

Molecular machinery that makes potent antibiotic revealed after decades of research

January 17 2019



The 3D structure of McbBCD, an enzyme (protein) that makes the potent antibiotic microcin B17 from a smaller protein known as a peptide, as revealed by X-ray crystallography. The red spheres show chemical "cycles" formed by the enzyme that are required for antibacterial activity. Credit: Dmitry Ghilarov

Scientists at Rutgers and universities in Russia, Poland and England have solved a nearly 30-year mystery—how the molecular machinery works in an enzyme that makes a potent antibiotic.



The findings, which appear in the journal *Molecular Cell*, provide the tools to design <u>new antibiotics</u>, anticancer drugs and other therapeutics.

The potent natural antibiotic, microcin B17, kills harmful *E. coli* bacteria. Microbial resistance to antibiotics- due to their overuse and misuse—is one of the biggest threats facing humanity, and there's an urgent need to find new drugs. Natural antibiotics that evolved over eons represent an attractive option to overcome resistance.

The scientists studied a molecular machine: an enzyme (protein) called McbBCD. The enzyme makes microcin B17 from a smaller protein known as a peptide. Scientists have known about microcin B17 and its unusual chemical structure for decades, but they did not understand the <u>molecular machinery</u> that makes it until now.

The scientists found that the enzyme triggers two <u>chemical reactions</u> that produce several chemical "cycles" required for antibacterial activity, according to senior author Konstantin Severinov, a principal investigator at the Waksman Institute of Microbiology and professor of molecular biology and biochemistry at Rutgers University-New Brunswick.

"Our research allows rational design of new peptide compounds that could become treatments ranging from antimicrobials to anticancer drugs," Severinov said. "There may be a trove of new antibiotics that could be made from peptides, using <u>enzyme</u> machines like McbBCD as a production tool."

The international team included scientists at the Skolkovo Institute of Science and Technology in Russia; Russian Academy of Sciences in Russia; Jagiellonian University in Poland; John Innes Centre in England; and Lomonosov Moscow State University in Russia.

More information: Molecular Cell (2019). DOI:



10.1016/j.molcel.2018.11.032

Provided by Rutgers University

Citation: Molecular machinery that makes potent antibiotic revealed after decades of research (2019, January 17) retrieved 28 April 2024 from <u>https://phys.org/news/2019-01-molecular-machinery-potent-antibiotic-revealed.html</u>

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