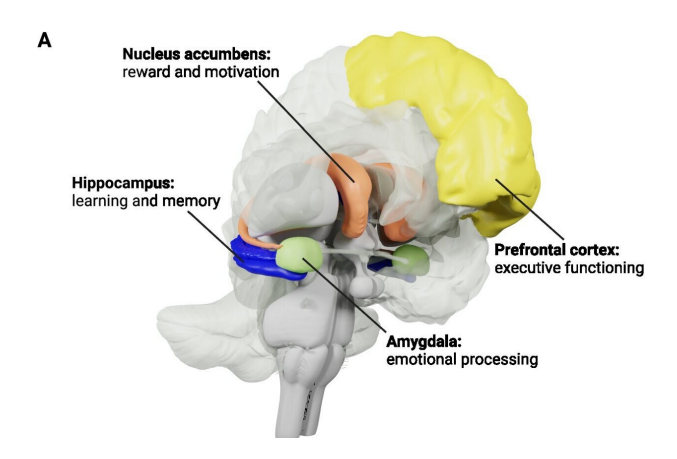


New study redefines how antidepressants aid in treating major depressive disorder

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Anatomy of key brain regions and network dysfunction involved in MDD. Credit: *Molecular Psychiatry* (2024). DOI: 10.1038/s41380-024-02625-2

Researchers from the University of Colorado Anschutz Medical Campus have established a new framework for understanding how classic



antidepressants work in treating major depressive disorder (MDD), reemphasizing their importance and aiming to reframe clinical conversation around their role in treatment.

The nature of the dysfunction at the root of MDD has been under investigation for decades. Classic antidepressants, like SSRIs (selective serotonin reuptake inhibitors, such as Prozac) cause an elevation in the levels of the brain chemical messenger, serotonin. This observation led to the idea that antidepressants work because they restore a chemical imbalance, such as a lack of serotonin.

However, subsequent years of research showed no significant decrease in serotonin in people with depression. While experts have moved away from this hypothesis due to lack of concrete evidence, this has led to a shift in public opinion on the effectiveness of these medications.

Antidepressants, such as SSRIs and serotonin and norepinephrine reuptake inhibitors (SNRIs) are still effective in alleviating depressive episodes in many patients, however. In a paper <u>published</u> in *Molecular Psychiatry*, researchers outline a new framework for understanding how antidepressants are efficacious in treating MDD. This framework helps clarify how antidepressants like SSRIs are still helpful, even if MDD isn't caused by a lack of serotonin.

"The best evidence of changes in the brain in people suffering from MDD is that some <u>brain regions</u> are not communicating with each other normally," says Scott Thompson, Ph.D., professor in the department of psychiatry at the University of Colorado School of Medicine and senior author. "When the parts of the brain responsible for reward, happiness, mood, <u>self-esteem</u>, even problem solving in some cases, are not communicating with each other properly, then they can't do their jobs properly.



"There is good evidence that antidepressants that increase serotonin, like SSRIs, all work by restoring the strength of the connections between these regions of the brain. So do novel therapeutics such as esketamine and psychedelics. This form of neuroplasticity helps release brain circuits from being 'stuck' in a pathological state, ultimately leading to a restoration of healthy brain function," said Thompson.

Thompson and colleagues liken this theory to a car running off the road and getting stuck in a ditch, requiring the help of a tow truck to pull the car out of its stuck state, allowing it to move freely down the road again.

Researchers are hoping <u>health care providers</u> will use their examples to bolster conversations with apprehensive patients about these treatments, helping them better understand their condition and how to treat it.

"We are hoping this framework provides clinicians new ways to communicate the way these treatments work in combating MDD," said C. Neill Epperson, MD, Robert Freedman endowed professor and chair of the department of psychiatry in the University of Colorado School of Medicine and co-author on the paper.

"Much of the public conversation around the effectiveness of antidepressants, and the role serotonin plays in diagnosis and treatment, has been negative and largely dangerous. While MDD is a heterogenous disorder with no one-size-fits-all solution, it is important to emphasize that if a treatment or medication is working for you, then it is lifesaving. Understanding how these medications promote neuroplasticity can help strengthen that message."

More information: Chloe E. Page et al, Beyond the serotonin deficit hypothesis: communicating a neuroplasticity framework of major depressive disorder, *Molecular Psychiatry* (2024). DOI: 10.1038/s41380-024-02625-2



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