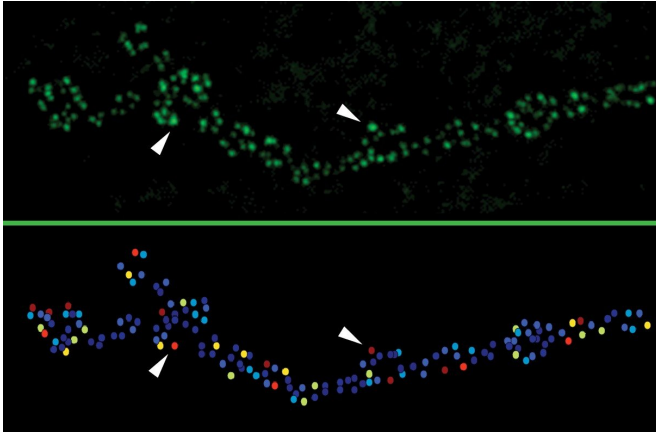


What separates the strong from weak among connections in the brain

10 July 2018



Top: Calcium channels (tagged green) along a neuron in a fly neuromuscular junction. Below: A map of active zone activity: Warmer colors indicate higher release probability with red being greater than 0.3 and deep blue being less than 0.1. Credit: Littleton Lab / MIT Picower Institute

To work at all, the nervous system needs its cells, or neurons, to connect and converse in a language of electrical impulses and chemical neurotransmitters. For the brain to be able to learn and adapt, it needs the connections, called synapses, to be able to strengthen or weaken. A new study by neuroscientists at MIT's Picower Institute for Learning and Memory helps to explain why strong synapses are stronger, and how they get that way.

By pinpointing the properties of synaptic strength and how they develop, the study could help scientists better understand how [synapses](#) might be made weaker or stronger. Deficiencies in synaptic development and change, or plasticity, have a role in many brain diseases such as autism or intellectual disability, said senior author Troy Littleton, Menicon Professor of Neuroscience in MIT's Department of Biology.

"The importance of our study is figuring out what are the molecular features of really strong synapses versus their weaker neighbors and how can we think about ways to convert really weak synapses to stronger ones," Littleton said.

In the study, published in *eLife*, Littleton's team used innovative imaging techniques in the model organism of the fruitfly *Drosophila* to focus on "active zones," which are fundamental components of synapses. The scientists identified specific characteristics associated with a strong connection on both sides of the synapse.

The team, led by postdoctoral researcher Yulia Akbergenova and graduate student Karen Cunningham, also studied how strong synapses and active zones grow, showing that those that have the longest to mature during a few critical days of development become the strongest.

Sources of strength

The team's study began with a survey of active zones at a junction where a motor neuron links up a muscle. About 300 active zones were present at the neuromuscular junction, which gave the team a rich diversity of synapses to examine.

Typically, neuroscientists study neural connectivity by measuring the electrical currents in the postsynaptic neuron after activation of the presynaptic one, but such measures represent an accumulation of transmission from many active zones. In the new study, the team was able to directly visualize the activity of individual active zones with unprecedented resolution using "optical quantal imaging."

"We optimized a genetically encoded calcium sensor to position it near active zones," Akbergenova said. "This allows us to directly visualize activity at individual release sites. Now we can resolve synaptic transmission at the level of

each individual release site."

Across many flies, the team consistently found that only about 10 percent of the active zones at the junction were strong, as measured by a high likelihood that they would release the neurotransmitter glutamate when the presynaptic neuron was stimulated. About 70 percent of the active zones were much weaker, barely ever releasing glutamate given the same stimulation. Another 20 percent were inactive. The strongest active zones had release probabilities as much as 50 times greater than weak ones.

"The initial observation was that the synapse made by the exact same neuron are not of the same strength," Littleton said. "So then the question became, what is it about an individual synapse that determines if it is strong or weak?"

The team ran several tests. In one experiment, for instance, they showed that it's not their supply of synaptic vesicles, the containers that hold their cache of glutamate. When they stimulated the presynaptic neurons over and over, the strong ones retained their comparatively higher likelihood of release, even as their synaptic vesicle supply was intermixed with those from nearby active zones.

The presynaptic tests that showed a difference had to do with measuring the rate of calcium influx into the active zone and the number of channels through which that calcium reaches the active zone. Calcium ions stimulate the vesicles to fuse to the membrane of the presynaptic cell, allowing neurotransmitters to be released.

At strong synapses, active zones had a significantly greater influx of calcium ions through a notably higher abundance of calcium ion channels than weak synapse active zones did.

Stronger active zones also had more of a protein called Bruchpilot that helps to cluster calcium channels at synapses.

Meanwhile, on the postsynaptic side, when the scientists measured the presence and distribution of glutamate receptor subtypes they found a dramatic difference at strong synapses. In the

typical weak synapse, GluRIIA and GluRIIB containing receptors were pretty much mixed together. But in strong synapses, the A subtype, which is more sensitive, crowded into the center while B was pushed out to the periphery, as if to maximize the receiving cell's ability to pick up that robust signal.

Might through maturity

With evidence of what makes strong synapses strong, the scientists then sought to determine how they get that way and why there aren't more of them. To do that, they studied each active zone from the beginning of development to several days afterward.

"This is the first time people have been able to follow a single active zone over many days of development from the time it is born in the early larvae through its maturation as the animal grows," Littleton said.

They did this "intravital imaging" by briefly anesthetizing the larvae every day to check for changes in the active zones. Using engineered GluRIIA and GluRIIB receptor proteins that glow different colors they could tell when a strong synapse had formed by the characteristic concentration of A and marginalization of B.

One phenomenon they noticed was that active zone formation accelerated with each passing day of development. This turned out to be important because their main finding was that synapse strength was related to active zone age. As synapses matured over several days, they accumulated more calcium channels and BRP, meaning that they became stronger with maturity, but only a few had the chance to do it for several days.

The researchers also wanted to know whether activity affected the rate of maturation, as would be expected in a nervous system that must be responsive to an animal's experience. By tinkering with different genes that modulate the degree of neuronal firing, they found that active zones indeed matured faster with more activity and slower when activity was reduced.

"These results provide a high resolution molecular and developmental understanding of several major factors underlying the extreme heterogeneity in release strength that exists across a population of active [zones](#)," Cunningham said. "Since the cohort of proteins that make up the presynaptic active zone in flies is largely conserved in mammalian synapses, these results will provide valuable insight into how active zone release heterogeneity might arise in more complex neural systems."

Provided by Massachusetts Institute of Technology

APA citation: What separates the strong from weak among connections in the brain (2018, July 10) retrieved 20 June 2019 from <https://medicalxpress.com/news/2018-07-strong-weak-brain.html>

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