Deciphering neuron-glia compartmentalization in cortical energy metabolism

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Energy demand is an important constraint on neural signaling. Several methods have been proposed to assess the energy budget of the brain based on a bottom-up approach in which the energy demand of individual biophysical processes are first estimated independently and then summed up to compute the brain’s total energy budget. Here, we address this question using a novel approach that makes use of published datasets that reported average cerebral glucose and oxygen utilization in humans and rodents during different activation states. Our approach allows us (1) to decipher neuron-glia compartmentalization in energy metabolism and (2) to compute a precise state-dependent energy budget for the brain. Under the assumption that the fraction of energy used for signaling is proportional to the cycling of neurotransmitters, we find that in the activated state, most of the energy (~80%) is oxidatively produced and consumed by neurons to support neuron-to-neuron signaling. Glial cells, while only contributing for a small fraction to energy production (~6%), actually take up a significant fraction of glucose (50% or more) from the blood and provide neurons with glucose-derived energy substrates. Our results suggest that glycolysis occurs for a significant part in astrocytes whereas most of the oxygen is utilized in neurons. As a consequence, a transfer of glucose-derived metabolites from glial cells to neurons has to take place. Furthermore, we find that the amplitude of this transfer is correlated to (1) the activity level of the brain; the larger the activity, the more metabolites are shuttled from glia to neurons and (2) the oxidative activity in astrocytes; with higher glial pyruvate metabolism, less metabolites are shuttled from glia to neurons. While some of the details of a bottom-up biophysical approach have to be simplified, our method allows for a straightforward assessment of the brain’s energy budget from macroscopic measurements with minimal underlying assumptions.

Keywords: astrocyte-neuron interactions, brain energy metabolism, brain energy budget, glucose, oxygen

INTRODUCTION
The energy requirements of the brain are amazingly high; indeed, while representing only about 2% of the body weight, its oxygen and glucose utilization account for approximately 20% of those of the whole organism, almost 10 times more than what would be predicted on a weight basis (Kety, 1957; Magistretti, 2008; Rolfe and Brown, 1997; Sokoloff, 1960). A similar mismatch is also observed for blood flow destined to the brain, which represents over 10% of cardiac output. A central feature of brain energy metabolism is that it is tightly coupled to neuronal activity. This tight coupling is at the basis of functional brain-imaging techniques, such as positron emission tomography and functional magnetic resonance imaging (Magistretti and Pellerin, 1999). However, the exact principles underlying this coupling and in particular the nature and role of neuron-glia metabolic interactions are still a subject of active research (Gordon et al., 2008; Rouach et al., 2008) and of controversy at times (Hertz et al., 2007). Similarly, the precise energy budget of the brain and in particular the cost associated with neuronal signaling is still not well understood (Attwell and Laughlin, 2001; Lennie, 2003).

Neuron-glia metabolic interactions have been the subject of several recent modeling studies (Aubert et al., 2005, 2007; Hyder et al., 2006). Most of these studies tend to support the presence of an activity-regulated lactate shuttle from astrocytes to neurons as originally proposed by Pellerin and Magistretti (1994). The presence of this shuttle has recently been reported in slices (Rouach et al., 2008). Similarly, several mathematical methods have been derived to assess the energy budget of the brain. These methods are traditionally based on a bottom-up approach in which the energy demand of individual biophysical processes are first estimated independently and then summed up to compute the brain's total energy budget (Attwell and Laughlin, 2001; Lennie, 2003; Nawroth et al., 2007). Most of these methods focus on energy requirements rather than on energy supply.

Here, we introduce a novel top-down approach that addresses the question of the brain’s energy budget from the opposite perspective. We re-analyzed datasets reporting average glucose and oxygen utilization in the human and rodent brains in different activation states (Gjedde, 2007). Using these figures, it is possible to evaluate the average tissue ATP production corresponding to the different
activation states. We then proceed to show that neuronal versus glial (i.e. astrocytic) oxygen and glucose metabolism can be evaluated by combining this first dataset with data reporting the rate of neuronal glucose-equivalent oxidation as a function of the rate of cycling of neurotransmitters (Hyder et al., 2006; Sibson et al., 1998). This can be achieved with minimal underlying hypotheses. It is then possible to compute the brain’s energy budget as well as the cycling rate of neurotransmitters corresponding to the different activation states. Finally, using our prediction of the compartmentalization of neuronal versus astrocytic glucose and oxygen utilization, it is possible to evaluate whether these two cell populations completely oxidize the glucose they take up or rather import or export glucose-derived metabolites.

Our results suggest that a significant part of glucose is taken up by astrocytes while oxygen is mostly consumed within the neuronal population. We find that this bias tends to increase with the level of activation. The higher the activation state, the higher the proportion of oxygen being used by neurons and the higher the proportion of glucose entering glia. Moreover, our approach yields an energy budget for different brain states. At high levels of activation, we find that about 80% of energy is produced to support neuronal signaling while glia only use up a small fraction of energy to recycle neurotransmitters (5–6%). Finally, we find that the glial population takes up more glucose than it can oxidize while the neurons oxidize more glucose-derived metabolites than the amount of glucose they take up. This is suggestive of a transfer of glucose-derived metabolites from the astrocytes to the neurons. Furthermore, the amplitude of this transfer is shown to be proportional to the level of activity as measured by the cycling of neurotransmitters, a finding consistent with the astrocyte-neuron lactate shuttle hypothesis (Pellerin and Magistretti, 1994). While lacking the details of a more biophysical approach (Attwell and Laughlin, 2001; Nawroth et al., 2007), our approach yields an energy budget that matches observed tissue glucose and oxygen utilization as well as the compartmentalization of glucose transport and oxygen utilization between neurons and astrocytes across different levels of brain activation. The resulting budget is in agreement with the one detailed in the work of Attwell and Laughlin (2001) and at the same time supports the view of all glial cells are pooled together and are hereafter referred to as astrocytes), (ii) that the system is at steady-state which was also assumed during the collection of the two datasets to be used and (iii) that energy measured as adenosine triphosphate production (ATP) is used in the compartment where it is produced.

Average CMR_{glc} and CMR_{o2} lead to the average tissue (T) ATP yield for each state S = 1,…,4 through the following stoichiometric relation (Gjedde, 2007; Gjedde et al., 2002)

\[ V_{ATP}^{T}(S) = 2CMR_{glc}(S) + \lambda CMR_{o2}(S) \]  

(1)

Theoretically, \( \lambda = 6 \) yielding a total of 38 ATP molecules per glucose molecule in the case of complete oxidation. Here, we use \( \lambda = 28/6 \) for a total of 30 ATP molecules, a yield closer to experimental measurements (Berg et al., 2006; Hinkle et al., 1991). It is worth noting however, that we find no significant difference in the results reported when using \( \lambda = 6 \).

As reviewed in Hyder et al. (2006), the total cycling of neurotransmitters \( V_{cyc}^{t} \), and the neuronal oxidative glucose utilization \( CMR_{glc}^{n} \) are linked by

\[ CMR_{glc}^{n} = AV_{cyc}^{t} + B \]  

(2)

with A and B parameters to be determined. As shown in Jolivet et al. (2009), assuming most of glucose entering the tricarboxylic acid (TCA) cycle in neurons is ultimately oxidized (Gjedde, 2007), Eq. 2 can be rewritten

\[ V_{cyc}^{t} = A^{-1}\left(\frac{1}{6}(1-r_{o2})CMR_{o2} - B\right) \]  

(3)

with \( r_{o2} \) the astrocytic contribution to oxygen utilization. All CMRs in our analysis describe rates measured in \( \mu \text{mol g}^{-1} \text{min}^{-1} \).

From these premises, state S = 1 (no function) can be readily identified to \( V_{cyc}^{t}(S) = 0 \). It follows that the total fraction of energy used for housekeeping metabolism (tot = n + a) is given by

\[ f_{hk,tot}(S) = \frac{V_{ATP}^{T}(S)}{V_{ATP}^{T}(S)} \]  

(4)

and as a consequence, the fraction of energy used for signaling is

\[ f_{tot}(S) = 1 - f_{hk,tot}(S). \]

To go one step further, we now make the hypothesis that the fraction of energy used for signaling is proportional to \( V_{cyc}^{t} \)

\[ f_{tot}(S) = k \frac{V_{cyc}^{t}(S)}{V_{ATP}^{T}(S)} \]  

(5)

with k a constant to be determined. Combining Eqs 3 and 5 leads to

\[ r_{o2}(S) = 1 - \frac{6}{CMR_{o2}(S)} \left( \frac{A_{f_{tot}(S)}V_{ATP}^{T}(S)}{k} + B \right) \]  

(6)

Choosing \( r_{o2} \) for a specific state S allows the determination of k. Inserting k back in Eq. 6 then determines \( r_{o2} \) for the other states. In other words, the hypothesis that the fraction of energy used for signaling is proportional to \( V_{cyc}^{t} \) (Eq. 5) makes the astrocytic contributions to oxygen utilization (\( r_{o2} \)) interdependent over different activation states. \( r_{o2}(S) \) can then be inserted into Eq. 3 to

**MATERIALS AND METHODS**

Our goal is to segment energy production in the cerebral gray matter in signaling (s) versus housekeeping or non-signaling components (hk) and further in astrocytic (a) versus neuronal (n) components. The detail of processes contained in these categories is discussed in the “Results” section. By construction, \( f_{hk,n} + f_{hk,a} + f_{a,n} + f_{a,a} = 1 \). To this end, we combine two preexisting datasets. The first dataset is the average tissue glucose (CMR_{glc}) and oxygen (CMR_{o2}) utilization computed for the different brain states as defined in Gjedde (2007). These four states S are: (1) no function, (2) low function/unconscious, (3) high function/awake and (4) maximum cortical activation. The second dataset is the measured empirical linear relation linking the total cycling of neurotransmitters \( V_{cyc}^{t} \) to the neuronal oxidative glucose utilization CMR_{glc}^{n} as recently reviewed in Hyder et al. (2006). The following analysis further assumes (i) that cells other than glia or neurons can be neglected (for simplicity,
compute $V_{\text{cyc}}(S)$. The fraction of energy expended by astrocytes for signaling is then given by

$$f_{\text{a}}(S) = \frac{N_f V_{\text{cyc}}(S)}{V_{\text{ATP}}(S)} \quad (7)$$

with $N_f$ the cost of recycling one neurotransmitter molecule and $f_{\text{a}}(S) = f_{\text{c}}(S) - f_{\text{b}}(S)$.

We then estimate the housekeeping metabolism of neurons from the value of $B$. Assuming that neurons are only fed on glucose in state 1, we write

$$f_{\text{b}}(S) = \frac{(6\lambda + 2) B}{V_{\text{ATP}}(S)} \quad (8)$$

and thus $f_{\text{b},a}(S) = f_{\text{b},a}(S) - f_{\text{b},a}(S)$. It is then possible to compute $r_{\text{gl}}$, the astrocytic contribution to glucose utilization by comparing $[f_{\text{a}}(S) + f_{\text{b},a}(S)]V_{\text{ATP}}(S)$ with the amount of energy produced by astrocytes computed from the fractions of glucose and oxygen that they use. Some algebra leads to

$$r_{\text{gl}}(S) = \frac{f_{\text{a}}(S) + f_{\text{b},a}(S)}{2CMR_{\text{gl}}(S)} \quad (9)$$

Finally, we compute the amount of non-oxidized carbons released or captured by neurons and astrocytes by comparing the utilization of glucose and oxygen in both cell types. To that end, we use Gjedde (2007)

$$J_{\text{carb,un}}(S) = 2[1 - r_{\text{gl}}(S)]CMR_{\text{gl}}(S) - \lambda[1 - r_{\text{gl}}(S)]CMR_{\text{O2}}(S) \quad (10)$$

for neurons and $J_{\text{carb,un}}(S) = 2r_{\text{gl}}(S)CMR_{\text{gl}}(S) - \lambda r_{\text{O2}}(S)CMR_{\text{O2}}(S)$ for astrocytes with $J_{\text{carb,un}}$ being positive if carbons are released or negative if carbons are required to satisfy for the observed oxygen utilization in that compartment.

Parameters in this study were chosen as follows. $A$ and $B$ were obtained by fitting Eq. 2 to the data in Figures 1A,B (least-square fit; data equally weighted). In the case of humans, because of the lack of data in the range $V_{\text{cyc}} = 0$, we imposed the value of $B$ to be three times smaller than the value of $B$ obtained for rodents to account for the fact that human glucose utilization in state 1 is one-third of the one in rodents (see Figures 1C,D) (Gjedde, 2007). This procedure yields $A = 0.86, B = 0.1$ (µmol g$^{-1}$ min$^{-1}$) for rodents and $A = 1.14, B = 0.03$ (µmol g$^{-1}$ min$^{-1}$) for humans. Finally, we used $N_f = 2.2$ assuming first that GABA constitutes 20% of the total neurotransmitter cycling (Patel et al., 2005) and second that recycling one glutamate molecule requires 2.33 ATPs (Attwell and Laughlin, 2001) while recycling one GABA molecule only uses 1.66 ATPs (Chatton et al., 2003).

RESULTS

We aim at computing an energy budget of cerebral gray matter for different states of activity (Gjedde, 2007). Rather than computing this budget in a bottom-up fashion by evaluating and then summing up the contribution of individual biophysical processes (Attwell and Laughlin, 2001; Nawroth et al., 2007), a useful approach, which however relies on a considerable number of estimates and approximations, we start from the total steady-state energy production and introduce a method to segment energy production of different compartments and categories of interest. More specifically, we aim at segmenting energy production in the gray matter in signaling versus non-signaling components (housekeeping) and further in glial versus neuronal components. To this end, we combine two preexisting datasets. The first dataset is the empirical linear relation linking the total cycling of neurotransmitters $V_{\text{cyc}}$ to the neuronal oxidative glucose utilization $CMR_{\text{glc}}$ (Hyder et al., 2006) (see Figures 1A,B). The second dataset is the average tissue glucose ($CMR_{\text{glc}}$) and oxygen ($CMR_{\text{O2}}$) utilization computed for the different brain states $S = 1,…,4$ (see Figures 1C,D). These four states $S$ are defined as: (1) no function, (2) low function/unconscious, (3) high function/awake and (4) maximum cortical activation (see Gjedde, 2007 for further details). It is shown in the “Materials and Methods” section that these two datasets are sufficient to compute a steady-state budget for gray matter under the hypothesis that the energy used for signaling is proportional to $V_{\text{cyc}}$, neglecting cells other than glia or neurons and assuming that energy measured as adenosine triphosphate production is used in the compartment where it is produced. For the sake of simplicity, all glial cells are pooled together and are hereafter indiscriminately referred to as astrocytes or glia. These two datasets are taken from the literature. All the other results presented in this section have been calculated on the basis of these data.

In order to segment energy production between the neuronal and the glial compartments, it is necessary to know their corresponding glucose and oxygen utilization. Direct and indirect measurements of the compartmentalization of glucose utilization are available for various conditions and systems but have yielded relatively different figures with estimates of the glial contribution to glucose utilization ($r_{\text{gl}}$) ranging from about 50% to 100% (Nehlig et al., 2004; Tsacopoulos et al., 1988; Végé et al., 2003; Zielke et al., 2007). So far, there is no direct measurement of the glial contribution to oxygen utilization ($r_{\text{O2}}$), however the relative activities of the TCA cycle could be used as a proxy (see Jolivet et al., 2009). It is however very difficult to relate these measurements to the abovementioned brain states $S = 1,…,4$. As demonstrated in the “Materials and Methods” section, the values of $r_{\text{O2}}$ become interdependent under the hypothesis that the fraction of energy used for signaling is proportional to $V_{\text{cyc}}$.

In other words, choosing the value of $r_{\text{O2}}$ for any given state determines the value of $r_{\text{O2}}$ for all the other states. In the following analysis, we consider two scenarios. In the first scenario, we assume that glial oxygen utilization is approximately constant across all states ($\Delta r_{\text{O2}} = 0\%$), consistent with the observation made in hippocampal slices by Kasischke et al. (2004) that NADH autofluorescence is constant in astrocytic mitochondria upon activation. In the second scenario, we assume that glial oxygen utilization increases by 15% from state 3 to state 4 ($\Delta r_{\text{O2}} = +15\%$) (Cruz et al., 2005).

Figure 2 shows $r_{\text{O2}}$ and $r_{\text{gl}}$ computed following the procedure detailed in the “Materials and Methods” section for rats (A) and humans (B) in the first (solid lines) and the second scenario (dashed lines). Our method yields results very similar for both species and consistent with experimental findings. We find that $r_{\text{gl}}$ is increasing with the level of activation and typically rests between 49% and 78% in humans and between 49% and 98% in rats in

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agreement with in vivo measurements (Nehlig et al., 2004) and other modeling efforts (Hyder et al., 2006). The increase of \( r_{\text{glc}} \) with the level of activation is less prominent in the second scenario than in the first scenario and this effect is more pronounced in humans with \( r_{\text{glc}} \) being almost constant in the latter case. In both species, we find the glial oxygen utilization fraction \( r_{\text{O2}} \) to be decreasing with the level of activation from about 49% in state 1 to approximately 10% in state 4. Again, these predictions are consistent with experimental findings showing that the relative contribution of glia to the total TCA cycle might be higher in anaesthetized animals (~41%) (Choi et al., 2002) than in awake animals (~21%) (Öz et al., 2004). Likewise, we find \( r_{\text{O2}} \) values in the range 6–17% in states 3 and 4 consistent with measurements in awake humans reporting ~8% (Shen et al., 1999), ~10% (Gruetter et al., 2001) and ~11% (Lebon et al., 2002) (calculated from the ratio of TCA cycles as summarized in Hyder et al., 2006). Fluctuations in the order of ±5% in the parameters convert into about ±5% fluctuations in the predicted value of \( r_{\text{O2}} \) and about ±19% fluctuations in the predicted value of \( r_{\text{glc}} \). However, the compartmentalization is not strongly affected with most of oxygen being used by neurons in both scenarios, most of glucose being used by astrocytes in the first scenario and a more balanced situation with about the same amount of glucose being used by both compartments in the second scenario. Finally, we observe that the values computed for \( r_{\text{O2}} \) in the activated state are remarkably close to the distribution of mitochondria between neurons and glia (Wong-Riley, 1989).

We formulated the two scenarios considered here as an increase of astrocytic oxygen utilization from state 3 to state 4 because it is numerically convenient. It is interesting to note however that in the second scenario, the astrocytic oxygen utilization \( (r_{\text{O2}} \cdot CMRO_2) \) continuously increases from state 1 to state 4 (Choi et al., 2002; Öz et al., 2004; Serres et al., 2008). We consider that GABA constitutes 20%
of the total neurotransmitter cycling (see Materials and Methods). If GABA is recycled by astrocytes, it seems surprising that \( r_{\text{go}} \) can be significantly lower than this value. However, each GABA recycled by astrocytes might lead to the production of NADH and FADH\(_2\) requiring a maximum of 1.5 O\(_2\) per GABA molecule. Hence, \( r_{\text{go}} \) can be smaller than 20% as long as \( 1.5 \times 20/100 \times V_{\text{cyc(tot)}} < r_{\text{go}} \times \text{CMR}_02 \) which is always the case in our results (data not shown). Similarly, it has been reported by several groups that the rate of glial anaplerotic glutamine synthesis might be as high as one-third of the rate of the glutamate-glutamine cycle (Lieth et al., 2001; Merle et al., 2002; Sibson et al., 2001; Xu et al., 2004). Glutamine generated in that way might lead to the production of NADH requiring a maximum of 1.5 O\(_2\) per glutamine molecule. Hence, \( r_{\text{go}} \) can be smaller than 33% as long as \( 1.5 \times 1/3 \times V_{\text{cyc(tot)}} < r_{\text{go}} \times \text{CMR}_02 \) which is always the case in our results (data not shown).

With \( r_{\text{go}}(S) \) and \( r_{\text{glc}}(S) \) available, it is possible to compute the energy budget for four categories of interest. They are the astrocytic housekeeping metabolism category, the neuronal housekeeping metabolism category, the astrocytic signaling category and the neuronal signaling category (see Materials and Methods). By construction, these categories cover the following biophysical processes: the housekeeping categories contain all energy demanding processes which are not related to signaling tasks, that is essentially maintaining the electrochemical gradient across the cell membrane by activating the Na\(^+\)/K\(^+\) pump as well as biochemical processes required to maintain structural integrity. The neuronal signaling category contains the energy expended after the activation of postsynaptic receptors and after the generation and propagation of action potentials, mostly due to the activation of the Na\(^+\)/K\(^+\) pump needed to restore the electrochemical gradients across the cell membrane. It also includes the energy required to package neurotransmitters into vesicles at presynaptic terminals. Finally, the astrocytic signaling category contains the energy used to capture and metabolically process neurotransmitters. **Figure 3** shows the energy budget computed (first scenario) for these four categories for rats (A) and humans (B) and details the exact percentages for state 4 (C–D). By construction, these four fractions sum up to 1 in all four states. As expected, the fraction of energy dedicated to housekeeping metabolism decreases with the level of activation while the fraction of energy dedicated to signaling increases with the level of activation to reach 84% and 86% in rodents and in humans respectively. Of these, the astrocytes only contribute for a small fraction of 7% in rodents and 5% in humans. The rest, about 80%, is used for neuronal signaling. Our approach yields figures for the energy budget remarkably similar to the results of Attwell and Laughlin (2001). Interestingly, the computed energy budget is not significantly dependent on the scenario considered, and therefore only the first scenario is shown.

Using the values of \( r_{\text{go}}(S) \) we have derived together with Eq. 3, it is now possible to compute \( V_{\text{cyc(tot)}}(S) \), i.e. the amplitude of the cycling of neurotransmitters corresponding to brain states \( S = 1, \ldots, 4 \). **Figure 4** shows \( V_{\text{cyc(tot)}}(S) \) calculated in this way for both humans and rodents in both scenarios. This procedure yields values in range of experimental measurements. We find values of \( V_{\text{cyc(tot)}} \) between 0 and 0.35 for humans and between 0 and 1.17 for rodents. Interestingly, these results closely match the ones reported by Gjedde, obtained with a different approach (Gjedde, 2007). He finds \( V_{\text{cyc(tot)}}(1) = 0, V_{\text{cyc(tot)}}(2) = 0.2 \) and \( V_{\text{cyc(tot)}}(3) = 0.5 \) for the rodent, whereas we can deduce \( V_{\text{cyc(tot)}}(1) = 0, V_{\text{cyc(tot)}}(2) = 0.2 \) and \( V_{\text{cyc(tot)}}(3) = 0.6 \). Note that the choice of a specific scenario does not affect the results significantly, nor do ±5% fluctuations in the free parameters (i.e. \( A, B \) and \( N_s \), the average cost of recycling one neurotransmitter molecule).

Finally, it is possible to evaluate the level of neuron-glia metabolic exchanges by comparing the balance of oxygen and glucose utilization in these two compartments. Indeed, complete oxidation of glucose requires six times more oxygen than glucose (Gjedde, 2007). Thus, combining the experimental values for \( \text{CMR}_02(S) \), \( \text{CMR}_{\text{glc}}(S) \) (**Figures 1C,D**) and the values we derived for \( r_{\text{go}}(S), r_{\text{glc}}(S) \) (**Figure 2**) into Eq. 10, it is possible to compute
$J_{\text{carbons}}(S)$. $J_{\text{carbons}}$ is positive if the compartment exports carbons (net transfer), that is, if the observed oxygen utilization in that compartment is smaller than six times the observed glucose utilization. On the contrary, $J_{\text{carbons}}$ is negative if the compartment imports carbons, that is, if the observed oxygen utilization in that compartment is larger than six times the observed glucose utilization. Doing so, we find that $J_{\text{carbons}}$ is systematically positive for astrocytes, systematically negative for neurons and very close to 0 in the case of state 1 for both the neuronal and the astrocytic compartments. This indicates that carbons are released from the glial compartment and are at least partially taken up by the neuronal compartment. In the case of state 1, glia and neurons are metabolically separated (no flux of carbons). The latter result is to be expected since the measured ratio $\text{CMR}_{\text{glu}}(1)/\text{CMR}_{\text{glu}}(1)$ is very close to 6 indicating a well balanced system. Moreover, we imposed that neurons use exclusively glucose in this state when computing $r_{\text{glu}}$, thus preventing glia-to-neuron carbon exchanges. Figure 5 shows the absolute value of $J_{\text{carbons}}$ versus the level of activation as embodied by the total cycling of neurotransmitters $V_{\text{cycling}}(S)$. $J_{\text{carbons}}$ is computed individually for neurons and glia as well as for both scenarios considered in this paper. In all cases, we observe a strong linear relationship ($R^2 \geq 0.99$) indicating that the amplitude of the glia-to-neuron carbon exchange is proportional to the activity in the tissue measured as the total cycling of neurotransmitters. The overall amplitude of this coupling significantly depends on the scenario considered. It is significantly stronger in the first scenario (constant astrocytic oxygen utilization) than in the second scenario (astrocytic oxygen utilization increases by 15% from state 3 to state 4). Finally, the fact that astrocytes always export more carbons than neurons capture suggests that a small fraction of carbons is simply released by the system to the bloodstream consistent with findings that lactate is being released by the brain in various states (Dalsgaard, 2006).

Our analysis relies on the choice of three parameters, $A$ and $B$ characterizing the linear relation between $\text{CMR}_{\text{glu}}^n$ and $V_{\text{cycling}}^n$, and $N$, representing the average cost of recycling one neurotransmitter molecule. In addition, we have arbitrarily selected two scenarios. In the first scenario, we assume constant astrocytic oxygen utilization ($\Delta \text{ox,a}^{34} = 0\%$), while in the second scenario we assume that the astrocytic oxygen utilization increases by 15% from state 3 to state 4.
state 4 ($\Delta_{\text{ox,a}}^{34} = +15\%$). None of the results presented in this paper was dramatically affected by ±5% fluctuations of the value of $A, B$ and/or $N_a$. On the contrary, the choice of a scenario for the relative increase of the astrocytic oxygen utilization has a significant impact on the results as already illustrated in Figures 2 and 5. In particular, $r_{\text{glc}}$ is significantly smaller with $\Delta_{\text{ox,a}}^{34} = +15\%$ than with $\Delta_{\text{ox,a}}^{34} = 0\%$ (Figure 2). Thus, increasing $\Delta_{\text{ox,a}}^{34}$ progressively shifts the balance of glucose utilization from being mostly glial to being mostly neuronal. As a consequence, the amplitude of the carbon shuttle from glia to neurons progressively decreases with increasing values of $\Delta_{\text{ox,a}}^{34}$ (Figure 5). Increasing $\Delta_{\text{ox,a}}^{34}$ beyond +15%, we find that the glia to neuron shuttle stops at $\Delta_{\text{ox,a}}^{34} = +22\%$ for humans and +23% for rodents yielding a situation where both cell types are metabolically independent and releasing excess carbons. Increasing $\Delta_{\text{ox,a}}^{34}$ further, we find that a shuttle from neurons to glia takes place for values $\Delta_{\text{ox,a}}^{34} = +27\%$ and higher.

**DISCUSSION**

The exact principles underlying the coupling of metabolic processes in register with neuronal activity and in particular the nature and role of neuron-glia metabolic interactions are still a subject of active debate. Here, we have introduced a novel computational approach to decipher neuron-glia compartmentalization in the context of brain energy metabolism. Based on measurements of tissue glucose and oxygen utilization in different brain states (Figure 1), our approach allows computing the fractions of oxygen and glucose taken up by glial cells ($r_{\text{O2}}$ and $r_{\text{glc}}$) and neurons ($1-r_{\text{O2}}$ and $1-r_{\text{glc}}$ respectively; Figure 2). Our approach suggests that a significant portion of glucose is taken up by astrocytes while oxygen is mostly used within the neuronal population and that this bias increases with the level of activation. The exact proportion of astrocytic glucose uptake is dependent on the scenario considered with most glucose being taken up by astrocytes in the first scenario and a more balanced situation in the second scenario. We then computed an activity-dependent energy budget for the brain (Figure 3) and assessed the balance of oxygen and glucose in both the neuronal and glial compartments (Figure 5). Finally, we found that there is a probable transfer of glucose-derived metabolites from astrocytes to neurons, the amplitude of which is proportional to the level of activity as measured by the cycling of neurotransmitters. Our approach is systematically applied to both human and rodent data yielding similar results for both species.
Our findings are in general agreement with results by Attwell and Laughlin (2001) on the cost of cortical computation. With our top-down approach it is not possible to distinguish between the cost of action potentials and the cost of postsynaptic potentials which are pooled together with other mechanisms in the neuronal signaling component \( f_{\text{act}} \). It is nevertheless possible to make a rough estimate of the cost of postsynaptic potentials using the same strategy as used in Eq. 7, that is allocating a fixed cost to postsynaptic potentials per recycled glutamate molecule while neglecting inhibition. After a quick calculation based on the study by Attwell and Laughlin using only the smallest reported value (that is 25.5 ATP/glutamate as a result of NMDA, non-NMDA and G-protein coupled receptors), we find that postsynaptic potentials would account for 47% of the total ATP utilization versus 35% for action potentials in humans and 58% versus 20% in rodents in the maximally activated state. These numbers suggest that postsynaptic potentials might contribute more to the energy budget than originally suggested by Attwell and Laughlin (2001). This is consistent with the finding that metabolic response is more correlated with local field potentials, reflecting principally post-synaptic potentials, that to spiking activity (Logothetis et al., 2001).

The findings presented here are also consistent with an activity-regulated transfer of glucose-derived metabolites from astrocytes to neurons as it was originally proposed by Pellerin and Magistretti (1994). Existence of this mechanism has been recently demonstrated in slices (Rouach et al., 2008). Our calculations show that such a shuttle from astrocytes to neurons is perfectly compatible with a relatively low cost associated to the recycling of neurotransmitters and with an upregulation of the astrocytic oxidative metabolism upon activation. Our approach does not allow us to determine the metabolic species involved in this shuttle but there is no doubt that lactate must be an important component of it, given the experimental evidence indicating lactate usage by neurons (Schurr and Payne, 2007). Of course the possibility that several metabolic species contribute to this shuttle cannot be excluded (see, e.g., Pfrieger, 2003). The potential roles of this shuttle beyond feeding neurons remains to be determined. Interestingly, lactate was recently demonstrated to play a direct role in the regulation of the local blood flow (Gordon et al., 2008; Yamanishi et al., 2006), of neuronal activity in the retrotrapezoid nucleus (Erlichman et al., 2008) as well as in the regulation of the local inhibitory activity in the subfornical organ (Shimizu et al., 2007).

Our approach relies on the division of brain activity in states as defined by Gjedde (2007). Average tissue glucose and oxygen utilization are provided for all these states (Figures 1C, D). While this division might be perceived as slightly arbitrary, it allows us to compute the astrocytic and neuronal fractions of glucose and oxygen utilization (Figure 2), the energy budget (Figure 3) and the net transport of glucose-derived metabolites (Figure 5) for a meaningful range of activation levels. These states should not be considered as fixed and well-defined categories but rather as individual samples along a continuum of brain activation states. For a better comparison with experimental results, it is then possible to convert these states into the corresponding amplitude of the cycling of neurotransmitters \( V_{\text{cyc}}(\text{neuro}) \) (Figure 4) as illustrated for the case of the net transport of glucose-derived metabolites (Figure 5). The same procedure can be applied to the glial and neuronal fractions of glucose and oxygen utilization and to the energy budget (not shown).

Our method is in part based on several assumptions that are rather common in the field. It is assumed that cells other than glia or neurons can be neglected since these two cell types constitute the vast majority of cells in the brain and occupy most of the cortical tissue volume (not counting extracellular space) (Verkhovsky and Butt, 2007). Even though ATP can be released as a neurotransmitter, notably by astrocytes, this release is unlikely to contribute significantly in comparison to the amount of ATP produced to fuel the system. Finally, like most modeling work and some experimental techniques in the field, our approach is restricted to the steady-state condition. It also relies on the hypothesis that the total energy used for signaling in the tissue is linearly proportional to the total cycling of neurotransmitters. While it is obvious that this hypothesis holds for the cost associated to neurotransmitters recycling, it is less clear that it holds for action potentials and postsynaptic potentials. However, the frequency versus current (\( f-I \)) curve of cortical neurons is close to linear in the presence of noise (Rauch et al., 2003). One can therefore assume that this hypothesis also holds within a physiologically relevant regime although it may not be valid in pathological cases or at either very low or very high levels of activation. Also note that our approach is not strictly restricted by this hypothesis and could easily be extended to non-linear relationships. The analysis would be essentially the same but would then require extra data in order to estimate the additional parameters entering into Eq. 5 and characterizing the relation between \( f_{\text{act}}(S) \) and \( V_{\text{cyc}}(\text{neuro})(S) \).

The results derived here rely on a steady-state or quasi steady-state condition. In the first few seconds that follow the increase or decrease in brain activation, the tissue undergoes fast metabolic transitions and the system is undoubtedly out of steady-state. However, upon activation, cerebral blood flow reaches a new stationary level after a few seconds only. Moreover, it was recently shown that sustained neuronal activation raises metabolism to a new steady-state level (Mangia et al., 2007). In particular, lactate quickly reaches a new steady-state and so does pyruvate because of the dynamic equilibrium between these metabolites. Similarly, recovery from activation is very slow, taking place over minutes to tens of minutes as illustrated by the slow decay of accumulated lactate after stimulation (Pritchard et al., 1991). Likewise, it seems reasonable to assume that different anesthetic levels correspond to different steady-states. Even though our formalism was developed under the assumption of steady-state conditions, we expect it to be valid except during the short transitions between different steady-states or quasi steady-states. Simulations of such non-steady-state transitions require alternative modeling methods.

As already discussed previously, our approach yields qualitatively robust results when we apply \( \pm 5\% \) fluctuations in the parameters. This is true except for \( \Delta_{\text{ox}} \) which characterizes how much the astrocytic oxygen utilization increases with activation (from states 3 to 4). In particular, increasing \( \Delta_{\text{ox}} \) progressively shifts the balance of glucose utilization from being mostly glial to being mostly neuronal. As a consequence, the amplitude of the carbon shuttle from glia to neurons progressively decreases with increasing values of \( \Delta_{\text{ox}} \) (Figure 5). While the value we chose for \( \Delta_{\text{ox}} \) is
in line with the study by Cruz et al. (2005) (\(\Delta_{\text{glc}}^{\text{ox,a}} = +16\%\)), some of us have reported larger values in the range of +90% in rodents and +60% in humans (Wyss et al., 2009). However, it should be noted that the increase in CMR_{glc} between rest and activation observed by Wyss and colleagues is about +130%, much larger than the values used in this study (+79% in humans and +75% in rats). When increasing CMR_{glc} so as to match such an increase, we could increase \(\Delta_{\text{glc}}^{\text{ox,a}}\) as high as \(\Delta_{\text{glc}}^{\text{ox,a}} = +60\%\) in rodents with no significant qualitative change in the results presented above (not shown). Nevertheless, the importance of \(\Delta_{\text{glc}}^{\text{ox,a}}\) as a control parameter in our approach suggests that the oxidative metabolism of astrocytes might play a very important role in the control of metabolic neuron-astrocyte interactions possibly inducing a change of regime from highly interdependent cell populations at the onset of activation when the astrocytic oxidative metabolism is not or only weakly upregulated (Cruz et al., 2005; Kasischke et al., 2004) to metabolically disconnected or weakly connected cell populations at later stages of stimulation (Wyss et al., 2009). The lactate produced in these late stages could diffuse away from the production site, possibly to other brain areas (Dienel and Cruz, 2004; Dienel and Hertz, 2001).

In our analysis, we have not considered glycogen, a potentially important energy substrate reservoir localized in astrocytes only. During strong activation, there is evidence that the stock of glycogen decreases (Swanson et al., 1992). Glycogen has been reported to be continuously degraded and resynthesized (Shulman et al., 2001; Walls et al., 2009). Glucose metabolism via this glycogen shunt only produces one ATP versus two generated via the glycolysis. Notwithstanding, our main results should not be significantly affected by this simplification even though it implies that our estimate of the astrocytic contribution to the total energy budget might be slightly overestimated. It is important however to keep in mind that most of the ATP is produced oxidatively. Moreover, the size of the glycogen reservoir is relatively small and thus may not contribute to an extent that would affect the calculated energy budget. Similarly, neglecting glycogen does not change our findings on the astrocyte-neuron transfer of glucose-derived metabolites. Glycogen utilization can be interpreted as an increase in the effective astrocytic fraction of the total glucose used to feed ATP synthesis. Furthermore, the end product of glycogenolysis in astrocytes has been reported to be lactate (Dringen et al., 1993). This process is consistent with the concept of the astrocyte-neuron lactate shuttle hypothesis rather than with the exchange of substrates in the opposite direction (Pellerin et al., 2007).

It should be further emphasized that our results do not depend on any assumption about the distribution of glucose and lactate transporters or on a specific scheme of stimulation during activation. Likewise, they do not depend on the exact route followed by metabolites, the extracellular space very likely being the place of exchanges between astrocytes and neurons. Furthermore, our results are independent from the exact pathways involved in glucose oxidation. More specifically, they hold equally well whether lactate or pyruvate is the end product of aerobic glycolysis in neurons (Schurr and Payne, 2007). Finally, our approach bypasses considerations on what drives the lactate shuttle, glutamate capture by astrocytes (Pellerin and Magistretti, 1994) or activation of astrocytic AMPA receptors (Caesar et al., 2008), while producing estimates of the cost of recycling glutamate consistent with other studies (Attwell and Laughlin, 2001). Future work is needed to experimentally test the predictions and conclusions drawn from the present modelling work.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any com- mercial or financial relationships that could be construed as a potential conflict of interest.