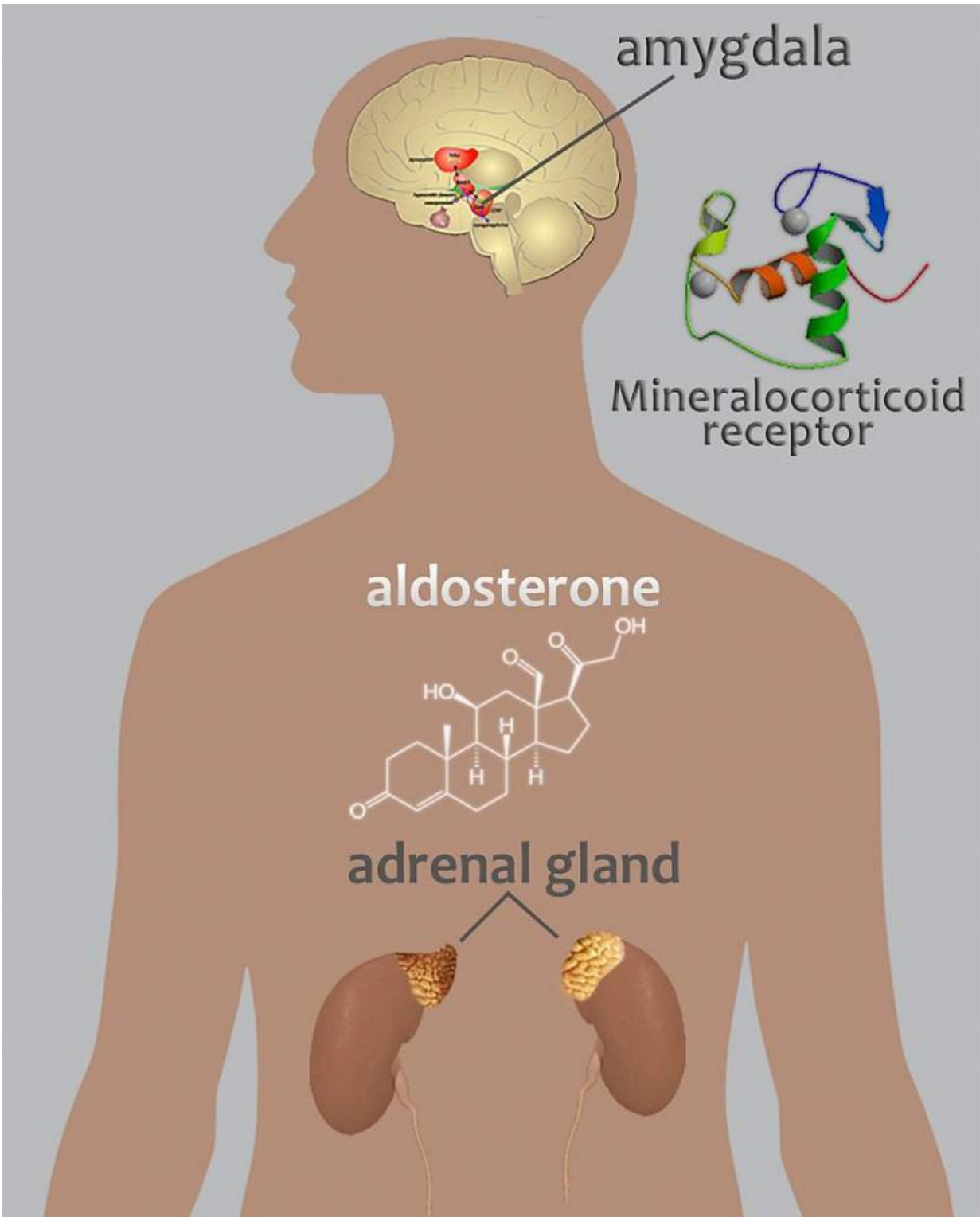


Findings link aldosterone with alcohol use disorder

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Aldosterone, a hormone produced in the adrenal glands, may contribute to alcohol use disorder (AUD) by binding to mineralocorticoid receptors in the

brain, particularly in the amygdala – a key brain area involved in the development and maintenance of AUD. Credit: NIAAA

A new study led by scientists at the National Institute on Alcohol Abuse and Alcoholism (NIAAA), part of the National Institutes of Health, demonstrates that aldosterone, a hormone produced in the adrenal glands, may contribute to alcohol use disorder (AUD). The novel research, conducted in collaboration with a team of investigators in the United States and Europe, appears in the journal *Molecular Psychiatry*.

" This intriguing work – conducted in humans as well as two other species—provides a compelling example of how basic and preclinical research is translated into studies with direct relevance to humans," said NIAAA Director George F. Koob, Ph.D., a co-author of the study. "It also demonstrates how interactions between the brain and the endocrine system may serve as potential targets for the development of new AUD medications."

Aldosterone helps regulate electrolyte and fluid balance by binding to mineralocorticoid receptors (MRs), which are located throughout the body. In the brain, MRs are mainly located in the [amygdala](#) and the [prefrontal cortex](#)—two key brain areas involved in the development and maintenance of AUD. In AUD, amygdala dysfunction heightens activation of brain stress systems resulting in anxiety and other negative emotions, while disruption of the prefrontal cortex impairs executive control systems involved in the ability to make decisions and regulate one's actions, emotions, and impulses.

"Previous studies, including a pilot clinical study that we published in 2008, illustrate the possible role for aldosterone in AUD," said senior author Lorenzo Leggio, M.D., Ph.D., chief of the Section on Clinical

Psychoneuroendocrinology and Neuropsychopharmacology, a NIAAA intramural laboratory jointly funded with the National Institute on Drug Abuse, also part of NIH. "Our overall hypothesis has been that aldosterone may play a role in AUD via its MR receptor and that this neuroendocrine pathway may be particularly important in anxiety, stress and stress-induced [alcohol drinking](#)."

The new report describes three separate studies, conducted with non-human primates, rats, and humans, that investigated the potential contribution of the aldosterone/MR pathway to AUD.

In a study conducted with non-human primates, the researchers found that animals which self-administered alcohol every day for six to 12 months had significantly higher blood aldosterone concentrations, compared with the concentrations measured prior to alcohol administration. In fact, the aldosterone increase observed at six months remained high after 12 months of continued [drinking](#), and did not increase further, suggesting that blood aldosterone concentrations become regulated at a new set-point under daily alcohol consumption. Furthermore, the researchers found that in the amygdala, lower MR gene expression – the process of making functional MR protein from the MR gene – was associated with increased alcohol drinking in the animals. Similar results were not found in the prefrontal cortex.

In a study conducted in a rat model of AUD, the researchers found that lower levels of MR gene expression in the amygdala (but not in the prefrontal cortex) were associated with increased anxiety-like behavior and compulsive drinking, compared to rats not exposed to alcohol. Conversely, higher levels of MR gene expression in the amygdala were associated with decreased anxiety and less compulsive alcohol drinking. Overall, these findings suggest that higher MR gene expression in the amygdala may protect against compulsive drinking or, conversely, that lower levels of MR gene expression may increase vulnerability for

anxiety-related compulsive drinking in the rat AUD model.

"The amygdala is a key regulator of emotion and stress and its adaptation to aldosterone signaling due to chronic alcohol drinking illustrates fundamental adaptations across organ systems that underlie the pathological state associated with AUD," said Kathleen Grant, Ph.D., study co-author and head of the Division of Neuroscience, Oregon National Primate Research Center at Oregon Health & Science University, Portland.

In a human study of about 40 individuals undergoing treatment for AUD, the researchers found that blood aldosterone concentrations were higher in individuals who continued drinking during the 12-week period, compared with those who were abstinent during the same time frame. For those who drank, the researchers found that aldosterone concentrations correlated with the amount of alcohol consumed during the study – higher drinking levels were associated with higher aldosterone concentrations. The researchers also found that increasing blood aldosterone concentrations correlated with increasing levels of both alcohol craving and anxiety.

Taken together, the researchers conclude that the findings provide support across three different species for a relationship between alcohol misuse, AUD, and specific changes in the aldosterone/MR pathway marked by increased circulating aldosterone and decreased mineralocorticoid receptor gene expression in the amygdala.

"Future studies should further investigate the mechanisms underlying the relationship between [alcohol](#) drinking and the aldosterone/MR pathway and whether this pathway might be targeted for the development of new pharmacotherapies for AUD," said Dr. Leggio.

More information: E G Aoun et al. A relationship between the

aldosterone–mineralocorticoid receptor pathway and alcohol drinking: preliminary translational findings across rats, monkeys and humans, *Molecular Psychiatry* (2017). [DOI: 10.1038/mp.2017.97](https://doi.org/10.1038/mp.2017.97)

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