Effects of Glutamate Receptor Activation on Local Oxygen Changes

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ABSTRACT: Glutamate is ubiquitous throughout the brain and serves as the primary excitatory neurotransmitter. Neurons require energy to fire, and energetic substrates (i.e., O2, glucose) are replenished via cerebral blood flow (CBF) to maintain metabolic homeostasis. Magnetic resonance brain functionality studies rely on the assumption that CBF and neuronal activity are coupled consistently throughout the brain; however, the origin of neuronal activity does not always coincide with signals indicative of energy consumption (e.g., O2 decreases) at high spatial resolutions. Therefore, relationships between excitatory neurotransmission and energy use must be evaluated at higher resolutions. In this study, we showed that both endogenously released and exogenously ejected glutamate decrease local tissue O2 concentrations, but whether hyperemic O2 restoration followed depended on the stimulus method. Electrically stimulating the glutamatergic corticostriatal pathway evoked biphasic O2 responses at striatal terminals: first O2 decreased, then concentrations increased above baseline. Using iontophoresis to locally eject ionotropic glutamate receptor antagonists revealed that these receptors only influenced the O2 decrease. We compared electrical stimulation to iontophoretic glutamate stimulation, and measured concurrent single-unit activity and O2 to limit both stimulation and recordings to <50 μm radius from our sensor. Similarly, iontophoretic glutamate delivery elicited monophasic O2 decreases without subsequent increases.

KEYWORDS: Glutamate, hyperemia, iontophoresis, ionotropic glutamate receptors, neurovascular coupling, single-unit electrophysiology

INTRODUCTION

Despite accounting for a small portion of total body mass, the brain uses a disproportionately large percentage of the body's resting energy. This discrepancy arises from the energetically expensive demands of neuronal communication and the maintenance of ionic gradients. Metabolic substrates (i.e., glucose and O2) are replenished after neuronal activity via increased regional cerebral blood flow (CBF), where they are consumed to generate adenosine triphosphate (ATP) and reestablish ion gradients. To maintain energetic homeostasis, neuronal activity is typically coupled with increased CBF in a process termed functional hyperemia. However, dysfunctional decoupling of neuronal activity and CBF can lead to chronic energy deficits where blood flow cannot replenish the energy consumed during neurotransmission. With insufficient energy, ionic gradients collapse, often leading to neuronal death. CBF delivery is not exclusively controlled by metabolic deficiencies, as the amplitudes of task-evoked CBF increases are maintained when the same task is performed under hypoglycemic or hypoxic conditions. Further, although CBF delivers excess O2 to overcompensate for energy loss, the O2 concentrations delivered vary depending on stimulus type and intensity. Together, these suggest that neuronal activity and metabolic deficits play complex roles in the regulation of energy throughout the brain.

Often, CBF is measured indirectly by monitoring extracellular O2 concentrations, where increases in O2 can be attributed to hyperemic increases in blood flow. This is the basis for blood oxygenation level-dependent (BOLD) functional magnetic resonance imaging (fMRI); however, positive BOLD fMRI signals (i.e., increases in oxygenated blood) only infer the presence of increased neural activity. Fast-scan cyclic voltammetry (FSCV) also measures O2 changes, but the carbon-fiber microelectrodes used with the technique limit analyte detection to a range within a 20 μm radius of the electrode surface. As with BOLD, these electrochemical measurements also do not provide information about neuronal activity. Previously, we used a multimodal sensor to perform simultaneous FSCV and single-unit electrophysiology. Here, we utilize a different voltammetric waveform at our sensor to study colocalized O2 dynamics in conjunction with cell firing. This approach uses a single, minimally invasive carbon-fiber microelectrode and enables simultaneous detection of both neuronal activity and O2 changes in localized environments. Further, we coupled iontophoresis, a local drug delivery method, to FSCV and electrophysiology at the same sensor to form a comprehensive suite of techniques suited for spatially focused functional hyperemia studies. Using iontophoresis to eject glutamate, an excitatory neurotransmitter, stimulated local neuronal firing in anesthetized subjects and gave us a high

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Figure 1. Electrically stimulated glutamatergic cell bodies in the PFC elicit biphasic O₂ changes in the striatum. (A) (i) The multimodal sensor ejected glutamate iontophoretically to determine proximity to glutamate-sensitive cells (e.g., medium spiny neuron, MSN). Sensor barrel diameters and carbon-fiber length not to scale. (ii) Four replicate perievent rasters, where each black tic represents a recorded single-unit action potential. In anesthetized subjects, glutamate ejections (gray box) evoked single-unit activity bursts that ceased soon after the ejection. (iii) Averaged single-unit waveform of the unit isolated in panel A part ii. (B) Sagittal view of the chosen stimulated glutamatergic pathway. A bipolar stimulating electrode was placed into the PFC (blue) and a multimodal sensor was inserted into the dorsal striatum (orange). (C) (top) Current taken at the O₂ reduction waveform of the unit isolated in panel A, part ii. (B) Sagittal view of the chosen stimulated glutamatergic pathway. A bipolar stimulating electrode was placed into the PFC (blue) and a multimodal sensor was inserted into the dorsal striatum (orange). (C) (top) Current taken at the O₂ reduction potential (−1.35 V, horizontal dotted line in color plot below) tracked the time course of O₂ changes from the representative color plot below. First, O₂ decreased during the stimulation (event 1) and subsequently increased above baseline following electrical stimulation (event 2). Cyclic voltammograms from each event (insets, each taken from horizontal white lines in color plot below) confirmed the identity of O₂, with background subtraction taken at the color plot black dashed line. Slight hysteresis was attributed to concomitant endogenous ion fluctuations. (bottom) Color plot representation of cyclic voltammetry data recorded around a 100 pulse, 20 Hz, 300 μA electrical stimulation as indicated by the gray bar. Abscissa, applied voltage; ordinate, acquisition time of the cyclic voltammograms. Background-subtracted currents are color coded. (D) Averaged O₂ changes evoked from 2 s glutamate iontophoresis ejections at the glutamate-sensitive striatal cells before the first electrical stimulation and 10 min after the last electrical stimulation (n = 6). Shading indicates ±SEM. (E) Experimental paradigm for assessing local striatal iGluR influence over O₂. PFC cell bodies were electrically stimulated after a cocktail of AP5 (50 mM) and CNQX (10 mM) was ejected at striatal terminals for 60 s. (F) Event 1 was significantly attenuated (***P = 0.001, Student’s t test) under iGluR blockade in the striatum. Error bars indicate ±SEM.

### RESULTS

**Electrically Stimulating PFC Glutamate Cell Bodies Evokes O₂ Responses at Striatal Terminals.** Glutamatergic neurons project from the PFC to medium spiny neurons (MSNs) in the striatum. We recorded O₂ changes at locations where glutamate iontophoresis elicited single-unit activity, that is, near the soma of glutamate-sensitive cells (Figure 1A). Unit activity was evoked during the glutamate ejection and ceased upon stimulation termination (Figure 1A). These cells were classified as MSNs, which account for over 90% of the cells in the striatum, based on their waveform shapes and low firing rates (Figure 1A). Next, we electrically stimulated the PFC to drive glutamate release in the striatum (Figure 1B) and recorded subsequent O₂ changes at glutamate-sensitive neurons. Electrical stimulations interfered with electrophysiological recordings, so we were unable to assess the single-unit activity during PFC stimulations. With FSCV, we observed robust, reproducible O₂ responses that consisted of two distinct phases: the first, event 1, was a decrease in O₂, followed by a second increase in O₂.

degree of control over the cell activity proximal to our sensor. Together, we used our multimodal sensor to stimulate and measure neuronal firing and the subsequent O₂ changes within a <50 μm radius of brain tissue. In this study, we compared the effects of electrically stimulated glutamate versus local glutamate iontophoresis on O₂ dynamics between the striatum and somatosensory cortex. We first elicited endogenous glutamate release via electrically stimulating the prefrontal cortex (PFC) and found that striatal O₂ concentrations both increased and decreased in a manner dependent on stimulation intensity and the availability of ionotropic glutamate receptors (iGluRs). In contrast, glutamate ejected locally within the striatum evoked either O₂ decreases or no change in local O₂ concentrations. The ejection method, controlled iontophoresis, uses current to deliver solution aliquots on the order of femto- to picomoles and that neuronal firing does not always couple to O₂ use throughout the cortex and striatum.
followed by an O₂ increase to above baseline values, termed event 2 (Figure 1C). These biphasic responses were consistent with poststimulus O₂ changes observed both locally using amperometry at microelectrodes⁸,²⁴ and globally with BOLD fMRI.⁸,¹⁰,²⁴ By comparison, 2 s glutamate iontophoresis ejections in the same locations evoked O₂ decreases of a similar magnitude as the electrical stimulations but failed to ever produce an event 2 (Figure 1D).

To confirm whether glutamatergic neurotransmission was responsible for the biphasic O₂ response, we iontophoresed iGluR antagonists for 60 s prior to electrical stimulation (Figure 1E). This was to achieve steady-state ejection conditions and saturate the tissue within ~100 μm of our sensor.²⁵ After collecting a predrug stimulation response, we delivered the NMDA receptor antagonist AP5 (50 mM) and the AMPA and kainate receptor antagonist CNQX (10 mM) for 60 s from a single iontophoresis barrel attached to the carbon-fiber electrode. After drug delivery, we stimulated the PFC again and compared the striatal O₂ changes before and after iGluR antagonism (Figure 1F). We chose a stimulation of 80 pulses delivered at 20 Hz, as this was the shortest stimulation train capable of reliably eliciting biphasic O₂ responses (Figure 2A,B). In all cases, the poststimulus O₂ decrease (event 1) significantly diminished under local iGluR blockade to 44.8% ± 11.1% of predrug values (\(t_{(2,8)} = 5.0, P = 0.001, n = 5\); Figure 1F), while differences in event 2 amplitudes were negligible. The attenuation of the evoked response with iGluR antagonists confirmed that PFC stimulations released endogenous glutamate into the striatum and indicated that local iGluR activation significantly contributed to O₂ decreases but not the overcompensating O₂ rebound.

**Biphasic O₂ Responses Depend on Electrical Stimulation Parameters.** We next investigated the dependence of the observed O₂ events on the stimulation parameters to explore what stimulus was sufficient to elicit a biphasic, hyperemic response. First, we kept stimulation frequencies and amplitudes constant at 20 Hz and 300 μA, respectively, and changed the number of stimulation pulses from 10–100 pulses (Figure 2A). The number of locations that responded with biphasic O₂ changes increased with the number of stimulus pulses (Figure 2B). Furthermore, the relationship was linear between both biphasic event amplitudes and the corresponding number of stimulation pulses (event 1, \(R^2 = 0.990\); event 2, \(R^2 = 0.975\), Figure 2C). All O₂ changes from 10 pulse stimulations were monophasic, so we omitted these data from the linear fit. This confirmed that both mono- and biphasic O₂ responses can be elicited at the same location, identified a pulse-dependent stimulus threshold that must be exceeded before biphasic responses occurred, and showed a strong linear relationship between the number of stimulation pulses and the resulting O₂ event amplitudes.

To ensure that the pulse-dependent stimulus effects were not caused by an increase in total stimulation length, we kept pulse numbers and amplitudes constant (120 pulses and 300 μA, respectively) and tested 5–60 Hz frequencies. These frequencies did not affect either event 1 or event 2 (when applicable) amplitudes or influence whether responses were mono- or biphasic (Figure 3A,B). Despite consistent amplitude changes across different frequencies, the time interval between stimulations and maximum O₂ event changes significantly increased from higher to lower frequency stimulations (one-way ANOVA, event 1, \(F_{(6,53)} = 6.26, P = 0.0002\); event 2, \(F_{(6,53)} = 5.47, P = 0.003\), Figure 3C). Bonferroni post hoc analysis revealed significant differences in event 1 minima times between both 5 and 10 Hz stimulations as compared to higher stimulation frequencies (5 Hz, 13.6 ± 2.5 s; 10 Hz, 13.1 ± 1.0 s; 30 Hz, 6.5 ± 0.5 s; 40 Hz, 6.4 ± 0.8 s; and 60 Hz, 6.2 ± 0.5 s; 5 Hz versus 30, 40, and 60 Hz: \(P < 0.01\) each, \(n = 6\) each; 10 Hz versus 30 and 40 Hz: \(P < 0.05\) each; 10 Hz versus 60 Hz: \(P < 0.01, n = 6\) each, Figure 3C). Time differences between stimulations and event 2 maxima (when present) also increased from 5 and 10 Hz stimulation frequencies to 60 Hz (5 Hz, 33.1 ± 7.5 s, \(n = 2\); 10 Hz, 30.8 ± 2.8 s, \(n = 4\); 60 Hz, 18.6 ± 0.5 s; 5 and 10 Hz versus 60 Hz: \(P < 0.05\) each, Figure 3C). These findings indicated that, without affecting O₂ event magnitudes,
low frequency stimuli increased the time elapsed between electrical stimulations and both the maximum event 1 and event 2 $O_2$ decreases and increases, respectively.

Glutamate Iontophoresis Provokes Local Single-Unit Firing and $O_2$ Decreases. We next delivered glutamate directly to the recording locations to exclude the confounding effects of nonglutamatergic neurotransmitters that could be released from electrical stimulation. Experiments started in the somatosensory cortex as proof of principle, then extended deeper within the brain through the striatum. We ejected glutamate from an iontophoresis barrel in 2 s pulses, until electrophysiology could resolve single-unit firing from background noise at the carbon fiber (Figure 4A,C). The amount of glutamate ejected was not quantified for each experiment, as the concentration gradient from a 2 s iontophoretic ejection overestimates the quantity ejected when using a standard calibration curve;26 instead, a steady-state glutamate concentration was used to estimate ejected quantities. A 120 s controlled iontophoresis ejection using 100 nA, with an assumed $175 \mu L s^{-1}$ ejection rate based on the ejection current,25 yielded a $3\times10^{-5}$ mM glutamate steady-state plateau. Therefore, glutamate iontophoresis for 2 s at 100 nA ejected glutamate on the order of $1\times10^{-2}$ pmol. Though steady-state ejections can diffuse solution to a radius of several hundreds of micrometers, a 2 s ejection is unlikely to diffuse much further than the barrel tip,26 especially with the efficiency of glial glutamate transporters.27

We investigated differences between the role of glutamate in superficial and deep brain environments, exploring the somatosensory cortex and the striatum. Cortical cells required $>100$ nA cathodic current to respond and were most often found at or below 0.9 mm D-V, corresponding to the most densely populated layers IV–VIb. These observations were consistent with published observations,16,17,28 and validated our approach to locally activate cells with iontophoresed glutamate.
Meanwhile, in agreement with the literature, we observed denser cell responses in the ventral striatum as compared to the dorsal striatum. Striatal cells responded using only 20 to 80 nA ejection currents, which was both less than the >100 nA ejection currents required for cortical cells and consistent with previous reports. The difference in ejection currents between brain regions is worth noting, as the amount of glutamate ejected per 2 s scales with ejection current. The consistent difference in the ejection currents required to elicit cell firing in the cortex versus the striatum suggested that glutamate sensitivity may differ between these two brain regions. However, cells also required different glutamate ejection currents within the same brain region, which we attribute to either cell proximity to the drug barrel, cell sensitivity to glutamate, or the availability of local excitatory receptors. These differences both between and within brain regions may be extremely subtle given that picomole or sub-picomole quantities of glutamate are ejected using currents above and below 100 nA, respectively.

In contrast to electrical stimulations, we observed one of two effects in the recorded environments following glutamate iontophoresis: either an exclusive monophasic O2 decrease (i.e., event 1 not followed by event 2; Figure 4A,B) or negligible O2 changes (Figure 4C,D). The monophasic O2 decreases typically returned to baseline within 40 s (Figure 4A,B). This contradicted a previous study that found a robust O2 increase following a local pressure ejection of 0.5 nmol glutamate aliquots. In contrast, our generous estimate of the glutamate delivered from a 2 s iontophoretic ejection was orders of magnitude less, 1–2 pmol. Overall, our sensor recorded one of four responses to glutamate at any given location. Either glutamate evoked cell firing or it did not, and the ejection was either succeeded by O2 decreases or did not perturb local tissue O2 (Figure 4), indicating that sensor responses were sensitive to the natural heterogeneity of the surrounding brain tissue.

**Glutamate Inhibits Fast-Spiking Interneuron Activity.** While characterizing glutamate-responsive neurons in the striatum, we serendipitously observed fast-spiking interneurons in ventral locations, offering us the unique opportunity to study them in vivo. These cells represent only 5–10% of striatal neurons and were identified as interneurons based on the observed high spontaneous firing rates (Figure 5A). Though we were unable to positively identify the type of fast-spiking interneuron from single-unit recordings alone, the 2 to 20 Hz spontaneous firing rates observed in deeply anesthetized subjects were in line with spontaneous firing rates reported for cholinergic interneurons. Up to 3 spontaneously active cells could be discerned per fast-spiking interneuron recording location (Figure 5A,B), consistent with interneuron clustering. Local glutamate ejections significantly inhibited, rather than augmented, single-unit firing rates in the vast majority of these spontaneously active cells (49.0% ± 8.2% of baseline activity, t(2,36) = 5.91, n = 19, P < 0.0001, Figure 5C); however, glutamate did not affect the single-unit activity of 3 fast-spiking cells that were included in these statistics. One fast-spiking cell responded with augmented single-unit firing and was excluded as a Grubbs statistical outlier (P < 0.05; cell no. 1 in Figure 5A).

In some instances, these ejections fully suppressed firing and were collected at both glutamate-sensitive and glutamate-insensitive locations in the cortex (n = 59 and 24, respectively) and striatum (n = 79 and 35, respectively, for MSNs). Additionally, we collected O2 data at fast-spiking interneuron locations (n = 16). For categorization purposes, we defined glutamate-sensitive cells as those that responded to glutamate to show this explicit chemical relationship between the majority of striatal, fast-spiking interneurons and glutamate in vivo in an intact brain.

**Transient O2 Events.** Finally, while characterizing glutamate responses in both cortical and striatal environments, we also observed spontaneous, discrete O2 decreases throughout both regions (Figure 6). These transient O2 events occurred without reliably concurrent cell activity, presented during baseline recordings in absence of glutamate, and persisted throughout data collections unaffected by glutamate ejections. The cyclic voltammograms revealed that the fluctuations were not noise because they only occurred at the O2 reduction potential. The overlapping relationships between these transients and glutamate-stimulated O2 decreases and single-unit activity between cortical and striatal environments are summarized in Figure 7. A total of n = 197 discrete data sets were collected at both glutamate-sensitive and glutamate-insensitive locations in the cortex (n = 59 and 24, respectively) and striatum (n = 79 and 35, respectively, for MSNs). Additionally, we collected O2 data at fast-spiking interneuron locations (n = 16). For categorization purposes, we defined glutamate-sensitive cells as those that responded to glutamate
ejections with increased single-unit activity, and transients as baseline-resolved $O_2$ changes that exceeded 3 standard deviations above baseline noise. Interestingly, 90.0% of all recorded locations with glutamate-sensitive cells presented with spontaneous $O_2$ transients. These transients appeared in 93.7% of striatal glutamate-sensitive recording locations (excluding interneurons) and 88.6% of striatal glutamate-insensitive locations (Figure 7). In contrast, spontaneous $O_2$ transients presented at 80.8% of glutamate-sensitive cortical cell locations but only 54.8% of cortical glutamate-insensitive locations (Figure 7). These data revealed the high probability of encountering spontaneous $O_2$ fluctuations in close proximity to glutamate-sensitive neurons, but further studies are needed to determine their significance.

**DISCUSSION**

In this study, we extended the use of an existing multimodal sensor to investigate the relationship between glutamatergic neurotransmission and $O_2$ responses in highly localized environments. We assessed the differences between stimulated (global, nonspecific) and iontophoresed (local, specific) glutamate responses. Endogenous glutamate release via electrical PFC stimulation elicited a biphasic $O_2$ pattern in the striatum, first decreasing and then increasing above baseline levels, consistent with the idea that $O_2$ is first locally consumed (event 1) following neurotransmission before being replenished via hyperemic CBF increases (event 2). In previous work, we showed that stimulated $O_2$ increases do not occur in brain slices devoid of CBF, supporting the idea that event 2 is hyperemic in origin. Striatal iGluR activation was responsible for a significant portion of the $O_2$ decrease amplitude, but its
negligible effect on the $O_2$ increase event (i.e., CBF) suggested that either stimulus strength, the release of other electrically evoked neurotransmitters, or both play this role in hyperemia. Local $O_2$ responses to glutamate iontophoresis resembled those from weak electrical stimulations, that is, monophasic $O_2$ decreases insufficient to provoke CBF-driven $O_2$ increases. Taken together, it is clear that our improved multimodal sensor provides a chemically and spatially selective method of studying exclusive glutameric influence on single-unit activity and local $O_2$ changes throughout the intact brain.

Here, we stimulated glutamergic cell bodies in the PFC to measure $O_2$ changes at terminals in the dorsal striatum. Typical electrical stimulation paradigms use biphasic pulses and optimize stimulation parameters through tuning curves. Studies have reported that below a threshold stimulation “strength”, both $O_2$ decreases and cell firing occurred without evoking CBF changes. We stimulated glutametric PFC projections to the striatum and evoked $O_2$ decreases that were only followed by $O_2$ increases past a threshold “strength” related to the number of stimulating pulses. The event amplitudes grew linearly with the number of stimulation pulses, as did the likelihood of the location responding with a biphasic $O_2$ pattern (i.e., first consuming $O_2$ then replenishing $O_2$ to increases above baseline). Together, these results support current functional hyperemia dogma that stimulating neuronal activity leads to increased energy consumption followed by overcompensating amounts of $O_2$ delivered via CBF. Thus, we verified that a stimulation threshold must be exceeded to elicit hyperemic $O_2$ responses in the corticostriatal pathway and identified that this threshold directly depends upon the number of electrical stimulus pulses.

In contrast to previous studies that show both CBF and neuronal activity as being dependent upon stimulation frequency, we found no relationship between higher stimulation frequencies and whether the locations expressed biphasic behavior. It is important to note that each of these studies kept the stimulation duration constant while modulating the frequency, which confounds frequency effects with the pulse numbers per stimulus. Further, we found no relationship between the event amplitudes and frequency, which we attribute to using a consistent number of pulses. Lowering the stimulation frequency only increased the amount of time between the stimulation onset and the maximal $O_2$ event responses, which again supported our conclusion that the number of pulses ultimately controls the magnitude of $O_2$ responses. Thus, we concluded that, for a constant number of electrical stimulus pulses, the PFC stimulation frequency has no influence over hyperemic $O_2$ amplitudes in the striatum.

Interestingly, we did not observe dopamine release at the sensor as a result of PFC stimulation, as has been reported in the ventral striatum; however, we positioned our electrodes in the dorsal striatum and optimized our sensor location based on robust glutamate-evoked single-unit activity. Additionally, the $O_2$ voltammetric waveform has poor sensitivity for dopamine (2.5 nA/μm at the dopamine oxidation potential [+0.6 V]). We observed catechol-like signals in some locations following stimulation, but the absence of a catecholamine reduction peak at −0.3 V and the presence of negative current at −0.1 V suggested that these signals indicated pH shifts.

While manipulating PFC stimulation parameters controlled whether striatal $O_2$ changes were mono- or biphasic, the specific role of glutamate as an excitatory neurotransmitter only influenced local $O_2$ decreases (i.e., event 1). Subsequent $O_2$ overcompensation (i.e., event 2) was insensitive to iGluR blockade at the cell terminals, consistent with studies that related neurovascular coupling to postsynaptic receptor activity. As a sufficient quantity of stimulation pulses was necessary to evoke $O_2$ overcompensation (i.e., CBF), the process responsible required either a larger recruitment of excited cells or robust, accumulative stimulation in order to release certain neurotransmitters. Meanwhile, glutamate iontophoresed at the same striatal locations as the PFC recordings only evoked $O_2$ decreases, even with high ejection currents (i.e., greater quantities of glutamate released). Extensive electrical PFC stimulations at similar frequencies have been shown to increase glutamate concentrations by several micromolar in the ventral striatum, orders of magnitude greater than what our iontophoretic ejections are estimated to release. The picomoles of glutamate iontophoresed with our 2 s ejection are unlikely to diffuse far enough to excite the same number of neurons as a stimulating electrode, but the magnitude of the $O_2$ decreases with iontophoresis were still comparable to “weaker” electrical stimuli (i.e., fewer pulses). Together, this suggests that local $O_2$ overcompensation at striatal terminals depended on PFC neurotransmission. However, the lack of specificity offered by electrical stimulation barred any conclusions about whether this hyperemic relationship was driven by neuronal activity or the stimulated release of vasoactive neurotransmitters, especially since the potential amount of striatal glutamate released versus ejected may differ by orders of magnitude.

It is well-known that electrical stimulations excite spatially extensive cell populations, so we adapted an existing multimodal sensor to excite local neuronal populations, record action potentials, and observe subsequent $O_2$ changes. These techniques have exceptionally high spatial resolutions; steady-state iontophoresis (i.e., ejections 120 s or longer) delivers drug to a radius in the hundreds of micrometers, extracellular single-unit recordings detect somatic firing within a ~50 μm radius, and FSCV at a carbon-fiber electrode probes $O_2$ changes within a 20 μm radius. A 2 s (i.e., far from steady-state) iontophoretic glutamate ejection, estimated to deliver 1–2 pmol of glutamate, evoked single-unit activity in highly localized cell populations, in agreement with previous studies. Nearly all recording locations at glutamate-sensitive cells (100+) showed either a monophasic $O_2$ decrease that returned to baseline or no $O_2$ change in response to the 2 s glutamate ejection (Figure 7; $O_2$ examples in Figure 4A,C). Glutamate iontophoresis evoked single-unit activity without subsequent $O_2$ changes and vice versa, in agreement with studies showing decoupling at high spatial resolution. However, the outstanding majority of glutamate-sensitive cells showed a decrease in $O_2$ after glutamate stimulation, similar to a "weak" electrical stimulus. Iontophoresed glutamate could not elicit biphasic responses, even with extended (>5 s) ejections using currents >500 nA (data not shown). Previous work established how to both quantify the drug ejected and track its spatial diffusion, which we used to confirm that a 2 s ejection released glutamate on the order of picomoles (vide supra). This is unlikely to be excitotoxic and establishes glutamate iontophoresis as a safe method of investigating local excitatory dynamics. It is unlikely that glutamate ejections of this magnitude could diffuse to a sufficient number of neurons and match the stimulation strength necessary to elicit a biphasic response, especially with the efficiency of glutamate uptake through neuronal and astrocytic transporters.
Cell firing is normally inhibited in deeply anaesthetized animals; however, we observed cells in the ventral striatum that fired spontaneously at high rates, a well-known characteristic of interneurons. Glutamate significantly and reversibly inhibited most of the recorded fast-spiking interneurons; however, glutamate had no effect on a small number of fast-spiking cells and increased cell firing only in one single case. While our sensor afforded us the unique advantage of recording interneuron responses to glutamate in an intact brain, technical limitations precluded our ability to definitively categorize the interneurons into their specific subtypes using single-unit activity waveforms alone.\(^{25,32}\) It is possible that the iontophoresed glutamate induces a temporary depolarization block, as has been previously observed with extended glutamate iontophoresis in dopamine neurons and ventral striatum neurons,\(^{45,44}\) which may explain why some neurons required longer ejections at stronger currents to quell firing (Figure 5B). The specific effect of glutamate on interneuron firing rates adds to the growing body of literature investigating the complex interactivity between neurotransmitters through striatal MSNs and fast-spiking interneurons.\(^{35,45,46}\)

Spontaneous O₂ events, found here in both the cortex and striatum, have been observed in vivo as early as 1957.\(^{47}\) Studies attributed these spontaneous events to factors ranging from the type of anesthetic used\(^{48}\) and to spontaneous astrocytic excitation,\(^{49,50}\) among other possibilities. Studies from Lowry et al. detected baseline current fluctuations when observing O₂ via amperometry, a technique that is also capable of determining absolute tissue O₂ concentrations.\(^{51}\) FSCV measures changes in O₂ rather than absolute concentrations; however, the unique voltammogram shape allowed us to confirm what may be mistaken for noise as O₂ (examples in Figure 1C). Though some transients were large (\(>20 \mu M\)), the short (<60 s) durations, continued neuronal sensitivity to glutamate excitation, and lack of concomitant ionic signals precluded these from representing spreading depolarizations.\(^{52}\)

When glutamate ejected during a transient event, we observed augmented O₂ decreases that returned to baseline on the same time scale as those without transients, suggesting that the transients are controlled by mechanisms that are independent of (or unlikely to compete with) neuronal glutamate receptor-dependent O₂ decreases. Indeed, spontaneous O₂ events did not always correspond with single-unit firing (Figure 6 and 7) despite the spatial resolution of O₂ detection being more confined than that of extracellular single-unit detection (\(\sim20\) and 50 \(\mu\)m radii, respectively).\(^{13,30}\) Nearly 30% of oxidative metabolism in the brain originates from astrocytes,\(^{53}\) undetectable with single-unit electrophysiology, making them attractive potential sources of these transients. That a larger percentage of transients were observed in deeper brain regions also indicated a possible environmental factor to their presence. The large overlap between locations with spontaneous transients and glutamate-sensitive neurons would make multiunit cellular recording an interesting future topic of exploration. Though we did not investigate any underlying chemical or physical source(s) of these transients, research from the Venton group has been investigating the relationship between adenosine and O₂ changes, both as stimulated events and as spontaneous transients.\(^{34,35}\)

As technology progresses to stimulate the brain more selectively and specifically, increasing evidence suggests that disconnects between functional hyperemia relationships exist in healthy subjects. Here, we elicited a hyperemic response from electrically stimulated glutamate release in the striatum, but targeted, exogenous glutamate excitation produced only decreases in recorded O₂. Modifying an existing multimodal sensor that paired iontophoresis with electrophysiology and FSCV permitted the simultaneous detection of highly localized O₂ and cell firing responses to selective glutamatergic stimuli in an intact, anaesthetized animal. This technique serves as a starting point for investigating spatially targeted cerebrovascular coupling as it relates to glutamate elicited neurotransmission and O₂ responses.

### Methods

#### Animal Care.

All animal protocols were approved by the Institutional Animal Care and Use Committee of the University of North Carolina at Chapel Hill (UNC) in accordance with the Guide for Care and Use of Laboratory Animals (eighth edition). Male Sprague–Dawley rats (300–450 g, Charles River, Wilmington, MA, USA) were pair-housed at UNC animal facilities, given food and water ad libitum, and kept on a 12 h light/dark cycle. Care was taken to reduce the number of animals used. A total of 6 animals were used for these experiments and glutamate iontophoresis data were pooled and analyzed from 70 animals originally used for other experiments. A single rat expired during a stimulation experiment, and was excluded from this study.

#### Surgery.

Animals were anesthetized with urethane (1.5 g/kg ip) and placed in a stereotaxic frame (Kopf, Tujunga, CA, USA). Coordinates are relative to bregma from the Paxinos and Watson (2007) atlas. Depths were measured relative to dura mater. Holes were drilled in the dorsal striatum (+0.7 or +2.0 mm A-P, +3.2 mm M-L, −2.5 to −3.5 mm D-V) for the carbon-fiber electrode and the PFC (+3.0 mm A-P, +0.8 mm M-L, −3.0 mm D-V) for the stimulating electrode. A Ag/AgCl reference electrode wire was placed in the contralateral hemisphere and served also as an iontophoresis ground. Holes were drilled for glutamate iontophoresis studies in the striatum (+2.2 mm A-P, +1.7 mm M-L, −6.4 to −7.8 mm D-V) and somatosensory cortex (+0.6 mm A-P, +2.8 mm M-L and +2.0 mm A-P, +2.6 mm M-L, −0.9 to −2.0 mm D-V for each).

#### Drugs and Solutions.

\(\text{dl-2-Amino-5-phosphonopentanoic acid sodium salt (APS, NMDA receptor antagonist), and 1,2,3,4-tetrahydro-7-nitro-2,3-dioxoquinoxaline-6-carbonitrile disodium salt (CNQX, AMPA receptor/kainate antagonist) were obtained from Abcam (Cambridge, MA, USA). All other chemicals were obtained from Sigma-Aldrich (St. Louis, MO, USA). Iontophoresis solutions of glutamate, APS, and CNQX were 200, 50, and 10 mM, respectively, with the latter two combined into a single cocktail solution. APS and CNQX concentrations were chosen based on existing literature, with a higher CNQX concentration to compensate for experimentally observed difficulties ejecting the drug at higher currents. Drugs were dissolved in 5 mM NaCl and ejected iontophoretically using cathodic currents (−25 to −400 nA).}

**Voltammetric O₂ Measurements and Iontophoresis.** Multi-modal sensors were fabricated from a four-barrelled glass capillary as described previously.\(^{15}\) Briefly, a ~5 \(\mu\)m diameter carbon fiber, pulled into a glass capillary, was inserted into a 4-barrel capillary and cut under a light microscope to an exposed length between 80–120 \(\mu\)m. The remaining iontophoresis barrels were filled with drug solutions. FSCV measurements were obtained and analyzed with the high-definition cyclic voltammetry (HDCV) computer program.\(^{14}\) The voltammetric waveform was 11 ms in duration, first scanned from a holding potential of 0.0 V to +0.8 V, then to −1.4 V, and finally back to the holding potential (vs Ag/AgCl reference electrode) at a scan rate of 400 V s\(^{−1}\). Carbon-fiber surfaces were conditioned in the brain by scanning the waveform at 60 Hz, followed by 10 Hz, each for 15 min. Simultaneous electrochemical and electrophysiological measurements were acquired with a 5 Hz repetition rate as described previously.\(^{14}\) Briefly, each 200 ms data acquisition period consisted of single-unit activity recordings for 179 ms, a voltammetric waveform lasting 11 ms, and a 0 V potential held both prior to and following
each waveform for 5 ms each. O2 current versus time traces were exported at the O2 reduction potential on the forward scan (−1.35 V) to exclude the nonfaradic current that immediately follows switching potentials from adsorbing species that disrupt the electrical double-layer capacitance (e.g., calcium).15 O2 concentrations were estimated from the current at the O2 reduction potential (−1.35 V) using a −0.35 nA μM−1 calibration factor, such that negative currents indicate O2 increases and positive currents indicate O2 decreases. The calibration methods used have been described elsewhere,12 where the O2 reduction current is measured from solutions of known O2 content passing through an airtight flow cell. Custom instrumentation controlled the alternating connection of the carbon fiber between a current transducer (cyclic voltammetry) and a voltage follower (electrophysiology measurements).24

Iontophoretic ejections were performed by applying constant current (NeuroPhone BH-2 System, Harvard Apparatus, Holliston, MA, USA). Ejection timing was controlled by HDCV. Iontophoretic barrels were primed at least 400 μm dorsal to the measurement region, outside the brain regions of interest, to test barrel functionality without affecting receptors at the actual recording locations.

**Electrically-Stimulated Endogenous Glutamate Release.** Stimulus isolators (NL800A, Neurolog, Digitimer, Hertfordshire, UK) generated electrical stimulations at a bipolar stainless steel electrode (Plastics One, West Lafayette, IN, USA) in the PFC glutamatergic cell bodies. A multimodal sensor was placed in the ipsilateral dorsal striatum. One drug barrel contained glutamate (200 mM) while another contained an APS (50 mM) and CNQX (10 mM) cocktail to antagonize iGluRs. Recordings were taken at locations where glutamate reponsive cells were present, defined as locations where single-unit activity accompanied a 2 s glutamate ejection using cathodic currents <100 nA. The electrode depth was optimized for the strongest biphasic response while maintaining sensitivity to iontophoresed glutamate. One location was used per animal. Dorsal striatum locations were verified with histology as described below.

Stimulations were optimized to maximize biphasic O2 responses while maintaining physiologically relevant stimulus lengths. Stimulation frequencies from 5 to 60 Hz were tested; all stimulations were comprised of 120 biphasic pulses of 0.2 ms width and 300 μA amplitude. These frequencies were chosen with the knowledge that high frequency corticostriatal stimulations raise striatal glutamate levels57 and are within the range of stimulations used in other studies.5,58 To measure the effect of stimulation duration, we held stimulation frequency and current amplitude constant (20 Hz and 300 μA, respectively) and varied pulse numbers from 10 to 100. A random number generator controlled the order in which different frequencies or pulse numbers were used.

Pharmacological studies investigated the role of local iGluRs on O2 events in the striatum following PFC stimulation. To antagonize NMDA, AMPA, and kainate receptors, a drug cocktail (50 mM APS and 10 mM CNQX) was iontophotically ejected into the dorsal striatum recording location for 60 s using cathodic current. The ejection duration was chosen to reach near-steady state diffusion, such that local (≈100 μm radius)25 receptors would become saturated with antagonists. After the drug ejections ended, 10–30 s elapsed before the postdrug PFC stimulation (20 Hz, 80 pulses) to measure O2 changes under local iGluR antagonism. Glutamate was iontophoresed before the first electrical stimulation in each subject and again 10 min after the end of the final drug cocktail stimulation to confirm drug clearance.

Local Glutamatergic Excitation with Iontophoresis. A multimodal sensor was placed in either the cortex or the striatum, with one barrel containing a glutamate solution (200 mM glutamate dissolved in 5 mM NaCl). Each probe was used for multiple locations, if background currents remained stable, to both avoid tissue damage and minimize the number of animals. Recording locations were at least 300 μm apart.25 Both the electrophysiological software and HDCV continuously display data in real time, which allowed us to locate glutamate-sensitive cells quickly. The sensor was lowered into the brain region of interest and glutamate was ejected (2 s). Locations were designated as cell locations if they produced reproducible single-unit firing that time-locked to the glutamate ejection. If no time-locked response occurred, the sensor was lowered further and the glutamate ejection repeated until a cell was located. We optimized the depth of cell recording locations for maximal single-unit activity responses to glutamate. Because multiple data sets could be collected per animal, data sets were also taken at locations that were insensitive to glutamate to characterize other metrics (i.e., O2 responses and spontaneous O2 transients, vide infra). Each recording location data set consisted of 3–6 replicates each of baseline O2 activity (nothing ejected) and 2 s glutamate ejections. Time between subsequent glutamate ejections was 120 s.

**Single-Unit Activity Electrophysiology.** Single-unit activity was measured at the carbon-fiber for 179 ms between each voltammetric scan. The data were amplified (×5000), fed through a bandpass filter (300–3000 Hz, Krohn-Hite Corp., Brockton, MA), and digitized using commercially available software (Digitizer, Plexon, Dallas, TX, USA). Single units were analyzed using Offline Sorter (Plexon, Dallas, TX, USA). Individual waveforms coinciding with a coupling artifact were excluded. After cell sorting, peri-event rasters depicting the firing rate of cells over time were compiled using commercially available software (NeuroExplorer, Nex Technologies, Madison, AL) and exported in 0.18 s bins for statistical analysis.

**Histology.** Following data collection, the probe location was lesioned by cyclically applying a ramp of 0–10 V DC potential three times over 20 s. Animals were euthanized with urethane cardiac puncture. Brains were removed and fixed in 10% formalin for >7 days. Brains were then cryoprotected in 30% sucrose for >48 h, before coronal sections (50 μm) were taken with a freezing microtome (Leica, Germany). Slices were mounted on microscope slides and viewed under a light microscope to confirm electrode placements.

**Statistical Analyses.** Statistical analysis was performed using GraphPad Prism 6 (GraphPad Software, San Diego, CA, USA). A repeated measures one-way analysis of variance (ANOVA) with post hoc Bonferroni’s test was used to evaluate significance between different electrical stimulation parameters. A Student’s t-test was used to determine significance between pre- and postdrug O2 events elicited by electrical stimulation. Differences were significant when *P < 0.05. Venn diagrams were generated using Venn Diagram Plotter software (Pacific Northwest National Laboratory, Richland, WA, USA; [https://omics.pnl.gov/](https://omics.pnl.gov/)).

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**Author Contributions**

L.R.W. drafted the manuscript, designed experiments, and carried out the acquisition, analysis, and interpretation of data and statistics. N.G.B. acquired and analyzed data. S.C. acquired data and designed instrumental setup. R.M.W. germinated the scope of research and designed experiments. All authors contributed to the editing of this manuscript.

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ABBREVIATIONS

ANOVA, analysis of variance; APS, DL-2-amino-5-phosphono-pentanoic acid; ATP, adenosine triphosphate; BOLD, blood oxygenation level-dependent; CBF, cerebral blood flow; CNQX, 1,2,3,4-tetrahydro-7-nitro-2,3-dioxiquinoxaline-6-carbonitrile; fMRI, functional magnetic resonance imaging; FSCV, fast-scan cyclic voltammetry; HDCV, high-definition cyclic voltammetry; iGluRs, ionotropic glutamate receptors; MSN, medium spiny neuron; PFC, prefrontal cortex; UNC, University of North Carolina at Chapel Hill

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