Boosting thalamic reticular nucleus function to treat ADHD and other disorders
24 March 2016, by Anne Trafton

The brain’s thalamic reticular nucleus (TRN), highlighted in the center, is responsible for blocking out distracting sensory input. "For autism, schizophrenia, and other neurodevelopmental disorders, it seems like TRN dysfunction may be involved in some patients," says Guoping Feng, the James W. and Patricia Poitras Professor of Neuroscience and a member of MIT's McGovern Institute for Brain Research and the Stanley Center for Psychiatric Research at the Broad Institute.

Feng and Michael Halassa, an assistant professor of psychiatry, neuroscience, and physiology at New York University, are the senior authors of the study, which appears in the March 23 online edition of Nature. The paper's lead authors are MIT graduate student Michael Wells and NYU postdoc Ralf Wimmer.

Paying attention

Feng, Halassa, and their colleagues set out to study a gene called Ptchd1, whose loss can produce attention deficits, hyperactivity, intellectual disability, aggression, and autism spectrum disorders. Because the gene is carried on the X chromosome, most individuals with these Ptchd1-related effects are male.

In mice, the researchers found that the part of the brain most affected by the loss of Ptchd1 is the TRN, which is a group of inhibitory nerve cells in the thalamus. It essentially acts as a gatekeeper, preventing unnecessary information from being relayed to the brain's cortex, where higher cognitive functions such as thought and planning occur.

"We receive all kinds of information from different sensory regions, and it all goes into the thalamus," Feng says. "All this information has to be filtered. Not everything we sense goes through."

If this gatekeeper is not functioning properly, too much information gets through, allowing the person to become easily distracted or overwhelmed. This
can lead to problems with attention and difficulty in learning.

The researchers found that when the Ptchd1 gene was knocked out in mice, the animals showed many of the same behavioral defects seen in human patients, including aggression, hyperactivity, attention deficit, and motor impairments. When the Ptchd1 gene was knocked out only in the TRN, the mice showed only hyperactivity and attention deficits.

**Toward new treatments**

At the cellular level, the researchers found that the Ptchd1 mutation disrupts channels that carry potassium ions, which prevents TRN neurons from being able to sufficiently inhibit thalamic output to the cortex. The researchers were also able restore the neurons' normal function with a compound that boosts activity of the potassium channel. This intervention reversed the TRN-related symptoms but not any of the symptoms that appear to be caused by deficits of some other circuit.

"The authors convincingly demonstrate that specific behavioral consequences of the Ptchd1 mutation—attention and sleep—arise from an alteration of a specific protein in a specific brain region, the thalamic reticular nucleus. These findings provide a clear and straightforward pathway from gene to behavior and suggest a pathway toward novel treatments for neurodevelopmental disorders such as autism," says Joshua Gordon, an associate professor of psychiatry at Columbia University, who was not involved in the research.

Most people with ADHD are now treated with psychostimulants such as Ritalin, which are effective in about 70 percent of patients. Feng and Halassa are now working on identifying genes that are specifically expressed in the TRN in hopes of developing drug targets that would modulate TRN activity. Such drugs may also help patients who don't have the Ptchd1 mutation, because their symptoms are also likely caused by TRN impairments, Feng says.

The researchers are also investigating when


Provided by Massachusetts Institute of Technology

This document is subject to copyright. Apart from any fair dealing for the purpose of private study or research, no part may be reproduced without the written permission. The content is provided for information purposes only.