

Researchers make significant advance in understanding calcium channel control of neurotransmitter release

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A, Cartoon depicting CaV2.1 α 1 subunit distal C-terminal interaction partners. B, left: Schematic view of stereotactic surgery to inject/coinject HdAd vectors expressing Cre + eGFP and CaV2.1 constructs + mCherry into the aVCN at age P1. Right: top: Experimental timeline from virus injection at P1 to electrophysiological recordings at P9-P11 prior to the onset of hearing (P12). Middle and bottom: schematic view of the viral constructs used, expressing either Cre + eGFP or CaV2.1 constructs + mCherry, respectively, driven by individual promotors. C, Calyx of Held terminals transduced with Cre + eGFP (top) and CaV2.1 + mCherry (middle). eGFP and mCherry signals overlap with those of a calyx of Held loaded with Lucifer Yellow via a patch pipette (bottom). Credit: Max Planck Florida Institute for Neuroscience

A new study conducted by researchers at the Max Planck Florida Institute for Neuroscience uncovers critical aspects of calcium channel function, overturning prevailing theories that explain variability in neurotransmitter release during neuronal communication.

Approximately eight years ago, former Research Group Leader at Max Planck Florida Institute for Neuroscience (MPFI) and current associate professor at the University of Iowa, Samuel Young Jr., Ph.D. set out to understand voltage-gated calcium (Ca²⁺) channel function, which determines the characteristics of neurotransmitter release and ultimately how the brain operates. To do so, Young and his team had to painstakingly develop new research methods to overcome severe technical challenges which were major roadblocks in the field. In a study published in the journal *eLife* on August 8, 2017, Matthias Lübbert, Ph.D. and R. Oliver Goral, Ph.D., post-doctoral researchers, and others in the Young Lab, describe how they manipulated voltage-gated Ca²⁺ channels and directly monitor their impact on Ca²⁺ signals in a presynaptic terminal for the first time, to test how they influence cellular



communication.

Neurons communicate with each other through electrical and chemical signals. An electrical signal called an action potential travels down the neuron to the synapse, a highly specialized structure at a point of contact between neurons, where information is transferred over a small space via chemical messengers. This causes voltage-gated Ca^{2+} channels - protein complex that form a pore in the cell membrane - to open, allowing Ca^{2+} to flow into the presynaptic compartment, the portion of the neuron that releases chemical messengers which impact neighboring cells. The influx of Ca^{2+} causes synaptic vesicles - packages of neurotransmitters - to merge with its membrane at specialized sites called active zones and release their contents. The neurotransmitters then travel across the gap and activate receptors on the second cell, influencing that cell's behavior.

The likelihood that an action potential will lead a neuron to release neurotransmitters, and the speed with which they are released, can vary considerably, depending on the quantity and location of the voltagegated Ca^{2+} channels in the presynapse. If voltage-gated Ca^{2+} channels are abundant and found close to the vesicles, the release will be more likely and more robust. The voltage-gated Ca^{2+} channels are made up of several subunits which combine to form a complex. The number and position of the channels is thought to depend on proteins acting on the alpha subunit, an individual component of the protein complex of the Ca^{2+} channel which contains many amino-acid sequences (motifs) that proteins can bind to.





A, Cartoons depicting CaV2.1 full transcript or mutants (top) with corresponding exemplary Ca2+ currents (bottom) triggered by 10ms voltage steps from ?70 mV to -40 mV in 10 mV steps and further to +60 mV in 5 mV steps. Credit: Max Planck Florida Institute for Neuroscience

Currently, the <u>molecular mechanisms</u> that regulate the number and organization of voltage-gated Ca^{2+} channels and their impact on information transmission in presynaptic terminals are not wellunderstood. "Voltage-gated Ca^{2+} channels are a major dictator of how information is transmitted in the brain, therefore it is imperative to understand the molecular mechanisms of Ca^{2+} channel function," explained Young. "Also, dysfunction of the Ca^{2+} channels is involved in wide array of neurological disorders such as migraine, epilepsy, and ataxia."

Scientists have identified certain proteins and sequences that they believed were responsible for direct interaction with the alpha subunit to control the number and organization of the channels in the presynaptic terminal. However, without the technology to make molecular perturbations and directly manipulate the alpha subunit, the studies that identified them relied on indirect observations.



Young and his team overcame these limitations. They modified Helper Dependent Adenoviral vectors (HdAd,) which are able to carry far more foreign DNA than commonly used viral vectors in the neuroscience field. These vectors express modified alpha subunit proteins while at the same time remove the endogenous alpha subunit of voltage-gated Ca²⁺ channels. By using these vectors in a mouse model, the team was able to disrupt the function of specific voltage-gated channels at specific times, without disrupting the endogenous alpha subunit expression in other areas, with unprecedented precision. The team focused on the Calyx of Held - a particularly large presynaptic terminal, which is a model system for understanding presynaptic function. Its size gives unparalleled experimental accessibility to record presynaptic activity compared to other presynaptic terminals in neurons. Using their cutting edge HdAd vectors in conjunction with their transgenic animals, and focusing on the Calyx of Held, they generated a platform to carry out a study which could simultaneously probe both the structure and function of these channels with unrivaled accuracy.





A, Cartoon depicting truncated regions in our CaV2.1 α 1 deletion mutants including the binding sites for CaV β 4, CASK, RBP, PXXP, RIM1/2 and Mint-1 along with the effects of C-terminal truncations on ICa. B, Averaged traces of RRP and total releasable pool measurements from mice expressing Cre + full transcript CaV2.1 rescue (grey), Δ 2365-2368 (cyan), Δ 2212-2368 (yellow), Δ 2061-2368 (purple), Δ 2042-2368 (green) or Δ 2016-2368 (blue). ICa (top) and EPSCs (bottom) triggered by 3ms and 30ms pulses, plotted on top of each other (n=10 for each group, except for Δ 2212-2368: n=8). Credit: Max Planck Florida Institute for Neuroscience

The new study overturned the popular theory which proposed crucial roles of certain proteins and binding motifs on the alpha subunit of voltage-gated Ca^{2+} channels for the regulation of abundance and neurotransmitter release. Moreover, it identified a completely different region of the alpha subunit which impacts positioning of the channels relative to synaptic vesicles but not the abundance, suggesting that the two characteristics are actually determined separately.

The results of the study will be of tremendous importance to understanding how neurons encode a broad diversity of information as well as causes, and possible treatments of Ca^{2+} channelopathies. "It's exciting," said Young, "our research has overturned current paradigms and possibly opened up a new way we think about the molecular mechanisms that control <u>neurotransmitter release</u>."

More information: Matthias Lübbert et al, A novel region in the CaV2.1 α 1 subunit C-terminus regulates fast synaptic vesicle fusion and vesicle docking at the mammalian presynaptic active zone, *eLife* (2017). DOI: 10.7554/eLife.28412



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