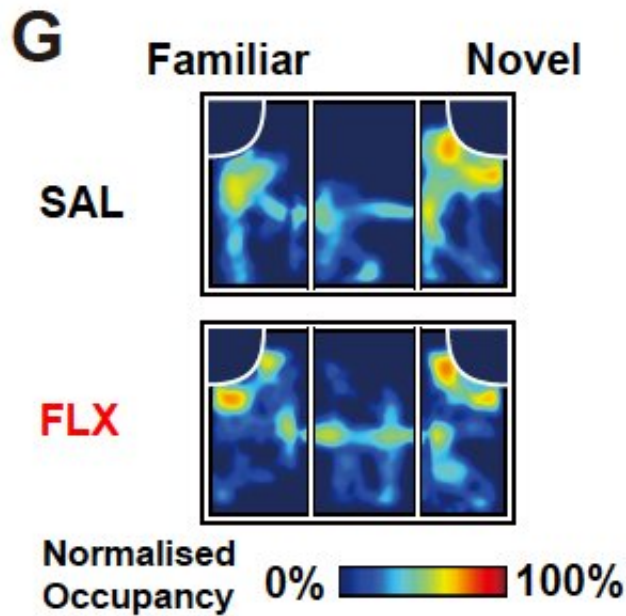


Possible link between autism and antidepressants use during pregnancy

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Representative heat map of results from the three-chamber social interaction test with a familiar mouse and novel mouse. Control animals show preference for new mice, but antidepressant-treated animals show no preference for familiar or new mice. Credit: Duke-NUS Medical School

An international team led by Duke-NUS Medical School has found a potential link between autistic-like behaviour in adult mice and exposure to a common antidepressant in the womb. They also identified a treatment that helped improve memory loss and social interactions, according to the new study published in the journal *Molecular Brain*.

Antidepressants are commonly prescribed for treating major depression and post-traumatic stress disorder, including in pregnant women. One of the most commonly prescribed antidepressants is fluoxetine, a [serotonin reuptake inhibitor](#). Fluoxetine can cross the placenta and is also detected in breast milk. Little is known about its safety during pregnancy, and not enough studies have been conducted on its long-term effects on offspring.

"Many human association studies have been conducted to investigate connections between antidepressant exposure during pregnancy and children with autism and attention deficit disorder (ADHD). But they have not been able to pinpoint a [causal relationship](#)," stated Associate Professor Hyunsoo Shawn Je, from Duke-NUS' Neuroscience and Behavioural Disorders (NBD) Programme, a senior and corresponding author of the study.

The team from Duke-NUS and their collaborators in South Korea and Singapore investigated [adult mice](#) born to mothers treated with fluoxetine (sold under the brand names Prozac and Sarafem) over a 15-day time period that corresponds to the [second trimester](#) in humans, in comparison with those born to mothers given normal saline as controls. They found key differences in behaviour. For example, the unexposed [mice](#) normally explored all three arms of a Y-shaped maze over a ten-minute time period and, over the courses of multiple arm entries, mice usually enter a less recently visited arm, while the fluoxetine-exposed ones were less inclined to explore unvisited arm.

In a second experiment, the mice were introduced to two juvenile mice, one after the other. When the second new mouse was introduced, mice that were not exposed to fluoxetine were more likely to only sniff the newly introduced mouse, recognizing that they had already met the first mouse. But the fluoxetine-exposed group sniffed both mice, indicating that they had impaired social novelty recognition.

The team then examined nerve signal transmission in the prefrontal cortex, a part of the brain involved in moderating social behaviour. They found impaired transmission caused by an overactive serotonin receptor. Treating fluoxetine-exposed mice with a compound that blocks the receptor alleviated their behavioural problems and improved their working memory.

The team next wants to examine autistic children born to mothers treated with [antidepressants](#) using [positron emission tomography](#) (PET) scans, an imaging technique used to observe metabolic processes in the body. If they also show enhanced serotonin receptor activity in the same area of the brain, the team plans to test whether FDA-approved serotonin receptor blockers can normalize their behaviours.

"The consensus among experts is that the rise in the number of people diagnosed with autism around the world is likely due to more awareness and testing rather than an increase in the prevalence of autism," noted Professor Patrick Casey, Senior Vice Dean for Research at Duke-NUS. "This collaborative study by our researchers offers a compelling case for a link between autism and antidepressant exposure in the womb in an animal model, and a possible mechanism that could potentially be exploited for future therapies."

More information: Weonjin Yu et al. Prenatal selective serotonin reuptake inhibitor (SSRI) exposure induces working memory and social recognition deficits by disrupting inhibitory synaptic networks in male mice, *Molecular Brain* (2019). [DOI: 10.1186/s13041-019-0452-5](https://doi.org/10.1186/s13041-019-0452-5)

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