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Researchers uncover brain pathway linked to impulsive behaviours

by Nanyang Technological University



Together with researchers from South Korea, Prof George Augustine from NTU's Lee Kong Chian School o...

Researchers from Singapore and South Korea have uncovered new details of a brain pathway that can cause impulsive behaviours.

Using mice, the research team led by Professor George Augustine from Nanyang Technological University, Singapore (NTU Singapore), discovered that impulsive behaviour is triggered when the brain signalling chemical dopamine is passed to an unexpected area of their brain.

The full route that dopamine signals take to produce an impulsive action has not previously been understood. To trace its pathway, Prof Augustine and his team used mice that had a specific set of dopamine receptors ("D2 receptors") genetically removed, effectively making their brains unable to detect dopamine signals.

The researchers artificially activated these receptors in specific parts of the brain, and the mice displayed impulsive behaviour when the signal was picked up by the amygdala – an almond-like structure deep in the brain.

What came next surprised the researchers: the amygdala's dopamine receptors in turn passed on the 'dopamine baton' to neurons that connect it with the 'bed nucleus of the stria terminalis' or BNST, a brain area not previously known to be involved in this pathway.

The BNST is a complex brain structure that orchestrates emotional and behavioural responses to stress. Its position in the pathway for impulsive behaviour is a new potential target for pharmaceutical developers, which could in turn lead to new treatments to manage this and other neuropsychiatric disorders.

"We have shown for the first time that impulsive behaviour in mice is only triggered when dopamine signals are received and passed on to an unexpected part of the brain – from the amygdala to the BNST," said Prof Augustine, a neuroscientist at NTU Singapore's Lee Kong Chian School of Medicine (LKCMedicine).

"This research shows that the amygdala serves as a key staging post in the dopamine pathway that triggers impulsive behaviour and confirms the role dopamine plays in regulating impulsivity," he said.

The research published in *Proceedings of the National Academy of Sciences (PNAS)*, last November, was conducted by Professor Augustine in collaboration with Professor Ja-Hyun Baik and colleagues from the Korea University. Researchers from Korea Institute of Science and Technology, and Institute of Molecular and Cell Biology in Singapore under the Agency for Science, Technology and Research (A*STAR) were also part of the team.

Commenting on the discovery, Assoc Prof Ong Say How, Senior Consultant and Chief, Department of Developmental Psychiatry, Institute of Mental Health, who was not involved in the research, said:

"This new finding potentially could usher in novel pharmacological treatments that specifically target the Dopamine D2 receptors located in the central amygdala and the BNST region, bringing about reductions in impulsivity and impulsive behaviours commonly seen in wide-ranging psychiatric conditions and disorders including conduct, anti-social and impulsive-control disorders. Reductions in impulsivity also imply more fore-planning and less risk-taking behaviours that often lead to accidental injuries as well as social, financial and legal problems later in life."

Prof Augustine plans to investigate further into the properties of the amygdala-to-BNST pathway, in order to pave the way for developing therapeutic strategies for controlling impulsive and compulsive behaviour.

"Through identification of a novel brain circuit for impulsivity we hope to encourage new research into treatment options for other neuropsychiatric disorders such as bipolar and attention deficit hyperactivity disorders (ADHD), and even depression," added Prof Augustine.

More information: Bokyeong Kim et al. Dopamine D2 receptor-mediated circuit from the central amygdala to the bed nucleus of the stria terminalis regulates impulsive behavior, *Proceedings of the National Academy of Sciences* (2018). DOI: 10.1073/pnas.1811664115

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