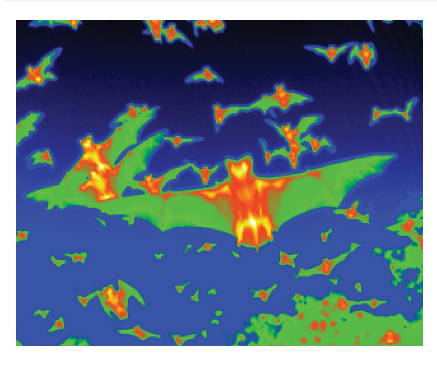


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A thermal image of bats flying in the evening in Blanco County, Texas.

PHOTOGRAPH BY NICK HRISTOV, UNIVERSITY OF CHICAGO, PBZ

SCIENCE

The controversial quest to make a 'contagious' vaccine

A new technology aims to stop wildlife from spreading Ebola, rabies, and other viruses. It could prevent the next pandemic by stopping pathogens from jumping from animals to people.

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Imagine a cure that's as contagious as the disease it fights—a vaccine that could replicate in a host's body and spread to others nearby, quickly and easily protecting a whole population from microbial attacks. That's the goal of several teams around the world who are reviving controversial research to develop self-spreading vaccines.

Their hope is to reduce infectious disease transmission among wild animals, thereby lowering the risk that harmful viruses and bacteria can jump from wildlife to humans as many experts believe happened with SARS-CoV-2, the virus that caused the COVID-19 pandemic.

The U.S. Centers for Disease Control and Prevention estimates that 60 percent of all known infectious diseases and 75 percent of new or emerging infectious diseases are zoonotic. Scientists cannot predict why, when, or how new zoonotic diseases will emerge. But when they do, these diseases are often deadly and costly to control. What's more, many researchers predict that climate change, biodiversity loss, and population growth will accelerate their spread.

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captured, vaccinated, and released. Self-spreading vaccines offer a solution.

Advances in genomic technology and virology, and a better understanding of disease transmission, have accelerated work that began in the 1980s to make genetically engineered viruses that spread from one animal to another, imparting immunity to disease rather than infection.

Researchers are currently developing self-spreading vaccines for Ebola, <u>bovine</u> tuberculosis, and Lassa fever, a viral disease spread by rats that causes upward of <u>300,000</u> infections annually in parts of West Africa. The approach could be expanded to target other <u>zoonotic diseases</u>, including rabies, West Nile virus, Lyme disease, and the plague.

Advocates for self-spreading vaccines say they could revolutionize public health by disrupting infectious disease spread among animals before a zoonotic spillover could occur—potentially preventing the next pandemic.

But others argue that the viruses used in these vaccines could themselves mutate, jump

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"Once you set something engineered and self-transmissible out into nature, you don't know what happens to it and where it will go," says Jonas Sandbrink, a biosecurity researcher at the University of Oxford's Future of Humanity Institute. "Even if you just start by setting it out into animal populations, part of the genetic elements might find their way back into humans."

The first, and only, self-spreading vaccine field trial

In 1999, veterinarian José Manuel Sánchez-Vizcaíno led a team of researchers to Isla del Aire, an island off the eastern coast of Spain, to test a self-spreading vaccine against two viral diseases: rabbit hemorrhagic disease and myxomatosis. Although neither disease infects humans, at the time they both had been decimating domestic and wild rabbit populations across China and Europe for several decades.

Traditional vaccines for both diseases were used in domestic rabbits, but trapping and vaccinating wild rabbits, which are notoriously fast-breeding, was an insurmountable task, Sánchez-Vizcaíno explains. He saw huge

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Health in Spain, and his team sliced out a gene from the rabbit hemorrhagic disease virus and inserted it into the genome of a mild strain of the myxoma virus, which causes myxomatosis. The final product was a hybrid virus vaccine that protected against both rabbit hemorrhagic disease and myxomatosis. Sánchez-Vizcaíno hypothesized that because the vaccine was similar enough to the original disease-causing myxoma virus, it would still spread among wild rabbits.

On the island the research team captured 147 rabbits, placed microchips in their necks, administered the vaccine to about half of them, and released them all back into the wild. For the next 32 days, the vaccinated and unvaccinated rabbits lived as they normally did. When researchers recaptured micro-chipped rabbits that had not been vaccinated originally, they found that 56 percent of them had antibodies to both viruses, indicating that the vaccine had successfully spread from vaccinated to unvaccinated animals.

The experiment was the first proof-of-concept field test for self-spreading vaccines—and it remains the only one ever attempted.

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approval for real-world use. The EMA noted technical issues with the vaccine's safety evaluation and requested that the team decode the myxoma genome, which had not been done before.

Although the team was given two years to comply, the funding body did not provide support for further work, recalls Juan Bárcena, then a Ph.D. student working under Sánchez-Vizcaíno. Bárcena no longer advocates for self-spreading vaccine technology, but he says that data from the laboratory and field trials showed the vaccine was safe and its spread remained confined to the rabbit populations.

Still, Bárcena doubts that the EMA would have ever approved their vaccine given the hesitancy and controversy around genetically modified organisms.

Scott Nuismer, a professor at the University of Idaho who conducts mathematical modelling studies of self-spreading vaccines today, noted that Sánchez-Vizcaíno's vaccine may have posed more risks than current technologies because the team used a myxoma virus, which is itself deadly, as its vehicle for the vaccine.

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in investing in research and development for a technology that, by design, would reduce its own profit margins, Sánchez-Vizcaíno speculates.

Vaccines in progress

Renewed interest and funding for the technology popped up around 2016, and today several research groups are developing self-spreading vaccines for animals.

Each of these new vaccines are so-called recombinant viruses. Researchers first identify a protein from the target microbe that serves as an antigen—a substance that triggers immune responses in vaccinated people or animals. Then the researchers select a virus to carry the vaccine and spread it. To do this, researchers capture a few animals from their target population—primates for Ebola, rats for Lassa fever—and isolate a virus that naturally infects those animals. Then they splice in genetic material from the target to create a vaccine.

Each of these vaccines uses a cytomegalovirus, or CMVs, a group that belongs to the herpes family.

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and can accommodate additional genes from the targeted microbe, says Alec Redwood, a principal research fellow at the University of Western Australia. He conducted self-spreading vaccine research in the early 2000s and is now part of a team developing a CMV-based Lassa fever vaccine.

CMVs also infect a host for life, induce strong immune responses yet do not often cause severe disease. Perhaps most importantly, CMVs are uniquely species-specific; the CMV that spreads among *Mastomys natalensis*, the rat species that spread Lassa fever, for example, cannot infect any animals other than *M*. *natalensis*.

Several small studies have demonstrated that the CMV-based Ebola and bovine tuberculosis vaccines are effective when delivered through traditional injections. Across two trials involving about 50 monkeys, the CMV-based tuberculosis vaccine reduced disease by 68 percent, researchers reported. In a separate study, three out of four monkeys vaccinated with the Ebola vaccine survived direct exposure to Ebola.

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allows researchers to control the number of times the vaccine can multiply, thereby limiting its lifetime, Redwood explains.

So far, no one has conducted any field or laboratory studies assessing the impact and safety of these vaccines delivered via the self-spreading mechanism. However, a recent mathematical modelling <u>study</u> reported that if it works as expected, releasing the Lassa fever vaccine could reduce disease transmission among rodents by 95 percent in less than a year.

"You can really see just how powerful the idea could be," says Nuismer, who was the senior author of the modelling study.

Risks of self-spreading vaccines

Despite the potential benefits, many experts warn that too little is known about zoonotic disease transmission and viral evolution to accurately predict what might happen if a selfspreading vaccine were released into the wild.

"Our understanding of infectious disease dynamics in wildlife remain for the most part too simple to meaningfully predict the outcome

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Australia and the president of the Wildlife Disease Association.

Bárcena's view of self-spreading diseases shifted after he saw how previous animal control strategies involving the intentional release of viruses had unforeseen consequences.

For instance, the myxoma virus that had become such a devastating challenge in Europe arose because a man in France intentionally released the virus in 1952 to keep rabbits out of his home garden. In 2018 Spanish researchers started noticing that a myxoma virus was killing wild hares, a species similar to rabbits. Scientists sequenced its genome and concluded that the myxoma virus had mixed with a poxvirus, enabling it to jump species.

"I don't know if a mathematical model would have said that 70 years later something like this can happen," says <u>Bárcena</u> who is now a senior scientist at the Center for Research in Animal Health in Spain.

Filippa Lentzos, a science and international security expert at King's College London, points

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in wild and domestic animal populations and, perhaps, even in humans.

Nuismer and Redwood both say it is highly unlikely that a CMV-based vaccine could ever jump species given the virus's biology.

Although the evolutionary factors underlying CMV's species-specificity are not entirely known, there has never been a documented case in the wild or in a laboratory of a successful cross-species CMV infection.

Another potential risk of self-disseminating vaccines is that ridding wild animals of infectious diseases could disrupt natural population control. The rodents that spread Lassa virus are pests that destroy crops and homes, contaminate stored food and drinking water, and create unsanitary living conditions. If the virus doesn't affect them anymore, their numbers could skyrocket.

"Say we cure these rodents of Lassa virus and that's good, that's great for humanity. Except what if that virus was controlling their population size or something? And then we get a wild expansion of the reservoir rodents," Nuismer says. "I see this as a much more

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In addition, there is an emerging understanding that viruses and bacteria exist in complex microbial ecosystems, perhaps keeping each other's populations in check. The impact of a self-spreading vaccine that wipes out one specific virus might have unknown consequences.

"Dramatically shifting the balance by attempting to eradicate or reduce an endemic virus in nature could risk the emergence of other pathogens which impact both the wildlife species themselves, as well as people and our domestic animals," says Peters.

To mitigate these risks, Nuismer and Redwood envision a progression of testing setups that moves slowly from lab-controlled trials to large-scale enclosures, perhaps on an island like Sánchez-Vizcaíno and his team did more than 20 years ago.

The long road ahead

Most researchers agree that self-spreading vaccines could never be applied to human populations, because universal informed consent would never be achieved.

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know, it's stuff of fantasy. It will never be used in humans," Redwood says.

But even using a self-spreading vaccine among animals faces regulatory and social hurdles.

"What are the political implications of such interventions, which do not recognize and cannot be contained by state or national borders?" Peters asks.

Sandbrink also points out that self-spreading vaccine research poses a biosecurity threat. Developing them and preventing some of their potential consequences involves fine-tuning transmissibility and altering genetic stability, techniques that "uniquely advance certain capabilities applicable to the creation of viruses for pandemics and as biological weapons," he says.

Scientific and global health communities and funding bodies should consider alternative solutions that provide the same benefit with less risk, Sandbrink urges. For instance, educating people about how to safely interact with wildlife may reduce the chance of viral spillover. Improving disease surveillance in

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Given the extremely high-risk and international nature of this work, and because the consequences are "potentially irreversible," Lentzos says stakeholders must engage in dialogue on how this research is regulated, and Nuismer and Redwood agree that there is still a long way to go.

"You don't need to be a Rhodes scholar to work out that people will be nervous about a disseminating viral vector. It's a concept that will scare people," says Redwood. "The way that I like to think about it is that it may never be used, but it's better to have something in the cupboard that can be used and is mature if we need it. And to say, 'Let's just not do this research because it's too dangerous,' to me, that makes no sense at all."



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