

HHS Public Access

Author manuscript *Nat Med.* Author manuscript; available in PMC 2024 July 02.

Published in final edited form as:

Nat Med. 2023 February ; 29(2): 317-333. doi:10.1038/s41591-022-02197-0.

New and emerging approaches to treat psychiatric disorders

Katherine W. Scangos^{1,≅}, Matthew W. State¹, Andrew H. Miller², Justin T. Baker³, Leanne M. Williams^{4,5}

¹Department of Psychiatry and Behavioral Sciences, UCSF Weill Institute for Neurosciences, University of California, San Francisco, San Francisco, CA, USA.

²Department of Psychiatry and Behavioral Sciences, Emory University School of Medicine, Atlanta, GA, USA.

³McLean Hospital Institute for Technology in Psychiatry, Belmont, MA, USA.

⁴Department of Psychiatry and Behavioral Sciences, Stanford University, Stanford, CA, USA.

⁵Mental Illness Research Education and Clinical Center (MIRECC), VA Palo Alto Health Care System, Palo Alto, CA, USA.

Abstract

Psychiatric disorders are highly prevalent, often devastating diseases that negatively impact the lives of millions of people worldwide. Although their etiological and diagnostic heterogeneity has long challenged drug discovery, an emerging circuit-based understanding of psychiatric illness is offering an important alternative to the current reliance on trial and error, both in the development and in the clinical application of treatments. Here we review new and emerging treatment approaches, with a particular emphasis on the revolutionary potential of brain-circuit-based interventions for precision psychiatry. Limitations of circuit models, challenges of bringing precision therapeutics to market and the crucial advances needed to overcome these obstacles are presented.

An estimated 970 million people worldwide are living with a mental health disorder¹. Unlike many other chronic diseases, mental disorders often arise in childhood, adolescence or young adulthood and persist throughout life, impairing function and quality of life for many decades². As a result, mental illness is cumulatively the leading cause of disability with global costs in the trillions^{3,4}. Although many somatic and psychological treatments are available, access to quality care is woefully inadequate, and the effect sizes for most

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[©] Correspondence should be addressed to Katherine W. Scangos. katherine.scangos@ucsf.edu.

Competing interests

K.W.S. receives salary and equity options from Neumora Therapeutics. L.M.W. has received advisory board fees from One Mind PsyberGuide and the Laureate Institute for Brain Research and declares US patent applications 10/034,645 and 15/820,338: systems and methods for detecting complex networks in MRI image data. J.T.B. has received consulting fees and equity options from Mindstrong Health as well as consulting fees from Verily Life Sciences. A.H.M. has received consulting fees from Cerevel Therapeutics and Sirtsei Pharmaceuticals.

Peer review information *Nature Medicine* thanks Edward Bullmore, James Murrough and the other, anonymous, reviewer(s) for their contribution to the peer review of this work. Primary Handling Editor: Karen O'Leary, in collaboration with the *Nature Medicine* team.

therapeutic approaches are small⁵. The general lack of an actionable understanding of disease pathophysiology and the paucity of reliable diagnostic or therapeutic biomarkers also limit clinical practice to a trial-and-error approach that can be a lengthy, frustrating process for patients and clinicians and still leaves up to 60% of individuals experiencing treatment resistance⁶.

Historically, the most common 'biological' conceptualization of psychiatric illness has emphasized molecular and cellular mechanisms and therapeutic targets; more recently, however, brain circuit organization and function have emerged as both an explanatory model and as a basis for designing interventions. The rapid development of technologies and advanced analytical tools (both in the laboratory and in the clinic) and collaborations across neuroscience disciplines, along with a decade of success in psychiatric genetics and genomics, are challenging traditional diagnostic boundaries and the notion of therapeutic specificity. What is emerging is a viable path forward in linking an individual's symptoms and symptom clusters to mechanisms and treatments in a personalized manner. In this review, we outline how brain-stimulation techniques are advancing our understanding of psychiatric disease. We discuss new and emerging approaches to treatment based on a circuit understanding of psychiatric illness and consider the factors impeding delivery of precision therapeutics to the clinical market.

Circuit-level understanding of psychiatric disease

In the past few decades, the concept of psychiatric illness has transitioned from the notion of a 'chemical imbalance' to one that focuses on genetic risks, altered molecular and cellular development and function and disordered circuits. Traditionally, the term 'neural circuit' has referred to neuronal communication through synaptic connections and neurotransmission⁷. Here, we use the term to describe interconnections of vast numbers of neurons that make up the 'connectome' of the brain⁸. Techniques such as functional magnetic resonance imaging (fMRI) and electroencephalography (EEG) can quantify the functionally correlated regions of activation that define such circuits (Table 1). With the use of these tools, a set of core brain circuits have been identified that are consistently observed in the task-free state or when engaged by affective or cognitive tasks. These circuits include the default mode network (DMN), involved in task-free self-reflective processes; the negative affect circuit (also known as the 'reward' circuit), engaged by responsiveness to social and learned rewards; and the cognitive control circuit, engaged by tasks that require inhibition and selective control (Fig. 1).

Studies analyzing dysfunction within these core brain circuits as well as countless wellpowered genomic studies (focused on either common alleles or rare large-effect mutations) do not support crisp diagnostic boundaries between psychiatric conditions, as codified by the Diagnostic and Statistical Manual (DSM). Partly in response to these types of findings, the US National Institute of Mental Health developed the Research Domain Criteria (RDoC)⁹, which emphasize multiple levels of analysis, including neural circuits, behaviors, self-report measures, physiology, cells, molecules and genes, with the hope of improving pathophysiological understanding and disentangling circuit dimensions underlying the

heterogeneity of psychiatric disorders. An overarching goal of RDoC is to facilitate efforts to translate a circuit-level understanding of psychiatric diseases across basic science through clinical application and, ultimately, to inform neural targets for a more mechanistic and selective approach to developing treatments. Some argue that the way in which RDoC models psychiatric disease as a continuum of 'normal' and the fact that it does not include the natural history of any disorder (just a static view) moves away from a 'disease' model of psychiatric illness and may be a flaw of this paradigm¹⁰. Progress toward using RDoC-informed neural targets for more selective mechanistic treatment strategies has been slow.

As datasets increase in size, new analytical approaches such as machine learning and graph theory (Table 1) are enabling the discovery and quantification of new circuit patterns in psychiatric patient populations and are also beginning to uncover new disease phenotypes. Research consortia are using imaging and other modalities to develop brain-based markers that help select among treatments and predict response outcomes for depression and other disorders. RDoC and RDoC-'themed' constructs are being used to enrich trials for emerging treatments with patients more suited to the mechanism of action of the new compound and are also being used as endpoints for trial outcomes. To date, the most consistent or cross-validated predictive markers of treatment outcomes are specifically defined neural circuit targets, such as regions of interest implicated in the disorder and/or drug action.

New brain-stimulation and recording techniques represent a remarkable approach that may directly probe disease etiology at the circuit level. These techniques enable direct access to the living human brain, coupled with the ability to modulate circuit activity, and have already identified new principles of circuit–behavior relationships and demonstrated the feasibility of personalized circuit-targeted therapy. The approach could further inform biomarkers of disease severity and treatment prediction and could help uncover disease mechanisms to develop new treatments.

A brain circuit model can advance our understanding of types of disease etiology that may benefit from particular treatments, but it may not be universally applicable to all diseases or therapies at this time. In the sections that follow, we discuss promising new treatment approaches with a particular focus on those that fit the circuit-based framework; we also discuss emerging approaches for which this model has not yet been sufficiently investigated but which are of widespread interest in the field.

Brain stimulation as a research tool and therapy

Brain stimulation is an effective alternative to pharmacological agents for the treatment of psychiatric disorders; several common approaches are shown in Fig. 2. While all these approaches use electrical activity to modulate brain function (connectivity, activity, oscillatory activity), they vary in the strength and focality of the electric fields. Repetitive transcranial magnetic stimulation (rTMS) and transcranial direct current stimulation (tDCS) are non-invasive approaches that target the cortex. Deep brain stimulation (DBS) enables the direct modulation of deeper (subcortical) structures with the highest spatial and temporal precision and holds tremendous potential for the treatment of psychiatric disorders because it enables precise modulation of activity at any node along a dysfunctional circuit.

In DBS, an electrode is surgically implanted directly into a disease-specific target brain structure and stimulation is regulated by a pulse generator typically placed under the skin in the chest wall¹¹. Stimulation is delivered continuously at an intensity and frequency selected in the clinic by testing several parameters and observing the effect on a patient's symptoms. This procedure is commonly used to treat Parkinson's disease and can dramatically restore motor function in this context. In 2009, the US Food and Drug Administration (FDA) granted humanitarian device exemption to DBS treatment of obsessive-compulsive disorder (OCD) after a series of studies showed substantial reduction of symptoms¹². The anterior capsule was first targeted for DBS in patients with OCD based on its surgical corollary, the anterior capsulotomy (creating a lesion within this region was shown to alleviate symptoms). In 2003, Helen Mayberg showed that DBS targeting the subcallosal cingulate cortex (SCC), a region involved in emotional behavior, could treat major depression, even in patients who were resistant to other treatments¹³. Additional open-label trials targeting the SCC¹⁴ or ventral capsule-ventral striatum (VC/VS)¹⁵ (a region innervated with dopaminergic neurons known to be involved in reward processing and error prediction) showed similar effects and led to excitement at the prospect of treating major depressive disorder (MDD) by reversing dysfunctional brain activity. However, several disappointing randomized controlled trials followed. A trial of SCC DBS for MDD was halted prematurely after a 6-month futility analysis in 2012, and a second randomized trial of VC/VS DBS failed to show a significant improvement in symptoms compared to sham stimulation^{16,17}. The reasons for the failures are likely in part due to the lack of precision targeting or personalized treatment.

Several compelling advances have since emerged and promise improvement over traditional DBS. One is the emerging concept of a connectomic target, where white matter tracts or their intersections (rather than isolated brain areas) are stimulated with electric current to target disease-related brain circuits. Imaging is used to identify the correct location based on the anatomy of each individual¹⁸. For MDD, it has been shown that stimulation at the intersection of white matter tracts traveling through the SCC best predict treatment response¹⁹, and prospectively targeting this region has yielded a 73% response rate at 6 months and an 82% response rate at 1 year²⁰. Similarly, in OCD, multiple targets along the corticostriatal–thalamocortical loop that projects from the anterior capsule have shown benefit, and a retrospective analysis demonstrated that stimulation of a subsection of the anterior capsule that directly connects areas of the prefrontal cortex with the subthalamic nucleus predicted clinical improvement, regardless of lead location²¹.

Yet preclinical and clinical research have shown that stimulation in different brain regions elicits a wide range of emotions, suggesting that there is unlikely to be a single brain target sufficient to treat heterogeneous psychiatric disorders. Frustration with a lack of reproducibility of small-scale imaging investigations focused on anatomical regions of interest and disappointment with an attendant failure of clinical psychiatric research to provide definitive insights into mechanisms underlying psychiatric disorders have reinvigorated interest in using experimental manipulation of brain circuits. The use of electrical stimulation to understand brain structure–function relationships is not new; Wilder Penfield used this approach to map the somatosensory homunculus many decades ago²², but complex and varied behavioral effects and the invasive nature of this procedure limited its use for a time²³. Nonetheless, it has now been shown by multiple groups that stimulating

different parts of the brain elicits different affective experiences in animal models $^{24-27}$ and human studies $^{28-36}$.

One recent study performed electrical stimulation mapping of brain function (Table 1) before DBS implantation to uncover dysfunctional circuit activity underlying core depression symptoms at the individual level. In doing so, key principles of the relationship between depression symptoms, mood state and neural stimulation were uncovered and suggested the importance of linking the timing of stimulation to the presence of a negative symptom state³⁴. Multi-site stimulation across the corticolimbic circuitry elicited a wide range of rapid-onset subjective experiences (that is, anxiety, pleasure, energy) and was found to be 'dose' responsive, sustained beyond the stimulation period itself and context dependent. The emotional state of the patient at the time of stimulation determined the clinical response in a reproducible manner, suggesting that there is an important interplay between a patient's current symptom profile and the therapeutic effect of stimulation. How a region is stimulated also plays a role in its effect. Another ongoing study is using stimulation–response mapping to identify optimal stimulation parameters based on both their network and clinical effects³⁷.

During electrical stimulation mapping, intracranial recordings provide feedback on network connectivity and response to stimulation. Intracranial EEG activity marks the presence of some neurological symptoms; for example, pathological beta frequency oscillations are correlated with motor symptoms in Parkinson's disease³⁸. While there is evidence that scalp EEG captures states of arousal or attention³⁹, biomarkers of dynamic symptom states and targeted deep sources of these signals have remained elusive in psychiatry. In one study, a multi-day brain-mapping period enabled researchers to correlate intracranial EEG activity with symptom severity and identify an electrographic marker of a negative mood state in a single individual: namely, high-frequency activity in the amygdala, a brain region involved in emotion processing⁴⁰. A structurally and functionally connected depression subnetwork composed of the amygdala and regions of the VC/VS was identified; stimulation in the VC/VS node consistently improved core depressive symptoms and reduced high-frequency activity in the amygdala.

These findings led researchers to rationally design personalized DBS treatment. Using a device capable of both sensing neural activity and delivering stimulation, neural activity was continuously recorded from the amygdala and focal stimulation was automatically delivered to the VC/VS only when the electrographic biomarker of a severe symptom state was detected⁴⁰. This real-time, biomarker-driven therapy, referred to as closed-loop DBS, led to rapid remission in a pilot patient with only 30 min of stimulation per day. Further work in larger patient groups will be needed to determine interindividual biomarker variability and therapeutic effects of personalized closed-loop DBS. Nonetheless, the concept of treating a chronic disorder by temporally targeting symptom states defined by objective brain markers offers a new framework for circuit-targeted precision therapeutics in psychiatry. A new clinically available DBS system that enables sensing and recording of neural activity (Medtronic's Percept) is an exciting advance that may reveal new biomarkers of symptom severity in larger populations⁴¹. Furthermore, the ability to explore cellular populations and

projections underlying such biomarkers and circuits using optogenetics (Table 1) in animal models promises a realm of exciting experimental discoveries^{42,43}.

Circuit targeting with the non-invasive brain-stimulation technique rTMS is also accelerating (Fig. 2). This technique does not have the spatial or temporal resolution of DBS but can provide targeted treatment non-invasively in a standard psychiatry clinic and is now approved by the FDA for treatment-resistant MDD and OCD and anxious depression^{44,45}. In this treatment, a time-varying magnetic field is applied to the scalp through a transcranial magnetic stimulation (TMS) coil and induces an electric field in underlying tissue that can depolarize cortical neurons. The type of coil, its positioning on the brain and the frequency of magnetic pulses lead to different clinical and circuit-level effects^{46–48}. A standard treatment course consists of 20-30 sessions over 4-6 weeks with a session duration of 30-40 min for standard 10-Hz rTMS. A new stimulation paradigm, intermittent theta burst stimulation, was found to have to have similar efficacy in about one-tenth of the treatment session time⁴⁹. The dorsolateral prefrontal cortex (DLPFC), a region involved in higher cognitive functions, is targeted in MDD and has traditionally been identified through scalp measurements that can be inaccurate. Recent evidence suggests that treatment outcome can be enhanced by using imaging to target the circuit comprising the region in the DLPFC that has the highest negative connectivity with the anterior cingulate cortex (ACC), a region with a key role in cognition and emotion⁵⁰. In one study, the entire TMS treatment course was compressed into 5 d by delivering ten high-dose intermittent theta burst stimulation sessions per day with individual functional connectivity MRI-guided targeting, and the authors reported an astonishing 86.4% remission rate⁵¹. Advances that may further improve circuit-targeted rational treatment design include new coils that can reach deeper brain structures⁴⁷, EEG-synchronized therapy⁵² and multi-target TMS⁵³.

An electrical intervention in which circuit targeting does not apply is electroconvulsive therapy (ECT), a treatment developed in the 1930s that involves applying electric current to the scalp to intentionally trigger a brief seizure in a safe and controlled manner (Fig. 2). Although its mechanism remains incompletely understood, it may induce neuroplastic changes, allowing the brain to form new connections⁵⁴. Despite its lack of circuit specificity, it remains the most effective therapy for MDD with a remission rate greater than 70% (compared to 30% with typical antidepressants)⁵⁵. Side effects and stigma have limited its use, however, and there is substantial interest in developing new, non-invasive treatments that can target deep brain structures without the requirement for anesthesia. One emerging treatment is low-intensity focused ultrasound (Table 1), which uses pulses of sound to reversibly alter neural activity and can target deep brain circuit structures with high precision⁵⁶. A pilot study in depression demonstrated a reduction in worry and improved mood⁵⁷, and further studies are underway.

Big data to map the human brain and identify diagnostic and susceptibility markers

A parallel and complimentary approach to identifying dysfunctional circuits at the individual level is to search for disease or symptom-related pathologic circuit patterns in large

neuroimaging datasets at the population level. Taking many brain-based features, such as the connectivity of each voxel (a three-dimensional 'pixel' that represents a tiny cube of brain tissue) of an fMRI with every other voxel, and deriving latent structure from the associations between them has been approached by mathematics through the field of machine learning. The push to collaborate to generate large datasets and apply these analytic approaches was felt first in the field of psychiatric genomics and was met with considerable success (Box 1). The goal of these efforts was to identify markers of diagnostic susceptibility. The frequency of small-effect common alleles in the population and successful application of large-scale genome-wide association studies allow for the calculation of polygenic risk scores, a cumulative measure of individual genetic risk. While still aspirational in the clinic, this type of genetic profiling has the potential to drive genetically informed, clinically relevant research into differential diagnosis, natural history of disorders and treatment response⁵⁸.

A critical lesson learned from the replication failure of small candidate gene association studies is that very large neuroimaging datasets may be critical to identifying diagnostic or susceptibility markers. Reproducibility, or lack thereof, is a well-known concern in neuroimaging for group-averaged data due to low statistical power, software errors, the lack of appropriate correction for multiple comparisons and incomplete standardization of analysis methods⁵⁹. Multiple large-scale meta-analyses of task activation and resting-state or structural imaging have found shared pathology across disorders but no circuit-specific effect of diagnosis or RDoC domain^{60–62}. A recent analysis combining multiple datasets of structural imaging and resting functional connectivity interrogated correlations between these imaging measures and individual variation in subclinical 'symptoms' in healthy participants, revealing much smaller effect sizes than expected⁶³. These results highlight the need for caution when seeking to identify individual differences in behavior and both resting-state and structural imaging⁶⁴.

Multimodal quantitative meta-analyses that combine data from several MRI modalities may help distinguish true disease-specific circuit pathology from the numerous false positives in the literature. One such study was able to successfully identify MDD-specific findings from pooled, spatially normalized structural and functional data⁶⁵. Further integration of data across additional MRI modalities and EEG using similar coordinate-based meta-analytic methods holds promise for increasing reproducibility⁶⁶.

Emerging therapies and circuit-based biomarkers

The use of predictive or response biomarkers for emerging treatments may help to identify which individuals and mechanisms are targeted by each treatment and thereby improve statistical power and enhance effect sizes in clinical studies. According to National Institutes of Health (NIH)–FDA guidelines, a predictive biomarker identifies individuals likely to experience a benefit from treatment and a response biomarker indicates the biologic effect of a treatment without necessarily drawing conclusions about links between these biological mechanisms and clinical efficacy⁶⁷. Consortia focused on predictive and response markers of treatment outcomes include PReDicT (Predicting Response to Depression Treatment), EMBARC (Establishing Moderators and Biosignatures of Antidepressant Response in Clinical Care), iSPOT-D (international Study to Predict Optimized Treatment in Depression)

and FAST-MAS (Fast-Fail Trials in Mood and Anxiety Spectrum Disorders). One emerging focus is the use of fMRI treatment response targets as clinical trial endpoints (relevant to targeting more selective drug mechanisms), rather than classically defined DSM diagnoses. Another is illustrated in machine learning clustering approaches, with EEG datasets or combined imaging and treatment data for⁶⁸ identification of new disease subgroups^{69–71}. Across studies, cross-validated predictive models based on dysfunction within neural circuits have demonstrated the promise of increasing the current trial-and-error chance of responding to available treatments^{72–74}. Below, we provide several illustrative examples that highlight the relationship of four core brain circuits with treatment outcomes, measured using fMRI, and the associated efforts to identify circuit-based predictive and response biomarkers (Fig. 1). Table 2 lists treatments associated with specific circuits that have resulted in positive phase 2 trials or are further along toward FDA approval.

Anhedonia and the reward circuit

Anhedonia, a reduced ability to experience pleasure, is a symptom observed in multiple mental disorders^{75,76} and implicates dysfunction within the corticostriatal circuits, interplay with mesolimbic dopamine systems and dysfunction in constructs described within the RDoC positive valence system^{9,75}. In patients treated with currently approved antidepressants, higher resting-state connectivity between reward circuit regions of interest (measured by fMRI) and greater behavioral sensitivity (measured by computational modeling of performance on a reward task) has been found to identify individuals who respond to the norepinephrine- and dopamine-reuptake inhibitor bupropion, following nonresponse to the selective serotonin receptor inhibitor (SSRI) sertraline⁷⁷. Machine learning using task-evoked reward circuit activation has been found to predict response specifically to bupropion versus placebo 78 . These findings indicate the potential promise of reward circuit measures as predictive biomarkers. Changes in activation of the striatum, a key region in the brain's positive affect-reward circuit, also show potential as a response biomarker of new selective treatments. Treatment with pramipexole, a selective dopamine 3 receptor antagonist, has been associated with changes in both striatal activation and accompanying improvements in anhedonia^{79,80}. Changes in both striatal activation and anhedonia have also been observed within a study design that links the striatal response biomarker to the putative mechanism of action using a κ -opioid receptor antagonist (that stimulates striatal dopamine release)^{81,82} and a potassium channel opener (KCNQ2–KCNQ3 subtype)^{83,84}.

Negative affect and the corticolimbic circuit

Alterations in amygdala reactivity and negative affect circuit connectivity cut across multiple mental disorders and involve an interplay with serotonergic systems⁸⁵. During the processing of negative affect stimuli, hyporeactivity of the amygdala, quantified using a standardized task-based fMRI method⁸⁶, has been found to predict differential response to SSRIs versus serotonin–norepinephrine receptor inhibitors in patients with MDD^{72,87}. Amygdala reactivity also shows early changes following treatment with the SSRI escitalopram, showing its promise as a response biomarker⁸⁸. Several emerging treatments target changes in negative affect circuitry as a response biomarker. BI 1358894, an inhibitor of the transient receptor potential canonical (TRPC)4 and TRPC5 ion channels⁸⁹, has completed phase 1 evaluation using fMRI response biomarkers of amygdala activation

and emotion processing and is in phase 2 trials for MDD⁹⁰, post-traumatic stress disorder (PTSD) and borderline personality disorder. An inability to downregulate negative affect circuitry during sleep may contribute to a subtype of MDD with sleep–wake problems, excessive arousal and stress-related physical symptoms. The orexin 2 antagonist seltorexant is being developed for sleep–wake and arousal features of depression⁹¹, and orexin may help downregulate central amygdala reactivity that underlies these features via mediation of serotonergic pathways⁹².

Executive function and cognitive control circuitry

Disruption of the cognitive control circuit and executive processes has been reported across schizophrenia, MDD, PTSD, anxiety and attention disorders and may be another neural hallmark and treatment target^{93,94}. Noradrenergic and dopaminergic neurotransmitter systems are implicated in the modulatory interplay with cognitive control circuitry, among others. Emerging treatments designed to enhance cognitive processes and underlying cognitive circuit function, assessed using behavioral performance on cognitive tests and fMRI include iclepertin (also known as BI 425809). Iclepertin is a glycine transporter 1 (GlyT1) inhibitor that has FDA breakthrough therapy designation for cognitive impairment in schizophrenia⁹⁵.

Depression and the DMN

Another prominent circuit is the DMN. There is a mixed pattern of findings for DMN and medication outcomes; consistency has been challenged by different study-design strategies and analytic methods⁹⁶. DMN hypoconnectivity has been found to characterize persistent depressive disorder and to normalize with duloxetine, a commonly prescribed serotoninnorepinephrine receptor inhibitor, compared to placebo⁹⁷. Frontal nodes of the DMN have been found to normalize with standard antidepressants in late-life depression comorbid with anxiety⁹⁸, and the posterior-node DMN has been found to correlate with symptom improvement following treatment with the SSRI escitalopram in geriatric depression⁹⁹, indicating the potential of DMN node features as response biomarkers. Pretreatment taskfree DMN hypoconnectivity has been found to predict subsequent non-remission across commonly prescribed antidepressants, while intact DMN connectivity predicts remission to antidepressant treatment⁷³. DMN connectivity has also been studied as a putative circuit biomarker for new and emerging therapeutics. Baseline SCC connectivity within the DMN has consistently been implicated in response to TMS^{96,100}, with responders tending to show lower pre-TMS connectivity and a greater inverse relationship of the SCC to frontal regions of cognitive control circuits. Zuranolone (SAGE-217)^{101,102}, currently in phase 3 development for MDD and post-partum depression (PPD), was found to modulate resting DMN connectivity, and the putative mechanism of action is alteration of the γ -aminobutyric acid (GABA)ergic excitatory-inhibitory balance^{96,100}.

Emerging therapies that lack predictive, response and mechanistic biomarkers

Despite the appeal of applying a circuit-based model to all psychiatric disorders and treatments, we are challenged by our lack of mechanistic understanding of these. While

it may turn out that all brain disorders can be treated at the level of the neural circuit, neural circuit dysfunction is unlikely to be the etiology of every disease phenotype and therefore may not be the only or the optimal treatment target. No review of new and emerging approaches to treatment in psychiatry would be complete without the mention of rapidly acting therapeutics, hallucinogens or immunotherapeutics. We know that these treatments affect the brain, but they are not circuit specific and no clear circuit markers of disease have yet predicted treatment response, although such studies are underway. In the following sections, we discuss these treatments and the surrounding literature involving circuit effects while acknowledging where the circuit model falls short.

Rapidly acting therapies

Rapidly acting therapies such as ketamine represent the first new class of pharmacotherapies in decades (Table 2). Similar to some forms of brain stimulation, clinical effects of these pharmacotherapies are observed within hours. This incredible near-immediate reversal of severe pathology suggests that circuit reorganization or other mechanisms of action may occur on multiple timescales and both rapid-onset and sustained changes could be targeted by new treatments. Ketamine is a synthetic dissociative anesthetic and N-methyl-daspartate (NMDA) receptor antagonist¹⁰³. A placebo-controlled study published in 2006 demonstrated that a single 40-min intravenous ketamine infusion significantly improved treatment-resistant MDD compared to placebo, beginning just 2 h after infusion¹⁰⁴. Subsequent studies have confirmed this finding for MDD¹⁰⁵ and explored efficacy for suicidal ideation¹⁰⁶, PTSD^{107,108}, OCD¹⁰⁹ and PPD¹¹⁰. The intranasal formulation of the S-enantiomer, esketamine (Spravato), was approved by the FDA for treatment-resistant MDD (in conjunction with an oral antidepressant) and for MDD with acute suicidal behavior in 2019 and 2020, respectively^{111–114}. Neuroplastic processes, mediated through a surge in glutamate, are critical for ketamine's antidepressant effects, although recent evidence has implicated agonism of opioid receptors as well^{115,116}. Although the clinical effects of ketamine are promising for treatment-resistant disorders, we currently lack knowledge about circuit-based predictive and response biomarkers and the mechanisms that mediate clinical response. Neuroimaging studies of ketamine have revealed mixed results. In MDD, lower pre-infusion resting-state connectivity involving the SCC and frontal regions of the DMN, followed by relative normalization of connectivity after ketamine infusion, has been found to characterize responders (for reviews, see refs. ^{117,118}). However, pretreatment differences have not been replicated across region-of-interest and whole-brain data-driven analyses¹¹⁷. Connectivity between the posterior region of the DMN and the insula, which is a core region in negative affect and other circuits, has also been positively associated with post-ketamine clinical response¹¹⁷. By contrast, in task (as opposed to resting) fMRI, higher baseline SCC activity elicited by positive incentive trials is correlated with anhedonia improvement¹¹⁹. In studies of task fMRI with negative emotion tasks, ketamine has been found to reduce pre-infusion amygdala overactivity within the negative affect circuit, correlating with a change in anhedonia and overall depression improvement^{117,118}. Other dissociative anesthetics including dextromethorphan¹²⁰ and nitrous oxide¹²¹ have also been shown to rapidly and significantly reduce MDD severity in randomized trials and are thought to have similar circuit effects. Together, these findings hold promise for identifying circuit-based response biomarkers that also shed light on the mechanisms by which ketamine

exerts its clinical effects. More research is needed to disentangle systematic effects across studies, especially given that work to date has varied widely in terms of the timing of imaging relative to ketamine administration, imaging conditions, analysis methodology and population characteristics. It is also unknown whether mixed results reflect different subtypes of disorder or interactive effects of differing oral antidepressants in patients.

Positive allosteric modulation of GABA type A (GABA_A) (an effect that enhances GABA inhibition) represents a new mechanism of action that also has rapid-onset effects¹²². In 2019, brexanolone received FDA approval for PPD, the first drug approved specifically for this indication¹²³. While it must be administered by intravenous infusion over a 60-h period, newer formulations are under study, including the orally administered compound zelquistenel¹²⁴.

Psychedelics

In the last 10 years, interest in psychedelic medicine has surged. On the one hand, the rise of psychedelics clashes with the goal of circuit-targeted precision and rational treatment design. These plant-based compounds, in use for thousands of years, affect the brain globally. Yet, the renewed excitement regarding their use may in part be due to rise of circuit-based understanding of psychiatric disease and the role of neuroplasticity in reshaping circuitry. Psilocybin, a classic hallucinogen derived from the Psilocybe mushroom, received FDA breakthrough designation for the treatment of MDD in 2019 (ref. ¹²⁵). In therapeutic trials, two doses spaced 3 weeks apart were shown to be safe and to have comparable efficacy to escitalopram treatment for 6 weeks¹²⁶, with antidepressant activity observed within 24 h of administration. Active metabolites of classic hallucinogens act primarily as full or partial agonists of serotonin (5-HT_{2A}) receptors¹²⁷. Downstream activation of glutamate receptors, NMDA and a-amino-3hydroxy-5-methyl-4-isoxazolepropionic acid promote rapid neurotrophic and neuroplastic effects^{128,129}. During the hyperplastic period, increased global connectivity with specific connectivity of sensory cortices has been reported^{130–132}, followed by context-dependent reorganization after exposure¹³². Evidence suggests that therapeutic efficacy requires psychotherapy before and after exposure and is correlated with the positive quality of the psychedelic experience^{133,134}. Similar network restructuring after exposure has also been observed for lysergic acid diethylamide (LSD)¹³⁰, N,N-dimethyltryptamine (DMT, N,N-DMT)¹³⁵, ketamine¹¹⁸ and 3,4-methylenedioxy-methamphetamine (MDMA)¹³⁶. Clinical trials for psychedelics are challenged by the lack of an adequate placebo, cost of on-site monitoring by medical professionals, schedule 1 substance designation and the realityaltering and potentially addictive properties^{137,138}. Identification of non-hallucinogenic plasticity-promoting antidepressants with greater circuit specificity are under investigation and may be critical for a scalable path to the clinical market¹³⁹.

Immunotherapeutics

Immunomodulatory therapies for psychiatric diseases are receiving increasing attention based on an emerging literature elucidating the impact of systemic inflammation on the brain and behavior^{140,141} (Box 2 and Fig. 3). Inflammatory biomarkers are observed in a host of psychiatric diseases including mood and anxiety disorders, PTSD, psychotic

disorders and OCD^{140,142–145}. They have been associated with non-response to conventional antidepressants and antipsychotics and can predict response to ECT and ketamine^{146–149}. Aided by biomarkers of inflammation that can identify relevant subgroups of psychiatric patients, the opportunity now exists to target inflammatory pathways related to pathology of psychiatric disorders.

Based on studies administering inflammatory stimuli to humans, inflammation reliably engages neurocircuits that regulate motivational processing, psychomotor speed and threat sensitivity^{150–156}. Changes in these neurocircuits map onto behaviors of anhedonia, psychomotor retardation and anxiety and therefore represent attractive targets for circuit-based therapeutics across psychiatric disorders¹⁴⁰. For example, exogenous administration of the inflammatory cytokine interferon a to patients with infectious diseases and cancer as well as administration of typhoid vaccination or endotoxin to healthy volunteers have shown inflammation-induced decreases in activity within reward-related brain regions including the ventral striatum, as well as increases in activity within threat-related regions, including the dorsal ACC, the insula and the amygdala^{150–153,157}. In addition, increased endogenous inflammation has been shown to affect functional connectivity in corticostriatal and corticolimbic circuits related to motivational processing and psychomotor speed, as well threat sensitivity and avoidance of loss^{150,154,156,158}.

Studies in humans and laboratory animals have also revealed neurotransmitter changes related to the effect of inflammation on neurocircuitry. Indeed, inflammation-induced decreases in dopamine availability and release in the striatum have been associated with decreased effort-based motivation^{153,159}. Moreover, increases in glutamate in the basal ganglia impact local coherence of neuronal activity and ultimately decrease functional connectivity in multiple neural networks, including those involved in reward processing¹⁶⁰.

Studies have examined the efficacy of compounds with anti-inflammatory properties including inhibitors of cyclo-oxygenase 2 (for example, celecoxib), minocycline, omega 3 fatty acids and cytokine-targeted therapies as well as drugs that target immunometabolism (such as pioglitazone in patients with mood disorders and schizophrenia)^{161,162} (Table 2). Although meta-analyses of this literature have suggested modest efficacy, results have been underwhelming. Moreover, two recent studies failed to find separation of either celecoxib or minocycline from placebo in the treatment of depression^{163,164}. Nevertheless, the results to date should be interpreted with caution because virtually all studies failed to stratify patients on the basis of inflammation, there was no assessment of target engagement in the immune system or brain, and primary outcome variables were not aligned with the known impact of inflammation on specific brain circuits and behaviors¹⁴⁰. Inclusion of these design elements in future clinical trials will help realize the potential of immunomodulatory strategies to treat inflammation-induced transdiagnostic symptoms in psychiatric diseases.

Precision psychiatry: challenges and future outlook

Drug discovery for central nervous system disorders in general is plagued by higher failure rates, longer development times and higher costs than other areas of medicine, which has led many pharmaceutical companies to abandon the effort entirely¹⁶⁵. While precision

psychiatry may be a solution, there are challenges with implementing the approach. The emerging genetics and neuroimaging findings that do not support crisp DSM diagnostic boundaries are misaligned with the current diagnostic indication system used by payors and regulatory agencies. The FDA and the European Medicines Agency are actively developing a framework for the use of symptom domains and circuits as drug targets and clinical trial endpoints, placing emphasis on phase 1–2 proof-of-concept biomarker studies and studies that demonstrate subtype specificity. Further validation or qualification of circuit targets is needed for their acceptance by these regulatory agencies. Focusing on well-defined brain regions that tie to behaviors of interest offers one means to develop tractable patient-level measures suited to surrogate endpoints. Continuing to work together as a scientific community to build and extend collaborative consortia and define methodological standards to improve replicability will be critical. Integration of multimodal data including genetics, EEG, fMRI, tasks of cognitive function, clinical data and patient history using computational approaches may also help capture the full heterogeneity of symptom clusters and identify subgroups that respond preferentially to one treatment option or another.

Another challenge is the evolutionary distance between commonly used animal model systems and human brains. Preclinical models of psychiatric disorders lack predictive validity¹⁶⁶. Recent advances are expanding the armamentarium to study circuit-level biology in the human brain, including the ability to implant human brain-like collections of cells, called organoids, into rodent circuits to control behavior¹⁶⁷ and the ability to perform ex vivo studies of circuit function in intact post-mortem human brains¹⁶⁸. The experimental manipulation of circuit function in humans with focal electrical stimulation or other emerging forms of neuromodulation and direct assessment of changes in symptoms could provide tremendous potential for precision psychiatry, especially if the established circuit–symptom findings can then be backtranslated into animal models to identify causal molecular and cellular effects.

Another recent advance that may improve the success rate of candidate molecules is the ability to focus drug discovery on behavioral outcomes of circuit dysfunction rather than specific mechanisms of action. A time series analysis approach called motion sequencing derives a 'language' of natural segments or 'syllables' of drug-induced behavior in the rodent, using thousands of hours of recorded video from hundreds of animals. These can be used to discern linkages of drug-induced behavioral 'units' to specific neural circuits underlying clinical symptoms, such as hyperactivity and anxiety, and are capable of identifying drug class and dose¹⁶⁹.

Currently accepted outcome measures in clinical trials of psychiatric disorders are selfreported measures or clinician rating scales that are subjective accounts of a patient's disease state, require a look-back period of a week or more and are known to lack coherence with behavioral measures¹⁷⁰. They may obscure relevant symptom patterns that occur on a finer temporal scale or behavioral dynamics critical for identification of diagnostic or treatment response biomarkers¹⁷⁰. Validated, standardized measures that examine symptom severity on shorter timescales would also benefit the development of new brain-stimulation and rapid-acting therapeutics. With advances in sensor technology and artificial intelligence, digitally collected behavioral measures may provide one solution for this challenge and

are already showing promise for predicting DBS treatment parameters at the individual level^{171–174} and symptom severity at the population level¹⁷⁵. A next step will be to determine to what extent the digital signals might act as surrogates of underlying brain circuits, particularly given the host of interpretability issues around distinguishing clinically significant variance from variance due to measurement or contextual factors¹⁷⁶. Indeed, the NIH's recent Brain Behavior Quantification and Synchronization effort seeks to bring together researchers across scientific disciplines to quantify high-density behavioral data with simultaneously recorded brain activity and contextual data from the environment (https://event.roseliassociates.com/bbqs-workshop/). Managing ethical and privacy questions surrounding deep dynamic phenotyping of healthcare data will be critical¹⁷⁷.

Given the remarkable complexity of the human brain and the nature of psychiatric symptoms involving the most distinctively human aspects of function, it is not surprising that precision psychiatry has been slow to emerge. Yet recent progress in technologies, treatments and biomarkers is bringing us closer to this goal at a time when increasing the therapeutic armamentarium in psychiatry could not be of greater need. This review is not comprehensive; we have focused on new and upcoming therapeutic approaches that emerge from a deeper understanding of neural circuits, but foundational advances in other fields of psychiatry such as data-driven mining of medical records, computational psychiatry models and therapeutic approaches combining mechanistic with digital interventions are also moving the field forward. Together, these advances promise an auspicious future for psychiatry and propel the field forward toward a better quality of life for patients and their families.

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Box 1

Advances in understanding psychiatric genetics

Rigorous large-scale genome-wide association studies, enome-wide studies of copy number variation and whole-genome and whole-exome sequencing studies emerged in the first decade of the new millennium and, together with new statistical approaches, have led to major advances in the understanding of psychiatric genetics. A key finding has been the varying contribution of common (greater than 1% of the population) versus rare mutations across categorical diagnoses. For example, progress in early-onset neurodevelopmental disorders (including autism spectrum disorders and intellectual disability) has mainly come from the identification of rare, often spontaneous (de novo) copy number variants¹⁹⁷⁻²⁰⁰ and protein-damaging sequence mutations²⁰⁰⁻²⁰⁶ carrying large biological effects. By contrast, for many later-onset disorders (such as schizophrenia or bipolar disorder), progress has primarily involved the identification of common polymorphisms, mainly in non-protein-coding regions, contributing individually small effects, with hundreds to thousands of genetic variations conspiring to determine individual risk^{207–211}. Overall, the move to collaborative large-scale studies has identified definitive risk genes and loci and clarified the genetic architecture of a wide range of categorical diagnoses. At the same time, this work has directly challenged the notion that genes 'breed true' by demonstrating tremendous overlap in common genetic risks across disparate diagnoses and widely differing outcomes, in contrast to what is seen with identical, large-effect rare mutations found in unrelated individuals.

Box 2

The role of inflammation in psychiatric disorders

- Early studies in laboratory animals indicated that systemic inflammation induces a prototypical behavioral response referred to as 'sickness behavior', characterized by symptoms of psychiatric diseases including apathy, anxiety, anorexia, hypersomnia and cognitive impairment²¹².
- Increased inflammation in psychiatric disorders is believed to arise from factors such as obesity, diabetes, cardiovascular disease, cancer and infectious diseases; childhood trauma; and disruption of homeostasis of the gut microbiota¹⁴⁰.
- Chronic stress and inflammation may disrupt integrity of the blood-brain barrier, allowing direct access of peripheral inflammatory cytokines to neuronal systems in specific brain regions including the VS²¹³. These findings complement data demonstrating immune cell trafficking to the brain, where local release of cytokines can lead to these entering the brain parenchyma and cerebrospinal fluid, as well as the glymphatic pathways that support immune regulation of brain function^{214–216}.
- The effect of inflammation on neural circuitry may arise from changes in neurotransmitter levels or their metabolism. Decreased dopamine availability in the striatum and increased glutamate in the basal ganglia may in part be related to the induction of the kynurenine pathway and the production of quinolinic acid, which increases release and decreases reuptake of glutamate from astrocytes while binding to excitotoxic extrasynaptic glutamate receptors²¹⁷.
- A shift from energy-efficient oxidative phosphorylation to the energyexpedient glycolysis in activated immune cells, referred to as the Warburg effect, may activate several signaling pathways associated with reward circuit connectivity and symptoms of anhedonia and psychomotor slowing^{192,193} including mammalian target of rapamycin (mTOR).

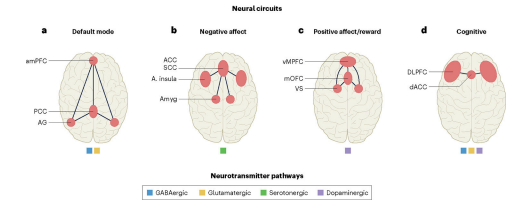


Fig. 1 |. Brain targets for selective pharmacotherapeutics.

Major circuits and neurotransmitter pathways that are disrupted in a wide range of psychiatric disorders and that represent targets for treatment, based on extant knowledge. **a**, The DMN is a major intrinsic circuit involved in self-reflective processes and defined by functional connectivity between regions in the anterior medial prefrontal cortex (amPFC), the posterior cingulate cortex (PCC) and the angular gyrus (AG) within the parietal cortex. Default mode connectivity implicates excitatory glutamatergic (yellow) and inhibitory GABAergic (blue) neurotransmitter pathways. b, The negative affect circuit is engaged by negative emotional stimuli such as threat and sadness and is involved in reactions to these stimuli, the experience of the emotional states that they induce and regulation of these reactions and experiences. Key regions are the amygdala (Amyg), the anterior insula (A. insula) and the ACC, encompassing the subcallosal cingulate cortex (SCC), also referred to as the subgenual anterior cingulate cortex. It is modulated by serotonergic neurotransmitter pathways (green). c, The positive affect circuit, also known as the reward circuit, is engaged by responsiveness to socially rewarding stimuli, learned rewards, anticipation of these stimuli and the motivation to expend effort to obtain such rewards. Key regions are the VS, encompassing the nucleus accumbens, the medial orbitofrontal cortex (mOFC) and the ventral medial prefrontal cortex (vMPFC). These corticostriatal regions of positive affect circuitry interdigitate with mesolimbic dopamine pathways (purple). **d**, The cognitive control circuit is engaged by higher cognitive functions such as working memory and is required to inhibit task-irrelevant responses under task demands. Cognitive control circuitry is centered on regions of the DLPFC and the dorsal ACC (dACC) and implicates, among others, dopaminergic, GABAergic, glutamatergic and noradrenergic (not shown) neurotransmitter pathways.

Brain-stimulation approaches

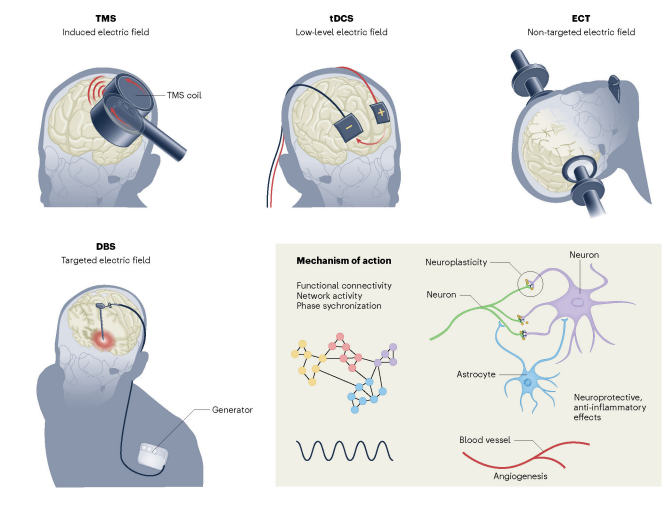
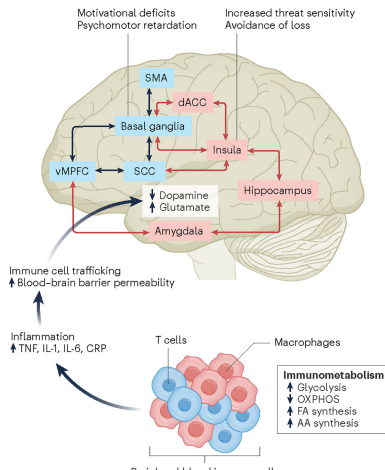


Fig. 2 |. Brain-stimulation techniques and mechanism of action.

In rTMS, a rapidly alternating magnetic field is used to induce a secondary electric field 2-4 cm deep in the cortical tissue, directly activating cortical neurons¹⁹⁵. The standard 'figure eight' coil is shown. In tDCS, a weak current (1-2 mA) is applied continuously to the scalp for 20-30 min, increasing (anodal tDCS) or decreasing (cathodal tDCS) excitability of the underlying cortical tissue¹⁹⁶. In ECT, an electric current is passed through the brain to safely induce a seizure while a patient is under general anesthesia. A standard course may include 12 ECT sessions over the course of 1-2 months⁵⁴. In DBS, electrodes are surgically implanted directly into the target brain structure, and stimulation is regulated by a pulse generator placed under the skin in the chest wall¹¹. Stimulation intensity, pulse width, frequency and polarity can be programmed to optimize clinical response. Standard continuous high-frequency stimulation may inhibit neuronal activity locally while activating inhibitory presynaptic terminals and axonal activity. The mechanism of action for brain stimulation (inset) remains incompletely understood but is hypothesized to be similar across modalities. It may change network activity, connectivity and phase-amplitude coupling, disrupt pathologic oscillations, induce neuroplasticity, neurogenesis and angiogenesis and alter inflammatory mediators.



Peripheral blood immune cells

Fig. 3 |. Inflammation as a therapeutic target in psychiatric disease.

There are many points along the pathway from the immune system to the brain that can serve as therapeutic targets. External and internal environmental factors can activate peripheral blood immune cells, which then undergo metabolic reprogramming that involves a shift from energy-efficient oxidative phosphorylation (OXPHOS) to energy-expedient glycolysis, allowing rapid growth and proliferation, supported by increased fatty acid (FA) and amino acid (AA) synthesis. Activated immune cells in turn produce inflammatory cytokines including tumor necrosis factor (TNF) that can increase permeability of the bloodbrain barrier by inhibiting production of claudin 5, a key protein in blood-brain barrier integrity, thus allowing direct access of inflammatory cytokines to the brain, produced in part by immune cells trafficking to perivascular spaces in the brain and the meninges. Once in the brain, inflammatory cytokines can decrease availability and release of dopamine in basal ganglia (for example, the striatum), while increasing the excitatory amino acid glutamate, whose activity may be further enhanced by activation of the kynurenine pathway. Altered neurotransmitter metabolism affects multiple brain regions, leading to disruption of neurocircuits that regulate motivation and motor activity (black arrows) as well as sensitivity to threat and loss (red arrows). These circuit-based behavioral biases in turn

contribute to symptoms of anhedonia, psychomotor slowing and anxiety, arousal and alarm. IL, interleukin; CRP, C-reactive protein; SMA, supplementary motor area.

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Table 1

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Tools to measure brain function	
fMRI	Measures changes in blood flow that occur with brain activity. Identifies brain regions and circuits activated during emotions and behaviors.
Diffusion tensor imaging	Measures diffusion of water in tissues to identify white matter tract architecture and connectivity.
Magnetic resonance spectroscopy	Measures brain metabolism and can identify the concentrations of chemical components within brain regions.
EEG	Measures spontaneous electrical activity of the brain from electrodes placed across the scalp. Can identify and characterize brain circuits in terms of their spectral content with high temporal resolution.
Magnetoencephalography	Measures the magnetic field produced by electrical currents inside neurons. Can be combined with an anatomical image of the brain to evaluate circuit function during emotions and behaviors.
Intracranial EEG	Measures spontaneous electrical activity of the brain from electrodes surgically placed within the brain. Provides direct measurement of brain activity with high spatiotemporal resolution.
Tools to manipulate brain function	
Intracranial brain mapping	Procedure where direct electrical stimulation of brain tissue is used to map the function of the brain. Behavior: brief electrical stimulation of cortical and subcortical regions at varying stimulation parameters elicits specific motor, cognitive or emotional responses. Links brain site to a particular behavior. Evoked potential: single pulses of stimulation are delivered to one region of the brain, and the response (evoked potential) is recorded in a distal region. Characteristics of the evoked potential indicate the strength and direction of brain connections.
TMS	A non-invasive form of brain stimulation that uses an alternating magnetic field to induce electric current in the underlying cortical tissue. Single TMS pulses can be used to map cortical activity and plasticity when paired with EEG.
Low-intensity focused ultrasound	A non-invasive and reversible treatment under study for psychiatric disorders. Neuronal activity is modulated through low-frequency sound waves focally concentrated on a deep brain structure.
Optogenetics	A preclinical research tool that uses light to control activity of neurons. Through genetic engineering, specific populations of neurons are modified to express light-sensitive ion channels (opsins). These neurons can then be optically activated in different behavioral settings to understand their function.
Tools to model brain function	
Machine learning	A field of mathematics that derives latent structure from associations within data. Supervised: prediction models trained to associate one set of inputs (for example, features of brain connectivity) with a desired set of labeled outputs (for example, treatment response). Robustness of findings are tested by splitting data into subsets for training and testing to quantify the extent to which hypotheses may generalize to unseen samples in the real world. Unsupervised: identification of patterns in data without a label. May be used to uncover subtypes of a psychiatric disorder.
Graph theory	A field of mathematics that studies the relationships between nodes (for example, brain regions) and their connections. Can identify and quantify connectivity patterns in brain circuits.

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Table 2 |

Emerging therapies and their putative circuit effects

	FDA-approved indication	Fast-track or breakthrough therapy designation	Indications in phase 3 trials	Additional indications (completed positive phase 2 trials)	Circuit effects
Selective dopamine 3 receptor	Selective dopamine 3 receptor modulation of reward circuit processes ^{178,179}	processes ^{178,179}			
D3R agonists	Restless leg syndrome, PD		MDD, TRD	Illness-related MDD, anhedonia in MDD and BD	Reward circuit including the VS, NAc, putamen, caudate, striatal interaction with the anterior insula ^{180,181}
Selective x -opioid receptor me	Selective κ -opioid receptor modulation of reward circuit processes 182	cesses ¹⁸²			
JNJ-67953964; BTRX-335140 (NMRA-140)	In development		MDD with anhedonia	MDD, smoking cessation	Reward circuit, including the VS ⁸²
KCNQ2-KCNQ3-selective potassium channel opener		modulation of reward circuit processes ^{83,84}	83,84		
Retigabine (ezogabine)	In development			MDD	Reward circuitry including the VS ⁸³
Selective inhibitor of TRPC4 and TRPC5 ion channels		implicating negative affect circuit processes 89	cesses ⁸⁹		
BI 135889	In development			MDD, PTSD, borderline personality disorder	Negative affect circuit, specifically the AMY^{90}
Selective orexin 2 receptor and	Selective or exin 2 receptor antagonism modulating negative affect circuit processes 91,92	affect circuit processes ^{91,92}			
Seltorexant (JNJ-42847922, MIN-202)	In development		MDD with insomnia	MDD, insomnia	Negative affect circuit, specifically the central AMY via orexin-mediated activation of 5-HT pathways ⁹²
Negative-allosteric modulator	· of the α-7 nicotinic acetylcholi	Negative-allosteric modulator of the a-7 nicotinic acetylcholine receptor modulating negative affect circuit processes ¹⁸³	affect circuit processes ¹⁸³		
BNC210		PTSD, fast-track designation (2019)	In development	PTSD, agitation in elderly	Negative affect circuit, including the AMY and AMY–ACC connectivity ¹⁸³
Glycine transporter 1 inhibite	or modulating glutamatergic pa	Glycine transporter 1 inhibitor modulating glutamatergic pathways through NMDA receptors and cognitive circuits and processes ¹⁸⁴	rs and cognitive circuits and F	processes ¹⁸⁴	
BI 425809		Cognitive impairment associated with SCZ, breakthrough therapy designation (2021)	SCZ, cognitive impairment in SCZ	Cognitive impairment in AD	Cognitive control circuitry
NMDA glycine-site partial ag	onist modulating glutamatergic	$NMDA$ glycine-site partial agonist modulating glutamatergic pathways through $NMDA$ receptors 185	otors ¹⁸⁵		
Rapastinel (zelquistinel)	In development		MDD	OCD	PFC, HPC ¹⁸⁶
Apimostinel				MDD	Unknown
Neuroactive steroid and \mathbf{GABA}_{A} receptor agonist modul	\mathbf{A}_{A} receptor agonist modulating e	lating excitatory-inhibitory GABAergic pathways and default mode connectivity ¹⁸⁷	pathways and default mode con	nectivity ¹⁸⁷	

	FDA-approved indication	Fast-track or breakthrough therapy designation	Indications in phase 3 trials	Additional indications (completed positive phase 2 trials)	Circuit effects
Brexanolone	PPD				DMN connectivity involving PCC ¹⁰²
Ganaxolone	Seizures (CDKL5)			PPD	and excitatory/inhibitory, GABAergic pathways
Zuranalone (SAGE-217)		Episodic MDD, breakthrough therapy designation (2021)	MDD, PPD	BD I/II	
Selective vasopressin V1a rece	ptor antagonist modulating pr	Selective vasopressin V1a receptor antagonist modulating prefrontal circuits involved in social cognitive processes and in regulating negative affective processes 188	al cognitive processes and in r	egulating negative affective J	processes ¹⁸⁸
SRX246				Intermittent explosive disorder	Social cognitive and threat regulation circuit, including mPFC, ACC and SCC ¹⁸⁸
NMDA receptor antagonist me	odulating DMN connectivity an	NMDA receptor antagonist modulating DMN connectivity and negative affect circuitry ^{117,118}			
Ketamine, esketamine	MDD in conjunction with oral antidepressant, MDD with acute SI		SI, PTSD, late-life MDD, PPD, SI in BD	MDD with alcohol use, OCD, PTSD, agitation, alcohol-use disorder	During treatment, disrupts hyperconnectivity of the DMN, decreases connectivity between the HPC
Dextromethorphan, AXS-05 (dextromethorphan-bupropion)		Breakthrough therapy designation for dextromethorphan–bupropion treatment of MDD (2019)	MDD, BD, SCZ, agitation in AD, irritability in HD, stimulant-use disorder	PTSD, smoking cessation	and other impot structures; may increase connectivity within the frontostriatal network ^{117,118} .
Nitrous oxide	In development		PPD	MDD, PTSD	
Serotonergic hallucinogens mo	dulating global internetwork (Serotonergic hallucinogens modulating global internetwork connectivity and social reward and negative affect processes ^{130–132,135}	nd negative affect processes ¹³	0-132,135	
Psilocybin		Breakthrough therapy designation for MDD (2019)	MDD	Illness-related anxiety/ depression	Global increase in internetwork connectivity during treatment, increased
TSD				Illness-related anxiety	connectivity within the DMIN following treatment
Ayahuasca/DMT				MDD	
Entactogen with serotonergic activity modulating fear		memory extinction and reconsolidation 136	n ¹³⁶		
МDМА		MDMA-assisted psychotherapy for PTSD, breakthrough therapy designation (2017)	MDMA-assisted psychotherapy for PTSD, PTSD	Illness-related anxiety, SAD in autistic adults	Facilitates fear memory extinction, modulates fear memory reconsolidation, altered activity in frontolimbic regions ¹³⁶
Brain stimulation with target-	specific network effects mediat	Brain stimulation with target-specific network effects mediated by direct or indirect activation of neural activity	n of neural activity		
rTMS	MDD, OCD, chronic migraine pain, smoking cessation, anxious depression		Alcohol-use disorder, SCZ, GAD, TBI, BD, SI	Panic disorder and MDD, PPD, Tourette's syndrome, PTSD, cocaine dependence	Decrease in SCC activity, change in SCC-DLPFC and DLPFC-VMPFC connectivity, decrease in SCC-DMN connectivity ¹⁸⁹
uDCS	In development		MDD, SCZ, BD, gambling disorder	Cocaine dependence, alcohol-use disorder	Increase in activation of VAN, increase in VAN-FPN connectivity during working memory task and DMN-FPN connectivity during language task ¹⁹⁰

Nat Med. Author manuscript; available in PMC 2024 July 02.

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		therapy designation	trials	(completed positive phase 2 trials)	
DBS	OCD, humanitarian device exemption (2009)		MDD	Tourette's syndrome	Stimulation-site-specific network modulation (OCD ¹⁸)
ECT	Labeled class II for catatonia, severe depressive episode of MDD or BD, 2018); class III (premarket approval) for SCZ, schizoaffective disorder, mania			SCZ	Hippocampal plasticity, modulation of corticolimbic circuitry via interaction with thalamocortical, cerebellar networks ¹⁹¹
Anti-inflammatory therapies targeting cognitive, rewa	rgeting cognitive, reward and	ird and DMNs ^{192,193}			
Rapamycin				MDD (with ketamine)	May modulate corticostriatal circuits and reward pathway
Celecoxib	In development		BD, MDD	SCZ	Alters HPC circuitry involved in spatial learning
Minocycline	In development		BD, SCZ, MDD		Alters HPC circuitry involved in spatial learning
Omega 3 fatty acids	In development		MDD, ADHD, SCZ, PTSD, alcohol dependence		Lower DMN and FPN hyperconnectivity
Anti-cytokine treatment (infliximab, losmapimod)				MDD, BD	Decrease in limbic activation (insula, cingulate cortex)
Pioglitazome				MDD, cocaine dependence, autism	Alters HPC circuitry involved in memory
Digital therapy ¹⁹⁴					
Digital pill			Substance-use disorder	MDD, SCZ, BD	Unknown
Digital biomarker-based detection and intervention system			Substance-use disorder	MDD	Unknown
Psychoeducation and intervention apps			SI	MDD	Unknown
Gaming/virtual reality	ADHD (marketing authorization)			MDD, autism	Unknown

vasopressin receptor 1A; AMY, amygdala; HPC, hippocampus; PFC, prefrontal cortex; mPFC, medial prefrontal cortex; VMPFC, ventromedial prefrontal cortex; NAc, nucleus accumbens; VAN, ventral attention network; FPN, frontoparietal network. IKD, treatment-resistant depression; BD, bipolar disorder; SCZ, schizophrenia; PD, Parkinson's disease; AD, Alzheimer's disease; SL, suicidal ideation; HD, Huntington's disease; SAD, social anxiety disorder; GAD, generalized anxiety disorder; TBI, traumatic brain injury; ADHD, attention-deficit hyperactivity disorder; D3R, dopamine receptor 3; CDKL5, cyclin-dependent kinase-like 5; VIa,

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