A big step toward treating tuberculosis without risking antibiotic resistance
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The mechanism of selectivity of evybactin. Evybactin (EVY) is transported into the bacterial cell by BacA (M. tuberculosis) and SbmA (E. coli) transporters. However, in E. coli, evybactin is removed by drug efflux pump AcrAB-TolC. As a result, evybactin selectively kills M. tuberculosis. Credit: Nature Chemical Biology (2022). DOI: 10.1038/s41589-022-01102-7

Tuberculosis, an infectious disease that killed 1.5 million people in 2020, is most commonly treated with a cocktail of four drugs that must be taken for up to six months. This treatment plan, however, can have the unintended consequence of leading to antibiotic resistance, both because people may not finish the full course of treatment and, more directly, because one of the drugs can act on bacteria other than Mycobacterium tuberculosis, including components of the intestinal microflora that are critical to health and well-being.

Recently, researchers identified an antimicrobial compound, which they named evybactin, which could selectively act against Mycobacterium tuberculosis.

Their findings were published in Nature Chemical Biology.

“Microorganisms such as actinomycetes and filamentous fungi have long been used as a source of antibiotics,” said first author Yu Imai, assistant professor at the Institute for Biomedical Sciences at Shinshu University, who previously discovered a new antibiotic while studying under Distinguished Professor Kim Lewis at the Antimicrobial Discovery Center, Northeastern University.

"However, in recent years, it has become difficult to find new antibiotics from these microorganisms, as known substances have been re-isolated. Recently, we have succeeded in discovering several antibiotics from the nematode symbiotic bacterium Photorhabdus and have shown that this bacterium is a useful source for the discovery of new antibiotics."

This discovery led to the idea that Photorhabdus may produce antibiotics that are active against M. tuberculosis, according to the researchers. In order to test if an antibiotic from Photorhabdus was active against M. tuberculosis and only M. tuberculosis—that is, it would not have the unintended side effect of acting against other, often healthy bacteria in the body—the researchers tested cultures prepared from Photorhabdus against both M. tuberculosis and S. aureus, a bacterium that causes staph infections.

"We used this method, called differential screening, [that] allows us to eliminate antibiotics that may exhibit broad-spectrum or cytotoxic activity in the early stages of screening," Imai said. "Thus, differential screening is an excellent method for finding antibiotics that exhibit species-specific activity."

The researchers discovered a novel antimicrobial, evybactin, which is a DNA gyrase inhibitor that they proved targeted M. tuberculosis only. They found that evybactin was transported inside the target bacteria across the cell membrane by BacA, a carrier of hydrophobic compounds, and then it
binds to a specific site of DNA gyrase.

This finding is significant not only because of the direct benefit of the discovery that evybactin can be used to treat tuberculosis without causing antibiotic resistance but also because of the implications of the method used to discover evybactin, as the method could potentially be used to discover even more antibiotics.

"In this study, we demonstrated that it is possible to find new antibiotics by using relatively untouched microorganisms as a search source for useful antibiotics," Imai said.

While simple animal experiments have shown that evybactin is active in vivo, according to the researchers, the next steps for this research include further testing in animals infected by M. tuberculosis to assess detail of the efficacy and long-term impacts.

"For accelerate development as an antituberculosis drug, it is necessary to evaluate the safety, pharmacokinetics and long-term administration of evybactin to M. tuberculosis patients," Imai said.

**More information:** Yu Imai et al, Evybactin is a DNA gyrase inhibitor that selectively kills Mycobacterium tuberculosis, *Nature Chemical Biology* (2022). DOI: 10.1038/s41589-022-01102-7

Provided by Shinshu University


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