RAPID COMMUNICATION

Differential Patterns of Subcortical Activity Evoked by Glial GLT-1 Blockade in Prelimbic and Infralimbic Cortex: Relationship to Antidepressant-Like Effects in Rats

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Abstract

Background: Glutamatergic neurotransmission has emerged as a novel target in antidepressant drug development, with a critical role of the ventral anterior cingulate cortex. We recently reported that blockade of the astrocytic glutamate transporter GLT-1 with dihydrokainic acid in infralimbic cortex (rodent equivalent of ventral anterior cingulate cortex), but not in the adjacent prefrontal cortex, evoked robust antidepressant-like effects through α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid receptor activation and increased serotonin release.

Methods: 2-deoxy-2-[18F]-fluoro-D-glucose-positron emission tomography and computed tomography in 36 male Wistar rats microinfused bilaterally in prelimbic cortex or infralimbic cortex with dihydrokainic acid or vehicle.

Results: Dihydrokainic acid microinfusion in infralimbic cortex and prelimbic cortex evoked dramatically different regional patterns of subcortical activity. In infralimbic cortex, dihydrokainic acid selectively affected midbrain areas, whereas in prelimbic cortex it affected the basal ganglia, the thalamus, and both superior and inferior colliculi.

Conclusions: These results highlight the differential connectivity of infralimbic and prelimbic cortex with subcortical brain regions and support the involvement of infralimbic cortex-midbrain pathway in the antidepressant-like effects of dihydrokainic acid.

Keywords: infralimbic, prelimbic, glutamate transporter-1, FDG-PET, dihydrokainic acid
Significance Statement

The glutamatergic system is a potential target to develop fast-acting antidepressants. On the other hand, glial cells tightly control glutamatergic synapses. We recently reported that boosting glutamatergic neurotransmission by inhibiting astrocytic glutamate uptake in a ventral area of the rat prefrontal cortex (infralimbic, IL; but not in the neighboring prelimbic cortex, PrL) evoked rapid and robust antidepressant-like effects. Using microPET scan, we show that the blockade of astrocytic glutamate uptake in IL and PrL evokes dramatically different patterns of brain activity, showing at the same time that IL-midbrain circuits (possibly involving serotonergic neurotransmission) underlie the reported antidepressant effects.

Introduction

The glutamatergic system is emerging as a promising venue for the development of fast-acting antidepressant treatments, given the immediate and persistent antidepressant effects of the noncompetitive N-methyl-D-aspartate receptor antagonist ketamine (Zarate et al., 2006). Its unique properties appear to result from the activation of α-amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) receptors by a metabolite (25,65,2R,6h)-hydroxynorketamine (Zanos et al., 2016) and by an increased synaptic plasticity evoked by mTOR signaling in rat medial prefrontal cortex (mPFC) (Li et al., 2010).

Ventral regions of the anterior cingulate cortex (vACC) seem to play a crucial role in the pathophysiology and treatment of major depressive disorder. Neuroimaging studies have reported controversial findings on the energy metabolism of that area in major depressive disorder patients. A reduced activity of the subgenual vACC was first described (Drevets et al., 1997; Ongür et al., 1998), whereas further studies reported an increased activity of the adjacent Brodmann area 25, which was normalized after effective treatment (Seminowicz et al., 2004; Mayberg et al., 2005). Likewise, optogenetic stimulation of the infralimbic cortex (IL, rodent equivalent of vACC) in rats mimicked the antidepressant-like effects of systemic ketamine administration (Fuchikami et al., 2015).

In line with these observations, we recently reported that the local single-point blockade of the astrocytic glutamate transporter GLT-1 with dihydrokainic acid (DHK) in rat IL evoked immediate and robust antidepressant-like effects in the forced swim and novelty suppressed feeding tests, associated with a marked elevation of serotonin (5-HT) release (Gasull-Camós et al., 2017). Behavioral and neurochemical effects were blocked and mimicked, respectively, by local microinfusion of the AMPA KA antagonist NBQX and S-AMPA. Interestingly, neither effect occurred when DHK or S-AMPA were locally applied in the adjacent prelimbic cortex (PrL) using exactly the same methodology and drug doses as in IL experiments (Gasull-Camós et al., 2017).

Using 2-deoxy-2-[18F]-fluoro-D-glucose ([18F]FDG) positron emission tomography (PET) in rats, we examined the changes in brain activity induced by DHK microinfusion in IL and PrL using the same application procedure than in our previous report. Given that antidepressant-like effects of DHK were associated to increased 5-HT release and were prevented by prior 5-HT synthesis inhibition with pCPA (Gasull-Camós et al., 2017), the working hypothesis was that DHK application in IL (but not in PrL) would increase the activity of midbrain 5-HT neurons, remarkably sensitive to changes of neuronal activity in the mPFC (Hajós et al., 1998; Celada et al., 2001; Warden et al., 2012).

Methods

Animals

Male Wistar rats (Charles River) weighing 280 to 330 g at the time of surgery were used. Rats were housed in a temperature- and humidity-controlled vivarium with a 12-h-light/dark cycle and with food and water ad libitum (unless otherwise stated). Experiments were performed according to the guidelines of the European Union Council Directive 2010/63/EU for care of laboratory animals and after approval by the Ethics Committee for Animal Experimentation of Hospital Gregorio Marañón, Madrid, Spain.

Surgery

Anesthetized rats (sodium pentobarbital, 60 mg/kg i.p.) were placed into a stereotaxic frame. Stainless-steel 22-gauge bilateral guide cannulae (Plastics One) were implanted in the cingulate cortex: AP +3.2; ML ±0.75; DV -2.4 ( Paxinos and Watson, 2005) as reported (Gasull-Camós et al., 2017). The coordinates were taken from bregma and the skull. Guide cannulae were fixed with 3 stainless-steel screws using dental acrylic. A dummy cannula was inserted inside the guide cannula and removed and reinserted daily to prevent occlusion. After surgery, rats were allowed 7 days of recovery.

Microinfusions

DHK was purchased from Tocris and dissolved in PBS 10x (Gasull-Camós et al., 2017). The microinfusion cannulae (28 gauge) extended 1.5 or 3 mm beyond the guide cannulae for PrL or IL local administration (DV: -3.9 or -5.4, respectively). The day previous to drug microinfusion and [18F]FDG-PET testing, a mock infusion was performed. On the testing day, an infusion/withdrawal pump (Harvard Apparatus) was used to bilaterally administer DHK or PBS into the PrL or IL cortex through two 100-μL Hamilton syringes connected to the microinfusion cannula via 0.28-mm ID polyethylene tubing. A volume of 0.5 μL/hemisphere was administered over 1 minute, and microinfusion cannulae were left in place for 3 minutes to allow drug diffusion.

In Vivo Imaging Studies

PET studies were acquired with a small animal PET/computed tomography (CT) scanner (ARGUS PET/CT, SEDECAL). [18F]FDG (37 MBq; IBA Molecular Spain S.A.) was injected through the tail vein 10 minutes after completing the DHK or PBS infusion. After 45 minutes of uptake, animals were scanned under anesthesia with sevoflurane (3% induction, 1.5% maintenance in 100% O2) for 45 minutes. Images were reconstructed by using a 2D-OSEM algorithm, full width half maximum of 1.45 mm, with a voxel size of 0.3875 × 0.3875 × 0.775 mm and an energy window of 400 to 700 keV. Decay and dead-time corrections were applied.

CT studies were acquired immediately before each PET scan with the same scanner to facilitate PET image registration. Acquisition parameters were 340 mA, 40 kV, 360 projections, 8 shots, and 200 μm of resolution. CT images were reconstructed.
using an FDK algorithm (isotropic voxel size of 0.121 mm) (Hadar et al., 2017).

**PET Preprocessing and Analysis**

PET scans were preprocessed as previously described (Hadar et al., 2017). Briefly, PET images were spatially co-registered to a random reference CT scan. A whole brain mask segmented on a magnetic resonance scan registered to the same reference CT scan was applied to all PET images to eliminate voxels outside the brain. Afterwards, PET images were smoothed with an isotropic Gaussian filter (2 mm full width half maximum).

Voxel value normalization consisted of standardizing PET intensity data to a brain region without statistically significant differences between groups (nonsignificant area [NSA]) obtained by an iterative method (Borghammer et al., 2009). First, PET data were normalized to global mean brain intensity and analyzed with a voxel-by-voxel analysis using SPM12 (http://www.fil.ion.ucl.ac.uk/spm/software/spm12/). Groups were compared using a 2-sample t test (P < .05, uncorrected). Then, a mask of the NSA was acquired from the resulting T-maps, excluding statistically significant clusters. Next, the NSA mask was used to normalize the original smoothed PETs, and the obtained images were reanalyzed with the same SPM protocol. Subsequently, an NSA mask from these second T maps was obtained, and we repeated the whole process until the fourth iteration, which was accepted as the final result of the analysis. Statistics were not corrected for multiple comparisons because of the lack of validation with rodent brain data of multiple comparison correction methods, such as familywise error rate or false discovery rate (Romero et al., 2011), and also because of the ethical demand to minimize the number of animals (Hadar et al., 2017). Thus, although this leads to a reduction in power, it prevents an underestimation of the statistical significance. Significant regions larger than 50 activated connected voxels were considered.

A region of interest analysis was performed to determine the intragroup global metabolic differences. Whole brain and NSA masks were used for this analysis. Whole brain data were normalized to the average NSA intensity. Global differences were assessed by means of a t test with a threshold for statistical significance set at P < .05.

**Results**

**In Vivo Study of the DHK Effects**

Measurements based on global alterations for whole brain metabolism showed no significant differences between treatment and vehicle groups, as shown by 2-sample t test. Thus, the microinfusion of DHK in PrL did not change the overall uptake of ⁴⁰FDG compared with controls (DHK: 1.0078 ± 0.0009; PBS: 1.0081 ± 0.0007; P = .825) and neither did the microinfusion of DHK in IL (DHK: 1.0078 ± 0.0003; PBS: 1.0082 ± 0.0004; P = .405).

Despite there were no differences in the average global uptake of ²⁰¹FDG, significant differences of the local metabolism were observed after DHK microinfusion. Thus, the microinfusion of DHK in PrL increased glucose metabolism in the prefrontal cortex (T = 6.42) and the cerebellum (T = 3.13). Conversely, PrL DHK application reduced glucose metabolism in the nucleus accumbens, the dorsal striatum (caudate-putamen), the thalamus, the ventral hippocampus, and the superior and inferior colliculi (T = 5.1) as well as in anterior cerebellar regions (T = 2.46) (Figure 1; Table 1). On the other hand, the microinfusion of DHK in IL produced an increase of glucose metabolism in the IL itself and the olfactory nucleus (T = 2.61) as well as in the temporal cortex (T = 5.16). A reduction of ²⁰¹FDG uptake occurred in the periaqueductal gray matter (PAG), the midbrain region (T = 3.77), the somatosensory cortex (T = 2.47), and the cerebellum (T = 2.42) (Figure 1; Table 1).

**Figure 1.** Changes in brain metabolic activity. Voxel-based SPM results in T-maps overlaid on a T2 magnetic resonance image, showing the changes in glucose metabolism due to dihydrokainic acid (DHK) administration in prefrontal (PrL, left) or infralimbic (IL, right). The color bars on the right represent the T values corresponding to lower (blue) and higher (red) 2-deoxy-2-[¹⁸F]-fluoro-D-glucose (⁴⁰FDG) uptake (P < .05 [unc.]; k = 50 voxels). Brain regions: Cerebellum (Cb), caudate-putamen (CPu), inferior colliculus (IC), infralimbic cortex (IL), nucleus accumbens (NAc), olfactory nucleus (ON), periaqueductual gray matter (PAG), prefrontal cortex (PFC), superior colliculus (SC), somatosensory cortex (S1), temporal cortex (Temp C), thalamus, ventral hippocampus (vHPC).
Discussion

The present study shows that blockade of the astroglial glutamate transporter GLT-1 with DHK in IL and PrL affects brain activity in a remarkably different manner, as assessed by microPET scan with $^{18}$FDG. In particular, the areas affected by DHK application in IL may reflect the brain circuitry responsible for the antidepressant-like effects and increased 5-HT release evoked by this procedure (Gasull-Camós et al., 2017).

The present and preceding observations add to previous studies supporting a crucial role of astrocytes in synaptic transmission and animal behavior (Oliveira et al., 2015), an effect due to their ability to control glutamatergic synapses (Perea and Araque, 2010). Hence, the astrocytic glutamate transporters GLT-1 and GLAST are responsible for the uptake of most synaptic glutamate, with a minor role of the neuronal transporter EAAC1 (Danbolt, 2001).

GLT-1 blockade markedly elevated energy metabolism in the application areas, an effect likely related to the increased glutamate outflow produced by DHK (Gasull-Camós et al., 2017) and the subsequent activation of local excitatory receptors, leading to an increased neuronal discharge and energy consumption. Despite that we used the same experimental procedure and that DHK elevates extracellular glutamate to the same extent in both subdivisions (Gasull-Camós et al., 2017), DHK application in PrL markedly activated neighboring PFC areas, such as cingulate and motor cortices, whereas DHK application in IL evoked a more restricted activation pattern. This difference may possibly be related to the inhibitory role of IL on PFC activity (Ji and Neugebauer, 2012), thus preventing an excitatory wave expanding outside the application site, as after PrL application.

In addition to their respective application sites, GLT-1 blockade affected several cortical and subcortical regions in a different manner, sometimes with opposite activity changes (increase and decrease in cortical/cerebellar and subcortical areas, respectively, as shown in Figure 1 and Table 1). DHK application in PrL mainly affected structures from basal ganglia circuits (e.g., dorsal and ventral striatum, thalamus) as well as the ventral hippocampus, the colliculi, and cerebellum. In contrast, GLT-1 blockade in IL mainly reduced metabolic activity in midbrain structures, such as the PAG, yet increased it in temporal cortex and olfactory nucleus. It is of note that some effects seem to be lateralized, like the reduced left S1 and the increased right temporal cortex glucose metabolism, after IL DHK. Interestingly, clinical studies in repeated transcranial magnetic stimulation showed that the left and right cortices require different frequency ranges of stimulation (high/excitatory and low/inhibitory, respectively) to achieve antidepressant efficacy (Chen et al., 2013). Moreover, greater activity of left PFC was predictive of favorable response to antidepressants (Hoehn-Saric et al., 2001). Although these studies focused on PFC, it would be interesting to study the contribution of brain activity asymmetry in other cortical regions for the antidepressant response.

The comparable elevation of extracellular glutamate concentration in IL and PrL after GLT-1 blockade (Gasull-Camós et al., 2017) indicates that differential effects of DHK in IL vs PrL in terms of animal behavior, elevation of 5-HT release, and brain areas affected are not due to a distinct control of synaptic glutamate by GLT-1 in each of these cortical areas. Most convincingly, these differences are due to the different role of IL and PrL in PFC functions (Dalley et al., 2004), resulting from their differential connectivity with subcortical structures involved in cognitive and emotional processing (Vertes, 2004; Gabbott et al., 2005). Additionally, a distinct reactivity of pyramidal neurons or local microcircuits in IL and PrL to the glutamate increase evoked by GLT-1 blockade might be involved, given the presence of reciprocal connections between both PFC subdivisions (Gabbott et al., 2003) and the inhibitory role of IL on PrL activity (Ji and Neugebauer, 2012).

The reduction of activity in subcortical structures after GLT-1 blockade in IL and PrL appears inconsistent with the excitatory nature of descending pyramidal inputs. However, some of these structures (e.g., dorsal striatum, nucleus accumbens, PAG) are composed essentially of GABAergic neurons, while others (e.g., thalamus) are tonically inhibited by GABAergic inputs from substantia nigra reticulata, ventral pallidum, and the thalamic reticular nucleus. Therefore, descending cortical excitatory inputs may activate inhibitory neurotransmission in input and output structures of the basal ganglia, thus leading to a reduced neuronal activity and energy consumption.

Likewise, the reduced energy metabolism in midbrain produced by DHK application in IL seems at variance with the elevated 5-HT release produced by this procedure (Gasull-Camós et al., 2017) and the increased metabolic activity in midbrain structures after GLT-1 blockade in IL and PrL. The GLT-1 blockade markedly reduced metabolic activity in midbrain structures, as shown in Table 1 and Figure 1. This might reflect the involvement of the dorsal and ventral striatum, the thalamus, and the ventral hippocampus in inhibitory neurotransmission in response to DHK application.
This 5-HT elevation, together with the cancellation of antidepressant-like effects of IL DHK application by prior 5-HT synthesis inhibition, led us to hypothesize an increased activity of raphe 5-HT neurons. Contrary to these expectations, the present data indicate a marked and overall reduction of energy metabolism in PAG (whose ventral part includes the dorsal raphe nucleus) after IL DHK application. One possible confounding factor is the use of anesthesia in the present study, the only experimental difference with respect to the previous study. However, this apparent contradiction may be partly explained by the complex neuronal connectivity between the mPFC and raphe 5-HT neurons, also including reciprocal PrL-IL connectivity, as stated above. Indeed, electrical stimulation at physiological rates (e.g., ~1 Hz) of the mPFC, particularly in its IL subdivision, led to a majority of inhibitory responses in dorsal raphe 5-HT neurons recorded in vivo in anesthetized rats. 

Inhibitory responses were mediated by local GABA<sub>α</sub> inputs (Hajós et al., 1998; Varga et al., 2001) as well as by self-inhibitory responses mediated by 5-HT<sub>1A</sub> autoreceptors (Celada et al., 2001). Actually, raphe 5-HT neurons receive local GABA contacts (Harandi et al., 1987), which inhibit 5-HT neurons (Liu et al., 2000). Contacts among cells of each neuronal subtype (GABA-GABA and 5-HT-5-HT) have also been reported (Harandi et al., 1987). Therefore, an IL-driven inhibition of PAG activity may attenuate GABA inputs onto 5-HT neurons, leading to an overall disinhibition of the 5-HT system and increased forebrain 5-HT release, as previously observed (Gasull-Camós et al., 2017). The activation of 5-HT neurons may have been masked by a greater reduction of PAG activity, given the limited resolution of the microPET technique.

In summary, the marked neurochemical (Gasull-Camós et al., 2017) and neuroimaging (present study) differences result in astroglial GLT-1 blockade in IL and PrL further enhance the relevance of astrocytes in modulating brain function and contribute to a better understanding of brain networks involved in fast antidepressant actions. The present data also warrant further investigation on the differential control of IL and PrL on serotonergic activity.

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Statement of Interest

F.A. has received consulting honoraria from Lundbeck A/S and has been PI of a grant from Lundbeck A/S. He is also a member of the scientific advisory board of Neurilaxis and a co-inventor of 2 patents on conjugated oligonucleotide sequences. The remaining authors declare no conflict of interest.

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