

Hydrogen sulfide critical to innate ability of bacteria to survive antibiotics

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Scanning electromicrograph of Staphylococcus aureus bacteria. Credit: NIAID

The signaling molecule hydrogen sulfide (H_2S) plays a critical role in antibiotic tolerance, the innate ability of bacteria to survive normally lethal levels of antibiotics, a new study finds.

Published online in the journal *Science* on June 11, the study revolves around tolerance, wherein bacteria in general have evolved to use common defense systems to resist <u>antibiotics</u>. Tolerance differs from antibiotic resistance, where one species happens to acquire a genetic change that helps them resist treatment.

In one defense mechanism, tolerant bacteria, also called "persisters," stop multiplying (proliferating), reducing their energy use (metabolism) to survive antibiotic treatment, but resuming growth when the treatment ends. Persisters are particularly abundant in biofilms, <u>bacterial colonies</u> that live in tough polymeric matrices which further prevent their eradication.

"The combined trends toward resistant infections and fewer new antimicrobials are projected to kill 10 million people annually by the year 2050," says corresponding study author Evgeny Nudler, Ph.D., the Julie Wilson Anderson Professor of Biochemistry at NYU Langone Health, and an investigator with the Howard Hughes Medical Institute. "New approaches are urgently needed to prevent this, and our study suggests that suppressing bacterial H_2S would make different antibiotics more potent."

In their prior work, the NYU Langone research team showed that H2S production is deployed against antibiotics by a wide variety of bacterial species, including two increasingly antibiotic-resistant pathogens



prevalent in hospital-borne infections: Staphylococcus aureus and Pseudomonas aeruginosa. S. aureus is gram-positive, while Pseudomonas aeruginosa is gram-negative, with the differing organizations of their outer layers demonstrating that H_2S production protects pathogens across the bacterial kingdom.

Remarkably, the research team team found that both species rely on the same enzyme, cystathionine γ -lyase (CSE), for the bulk of H₂S production. Blocking its action would represent then a way to remove an important defense against antibiotics, but available CSE inhibitors have a low potency against bacterial CSE and a high probability of causing side effects in human tissue, says Nudler.

To find better inhibitors, the research team obtained an X-ray structure of S. aureus CSE and used it to "virtually screen" millions of drug-like compounds looking for those with the right shape and properties to block the enzyme's action without side effects. The team selected lead compounds, NL1, NL2, and NL3, which inhibited the bacterial CSE, blocked H2S production by both S. aureus and P. aeruginosa, and strengthened the effect of bactericidal antibiotics from different classes. Furthermore, NL1 increased the potency of antibiotic effect in mouse models of S. aureus and P. aeruginosa infection.

Unexpectedly, further testing revealed that the NL compounds markedly diminished persisters, and suppressed biofilm formation in both pathogens. How exactly H2S contributes to tolerance remains to be established, but there are some hints.

"Bacteria appear to use controlled, self-poisoning with H_2S to slow down their metabolism, preventing the antibiotics from using the bacteria's energy production system to kill them," says Nudler. "Interfering with the H_2S -based defenses represents a largely unexplored alternative to the traditional antibiotic discovery. Our results suggest that a new kind of



small molecule potentiator can strengthen the effect of major classes of clinically important antibiotics."

The authors note several opportunities for designing conceptually novel antimicrobial therapeutics by combining H_2S -blocking potentiators with antibiotics. Such combinations may have better efficacy against bacterial biofilms. Other potential applications include overcoming intermediate-level antibiotic resistance; reducing antibiotic dose and related toxicity while maintaining efficacy; and enhancing the bacteria-killing (bactericidal) effect at the same antibiotic dose.

More information: K. Shatalin el al., "Inhibitors of bacterial H_2S biogenesis targeting antibiotic resistance and tolerance," *Science* (2021). <u>science.sciencemag.org/cgi/doi ... 1126/science.abd8377</u>

Provided by NYU Langone Health

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