

Discovery of critical inhibitory site in an energy-producing enzyme could lead to the rational design of antibiotics

December 9 2022



(A) Deaths from AMR are predicted to rise globally in the next 35 years.
(O'Neill J, Chair. Tackling Drug-Resistant Infections Globally: Final Report and Recommendations. London, UK: Review on Antimicrobial Resistance; 2016; p. 1-84.) (B, C) Allosteric sites (red shadow) of mammalian mtHCO (A: PDB 7XMB) and bacterial HCO (B: PDB 6FWF). Helices of subunit I are shown as blue, and a mammalian-specific subunit, COX7C (shown as red), covers the



surface of the inhibitor. The other helices are shown in yellow. (D, E) The surface of the allosteric site is distinct from each other (C: mtHCO, D: bacterial HCO), which makes fitting inhibitors to each corresponding site must have a different character/profile. (F) Q275 is a specific Neisseria HCO inhibitor (filled circle), did not inhibit mammalian mtHCO (open circle). Credit: *Nature communications/* National Cerebral and Cardiovascular Center

Antimicrobial resistance (AMR) is a global health problem. Species of drug-resistant pathogens, including Neisseria gonorrhoeae, could become untreatable due to a high degree of AMR.

A team of researchers has discovered a new mechanism to allosterically inhibit the <u>critical enzyme</u> in the <u>respiratory chain</u>, widely conserved in all three domains of life: bacteria, archaea, and eukaryotes. The team has succeeded in identifying an antibiotic based on their findings, which is effective against a super drug-resistant strain of Neisseria gonorrhoeae.

The team published their findings in the journal Nature Communications.

Antimicrobial resistance (AMR) is a global health problem. Many efforts have been made to reduce the burden of AMR perils globally since 2013. Yet, threats from some species continue to rise regardless: drug-resistant Neisseria gonorrhoeae is one of five urgent threats. Resistance to ceftriaxone, the last option for an empirical first-line antibiotic against Neisseria gonorrhoeae in most countries, has been reported and continues to emerge globally.

The gonococcal infection could become untreatable due to a high degree of AMR, which would increase serious complications: infertility, <u>ectopic</u> <u>pregnancy</u>, and increased transmission of HIV, and neonatal keratoconjunctivitis that can lead to blindness. The emergence of



resistant pathogens to currently available antibiotics is very alarming; thus, the development of antibiotics with a novel mechanism of action is seriously required.

The respiratory chain has recently garnered considerable scientific attention as a potential target for antibiotics. As respiratory enzymes are essential for life, their core structure is generally conserved from bacteria to mammals. Thus, the surface of the substrate binding pocket is quite similar among species, making it challenging to develop a competitive inhibitor for the substrate binding pocket.

Another type of inhibitor for an enzyme is an allosteric inhibitor. This type of inhibitor causes a structural change of the enzyme, leading to the inhibition of its activity. Allosteric sites are evolutionarily less conserved in <u>amino acid sequence</u> than substrate binding sites, theoretically improving selectivity and reducing toxicity. However, a systematic and strategic search for allosteric inhibitors has yet to be established.

The team identified an allosteric inhibitory site buried inside mammalian mitochondrial heme-copper oxidases (mtHCOs), the essential respiratory enzymes for life. The steric conformation around the binding pocket of HCOs is highly conserved among bacteria and mammals, yet the latter has an extra helix.

The existence of an additional helix in mammalian mtHcO makes surface of the pockets distinct from bacterial HCOs. Thus, fitting inhibitors to each corresponding site must have a different character/profile. This structural difference in the conserved allostery enabled us to rationally identify bacterial HCO-specific inhibitors: an antibiotic compound, Q275, against ceftriaxone-resistant Neisseria gonorrhoeae.

Future perspective for finding allosteric modulators



This approach can be applied to finding allosteric modulators in other therapeutic targets. Enzymes generally acquire additional domains or subunits along molecular evolution, bigger in eukaryotes than in their bacterial counterparts. They could likely contain allostery inside the protein at the boundary of the structures between eukaryotes and bacteria, leading to the development of novel antibiotics, as the respiratory chain is a proven target for antibiotics.

Furthermore, any fundamental molecule essential for life and conserved among species could be a potential target. Also, the additional peptides might contain a positive allosteric site at the border of their core structure; a positive allosteric modulator for the loss-of-function human disease could be a therapeutic direction. Thus, in conclusion, this study will open new avenues in protein science and therapeutic development, especially for antibiotics with a novel mechanism of action.

More information: Yuya Nishida et al, Identifying antibiotics based on structural differences in the conserved allostery from mitochondrial heme-copper oxidases, *Nature Communications* (2022). <u>DOI:</u> <u>10.1038/s41467-022-34771-y</u>

Provided by National Cerebral and Cardiovascular Center

Citation: Discovery of critical inhibitory site in an energy-producing enzyme could lead to the rational design of antibiotics (2022, December 9) retrieved 26 April 2024 from <u>https://phys.org/news/2022-12-discovery-critical-inhibitory-site-energy-producing.html</u>

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