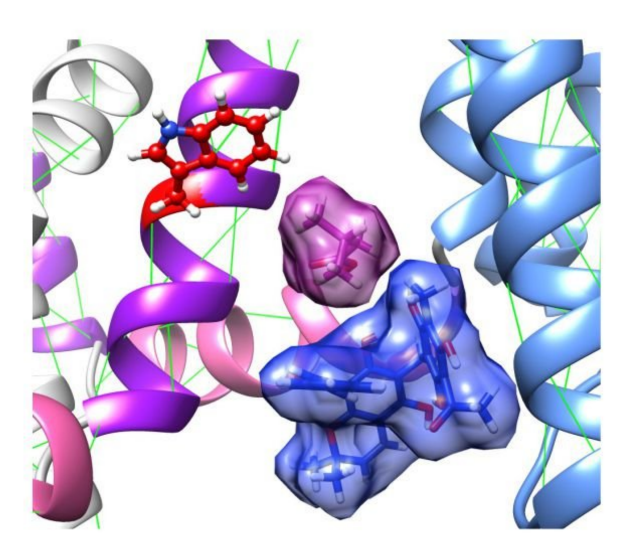


Researchers discover molecular mechanisms of ancient herbal remedies

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Computer model of the two herbal components, an isovaleric acid molecule (violet, upper left) and mallotoxin (blue, lower right), occupying neighboring



binding sites to synergistically activate KCNQ2/3 channels. Credit: Geoff Abbott, PhD, UCI School of Medicine

Researchers in the Department of Physiology & Biophysics at the University of California, Irvine School of Medicine have discovered the molecular basis for a therapeutic action of an ancient herbal medicine used across Africa to treat various illnesses, including epilepsy.

The herbal medicine, a leaf extract from the shrub *Mallotus oppositifolius*, was previously found to be effective in controlling seizures but the mechanism was unknown. The discovery, published in *Nature Communications*, found that two components of the Mallotus leaf extract activate KCNQ2/3, a <u>potassium ion channel</u> essential for controlling <u>electrical activity</u> in the brain. The two components were somewhat effective alone, but in combination were highly effective both at activating KCNQ2/3 channels and at preventing life-threatening seizures.

The UCI research team, comprising postdoctoral fellow Rían Manville, Ph.D. and principal investigator Geoffrey Abbott, MSc, Ph.D., screened individual compounds from the leaf extract for channel opening activity, and then combined the two most active compounds to discover the therapeutic synergy contained in an African folk remedy used for centuries. Strikingly, one of the two compounds identified, isovaleric acid, is also a main component of valerian root, an herb used in ancient Greece as an insomnia sleep remedy, and for centuries by the English and also native Americans as an anticonvulsant. Valerian root is still used by as many as 2 million people each week in the United States as an herbal remedy for anxiety and insomnia.

"We are very interested in taking a molecular approach to



ethnobotany—the study of plants and their use by local populations—to discover the molecular mechanisms for ancient remedies and to use this knowledge to create safer and more effective drugs. The KCNQ channels we study are typically opened by electrical activity, but we know that they are also incredibly sensitive to the presence of small molecules, including neurotransmitters, but also molecules from outside, such as drugs, and constituents of food and herbal extracts," said Abbott. "Some folk medicines are in danger of being lost, either because traditional practices are being forgotten, or because the plant species used are endangered. Species loss can arise from over-collecting, habitat destruction, or climate change. There is a race against time to prevent this incredible resource being lost forever."

The UCI team found that the <u>herbal extract</u> they studied had different channel subtype preferences than modern drugs that activate the KCNQ2/3 channel, such as the anticonvulsant <u>drug</u>, retigabine. Because of this, by combining the herbal compounds with retigabine, they were able to completely lock open the channel, a feat not previously achieved.

"Locking open the <u>channel</u> is a neat trick, but it could also have clinical implications. Retigabine was removed from the market last year because of a surprising side effect: it turns the skin and whites of the eyes blue. However, by combining retigabine with the herbal components, we found we could greatly reduce the retigabine dosage required for activity. This type of strategy might one day enable us to use drugs like retigabine at dosages low enough to be safe, whilst retaining or even enhancing their efficacy by combining them with natural booster compounds derived from plants," said Abbott.

In addition to the booster effects of the herbal extract, identification of the ability of specific chemicals within plants to activate influential ion channels such as KCNQ2/3 may lead one day to new epilepsy, anxiety and pain drugs that exploit the alternative chemical spaces offered by the



molecular constituents of ethnobotanicals.

More information: Rían W. Manville et al, Ancient and modern anticonvulsants act synergistically in a KCNQ potassium channel binding pocket, *Nature Communications* (2018). DOI: 10.1038/s41467-018-06339-2

Provided by University of California, Irvine

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