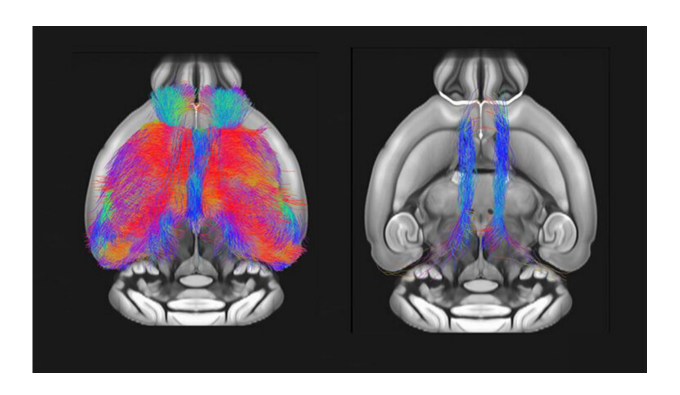


Autism gene linked to brain and behavior deficits in mice

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Shank3 mutants exhibit prefrontal mis-wiring and gray matter defects. Credit: Pagani et al., *JNeurosci* (2019)

Mice lacking the gene Shank3 display structural and functional deficits in the prefrontal cortex, finds a study published in *JNeurosci*. The research advances our understanding of one of the most common genetic risk factors for autism spectrum disorders.



Shank3 mutations are commonly observed in humans with autism and related developmental disorders. Previous animal research documents associations between Shank3 and basal ganglia dysfunction, which may contribute to repetitive behaviors characteristic of autism. In humans, Shank3 mutations have also been linked to intellectual and <u>language</u> <u>difficulties</u>.

Using magnetic resonance imaging in adult male mice, Marco Pagani, Alessandro Gozzi and colleagues demonstrate reduced <u>prefrontal cortex</u> connectivity and gray matter volume in Shank3-deficient animals.

The researchers report these brain differences are tightly linked to impaired social interaction with female mice.

These findings establish a role of Shank3 in maintaining prefrontal cortex connectivity, which may increase autism risk when disrupted.

More information: Deletion of Autism Risk Gene Shank3 Disrupts Prefrontal Connectivity, *JNeurosci* (2019). DOI: 10.1523/JNEUROSCI.2529-18.2019

Provided by Society for Neuroscience

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