Common blood thinner heparin shows promise as cobra bite antidote

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Nubian spitting cobra (Naja nubiae). Credit: The Trustees of the Natural History Museum, London + Callum Mair

Scientists at the University of Sydney and Liverpool School of Tropical Medicine have made a remarkable discovery: a commonly-used blood
thinner, heparin, can be repurposed as an inexpensive antidote for cobra venom.

Cobras kill thousands of people a year worldwide and perhaps a hundred thousand more are seriously maimed by necrosis—the death of body tissue and cells—caused by the venom, which can lead to amputation.

Current antivenom treatment is expensive and does not effectively treat the necrosis of the flesh where the bite occurs.

"Our discovery could drastically reduce the terrible injuries from necrosis caused by cobra bites—and it might also slow the venom, which could improve survival rates," said Professor Greg Neely, a corresponding author of the study from the Charles Perkins Center and Faculty of Science at the University of Sydney.

Using CRISPR gene-editing technology to identify ways to block cobra venom, the team, which consisted of scientists based in Australia, Canada, Costa Rica and the UK, successfully repurposed heparin (a common blood thinner) and related drugs and showed they can stop the necrosis caused by cobra bites.

The research is published on the front cover of *Science Translational Medicine*.

Ph.D. student and lead author, Tian Du, also from the University of Sydney, said, "Heparin is inexpensive, ubiquitous and a World Health Organization-listed Essential Medicine. After successful human trials, it could be rolled out relatively quickly to become a cheap, safe and effective drug for treating cobra bites."

The team used CRISPR to find the *human genes* that cobra venom needs to cause necrosis that kills the flesh around the bite. One of the required
venom targets are enzymes needed to produce the related molecules heparan and heparin, which many human and animal cells produce.

Heparan is on the cell surface and heparin is released during an immune response. Their similar structure means the venom can bind to both. The team used this knowledge to make an antidote that can stop necrosis in human cells and mice.

Unlike current antivenoms for cobra bites, which are 19th century technologies, the heparinoid drugs act as a 'decoy' antidote. By flooding the bite site with 'decoy' heparin sulfate or related heparinoid molecules, the antidote can bind to and neutralize the toxins within the venom that cause tissue damage.

Joint corresponding author, Professor Nicholas Casewell, Head of the Center for Snakebite Research & Interventions at Liverpool School of Tropical Medicine, said, "Snakebites remain the deadliest of the neglected tropical diseases, with its burden landing overwhelmingly on rural communities in low- and middle-income countries.

"Our findings are exciting because current antivenoms are largely ineffective against severe local envenoming, which involves painful progressive swelling, blistering and/or tissue necrosis around the bite site. This can lead to loss of limb function, amputation and lifelong disability."
Snakebites kill up to 138,000 people a year, with 400,000 more experiencing long-term consequences of the bite. While the number affected by cobras is unclear, in some parts of India and Africa, cobra species account for most snakebite incidents.

The World Health Organization has identified snakebite as a priority in its program for tackling neglected tropical diseases. It has announced an ambitious goal of reducing the global burden of snakebite in half by 2030.

Professor Neely said, "That target is just five years away now. We hope that the new cobra antidote we found can assist in the global fight to
reduce death and injury from snakebite in some of the world's poorest communities."

Working in the Dr. John and Anne Chong Laboratory for Functional Genomics at the Charles Perkins Center, Professor Neely's team takes a systematic approach to finding drugs to treat deadly or painful venoms.

It does this using CRISPR to identify the genetic targets used by a venom or toxin inside humans and other mammals. It then uses this knowledge to design ways to block this interaction and ideally protect people from the deadly actions of these venoms.

Artist's impression. Naja pallida spitting at a shield made of heparin sulfate. Credit: Hassan Tahini/ScienceBrush
This approach was used to identify an antidote to box jellyfish venom by the team in 2019.

Professor Casewell leads the Center for Snakebite Research & Interventions at the Liverpool School of Tropical Medicine (LSTM). The center has conducted a diverse portfolio of research activities to better understand the biology of snake venoms and improve the efficacy, safety and affordability of antivenom treatment for tropical snakebite victims for more than 50 years. It boasts some of the world's leading snakebite experts and has access to LSTM's herpetarium, the largest and most diverse collection of tropical venomous snakes in the UK.


Provided by University of Sydney


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