


# New universal Ebola vaccine may fight all four virus species that infect humans

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The highly infectious and deadly Ebola virus is shown in this research photo from the National Institute of Allergy and Infectious Diseases. Ebola virus disease has an average fatality rate of about 50%, although rates in certain recent

outbreaks have reached close to 90%, according to the World Health Organization. Researchers report in the *Journal of Virology* early development of a potential universal vaccine for Ebola. Preclinical tests show it may neutralize all four species of the virus infecting people. Credit: NIAID

Infectious disease scientists report early development of a potential universal vaccine for Ebola viruses that preclinical tests show might neutralize all four species of these deadly viruses infecting people in recent outbreaks, mainly in Africa.

Scientists at Cincinnati Children's Hospital Medical Center report their preclinical results in the *Journal of Virology*, published by the American Society for Microbiology.

Although still in early preclinical testing, researchers report that their data indicate that the prospective vaccine has potential to be a stand-alone protection from Ebola. It also could broaden and extend the durability of protective immunity induced by current live vaccines already being tested in clinical trials against individual Ebola virus species, said Karnail Singh, Ph.D., the study's co-principal investigator in the Division of Infectious Diseases.

"This could be a significant advancement in the global effort to prevent or manage Ebola outbreaks, especially if this vaccine used alone or in combination with another Ebola vaccine results in long-term and durable protective immunity against different Ebola viruses," Singh said.

A deadly Ebola outbreak in West Africa between 2013 and 2016 accelerated international efforts to develop vaccines for these highly infectious and harmful viruses. This led to development of recombinant Ebola vaccines in which glycoprotein from Zaire Ebola virus is

engineered into another modified live viral vector. When administered, these live vaccines induce immune responses against the Ebola glycoprotein that, in turn, protect against any subsequent attack by the Ebola virus.

Singh and colleagues report that while the live-vector vaccines are producing encouraging results in clinical trials, until the current study none of the new vaccines under development have been shown to induce immune responses that cross-react against multiple Ebola virus species that cause the deadly disease in humans.

## **A Different Approach**

The new vaccine takes a novel approach, according to the study. The researchers designed a bivalent, spherical Ebola virus-like particle (VLP) that incorporates two genetically diverse glycoproteins (one each from the Zaire Ebola virus and Sudan Ebola virus) on a spherical core.

This approach will not cause illness in the recipient as the VLPs lack the genetic material and do not multiply. The vaccine works by stimulating immune responses against Ebola that generate virus-fighting antibodies to attack the different virus species.

When the researchers administered their new Ebola VLP vaccine to appropriate animal models, it produced robust immune responses against Ebola virus species known to be pathogenic in humans.

Although the new vaccine uses glycoproteins from two Ebola virus species, Singh said it might work against all four known pathogenic Ebola viruses as responses to one of the glycoproteins generates cross-reactive responses against two other Ebola virus species.

## More Testing Needed

The researchers emphasize that extensive additional preclinical testing of the prospective Ebola VLP vaccine is needed before it could potentially be tested in [clinical trials](#).

A key collaborator on the multi-institutional study—which included the University of Cincinnati College of Medicine, the Emory University School of Medicine, and the University of Louisiana's New Iberia Research Center—was Paul Spearman, MD, Division Director of Infectious Diseases at Cincinnati Children's.

Spearman said at the moment, vaccine challenge experiments are in the planning stages. They will involve working in collaboration with an institution that has Level 4 biosafety facilities and will require additional external funding to move this promising research forward.

"If the data from those studies is equally encouraging, the vaccine should be ready to progress to generation of clinical grade material for human trials," he said.

The study was funded in part by a pilot grant to Singh and Spearman by Innovation Ventures, the technology commercialization arm of Cincinnati Children's, the Cincinnati Children's Research Foundation and support from the New Iberia Research Center, University of Louisiana at Lafayette. Partial support for the study's use of [virus](#)-like-particle (VLP) platforms to conduct Ebola [vaccine](#) research was provided by the National Institutes of Health.

**More information:** Karnail Singh et al, A Bivalent, Spherical Virus-Like Particle Vaccine Enhances Breadth of Immune Responses against Pathogenic Ebola Viruses in Rhesus Macaques, *Journal of Virology* (2020). [DOI: 10.1128/JVI.01884-19](https://doi.org/10.1128/JVI.01884-19)

Provided by Cincinnati Children's Hospital Medical Center

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