Predominant Enhancement of Glucose Uptake in Astrocytes versus Neurons during Activation of the Somatosensory Cortex

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Glucose is the primary energetic substrate of the brain, and measurements of its metabolism are the basis of major functional cerebral imaging methods. Contrary to the general view that neurons are fueled solely by glucose in proportion to their energetic needs, recent in vitro and ex vivo analyses suggest that glucose preferentially feeds astrocytes. However, the cellular fate of glucose in the intact brain has not yet been directly observed. We have used a real-time method for measuring glucose uptake in astrocytes and neurons in vitro and in vivo and ex vivo in male rats by imaging the trafficking of the nonmetabolizable glucose analog 6-deoxy-N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl)-aminoglucose (6-NBDG) using two-photon microscopy. During resting conditions we found that astrocytes and neurons both take up 6-NBDG at the same rate in the barrel cortex of the rat. However, during intense neuronal activity triggered by whisker stimulation, astrocytes rapidly accelerated their uptake, whereas neuronal uptake remained almost unchanged. After the stimulation period, astrocytes returned to their preactivation rates of uptake paralleling the neuronal rate of uptake. These observations suggest that glucose is taken up primarily by astrocytes, supporting the view that functional imaging experiments based on glucose analogs extraction may predominantly reflect the metabolic activity of the astrocytic network.

Introduction
Glucose is the main source of energy for the brain (Sokoloff, 1977). The spatiotemporal coincidence of glucose utilization with localized neuronal activation, or “neuro-metabolic coupling” underlies the interpretation of widely used functional imaging methods of the human brain (Raichle and Mintun, 2006). Thus, a significant portion of our knowledge of intact brain function depends on an understanding of localized glucose processing by brain cells. Because clinically used imaging tools do not possess cellular-scale resolution, studying the fate of glucose in the brain parenchyma after it has left the vascular compartment represents a challenging task (Nehlig and Coles, 2007). The traditional method for measuring glucose metabolism is the autoradiography of brain slices obtained from animals perfused in vivo by systemic injection of 2-deoxyglucose (2-DG) (Sokoloff, 1977), which at best provides submillimeter resolution. The observation that 2-DG labeling in the cortex is less intensely present in neuronal somata than in the neuropil, which is rich in synapses and astrocytes, has led to the proposition that astrocytes, despite their apparent passivity, may be important users of glucose (McCasland, 1996). Astrocytes’ avidity for glucose has been observed in tissue culture where the basal rate of glucose utilization is 3–10 times higher than in neurons (Lopes-Cardozo et al., 1986; Magistretti and Pellerin, 1996a). These ex vivo or in vitro findings converge on the view that neurons, despite their higher energetic needs (Attwell and Laughlin, 2001), may have a much lower glucose utilization than astrocytes (Herrero-Mendez et al., 2009). However, this is contradicted by other studies showing that neurons are the primary site of glucose uptake and utilization (Brookes and Yarowsky, 1985; Itoh et al., 2004). Yet, direct in vivo evidence of such a compartmentalization of glucose utilization, if any, is still lacking (Hyder et al., 2006; Nehlig and Coles, 2007). 2-DG analyses, using higher spatial-resolution methods combined with immunohistochemical labeling, indicate that neurons and glia may have at rest an equal share of glucose utilization (Nehlig et al., 2004).

To address the role of glucose in fueling neuronal activity, more direct methods in the living brain are needed (Nehlig and Coles, 2007). We hypothesized that the nonmetabolizable fluorescent glucose analog 6-deoxy-N-(7-nitrobenz-2-oxa-1,3-diazol-4-yl)-aminoglucose (6-NBDG) (Speizer et al., 1985), combined with high spatial-resolution, real-time two-photon microscopy, would be effective in addressing this issue in the living brain. Fluorescent glucose analogs have been increasingly used for studying glucose transport and metabolism in many cell types besides astrocytes and neurons (Shimada et al., 1994; Porras et al., 2004; Rouach et al., 2008). Here, we examine the kinetics of 6-NBDG uptake into astrocytes and neurons in the rat barrel cortex, during both resting conditions and sensory stimulation.

Materials and Methods
Surgical procedure. All experimental procedures were performed according to the animal use and welfare guidelines of Rutgers University. Male...
Sprague Dawley rats (50–100 g) were anesthetized with urethane (1.7 g/kg, i.p.). Animals were tracheotomized and mechanically ventilated. The pulmonary volume was set, and blood gas components were monitored (PaCO2 between 35 and 45 mmHg, PaO2 between 100 and 200 mmHg, pH between 7.35 and 7.50). The femoral artery was cannulated to permit vascular access and the monitoring of the mean arterial pressure (maintained between 75 and 100 mmHg). The rectal temperature was monitored and kept close to 37.5°C with a heating pad (Harvard Apparatus). A steel frame (1.5 cm × 6 cm) with a hole (4 mm diameter) centered over the right barrel cortex (2.5 mm posterior to the bregma and 4.9 mm lateral from midline) was attached on the skull with dental cement. A craniotomy was performed, and the dura mater was removed. After the labeling procedure for astrocytes, the craniotomy was filled with agarose (Sigma; type III-A, 0.75% in a HEPES-buffered solution) and covered with a coverslip immobilized with dental cement.

Labeling of astrocytes and 6-NBDG administration. Astrocytes were labeled by bathing the cortical surface with SR-101 (1 μM) in HEPES buffer, pH 7.42 (Sigma), for 10 min (Nimmerjahn et al., 2004). 6-NBDG (Invitrogen) was dissolved in a solution of 55% of HEPES buffer and 45% of DMSO, pH 7.42, to a concentration of 5 mg/ml. The femoral vein was cannulated, and 6-NBDG was injected with an infusion pump at a rate of 100 μl/min/100 g for bolus experiments or 20 μl/min/100 g for continuous infusion experiments.

Whisker stimulation. The whiskers on the left of the animal’s snout were stimulated with puffs of air produced by a picospritzer (Harvard Apparatus) coupled to a stimulator (Master 8; DL Instrument International). Air was ejected at 1 bar via plastic tubing (1.5 mm diameter) positioned ~1 cm lateral and anterior to the animal’s nose to stimulate the whole whisker pad. Whiskers were stimulated at 5 Hz for 30 s (pulse width ~100 ms) repeated five times with periods of 30 s of rest between each block of stimulation. Online display of the local field potential (LFP) was used to monitor the presence of increased spindle-like activity during the whisker stimulations.

EEG recording. A tungsten wire (50 μm in diameter) was inserted close (≤500 μm) to the imaging area and sealed underneath the coverslip window to record the LFPs in layer II–III of the barrel cortex. The neurophysiological signal was acquired at 2 kHz with a DAM-80 AC amplifier (WPI) and Data-Trax acquisition software (WPI). The wide-band signal was downsampled to 1250 Hz and used as LFP signal. Data analysis was performed with custom-written Matlab software (The Mathworks). The LFP displayed a slow oscillation activity during anesthesia, and the amount of urethane was adjusted for keeping the plane of anesthesia at constant level. Spindle-like activity was observed spontaneously and was associated with a deflection of the contralateral whiskers to craniotomy but not in response to the stimulation of the ipsilateral whiskers (Fig. 1c).

Two-photon laser scanning microscopy. Imaging was performed with a custom-built two-photon laser (Chameleon; Coherent) scanning microscope consisting of an Olympus trinocular, Cambridge scan mirrors, and a XLUMPFLN-20XW Olympus objective. Fluorescence emission was directed by a 700-LP dichroic, split with a 565-LP dichroic into green and red channels, and further restricted with BGG22 and HQ607/45 filters placed before the green and red photomultipliers (Hamamatsu), respectively. Frames (512 × 512 pixels) were collected at a frequency of 0.016 Hz (bolus experiment) or 0.32 Hz (activation experiment).

Figure 1. 6-NBDG two-photon microscopy experimental set-up. a, Detail of the cranial window preparation mounted over the barrel cortex. A tungsten wire (50 μm diameter) was inserted close to the imaging area to record the LFP in layer II–III of the barrel cortex. Agarose filled the chamber to stabilize the brain from cardiorespiratory movements. Astrocytes were labeled using SR-101. b, Characterization of 6-NBDG two-photon excitation in vivo. c, Experimental arrangement for simultaneous LFP recording and two-photon imaging during whisker stimulation. 6-NBDG was infused to the femoral vein. The whiskers of the contralateral side were mechanically stimulated by air puffs generating sustained network activity in the LFP. The time-frequency spectrogram of the LFP showed an increase in the LFP power in the 7–20 Hz range during each period of 30 s of whisker stimulation.
Data analysis. Neurons and astrocytes were distinguished by using the red signal emitted by SR-101. Astrocytes concentrate SR-101, and their somata appear intensely bright. In contrast, neurons exclude the dye and appear as black holes (Nimmerjahn et al., 2004). Each image of the sequence was aligned over the previous image with Align Slice (ImageJ, National Institutes of Health) to correct x–y deviation caused by possible drift of the tissue. Fluorescence intensity was measured in a region of interest (average intensity over 250 pixels) strictly limited to the somatic area. Signals were expressed as relative fluorescence changes \( \frac{F}{F_0} \), where \( F_0 \) was the mean of the baseline period. Slope was determined after the equation of the linear regression line through the data points.

Statistics. The results are presented as mean ± SEM. Statistical analyses were performed with an unpaired Student’s t test after the demonstration of homogeneity of variance with a one-way ANOVA or a repeated-measure ANOVA. The Bonferroni test was used for post hoc testing. The significance level was set at \( p = 0.05 \) (two-tailed).

Results

Uptake of 6-NBDG in resting condition

6-NBDG in solution was best excited at wavelengths <820 nm (Fig. 1b). We set the excitation wavelength at 810 nm to stray from the autofluorescence zone around 830 nm. The barrel somatosensory cortex was imaged at a depth of 250–350 μm on a field containing several astrocytes and neurons (2.1 ± 0.6 and 2.1 ± 0.5 cells per field of imaging, respectively) and at least one blood vessel to control for the 6-NBDG arrival into the local microcirculation (Fig. 2a). A bolus of 2 mg/100 g of 6-NBDG was injected intravenously at a rate of 100 μl/min/100 g. Considering that the blood volume (in milliliters) is 7–8% of body weight and assuming a normal glycemia (0.38–0.55 mmol/100 ml of blood), the equivalent increase of blood glucose at the end of the injection of 6-NBDG was 0.07 mmol/100 ml. Once the bolus started, fluorescence could readily be seen filling up the local vascular tree. For the measurement of 6-NBDG fluorescence intensity, a square region of interest was chosen inside one or more vessels; for both cell types this square was limited strictly to the somatic area. As NBDG fluorescence progressively invaded the brain parenchyma, the whole field of observation appeared in a homogeneous manner without a visually obvious “hot spot” (supplemental video, available at www.jneurosci.org as supplemental material, and Fig. 2a). Astrocytes and neurons (six animals, 13 astrocytes, and 13 neurons) showed similar kinetics of uptake over 45 min observation period (Fig. 2c).

Uptake of 6-NBDG during whisker stimulation

The aim of the second set of experiments was to analyze the kinetic of 6-NBDG uptake in astrocytes and neurons during physiological activation (Fig. 1c). After a stabilization period of 15 min, 6-NBDG (5 mg/ml) was injected into the femoral vein at the constant flow of 20 μl/min/100 g of body weight until the end.
We present a novel approach for studying brain glucose uptake \textit{in vivo} with high spatial and temporal resolution. We show that the glucose analog 6-NBDG is detectable in the brain at a subcellular scale, seconds after its intravenous administration at physiologically relevant concentrations. Our study is the first real-time imaging of the cellular supply of a glucose analog in the intact brain. We found that 6-NBDG homogeneously invades the parenchymal compartment and reaches astrocytes and neurons at the same rate in the absence of triggered neuronal work. This is in agreement with previous \textit{ex vivo} observations showing that 45 min after its systemic injection the radioactive glucose analog 2-DG is detected equally in astrocytes and neurons (Nehlig et al., 2004). 6-NBDG is a nonmetabolizable analog of glucose whose uptake is affected only by glucose transporter activity. Fluorescent glucose analogs enter cells through glucose transporters with an affinity 300 times higher than D-glucose, albeit at a much slower rate (Yamada et al., 2000; Barros et al., 2009). It has been demonstrated \textit{in vitro} that glucose inhibits 6-NBDG uptake with the same efficiency in astrocytes and neurons (Aller et al., 1997), suggesting that 6-NBDG would compete with glucose equally for transport into the two cell types. However, the concentration of 6-NBDG used here represents only 5% of the glycemia. It is therefore unlikely that 6-NBDG would compete with blood-borne glucose and would modify the kinetic of its transport and metabolism.

The barrel cortex is among the best characterized sensory areas of the rat brain. Activation of this cortical area generates a well understood neuronal network activity (Contreras and Steriade, 1996). Because this system produces attenuation of neuronal responses to sustained whisker deflection (Chung et al., 2002), we avoided habituation by stimulating the whiskers intermittently at a low frequency. This approach produced a steady level of activation as evidenced by spectral analysis of LFP and increased firing of experiment to create a steady delivery of 6-NBDG to the brain (Fig. 3a). As in the bolus experiment described above, the two cell types showed similar uptake rate with no significant difference between slopes (0.06 ± 0.04 for astrocytes and 0.05 ± 0.03 for neurons; \(p > 0.05\)) (Fig. 3a,c) and between fluorescence amplitude (3.57 ± 0.68% for astrocytes and 2.15 ± 0.57% for neurons; \(p > 0.05\)) (Fig. 3d) in the baseline condition. Whisker stimulation started 10 min after the beginning of 6-NBDG infusion, whereas the loading rate in the parenchyma showed a steady slope (slope changes not significantly different from 0 after 6 min of infusion; \(n = 4\) animals). Air puffs triggered a sustained evoked responses in the LFP recorded in layer II–III of the barrel cortex, also reflected by an increased LFP power in the 7–20 Hz range (Fig. 1c). Such an increase of network activity was strictly observed during stimulation of the contralateral whisker pad relative to the recording site, but not during the stimulation of the ipsilateral whisker pad. Arterial pressure did not show consistent changes during the activation. During the stimulation of the whisker pad, 6-NBDG uptake slope significantly increased in astrocytes (from 0.06 ± 0.04 to 0.22 ± 0.04; \(p < 0.001\)) but nonsignificantly in neurons (from 0.05 ± 0.03 to 0.11 ± 0.03; \(p > 0.05\); six animals, 31 astrocytes, and 31 neurons). After the stimulation, both slopes decreased to their prestimulation level. There was no difference between the fluorescence intensity level of astrocytes and neurons just before the stimulation (mean >75 s) (see Fig. 3d). However, just after the stimulation, astrocytes showed a much greater accumulation of 6-NBDG than neurons (mean >75 s; \(p < 0.001\)) (Fig. 3b).

**Discussion**

We present a novel approach for studying brain glucose uptake \textit{in vivo} with high spatial and temporal resolution. We show that the...
mediated differential enhancement of glucose uptake in astrocytes essentially took place in astrocytes, whereas the neuronal extraction rate displayed only a tendency to increase. This activation-mediated differential enhancement of glucose uptake in astrocytes and neurons has been reported previously in several in vitro studies. As pioneers in the use of fluorescent glucose to investigate brain glucose metabolism, Barros and coworkers have shown in vitro that glutamate-induced neuronal firing surprisingly mediates acute inhibition of glucose transport in neurons while stimulating astrocytic glucose uptake (Loaiza et al., 2003; Porras et al., 2004). Thus, it seems likely that during neuronal activation there is a redistribution of glucose away from neurons and toward astrocytes. In line with this, a novel computation of experimental datasets from rodents and humans predicted that when the activity level of the brain increases the proportion of glucose entering into glia also increases (Jolivet et al., 2009). In addition, further work suggests that in acute cerebellar slices Bergmann glial cells transport and metabolize fluorescent hexoses faster than both Purkinje and granule cells (Barros et al., 2009). Therefore, during activation, in addition to an increase in the delivery of glucose caused by local hyperemia, those results are suggestive of an active regulation of the transport of glucose into astrocytes and neurons.

We have to point out two potential caveats in the interpretation of our results. First, it is possible that anesthesia modifies the rules of glucose extraction and trafficking. Future studies should be conducted either with a different class of anesthetic agents or, preferably, without anesthesia to confirm the present findings (Oz et al., 2004). Second, most energy metabolism may take place at the level of synapses rather than at or around the cell bodies of astrocytes and neurons, where our measurements were taken. However, given its high diffusion rate, 6-NBDG is expected to distribute homogeneously in all compartments of the cell (Barros et al., 2009).

Whatever the mechanism of glucose distribution between astrocytes and neurons, our data raise the question of whether brain cells take up glucose in proportion to their energetic needs. The energetic needs of astrocytes and neurons is still an open debate because calculations critically depend on various assumptions and experimental conditions. At rest we found fluorescence intensity of 6-NBDG to be similar in astrocytes and neurons, suggesting the need for glucose is the same in both cell types. In rats, the volume occupied by astrocytes in vivo is estimated to be ~30% of the cortical gray matter cellular volume (not counting extracellular space) (Norenberg, 1994; Hertz, 2008). According to the measurement of acetate oxidation, which is selectively oxidized by astrocytes, oxidative metabolism of astrocytes also accounts for ~30% of total brain cell oxidative metabolism in rodents (Sibson et al., 2001; Oz et al., 2004; Cruz et al., 2005; Patel et al., 2005). Therefore, the level of oxidative metabolism in astrocytes and neurons may not differ greatly in resting conditions consistent with similar 6-NBDG uptake found here in both cell types. During stimulation, the parallel uptake of glucose analog by astrocytes and neurons is no longer observed: a vast majority of the additional glucose taken up by astrocytes. A first possible interpretation is that astrocytes take up more glucose during functional activation because they spend more energy than active neurons. In support of this, it has been shown that the oxidative metabolism of astrocytes increases during brain activation, suggesting the astrocytic tricarboxylic acid cycle is not saturated in resting condition and may support the extra work resulting from neuronal activation (Cruz et al., 2005; Wyss et al., 2009). Interestingly, it has been proposed that the multitude of astrocytic tasks unraveled such as active clearance of K⁺ (Somjen et al., 2008) or fast and local calcium signaling (Chuquet et al., 2010) could altogether account for a high energetic expense and therefore a high need of glucose by astrocytes (Hertz et al., 2007).

However, although astrocytic energy needs could have been underestimated, it is unlikely that their energy expenditure overcomes the neuronal one. A general consensus is that astrocytes contribute significantly less to ATP production than neurons during spiking activity (Attwell and Laughlin, 2001; Hyde et al., 2006; Jolivet et al., 2009). Therefore, our demonstration that glucose is preferentially taken up by astrocytes during enhanced neuronal activity represents a paradox. Considering that glucose is the almost exclusive energy substrate used by the brain, one explanation to resolve this paradox lies in the proposal that lactate, derived from the glucose extracted by astrocytes, may be synthesized and delivered to neurons to complement their energy budget (Magistretti and Pellerin, 1999; Pellerin et al., 2007). The corollary of this is that neurons may use less glucose than predicted by their energy requirement in agreement with the present finding. Indeed, neurons have a lower glycolytic rate than astrocytes (Almeida et al., 2001), which has been explained by the finding that the enzyme 6-phosphofructo-2-kinase/fructose-2,6-biphosphatase-3, a key activator of glycolysis, is active only in astrocytes (Herrero-Mendez et al., 2009). The present data are compatible with the astrocyte-to-neuron shuttle hypothesis originally conceptualized by Magistretti and Pellerin (1999) (Hyder et al., 2006) and provide the first in vivo evidence of a preferential transport of glucose to astrocytes when neuronal activity is increased. Nevertheless, the contribution of astrocytes to support neuronal energetics is likely to depend on the stimulus paradigm. The flux of lactate from astrocytes to neurons has been suggested to be reversible, depending on the redox state of each cell (NAD+/NADH) (Cerdán et al., 2006; Mangia et al., 2009).

In summary, in vivo 6-NBDG two-photon microscopy represents an effective approach for the study of glucose trafficking in the intact brain. The present study described the uptake kinetics of a glucose analog in astrocytes and neurons in vivo. Under “basal” cortical activity both cell types take up 6-NBDG at the same rate. During intense firing triggered by a sensory stimulation, the uptake is accelerated mainly in astrocytes, supporting the hypothesis that astrocytes play a central role in brain energetics (Voutsinos-Porche et al., 2003). An indirect implication of our findings is that signals detected in functional imaging experiments, using glucose analogs such as in positron emission tomography or 2-DG autoradiography, may predominantly reflect the metabolic activity of the astrocytic network (Magistretti and Pellerin, 1996b).

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