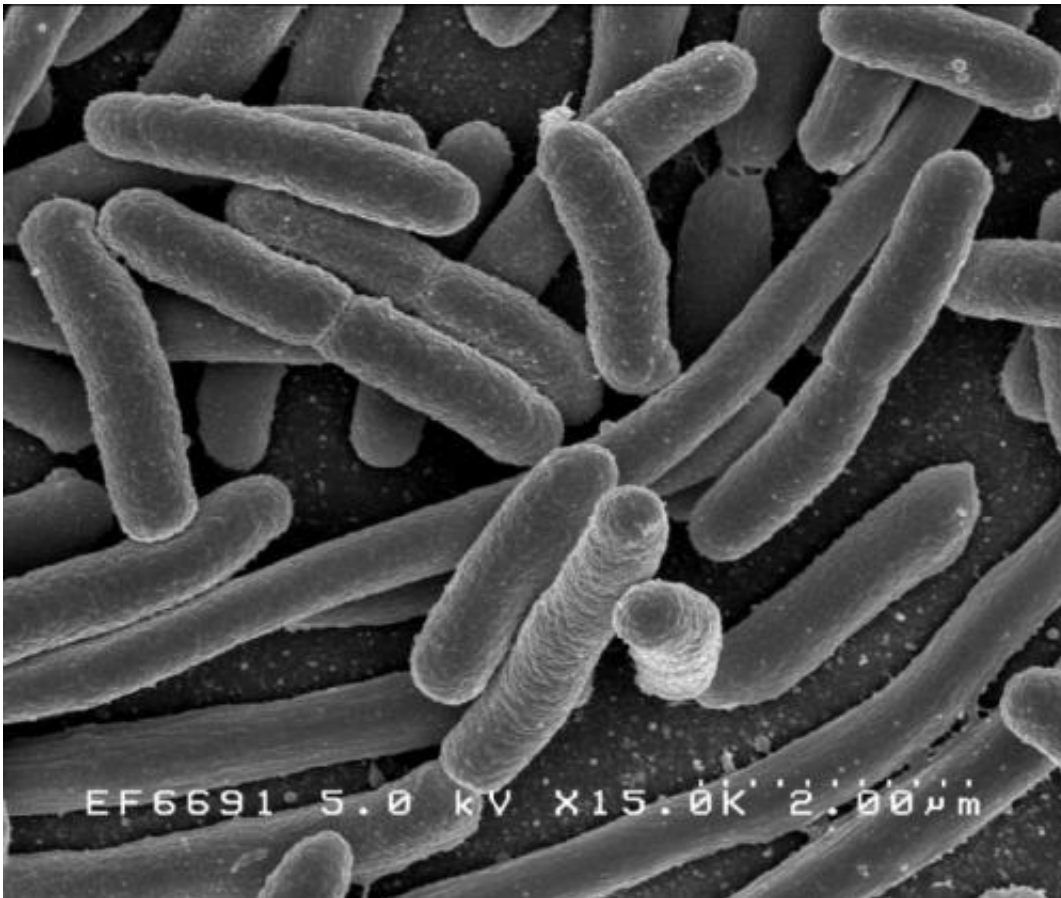


Engineered viruses could fight drug resistance

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Escherichia coli. Credit: Rocky Mountain Laboratories, NIAID, NIH

In the battle against antibiotic resistance, many scientists have been trying to deploy naturally occurring viruses called bacteriophages that can infect and kill bacteria.

Bacteriophages kill bacteria through different mechanisms than antibiotics, and they can target specific strains, making them an appealing option for potentially overcoming [multidrug resistance](#). However, quickly finding and optimizing well-defined bacteriophages to use against a bacterial target is challenging.

In a new study, MIT biological engineers showed that they could rapidly program bacteriophages to kill different strains of *E. coli* by making mutations in a viral protein that binds to host cells. These engineered bacteriophages are also less likely to provoke resistance in bacteria, the researchers found.

"As we're seeing in the news more and more now, bacterial resistance is continuing to evolve and is increasingly problematic for public health," says Timothy Lu, an MIT associate professor of electrical engineering and computer science and of biological engineering. "Phages represent a very different way of killing bacteria than antibiotics, which is complementary to antibiotics, rather than trying to replace them."

The researchers created several engineered phages that could kill *E. coli* grown in the lab. One of the newly created phages was also able to eliminate two *E. coli* strains that are resistant to naturally occurring phages from a skin infection in mice.

Lu is the senior author of the study, which appears in the Oct. 3 issue of *Cell*. MIT postdoc Kevin Yehl and former postdoc Sebastien Lemire are the lead authors of the paper.

Engineered viruses

The Food and Drug Administration has approved a handful of bacteriophages for killing harmful bacteria in food, but they have not been widely used to treat infections because finding naturally occurring

phages that target the right kind of bacteria can be a difficult and time-consuming process.

To make such treatments easier to develop, Lu's lab has been working on engineered viral "scaffolds" that can be easily repurposed to target different bacterial strains or different resistance mechanisms.

"We think phages are a good toolkit for killing and knocking down bacteria levels inside a complex ecosystem, but in a targeted way," Lu says.

In 2015, the researchers used a [phage](#) from the T7 family, which naturally kills E.coli, and showed that they could program it to target other bacteria by swapping in different genes that code for tail fibers, the protein that bacteriophages use to latch onto receptors on the surfaces of host cells.

While that approach did work, the researchers wanted to find a way to speed up the process of tailoring phages to a particular type of bacteria. In their new study, they came up with a strategy that allows them to rapidly create and test a much greater number of tail fiber variants.

From previous studies of tail fiber structure, the researchers knew that the protein consists of segments called beta sheets that are connected by loops. They decided to try systematically mutating only the amino acids that form the loops, while preserving the beta sheet structure.

"We identified regions that we thought would have minimal effect on the protein structure, but would be able to change its binding interaction with the bacteria," Yehl says.

They created phages with about 10,000,000 different tail fibers and tested them against several strains of E. coli that had evolved to be

resistant to the nonengineered [bacteriophage](#). One way that *E. coli* can become resistant to bacteriophages is by mutating "LPS" receptors so that they are shortened or missing, but the MIT team found that some of their engineered phages could kill even strains of *E. coli* with mutated or missing LPS receptors.

Other targets

Lu and Yehl now plan to apply this approach to targeting other resistance mechanisms used by *E. coli*, and they also hope to develop phages that can kill other types of harmful bacteria. "This is just the beginning, as there are many other viral scaffolds and bacteria to target," Yehl says. The researchers are also interested in using bacteriophages as a tool to target specific strains of [bacteria](#) that live in the human gut and cause health problems.

"Being able to selectively hit those nonbeneficial [strains](#) could give us a lot of benefits in terms of human clinical outcomes," Lu says.

More information: Kevin Yehl et al, Engineering Phage Host-Range and Suppressing Bacterial Resistance through Phage Tail Fiber Mutagenesis, *Cell* (2019). DOI: 10.1016/j.cell.2019.09.015 , [dx.doi.org/10.1016/j.cell.2019.09.015](https://doi.org/10.1016/j.cell.2019.09.015)

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