

Researchers identify key brain circuits for reward-seeking and avoidance behavior

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Researchers mapped new connections in the extended amygdala related to reward-seeking and aversion. Here neurons in that pathway are labeled in fluorescent green in a cross-section of a rodent brain. Credit: Stanford University - de Lecea lab

Researchers have identified connections between neurons in brain systems associated with reward, stress, and emotion. Conducted in mice,



the new study may help untangle multiple psychiatric conditions, including alcohol use disorder, anxiety disorders, insomnia, and depression in humans.

"Understanding these intricate <u>brain systems</u> will be critical for developing diagnostic and therapeutic tools for a broad array of conditions," said George F. Koob, Ph.D., director of the National Institute on Alcohol Abuse and Alcoholism (NIAAA), which contributed funding for the study. The National Institute of Mental Health (NIMH) also provided major support for the research. NIAAA and NIMH are parts of the National Institutes of Health.

A report of the study, by first author Dr. William Giardino and colleagues at Stanford University, appears in the August 2018 issue of *Nature Neuroscience*.

Responding appropriately to aversive or rewarding stimuli is essential for survival. This requires fine-tuned regulation of brain systems that enable rapid responses to changes in the environment, such as those involved in sleep, wakefulness, stress, and reward-seeking. These same brain systems are often dysregulated in addiction and other psychiatric conditions.

In the new study, researchers looked at the extended amygdala, a brain region involved in fear, arousal, and emotional processing and which plays a significant role in drug and alcohol addiction. They focused on a part of this structure known as the bed nucleus of stria terminalis (BNST), which connects the extended amygdala to the hypothalamus, a brain region that regulates sleep, appetite, and body temperature.

The hypothalamus is also thought to promote both negative and positive emotional states. A better understanding of how the BNST and hypothalamus work to coordinate emotion-related behavior could shed



light on the emotional processes dysregulated in addiction.

"These circuits, also implicated in binge drinking, are likely to be elements in understanding the detailed mechanisms driving stress-related alcohol- or drug-seeking and consummatory behaviors," said first author Dr. Giardino.

To map the brain circuitry between the BNST and the hypothalamus, Dr. Giardino and his colleagues exposed mice to rewarding and aversive stimuli, and then visualized and manipulated the activity of neurons using fiber optic techniques.

The scientists identified two distinct subpopulations of neurons in the BNST that connect to separate populations of neurons in the lateral hypothalamus. These parallel circuits drove opposing <u>emotional states</u>: avoidance (aversion) and approach (preference). Different neurotransmitters were linked to aversion and preference within these neurocircuits – corticotropin releasing factor was involved in aversion, while cholecystokinin played a role in preference.

"The discovery of the unique and opposing influences of these specific BNST to LH circuits may be broadly relevant to the fields of neuroscience and <u>mental health</u>," said senior author Luis de Lecea, Ph.D., Professor of Psychiatry and Behavioral Sciences at Stanford University.

More information: William J. Giardino et al. Parallel circuits from the bed nuclei of stria terminalis to the lateral hypothalamus drive opposing emotional states, *Nature Neuroscience* (2018). DOI: 10.1038/s41593-018-0198-x



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