

Complexity of NMDA receptor drug discovery target revealed

June 28 2018



Credit: Human Brain Project

Know your target. Especially if your target is coming into focus for treating diseases such as schizophrenia and treatment-resistant depression.

NMDA receptors, critical for learning and memory, are sensors in the



brain. Studying them in molecular detail is challenging, because they usually come in four parts, and the parts aren't all the same.

Researchers at Emory have been probing one variety of NMDA receptor assembly found in the cerebellum, and also in the thalamus, a central gateway for sensory inputs, important for cognition, movement and sleep. This variety includes a subunit called GluN2C—together with two partners, GluN1 and GluN2A.

The results are scheduled for publication in Neuron.

Outside of a living brain, NMDA receptor assemblies are typically studied with either two copies of GluN2C or two of GluN2A, but not with one of each, says senior author Stephen Traynelis, Ph.D., professor of pharmacology at Emory University School of Medicine.

"Our data suggest that GluN2C is rarely by itself," Traynelis says. "It's typically paired up with another GluN2 subunit. This means we really don't know what the properties of the main NMDA receptor in the cerebellum or the thalamus are."

Psychiatrists have become interested in GluN2C because it appears to decline in the brains of schizophrenia patients. Mice without adequate levels of GluN2C display abnormalities in <u>learning, memory and sensory</u> processing, which together resemble schizophrenia in humans. In addition, GluN2C appears to be important for the mechanism of ketamine, a drug being studied for its rapid anti-depressant effects.

Using drugs that are selective for particular combinations of NMDA receptor subunits, Traynelis' laboratory showed that an assembly of GluN2A and GluN2C is the dominant form in the cerebellum. When GluN2C is introduced into cortical neurons, it prefers to pair up with GluN2A, the researchers found. This raises the question in regions such



as the thalamus of whether GluN2C also appears with a partner GluN2 subunit. They also observed that the GluN2A-GluN2C assembly has distinct electrochemical properties.

Traynelis' laboratory, working with colleagues from Emory's Department of Chemistry, has <u>recently identified compounds that are selective</u> for GluN2C-containing assemblies, and determined which compounds are likely to be active against GluN2C when it is paired up with GluN2A.

"For GluN2C, we believe that the assembly that contains GluN1 and two different GluN2 subunits will be the receptor to study, whether it comes to pharmacology, biophysics, or clinical relevance," he says. Similar efforts are underway to identify the subunit composition of GluN2Dcontaining receptors, which may also be paired with other subunits.

More information: *Neuron* (2018). DOI: 10.1016/j.neuron.2018.06.010

Provided by Emory University

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