Molecular imaging offers unprecedented opportunities for investigating dynamic changes underlying neuropsychiatric conditions. Here, we evaluated whether \[^{11}C\]ABP688, a positron emission tomography (PET) ligand that binds to the allosteric site of the metabotropic glutamate receptor type 5 (mGluR5), is sensitive to glutamate fluctuations after a pharmacological challenge. For this, we used ceftriaxone (CEF) administration in rats, an activator of the GLT-1 transporter (EAAT2), which is known to decrease extracellular levels of glutamate. MicroPET \[^{11}C\]ABP688 dynamic acquisitions were conducted in rats after a venous injection of either saline (baseline) or CEF 200 mg/kg (challenge). Binding potentials (BP\textsubscript{ND}) were obtained using the simplified reference tissue method. Between-condition parametric maps indicating brain regions showing the highest CEF effects guided placement of microdialysis probes for subsequent assessment of extracellular levels of glutamate. The CEF administration increased \[^{11}C\]ABP688 BP\textsubscript{ND} in the thalamic ventral anterior (VA) nucleus bilaterally. Subsequent microdialysis assessment revealed declines in extracellular glutamate concentrations in the VA. The present results support the concept that availability of mGluR5 allosteric binding sites is sensitive to extracellular concentrations of glutamate. This interesting property of mGluR5 allosteric binding sites has potential applications for assessing the role of glutamate in the pathogenesis of neuropsychiatric conditions.

**Keywords:** \[^{11}C\]ABP688; ceftriaxone; glutamate; GLT-1; mGluR5; positron emission tomography

**INTRODUCTION**

Glutamate is the major excitatory neurotransmitter in the mammalian brain, acting on ionotropic (AMPAr, KA) and metabotropic (mGluRs) receptors. The ionotropic receptors produce fast-acting excitatory effects, while mGluRs (G protein-coupled receptors) have a crucial role in the modulation of glutamatergic neurotransmission.\(^1\) Since glutamate is not degraded in the extracellular compartment, two major astrocytic transporters, GLT-1 (EAAT2) and GLAST (EAAT1), remove glutamate from the synapses and provide the regulation necessary to orchestrate receptor excitability.\(^2\) The disruption of this fine-tuning mechanism causes hyperactivation of glutamatergic receptors, a well-established detrimental scenario termed excitotoxicity, which is described as a fundamental player in the pathophysiology of several neurologic and psychiatric disorders.\(^3,4\)

Due to the glutamatergic system’s involvement in a large array of neuropsychiatric conditions, techniques allowing for noninvasive *in vivo* estimation of glutamatergic neurotransmission are highly desirable.

Positron emission tomography (PET) is an imaging technique that allows for the *in vivo* measurement of biologic processes such as receptor availability or fluctuations of endogenous neurotransmitters.\(^5\) Advances in pharmacology have allowed for the development of PET radiopharmaceuticals targeting allosteric binding sites of the metabotropic glutamate receptor type 5 (mGluR5).\(^6\) Previous PET studies showed that pharmacological challenges with N-acetylcysteine, a facilitator of the cysteine-glutamate antiporter that increases the release of nonsynaptic glutamate, produced a decrease in the binding of the PET radiopharmaceutical \[^{11}C\]ABP688, an allosteric and highly selective mGluR5 antagonist, in baboons\(^7\) and in rhesus monkeys.\(^8\) In contrast, a recent study showed no effect of N-acetylcysteine on \[^{11}C\]ABP688 binding in rats.\(^9\) In fact, little is known regarding noncompetitive interactions between glutamate and the availability of mGluR5 allosteric binding sites. Although important, none of the previous studies tested whether pharmacologically induced declines of extracellular glutamate concentrations increased the availability of mGluR5 allosteric binding sites.

With this in mind, we used ceftriaxone (CEF)—a potent GLT-1 activator, which decreases extracellular levels of glutamate\(^10,11\)—to investigate whether the PET radiopharmaceutical \[^{11}C\]ABP688 is sensitive to declines in extracellular glutamate concentrations.

**MATERIALS AND METHODS**

**Animals**

Sprague-Dawley rats (n = 5, 392.10 ± 58.69 g) were scanned twice within 2 to 7 days (baseline and challenge). The scanning was conducted between
1030 h and 1330 h. Additional Sprague-Dawley rats (n = 6, 452.10 ± 68.69 g) were used for microdialysis. The microdialysis was conducted between 1030 h and 1350 h. Animals were kept in a room with controlled temperature (21°C) under a 12-hour light/12-hour dark cycle (lights on 0800 h), with ad libitum access to food and water. All procedures were performed according the Guide to the Care and Use of Experimental Animals (Ed2) of the Canadian Council on Animal Care. The protocols for PET imaging, surgery, and microdialysis were approved by the Animal Care Committee of McGill University (Montreal, QC, Canada).

Radiosynthesis, Pharmacologicals, and Imaging Procedures

\([^{11}C]ABP688\) was synthesized as described previously.\(^3\) The mean specific radioactivity for the baseline was 12.35 ± 5.75 TBq/mmol and for the challenge 12.74 ± 2.49 TBq/mmol (mean ± s.d.). The PET acquisitions were performed using a Concorde MicroPET R4 scanner (Siemens-CTI, Knoxville, TN, USA), as previously described by Elmenhorst.\(^1\) Rats were placed in a prone position with the head immobilized by both the body holder and the nose cone of the anesthesia system (2% isoflurane at 0.5 L/min oxygen flow). The brain was positioned in the center of the field of view. A 10-min transmission scan with a rotating \([^{57}Co]\) point source was acquired to correct for attenuation followed by the 60-minute emission scan. For the baseline, they received a tail vein injection of saline 30 minutes before the scan (Figure 1A). For the challenge, they received a tail vein injection of CEF 200 mg/kg 30 minutes before the scan (Figure 1B). Subsequently, \([^{11}C]ABP688\) was administered as a bolus injection (0.2 mL) into the tail vein. Respiratory rate, heart rate, and body temperature were monitored throughout the scan (Biopac Systems MP150, Goleta, CA, USA). Mean injected radioactivity was 17.02 ± 2.89 MBq for baseline and 16.62 ± 4.31 MBq for challenge (mean ± s.d.). Ceftriaxone was dissolved in saline (200 mg/kg).

Image Reconstruction, Analysis, Processing, and Estimation of Binding Parameters

Images were reconstructed using an MAP (maximum a posteriori) algorithm, normalized and corrected for scatter, dead time, and decay. Imaging analysis was conducted using minctools (http://www.bic.mni.mcgill.ca/ServicesSoftware/). Time-averaged tissue-radioactivity images were manually coregistered to a standard rat histologic template.\(^1\) The image outcome measure, nondisplaceable binding potential (BP\(_{ND}\)), was estimated using the simplified reference tissue model,\(^1\) with the cerebellum as a reference region due to its low \([^{11}C]ABP688\) binding.\(^1\) \([^{11}C]ABP688\) BP\(_{ND}\) was estimated for every dynamic scan, with resulting images convolved using a 2.4-mm Gaussian kernel. The PET images were corrected for motion by realigning each frame to a reference frame using mutual information as the cost function.\(^1\) For detailed information regarding PET methodology (including error analysis, parameter sensitivity analysis, and scan duration) and simplified reference tissue model analytical details, see Elmenhorst.\(^1\)

Microdialysis Surgery, Procedure, and Histology

Rats were anesthetized with sodium isoflurane (2% isoflurane at 0.5 L/min oxygen flow), and stereotaxically implanted with 22 gauge stainless steel guides into the left frontal cortex (FC), anteroposterior (AP): 4.20 mm, mediolateral (ML): 1.0 mm, dorsoventral (DV): 5.0 mm) and into the right thalamic ventral anterior (VA) (AP: 1.88 mm, ML: 2.0 mm, DV: 6.5 mm) following Paxinos coordinates.\(^1\) These cannulas were used to insert the microdialysis probes into the target sites. Cannulas were secured with acrylic dental cement and a single anchor screw threaded into the cranium. Carprofen (5 mg/kg, subcutaneously) was used for postoperative analgesia (once daily for 2 days). Animals were allowed 1 week for recovery (housed one per cage) before testing. Microdialysis was performed as previously described by Lupinsky.\(^1\) Probes were calibrated in artificial cerebrospinal fluid (26 mmol/L NaHCO\(_3\), 3 mmol/L NaH\(_2\)PO\(_4\), 1.3 mmol/L MgCl\(_2\), 2.3 mmol/L CaCl\(_2\), 3.0 mmol/L KCl, 126 mmol/L NaCl. 0.2 mmol/L L-aspartic acid) containing 100 ng/ml aspartate, GLU, and GABA. In vitro probe recovery ranged from 14% to 19% at a flow rate of 2 μL/min. Computer-controlled microinfusion pumps were used to deliver perfusate to the probes, and the dialysate was collected from the fused silica outlet line (dead volume: 0.79 μL).

Microdialysis was performed under isoflurane anesthesia (2% isoflurane at 0.5 L/min oxygen flow). Two microdialysis probes (see Figures 4A and 4B) were inserted into the animals’ indwelling guide cannula and perfused with sterile artificial cerebrospinal fluid (flow rate set at 1 μL/min). Samples were then taken at 20-minute intervals for 1 hour (baseline), followed by CEF (200 mg/kg) injection in the tail vein, while still collecting samples for 4 hours (challenge, Figure 1C). Each 20-μL dialysate sample was collected in a fraction vial preloaded with 1 μL of 0.25 mol/L perchloric acid to prevent analyte degradation and immediately stored at 4°C for subsequent analysis. After microdialysis, animals brains were dissected and stored in 4% paraformaldehyde and subsequently cryoprotected in a 30% sucrose solution. Brains were sliced in 20-μm-thick samples (coronal) and stained with cresyl violet (Nissl staining) for confirmation of probes placement.

Lysate High-Performance Liquid Chromatography Analysis

Glutamate levels were determined as previously described by Lupinsky.\(^1\) A high-performance liquid chromatography precolumn derivatization with electrochemical detection was used to determine glutamate levels. The chromatographic system consisted of an ESA pump (model 582) and an ESA injector (model 542) coupled to a Waters Xterra MS C18 3.0 × 50 mm 5 μm analytical column. The mobile phase was prepared as needed and consisted of 3.5% acetonitrile, 20% methanol, and 100 mmol/L sodium acid to prevent analyte degradation and immediately stored at 4°C for subsequent analysis. After microdialysis, animals brains were dissected and stored in 4% paraformaldehyde and subsequently cryoprotected in a 30% sucrose solution. Brains were sliced in 20-μm-thick samples (coronial) and stained with cresyl violet (Nissl staining) for confirmation of probes placement.
phosphate dibasic (Na₂HPO₄) adjusted to pH 6.7 with 85% phosphoric acid. The flow rate was set at 0.5 mL/min, and the electrochemical detector ESA Coularray 5600 A, Chelmsford, MA, USA) was set at potentials of +150 mV and +550 mV. Working standards (100 ng/mL) and derivatization reagents were prepared fresh daily from stock solutions and loaded with samples into a refrigerated (10°C) ESA autosampler (model 542, Chelmsford, MA, USA). Before injection onto the analytical column, each fraction was sequentially mixed with 20 μL of o-phthaldehyde (0.0143 mol/L) diluted with 0.1 mol/L sodium tetraborate and 20 μL of 3-mercaptopropionic acid (0.071 mol/L) diluted with H₂O and allowed to react for 5 minutes. After each injection, the injection loop was flushed with 20% methanol to prevent contamination of subsequent samples. Under these conditions, the retention time for GLU was 2.4 minutes with a total run time of 30 minutes/sample. Chromatographic peak analysis was accomplished by

Figure 2. [¹¹C]ABP688 time-activity curves (TACs) obtained from baseline and challenge conditions. (A) Baseline and post ceftriaxone TACs from the cerebellum (reference region) and (B) the thalamic VA nucleus (high-binding region) of a single rat. Blue squares represent baseline condition and red squares the challenge condition.

Figure 3. Baseline and ceftriaxone (CEF) challenge averaged [¹¹C]ABP688 nondisplaceable binding potential (BPND). Averaged [¹¹C]ABP688 BPND images obtained during the baseline overlaid on a histologic template are shown in axial (A), sagittal (B), coronal (C), as well as dorsal (D) and mid-sagittal surface projections (E). Averaged [¹¹C]ABP688 BPND images obtained after CEF challenge overlaid on a histologic template are shown in axial (F), sagittal (G), coronal (H), as well as dorsal (I) and mid-sagittal surface projections (J). Statistical parametric images (t-stats contrast (CEF challenge < baseline)) overlaid on a histologic template are shown on axial view (K); sagittal view (L); coronal view (M); and rat brain 3D surface showing (N) peak t-stat region (the gray object represents hippocampal position in the 3D surface). Note group differences showing a symmetric cluster in the anterior thalamus, encompassing basal forebrain and posterior striatum with the local maxima at the thalamic VA. [¹¹C]ABP688 BPND in the local maxima, during baseline and challenge microPET scans (O). Schematic representation of [¹¹C]ABP688 BPND binding in the metabotropic glutamate receptor type 5 (mGluR5) allosteric site (P). Images represented as [¹¹C]ABP688 BPND and t-value. Graph represented as mean ± s.d., n = 5. **P < 0.01. PET, positron emission tomography.
Identification of unknown peaks in a sample matched according to retention times from known standards using ESA’s CoulArray software.

Statistical Analysis
A paired Student’s t-test was used to calculate differences in the $[^{11}C]ABP688$ BPND between challenge and baseline. Comparisons between baseline and challenge were conducted at the voxel level using t-statistic analyses (RMINC, https://launchpad.net/rminc). Regions showing high specificity binding were adjusted for a statistical cluster-wise threshold of $P < 0.05$, and corrected for multiple comparisons using random field theory.19 For microdialysis, two-way ANOVA with repeated measures followed by Bonferroni correction was used. Differences were considered statistically significant at $P < 0.05$.

RESULTS
Ceftriaxone Challenge Increased $[^{11}C]ABP688$ BPND with Peak Effect in the Thalamic Ventral Anterior Nucleus
Figure 2 shows the representative time-activity curves of reference region (cerebellum, Figure 2A) and of a high-binding region (thalamic VA nucleus, Figure 2B). Averaged BPND images showed consistent brain uptake of $[^{11}C]ABP688$, and no global differences between baseline and challenge (baseline: $1.06 \pm 0.24$; challenge: $1.14 \pm 0.26$; $t(4) = 0.5135$, $P = 0.6347$; Figures 3A and 3J). Voxel-based analysis showed a large bilateral symmetric cluster with increased $[^{11}C]ABP688$ BPND in the thalamus, with peak effect in the thalamic ventral nucleus (VA) (peak $t(4) = 6.78$, $P = 0.0025$, Figures 3K–3N) after the challenge with CEF. Paired t-test showed a significant increase in the $[^{11}C]ABP688$ BPND in the VA (local maxima) after challenge with CEF ($t(4) = 5.295$, $P = 0.0061$, baseline: $0.9781 \pm 0.24$; challenge: $1.379 \pm 0.08$ Figure 3O).

Ceftriaxone Evoked Glutamate Reductions in the Thalamic Ventral Nucleus
Repeated measures two-way ANOVA revealed an effect of drug ($F(13,130) = 5.379$, $P = 0.0001$) but not of region ($F(1,10) = 1.373$, $P = 0.2685$). A significant interaction between drug and region ($F(13,130) = 2.109$, $P = 0.0175$) was observed. Bonferroni post hoc analysis revealed decreased glutamate concentrations after challenge in the VA ($P_{(range)} = 0.0004$ to 0.0314) but not in the FC ($P_{(range)} = 0.2526$ to 0.999; Figure 4C). Comparison between averages of baseline and CEF challenge during the scan period also showed decreased extracellular levels of glutamate in the VA ($t(5) = 4.183$, $P = 0.0086$), but not in the FC ($t(5) = 1.482$, $P = 0.20$; Figure 4D).

DISCUSSION
The present study showed for the first time the effect of the reduction of extracellular glutamate levels, using a GLT-1 pharmacological challenge with CEF, on PET $[^{11}C]ABP688$ binding. In a previous study, we validated the cerebellum as a suitable reference region for the quantification of mGluR5 availability using $[^{11}C]ABP688$ in rats, via in vivo (microPET) and in vitro (autoradiography) techniques. Additionally, we performed a blood-based pharmacokinetics analysis and a blocking study with 1,2-methyl-6-(phenylethynyl)-pyridine, an antagonist of mGluR5 allosteric binding site and a competitor of $[^{11}C]ABP688$.13 Furthermore, we evaluated the reproducibility of $[^{11}C]ABP688$ binding in a test-retest study and examined several pharmacokinetic models for determination of binding potential (BPND), showing that the simplified reference tissue model is the pharmacokinetic model that presents lowest variability.20 These validation studies allowed us to proceed with a pharmacological challenge aiming to identify the impact of reduced...
glutamate concentrations on \(^{11}C\)ABP688 binding in rats. Here, we show that the acute CEF challenge increased \(^{11}C\)ABP688 binding in the thalamic VA. Beyond the considerable expression levels of GLT-1, \(^{11}C\)ABP688 binding is highly dependent on the tertiary and quaternary receptor conformations.\(^{25}\) Additionally, mGLR5 can assume oligomeric or heteromeric forms,\(^{26–28}\) which may have an impact on the availability of the allosteric site. One could also argue that glutamate levels alter mGLR5 conformational states. In fact, affinity shift in the receptor–radioligand interactions is a phenomenon already described in the case of dopamine D2 receptors, where an amphetamine challenge altered the affinity of a D2 PET radiopharmaceutical.\(^{29,30}\)

Interestingly, \(^{11}C\)ABP688 binding to the transmembrane allosteric site instead of orthosteric binding site of glutamate (Figure 3P) is highly dependent on the tertiary and quaternary receptor conformations.\(^{25}\) Additionally, mGLR5 can assume oligomeric or heteromeric forms,\(^{26–28}\) which may have an impact on the availability of the allosteric site. One could also argue that glutamate levels alter mGLR5 conformational states. In fact, affinity shift in the receptor–radioligand interactions is a phenomenon already described in the case of dopamine D2 receptors, where an amphetamine challenge altered the affinity of a D2 PET radiopharmaceutical.\(^{29,30}\)

Guided by \(^{11}C\)ABP688 microPET data, a microdialysis study was performed to confirm the reduction of glutamate levels in the VA. As expected, since \(^{11}C\)ABP688 binding was not altered in the cerebral cortex, levels of glutamate in the FC (used as a control region) remained stable after CEF injection. In contrast, 20 minutes after CEF injection glutamate levels were reduced in the VA. However, the reason why this thalamic region is the first to be affected by CEF since GLT-1 and mGLR5 are expressed in almost all brain regions remains unknown and requires further investigation. One could argue that the differential expression and splice variants of GLT-1 in diverse brain regions may yield distinct activation by CEF.\(^{32,33}\) Further studies are necessary to validate this observation; however, it seems unlikely that the changes in \(^{11}C\)ABP688 binding are the result of a direct displacement by CEF, since CEF does not bind to the mGLR5 allosteric site. In fact, further methodological developments are necessary to better understand noncompetitive interactions using PET. Altogether, our results support a theoretical framework in which synaptic glutamate concentrations antecorrelate with \(^{11}C\)ABP688 binding (i.e., low levels of glutamate will produce increased \(^{11}C\)ABP688 binding and high levels of glutamate decreased \(^{11}C\)ABP688 binding). Interestingly, two recent studies showed that \(^{11}C\)ABP688 PET was capable of identifying reductions in mGLR5 availability in patients with major depressive disorder,\(^{34}\) and in an epilepsy rat model.\(^{35}\) On the basis of our assumptions, reduced \(^{11}C\)ABP688 PET binding in this study may indicate increased glutamate levels, a possible sign of early excitotoxicity. This seems reasonable since excitotoxicity is an important pathological entity in both major depression\(^{36}\) and epilepsy.\(^{37}\)

In summary, molecular imaging using \(^{11}C\)ABP688 seems to constitute a valuable tool for noninvasive and in vivo assessment of glutamatergic neurotransmission. Studies encompassing a wide range of clinical populations displaying glutamate-related neurologic and neuropsychiatric conditions are necessary to fully evaluate the feasibility and potential of \(^{11}C\)ABP688 for measuring clinically-relevant glutamatergic fluctuations. If these clinical studies succeed, then \(^{11}C\)ABP688 PET imaging will offer the opportunity of identifying glutamate abnormalities (e.g., brain excitotoxicity), which may allow for the development of glutamate-based disease signatures and for the evaluation of glutamate focused therapies.

**AUTHOR CONTRIBUTIONS**

ERZ and PRN were responsible for the design, drafting and revising the manuscript. ERZ and MJP were responsible for imaging processing and analysis. AL was responsible for drafting and revising the manuscript. ERZ, Arturo A and AA were responsible for microPET procedures and histology. ERZ, Arturo A and LM were responsible for microdialysis procedures. ESS and JPS were responsible for the radiochemistry. AG, SG and PRN critically revised the final version of the manuscript. PRN and AG were responsible for overseeing the entire PET and microdialysis procedures, respectively.

**DISCLOSURE/CONFLICT OF INTEREST**

The authors declare no conflict of interest.

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