EAAT2 as a Research Target in Bipolar Disorder and Unipolar Depression: A Systematic Review

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Abstract
Glutamate is implicated in the neuropathology of both major depressive disorder and bipolar disorder. Excitatory amino acid transporter 2 (EAAT2) is the major glutamate transporter in the mammalian brain, removing glutamate from the synaptic cleft and transporting it into glia for recycling. It is thereby the principal regulator of extracellular glutamate levels and prevents neuronal excitotoxicity. EAAT2 is a promising target for elucidating the mechanisms by which the glutamate-glutamine cycle interacts with neuronal systems in mood disorders. Forty EAAT2 studies (published January 1992–January 2018) were identified via a systematic literature search. The studies demonstrated that chronic stress/stereoids were most commonly associated with decreased EAAT2. In rodents, EAAT2 inhibition worsened depressive behaviors. Human EAAT2 expression usually decreased in depression, with some regional brain differences. Fewer data have been collected regarding the roles and regulation of EAAT2 in bipolar disorder. Future directions for research include correlating EAAT2 and glutamate levels in vivo, elucidating genetic variability and epigenetic regulation, clarifying intracellular protein and pharmacologic interactions, and examining EAAT2 in different bipolar mood states. As part of a macromolecular complex within glia, EAAT2 may contribute significantly to intracellular signaling, energy regulation, and cellular homeostasis. An enhanced understanding of this system is needed.

Introduction
Mood disorders cause a significant burden of disease worldwide. Globally, depression has a prevalence of 4.4\% [1], affecting approximately 322 million people, with significant loss of quality-adjusted life years and productivity, and increased risk of suicide. Bipolar disorder (BD) has an estimated prevalence of up to 4.4\% in the US population [2] and is associated with elevated suicide risk and significant morbidity. Depression and BD are clinically heterogeneous mood disorders which, in mania or severe depression, can present with psychotic symptoms. As the major excitatory neurotransmitter of the mammalian brain, glutamate has been associated with both mood and...
psychotic disorders [3, 4]. Released by presynaptic neurons, glutamate is transported into surrounding glial cells and recycled back to glutamine. Glutamine is the precursor molecule to both glutamate and γ-aminobutyric acid (GABA), the major inhibitory neurotransmitter in the central nervous system. This glutamate-glutamine cycle helps regulate neuronal excitability [5].

The main glutamate transporter in the mammalian brain is the excitatory amino acid transporter 2 (EAAT2), also called glutamate transporter 1 (Glt-1) in rodent literature, or solute carrier family 1 member 2 (SLC1A2). It is encoded by the SLC1A2 gene and inhibited by dihydrokainate (DHK) [6]. EAAT2 plays a crucial role in preventing extracellular glutamate concentrations from reaching neurotoxic levels [7] and in recycling glutamate at synapses by transporting glutamate into astrocytes for conversion to glutamine (Fig. 1). This figure also illustrates an overview of the multiple dynamic interactions between glutamate receptors and transporter subtypes within the glutamate cycle. There are five mammalian EAAT isoforms (EAAT1–5) of the solute carrier 1 (SLC1) glutamate transporter family [8]. EAAT isoforms have different tissue localizations, amino acid sequences, and pharmacological profiles. Only EAAT1 (also known as glutamate-aspartate transporter, GLAST) and EAAT2 are expressed in astrocytes. EAAT2 is additionally expressed on axon terminals [9]. EAAT3 is fairly ubiquitously expressed in neuronal tissue, with high intracellular and postsynaptic terminal concentrations, possibly for rapid buffering of local glutamate changes [10].

Glutamate transporters share 25–30% homology [11], and the structure of EAAT2 is partially extrapolated from homologs. EAAT2 is a bowl-shaped homotrimer with a large central water-filled cavity extending into the cell membrane [12]. Each monomer contains glutamate-binding sites and transport pathways and can probably bind and transport glutamate independently of the other monomers [13]. Each monomer core contains substrate- and ion-binding sites and is comprised of two hairpin loops (HP1, HP2) which may comprise the internal and external gates of the transporter and multiple transmembrane (TM1–8)
segments. Glutamate and kainate induce distinct conformational changes in the external gate of EAAT2 [14]. EAAT2 is believed to co-transport glutamate with three sodium ions and one hydrogen ion down concentration gradients into the cell, with sequential counter-transport of one potassium ion; each monomer also contains a chloride channel, possibly a feedback system reducing glutamate release [15]. EAAT2 is part of a macromolecular complex that includes aquaporin-4 (AQP4) [16], a water channel that plays a key role in ion and fluid homeostasis within the brain [17]. This complex further includes other membrane proteins, mitochondria, and glycolytic enzymes, suggesting complex regulatory and homeostatic functions [18]. AQP4 may buffer astrocytic water volume affected by EAAT2 activation [19]. AQP4 is of clinical interest in a wide variety of neurological conditions, including neuromyelitis optica, epilepsy, Alzheimer’s dementia, traumatic brain injury, and Parkinson’s disease.

EAAT2/Glt-1 displays transcript variants [20], and Glt-1a and Glt-1b are the most commonly described. The two splice variants have different cellular and subcellular expression profiles [21]. EAAT2 is differentially expressed across anatomical regions in different psychiatric conditions. For example, rodent depression models demonstrate increased hippocampal Glt-1a mRNA/protein, but schizophrenic humans have decreased parahippocampal and dorsolateral prefrontal cortex EAAT2 [7]. Pharmacologic studies also find associations between EAAT2 and neuropsychiatric disorders: riluzole, ceftriaxone, and steroids are medications that alter EAAT2 expression and may have antidepressant effects in rodents [22, 23]. Riluzole is a sodium channel blocker that upregulates EAAT2 protein levels, reduces synaptic glutamate release, and inhibits ionotropic glutamate receptors [22]. Ceftriaxone is a β-lactam antibiotic which upregulates EAAT2 levels and glial glutamate uptake, and has been investigated for the prevention of alcohol withdrawal [24]. Glucocorticoids cause immediate hippocampal Glt-1 mRNA/protein increases in rodent stress models [25, 26], followed by gradual decreases after chronic exposure [22].

The role of EAAT2 in selected disease processes has been summarized previously [8]. This review summarizes the current state of knowledge of the roles of EAAT2 in unipolar and bipolar mood disorders. For precision, we refer to Glt-1 in rodent studies, and EAAT2 in human studies.

**Literature Search**

We conducted a comprehensive literature search of PubMed, PsycInfo, and Web of Science from 1 January 1992 [27] through 15 January 2018. Database-specific
truncations ensured that searches included singulars/plurals and alternate suffixes or spellings. Boolean search string was: Title, Abstract, Keywords/MeSH terms (EAAT2 OR SLC1A2 OR Glt-1 OR “excitatory amino acid transporter” OR “solute carrier family 1 member 2” OR “glutamate transporter 1”) AND Title, Abstract, Keywords/MeSH terms: (bipolar OR depress* OR unipolar).

Articles were collated in EndNote X8 (Clarivate Analytics, Philadelphia, PA, USA). Duplicates were removed by the software via author, year, title, and journal comparison, and any remaining duplicates were removed manually by first author. Each abstract was reviewed independently by first and final authors. Sixty-seven articles were selected for full text review (Fig. 2).

Literature was eligible for inclusion: (1) if it discussed the roles, functions, interactions, or regulation of EAAT2 in human patients with BD or unipolar depression, or mammalian models (behavioral, genetic, or pharmacological) of BD or unipolar depression, or models of molecular interactions in BD or unipolar depression; (2) if the full text was available in English; and (3) if it was published in print or electronically between January 1992 and January 2018. Exclusion criteria were: (1) studies inducing mood symptoms through brain tissue damage (ischemia, surgical resection, etc.); (2) genetic models of non-mood disorder diseases; and (3) in silico models of molecular interactions. Forty articles met all the inclusion criteria. Twenty-six described animal models of depressive or bipolar symptoms [17, 22, 25, 26, 28–47]. Sixteen described human studies of major depressive disorder (MDD) and/or BD [48–63]. The papers are summarized in Tables 1 and 2. Each experimental paper was reviewed for design, sample size, description and duration of experiment, types of controls, how results were obtained and analyzed, and experimental findings. Data were extracted independently and in duplicate. Studies were appraised for quality using a modification of a published review process [64]; all papers met a quality score of moderate or high.

**Findings**

**Glt-1 Expression in Animal Models of Mood Disorders**

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EAAT2 in Bipolar Disorder and Unipolar Depression

Rodent Stress Models of Depression

Studies of Glt-1 in rodent mood models have produced conflicting results. Earlier studies reported that stress upregulated hippocampal Glt-1 expression [25, 26, 37], possibly as a neuroprotective response to stress-induced synaptic glutamate increases [39]. Later studies demonstrated that chronic stress downregulated Glt-1 in both rats [29, 32, 35, 36] and mice [37, 42].

In rats, chronic unpredictable stress decreased hippocampal Glt-1 protein without changing EAAT1/GLAST [45]. Chronic stress also significantly decreased hippocampal EAAT2/Glt-1 and EAAT3 protein levels (encoded by SLC1A2 and SLC1A1, respectively), in addition to associated depressive behaviors [46]. Stress may decrease Glt-1 partially via phosphodiesterases, since the phosphodiesterase 4D inhibitor (GEBR-7b) prevented stress-induced hippocampal Glt-1 protein decreases in rats [36].

Sex-specific effects in Glt-1 expression have also been reported. In stressed rats, compared to controls, females demonstrated reduced Glt-1 protein in the PFC, while males showed reduced Glt-1 in the striatum [40]. Prenatal stress administered to pregnant rats significantly reduced hippocampal, frontal cortex, and striatum Glt-1 levels in male rat offspring and was associated with increased depressive behaviors [43, 44]. Female offspring were not assessed. Prenatal stress reduces glial cells in animals and humans, and sex differences in hippocampal glial numbers have been reported, with female offspring in a prenatal stress model demonstrating significant glial reduction, depressed behaviors, and cognitive deficits compared to male offspring [67]. Since Glt-1 is preferentially expressed in glia, it is important to consider whether Glt-1 decreases are due to changes in transcription/translation, or to a loss of glial cells expressing Glt-1. Correlating Glt-1 levels and glial numbers across species and between brain regions would help interpret these results.

Strain-specific responses were seen both within mice [37] and rats [41]. Chronic stress increased hippocampal Glt-1 levels in DBA/2J mice (a strain bred for extreme intolerance to alcohol/morphine and susceptibility to seizures) but decreased Glt-1 levels in seizure-resistant C57BL/6J mice and other strains [37]. DBA/2J astrocytes have previously demonstrated much higher rates of GABA synthesis than C57BL/6J and higher glutamate-to-GABA ratios [68]. This suggests the distinct Glt-1 stress...
Table 1. Summaries of animal studies included in the systematic analysis

| Reference | Description | Species | Tissue examined | Subject characteristics | Findings |
|-----------|-------------|---------|-----------------|-------------------------|----------|
| Reagan et al. [25], 2004 | Examine the chronic restraint stress effect on Glu-1 mRNA and protein expression in rat hippocampus, and modulating effects of the antidepressant tianeptine | Sprague-Dawley rats, CD1 hippocampus strain | 40 adult male rats: 10 CRS + saline | Stress increases Glu-1a mRNA and protein in CA3; tianeptine prevents Glu-1a mRNA increases in CA3; neither stress nor tianeptine alter Glu-1b mRNA in CA3; stress increases Glu-1b protein across hippocampus; tianeptine does not prevent Glu-1b protein increases |
| Wood et al. [26], 2004 | Measure hippocampal Glu-1 mRNA expression and neuronal remodeling in chronic restraint stress models of depression, and effect of lithium on Glu-1 expression and histological findings | Sprague-Dawley rats | 30 adult male rats: 7 CRS + lithium | Stress increased Glu-1 mRNA CA3 expression; lithium prevents this stress-induced Glu-1 increase; lithium also prevents stress-induced decrease in CA3 pyramidal apical dendritic length |
| Bechsholt-Gompf et al. [28], 2010 | Examine the effect of Glu-1 inhibition by DHK on behavioral tests of mood and cognition | Sprague-Dawley rats | Whole brain coronal sections for immunohistochemistry | DHK inhibition of Glu-1 increases intracranial self-stimulation; DHK inhibition of Glu-1 decreases spatial navigation impairments; DHK increases c-Fos in reward/mood brain regions |
| Zink et al. [47], 2010 | Glutamate transporter expression in behaviorally characterized lines of rats with congenital liability to or resistance to stress-induced helplessness | Sprague-Dawley rats; subsets inbred for congenital learned helplessness | Hippocampus, occipital cortex | Glu-1 mRNA and protein decrease in helpless CLH cohort |
| Gourley et al. [22], 2012 | Effect of riluzole on Glu-1 expression and behaviors in rodent acute stress and chronic acute corticosteroid administration | C57BL6 mice | Hippocampus | Corticosteroids reduce Glu-1 protein expression in hippocampus; riluzole restores steroid-suppressed protein Glu-1 expression; riluzole increases Glu-1 protein expression in steroid-naive mice; riluzole prevents despair behavior in forced swim, but not tail suspension |
| Mali et al. [37], 2012 | RNA expression study in hippocampus across inbred mouse strains to evaluate antidepressant effects of nor-triptiline and escitalopram on gene expression in chronic stress-induced (forced swim, maternal separation, chronic mild stress) models of depression | 4 inbred mouse strains: C57BL/6J, DBA/2J, 129S1/SvEv, FVB/NJ | 72 male and 72 female mice | SLCA12 mRNA hippocampal expression decreased in C57BL/6J, 129S1/SvEv, FVB/NJ strains; SLCA12 mRNA hippocampal expression increased in DBA/2J strain; escitalopram and nortriptyline altered SLCA12 in strain-specific manner, reversing stress-induced SLCA12 mRNA changes in each strain |
| Martirosa et al. [39], 2012 | Explore the effect of chronic corticosterone administration on EAAT2 hippocampal protein levels | Wistar rats | Hippocampus | Corticosterone increased EAAT2 hippocampal protein expression |
| Zhang et al. [43], 2013 | Examine EAAT2 mRNA expression in hippocampus, striatum, and frontal cortex of rat after prenatal stress | Sprague-Dawley rats | Hippocampus, striatum, frontal cortex | Prematernal stress significantly reduced EAAT2 mRNA expression in hippocampus, frontal cortex, and striatum of juvenile offspring, with associated depressive behavior |
| Chen et al. [29], 2014 | Evaluate the effects of chronic unpredictable stress and 10 mg/kg fluoxetine on depressive behaviors in rats and Glu-1 protein levels in rat hippocampus | Sprague-Dawley rats | Hippocampus | CUS associated with depressive behaviors, and decreased Glu-1 protein; fluoxetine reversed both |
| Cui et al. [30], 2014 | DHK and genetic inhibition of Glu-1 transport in mouse LHB, testing for LHB neuronal activity, behavioral response (with and without 20 mg/kg fluoxetine), and sleep EEG/EMG | C57BL/6J mice Flxed Glu-1 mice | Lateral habenula | Glu-1 inhibition increased excitability in LHB; Glu-1 inhibition displayed worse depressive behavior and increased susceptibility to chronic stress; Glu-1 inhibition had disinhibited REM sleep; fluoxetine improved depressive behavior on tail suspension despite Glu-1 inhibition |
| Kong et al. [17], 2014 | AQP4 wild-type vs. knockout mice administered chronic corticosterone, EAAT1 and EAAT2 hippocampal protein expression evaluated | CD1 mice | Hippocampus, Dentate gyrus | AQP4 knockout mice had exacerbated depressive behaviors; corticosterone inhibited hippocampal EAAT2 protein expression only in AQP4−/− mice; corticosterone inhibited hippocampal EAAT1 protein expression in both wild-type and knockout mice |
| Veressah et al. [42], 2014 | Mice exposed to chronic social defeat to induce depressive phenotype, subsequent evaluation of sucrose preference, social interaction, and prefrontal cortex markers including EAAT2 mRNA | C57BL6 mice | Prefrontal cortex | EAAT2 mRNA levels significantly lower in PFC of depressed mice |
| John et al. [33], 2015 | DHK blockade of Glu-1 in the amygdala assessed by ICSS (as an index of hedonism and anxious behavior) and behavior tests | Sprague-Dawley rats | Central nucleus of amygdala | Amygdala Glu-1 inhibition by DHK induces anxiety and anhedonic depressive behaviors |
Table 1 (continued)

| Reference | Description | Species | Tissue examined | Subject characteristics | Findings |
|-----------|-------------|---------|----------------|-------------------------|----------|
| Zhu et al. [45], 2015 | Examine the effect of propofol and ECS on glutamate uptake, and EAAT1/2 protein levels, in rats stressed through chronic unpredictable mild stress | Sprague-Dawley rats | Hippocampus | Male rats: Control + vehicle | CUS decreased hippocampal Glt-1 protein expression; ketamine significantly attenuated this reduction; K252α pretreatment antagonized the effects of ketamine |
| Liu et al. [35], 2016a | Examine the effect of K252α (a TrkB receptor inhibitor) on chronically stressed rats as measured behaviorally and by Glt-1 protein levels | Swiss mice | Hippocampus | Adult male rats: 8 nonstressed + vehicle 8 CUS + vehicle 8 CUS + K252α (0.05 mg/kg) | CUS decreased rat exploration and movement velocity; CUS decreased hippocampal Glt-1 protein expression; GEBR-7b significantly reversed this reduction |
| Liu et al. [36], 2016b | Examine the effect of GEBR-7b (a specific depolarizing agent), NBQX (AMPA antagonist), s-AMPA (AMPA agonist), citochrom, and pCPA (5-HT synthesis inhibitor) on depression-related behaviors and synaptic molecules in mice | albino Wistar rats | Prefrontal cortex, doral striatum, nucleus accumbens | 51 male and 72 female adult Long-Evans rats: Female nonstressed Male nonstressed | CND5 reduced Glt-1 protein levels in female PFC; CDS5 reduced Glt-1 protein levels in male striatum |
| Gasull-Camós et al. [31], 2017 | Investigate the behavioral and molecular effects in rats of: DHK (Gl-1 blocker), veratridine (nonspecific depolarizing agent), NRQX (AMPA antagonist), s-AMPA (AMPA agonist), citochrom, and pCPA (5-HT synthesis inhibitor) | Wistar rats | DHK Gl-1 blockade of: infralimbic cortex, prelimbic cortex | Male rats: Intraperitoneal injection of pCPA Intraperitoneal injection of DHK or veratridine Intraperitoneal injection of NRQX or s-AMPA or citochrom | DHK infusion to infralimbic cortex induced behavioral antidepressant response; DHK infusion to prelimbic cortex induced no behavioral response; veratridine infusion to infralimbic cortex induced rapid behavioral antidepressant response; veratridine infusion to prelimbic cortex induced no behavioral change; DHK-induced antidepressant response prevented by NRQX and pCPA, mimicked by s-AMPA and citochrom |
| Zhang et al. [44], 2017 | Effect of sciarin on behavior and glutamate system in prenatally stressed male rats | Sprague-Dawley rats | Hippocampus | Pregnant female rats: CBS vs. control 2 male offspring from each mother: 8 control 8 prenatal stress 8 PFS + saline 8 PFS + icariin 80 mg/kg 8 PFS + icariin 40 mg/kg | Prenatal stress decreased EAAT2 protein expression in male offspring; prenatal stress increased mGluR1 and mGluR5 protein expression in male offspring; prenatal stress offspring displayed decreased sucrose preference, increased immobility, chronic postnatal sciarin administration increased sucrose preference, reduced immobility time, chronic postnatal sciarin administration increased EAAT2, and decreased mgluR1/mgluR3 protein levels |
| Zhu et al. [46], 2017 | Assess the effect of subanesthetic doses of ketamine on depressed behavior and glutamate transporter hippocampal expression in a stress model of depression | Sprague-Dawley rats | Hippocampus | 60 adult male rats: 12 control 12 CUMS + saline 12 CUMS + 10 mg/kg ketamine 12 CUMS + 25 mg/kg ketamine 12 CUMS + 50 mg/kg ketamine | CUMs significantly downregulated EAAT2 hippocampal protein levels; ketamine reversed CUMs-induced deficits; EAAT2 induction was dose-dependent; CUMs decreased sucrose preference, locomotion, exploratory behavior; ketamine reversed CUMs-induced behaviors; CUMs increased extracellular hippocampal glutamate concentrations; ketamine reduced CUMs-induced hippocampal glutamate concentrations |
| Reference Description Species Tissue examined Subject characteristics Findings |
| Zhu et al. [41], 2015 | Examine chronic mild stress in high and low anxious phenotype rats; effect on DNA damage, NMADA, and Glu-1 levels in PFC, hippocampus, amygdala, and NA | Carioca High-Freezing rats Carioca Low-Freezing rats Albino Wistar rats as controls | Prefrontal cortex, hippocampus, amygdala, nucleus accumbens | 6 groups: Wistar + control conditions Wistar + stress GlF + control conditions GlF + stress CUMs + control conditions CUMs + stress | EAAT2 decreased in PFC of stressed Wistars relative to unstressed Wistars; EAAT2 increased in PFC of unstressed CUMs relative to unstressed Wistars; EAAT2 increased in hippocampus of unstressed CUMs relative to unstressed Wistars; EAAT2 decreased in hippocampus of stressed CUMs relative to unstressed CUMs; EAAT2 decreased in amygdala of stressed CUMs relative to unstressed Wistars; EAAT2 decreased in amygdala of stressed CUMs relative to unstressed Wistars; EAAT2 did not change in NA of any group relative to control |
| Gómez-Galán et al. [40], 2016 | Genetic depression model in rats subjected to social isolation to assess molecular effects, then assessed for effects of physical exercise on social isolation-induced molecular effects | FSL rats | Sprague-Dawley rats | Rats: 7 of each strain: group-housed 7 of each strain: social isolation 7 of each strain: social isolation with exercise | Social isolation decreased Glt-1 protein levels, exercise reversed the social isolation-induced Glt-1 protein decrease |
| Réus et al. [38], 2016 | Examine the effect of pCPA (5-HT synthesis inhibitor) and citalopram, and (AMPA antagonist), s-AMPA specific depolarizing agent), NBQX (AMPA antagonist), s-AMPA (AMPA agonist), citochrom, and pCPA (5-HT synthesis inhibitor) on depression-related behaviors and synaptic molecules in mice | Sprague-Dawley rats | Hippocampus | Intraperitoneal infusion of pCPA Infralimbic infusion of DHK or veratridine | CUS decreased hippocampal Glt-1 protein expression; ketamine significantly attenuated this reduction; K252α pretreatment antagonized the effects of ketamine |
| Liu et al. [32], 2016 | Examine the effect of K252α (a TrkB receptor inhibitor) on whether BDNF-TrKB signaling is part of the antidepressant effect of ketamine in a chronic unpredictable stress model of depression in rats | Sprague-Dawley rats | Hippocampus | Male rats: Control + vehicle CUS + vehicle CUS + ketamine (10 mg/kg) CUS + K252α (1 μL, 2 mM) + ketamine (10 mg/kg) | CUS decreased depression model in rats subjected to social isolation to assess molecular effects, then assessed for effects of physical exercise on social isolation-induced molecular effects |

EAAT2 in Bipolar Disorder and Unipolar Depression  
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may be related to an underlying difference in the glutamate/glutamine/GABA system, though it remains unclear whether this involves altered gene transcription, protein expression, transport activity, or GABA synthesis. Different Glt-1 responses to stress have also been seen in rats [41]. Rats bred for helpless behavioral responses to chronic stress were examined within strain, to see whether they displayed helplessness or stress resistance. Their behavior was also compared to that of control rats [47].

The stress-susceptible/behaviorally resilient strain and the control rats retained higher levels of Glt-1, while the stress-susceptible/behaviorally helpless strain had decreased Glt-1 mRNA/protein. Reduced astroglial glutamate uptake and increased glutamate levels were seen in these genetically and behaviorally helpless rats.

Pharmacological Models of Depression and Glt-1 Modulation

Chronic corticosteroid administration induced depressive behaviors in rodents [17, 22, 39]. In rats, corticosterone treatment increased hippocampal Glt-1 [39]. In mice, steroids reduced hippocampal BDNF and GLAST, glial neurotrophic factor, and neurogenesis in mice, but did not significantly change Glt-1 [17, 22]. However, in AQP4 knockout mice, 20 mg/kg corticosterone daily for 21 days significantly downregulated Glt-1 protein [17]. Differences in Glt-1 expression within these mouse strains may reflect a effect of AQP4 function. AQP4 knockouts have a reduction in astrocyte number and greater astrocytic vulnerability to steroid-induced injury [17]; their loss could account for the Glt-1 protein level decrease.

In mouse models of steroid-induced depression, riluzole reversed steroid-induced hippocampal Glt-1 suppression and prevented despair behaviors [22], while increasing Glt-1 hippocampal protein in controls. Exercise also reversed social isolation-induced hippocampal Glt-1 protein decrease in rats [32]. A complex study of the molecular and behavioral effects of alcohol withdrawal in rats found that depressive and anxious behaviors could be reversed by ceftriaxone (which upregulates EAAT2 [24]), but they reappeared after DHK treatment [34]. Alcohol withdrawal was associated with significant Glt-1 protein reduction in lateral habenula, which was normalized by ceftriaxone.

Four papers examined the behavioral and molecular consequences of Glt-1 inhibition in rodents [28, 30, 31, 33]. DHK is a specific Glt-1 inhibitor [6]. In Sprague-Dawley rats, DHK infusion to the lateral ventricles [28] and amygdala [33] resulted in anhedonic/dysphoric behaviors as measured by intracranial self-stimulation. DHK infusion increased c-Fos (a proto-oncogene expressed by neurons after depolarization, and a proxy for neuronal activity) in reward and mood brain regions, including the nucleus accumbens, amygdala, and infralimbic cortex [28]. Mice with acute Glt-1 inhibition (via DHK) and chronic Glt-1 inhibition (flexed Glt-1 knockout) displayed depressive behaviors, reduced resilience to chronic stress, and sleep changes, with increased excitability of their lateral habenula neurons, which are typically activated by unpleasant stimuli [30]. In Wistar rats, DHK infusion to the infralimbic cortex prevented depression-like behavioral responses in stressful scenarios, though DHK had no effect when administered to the prelimbic cortex [31]. The infralimbic result differs from previously reported DHK-induced depressive behaviors. It is unclear whether the difference is related to strain differences between Sprague-Dawley and Wistar rats, or Glt-1 function in the infralimbic cortex. Although DHK is a specific inhibitor of Glt-1 and therefore acts on the glutamate system, the same study found that its antidepressant effect was prevented by serotonin depletion and AMPA glutamate receptor antagonism, while an antidepressant response was seen with the AMPA agonist (s-AMPA). These results probably reflect the complex interplay of multiple neurotransmitter systems in depression.

### Table 1 (continued)

| Reference | Description | Species | Tissue examined | Subject characteristics | Findings |
|-----------|-------------|---------|-----------------|-------------------------|----------|
| Kang et al. [34], 2018 | Explore the correlation between Glt-1 function, LHb activity, and alcohol withdrawal behaviors | Sprague-Dawley rats | Lateral habenula | Glt-1 protein levels reduced in LHb of alcohol withdrawal rats; Glt-1 protein levels normalized in LHb of alcohol withdrawal rats by ceftriaxone; alcohol withdrawal rats have higher depressive and anxious behaviors – these are reversed by ceftriaxone, and restored by DHK blockade of Glt-1 |

AQP4, aquaporin-4; CA3, cornu ammonis subregion 3; CEA, central amygdala; CRS, chronic restraint stress; CHF, Carioca High-Freezing rats; CLF, Carioca Low-Freezing rats; cLH, congenital learned helpless; cNLH, congenital not-learned helpless; CSDS, chronic social defeat stress; CUMS, chronic unpredictable mild stress; DHK, dihydrokainic acid; ECS, electroconvulsive shock; EEG, electroencephalogram; EMG, electromyogram; FSL, Flinders sensitive line; ICSS, intracranial self-stimulation; LHb, lateral habenula; NA, nucleus accumbens; PFC, prefrontal cortex.
EAAT2 mRNA expression was significantly decreased in MDD compared to controls.

Shao et al. [62], 2008

RNA extracted from postmortem brains of patients with BD, MDD, and controls for microarray analysis and in situ hybridization.

Table 2. Summary of human studies included in the systematic analysis

| Reference | Description | Subject characteristics | Tissue examined | Findings |
|-----------|-------------|-------------------------|----------------|---------|
| McCullumsmith et al. [54], 2002 | *In situ* hybridization of EAAT1/2/3/4 mRNA in postmortem brains of patients with SZ, BD, MDD, and controls | 60 postmortem brains: 15 SZ (M/F: 6/9, 5.8–6.6, age 44.2 [25–62]) 15 BD (9 M/F: pH 6.2 [5.8–6.6], age 42.3 [25–61]) 15 MDD (9 M/F: pH 6.2, [5.6–6.5], age 46.4 [30–65]) 15 control (9 M/F: pH 6.3 [5.8–6.6], age 48.1 [29–68]) 56 Caucassian, 2 Asian, 2 African-American | Striatal tissue: Caudate, Putamen | Ventral striatum EAAT2 mRNA was expressed in all striatal tissues; significantly lower EAAT2 mRNA levels in MDD compared to SZ; significantly lower EAAT3 mRNA levels in BD/SZ compared to controls; significantly lower EAAT4 mRNA levels in MDD compared to controls; significantly lower EAAT2 mRNA levels in ventral striatum than caudate/putamen across all diagnoses and controls; insufficient sample sizes to analyze effect of medications |
| Choudary et al. [50], 2003 | *In situ* hybridization of EAAT2 mRNA in glia of LC of men with MDD | 22 postmortem brains: 7 control (1 F/M; pH 7.0±0.1, age 50±16.7) 9 MDD (2 F/M; pH 6.98±0.1, age 48.0±14.2) 6 BD (1 F/M; pH 6.97±0.1, age 47.5±18.7) | Left DLPFC (BA49 and 46) | Anterior cingulate gyrus (BA24) EAAT2 mRNA expression was significantly decreased in AnCG and DLPFC in MDD compared to controls; EAAT2 mRNA levels were higher in AnCG than DLPFC in controls, BD, and MDD; EAAT3 mRNA levels decreased in AnCG and DLPFC in MDD compared to controls |
| Shao et al. [62], 2008 | RNA extracted from postmortem brains of patients with BD, MDD, and controls for microarray analysis and PCR validation of candidate genes; effects of age, sex, and pH controlled for by ANCOVA | 105 postmortem brains: 35 SZ (19 M/F; pH 6.8±0.25, age 42.9±9.6) 35 BD (15 M/F; pH 6.5±0.23, age 45.3±9.8) 35 control (23 M/F; pH 6.6±0.26, age 44.4±6.5) | DLPFC (BA46) | ANCOVA adjusted for pH and age, and restricted to above-mentioned pools of genes identified with qRT-PCR; EAAT2 mRNA expression decreased to statistical significance on microarray validation but did not validate, unclear whether due to targeting wrong region of gene; EAAT2 mRNA expression significantly increased in proportion to lifetime antidepressive exposure (fluoxetine equivalents) |
| Miguel-Hidalgo et al. [57], 2010 | Comparison of EAAT2 protein levels across cortical layers in left orbital frontal cortex of postmortem patients with major depression, alcohol use disorder, major depression + alcohol use disorder, and controls | 49 postmortem brains: 13 MDD (9 M/F; pH 6.6±0.18, age 48.5±18.89) 10 MDD + ALC (4 M/F; pH 6.4±16.23, age 50±16.1±20) 13 AD (11 M/F; pH 6±16.1±16, age 48.0±14.17) 13 control (6 M/F; pH 6.4±3±8, age 51.1±14.44) | Left orbitofrontal cortex (BA47) | EAAT2 protein levels significantly lower in MDD and MDD+ALC than controls, EAAT1 protein levels lower in MDD and MDD+ALC than controls; EAAT2 immunostaining highest in layer II, affected by age, postmortem delay, pH |
| Bernard et al. [48], 2011 | Microarray study of gene expression in LC of postmortem patients with major depression, BD, and controls; genes interrogated with ISH | 27 postmortem brains: 9 control (8 M/F; pH 6.94±0.04, age 50.7±5.65) 12 MDD (11 M/F; pH 7±3±0.04, age 49.6±4.92) 6 BD (5 M/F; pH 6.9±0.04, age 43.6±7.4) 25 Caucasian, 1 Island Pacific Islander, 1 African-American | LC | Glutamate signaling pathway most significantly altered in MDD; SLC1A2 (EAAT2) gene expression significantly decreased in midrostral LC in MDD on microarray and qR-PCR; SLC1A1 (EAAT1) gene expression significantly decreased in midrostral LC in MDD on microarray and qR-PCR; SLC1A2 (EAAT2) gene expression not decreased to statistical significance on ISH; SLC1A3 (EAAT3) gene expression significantly increased on ISH |
| Murphy et al. [58], 2011 | Genotyping for SNPs associated with suicide attempt or blood samples of 76 suicide attempters and 83 nonattempters, with a variety of diagnoses including BD, MDD, SZ, other psychotic disorders, substance/ alcohol use disorders, anxiety disorders, and eating disorders | 159 blood samples including: 9 BD (5 M/A nonattempt) 85 MDD (48/53 nonattempt) Comorbid diagnoses present for significant but unspecified numbers of patients | White blood cells | Suicide attempters significantly younger than nonattempters; genotypic associations found in blood samples of suicide attempters and nonattempters; no significant differences in genotype distribution at SNPs associated with suicide attempt or blood samples of suicide attempters and nonattempters from BD, MDD, and controls |
| Dallagoziania et al. [51], 2012 | DNA extracted from whole blood of 130 BD type 1 patients to examine effect of SLC1A2/SNPs rs4354868 (-181A>C) on glutamate function, illness recurrence, and effect of lithium treatment | 110 BD type 1 blood samples: 56 not taking lithium, or not taking for ≥ 26 months 54 taking lithium for ≥26 months All patients Caucasian, 76 F/M | White blood cells | Allicic frequencies similar to those of general population; rs4354868 TT (n = 37) gene expression significantly lower mood episode frequency; rs4354868 G/G (n = 16; reduced EAAT2 transcription) significantly associated with more manic episodes, greater episode frequency; treatment countered the G/G polymorphism effect and reduced recurrence rate |
| Chandler et al. [49], 2013 | Gene expression analysis of SLC1A2 in glia of LC of men with MDD | 49 postmortem brains: 20 controls (20 M; pH 6.59±0.08, age 42±4) 19 MDD (19 M; pH 6.62±0.06, age 43±4) | Pontine LC, right PPC (BA10) of subset | Significantly decreased SLC1A3 (EAAT1) in MDD homogenized LC than controls; no significant difference in SLC1A2 (EAAT2) expression between MDD and controls in homogenized LC; significantly lower SLC1A3 (EAAT1) in astrocytes of MDD compared to controls; significantly decreased SLC1A2 (EAAT2) expression in astrocytes of LC in MDD compared to controls; no significant difference in SLC1A2 expression in PPC between MDD and controls; insufficient protein to compare SLC1A3/EAAT1 gene expression with EAAT1/2 protein levels |
| Medina et al. [55], 2013 | *In situ* hybridization for genes for EAAT2 and VGLUT1 from hippocampus of patients with MDD and BD | 25 postmortem brains: 11 control (11 M; pH 6.6–7.2, age 49–78) 15 MDD (15 M/F; pH 6.1–7.1, age 19–72) 7 BD (2 M/F; pH 6.5–7.1; age 23–69) | Hippocampus | SLC1A3 (EAAT3) gene significantly downregulated in MDD, nonsignificantly downregulated in BD, significantly upregulated in MDD, and nonsignificantly upregulated in BD; SLC1A2 gene expression significantly downregulated in MDD but not BD |
and suggest that a Glt-1 inhibitor cannot reverse despair behavior in a serotonin-depleted or glutamatergically inhibited brain.

Several papers explored the pharmacological modulation of Glt-1 expression in depression models [22, 25, 26, 29–31, 35, 37, 38, 44–46], including antidepressant agents (fluoxetine, citalopram, escitalopram, ketamine, nortriptyline, and tianeptine) and their effect on Glt-1. In rats, fluoxetine reversed stress-induced depressive behaviors even in the presence of a Glt-1 inhibitor. In mice, fluoxetine reversed stress-induced depressive behaviors in animals with ketamine, nortriptyline, and tianeptine. In rats, tianeptine improved depressive behaviors even in the presence of a Glt-1 inhibitor.

| Reference | Description | Subject characteristics | Tissue examined | Findings |
|-----------|-------------|-------------------------|----------------|---------|
| Oh et al. [59], 2014 | Data-mined neuropathological data from multiple brain regions in online dataset from Stanley neuro-pathology consortium (SNICD) | 60 postmortem brains (15 SZ [M:6,F:9]; pH 5.8–6.6, age 25–62; 13C/2A) | DLPCF (BA46) | SLC1A2 mRNA expression significantly increased in DLPCF white matter of MDD compared to controls. SLC1A2 mRNA expression slightly increased in DLPCF deep gray matter in MDD compared to controls, lower SLC1A2 mRNA expression significantly correlated with decreased GAD1 mRNA expression in MDD but not controls |
| Poletti et al. [61], 2014 | Comparing effect of SLC1A2 SNP rs4354668 alleles on hippocampal gray matter volume in BD patients exposed to high scores of ACE vs low scores | 86 BD type 1 blood samples (56 F/30 M; none on lithium) | White blood cells | SLC1A2 SNP rs4354668 significantly influences gray matter volume in BD subjects with low ACE scores, with T/T homozygotes associated with significantly decreased gray matter volume. SLC1A2 SNP rs4354668 alleles made no differences in high ACE group; high ACE associated with hippocampal gray matter volume reduction |
| Florentino et al. [52], 2015 | Two data sets from UCL1 and UCL2 cohorts were screened for SLC1A2 variants associated with BD and compared with NHS samples of patients with SZ and with healthy European controls | 1,099 BD, 630 SZ, 1,095 control: 506 BD1 (UCL1, blood); 593 BD1 (UCL2, saliva); 630 SZ (NHS patients, blood) | Blood | No SLC1A2 gene variant associated with BD or SZ. Rare variants detected only in cases: rs1485827:57G>A found only in BD1 cases; no controls; rs19959866G>A found in 1 BD and 1 SZ case; no controls; rs3436686A>C failed to replicate association with lithium efficacy; rs475484G>T failed to replicate association with attempted suicide |
| Poletti et al. [60], 2015 | Assessed depressed BD type 1 impulsivity for cognitive performance; compared results to rs4354668 variants in the EAAT2 gene promoter, and rs2373380 variants in the EAAT1 gene promoter | 60 BD type 1, depressed, no ECT in previous 6 months; no controls | White blood cells | EAAT1 genotype distribution respected Hardy-Weinberg equilibrium; EAAT2 genotype distribution did not respect Hardy-Weinberg equilibrium; EAAT1 polymorphisms were significantly associated with differences in working memory, verbal fluency, and selective attention; EAAT2 polymorphisms showed no association with cognitive functions |
| Medina et al. [56], 2016 | Hippocampal RNA microarray analysis in MDD patients vs. controls | 23 postmortem brains: 13 MDD (13 M; pH 6.5–7.1; age 19–65) | Hippocampus | SLC1A2 (EAAT2) downregulated in MDD compared to controls; SLC1A3 (EAAT1) downregulated in MDD compared to controls; AQ4P downregulated in MDD compared to controls |
| Zhao et al. [63], 2016 | Comparison of RNA expression of components of glutamate metabolism in young MDD patients and controls, and elderly depressed patients and controls | 36 postmortem brains from Stanley Medical Research Institute (SMRI): 17 MDD (10 M/7 F; 16C/1 Hispanic; pH 6.3–6.8; age 24–63); 7 MDD (3 M/4 F; pH 6.3–6.9; age 36–56); 12 control (8 M/4 F; 11/1 Hispanic; pH 6.3–6.9; age 24–63); 37 postmortem brains from Netherlands brain bank (NBB): 22 elderly control (pH 5.80–7.39; age 56–96); 5 elderly MDD, 10 BD depressed (pH 6.26–6.82; age 45–93) | ACC (BA24); DLPCF (BA46 SMRI; BA9 NBB) | SMRI (younger patient group): EAAT1 and EAAT2 gene expression significantly decreased in DLPCF of suicide patients compared to non-suicidal MDD; EAAT1 and EAAT2 transcript levels significantly increased in the DLPCF of non-suicidal MDD patients compared to controls; no differences in gene transcription found between violent and non-violent suicide methods; no statistically significant differences in EAAT2 expression in ACC; EAAT1 and EAAT2 gene expression positively correlated with BDNF gene expression in MDD+ suicide patients NBB (elderly patient group): No suicides in this population; no significant differences in EAAT2 expression in ACC; no significant differences in EAAT2 expression between depressed group between those with and without suicidal ideation |
| Jia et al. [53], 2017 | Examine DNA methylation of SLC1A2 promoter in BD patients with and without binge-eating and substance use disorders | 30 BD type 1 blood samples: 30 BD (14 M/16 F; pH 42.6±15.4); 30 BD + comorbid BE (7 M/23 F; pH 45.2±14.8); 30 BD + comorbid ND (14 M/16 F; pH 35.2±13.1) | White blood cells | SLC1A2 promoter region hypermethylated in BD without addictions; SLC1A2 promoter region hypomethylated in BD with nicotine addiction and binge eating |

A. Asian; AA. African-American; ACC. anterior cingulate cortex; ACE. adverse childhood experiences; AD. alcohol dependence; ANCOVA. analysis of covariance; BA. Bredemann Area; C. Caucasian; BD. bipolar disorder; BD1. bipolar disorder type 1; or type 2; DLPCF. dorsolateral prefrontal cortex; GAD1. glutamic acid decarboxylase 1, SH, in situ hybridization; NHS. National Health Service of the United Kingdom; MDA. major depressive disorder with alcohol dependence; MDD. major depressive disorder; MDD-A. major depressive disorder – died by suicide; LC. locus coeruleus; PFC. prefrontal cortex; qRT-PCR. quantitative real-time polymerase chain reaction; Q-PCR. quantitative real-time polymerase chain reaction; SA. suicide attempt; SZ. schizophrenia; UCL1/2. University College of London Bipolar Affective Disorder genome wide association study cohorts 1 and 2.
on hippocampal Glt-1 protein levels [38]. In summary, fluoxetine improved depression-like behaviors in mice and rats, but only normalized hippocampal Glt-1 levels in rats without altering reduced hippocampal Glt-1 levels in mice. Various explanations for differential Glt-1 expression in response to fluoxetine include dose response and acuity (rats received fluoxetine 10 mg daily for 4 weeks [29], while mice received fluoxetine 20 mg/kg as a single dose [30] or daily for 7 days [38]), or species response differences. In rats, stress increased hippocampal Glt-1a and Glt-1b, and the tricyclic antidepressant (tianeptine) prevented Glt-1a increases, but not Glt-1b [25]. There were also marked differences in the effect of antidepressants across mouse strains [37]. In three strains (C57BL/6J, 129S1/SvemJ, FVB/NJ) chronic stress increased Glt-1 mRNA, while the antidepressants escitalopram and nor- triptyline decreased Glt-1, but in the fourth strain (DBA/2J) the opposite was demonstrated: stress decreased Glt-1 mRNA, while antidepressants increased it.

Ketamine is used off-label for rapid treatment of depression [69]. In rats, ketamine significantly reversed stress-induced decreases of hippocampal Glt-1 mRNA, decreased stress-induced hippocampal glutamate increases, and reversed depression-like behaviors (spontaneous locomotion, sucrose preference) [35, 46]. The effect of ketamine on Glt-1 may be modulated through BDNF-TrkB signaling, since the ketamine-induced Glt-1 changes were prevented when rats were pretreated with a TrkB receptor inhibitor [35]. Lithium is a mood stabilizer used in BD and (without Food and Drug Administration indication) in depression treatment and prophylaxis. In rats with stress-induced increased hippocampal Glt-1 mRNA, lithium prevented both this and loss of dendrite length [26].

Electroconvulsive shock (ECS) is an animal model of electroconvulsive therapy (ECT) which is a clinical treatment for depression in humans. Unlike ECT, however, ECS induces seizures in typically awake animals and therefore is itself a potential stressor. In chronically stressed rats, ECS further downregulated Glt-1, increased hippocampal glutamate, and worsened behavioral markers of depression [45]. The anesthetic agent propofol significantly normalized stress-induced and ECS-induced Glt-1 downregulation, glutamate concentration, and behavior [45].

Icariin is a flavonoid extracted from Horny Goat Weed and used extensively in Chinese traditional medicine, with antidepressant and neuroprotective effects in rodents. In prenatally stressed rats, icariin administered to male offspring reduced stress-elevated group I metabotropic glutamate receptors (mGluR1/5) mRNA and protein levels, and increased stress-reduced EAAT2 mRNA and protein levels. Zinc or fluoxetine decreased stress-induced depressive behaviors, but neither altered hippocampal Glt-1 levels [38].

Human Studies

Genetic Findings

Although several studies have analyzed genetic associations of SLC1A2 with mood disorders, no SLC1A2 single nucleotide polymorphism (SNP) has a confirmed diagnostic association with MDD or BD. In the currently available GWAS data from the Psychiatric Genomics Consortium (PGC) no SNP within a 20-kb window of SLC1A2 reaches GWAS significance for MDD [70] or BD [71], or schizophrenia [72] (no association having a p value lower than 5 × 10^{-8}). It is possible that rare variants may be detected in future sequencing studies, and for now this remains speculative. However, positive genetic correlations have been reported for subphenotypes of mood disorders. For example, minor alleles within SLC1A2 have been associated with the rapid cycling subtype of BD [73]. Additionally, SLC1A3 and SLC1A2 polymorphisms have been studied in mood and psychotic disorders, and found to be associated with differences in cognition, memory, executive function, suicide completion, and medication responses [74, 75]. The effects of 28 SNPs in 18 candidate genes were studied in a mixed population of 159 psychiatric patients including 85 MDD and 9 BD patients [58]. rs4755404 (an SNP in an intronic region of SLC1A2) was significantly associated with suicide attempts. Testing for associations with other clinical variables such as diagnosis, an association with schizophrenia and other psychotic disorders was found in a dominant model, but no effect was seen for MDD or BD, though associations with diagnoses was not the primary focus of the study. A functional SNP, rs4354668 (SLC1A2–181 A>C), affecting protein expression and plasma glutamate levels [76], was studied for its effect on history of illness in BD (n = 110) [51]. This study found that T/T homozygotes had a significantly lower mood episode frequency, while G/G homozygotes experienced higher mood episode recurrence (depressive, manic, mixed-state) and an increased mania-to-depression ratio. Furthermore, a multivariate analysis revealed an interaction between the genotype and lithium treatment with the effect of the genotype on episode recurrence rate only being present in lithium-untreated patients. The G allele of SNP rs4354668 (within the promoter region of SLC1A2) is associated with decreased EAAT2 activity and increased plasma
glutamate concentrations [60]. The minor allele T of rs43534668 was associated with reduced hippocampal gray matter volume, but only in patients with low numbers of adverse childhood experiences (ACEs) [61]. Patients with high ACEs displayed an overall reduction in gray matter volume compared to patients with low ACEs that was not further reduced in subjects with the T allele. Complete histories of psychopharmacological treatments were limited, but lifetime lithium exposure did not alter allelic associations. An association between brain volume and number of mood episodes was not reported. The rs43534668 G allele has been associated with worse cognition (executive function and working memory) in schizophrenia, but this effect was not observed in BD [60].

The largest genetic study of SLC1A2 in BD included 1,099 BD patients and 1,095 controls [52]. Its primary analysis included nonsynonymous variants, intron/exon junction variants, and SNPs in the putative regulatory region, whose minor allele frequency was unknown or <0.01. Two recurrent missense variants (rs145827578: G>A, rs199599866: G>A) and one recurrent 5′-UTR variant (ss825678885: G>T) were found in BD cases only (3, 1, and 2, respectively) and not controls; none of the analyzed SNPs reached statistical significance alone. Taking advantage of the large sample size, the authors also tried to replicate previous studies but found no association of rs4755404 with attempted suicide [58] and no association of rs4354668 with lithium efficacy, defined as a reduction in mood episode number [51].

Molecular Differences by Brain Region

Prefrontal Cortex. The prefrontal cortex (PFC) is an association cortex crucial to human cognition, organizing input from diverse sensory modalities, maintaining attention, working memory, and coordination of goal-directed behaviors. It comprises several Brodmann areas (BA) anterior to primary motor and premotor cortices, responsible for executive functioning (BA9/10/46), language processing (BA44/45), and emotional processing and sociability (BA47/10/11/13) [77]. The PFC has been studied in many psychiatric disorders, with differences found in conditions including autism, schizophrenia, MDD, and BD. Six articles studied EAAT2 expression for this brain region in MDD and BD [49, 50, 57, 59, 62, 63]. Microarray analysis of gene expression in the dorsolateral PFC (DLPFC: BA9/46) in MDD, BD, and controls found SLC1A2 downregulated in MDD (fold change: 0.71, p < 0.05) but not in BD [50]. Attempted verification by in situ hybridization (ISH) observed a nonsignificant downward trend. A partial corroboration [59] reported that SLC1A2 expression was significantly reduced in DLPFC white matter (BA46) of MDD compared to controls and found a trend for reduced expression of SLC1A2 in superficial and deep gray matter. However, the differences were very small and the p values were not corrected for multiple testing. Furthermore, the authors report a correlation between GAD1 (glutamate decarboxylase 1) and SLC1A2 gene expression in the deep gray matter of BA46 in MDD but not in controls. No significant results were reported for BD. Adding to gene expression data, EAAT2 protein expression in the left orbitofrontal cortex (BA47) of patients with MDD, MDD + alcoholism (MAD), alcoholism, and controls was examined [57]. Immunohistochemistry found significantly reduced EAAT2 protein levels in MDD and MAD but not in alcoholism alone, when compared with controls. The same trend was observed in Western blot, but the results were not statistically significant. Another study based on microarray analysis of gene expression in DLPFC in BD reported elevated EAAT2 levels after correcting for pH and age. However, this increase could not be validated by qPCR, with the authors speculating this was due to targeting the wrong region of the gene [62]. In contrast to these findings are reports of no differences in gene expression in homogenized PFC (BA10) between matched MDD and controls [49]. Grouping patients by age and suicide status found young MDD patients without suicide had elevated DLPFC (BA9) EAAT2 levels on qPCR when compared to controls and young MDD suicides [63]. No difference in gene transcription was found between violent and nonviolent suicide methods and no differences were reported between MDD and controls, or between MDD and controls in older patients, even when grouped according to suicidal ideation.

Anterior Cingulate Cortex. The anterior cingulate cortex (ACC) is part of the limbic system and surrounds the anterior part of the corpus callosum, comprising BA24, 32, and 33. It appears to play important roles in both autonomic and higher-level cognitive functions, including pain processing, performance monitoring, value encoding, decision making, emotions, learning, and motivation [78]. The ACC has been heavily studied in psychiatric disorders and has been associated with MDD, schizophrenia, and ADHD. One study described a trend in ACC similar to DLPFC: significant downregulation of SLC1A2 in microarray with a downward trend strongest in layer 2 [50]. In a direct comparison between the two brain regions between all patients and controls, ACC had higher SLC1A2 expression than DLPFC. A separate study found no differences in ACC SLC1A2 levels in MDD compared...
to controls, nor in any of the other comparisons (young suicidal vs. nonsuicidal vs. controls; older patients vs. controls; older patients with suicidal ideation vs. controls) [63]. However, a weak correlation ($p = 0.036$) between levels of $SLC1A2$ and TrkB.T1 (the truncated receptor of BDNF) was reported in MDD patients regardless of suicidal ideation.

$Locus coeruleus$. The locus coeruleus (LC) is the primary site of central nervous system norepinephrine synthesis and is involved in autonomic responses (arousal, sleep/wake cycles, stress response) and higher cognition (attention, memory, perception, motivation) [79]. Differential expression in the midrostral LC of genes involved in glutamate signaling was studied in healthy controls and MDD [48]. $SLC1A2$ was decreased in microarray and qPCR, but not ISH, with no differences between BD and controls. A separate study of glutamatergic and astrocytic markers in postmortem homogenized pontine LC and microdissected astrocytes/oligodendrocytes found no differences in LC $SLC1A2$ expression between MDD and controls [49]. However, there was a significant reduction of $SLC1A2$ in astrocytes ($p < 0.001$) but not oligodendrocytes in MDD compared to controls.

$Striatum$. The striatum is part of the subcortical basal ganglia and plays an important role in motor and reward systems, including reward cognition, reinforcement, and motivation. It receives glutamatergic and dopaminergic inputs. One study reported no difference in EAAT2 mRNA levels measured by ISH in the ventral striatum, caudate, or putamen between MDD and BD diagnoses [54]. However, EAAT2 mRNA levels were significantly lower in the ventral striatum than in the caudate nucleus and putamen across all diagnoses and controls.

$Hippocampus$. This brain region is primarily involved in memory formation. A study of microdissected hippocampus in MDD, BD, and healthy controls found a significant downregulation of EAAT2 in MDD but not BD patients by microarray and ISH [55]. These results were confirmed when measuring samples from the same cohort by qPCR ($p = 0.002$) [56].

**Discussion**

EAAT2 may play a role in the etiology and phenotype of mood disorders but the data are still contradictory. Possible mechanisms include preventing glutamate excitotoxicity, regulation of glutamate release, modulating glutamatergic receptor activation, and (as part of a macromolecular complex) intracellular water/ion homeostasis and energy metabolism. Gross neuronal injury results in a massive release of glutamate [80, 81], activating glutamate receptors and causing an excitotoxic cascade [82]. Excitotoxicity causes injury and death of all three components of the neurovascular unit (neuron, glia, vascular cells) [83], further reducing EAAT2 levels while releasing additional glutamate. Glutamate neurotoxicity as a mechanism in psychiatric disorders remains hypothetical [84] but is already being explored in vivo via magnetic resonance spectroscopy (MRS) [85]. MRS combines glutamate + glutamine (the nonexcitotoxic precursor) as a Glx measurement, and studies have found that MDD is typically associated with decreased Glx, while BD is associated with increased Glx. Such data could be combined with EAAT2 studies to determine whether EAAT2 adapts to glutamate, or whether differential EAAT2 expression shifts the glutamatergic signaling system [86]. Arguably, induced animal models of depression best mimic environmentally induced adjustment disorders in humans and may not reflect the same biological mechanisms as primary mood disorders in human subjects. However, the most common findings in human and animal depression models were decreased EAAT2/Glt-1. A small number of animal studies reported Glt-1 upregulation in response to stress/corticosterone, perhaps reflecting an adaptive Glt-1 response to glutamate neurotoxicity. EAAT2 increases and decreases may reflect different experimental chronic stress duration, time of measurement since last stress exposure (i.e., acute stress response superimposed on chronic stress), and in at least one study, strain-specific differences in glutamatergic systems [37]. MRS studies are scarce in rodents, but comparison of glutamate levels with Glt-1 mRNA/protein levels would help clarify whether Glt-1 function and expression adapt to glutamate fluctuations as hypothesized.

A recent functional brain imaging study uncovered an interaction between $SLC1A2$ polymorphisms and ACE severity upon axial diffusion (along axonal fiber axis) in white matter tracts of BD patients [87]. This is important evidence for the role of glutamate in brain structural changes in response to stress. If stress and depression are associated with decreased glutamate concentrations, as human MRS studies have suggested, EAAT2/Glt-1 downregulation may be an adaptive glial transcriptional and/or translational response to reduced synaptic glutamate concentrations and subsequent reduced need for glutamate clearance. Glutamate reduction may be secondary either to absolute decreased glutamate levels (such as recycling back to glutamine and/or reduced glutamate synthesis) or decreased presynaptic glutamate release. An-
other mechanism of EAAT2/Glt-1 reduction might involve decreased glial fibrillary acidic protein (GFAP), a filament protein within astrocytes associated with EAAT2 cell membrane trafficking [88]. Consistent with this mechanism, elderly depressed and suicide subjects have demonstrated decreased GFAP [89, 90].

EAAT2 and AQP4 form a macromolecular complex and co-express with multiple other membrane proteins, as well as mitochondria and glycolytic enzymes [18]. In addition to regulating water and ion homeostasis of the neurovascular unit [91], AQP4 may significantly modulate astrocytic function and adult neurogenesis in depression. It has been previously noted that AQP4 rodent knockouts lose both the fluoxetine-induced antidepressant effect and hippocampus neurogenesis [92] and have greater depressive behaviors in response to chronic corticosterone administration [17]. The co-regulation of EAAT2 and AQP4 in mood disorders needs future investigation.

We found no study that reviewed the effect of antipsychotics on EAAT2 expression in human mood disorders. This is relevant because chronic antipsychotic exposure in primates has been associated with decreased brain volume [93] and decreased glial numbers [94]. Chronic antipsychotics in rodents have been associated with decreased Glt-1 [95], which might be secondary to glial decreases. These data are pertinent to future human studies of EAAT2 in mood disorders, as antipsychotics are clinically used for mood stabilization and as adjunctive therapy of depressive and anxious symptoms.

Individual SNPs are difficult to associate with mood disorders across genetic studies, partly reflecting the complexity of the pathophysiology of these conditions and partly the need for very large samples for adequate power. Human studies have suggested genetic and epigenetic contributions to BD and MDD, including differential expression of the SLC1A2 gene. Our group, for example, has studied epigenetic modifications of CpG islands in SLC1A2 promoter regions in BD with and without comorbid substance or food addictions [53]. BD was associated with hypermethylation, while BD + comorbid addictions were associated with hypomethylation, and the effect of addiction on epigenetic modulation of psychiatric disorders remains an area of ongoing research. Human studies have generally reported that MDD is associated with decreased SLC1A2 gene expression in most brain regions, but this has not been replicated in all brain regions, patient cohorts, or cortical layers. One study reported EAAT2 immunostaining was most pronounced in cortical layer II [57], a layer with high intracortical connectivity; the significance of this relationship requires further exploration. Studies have not reported significant associations between BD and SLC1A2 expression, except one reporting increased SLC1A2 [62]. A difficulty with the human studies reviewed is that they report weak associations and are frequently not corrected for multiple testing. This is especially important in studies of microarray data. Additionally, our systematic process found no studies of gene expression in similar populations/brain regions explicitly reporting negative findings of SLC1A2, leading to a potential bias towards positive or false-positive findings.

While unipolar depression seems to be associated with decreased SLC1A2 gene and EAAT2/Glt-1 protein expression, no clear conclusion on the role of EAAT2 in BD can be drawn from the current data, pointing towards varying contributions of EAAT2 to the pathophysiology of different psychiatric disorders. Significant mechanistic questions remain, including the effect of brain regions and regional neural networks, genetic influences on glutamatergic regulation, and glial number. The involvement of EAAT2, not only in preventing excitotoxicity but also as part of a localized macromolecular complex, suggests that EAAT2 may be part of a complicated regulatory system, affecting not just glutamatergic pathways but also glial and neuronal cell homeostasis [96].

**Future Perspectives**

An increase in the understanding of the role of EAAT2 in mood disorders is highly warranted. Future studies could include the following areas of research: correlating EAAT2 with glutamatergic markers in vivo, comparing EAAT2 protein/mRNA expression in unipolar depressive disorders during depressive episodes and in remission, and in BD between manic/hypomanic, euthymic, and depressive mood states. Further understanding of the roles of the components of the macromolecular complex may also prove highly beneficial to elucidating roles and mechanisms of action of EAAT2.

**Statement of Ethics**

The authors have no ethical conflicts to disclose.

**Disclosure Statement**

The authors have no conflicts of interest to declare.
EAAT2 in Bipolar Disorder and Unipolar Depression

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Author Contributions

C.J.B. and M.V. conceived and designed the study. C.J.B., L.M.W., V.M., A.M.C.H., and M.V. performed data extraction. C.J.B. and L.M.W. undertook quality analysis. All authors provided critical feedback and participated in analysis and manuscript content. M.S., M.A.F., and M.V. supervised the project.

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