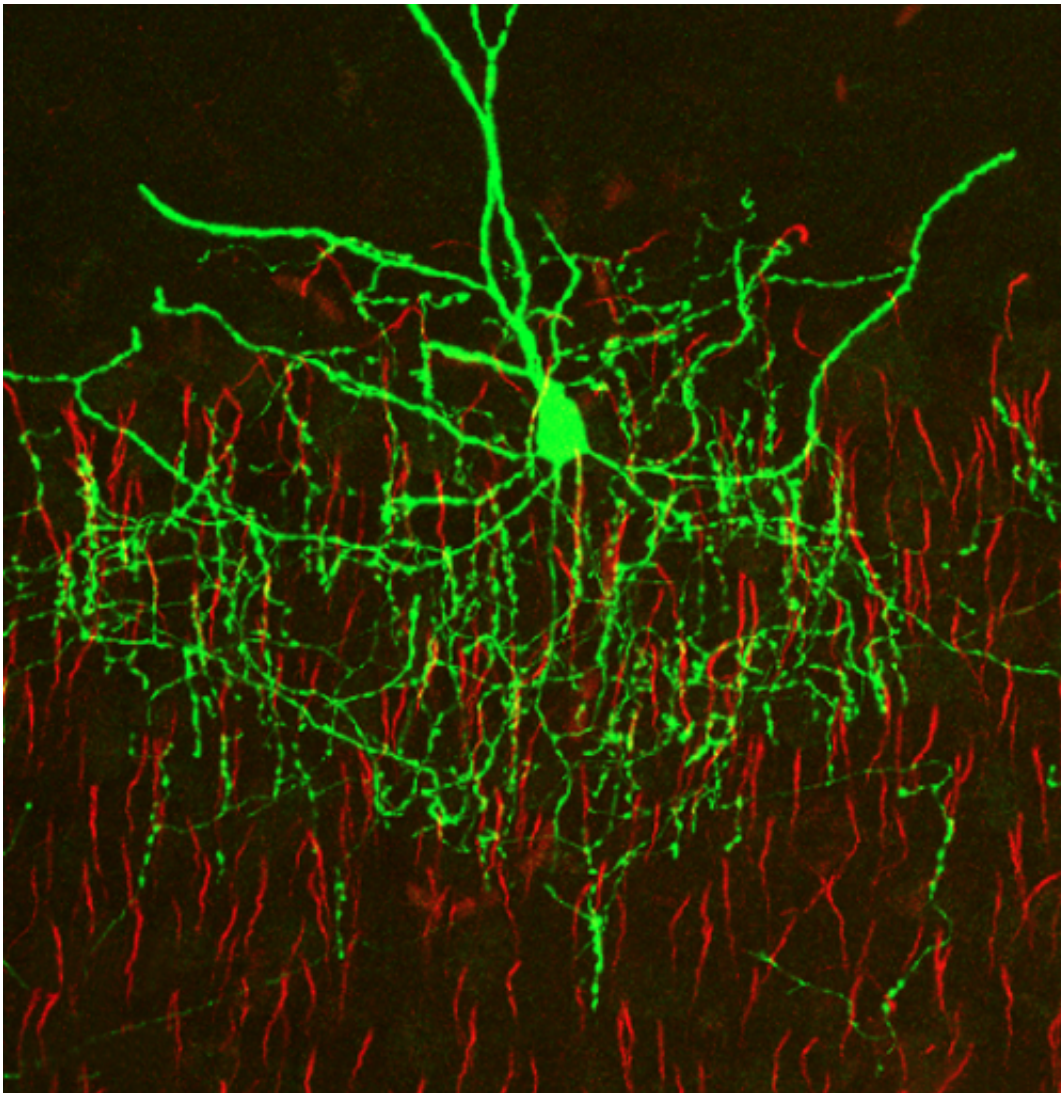


Scientists discover two proteins that control chandelier cell architecture

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Chandelier cells use their remarkable branching architecture to inhibit hundreds of excitatory neurons in the brain. CSHL researchers have identified two proteins that control the structure of chandelier cells, offering insight into how

these cells are regulated. Credit: Linda Van Aelst, Cold Spring Harbor Laboratory

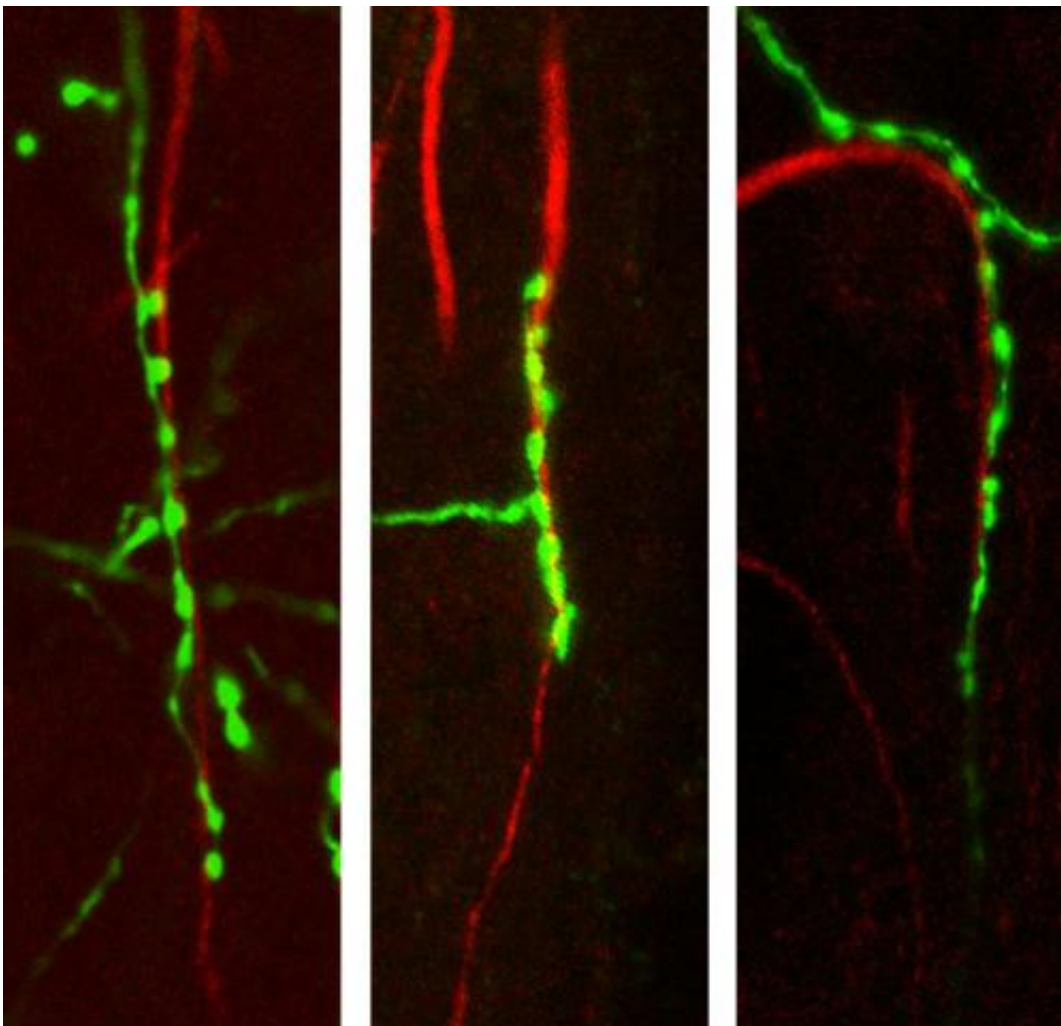
Chandelier cells are neurons that use their unique shape to act like master circuit breakers in the brain's cerebral cortex. These cells have dozens, often hundreds, of branching axonal projections – output channels from the cell body of the neuron – that lend the full structure of a chandelier-like appearance. Each of those projections extends to a nearby excitatory neuron. The unique structure allows just one inhibitory chandelier cell to block or modify the output of literally hundreds of other cells at one time.

Without such large-scale inhibition, some circuits in the brain would seize up, as occurs in epilepsy. Abnormal chandelier cell function also has been implicated in schizophrenia. Yet after nearly 40 years of research, little is known about how these important inhibitory neurons develop and function. In work published today in *Cell Reports*, a team led by CSHL Professor Linda Van Aelst identifies two proteins that control the structure of chandelier cells, and offers insight into how they are regulated.

To study the architecture of chandelier cells, Van Aelst and colleagues first had to find a way to visualize them. Generally, scientists try to find a unique marker, a sort of molecular signature, to distinguish one type of neuron from the many others in the brain. But no markers are known for chandelier cells. So Van Aelst and Yilin Tai, Ph.D., lead author on the study, developed a way to label chandelier cells within the mouse brain.

Using this new method, the team found two proteins, DOCK7 and ErbB4, whose activity is essential in processes that give chandelier cells their striking shape. When the function of these proteins is disrupted,

chandelier cells have fewer, more disorganized, axonal projections. Van Aelst and colleagues used a series of biochemical experiments to explore the relationship between the two proteins. They found that DOCK7 activates ErbB4 through a previously unknown mechanism; this activation must occur if chandelier cells are to develop their characteristic architecture.



Chandelier cells have traditionally been very difficult to image. CSHL Professor Linda Van Aelst and her colleagues developed a new method to visualize these cells in the brain. As seen here, the end of one axonal projection from a chandelier cell, in green, directly contacts a nearby excitatory neuron, shown in red. Credit: Linda Van Aelst, Cold Spring Harbor Laboratory

Moving forward, Van Aelst says she is interested in exploring the relationship between structure and function of chandelier cells. "We envisage that morphological changes are likely to impact the function of chandelier cells, and consequently, alter the activity of cortical networks. We believe irregularities in these networks contribute to the cognitive abnormalities characteristic of schizophrenia and epilepsy. As we move forward, therefore, we hope that our findings will improve our understanding of these devastating neurological disorders."

More information: "Regulation of Chandelier Cell Cartridge and Bouton Development via DOCK7 Mediated ErbB4 Activation" appears online in *Cell Reports* on January 16, 2014. The authors are: Yilin Tai, Justyna Janas, Chia-Lin Wang, and Linda Van Aelst.

Provided by Cold Spring Harbor Laboratory

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