Ketamine Modulates Hippocampal Neurochemistry and Functional Connectivity – A Combined Magnetic Resonance Spectroscopy and Resting State fMRI Study in Healthy Volunteers

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Abstract

A growing body of evidence suggests glutamate excess in schizophrenia and that N-methyl-d-aspartate receptor (NMDAR) hypofunction on γ-aminobutyric acid (GABA) interneurons disinhibiting pyramidal cells may be relevant to this hyperglutamatergic state. To better understand how NMDAR hypofunction affects the brain, we used Magnetic Resonance Spectroscopy and resting state functional MRI to study the effects of ketamine on hippocampal neurometabolite levels and functional connectivity in 15 healthy human subjects. We observed a ketamine induced increase of hippocampal Glx (glutamate+glutamine; F= 3.76; p= 0.04), a decrease in fronto-temporal (t=4.92, pFDR< .05, kE= 2198, x= -30, y= 52, z= 14) and temporo-parietal functional connectivity (t=5.07, pFDR< .05, kE= 6094, x= -28, y= -36, z= -2), and a possible link between connectivity changes and elevated Glx. Our data empirically support that hippocampal glutamatergic elevation and resting state network alterations may arise from NMDAR hypofunction and establish a proof of principle whereby experimental modelling of a disorder can help mechanistically integrate distinct neuroimaging abnormalities in schizophrenia.

Keywords

schizophrenia; Glx (glutamate+glutamine); N-methyl-d-aspartate receptor (NMDAR); GABA; psychosis; fronto-temporal; temporo-parietal
Although the dopamine model of schizophrenia remains heuristically valuable, this phenotypically complex illness cannot be explained by dopaminergic dysfunction alone. A growing body of evidence suggests glutamate excess in schizophrenia\(^1\)-\(^3\) and that N-methyl-D-aspartate receptor (NMDAR) hypofunction on \(\gamma\)-aminobutyric acid (GABA) interneurons disinhibiting pyramidal cells may be relevant to this hyperglutamatergic state\(^4\)-\(^6\).

Due to its unique composition, with approximately 90% of its neurons being pyramidal cells, the hippocampus is a region that may be especially vulnerable to shifts in the excitation/inhibition balance secondary to NMDAR hypofunction as opposed to the cerebral cortex where the number of pyramidal neurons and interneurons is more similar\(^7\). Post-mortem and neuroimaging studies suggest extensive hippocampal pathology in schizophrenia\(^7\)-\(^11\). We recently reported evidence of glutamatergic excess in the hippocampus in unmedicated patients\(^1\) using Magnetic Resonance Spectroscopy (MRS), a technique that allows neurometabolite quantification \(in\ \text{vivo}\). Glutamatergic hyperactivity has been shown to result in disorganized neuronal activity in animal models\(^4\). Resting state functional Magnetic Resonance Imaging (fMRI) allows approximation of neuronal activity by measuring the temporal covariation of low frequency fluctuations of the blood oxygen level dependent (BOLD) signal\(^12\). In schizophrenia, aberrant hippocampal resting state connectivity to frontal areas including the anterior cingulate cortex (ACC), medial prefrontal cortex (MPFC), parietal areas including the posterior cingulate cortex (PCC) and precuneus, as well as the thalamus\(^13\),\(^14\) and striatum\(^15\)-\(^18\) suggests disorganized neuronal activity, possibly related to NMDAR hypofunction\(^15\). Unfortunately, even contemporary imaging techniques cannot readily quantify NMDAR function \(in\ \text{vivo}\). Alternatively, experimental manipulation of the NMDAR could provide proof of principal and advance our mechanistic understanding of the illness.

Subanesthetic doses of ketamine preferentially block NMDAR on GABAergic interneurons\(^19\)-\(^21\) and have been shown to transiently induce a behavioral phenotype similar to that seen in schizophrenia\(^22\)-\(^26\), without long term adverse effects\(^27\). However, previous neuroimaging studies have not mapped ketamine’s effect on hippocampal glutamate and functional connectivity in the same population, which could inform a mechanistically based and clinically relevant model integrating different patterns of brain abnormalities. To programmatically build upon our recent studies in schizophrenia\(^1\),\(^10\),\(^15\),\(^16\),\(^28\),\(^29\), we enrolled a group of healthy volunteers to contrast (1) hippocampal Glx (glutamate +glutamine) using MRS and (2) hippocampal connectivity using resting state fMRI during a saline infusion and during a ketamine challenge. We hypothesized that experimentally induced NMDAR blockage would result in an increase of hippocampal Glx and in fronto-temporal and temporo-parietal functional dysconnectivity, resembling abnormalities seen in the disorder. Additionally, we conducted exploratory analyses investigating relationships between Glx, functional connectivity, and psychosis severity.
METHODS

Subjects

We recruited 19 healthy volunteers who gave written informed consent for this Institutional Review Board approved study. Exclusion criteria were a history of an Axis I disorder or a psychotic disorder in a first-degree family member, major medical or neurological conditions, lifetime use of psychotropic medications, prior exposure to ketamine, and pregnancy or breastfeeding.

Subjects meeting eligibility criteria during a phone screen were invited to complete a Diagnostic Interview for Genetic Studies and a psychiatric assessment and physical exam conducted by a board certified psychiatrist (NVK). Urine drug screens and, if applicable, pregnancy tests were completed during the screen and before each ketamine infusion. The Hamilton Rating Scale for Depression (HRSD)\(^{30}\) and Young Mania Rating Scale (YMRS)\(^{31}\) were used to assess mood symptoms before each infusion. The Clinician Administered Dissociative States Scale (CADSS)\(^{32}\) and the Brief Psychiatric Rating Scale (BPRS)\(^{33}\) were completed before and after the ketamine challenge (subjects were asked to retrospectively report symptoms at the time drug effects were the most prominent).

To ensure drug tolerability and minimize novelty, subjects received an intravenous racemic ketamine challenge (0.27mg/kg bolus over 10 minutes, followed by a continuous infusion of 0.25mg/kg/hour for 50 minutes) in the Clinical Research Unit at least one week prior to imaging. Ten milliliters of blood were collected immediately after completion of the bolus and 50 minutes after start of the challenge during the drug tolerability assessment rather than scanning to avoid potential delays in our scanning timeline due to complications during the blood draws. Blood samples were centrifuged to obtain plasma and stored at -40°C. Ketamine plasma levels were assayed (Nathan Klein Institute) using a liquid chromatographic procedure. During the ketamine challenge, vital signs including heart rate, blood pressure, peripheral oxygen saturation, and respiratory rate (CO\(_2\) monitoring during scanning) were monitored by an anaesthesiology fellow under supervision of a board certified anaesthesiologist according to the standards for basic anaesthetic monitoring. Monitoring was continued for one hour after infusion completion. Prior to discharge into the care of an accompanying driver, subjects were medically cleared by the fellow and psychiatrist.

Two subjects withdrew from the study because they developed emesis, and two subjects revoked consent after the initial ketamine challenge, citing time constraints as their reason (both denied adverse drug effects); 15 subjects completed imaging.

Following anatomical, spectroscopy, and resting state fMRI scans during a saline infusion (flow rate of 0.01ml/s), subjects were given a short break. After repeat anatomical scans, subjects received a ketamine challenge consisting of a bolus (0.27mg/kg over 10 minutes), followed by a continuous infusion (0.25mg/kg/hour, flow rate of 0.01ml/s). Spectroscopy acquisition was started approximately 13 minutes after start of the challenge (10 minutes for bolus administration and 3 minutes for shimming, this also allowed for stabilization of ketamine plasma levels) and lasted for 21.3 minutes and followed by resting state scans that
were started approximately 45 minutes after the start of the challenge and lasted for 7.5 minutes.

**Image acquisition**

Imaging was performed on a 3T head-only scanner (Magnetom Allegra, Siemens). A high-resolution structural scan was acquired (MPRAGE, TR/TE/TI=2300/3.93/1100msec, 1mm isotropic voxels).

A series of T1-weighted anatomical scans (TR/TE= 250/3.48ms, 5mm slice thickness) were acquired to aid placement of the left hippocampus spectroscopy voxel (2.7x1.5x1cm; Figure 1). Voxel placement for the second acquisition was guided by an image of the voxel placement during the first scan. After manual shimming, CHESS pulses were used to suppress the water signal. Spectra were acquired using a PRESS sequence (TR/TE= 2000/80ms to optimize the glutamate signal and minimize macromolecule contribution; 1200 Hz spectral bandwidth; 1024 points; 640 averages, and 8 averages without water suppression).

Resting state fMRI scans were acquired during a gradient recalled echo-planar imaging sequence (TR/TE= 2000/30msec, flip angle= 70°, field of view= 192x192mm$^2$, 64x64 matrix, 6mm slice thickness, 1mm gap, 30 axial slices, 225 acquisitions). Subjects were instructed to keep their eyes open and stare passively ahead. No subject reported falling asleep during the resting state scan.

**MRS data processing**

MRS data were quantified in the time domain with AMARES in jMRUI (version 5.2); prior knowledge derived from in vitro and in vivo metabolite spectra was included in the model, as reported elsewhere. Exclusion criteria were (1) line width of the magnitude signal during manual shimming >20 Hz at FWHM and (2) Cramer-Rao lower bounds (CRLB) >20%. No spectra were excluded. Metabolites were quantified with respect to creatine (Cr/ internal water (W) did not differ between conditions; F= 0.517; p=.48); for the sake of brevity they will be referred to as Glx, NAA (N-acetyl-aspartate+ N-acetylaspartylglutamate), and Cho (glycerophosphocholine+ phosphocholine) hereafter. Structural scans were segmented into grey matter (GM), white matter (WM), and cerebrospinal fluid to calculate voxel tissue fractions, using in house Matlab codes.

**Resting state data processing**

After removing the first five frames from each run to allow for signal equilibration, data were slice timing corrected, realigned, co-registered, normalized, and spatially smoothed with a 6mm Gaussian kernel using the conn toolbox in SPM8. Physiological and other spurious sources of noise were estimated with aCompcor, and the first five principal temporal components each from white matter and CSF were removed. No global signal regression was performed. Frames with movement >0.5mm or global signal z score change >3 were identified with artifact detection (ART), scrubbed, and excluded from analyses. Residual data were band-pass filtered (0.009-0.08 Hz). The entire time series was excluded if >50% of frames were contaminated by movement, or if mean framewise displacement
exceeded 0.5 mm. No scans met these criteria. We observed no differences in mean framewise displacement (saline: 0.18 +/- 0.06 mm; ketamine: 0.20 +/- 0.08 mm; t = -1.17; p = 0.26), or mean proportion of frames scrubbed (saline: 6.71 +/- 5.46%; ketamine: 8.31 +/- 7.71%; t = -0.70; p = 0.50) across time points. The first eigenvariate of the BOLD time series from a left hippocampus seed region, defined by the AAL atlas, was extracted and correlated to the time series of all other voxels to create seed-to-voxel correlation maps for each subject.

**Statistical Analyses**

To investigate change in metabolites between saline and ketamine infusions, we used a mixed repeated measures design with neurometabolites as dependent variables, experimental condition (saline vs ketamine) as fixed factor, and voxel GM fraction (GM% / GM% + WM %) as covariate. Given directionality in hypothesis for changes in Glx, we conducted a one-sided test; changes in other metabolites were examined with two-sided tests. Results were considered statistically significant at p < .05. In an exploratory fashion, we also examined relationships between Glx and clinical observations using Pearson’s correlations.

To investigate change in functional connectivity between saline and ketamine infusions, we did paired-sample t-tests on the subject level connectivity maps. Analyses were limited (masked) to include only regions that were functionally connected to the hippocampus in at least one of the conditions (saline/ ketamine). These masks were computed by thresholding ($P_{FDR} < .05$), binarizing, and then combining each within-group analysis (both conditions) producing masks of the combined connected areas. Analyses were corrected for multiple comparisons using the false discovery rate (FDR) and are reported at $P_{FDR} < 0.05$. To assess the relationship between resting state connectivity changes and Glx, we created connectivity change maps using the ImCalc function in SPM8, and extracted the first eigenvariate for each cluster that showed decreased connectivity during the ketamine challenge, which was then correlated with Glx during the ketamine challenge.

In an exploratory fashion, we used regression analyses to examine relationships between connectivity during the saline infusion and ketamine induced severity of psychosis (to assess to what extent baseline functional connectivity may be related to the brain’s propensity to develop psychosis) and between connectivity during the ketamine challenge and ketamine induced severity of psychosis, defined as BPRS total score. To explore if any associations unique to symptom dimension exist, we then performed regression analysis using both BPRS positive and negative symptom scores as regressors in the same analysis.

**RESULTS**

**Demographics, clinical observations, and laboratory results**

Fifteen subjects (10 male/ 5 female) aged 24.80 +/- 3.49 years completed scanning. One subject smoked 3 cigarettes per day; all others denied smoking. None had HDRS or YMRS scores in the clinical range. BPRS and CADSS scores were significantly higher during the ketamine challenge compared to the saline infusion (Table 1). Ketamine plasma levels were
74.27 +/- 22.08ng/ml and 97.47 +/- 19.59ng/ml immediately after completion of the bolus and 50 minutes after start of infusion, respectively.

**MRS measurements**

We found a significant increase in Glx during the ketamine challenge compared to the saline infusion, Glx/W increased at trend level (saline: 0.18, ketamine: 0.21; F 1.977 p= .09); NAA and Cho did not differ between measurements (Table 2, Figure 1). We observed no correlations between BPRS or CADSS scores and Glx at endpoint or percent change in Glx between saline and ketamine infusions (all p > .05).

**Resting state functional connectivity**

Left hippocampus functional connectivity patterns during the saline infusion were consistent with prior reports in healthy subjects (supplemental Figure). During the ketamine challenge, we found two clusters of decreased connectivity, a frontal cluster spanning across the ACC, MPFC, and middle cingulate (t=4.92, k_E [cluster extent]= 2198, x= -30, y= 52, z= 14) and a largely tempo-parietal cluster spanning from the bilateral hippocampus to the right insula, bilateral precuneus, PCC, lingual gyrus, and calcarine sulcus (t=5.07, k_E= 6094, x= -28, y= -36, z= -2). No areas showed increased hippocampus connectivity during the ketamine challenge. We observed a significant negative correlation between Glx during the ketamine challenge and hippocampus connectivity change with ketamine in the frontal cluster (r= 0.52; p<.05) and a trend-level negative correlation between Glx during the ketamine challenge and hippocampus connectivity change with ketamine in the temporo-parietal cluster (r= 0.51; p= .05; Figure 2). No correlations between connectivity change and change in Glx were observed.

BPRS total score during the ketamine challenge was negatively correlated with hippocampus connectivity during saline infusion to a cluster spanning across the ACC, MPFC, insula and rolandic operculum (t=5.68, k_E= 2626, x= -6, y= 44, z= -18) and to a cluster spanning across the temporal lobe and cerebellum (t=5.50, k_E= 4053, x= -66, y= -38, z= -22). BPRS total scores during the ketamine challenge were negatively correlated with hippocampus connectivity during the ketamine challenge to a cluster spanning across the superior frontal gyrus, middle cingulum, PCC, and precuneus (t=7.06, k_E= 5916, x= 4, y= 10, z= 42) and to a cluster spanning across the bilateral parahippocampus, superior/ inferior parietal cortices, superior and middle occipital cortices, and lingual gyrus (t=5.07, k_E= 6094, x= -28, y= -36, z= -2; Figure 3). None of the associations were unique to a specific symptom dimension.

**DISCUSSION**

To our knowledge, this is the first study to date combining MRS and resting state fMRI to probe the effects of experimentally induced NMDAR hypofunction in vivo in healthy human subjects, and the first to report elevated subcortical glutamatergic levels after ketamine administration. We observed a ketamine induced increase of hippocampal Glx, a decrease in fronto-temporal and temporo-parietal functional connectivity, and a possible link between connectivity changes and elevated Glx. It is important to note that alterations closely resemble hippocampal neurometabolic and network-level abnormalities we have previously...
reported in unmedicated patients with schizophrenia\textsuperscript{1, 15}. Our experiment adds to the efforts in bridging the gap between contemporary mechanistic models and findings from \textit{in vivo} neuroimaging studies in schizophrenia, suggesting that hippocampal glutamate excess and functional network disturbances may reflect a disrupted excitation/inhibition balance.

In agreement with this, prior studies have reported elevated hippocampal rCBF in unmedicated patients\textsuperscript{41} and elevated hippocampal blood volume in medicated patients\textsuperscript{42}. Linking cellular-level mechanisms and neuroimaging findings, Schobel and colleagues reported a series of experiments in a mouse model of schizophrenia, showing that ketamine administration causes an increase in extracellular glutamate, hippocampal hypermetabolism, and atrophy. In parallel, they identified a relationship between hippocampal hypermetabolism and structural deficits in patients transitioning from a prodromal state to syndromal psychosis\textsuperscript{43}. The authors concluded that glutamate acts as a pathogenic driver of hippocampal pathology.

Because neuronal composition is variable across the brain, it is conceivable that disparate regions express differential vulnerability to disease mechanisms. Probing sensitivity to ketamine induced NMDAR hypofunction, two prior studies reported increases of glutamatergic indices in the ACC\textsuperscript{44, 45}, but others found no change in the ACC\textsuperscript{46} and occipital cortex\textsuperscript{47}. Like us, Rowland and colleagues reported no relationship between glutamatergic indices and BPRS or CADSS scores, which is consistent with many studies in schizophrenia that fail to establish a direct link between glutamate excess and symptom severity\textsuperscript{48}. While it is possible that these indices reflect a more general marker of psychopathology that is not readily captured by behavioral scales, an alternative interpretation is that clinical symptoms may emerge from disrupted network-level functional connectivity patterns related to glutamate excess. In other words, it is possible that glutamate acts as a moderating variable between resting state connectivity and behavioral symptoms.

Ketamine induced NMDAR hypofunction on GABA interneurons has been shown to disrupt gamma oscillations\textsuperscript{49}, which are thought to regulate synchronized brain activity\textsuperscript{50}, and in turn could also affect resting state connectivity. Several studies have reported ketamine induced connectivity changes, including increased whole brain global connectivity\textsuperscript{51}, increased cortico-thalamic connectivity\textsuperscript{52}, decreased default mode network connectivity\textsuperscript{53}, and mixed increases and decreases among eight large scale networks\textsuperscript{54}. Examining prefrontal-hippocampal connectivity using region-of-interest masks in the dorsolateral prefrontal cortices and bilateral hippocampi, Grimm and colleagues reported hyperconnectivity after a ketamine challenge\textsuperscript{55}. While this may appear to be inconsistent with our finding of a decrease in hippocampus connectivity, we may have simply captured a different window in the temporal dynamic\textsuperscript{56} of ketamine induced network changes (we scanned during the ketamine administration, whereas Grimm scanned 20 minutes after completion of the challenge). Preclinical models suggest a dose dependent ketamine induced extracellular glutamate increase of up to 200-300% of baseline levels within 20 minutes of administration with the effect subsiding between 60 and 100 minutes of administration\textsuperscript{57}. Similarly, an MRS study examining ketamine effects in the MPFC suggests a 17+-25% increase of Glx within 30 minutes of administration that subsided within one hour\textsuperscript{58}, which is consistent findings of a \textsuperscript{13}C-MRS study reporting an increase of \textsuperscript{3}C enrichment of...
glutamate-C4 of approximately 15%\textsuperscript{59}, and similar to our report of a mean increase of 15.9±28% of hippocampal Glx.

Ketamine induced decreases in cortico-hippocampal functional connectivity spanning to the ACC, MPFC, insula, precuneus, and lingual gyrus reported here recapitulate preclinical findings of NMDAR hypofunction related hippocampal connectivity changes to homologous areas of the rat brain\textsuperscript{60}. In parallel, ketamine modulates cerebral blood flow (rCBF) in the hippocampus, ACC, MPFC, and lingual gyrus in both healthy controls and volunteers with schizophrenia\textsuperscript{41, 42}, again implicating vulnerability to NMDAR blockage in this functional circuit.

It is intriguing that ketamine induces cortico-hippocampal circuitry alterations that are similar to those seen in schizophrenia\textsuperscript{15-18} but does not replicate hippocampal connectivity abnormalities to the thalamus and to the striatum\textsuperscript{13-15, 61}, an area rich in dopaminergic innervation. This possibly reflects a dissociation of glutamate and dopamine system abnormalities, which is in agreement with prior studies failing to reverse ketamine induced psychosis with haloperidol, a dopamine D\textsubscript{2} receptor antagonist, in healthy subjects and volunteers with schizophrenia\textsuperscript{22, 62}. Alternatively, it may be related to the ketamine dose\textsuperscript{63} or to the statistical threshold set for image analyses, since fronto-striatal-thalamic connectivity has been shown to increase with ketamine\textsuperscript{64}, and preclinical models have established a link between hippocampal glutamate and dopamine firing in the ventral tegmental area\textsuperscript{65}. Taken together, findings argue against a simplistic interpretation of ketamine as a schizophrenia model but rather for a model of a hyperglutamatergic stage\textsuperscript{66} or subtype of the illness. A recent report of elevated cortical glutamate (though no data exists for the hippocampus) in patients with treatment-resistant schizophrenia compared to treatment-responsive patients\textsuperscript{67} underscores the value of this pharmacological neuroimaging phenotype as translational target for novel drug development.

Interestingly, lower hippocampus connectivity during saline infusion to the ACC and insula, both regions part of a network responding to behaviourally salient events and reported abnormal in schizophrenia\textsuperscript{68, 69}, was related to higher symptom severity during the ketamine challenge. Putting this in context of NMDAR blockage induced reductions in connectivity, it is tempting to interpret findings akin to a cognitive reserve model\textsuperscript{70}, where greater hippocampal functional connectivity during saline infusion to the salience network may provide a higher threshold for clinical symptoms to emerge and thus reflect the brain’s resilience to developing psychosis. The possibility of leveraging neuroimaging data to predict behaviour, disease vulnerability, or propensity to treatment response is of great interest to the field\textsuperscript{71}. For example, the predictive value of imaging biomarkers for antipsychotic response in schizophrenia is beginning to be explored\textsuperscript{68, 72}. As this area of research matures, we can hope to be able to derive illness models that can bridge mechanisms of brain function and pathology at the macroscopic scale to disease vulnerability and behavioral phenotypes.

Several strengths and limitations need to be considered. To be able to compare spectroscopy and resting state data with our previous studies, imaging and data analysis parameters replicated those we had implemented previously (except for more stringent movement...
correction of resting state data here). We did not include a placebo control. Because of ketamine’s unique side effect profile, including dissociative effects and nystagmus, the risk of “unblinding” is substantial. We acknowledge that it would be ideal to include a study arm with an active placebo that could mitigate these concerns, but any drug producing similar side effects could also confound neurometabolite measurements and resting state connectivity, making interpretation of findings difficult. Additionally, a counterbalance design and randomization of subjects to the different conditions could have better controlled for anticipation and expectancy effects. We reported a significant increase in Glx using a one sided t-test given the directionality of our hypothesis based on our report in schizophrenia, but we were underpowered to detect significant abnormalities in Glx/W and in Glx assuming the directionality of the change had been unknown a priori. Also, MRS data was collected only once, which precludes us to directly comment on the temporal dynamics of hippocampal Glx changes.

In summary, our data empirically support that hippocampal glutamatergic elevation and resting state network alterations may arise from NMDAR hypofunction and establish a proof of principle whereby experimental modelling of a disorder can help mechanistically integrate distinct neuroimaging abnormalities in schizophrenia. The results of this study also have potential clinical implications. Medications attenuating the impact of hippocampal glutamatergic hyperactivity could conceivably diminish functional dysconnectivity and perhaps alleviate disease burden across symptom dimensions in this profoundly disabling neuropsychiatric disorder. Future studies investigating the utility of these translational biomarkers to assess target engagement, circuit-relevant pharmacodynamic activity, and efficacy of novel pharmacotherapies will be important to inform this question.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

Acknowledgments

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Figure 1.
(A) Example of magnetic resonance spectroscopy (MRS) voxel placement in the left hippocampus (2.7×1.5×1 cm). The image is displayed in radiological convention (right side of image is subject’s left side). (B+C) Example spectra of the same subject during (B) saline infusion and (C) ketamine infusion. The black line is a spectrum (640 averages) obtained from the left hippocampus voxel, the red line is an overlay of spectral fit. (D) Hippocampal Glx during saline and ketamine infusion (n=15). Squares represent individual measurements. Lines connect saline and ketamine measurements of individual subjects. Boxplots indicated that Glx measures during saline the infusion for three subjects were outliers (no outliers detected during ketamine challenge). Repeating analyses excluding these subjects did not alter results, Glx remained significantly higher during ketamine challenge compared to saline infusion (n=12; F= 9.408; p< 0.01). Grey indicates that Glx value during saline infusion was an outlier. Cho, choline; Cr, creatine; Glx, glutamate + glutamine; NAA, N-acetyl-aspartate; ppm, parts per million.
Figure 2.
Left: Areas of decreased functional connectivity during a ketamine challenge compared to a saline infusion (p_{FDR} < .05). No areas of increased connectivity during the ketamine challenge were detected. Clusters were overlaid on the Xjview single subject T1 template. Numbers indicate MNI coordinates. Color bar indicates t values. Right: Correlations between ketamine induced connectivity change (first eigenvariate) and Glx during a ketamine challenge. ACC: anterior cingulate cortex, MPFC: medial prefrontal cortex, Glx: glutamate+glutamine, PCC: posterior cingulate cortex.
Figure 3.
Resting state functional connectivity in relation to symptom severity. Top row: Functional connectivity during a saline infusion in relationship to Brief Psychiatric Rating Scale (BPRS) total scores during a ketamine challenge. Clusters indicate regions where connectivity was negatively correlated with symptom severity ($p_{FDR} < .05$). No clusters were detected that showed positive correlations between connectivity and symptom severity. Bottom row: Functional connectivity during a ketamine challenge in relationship to BPRS total scores during the ketamine challenge. Clusters indicate regions where connectivity was negatively correlated with symptom severity ($p_{FDR} < .05$). No clusters were detected that showed positive correlations between connectivity and symptom severity. Clusters were overlaid on the Xjview single subject T1 template. Numbers indicate MNI coordinates. Color bar indicates $t$ values. ACC: anterior cingulate cortex, MPFC: medial prefrontal cortex.
|                    | Saline     | Ketamine   | t-score  | p value |
|--------------------|------------|------------|----------|---------|
| **BPRS**           |            |            |          |         |
| Total              | 20.60 (0.74) | 32.73 (4.94) | -9.742   | < .01   |
| Positive           | 3.00 (0.00)  | 5.87 (1.69)  | -6.590   | < .01   |
| Negative           | 3.13 (0.35)  | 6.87 (1.96)  | -7.047   | < .01   |
| **CADSS**          |            |            |          |         |
| Total score        | 0.07 (0.02)  | 13.60 (6.50) | -8.049   | < .01   |
| Amnesia            | 0.07 (0.02)  | 2.07 (1.71)  | -4.472   | < .01   |
| Derealization      | 0.00 (0.00)  | 7.27 (3.92)  | -7.183   | < .01   |
| Depersonalization  | 0.00 (0.00)  | 3.47 (2.10)  | -6.394   | < .01   |
| Confusion          | 0.00 (0.00)  | 0.13 (3.52)  | -1.468   | 0.16    |
| **HRSD**           | 0.20 (0.40)  |            |          |         |
| **YMRS**           | 0.20 (0.56)  |            |          |         |

Abbreviations: BPRS Brief Psychiatric Rating Scale; CADSS Clinician Administered Dissociative States Scale; HRSD Hamilton Rating Scale for Depression; YMRS Young Mania Rating Scale

1 Mean (SD) unless indicated otherwise; n= 15
2 Brief Psychiatric Rating Scale (1 – 7 scale); positive (conceptual disorganization, hallucinatory behavior, and unusual thought content); negative (emotional withdrawal, motor retardation, and blunted affect)
Table 2

Neurometabolites and spectral quality indices $^1$

|                          | Saline            | Ketamine         | t/F   | p value |
|--------------------------|-------------------|------------------|-------|---------|
| **Neurometabolite measures** |                  |                  |       |         |
| Glx                      | 0.62 (0.13)       | 0.69 (0.08)      | 3.756 | 0.04$^2$|
| NAA                      | 1.35 (0.10)       | 1.38 (0.16)      | 0.346 | 0.57    |
| Cho                      | 0.94 (0.12)       | 0.96 (0.16)      | 0.151 | 0.70    |
| **Spectral quality indices** |                  |                  |       |         |
| Glx CRLB                 | 0.09 (0.02)       | 0.10 (0.02)      | -2.182| 0.05    |
| NAA CRLB                 | 0.04 (0.01)       | 0.04 (0.01)      | -1.799| 0.09    |
| Cho CRLB                 | 0.03 (0.01)       | 0.03 (0.01)      | -1.713| 0.11    |
| FWHM                     | 7.42 (1.32)       | 7.63 (1.46)      | -0.694| 0.50    |
| SNR                      | 12.57 (1.76)      | 11.89 (1.85)     | 1.260 | 0.23    |
| **Voxel tissue fraction** |                  |                  |       |         |
| grey matter (%)          | 63.67 (4.64)      | 63.11 (6.00)     | 0.527 | 0.61    |
| white matter (%)         | 33.64 (4.81)      | 34.42 (6.45)     | -0.701| 0.50    |
| cerebrospinal fluid (%)  | 2.68 (0.14)       | 2.47 (0.16)      | 1.009 | 0.33    |

Abbreviations: Cho Choline; CRLB, Cramer Rao Lower Bounds; FWHM Full width at half maximum; Glx glutamate+glutamine; NAA N-acetyl-aspartate; SNR Signal to noise ratio

$^1$ Mean (SD) unless indicated otherwise, n= 15

$^2$ one-sided F test