A selective review of glutamate pharmacological therapy in obsessive–compulsive and related disorders

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Abstract: Glutamate, an excitatory central nervous system neurotransmitter, is emerging as a potential alternative pharmacological treatment when compared to gamma-aminobutyric acid (GABA)-, dopamine-, and serotonin-modulating treatments for neuropsychiatric conditions. The pathophysiology, animal models, and clinical trials of glutamate modulation are explored in disorders with underlying inhibitory deficits (cognitive, motor, behavioral) including obsessive–compulsive disorder, attention deficit hyperactivity disorder, Tourette syndrome, trichotillomania, excoriation disorder, and nail biting. Obsessive–compulsive disorder, attention deficit hyperactivity disorder, and grooming disorders (trichotillomania and excoriation disorder) have emerging positive data, although only scarce controlled trials are available. The evidence is less supportive for the use of glutamate modulators in Tourette syndrome. Glutamate-modulating agents show promise in the treatment of disorders of inhibition.

Keywords: glutamate, obsessive–compulsive disorder, attention deficit hyperactivity disorder, trichotillomania, excoriation disorder, modulation

Introduction

The use of glutamate-modulating drugs for neuropsychiatric conditions is emerging as an alternative to three decades of use of dopamine-, serotonin-, and gamma-aminobutyric acid (GABA)-modulating drugs in neuropsychiatry.1–4 While such use is still nascent in clinical settings, there is promise of significant impact on improved outcomes for patients. Research in genetic epidemiology and neurophysiology is pinpointing glutamate, the essential excitatory central nervous system (CNS) neurotransmitter, as a direct target for pharmacologic manipulation. In this selective review, we present current drug trial data in disorders that exhibit features of a lack of inhibitory control, including cognitive (obsessive–compulsive disorder [OCD]), motor (Gilles de la Tourette syndrome [TS]), behavioral (attention deficit hyperactivity disorder [ADHD]), and grooming (trichotillomania [TTM], excoriation disorder [ExD]) domains.

Obsessive–compulsive disorder

OCD is characterized by the recurrence of unwelcome and intrusive ideations, urges, or images (obsessions), as well as repetitive, rigid behaviors (compulsions).4 Obsessions can include fears related to contamination, religious, and sexual themes. Compulsions include behaviors such as washing, checking, counting, ordering, and hoarding. Obsessions and compulsions exist in symptom groupings or dimensions, with the four most commonly reported OCD symptom dimensions being: 1) contamination/washing; 2) aggressive/sexual/religious obsessions; 3) counting/repeating/checking compulsions;
and 4) hoarding tendencies.\textsuperscript{6} The prevalence rate of OCD in adults in the United States is 2.3%,\textsuperscript{7} while it is lower in pediatric populations. In one study of 10,438 children in the United Kingdom, the prevalence rate of OCD was 0.25%.\textsuperscript{8} Standard pharmacologic treatment consists of serotonin reuptake inhibitors (SRIs), with moderate efficacy in children. A primary treatment approach in children with OCD is cognitive behavioral therapy (CBT) with exposure to OCD-related fears (exposure–response prevention [ERP]). The exposure paradigm is thought to be dependent upon glutamate-facilitated extinction learning.\textsuperscript{9} With medication and CBT/ERP, up to 54% of children with OCD experience clinical remission, but only 39% experience it with CBT alone and 21% with medication alone.\textsuperscript{10}

A developmentally-mediated brain dysplasia in cortico–subcortico–cortical networks has been postulated to underlie pediatric OCD.\textsuperscript{11} Given that glutamate signaling is critical in early brain development through the facilitation of neuronal proliferation, migration, and differentiation, the formulation of the glutamate hypothesis of OCD relies on a dysregulation in developmental glutamate signaling homeostasis.\textsuperscript{12} A rich expression of glutamate and GABA neurotransmitter activity in the fetal and postnatal brain supports early neuronal growth and maturation. Metabotropic glutamate receptor (mGluR)1, for example, has high levels of expression in the fetal brain, especially in the striatum, with subsequent progressive decreases, while mGluR4 is expressed mostly in the adult brain.\textsuperscript{13} In a synchronized manner, both ionic and mGluRs are expressed in specific brain regions at specific times during brain development.\textsuperscript{14} The excitatory effect of glutamate neuronal networks makes it important for effective efferent activity, but it also creates the potential for excitotoxicity.\textsuperscript{15} In line with the glutamate hypothesis, functional imaging studies of OCD have shown an overactive cortico–striato–thalamo–cortical (CSTC) loop, with glutamate playing a central excitatory role (cortex, thalamus), while GABA and other inhibitory neurotransmitter systems are operant in the striatum and other subcortical regions.\textsuperscript{16} Overactivity of the right caudate in OCD and the limbic CSTC\textsuperscript{17,18} is postulated to be a parallel dysfunction in glutamate homeostatic systems, such as the glutamate transporter family. Among members of this family of transporters, SLC1A1 in chromosome region 9p24, has emerged as an intriguing positional candidate gene for OCD. Multiple studies support the association of OCD with SLC1A1 variants,\textsuperscript{19–21} including specific possible functional mutations,\textsuperscript{22} but a more recent meta-analytic study has lent less support to its impact.\textsuperscript{23} In the postsynaptic membrane, receptor conformation variants are plausibly linked to OCD. The NMDAR2B component of the N-methyl-D-aspartate (NMDA) receptor is codified by GRIN2B in chromosome region 12p12. GRIN2B variants have been linked to OCD\textsuperscript{24,25} and following the glutamate pathway’s downstream connections, variants in BDNF have also been linked to OCD. These BDNF variants include SNP rs2883187,\textsuperscript{26} SNP rs1519480,\textsuperscript{27} and the Met allele of Val66Met.\textsuperscript{28} The evidence is thus mounting for a direct role of glutamate in the pathophysiology of OCD (Table 1); however, it is too early to determine whether glutamate dysregulation is a sign of intracortical excitability due to aberrant CSTC aberrant inputs,\textsuperscript{29} another unknown glutamatergic abnormality, or a reactive effect of the disorder itself on brain function.\textsuperscript{30}

Animals have been heuristically useful in understanding the pathophysiologic mechanisms in OCD that may lead to drug discovery. The inhibition of marble-burying behavior has been used as a traditional model to test for anxiolytic drugs and OCD-related hoarding behaviors. In a test specific to anxiolytic effects, the mGluR5 (group 1) mediates marble-burying behavior in mice, and the mGluR5 antagonist, MPEP, able to inhibit these behaviors.\textsuperscript{31} Similarly, the marble-burying behaviors are also mediated by group 2 mGluRs (mGluR2 and mGluR3).\textsuperscript{32} Additional support for glutamate-related animal models in OCD is provided by Sapap3 mutant mice, a knockout (KO) mouse that alters a synaptic density-associated protein. Sapap3 mutant mice demonstrate an excess of grooming behaviors, which is decreased significantly with the use of the selective SRI (SSRI) fluoxetine.\textsuperscript{33} Additionally, in these KO mice, the surface expression and activity of mGluR5s are increased, suggesting that glutamate-driven overactive mGluR5 transmission unfavorably alters synaptic plasticity. In effect, a positive allosteric modulator of mGluR5 reproduces this effect in wild-type mice.\textsuperscript{34} Interestingly, mGluR5 receptors have functional connections to BDNF, DLGAP1, and GRIN2B, as shown by a bioinformatics approach (http://www.string-db.org). In the rat attenuation model of OCD, decreasing rewards to lever pressing is used to identify those animals who “perseverate” in pressing behavior despite lower rewards. The excessive response can be abolished with fluoxetine and other serotonin-enhancing agents, but not by haloperidol or benzodiazepines.\textsuperscript{35} In a follow-up paradigm, the NMDA agonist D-cycloserine (DCS), but not an NMDA antagonist, helped attenuate the perseverative response, supporting a role for NMDA-facilitated learning of the extinction process.\textsuperscript{36}
Clinical trials of glutamate-modulating drugs in OCD

Amantadine

Amantadine (1-adamantanamine hydrochloride) is an antiviral drug that was formerly, but no longer, used for influenza prophylaxis. Its current uses are to improve alertness and arousal in post-traumatic brain injury in children, improve executive dysfunction in patients with Alzheimer’s dementia, and treat the early stages of Parkinson’s. The mechanism of action is thought to consist of releasing dopamine from the presynapse, in addition to possible negative modulation of the NMDA receptor. While it can be neuroprotective due to its glutamate antagonism properties, amantadine has also resulted in acute neuropsychiatric side effects such as hallucinations and confusion. A case study reported that a treatment-refractory patient with OCD responded to amantadine (200 mg/day) added to clomipramine (225 mg/day). In an open-label study of amantadine in eight patients with OCD who had failed one SSRI trial, Yale–Brown Obsessive Compulsive Scale (Y-BOCS) scores improved for compulsions (15.3±3.2 versus 10.6±4.7; P<0.02; degrees of freedom [df]=7; t=2.36) and obsessions (12.7±3.3 versus 8.1±5; P<0.05; df=7; t=2.36).

D-cycloserine

DCS is an analog of D-alanine and an NMDA receptor partial agonist at the glycine-binding site. It has been well established that DCS has the ability to contribute to fear extinction learning. DCS facilitates conditioned fear extinction in rats when injected in the amygdala or systemically, suggesting that treatments that activate NMDA receptors can promote the learning of extinction learning. The sum of preclinical studies thus supports the blocking of extinction learning by NMDA antagonists and its facilitation by agonists such as DCS. In the treatment of OCD, DCS has been combined with ERP as a treatment option for nonresponders to traditional ERP. DCS administered 2 hours prior to ERP can produce an accelerated decrease in obsession-related fear ratings in comparison to the placebo across four ERP therapy sessions. This acceleration of response was also seen in another study of 16 adults with OCD, nine of whom received DCS + ERP, while seven received placebo. Midtreatment, the active group had significantly lower Y-BOCS scores (t=2.87; df=21; P=0.009) with a large effect size (Cohen’s d =1.17). At 1-month follow-up, differences were no longer apparent. In children with OCD, a negative study suggested that the differences in methodology may affect results. A meta-analysis of 13 studies, only one using pediatric OCD subjects, showed a small-to-moderate effect size for the use of DCS augmentation for exposure therapy (Cohen’s d =−0.34).

Glycine

Glycine is biosynthesized from serine in a reversible folate-dependent reaction to form a nonessential amino acid (NH2CH2COOH), which can act as a neurotransmitter. In the nervous system, glycine is inhibitory in the spinal cord, playing a role in inhibitory postsynaptic potentials, but excitatory in the brain and cerebellum, where it is a glutamate coagonist at the NMDA receptors. Originally studied as a neurotransmitter for pain signaling in the spinal cord, it has gained importance in CNS disorders due to its NMDA coagonist properties and its possible role in determining the number and positioning of cortical interneurons. For example, activation of the NMDAR D-serine/glycine site or inhibition of glycine transport has been pharmacologically induced to benefit animal models and clinical trials in schizophrenia. In a placebo-controlled double-blind study of OCD, 24 subjects were assigned glycine 60 g/day or placebo. Glycine noncompliance, partly due a lack of palatability of the compound, led to ten dropouts. Follow-ups at 4-weeks, 8-weeks, and 12-weeks showed a marginally greater decrease of Y-BOCS OCD severity (P=0.053) in the glycine group (6.04 mean decrease; number [n]=5) compared to the placebo group (1.0 mean decrease; n=9). There were two out of five responders in the glycine group and zero out of nine responders in the placebo group (P=0.11).

Ketamine

Ketamine (2-[2-chlorophenyl]-2-[methylamino] cyclohexanone) is a noncompetitive, nonselective, high-affinity NMDA receptor antagonist with psychotomimetic properties. Related to the more potent phencyclidine, its profile additionally includes a short half-life and weak binding to the mu opiate receptor, monoamine transporter sites, and acetylcholinesterase receptors. Based on preclinical research and the emerging glutamate hypothesis of mental illness, ketamine was used at 0.5 mg/kg to treat refractory depression with positive results. More recent overviews of this use of ketamine overall confirm its utility, but unanswered questions still remain about the precise mechanism of action and long-term benefits. The treatment of OCD with ketamine is in the initial stages of validation. An isolated case report used two intravenous (IV) ketamine infusions (0.5 mg/kg) 1 week apart on a 24-year-old female with OCD. No symptom reduction was observed after placebo; on ketamine infusion, almost
complete reduction of obsessions was observed up to 7 days postinjection. In an open-label trial of IV ketamine 0.5 mg/kg in ten patients with refractory OCD, the initial symptom reduction of OCD symptoms was observed, along with decreases in the Hamilton Depression Rating Scale-17, a clinician-administered Dissociative States Scale, and the Clinical Global Impression (CGI) scale. Though the decrease in symptoms was maintained at 50% in four out of seven depressed patients, none of the OCD patients (n=10) sustained significant improvements. In a recent randomized, double-blind, placebo crossover study, adults with OCD (n=15) received two IV ketamine infusion treatments. The comparison groups were saline (n=7) and low-dose ketamine (0.5 mg/kg) (n=8). Those in the full-dose experimental group reported substantial improvements in OCD symptom severity, with 50% meeting the criteria for treatment response (≥35% Y-BOCS reduction) using the OCD visual analog scale and the Y-BOCS. The residual effects of ketamine lasted longer than the expected 1-week postinfusion.

**Lamotrigine**

Lamotrigine (6-[2,3-dichlorophenyl]-1,2,4-triazine-3,5-diamine) is an anticonvulsant, antiepileptic drug with voltage-gated Na-channel blocking properties. In addition, lamotrigine also facilitates GABA release and reduces the presynaptic release of glutamate. Its anticonvulsant and mood-stabilizing properties make it a treatment of choice for the management of seizures and manic–depressive illness, in particular through putative neuroprotective effects. It is especially effective in the prevention of the depressive component of bipolar disorder, bipolar relapse, and unipolar depression resistant to treatment. One double-blind trial has also shown preliminary efficacy in the treatment of the re-experiencing and numbing symptoms of post-traumatic stress disorder.

The glutamate-modulating properties of lamotrigine, based on the property of inhibiting Na+ channel-opening glutamate release, suggests that it may have some utility in the treatment of OCD. In a single case study of a 59-year-old woman with treatment-resistant OCD, the combination of a stable dose of clomipramine (225 mg/day) and lamotrigine (up to 150 mg/day) yielded a significant reduction in OCD Y-BOCS severity scores after 10 weeks. A subsequent 16-week, double-blind, placebo-controlled study of lamotrigine in 33 SSRI-resistant OCD patients showed that lamotrigine, added to standard SSRI treatment, substantially improved obsession and compulsion severity (P<0.0001).

**Memantine**

Memantine (3,5-dimethyladaman-1-amine) is a noncompetitive NMDA receptor antagonist with neuroprotective properties in cortical neuron cultures and in humans. These properties propelled its use in Parkinson’s disease, Huntington’s disease, and Alzheimer’s disease. The same glutamate-modulating property has motivated the assessment of memantine in OCD.

A reported case study of a 15-year-old female with severe OCD used memantine 5 mg twice daily (bid) in addition to preexisting citalopram. Obsessions and rituals were soon controlled for the first time in 9 months. An additional case study yielded similar results in a 29-year-old male, wherein 3 weeks of 15 mg/day of memantine produced over a 40% reduction in the Y-BOCS severity score, although another patient did not benefit. Next, an open-label trial enrolled 15 SRI-resistant OCD patients for a 12-week augmentation using memantine 10 mg bid. Data from 14 of the 15 subjects yielded significantly lower Y-BOCS scores compared to baseline (P<0.05) and 42.9% of the participants were classified as responders (≥25% reduction in Y-BOCS scores and a CGI rating of “much” or “very much” improved).

Another open-label trial in ten OCD and seven generalized anxiety disorder subjects also used memantine 10 mg bid as a standalone or add-on psychopharmacological treatment. Patients with OCD experienced a 41% reduction of Y-BOCS severity scores compared to baseline (P<0.001), with three of ten subjects classified as responders. Finally, an 8-week randomized, double-blind, placebo-controlled study of memantine 10 mg bid was conducted on 42 patients with moderate-to-severe OCD. Of 38 completers, memantine was titrated to 10 mg bid. By 8 weeks, 100% of those on memantine and 32% of those on placebo achieved partial or complete response (P<0.001).

**N-acetylcysteine**

N-acetylcysteine (NAC), a derivative of cysteine, has been used as a mucolytic in bronchitis and as an anti-inflammatory antioxidant. Administration of NAC within 16 hours postingestion is a primary treatment of acute acetaminophen toxicity; it can also be useful in the management of toxicity when hepatic failure has been established. Based on its ability to replenish the endogenous active antioxidant glutathione and direct scavenging of reactive oxygen species, NAC has been targeted to decrease doxorubicin cardiotoxicity in rat animal models and the oxidative damage in acute respiratory distress syndrome. Its thiol properties have supported its use in the protection of chemotherapy-induced cytotoxicity, if...
administered shortly after cisplatin, and its microcirculatory effects and inhibition of neutrophil aggregation in ischemia–reperfusion cardiac injury for coronary artery bypass graft surgery. Finally, even dietary NAC can replenish the antioxidant glycine pool, with a resultant increase in cysteine, glutamine, and oxidized glutathione.

In a single study that evaluated the total antioxidant and antioxidant status in serum, there was increased oxidant status and lower antioxidant status, with a resulting high oxidative stress index in children with OCD. Another study in adults with OCD measured erythrocyte malondialdehyde, a product of lipid peroxidation, and found that OCD was associated with higher levels of malondialdehyde, suggesting that a higher oxidation level was present. In this study, the antioxidant vitamin E, but not vitamin C, was also significantly lower in patients with OCD. Clinically, only a single case study of a 58-year-old woman with SRI-refractory OCD, in whom 3 g/day of NAC augmented fluvoxamine, reported a decrease of her Y-BOCS score from 32 (severe OCD) to 9 (no OCD) over 12 weeks.

Riluzole (2-amino-6-trifluoromethoxy benzothiazole) is a negative regulator of CNS glutamate activity. Animal models suggest that riluzole is not specific to glutamate – it also decreases the release of acetylcholine and dopamine-trough mechanisms independent of NMDA or AMPA receptors. Other cell experiments support that in cortical neurons, riluzole inhibits voltage-dependent Na+ channels, resulting in nonglutamate-mediated anticonvulsant properties. The neuroprotective properties of riluzole made it an attractive option for the treatment of amyotrophic lateral sclerosis (ALS), a neurodegenerative disease of cortical, brainstem, and spinal cord motor neurons. Based on successful double-blind trials, in 1996, riluzole was approved for the treatment of ALS. Clinical experience with riluzole in ALS suggests that it prolongs survival, along with an adequate long-term use safety profile.

Riluzole has been used in open and randomized clinical trials for the management of OCD symptoms. An early open-label study on 13 treatment-resistant patients with OCD ages 18–65 years used riluzole 50 mg bid to augment therapy. Of the 13 patients, seven out of 13 (54%) reported a >35% reduction in the Y-BOCS severity score. Of these, five out of 13 (38%) were treatment responders. Depression and anxiety measures also improved, and no concerning side effects were reported. Another open-label study of six children with OCD demonstrated that four out of six (67%) had a mean 40% decrease in OCD symptom severity. However, a recent double-blind randomized trial on 60 children with OCD, using riluzole 100 mg/day for augmentation, did not demonstrate an advantage over placebo.

Topiramate

Topiramate (C12H21NO8S) is an anticonvulsant agent with linear kinetics and low binding to plasma proteins. Its glutamate-modulating properties are mediated through AMPA receptor antagonism, in addition to the blockage of voltage-dependent Na+ channel and the potentiation of GABA-A neurotransmission. It also inhibits AMPA and kainate receptor facilitation of Ca2+ influx, thus providing a potential buffer to glutamatergic excitotoxicity. Topiramate’s anorectic properties have motivated its use for weight control in combination with phentermine, and its pain control properties support topiramate’s use in the prophylaxis of chronic migraine.

A case report of a 45-year-old paroxetine-resistant patient with OCD highlighted the use of topiramate 150 mg/day. After 9 weeks, the Y-BOCS OCD severity score decreased significantly. A more extended review of 16 consecutive adult patients with OCD, who had only partial response to SRI with or without neuroleptic augmentation, were reviewed after topiramate was added. Upon topiramate addition at 250 mg/day over an average of 9 weeks, there was a positive response in eleven out of 16 (69%) patients. A 12-week, double-blind, placebo-controlled trial of topiramate was conducted in 36 patients with OCD. Eighteen patients received a 12-week SSRI trial in addition to topiramate (mean dose: 180 mg); the other 18 patients received placebo in addition to the SSRI. Only the compulsions subscale improved (P=0.01), not the obsessions or total Y-BOCS scales. However, many patients had discontinued treatment due to side effects in the active arm (5/18; 28%). One case study of OCD emerging after topiramate treatment remains as an isolated report.

Pregabalin

Pregabalin, (S)-(+)-3-aminomethyl-5-methylhexanoic, was designed as a blood–brain barrier-permeable GABA analog related to gabapentin. Originally designed as an anticonvulsant, it has been found to be effective in the treatment of hyperalgesia and pathologic anxiety. In hippocampal neurons, pregabalin decreases excytosis, which opens neurotransmitter vesicles from presynaptic sites, a process that appears to depend on NMDA activation. Based on its anxiolytic properties, pregabalin was used in an 8-week open-label trial in ten OCD SRI–neuroleptic-resistant patients.
Adjunctive pregabalin was administered at 225–675 mg/day, resulting in a significant reduction of OCD Y-BOCS severity scores from 27.1 (standard deviation [SD] = 6.5) to 12.3 (SD = 7.1). Overall, a 35% symptom improvement was seen in eight of the patients’ Y-BOCS scores.119

**Tourette syndrome**

TS is a childhood onset neurobehavioral disorder characterized by >12 months of multiple motor tics and at least one vocal tic during the course of the disorder. Tics are stereotyped, suppressible, and suggestible repetitive movements of single muscle groups (simple tics) or multiple muscle groups (complex tics).120 Involuntary and detrimental to daily functioning, tics are commonly treated with behavioral therapy with an emphasis on tic suppression121 to improve psychiatric and psychosocial functioning in children122 and adults.123 Clinically, TS is often comorbid with ADHD,124 OCD,125 as well as anxiety and mood disorders. Comorbidities significantly impact the quality of life of TS patients.126

Pathophysiologically, TS has been associated with the CSTC with a key focus on synaptic neurotransmission abnormalities.127,128 One of the main proposed mechanisms in TS is the increased transmission of dopamine,129 possibly due to abnormally high dopamine receptor prefrontal density.130 An alternative view is illustrated by the putative involvement of glutamate in neurotransmission in CSTC circuits,131 extensive interaction of dopamine and glutamate systems (Table 2),132 and postmortem brain studies.133

Animal models in TS have been fraught with the innate complexity of modeling specific motor movements (tics) in nonhuman species. A working animal model will depend on the physiological assumptions about tics; for example, deficient gating has been proposed as a mechanism for propensity to tic expression based on premonitory urges associated with tics. The prepulse inhibition (PPI) of the acoustic startle reflex-deficit animal model is used to demonstrate deficient gating, and although not specific to TS, it can be used to explore pharmacologic options for tics.134 An alternate animal model relies on D1 receptor variants that drive glutamatergic neurocircuitry excitability; a mouse model displays compulsive-like repetitive behaviors.135,136 Interestingly, in these animal models, PPI is disrupted by the chemically-induced antagonism of D1 receptors in the medial prefrontal cortex (dopamine mechanism) and NMDA receptors in the ventral hippocampus (glutamate mechanism).137 These models suggest complex interactions in the cortical–subcortical neurocircuitry that involve dopamine and glutamate influences, along with a decrease in PPI.

**Clinical trials of glutamate-modulating drugs in TS**

**Ketamine**

Ketamine, a high-affinity NMDA receptor antagonist, has been found to increase PPI levels, making it a plausible drug to ameliorate tics. No direct trials of ketamine in TS are available; therefore, data on the effect of ketamine on PPI are reviewed. To measure the cognitive and PPI effects of ketamine, 20 ketamine-naive subjects with negligible baseline startle responses were given either a placebo or a dose of ketamine of 0.23 mg/kg over 5 hours, followed by ketamine 0.5 mg/kg over 1 hour. A significant increase in PPI levels was detected in the active drug group.138 In another study with 16 healthy patients, researchers found that subanesthetic doses of ketamine significantly increased PPI.139 In both trials, experienced cognitive effects resembled those seen in schizophrenia, but PPI levels were increased by ketamine rather than decreased, as seen in schizophrenic patients, suggesting utility in disorders with abnormal levels of PPI.

**Lamotrigine**

The efficacy of lamotrigine in TS has been variable. A single case study reported decreased TS symptoms in a 26-year-old woman with a clinical diagnosis of bipolar II disorder, TS, migraine, and complex partial seizures.140 An initial Yale Global Tic Severity Scale (YGTSS) score of 39/50 was decreased over 6 months after a 12-week titration of lamotrigine to 300 mg/day.140 Another case report documented increased tics after lamotrigine 200 mg/day for mood stabilization in a 55-year-old female with bipolar disorder.141 Tics included shrugging her right shoulder, wagging her hips, pawing her feet on the ground, and vocal tics. After tapering lamotrigine dosage, her tic symptoms faded and disappeared in 2 weeks’ time.141 TS symptom emergence, which improved on lamotrigine withdrawal, has been reported in five children using lamotrigine for the control of seizure disorders.142,143

**Pregabalin**

Motor tics that accompany TS are self-injurious ∼4%–43% of the time.144 In one case study, a 54-year-old man with TS with self-injury presented with fibromyalgia, comorbid OCD, and self-injurious behaviors (tongue self-mutilation).145 The patient was resistant to SSRIs, clomipramine, and haloperidol. Pregabalin monotherapy 600 mg/day was administered given the history of fibromyalgia and failure of other approaches; it was successful in improving self-injurious behaviors over 52 weeks of use.145
Topiramate
A retrospective chart review of 41 patients with TS who were administered topiramate (mean dose: 150 mg/day) for an average of 9 months, suggested that this drug could be effective for tic management. In this case series, topiramate was used as the primary monotherapy in 21 of 41 patients, while concomitant medications included neuroleptics (11/41), clonidine or guanfacine (6/41), tetrabenazine (3/41), and botox (3/41). Over 75% of subjects had “moderate to marked improvement” of tics. The main adverse effects were those common to topiramate: cognitive–language problems (25%) and aggression/mood swings (10%). A single 10-week, randomized, double-blind, placebo-controlled, parallel group study compared topiramate versus placebo in 29 patients with TS. The main outcome, “total tic score”, significantly improved in the active drug group (14.3 [SD =10.5]) compared to placebo (5.0 [SD =9.9]) (P<0.05). YGTSS motor, vocal, and total tic scores were significantly improved in the active drug group compared to placebo. No difference in adverse events was noted between the two groups. To date, topiramate constitutes a second-line agent in the management of clinically impairing tics.

Attention-deficit hyperactivity disorder
ADHD is a neurobehavioral disorder characterized by clinically impairing inattention and/or hyperactivity–impulsivity, which arise during early development. ADHD is categorized into three subtypes: 1) predominantly inattentive; 2) predominantly hyperactive–impulsive; and 3) combined. However, the full characterization of the syndrome should also emphasize related core deficits: an inability to inhibit the prepotent response, working memory developmental lags, and executive dysfunction. In a 2011 US survey, up to 11% of children and adolescents 4–17 years of age were affected, and less than two-thirds of diagnosed children were taking medication.

Anatomical abnormalities in ADHD include reductions in total cerebral, prefrontal cortex, basal ganglia, dorsal anterior cingulate cortex, corpus callosum, and cerebellar volumes. Functional magnetic resonance imaging (fMRI), magnetic resonance spectroscopy (MRS), and resting-state fMRI also detect abnormalities in ADHD when compared to normotypicals. Specific to glutamate, a proton MRS ([1H]MRS) study of 40 adults with ADHD (24 medication-naïve), found that glutamate + glutamine (Glx) was lower in the basal ganglia and dorsolateral prefrontal cortex when compared to a control region (medial parietal cortex). Medication status did not alter the results, and the level of symptoms significantly correlated with Glx levels in the basal ganglia. Adults with autism spectrum disorder have also displayed decreased Glx concentrations in the basal ganglia compared to a parietal cortex control region. Other target regions in adults with ADHD, the right anterior cingulate cortex (decreased Glx) and the left cerebellar hemisphere (increased Glx), also have alterations in Glx (glutamate/glutamine) resonance. An additional study in children with ADHD found that particularly Glx – and not N-acetylaspartate, creatine/phosphocreatine (Cr), or choline – was increased in the right prefrontal cortex and left striatum. An increased Glx ratio was seen in the right prefrontal cingulate in another study of patients with ADHD aged 8–54 years compared to controls. The alterations in the Glx balance in different brain regions in ADHD are still of unknown valence. Although ADHD is known to be highly influenced by dopamine-linked neurocircuitry, extensive interactions between the glutamate and dopamine neural systems support a role for glutamate in ADHD pathophysiology (Table 3). For example, in the prefrontal cortex, D1 and D2 receptors modulate glutamate transmission at the NMDA and AMPA receptor sites. In turn, the downregulation of glutamate transmission for NMDA receptors is affected by GABAAergic interneurons via D1 receptors. Following the demonstration of increased Glx content in ADHD, the effect of medication on brain regions has been examined. Methylphenidate and atomoxetine effect a 56% reduction in Glx in the left striatum in parallel to decreasing ADHD symptoms. In an analogous subsequent study in 13 treatment-naïve children with ADHD compared to controls, both Glx and Cr were increased pretreatment in the left striatum. After methylphenidate treatment, symptoms were improved and only striatal creatine was found to be reduced. A more recent MRS study in adolescents with ADHD found increased ratios of Glutamine/Inositol, Glutamate/Inositol, and Glx/Ino in the anterior cingulate – a brain region targeted for its role in attention in relation to error detection in subjects with ADHD compared to controls. However, these differences were not statistically significant. Additionally, no significant differences were seen in these metabolite ratios pre- and post-treatment with a methylphenidate long-acting preparation.

Animal models have been instrumental in understanding the biological underpinnings of ADHD, as well as the interaction of the dopamine and glutamate systems influencing attentional and hyperactivity behavioral domains. An early model located the dopamine receptor (DR) D1 as a potential target for genetic manipulation. The DRD1 KO mouse
exhibits excessive motor activity, as well as reduced striatum dynorphin.172 Later experiments reaffirm that DRD1 plays a role in attentional performance with D1 agonists improving, and D1 antagonists worsening, attentional performance in mouse models. D2 antagonists have no effect on attentional performance in these models.173 A direct measure of the glutamate-stimulated release of dopamine in key ADHD-related brain regions used the spontaneously hypertensive rat model of ADHD (SHR). Compared to control rats, SHR showed increased glutamate-stimulated dopamine release in the substantia nigra, implicating altered glutamate regulation of dopamine neurons in this ADHD rodent model.174 A recent extension of these findings confirmed increased evoked glutamate release in the cingulate and infralimbic cortices, as well as in the striatum, using the SHR model of ADHD.175

Genetic lines of research are also pointing to the influence of glutamate in ADHD. A recent genome-wide study implicates glutamatergic synaptic adhesion molecules and mGluRs as etiologic in ADHD.176 Another genome-wide study using copy-number variants identified copy-number variants spanning GRM genes.177 DRD4 variants have also been associated with ADHD risk.159 It had been shown earlier that DRD4-deficient mice display cortical hyperexcitability, suggesting that DRD4 stimulation negatively regulates glutamate activity, thereby curbing neurotoxicity.178 In addition, manipulation of DRD4 activity in pyramidal cortical neurons resulted in reversible decreases of NMDA receptor-mediated currents, in association with inhibition of protein kinase A (PKA), activation of protein phosphatase 1, and inhibition of calmodulin-dependent protein kinase II (CAMKII).179 Conversely, in DRD4-deficient mice, there was increased expression of DRD1 and NMDA receptors in the nucleus accumbens and hippocampal CA1 tissue, suggesting a functional relationship between DRD4, DRD1, and NMDA receptors.180 These relationships carry over into the striatum, following more recent experiments with DRD4-deficient mice, which show that resting extracellular levels of glutamate are increased in the striatum, and glutamate clearance kinetics are significantly decreased in the dorsal striatum.181 PKA, an intracellular target of DRD1 stimulation, is also implicated in attentional performance. Inhibition of PKA both reduces attentional performance and increases locomotor activity. In turn, PKA activates cyclic adenosine monophosphate (cAMP) responsive element binding protein (CREB), a transcription factor that binds to DNA “responsive elements” intracellularly, increasing the transcription of c-Fos, BDNF, somatostatin, VGF, enkephalin, and corticotrophin releasing hormone, among others.182 Inhibition of PKA reduces the accuracy of a serial reaction time task in rodents, a test that parallels the continuous performance test in humans. The same effect is observed with the disruption of CREB.183 The interplay of glutamate signaling and CREB influences attentional and locomotor systems in animal models. Among these models, glutamate homeostasis in the ventromedial prefrontal cortex (vmPFC) is an important framework for exploring the symptoms of ADHD. Infusion of rat vmPFC with the NMDA antagonist 3-((R)-2-carboxyipiperazin-4-propyl-1-phosphonic acid ([R]-CPP) decreases attentional performance, while increasing impulsivity and perseverative responses. These changes are accompanied by increased excitotoxic glutamate efflux in the vmPFC and changes in CREB expression in the vmPFC (increased) and striatum (decreased). Counteraction of the glutamate imbalance produced by (R)-CPP is achieved with the mGluR2–3 agonist agent, LY379268, which restores attentional performance and impulsivity, but not perseverative responses. This model suggests that the NMDA and mGluR2–3 pathways may underlie the core components of ADHD.184 Another rodent model of ADHD is the Naples high-excitability rat, which is bred to display behavioral arousal to novelty.185 Forebrain levels of glutamate, as well as d- and l-aspartate, are elevated in Naples high-excitability rats, suggesting that these levels may be causing neurotoxicity in the fronostriatal pathways.186

Clinical trials of glutamate-modulating drugs in ADHD

Amantadine

Recent trials of amantadine in pediatric populations have included its use in autism and ADHD.41 In an early open-label trial of amantadine, eight children with ADHD 10–13 years of age were taken off stimulant medication and provided with amantadine 100 mg bid for 1 month. While two children had similar responses as they had to methylphenidate, most only had an intermediate response compared to methylphenidate.186 A 6-week, open-label trial187 using amantadine 50–150 mg on 24 stimulant-naïve children with ADHD aged 5–13 years showed superiority over placebo. Only one child could not complete the trial due to headaches, and another had a transient decrease in appetite. Up to 60% of children had a >25% response in parent ADHD ratings, and 50% by teacher ratings.187 A head-to-head 6-week comparison trial188 of amantadine (100–150 mg/day) versus methylphenidate (20–30 mg/day) among 28 boys and 12 girls with ADHD aged 6–14 years showed no difference between the two drugs. Parent ratings showed 55% were responders on methylphenidate and 50% on amantadine; the teacher ratings were 35% responders for methylphenidate and 30% for amantadine.188
Table 1: Glutamate-modulating pharmacotherapy of obsessive–compulsive disorder

<table>
<thead>
<tr>
<th>Medication</th>
<th>Author</th>
<th>Year</th>
<th>Duration</th>
<th>N</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amantadine</td>
<td>Pasquini 16</td>
<td>2010</td>
<td>NA</td>
<td>1</td>
<td>Improvement on 200 mg/day augmentation for clomipramine 225 mg/day</td>
</tr>
<tr>
<td></td>
<td>Stryjer 44</td>
<td>2014</td>
<td>6 weeks</td>
<td>8</td>
<td>Improvement on 200 mg augmentation for stable SSRI</td>
</tr>
<tr>
<td>D-cycloserine</td>
<td>Kushner 26</td>
<td>2007</td>
<td>12 weeks</td>
<td>25</td>
<td>Improvement on 125 mg/day + ERP sessions</td>
</tr>
<tr>
<td></td>
<td>Wilhelm 33</td>
<td>2008</td>
<td>5 weeks</td>
<td>23</td>
<td>Improvement on 100 mg/day + behavior therapy</td>
</tr>
<tr>
<td></td>
<td>Storch 31</td>
<td>2007</td>
<td>12 weeks</td>
<td>24</td>
<td>No improvements on 250 mg/day</td>
</tr>
<tr>
<td>Glycine</td>
<td>Greenberg 46</td>
<td>2009</td>
<td>12 weeks</td>
<td>24</td>
<td>Improvement on 60 g/day</td>
</tr>
<tr>
<td>Ketamine</td>
<td>Rodriguez 60</td>
<td>2011</td>
<td>2 weeks</td>
<td>1</td>
<td>Improvement on IV ketamine 0.5 mg/kg</td>
</tr>
<tr>
<td></td>
<td>Bloch 31</td>
<td>2012</td>
<td>1–3 days</td>
<td>10</td>
<td>Initial improvement on IV ketamine 0.5 mg/kg</td>
</tr>
<tr>
<td></td>
<td>Rodriguez 62</td>
<td>2013</td>
<td>Twice</td>
<td>15</td>
<td>Improvement on IV ketamine 0.5 mg/kg</td>
</tr>
<tr>
<td>Lamotrigine</td>
<td>Uzun 12</td>
<td>2010</td>
<td>10 weeks</td>
<td>1</td>
<td>Improvement on 150 mg/day + clomipramine 225 mg/day</td>
</tr>
<tr>
<td></td>
<td>Bruno 71</td>
<td>2012</td>
<td>16 weeks</td>
<td>33</td>
<td>Improvement on 100 mg/day + standard SSRI</td>
</tr>
<tr>
<td>Memantine</td>
<td>Hezel 79</td>
<td>2009</td>
<td>4 weeks</td>
<td>1</td>
<td>Improvement on 5 mg bid + citalopram</td>
</tr>
<tr>
<td></td>
<td>Pasquin 85</td>
<td>2006</td>
<td>3 weeks</td>
<td>1</td>
<td>Improvement on 15 mg bid</td>
</tr>
<tr>
<td></td>
<td>Aboujaoude 83</td>
<td>2009</td>
<td>12 weeks</td>
<td>15</td>
<td>Improvement on 10 mg bid</td>
</tr>
<tr>
<td></td>
<td>Feusner 82</td>
<td>2009</td>
<td>12 weeks</td>
<td>10</td>
<td>Improvement on 10 mg bid</td>
</tr>
<tr>
<td></td>
<td>Ghaleiha 83</td>
<td>2013</td>
<td>8 weeks</td>
<td>42</td>
<td>Improvement on 10 mg bid</td>
</tr>
<tr>
<td>NAC</td>
<td>Lafleur 97</td>
<td>2006</td>
<td>12 weeks</td>
<td>1</td>
<td>Improvement 3 g/day + fluvoxamine</td>
</tr>
<tr>
<td>Pregabalin</td>
<td>Oulis 19</td>
<td>2011</td>
<td>8 weeks</td>
<td>10</td>
<td>Improvement on 225–675 mg/day augmentation</td>
</tr>
<tr>
<td>Riluzole</td>
<td>Coric 35</td>
<td>2005</td>
<td>6–12 weeks</td>
<td>13</td>
<td>Improvement on 50 mg bid augmentation</td>
</tr>
<tr>
<td></td>
<td>Grant 76</td>
<td>2007</td>
<td>12 weeks</td>
<td>6</td>
<td>Improvement on 120 mg bid</td>
</tr>
<tr>
<td></td>
<td>Grant 77</td>
<td>2014</td>
<td>12 weeks</td>
<td>60</td>
<td>Limited improvement on 100 mg/day augmentation</td>
</tr>
<tr>
<td>Topiramate</td>
<td>Hollander 113</td>
<td>2006</td>
<td>9 weeks</td>
<td>1</td>
<td>Improvement on 150 mg/day augmentation</td>
</tr>
<tr>
<td></td>
<td>Van Ameringen 114</td>
<td>2006</td>
<td>9 weeks</td>
<td>16</td>
<td>Improvement on 250 mg/day augmentation</td>
</tr>
<tr>
<td></td>
<td>Berlin 115</td>
<td>2011</td>
<td>12 weeks</td>
<td>36</td>
<td>Limited improvement on 180 mg/day augmentation to SSRI</td>
</tr>
</tbody>
</table>

Abbreviations: N, number of patients; NA, not applicable; SSRI, selective serotonin reuptake inhibitor; ERP, exposure–response prevention; IV, intravenous; bid, twice per day; NAC, N-acetylcysteine.

From these trials, amantadine has a modest effect in treating ADHD. Given its primary glutamate-modulating action, amantadine may be useful in treating symptoms of ADHD in specific subgroups of affected children.

Lamotrigine

A retrospective case series of 40 adult patients aged 16–55 years with ADHD comorbid with bipolar II or recurrent depression was conducted to gauge the effect on ADHD symptoms. Most patients (38/40) were on methylphenidate as well. With lamotrigine doses ranging from 25–250 mg, CGI scores were improved over the 48-month follow-up span. Antiepileptics can worsen attentional performance, and surveys can be conducted to assess these detrimental effects. In one survey on newly diagnosed childhood absence epilepsy, valproate, but not lamotrigine or ethosuximide, worsened attentional performance. Overall, lamotrigine improved cognition and attention in 24 of 45 children receiving lamotrigine for seizure control in one study, and the improvement of ADHD symptoms with lamotrigine does not appear to be dependent on ameliorating seizure activity. Given its benign to favorable cognitive profile, clinicians have advocated using lamotrigine as the antiepileptic of choice when seizures are comorbid with ADHD. Currently, only indirect evidence supports the use of lamotrigine for the management of ADHD symptoms in children.

Table 2: Glutamate-modulating treatments for Tourette syndrome

<table>
<thead>
<tr>
<th>Medication</th>
<th>Author</th>
<th>Year</th>
<th>Duration</th>
<th>N</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lamotrigine</td>
<td>Ortiz 140</td>
<td>2012</td>
<td>12 weeks</td>
<td>1</td>
<td>Improvement on 300 mg/day</td>
</tr>
<tr>
<td></td>
<td>Seemüller 141</td>
<td>2006</td>
<td>3 months</td>
<td>1</td>
<td>Tics on 200 mg/day</td>
</tr>
<tr>
<td></td>
<td>Sotero de Menezes 143</td>
<td>2000</td>
<td>10 months</td>
<td>8</td>
<td>Tics on 4–17 mg/kg/day</td>
</tr>
<tr>
<td>Pregabalin</td>
<td>Fornaro 45</td>
<td>2012</td>
<td>52 weeks</td>
<td>1</td>
<td>Improvement on 600 mg/day</td>
</tr>
<tr>
<td>Topiramate</td>
<td>Kuo 46</td>
<td>2010</td>
<td>9 months</td>
<td>41</td>
<td>Improvement on 150 mg/day monotherapy and augmentation</td>
</tr>
<tr>
<td></td>
<td>Jankovic 147</td>
<td>2010</td>
<td>10 weeks</td>
<td>29</td>
<td>Improvement on 118 mg/day</td>
</tr>
</tbody>
</table>

Abbreviation: N, number of patients.
Table 3 Glutamate-modulating pharmacotherapy for attention deficit hyperactivity disorder

<table>
<thead>
<tr>
<th>Medication</th>
<th>Author</th>
<th>Year</th>
<th>Duration</th>
<th>N</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>Amantadine</td>
<td>Öncü²⁸</td>
<td>2014</td>
<td>48 months</td>
<td>40</td>
<td>Improvement on 25–250 mg/day</td>
</tr>
<tr>
<td>Lamotrigine</td>
<td>Finding²⁷</td>
<td>2007</td>
<td>8 weeks</td>
<td>16</td>
<td>Improvement on 10–20 mg/day</td>
</tr>
<tr>
<td>Memantine</td>
<td>Surman²⁴</td>
<td>2013</td>
<td>12 weeks</td>
<td>28</td>
<td>Improvement on 10 mg/day</td>
</tr>
<tr>
<td>NAC</td>
<td>García²⁶</td>
<td>2013</td>
<td>3 months</td>
<td>49</td>
<td>Improvement on 2.4–4.8 g/day</td>
</tr>
</tbody>
</table>

Abbreviations: N, number of patients; NAC, N-acetylcysteine.

Memantine

Memantine is related to amantadine, with both increasing striatal dopamine via NMDA antagonism.²⁹ Memantine 20 mg/day was compared to placebo in 16 children ages 6–12 years to test for efficacy in ADHD. In this pilot study, memantine was found to be efficacious, with no significant adverse effects.³⁰ A more recent open-label 12-week trial³¹ on 28 adults with ADHD or ADHD not otherwise specified used memantine 10 mg bid. Exclusion criteria decreased the subject pool to 28 participants. The Adult ADHD Investigator Symptom Report yielded a reduction in total symptoms (−17.5; P<0.001), inattentive symptoms (−10.6; P<0.001), and hyperactive symptoms (−6.9; P<0.01) at the 12-week assessment. A total of 44% of participants had CGI ratings of much or very much improved.³¹

N-acetylcysteine

A trial of NAC was conducted in 49 adults with systemic lupus erythematosus over 3 months.³² Using an ADHD rating scale, patients with systemic lupus erythematosus had significantly greater ADHD symptoms than controls. The ADHD symptoms significantly improved on NAC 2.4–4.8 g/day (P<0.001).³²

Table 4 Glutamate-modulating pharmacotherapy in grooming disorders and OCD-related disorders

<table>
<thead>
<tr>
<th>Medication</th>
<th>Author</th>
<th>Year</th>
<th>Duration</th>
<th>N</th>
<th>Results</th>
</tr>
</thead>
<tbody>
<tr>
<td>NAC</td>
<td>Grant²⁴</td>
<td>2009</td>
<td>12 weeks</td>
<td>50</td>
<td>Improvement on 1.2–2.4 g/day (TTM)</td>
</tr>
<tr>
<td>NAC</td>
<td>Rodrigues-Barata²⁶</td>
<td>2012</td>
<td>2–6 months</td>
<td>2</td>
<td>Improvement on 1.200 mg/day (TTM)</td>
</tr>
<tr>
<td>NAC</td>
<td>Taylor²⁰</td>
<td>2014</td>
<td>10 weeks</td>
<td>1</td>
<td>Improvement on 1.200 mg/day (TTM)</td>
</tr>
<tr>
<td>NAC</td>
<td>Bloch²⁵</td>
<td>2013</td>
<td>12 weeks</td>
<td>39</td>
<td>Limited improvement on 1.200 mg twice daily (TTM)</td>
</tr>
<tr>
<td>NAC</td>
<td>Odlaug²¹</td>
<td>2007</td>
<td>2 weeks</td>
<td>5</td>
<td>Improvement on 1.800 mg/day (ExD and TTM)</td>
</tr>
<tr>
<td>NAC</td>
<td>Miller²⁷</td>
<td>2014</td>
<td>12 weeks</td>
<td>35</td>
<td>Improvement on 450–1,200 mg/day (ExD)</td>
</tr>
<tr>
<td>NAC</td>
<td>Grant²¹¹</td>
<td>2007</td>
<td>12 weeks</td>
<td>24</td>
<td>Improvement on 25–300 mg/day (ExD)</td>
</tr>
<tr>
<td>NAC</td>
<td>Grant²⁶</td>
<td>2010</td>
<td>12 weeks</td>
<td>29</td>
<td>Limited improvement on 12.5–300 mg/day (ExD)</td>
</tr>
<tr>
<td>NAC</td>
<td>Lochner²¹²</td>
<td>2006</td>
<td>16 weeks</td>
<td>9</td>
<td>Improvement on 50–250 mg/day (TTM)</td>
</tr>
<tr>
<td>NAC</td>
<td>Shapiro²¹³</td>
<td>2002</td>
<td>8 weeks</td>
<td>3</td>
<td>Improvement on 50–150 mg/day augmentation (ExD)</td>
</tr>
<tr>
<td>NAC</td>
<td>Jaffery²¹⁴</td>
<td>2010</td>
<td>12 weeks</td>
<td>2</td>
<td>Improvement on 200 mg/day augmentation (ExD)</td>
</tr>
<tr>
<td>NAC</td>
<td>Coric²¹⁵</td>
<td>2007</td>
<td>3 months</td>
<td>1</td>
<td>Improvement on 100 mg/day augmentation for fluoxetine (ExD)</td>
</tr>
</tbody>
</table>

Abbreviations: OCD, obsessive–compulsive disorder; N, number of patients; NAC, N-acetylcysteine; TTM, trichotillomania; ExD, excoration disorder.

Trichotillomania, excoration disorder (OCD-related disorders)

TTM is an OCD-related disorder (OCD-RD) characterized by repetitive hair pulling resulting in noticeable hair loss. Individuals may develop rituals around the selection and removal of hair, and may engage in manipulations like rolling, examining, biting, or swallowing the hair after removal.¹⁴ Children with TTM engage in automatic hair pulling, while older children appear to have more focused pulling and are more aware of associated urges, with increased frequency of hair pulling.¹⁴

Dermatillomania or ExD is characterized by recurrent skin picking resulting in skin lesions. The age at onset can be in childhood and triggers include stress, boredom, or the feel or look of the skin.¹⁵ A striking observation is the high degree of comorbidity associated with skin-picking behaviors, including OCD, body dysmorphic disorder, substance use disorders, eating disorders, TTM, kleptomania, and compulsive buying.¹⁶ The prevalence of skin-picking disorder ranges from 1.4%–5.4% based on two recent community samples.¹⁷,¹⁸ TTM and ExD are classified by the Diagnostic and Statistical Manual of Mental Disorders, 5th edition, as OCD-RD. These body-focused repetitive behaviors may be preceded by a sense of increasing tension and followed by a sense of gratification.
pleasure, or relief. A continuum of conscious awareness may be associated with TTM and ExD, and the behaviors may be preceded by stress, anxiety, or boredom.151

The pathophysiology of OCD-RD has been investigated; however, much work needs to be done to further clarify their relationship to OCD. There are indications of increased familiality in TTM199 and ExD,200 suggesting a genetic diathesis. A lack of motor inhibitory control, such as in the Go–NoGo task in TTM201 and in the stop–signal task in ExD,160 point to an underlying neural inhibitory network deficit. Positron emission tomography studies show an increase in mean global metabolism in TTM compared to controls, as well as normalisation of increased orbitofrontal and anterior cingulate regions with clomipramine treatment.202 More recently, and paralleling neuropsychological inhibitory deficit findings, a white matter tract study of subjects with TTM found reduced fractional anisotropy (white matter disorganization) in the anterior cingulate, pre-supplementary motor area, and temporal cortices,161 as well as increased mean diffusivity in the frontal–striato–thalamic white matter tracts.203 In ExD, analogous white matter disorganization is also found in areas associated with inhibitory control, such as the tracts emerging from the inferior right prefrontal regions.204 These pathophysiologic changes suggest a neurocircuitry imbalance in inhibitory neural control systems (Table 4).

Animal models of OCD-RD have sought to simulate repetitive behaviors that include animal self-grooming motor sequences. Hox genes are developmental homeobox genes that direct somatic body structure,153 one of which is hoxb8, a transcription factor directing anterior limb development.205 The hoxb8 mutant mouse exhibits an abnormally high levels of grooming behaviors, causing skin lesions and hair removal.206 Another notable example of a mouse model of grooming behaviors is the SAP90/PSD95-associated protein 3 (SAPAP3) KO mouse, which exhibits heightened anxiety-like behaviors and excessive self-grooming behaviors leading to facial hair loss and skin lesions.33 SAPAPs are SAP90/PSD95-associated proteins that act as synaptic linker proteins between glutamate receptor-binding proteins and the cytoskeleton, while SAPAP3 is particularly highly expressed in glutamatergic synapses of the striatum.207 In humans, SAPAP3 is expressed in dendrites, and SAPAP3 genetic variants have been associated with TTM and ExD.208,209

Clinical trials of glutamate-modulating drugs in TTM and ExD

N-acetylcysteine

A 12-week, double-blind, placebo-controlled trial of NAC 1.2–2.4 g/day in 50 adults with TTM showed a greater reduction in hair-pulling behaviors in the active drug arm.163 Up to 56% of patients taking NAC were much or very much improved, while only 16% of subjects receiving placebo had the same benefit. NAC produced no adverse events in the treated group.163 Following the randomized trial, additional case reports of NAC in TTM have been reported. A case report of two females aged 23 years and 19 years showed benefit of NAC 1,200 mg/day, with complete hair regrowth at 3 months and 2 months, respectively, after long courses of illness.156 Another case report of a 58-year-old female with TTM used NAC 1,200 mg/day, with a gradual improvement of hair loss observed over a 10-week period.150 In contrast to positive trials in adults, a double-blind placebo randomized trial of NAC in TTM, which included 39 children ages 8–17 years, showed no difference between NAC (25% improved) when compared to placebo (21% improved).155

The use of NAC in ExD is highlighted by a case report of five patients with TTM or ExD who were given NAC for symptom management. One of the reported patients had ExD, which responded to NAC 1,800 mg/day after partial response to 600 mg/day, and then 1,200 mg/day.210 A more recent report highlighted the use of NAC in 35 patients with Prader–Willi syndrome (PWS) and skin-picking behaviors. Patients diagnosed with PWS were aged 5–39 years (23 females/12 males) and treatment consisted of NAC at 450–1,200 mg/day for 12 weeks.175 At treatment conclusion, all patients had reduced (n=10) or absent (n=25) skin-picking behaviors.175

Lamotrigine

An open-label trial of lamotrigine 25–300 mg/day for the management of skin-picking behaviors showed that 67% of subjects were much or very much improved, prompting a randomized controlled trial.211 A double-blind, placebo-controlled, randomized trial of lamotrigine for ExD showed a nonsignificant benefit of active drug (44% responders) compared to placebo (31% responders). Only those patients with difficulties in a test of shifting (cognitive flexibility) benefited maximally from the use of lamotrigine.162

Topiramate

An open-label, 16-week trial of topiramate 50–250 mg/day in 14 adults with TTM resulted in nine adults completing the study.212 Trial completers had a significant reduction of hair-pulling behaviors, with 6/9 classified as responders. Five patients dropped out due to adverse effects.212 In a model of ExD using patients with PWS, topiramate 50–150 mg/day was administered as an add-on for the management of self-injury. All three patients experienced a significant resolution.
of ExD within 8 weeks of initiating the topiramate trial.\textsuperscript{213} Another case report\textsuperscript{214} noted that two adolescents, aged 15 years and 16 years, with autism spectrum disorder and significant ExD symptoms benefited from topiramate addition to their psychotropic regimen in doses up to 200 mg/day for 12 weeks.

## Riluzole

A 52-year-old female with OCD, anorexia, and major depression, who also exhibited ExD, was treated with fluoxetine, to which riluzole 100 mg bid was added. Scores on OCD rating, eating disordered behaviors, and pathologic skin picking all improved on this regimen. It was noted that urges and compulsive behaviors had diminished notably with the addition of riluzole.\textsuperscript{215}

## Conclusion

CNS homeostasis of inhibition–disinhibition signaling ultimately depends on a well-regulated glutamate–GABA balance, in conjunction with other neurotransmitter systems that impact on this final effector pathway, which highly impacts neuronal health.\textsuperscript{216} Future drug design approaches will benefit from a better understanding of these pathways (which also impact on other biologic systems, including immune and developmental networks) in OCD and related disorders, tics and ADHD, in order to provide a paradigmatic framework to better understand the imbalance in inhibition–disinhibition from the molecular level (glutamate–GABA) to the macrobehavioral level (obsessions, compulsions, tics, hyperactivity, and grooming behaviors). The heuristic value of considering these cross-disorder clinical manifestations in toto in relation to glutamate awaits future drug discovery to address these disinhibitory phenomena. In summary, drugs that impact the glutamatergic balance in the CNS are emerging as a therapeutic alternative for neuropsychiatric disorders, which implicate abnormal inhibitory control in cognitive, motor, behavioral, and grooming domains. OCD has the most support at present for the use of glutamate modulators, with ADHD and grooming disorders also showing promise. TS appears to be less affected by glutamate modulators, but the research is quite sparse in this condition. Future research into the pathophysiology of glutamate actions, and resultant drug discovery, in the CNS should include its role in early development, neurogenesis, and trophic mechanisms.\textsuperscript{219}

## Disclosure

The authors report no conflicts of interest in this work.


\textbf{References}


