

New use for an old drug: How does ketamine combat depression?

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A group of proteins called 4E-BPs, involved in memory formation, is the



key to unlocking the antidepressant effect of ketamine in the brain, according to researchers from three Canadian universities. The discovery could lead to better and safer treatments for certain patients suffering from major depression.

Because more than 30% of patients are resistant to <u>selective serotonin</u> <u>reuptake inhibitors</u> (SSRI), the most commonly-prescribed antidepressants, finding an effective treatment for major depressive disorder is challenging.

Initially, ketamine was approved for anesthesia and pain relief. Since its discovery, researchers have been studying new uses for this drug, and ketamine was approved last year for patients with <u>major depression</u> who are treatment-resistant. Unlike standard antidepressants, which can take several weeks to have an effect, ketamine works within hours. Until now, little was known about the molecular mechanism that triggers the antidepressant effect of ketamine on the brain.

In the study published in *Nature*, researchers from McGill University, Université de Montréal, and Carleton University investigated the effect of ketamine on behavior and neuronal activity in mice. Using genetic tools to remove proteins from specific brain cells, the team found that when 4E-BPs are absent in the brain, specifically in neurons, ketamine cannot produce its antidepressant effect. 4E-BPs act as a switch to turn on or off the process of protein synthesis—an essential component of memory formation.

"This is yet another prime example of how basic research, in this case the control of protein synthesis, leads to major discoveries in understanding disease, and the hope of curing it," says co-author Nahum Sonenberg, a professor at the Department of Biochemistry at McGill University.



The researchers examined the role of 4E-BPs on ketamine's effect in two major types of neurons: excitatory neurons, which make up most of the neurons in certain parts of the brain, and inhibitory neurons, which control excitatory neurons and have important effects on behavior.

"We were expecting that 4E-BPs would only be important in excitatory cells, but surprisingly, removing 4E-BPs from inhibitory cells was sufficient to block the effect of ketamine," says co-author Jean-Claude Lacaille, a professor at the Department of Neurosciences at Université de Montréal.

Medicine is not one-size-fits-all

The discovery and approval of ketamine for treatment-resistant patients was considered a major advance in modern psychiatry. Despite its promise, ketamine remains a less-than-perfect therapy because it can be addictive. The researchers hope that their findings will pave the way for better and safer antidepressant therapies for patients with <u>major</u> <u>depressive disorder</u>.

"Too many decisions continue to be made by a trial-and-error approach that can prolong the suffering of patients and affect their quality of life," says co-author Aguilar-Valles, a former research associate at McGill University and now an assistant professor at Carleton University. "Our discovery has the potential to bring us closer to find a safer alternative to ketamine, and ultimately to a personalized medicine approach, where medical treatments are tailored to the individual characteristics of each patient."

The study also involved a clinician-researcher, Dr. Gabriella Gobbi, from the Department of Psychiatry at McGill University, who works with individuals affected with depression and other psychiatric illnesses.



As next steps, the researchers will examine whether males and females have different responses to <u>ketamine</u>. This could have important implications for treatment for people with depressive disorders, among which women are significantly overrepresented.

More information: Argel Aguilar-Valles et al, Antidepressant actions of ketamine engage cell-specific translation via eIF4E, *Nature* (2020). DOI: 10.1038/s41586-020-03047-0

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