Unique prefrontal GABA and glutamate disturbances in co-occurring bipolar disorder and alcohol dependence

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Bipolar disorder (BD) and alcohol dependence (AD) frequently co-occur, and co-occurring BD and AD are associated with devastating public health costs. Minimal neurobiological research exists to guide the development of effective treatments for this treatment-resistant population. We believe the present study represents the first investigation of prefrontal gamma-aminobutyric acid (GABA) and glutamate levels in co-occurring BD and current AD. The participants were 78 individuals who met DSM-IV criteria for BD I/II and current AD (n = 20), BD I/II alone (n = 19), current AD alone (n = 20) or no diagnosis (n = 19). The participants completed a baseline diagnostic visit, then returned approximately 4 days later for a two-dimensional J-resolved proton magnetic resonance spectroscopy (1H-MRS) acquisition in dorsal anterior cingulate cortex (dACC). All participants were required to demonstrate ≥ 1 week of abstinence from alcohol/drugs via serial biomarker testing before 1H-MRS. A 2 × 2 factorial analysis of variance of cerebrospinal fluid (CSF)-corrected GABA/water concentrations demonstrated a significant BD × AD interaction (F = 2.91, P < 0.05), signifying uniquely low levels of GABA in BD+AD; this effect doubled when the sample was restricted to individuals who consumed alcohol within 2 weeks of 1H-MRS. There were no overall effects of BD/AD on CSF-corrected glutamate/water levels. However, the BD × AD interaction, signifying uniquely low levels of glutamate in BD+AD, approached statistical significance (F = 3.83, P = 0.06) in individuals who consumed alcohol within 2 weeks of 1H-MRS. The dACC GABA levels were significantly, negatively associated with Barratt Impulsiveness Scale (r = −0.28, P = 0.02) and Obsessive Compulsive Drinking Scale (r = −0.35, P < 0.01) scores. If replicated, these results may suggest that future treatment studies should preferentially evaluate therapeutics in BD+AD known to increase prefrontal GABA and glutamate levels.

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INTRODUCTION

Individuals with bipolar disorders (BD) have disproportionately high rates of alcohol dependence,¹,² and the combination of BD and AD (BD+AD) is associated with substantially worse outcomes than either disorder alone, including poorer clinical course,³ treatment resistance,⁴ suicidality,⁵ violent crime,⁶ disability⁷ and hospitalization.⁸ Unfortunately, treatment research for co-occurring BD and AD is remarkably limited. Only seven double-blind, placebo-controlled studies of medications for BD+AD have been conducted to date,⁹–¹⁵ and only one of these studies demonstrated a medication (that is, valproate) effect on drinking.¹² Better understanding the mechanisms that give rise to co-occurring BD and AD could help to identify differences in genetic risk, prevention strategies and novel therapeutic targets.¹⁶ The BD and AD research literatures support the hypothesis that dysregulated glutamate and γ-aminobutyric acid (GABA) transmission may partially explain the high observed diagnostic co-occurrence between BD and AD, both directly and through their impact on salient neurobehavioral constructs (impulsivity, drug craving) central to BD and AD.

Glutamate and GABA, the main excitatory and inhibitory neurotransmitters in mammals, respectively, are amino acids that are principally involved in the coordination of cortical activity, synaptic plasticity and modulation of most other neurotransmitter systems.¹⁷,¹⁸ Data from animal studies suggest that acute alcohol exposure augments the effects of GABA on the GABA_A receptor and antagonizes the effects of glutamate on the N-methyl-D-aspartate (NMDA) receptor; prolonged ethanol exposure results in compensatory downregulation of GABA_A receptors and upregulation of NMDA receptors, culminating in a hypoGABAergic/hyperglutamatergic state in alcohol withdrawal.¹⁹,²⁰ Proton magnetic resonance spectroscopy (1H-MRS) studies have demonstrated fluctuating brain glutamate levels in individuals with AD alone, depending on time of drinking; low prefrontal glutamate levels have been observed during active drinking¹¹ and 2–50 days since last drink,²²–²⁴ whereas abnormally high glutamate levels observed during acute withdrawal.²⁵ A recent investigation demonstrated that alcohol craving, measured by the Obsessive Compulsive Drinking Scale (OCDS),²⁶ was positively correlated with glutamate+glutamine levels in the nucleus accumbens and anterior cingulate cortex (ACC) in AD participants.²⁵ Conversely, 1H-MRS investigations of GABA in AD have been rare and have produced inconsistent findings.²⁷,²⁸,²⁹ Studies of adults with BD have demonstrated links between genes responsible for coding ionotropic glutamate receptor subunits and risk of BD,³⁰ as well as response to lithium,³¹ the gold standard maintenance treatment for BD. Several lines of evidence suggest altered GABA neurotransmission in BD as well, including reduced levels of cerebrospinal fluid and plasma GABA,³²,³³ and differences in GABA receptor genes, GABRA₁,³⁴
GABRA3 (ref. 35) and GABRA5.36 In animals, mood-stabilizing medications used to treat bipolar disorder, such as lithium and valproate, have been shown to decrease intrasynaptic glutamate via a number of mechanisms (for example, inhibition of NMDA receptor-mediated Ca2+ influx).37 H-MRS studies in BD have demonstrated that individuals with BD alone have elevated prefrontal glutamate across mood states.38–44 In contrast, H-MRS investigations of GABA in BD have produced mixed,45–47 and largely null,48–50 findings. Only one H-MRS study of co-occurring BD+AD has been published to date,51 despite the clinical importance of this population. Male participants with BD alone (that is, no history of AD) were found to have higher glutamate concentrations in the left dorsolateral prefrontal cortex relative to female participants with BD alone, participants with co-occurring BD and AD, and healthy controls. This study was limited to the BD+AD participants in remission (that is, 6 months to 22 years of abstinence) only, and did not include a comparator group of non-bipolar AD participants, making it difficult to distinguish the extent of contribution of alcohol to the finding.

Impulsivity, often defined as the tendency to respond without forethought,52 has been hypothesized as a critical ‘link’ between BD and SUD.53 Impulsivity has at least two dimensions (impaired response inhibition, impulsive choice), both are elevated in individuals with AD54,55 and BD53,56 regardless of mood state.57 Preclinical studies have shown that injections of NMDA receptor antagonists are associated with impulsive behavior; they appear to increase extracellular levels of glutamate due to their disinhibition of GABAergic inputs.59 Similarly, a recent study demonstrated increased impulsive behavior with infusion of the GABAβ receptor agonist muscimol into rat ventromedial prefrontal cortex.60 Finally, a number of 1H-MRS studies have demonstrated a positive association between ACC glutamate and impulsivity61–63 and a negative association between dorsolateral prefrontal cortex64 and ACC63,65 GABA and impulsivity across clinical populations. Elevated impulsivity might provide some basis for substance use and relapse potential in both BD and AD.

The present study was designed to provide a first 1H-MRS study of prefrontal GABA and glutamate levels in individuals with co-occurring BD and current AD, BD alone, current AD alone and healthy control (HC) subjects. We hypothesized a priori that individuals with co-occurring BD and AD would have significantly different prefrontal glutamate and GABA levels relative to individuals with BD or AD alone or healthy controls. We also hypothesized a priori that individuals with co-occurring BD and AD would have significantly higher scores on self-report measures of trait impulsivity and alcohol craving relative to individuals with BD or AD alone or healthy controls, and that impulsivity and alcohol craving would be associated with prefrontal GABA and glutamate levels, across participants.

MATERIALS AND METHODS

Participants

Seventy-eight individuals who met DSM-IV criteria for BD I or II and ‘current’ AD (that is, full criteria met within the past 6 months) (BD+AD, n = 20), BD I or II alone (BD, n = 19), current AD alone (AD, n = 20), or no psychiatric diagnosis (HC, n = 19) were recruited from inpatient and outpatient clinical settings and community advertisements. The sample size was chosen to provide approximately 80% power to detect large (that is, 0.8–1.0) effect sizes. Written informed consent was obtained from every participant. To avoid confounding with the effects of acute alcohol or drug intoxication or withdrawal, all the participants were required to demonstrate ≥ 1 week of abstinence from alcohol and drugs via serial self-report (Timeline Followback, TLFB69,70), the sensitive urine marker of recent alcohol use, ethyl glucuronide (EtG62) and urine drug screen (UDS) testing. Participants who reported any alcohol or drug use or provided positive EtG or UDS tests over the 1-week abstinence period were given two additional chances to achieve 1 week of abstinence before exclusion. In the case of conflicting TLFB and cannabinoid-UDS data, we used a validated algorithm to distinguish new cannabis use from residual cannabinoid excretion over the course of the 1-week monitored abstinence period.68 Participants whose positive cannabinoid-UDS was determined to be due to residual cannabinoid excretion (n = 4) were retained. General exclusions included serious medical illness or history of head injury, psychotic disorder, recurrent major depressive disorder, past month posttraumatic stress disorder, nonspecific compulsive disorder or eating disorder (assessed with the Structured Clinical Interview for DSM-IV Axis I Disorders, SCID-I69), current use of benzodiazepines or antidipsotropics (that is, naltrexone, acamprosate, disulfiram), past month electroconvulsive therapy, history of delirium tremens or >1 lifetime alcohol withdrawal seizure, acute alcohol withdrawal (that is, Clinical Institute Withdrawal of Alcohol Scale, CIWA-Ar45) or daily drug use in the past month. Co-occurring alcohol use disorder was exclusionary for BD and HC, but not BD+AD and AD, participants. BD+AD and AD participants were required to exceed healthy weekly drinking levels (that is, >7 drinks per week for females, >14 drinks per week for males71) in the past month (or, alternatively, the month preceding abstinence), whereas the BD and HC participants exceeding healthy weekly drinking levels in the past month were excluded. Finally, the BD+AD or BD participants with substantial (that is, >20%) medication dose changes <1 week before magnetic resonance imaging (MRI) were excluded.

Procedure

The study was conducted at the Medical University of South Carolina (MUSC) from 17 April 2013 to 24 August 2015. The study protocol was approved by the MUSC institutional review board. The participants completed a baseline diagnostic visit including the SCID-IV,72 Montgomery-Asberg Depression Rating Scale,73 Young Mania Rating Scale,72 TLFB, EtG, UDS, Barratt Impulsiveness Scale (BIS 1174), OCDS,75 demographic interview and brief abstinence counseling with a licensed PhD/MD clinician. The participants then returned approximately 4 days later (M = 3.60, s.d. = 0.83) for an MRI visit. Before scanning, the participants repeated the Montgomery-Asberg Depression Rating Scale, Young Mania Rating Scale, TLFB, EtG and UDS. Following these procedures, and with evidence of abstinence from alcohol and drugs, the participants completed the H-MRS scan in a Siemens 3.0 T Trio MR scanner with actively shielded magnet, high-performance gradients (45 mT m−1, 200 mT m−1 s−1), and a 32-channel head coil. A structural scan was taken for 1H-MRS voxel placement and tissue segmentation (TR/TE/repetition time/echo time) = 1900/2.26 ms; field of view = 256 mm2; flip angle = 9°; spatial resolution = 1 × 1 × 1 mm). The 1H-MRS voxel was placed in the dorsal ACC (dACC) on midsagittal T1-weighted images, posterior and superior to the genu of the corpus callosum, with the ventral edge of the voxel aligned with the dorsal edge of the callosum.25 A dACC voxel was chosen because most 1H-MRS studies in AD or BD have focused on this region, and because dACC has been associated with impulsivity76 and alcohol craving77 in 1H-MRS studies. A voxel size of 2.5x2.5x3 cm was selected to ensure adequate signal to noise properties.78 Following placement of six outer-volume suppression bands approximately 1 cm away from the voxel faces and auto-shimming, single-voxel, water-suppressed 1H-MRS spectra were acquired using a two-dimensional J-resolved Spectroscopy sequence: TR/TE = 2400/329 ms; ΔTE = 2 ms; four signal averages per TE step with online averaging; two-dimensional (2D) spectral width = 2000–500 Hz; 2D matrix size = 2048 × 100; total acquisition time = 13:28 min (ref. 74) water unsuppressed 2D 1H-MRS data were also acquired from the dACC voxel with two signal average recorded for each TE step (total acquisition time = 3:28 min). See Figure 1 for a representative voxel placement and sample spectrum.

Analysis plan

Skull stripping and whole-brain tissue-type segmentation were performed on MP-RAGE images using the FSL BET and FAST tools.79 In-house MATALAB functions were used to extract the three-dimensional volume corresponding to the positioned MRS voxel to obtain within-voxel gray matter, white matter and cerebrospinal fluid tissue content for each subject. Eddy currents and residual water were removed using in-house MATALAB functions. Subsequently, the Profit algorithm was applied using software-supplied 2D basis sets (which contain 19 metabolites total75). Before Fourier transformation, the raw 2D matrix was zero-filled. CRLB (Cramer-Rao Lower Bound) values, which reflect the uncertainty of estimated model parameters, were provided by the ProFit software. Estimated GABA and Translational Psychiatry (2017), 1–8
glutamate peak areas were normalized to the unsuppressed water signal. Finally, GABA/water and glutamate/water ratios were corrected for (that is, multiplied by) within-voxel cerebrospinal fluid fraction (that is, cerebrospinal fluid proportion+1).74 Participant characteristics were compared between groups using one-way analysis of variance for continuous variables (followed by pairwise independent samples t-tests in the case of significant omnibus tests) and chi-square for categorical variables (followed by pairwise chi-square tests in the case of significant omnibus tests). The associations between select participant characteristics and GABA and glutamate levels were tested using independent samples t-tests. The primary dependent variables (GABA and glutamate) were evaluated for potential violations of statistical test assumptions. The analysis of GABA and glutamate levels was accomplished via 2 (BD) × 2 (AD) factorial analyses of variance. These models were run both in the full sample, as well as in the restricted sample of participants who consumed alcohol within 2 weeks of MRI due to our previous finding of associations between recent drinking and dACC glutamate in AD.77 The levels of additional estimated metabolites (for example, creatine, N-acetyl aspartate) were not examined as part of planned analyses to control alpha inflation due to multiple comparisons. However, results from exploratory 2×2 factorial analyses of variance for other estimated metabolites are provided in Supplementary Table 1.

RESULTS

Ten participants (BD+AD n=3, BD n=1, AD n=5, HC n=1) required one to two additional visits between baseline and MRI to demonstrate ≥1 week of abstinence from alcohol/drugs. Across groups, 40% of the participants were female, 30% were smokers and 40% reported consuming alcohol ≤2 weeks before the MRI; mean (s.d.) age = 38.5 (11.8).

Preliminary evaluation of potential covariates

See Table 1 for a detailed comparison of participant characteristics by participant group. There were marginal group differences in participant group demographics, as well as in levels and percent heavy drinking days (≥4 drinks per day for men, ≥3 drinks per day for women) reported via Timeline Followback. There were no significant differences in sex, age, or alcohol consumption between groups for alcohol dependence and bipolar disorder. However, there were significant differences in smoking status, drug dependence, anxiety disorders, and BD subtype between groups. There were also significant differences in medication use between groups, with a higher proportion of participants with bipolar disorder and alcohol dependence receiving lithium, antipsychotics, anticonvulsants, and antidepressants compared to healthy controls. Additionally, there were significant differences in self-reported alcohol consumption between groups, with participants with bipolar disorder and alcohol dependence reporting higher levels of alcohol consumption compared to healthy controls.

Table 1. Patient characteristics by participant group

| Participant group | P         |
|-------------------|-----------|
| BD and AD¹ (n=20) | AD alone² (n=19) | BD alone³ (n=20) | Healthy controls⁴ (n=19) |
| Age (M, s.d.)     | 36.8 (12.5) | 43.1 (12.0) | 36.3 (11.4) | 38.0 (11.1) | 0.267 |
| Sex (% female)    | 35.0       | 36.8       | 45.0       | 42.1       | 0.912 |
| Drank w/in 2 wk (%) | 40.0       | 36.8       | 45.0       | 36.8       | 0.947 |
| %HDD (M, s.d.)    | 41.7 (25.3)⁴ | 50.2 (23.4)⁴ | 1.8 (5.0)¹,² | 1.3 (3.3)¹,² | 0.000 |
| Smoking status (%) | 50.0¹     | 31.6       | 15.0¹     | 21.1       | 0.080 |
| Drug dependence (%) | 40.0³,⁴  | 36.8³,⁴   | 0.0³,⁴   | 0.0³,⁴   | 0.000 |
| Anxiety disorder (%) | 45.0³      | 21.1⁴      | 40.0³      | 0.0³,⁴   | 0.000 |
| BD subtype (% I)  | 70.0²,⁴   | 0.0²,³    | 70.0²,⁴   | 0.0²,³    | 0.000 |
| MADRS (M, s.d.)   | 9.8 (8.7)²,⁴ | 3.5 (4.2)¹,³ | 9.3 (6.7)²,⁴ | 1.1 (2.1)¹,³ | 0.000 |
| YMRS (M, s.d.)    | 2.4 (3.9) | 1.4 (1.7) | 1.8 (2.9) | 0.4 (0.9) | 0.113 |
| ADS (M, s.d.)     | 18.4 (8.3)²,⁴ | 17.4 (8.8)³,⁴ | 1.9 (2.2)¹,² | 0.7 (1.4)¹,² | 0.000 |

Abbreviations: AD, alcohol dependence; ADS, Alcohol Dependence Scale; BD, bipolar disorder; %HDD, % of the 90 days preceding MRI that NIAAA-defined heavy drinking days (≥4 drinks per day for men, ≥3 drinks per day for women) were reported via Timeline Followback; MADRS, Montgomery-Asberg Depression Rating Scale; YMRS, Young Mania Rating Scale. Drank w/in 2 wk indicates reported consuming alcohol within 2 weeks of MRI. Drug Dependence represents past year drug dependence. P < 0.10 omnibus F (continuous variables) and χ² (categorical variables) tests were followed up with pairwise tests. Within-cell superscript numbers denote which participant groups a given group was significantly different from (P < 0.05) in these follow-up comparisons.
smoking ($P = 0.08$), but smoking was not independently associated with dACC glutamate ($P = 0.70$) or GABA ($P = 0.19$) levels across the sample. Recency of alcohol consumption (that is, number of days of abstinence preceding the MRI visit) did not differ between BD and AD ($M = 32.4$, s.d. = $33.3$) and AD ($M = 29.5$, s.d. = $23.0$) participants ($P = 0.75$). There were marginal differences in anticonvulsant use ($P = 0.06$; lamotrigine $n = 11$, divalproex sodium $n = 2$, gabapentin $n = 6$, topiramate $n = 1$, oxcarbazepine $n = 1$), with more BD+AD (65%) than BD (35%) participants taking anticonvulsants, but anticonvulsant use was not significantly independently associated with glutamate (BD+AD $P = 0.77$, BD $P = 0.52$) or GABA (BD+AD $P = 0.92$, BD $P = 0.69$) levels. Co-occurring past year drug dependence consisted primarily of cannabis dependence (76.9% of total drug dependence diagnoses; 15.4% stimulant and 7.7% cocaine dependence), and co-occurring drug dependence was not independently associated with glutamate (BD+AD $P = 0.77$, BD $P = 0.52$) or GABA (BD+AD $P = 0.92$, BD $P = 0.69$). Very few participants met criteria for drug dependence within the month preceding the study that is, BD+AD $n = 4$, AD $n = 2$ and few participants reported any illicit drug use within 2 weeks of MRI (that is, BD+AD $n = 1$, AD $n = 4$, BD $n = 3$). Co-occurring anxiety disorder diagnoses consisted primarily of social phobia (53.8% of total anxiety disorder diagnoses) and GAD (19.2%; 15.4% specific phobia, 7.7% NOS, 3.8% panic disorder), and co-occurring anxiety disorder was not independently associated with glutamate ($P = 0.24$) or GABA ($P = 0.56$) levels.

Primary analysis

The mean water line-width was 6.57 Hz (full width at half maximum; s.d. = 0.79 Hz, maximum = 9.07 Hz). Average gray matter to brain matter (brain matter = gray matter + white matter) ratio was 61.12% (s.d. = 3.24%) and did not differ between the participant groups ($F = 0.27$, $P = 0.85$). Mean glutamate CRLB % was 2.22 (s.d. = 1.39, maximum = 6.29). Seven participants ($n = 4$ HC, $n = 2$ BD, $n = 1$ AD) were excluded from analyses involving GABA, due to GABA CRLB > 20%. Mean GABA CRLB % for the remaining participants was 8.20 (s.d. = 3.88, maximum = 19.99). A 2 × 2 factorial, general linear modeling analysis of dACC GABA/water concentrations demonstrated a significant main effect of AD ($F = 5.36$, $P = 0.02$) that was qualified by a significant BD × AD interaction ($F = 2.91$, $P = 0.05$), signifying uniquely low levels of dACC GABA/water in BD+AD relative to BD, AD and HC (Cohen’s $d$ for BD+AD vs HC = 0.78). This effect nearly doubled when the sample was restricted to individuals who reported consuming alcohol within the 2 weeks before MRI ($n = 28$, BD × AD interaction $F = 2.15$, $P = 0.03$, Cohen’s $d$ for BD+AD vs HC = 1.53); see Figure 2. In the overall sample, there were no significant effects of BD and/or AD on dACC glutamate/water levels (BD × AD interaction $F = 1.46$, $P = 0.23$, Cohen’s $d$ for BD+AD vs HC = 0.33). However, when the sample was restricted to individuals who reported consuming alcohol within the 2 weeks preceding their scan ($n = 31$), the BD × AD interaction, signifying uniquely low levels of dACC glutamate/water in BD+AD (Cohen’s $d$ for BD+AD vs HC = 1.01), approached significance ($F = 3.83$, $P = 0.06$) and reached significance ($F = 4.21$, $P = 0.05$) once within-voxel gray matter/brain matter composition was statistically controlled; see Figure 2. To further explore this finding, we estimated Pearson correlations between glutamate/water levels and number of days of abstinence from alcohol. We restricted this analysis to individuals who reported drinking within 30 days of their scan ($n = 52$) to reduce the influence of outliers on results while retaining enough cases to estimate stable within-group correlations. Glutamate/water levels and days of abstinence were not significantly correlated in the full sample ($r = 0.18$, $P = 0.19$). However, examination of within-group associations revealed that glutamate/water levels and days of abstinence were significantly correlated in the BD+AD group ($r = 0.63$, $P = 0.02$) but not in any of the other groups (BD $r = 0.19$, $P = 0.58$; AD $r = 0.02$, $P = 0.96$; HC $r = 0.14$, $P = 0.63$).

As hypothesized, scores on the BIS were highest in BD+AD relative to BD, AD or HC individuals, with both BD and AD demonstrating significant main effects on BIS (BD $F = 24.25$, $P < 0.01$; AD $F = 9.13$, $P < 0.01$; BD × AD $F = 1.07$, $P = 0.30$); see Figure 3. BD+AD also scored highest on the OCDS followed by AD, despite similar recent alcohol consumption between BD+AD and AD participants (for example, drinks per drinking day, $P = 0.44$); BD and HC had similarly low OCDS scores (BD $F = 4.25$, $P = 0.04$; AD $F = 49.86$, $P < 0.01$; BD × AD $F = 3.36$, $P = 0.07$); see Figure 3. The dACC GABA/water levels were significantly negatively correlated with BIS ($r = 0.28$, $P = 0.02$) and OCDS ($r = 0.35$, $P < 0.01$) scores across groups; see Figure 4. Although glutamate/water concentrations were not significantly associated with BIS ($r = 0.03$, $P = 0.83$) or OCDS ($r = 0.18$, $P = 0.11$) scores in the full sample, glutamate/water levels were significantly negatively associated with OCDS ($r = 0.35$, $P = 0.05$) scores in individuals who reported consuming alcohol within 2 weeks of MRI; association between glutamate/water and BIS in this subsample was similar in magnitude but not significant ($r = 0.22$, $P = 0.25$). Although BIS was normally distributed, OCDS was not, because HC and BD without AD had predictably low OCDS scores (that is, all under ‘5’). However,
correlations involving OCDS were very similar when HC and BD were removed from the sample (GABA $r = -0.36$, glutamate $r = -0.20$; past 2 week drinkers: GABA $r = -0.30$, glutamate $r = -0.48$). In sum, these results demonstrate that BD+AD have abnormally low levels of dACC GABA and glutamate, and that low levels of GABA and glutamate are associated with assessments of salient neurobehavioral constructs (that is, impulsivity and craving) in BD+AD.

**DISCUSSION**

The present study demonstrated that individuals with co-occurring BD and current AD had uniquely low levels of prefrontal GABA and glutamate, and that low levels of GABA and glutamate were associated with both impulsivity and craving. These effects were particularly strong in individuals who drank within 2 weeks of the $^{1}$H-MRS scan. Despite different $^{1}$H-MRS methods, brain areas, and populations, extant studies have consistently demonstrated abnormally low levels of prefrontal glutamate in AD alone individuals$^{21-24}$ not experiencing acute alcohol withdrawal. Although studies of BD alone have reported elevated prefrontal glutamate levels,$^{38-44}$ $^{1}$H-MRS research on co-occurring BD and AD (that is ref. 51 and the present study) suggests that when BD and AD are combined within the same individual, the effects of AD may modify the effects of BD on prefrontal glutamate levels leading to a net decrease in glutamate to levels even less than in AD alone. Though evidence for prefrontal GABA abnormalities in $^{1}$H-MRS studies of AD$^{23,28,29}$ or BD$^{45-50}$ alone has been inconsistent, our data suggest that individuals with co-occurring BD and AD have uniquely low levels of prefrontal GABA.

Although there are several points of convergence between our findings and the extant literature, there are also notable points of divergence. For example, although the extant literature has demonstrated increased prefrontal glutamate in BD alone, and decreased prefrontal glutamate in AD alone not experiencing acute alcohol withdrawal, relative to healthy controls, individuals with BD or AD alone in the present study did not have statistically different prefrontal glutamate levels than controls. One potential explanation for the lack of differences between AD alone and controls could be that prefrontal glutamate levels may have normalized during the $\geq$ 1-week monitored period of abstinence that preceded the $^{1}$H-MRS scan$^{23,25,77}$ only 35–40% of AD alone and co-occurring BD and AD participants reported drinking in the 2 weeks preceding $^{1}$H-MRS (that is, in the week preceding the monitored 1-week abstinence period). Another potential explanation is that we carefully excluded AD with potentially confounding clinical characteristics, including history of recurrent depression or complicated alcohol withdrawal including seizures, current post-traumatic stress disorder or obsessive-compulsive disorder or prescription of psychotropic medications. Similar potential explanations could be invoked concerning the lack of observed
difference in prefrontal glutamate between BD alone and controls. For example, unlike most prior BD 1H-MRS studies, we carefully excluded individuals with any history of SUD as well as any alcohol/drug consumption in the week preceding MRI. Furthermore, our BD, and co-occurring BD and AD, participants were not severely depressed or manic, and most BD and co-occurring BD and AD participants were taking mood-stabilizing medications (that is, lithium, anticonvulsant medications, atypical antipsychotic medications), which may have reduced potential effect sizes of BD on glutamate levels. The present investigation represents a foundation for future studies to follow-up on these potential explanations by, for example, scanning participants without a preceding period of abstinence or focusing on unmedicated BD individuals.

If replicated, the findings from the present study may support preferential evaluation of therapeutics known to increase prefrontal GABA and glutamate in co-occurring BD and AD. Traditionally, treatment trials for co-occurring BD and AD have investigated medications FDA-approved to treat either BD or AD in hopes that such medications would prove efficacious in individuals with both disorders. These hopes have largely not been realized. A different approach to selecting and developing medications for clinical trials would be to target neurobiological dysfunctions characteristic of individuals with both BD and AD. Our findings not only suggest that individuals with co-occurring BD and AD have unique deficits in prefrontal GABA and glutamate, but that these deficits may be associated with salient neurobehavioral constructs including impulsivity and alcohol craving. Future studies can expand this research further by evaluating more objective/behavioral measures of impulsivity and craving and by including 1H-MRS imaging in clinical trials with specific glutamate/GABA-acting medications.

Though our findings may provide potential avenues for treatment development in individuals with co-occurring BD and AD, a mechanistic understanding of these findings is, at present, limited. The glutamate–GABA cycle is complex, involving several important enzymes (glutamic acid decarboxylase, glutamine synthetase, phosphatase-activated glutaminase) as well as transporters for glutamate, glutamine and GABA selectively localized in astrocytes and glutamatergic/GABAergic neurons, respectively. Further, overall homeostasis of glutamate–GABA biosynthetic coupling is regulated by more than one mechanism simultaneously; for instance, upon uptake of synthetically released glutamate into astrocytes, glutamine may be converted to glutamine either by glutamine synthetase or by oxidative metabolism, and the extent to which these pathways are engaged appears to depend in part upon glutamatergic tone. Although not the main focus of this investigation, we found no significant differences in glutamine levels between study groups. How the myriad enzymes and transporters involved in the glutamate–GABA cycle are regulated by chronic alcohol use or duration of abstinence is poorly understood in humans, much less in those with both BD and AD. Last, 1H-MRS cannot at present distinguish neuronal vs glial glutamate neurotransmitter pools even under normal physiological conditions in healthy subjects. Nonetheless, each of these macromolecular sites in the tripartite synapse represent testable potential therapeutic targets that may be better characterized in the future using selective pharmacologic agents and improving MRS technology (for example, carbon-13 MRS).

The results of the present study should be interpreted in light of several limitations. First, a number of potentially confounding clinical characteristics (for example, concomitant medications, co-occurring drug dependence) were not exclusionary. Allowing these potential sources of variability into the study was by design, to balance feasibility of recruitment/retention and scientific concerns. Striking this balance is particularly important in research involving co-occurring BD and AD, a notoriously difficult to recruit/retain population for research. Though imperfect, we used a statistical approach to evaluating and controlling for the potential impact of these characteristics on glutamate and GABA levels and we found no evidence that such factors were independently responsible for our findings; sample sizes were too small to evaluate potential interactive effects. Second, our findings demonstrated that prefrontal GABA and glutamate differences between co-occurring BD and AD and all other participant groups were particularly strong in participants who drank within 2 weeks of MRI. Given that only 40% of participants reported drinking within 2 weeks of MRI, these findings were based on small samples (that is, n = 28–31), and should be considered tentative until replicated. This concern is particularly salient for findings involving glutamate, as findings involving GABA were significant in both the full and restricted (recent drinking) samples. Third, because we excluded participants with a history of delirium tremens or >1 lifetime alcohol withdrawal seizure, it is unclear whether our findings generalize to individuals with very severe AD. Fourth, the ProFit algorithm used to analyze our 2D J-PRESS 1H-MRS data did not allow for separation of co-edited macromolecules (for example, proteins) from the observed GABA signal. Although it is very unlikely that between-group differences in macromolecules represent a viable alternative explanation for our GABA findings, future studies should evaluate macromolecule-suppressed GABA acquisitions in co-occurring BD and AD. Finally, given that we did not acquire data from brain regions other than dACC, we cannot comment on the regional specificity of our findings.

These limitations notwithstanding, the present study extends the literature by demonstrating uniquely low levels of prefrontal GABA and glutamate in individuals with co-occurring BD and current AD, along with associations between prefrontal GABA and glutamate levels and impulsivity and alcohol craving. If replicated, these findings support preferential evaluation of therapeutics known to ‘normalize’ prefrontal GABA and glutamate levels in co-occurring BD and AD.

CONFLICT OF INTEREST

PFR consults for Kyowa Hallo and Tai Medical. RFA has been a consultant in the recent past for Lilly, Lundbeck, Novartis, Indivior and Alkermes, and on Advisory Boards for Alkermes, Indivior and Lundbeck. He has received grant funds from Lilly. He is a member of the Alcohol Clinical Trials Initiative (ACTIVE) sponsored by the American Society of Clinical Psychopharmacology (ASCP) which has been supported in the last 3 years by AbbVie, Alkermes, Ethypharm, Indivior, Lilly, Lundbeck, Otsuka, Pfizer and XenoPort. The remaining authors declare no conflict of interest.

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