Research Report

Probing ischemic tissue fate with BOLD fMRI of brief oxygen challenge

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Abstract

It has been recently shown that at-risk tissue exhibits exaggerated T2⁎-weighted MRI signal increases during transient oxygen challenge (OC), suggesting that the tissue is still metabolically active. This study further characterized the effects of transient OC on T2⁎-weighted MRI in permanent focal stroke rats (N=8) using additional quantitative measures. The major findings were: i) the ischemic core cluster showed no significant response, whereas the mismatch cluster showed markedly higher percent changes relative to normal tissue in the acute phase. ii) Many of the mismatch pixels showed exaggerated OC responses which became hyperintense on T2-weighted MRI at 24 h. The area with exaggerated OC responses was larger than the mismatch, suggesting that some tissue with reduced diffusion were potentially at risk. iii) Basal T2⁎-weighted intensities on the perfusion–diffusion contourplot were high in normal tissue and low in the core, with a sharp transition in the mismatch. iv) OC-induced changes on the perfusion–diffusion contourplot dropped as perfusion and diffusion values fell below their respective viability thresholds. v) Basal T1 increased slightly in the ischemic core (P<0.05). OC decreased T1 in normal (P<0.05) but not in mismatch and core pixels. vi) OC decreased CBF in normal (P<0.05) but not in mismatch and core pixels. T2⁎-weighted MRI of OC has the potential to offer unique clinically relevant data.

Keywords: MRI, Perfusion, Diffusion, Oxygen inhalation, Penumbra

1. Introduction

The ability to distinguish reversible from irreversibly damaged ischemic brain tissue is of clinical importance in deciding treatment strategies for acute stroke patients. The gold standard to define ischemic penumbra (i.e., salvageable tissue) is by imaging oxygen extraction fraction using positron emission tomography which unfortunately is not widely available and involves ionized irradiation (Baron, 1999). A more commonly used imaging modality to define the ischemic penumbra is the perfusion–diffusion mismatch by MRI (Schlaug et al., 1999). However, some mismatch tissue is oligemic, salvageable and other is not salvageable, depending on the duration and nature of ischemic injury, and proximity of patent vessels, among
others. Thus, the perfusion–diffusion mismatch only approximates the ischemic penumbra. Despite its shortcomings (Kidwell et al., 2003), the perfusion–diffusion mismatch remains widely utilized to guide acute stroke treatment in the clinical setting. Numerous studies have attempted to probe the fate of the mismatch to improve prediction accuracy of the ischemic tissue fate.

A novel approach was recently introduced to further probe tissue at risk by using $T_2^*$-weighted MRI of transient oxygen challenge in ischemic stroke (Dani et al., 2010; Santosh et al., 2008). $T_2^*$-weighted signal intensity is sensitive to relative concentration of deoxyhemoglobin (Ogawa and Lee, 1990; Ogawa et al., 1993). The infarct core showed little or no change in $T_2^*$-weighted signal intensity during oxygen challenge (OC). The at-risk regions surrounding the infarct core showed an exaggerated increase in $T_2^*$-weighted signal intensity compared to the homologous region in the contralateral hemisphere. OC brings in oxygenated blood displacing the high deoxyhemoglobin concentration in the at-risk region where cerebral blood flow (CBF) is partially compromised but metabolic activity remains significant. It was thus hypothesized that tissue with exaggerated increase in $T_2^*$-weighted signal intensity during OC is potentially salvageable (Dani et al., 2010; Santosh et al., 2008).

The goal of this study was to further characterize the OC $T_2^*$-weighted signal responses in the permanent focal stroke rats by using additional and quantitative measures. The ischemic core, normal and mismatch were objectively classified pixel-by-pixel using automated cluster analysis based on quantitative apparent diffusion coefficient (ADC) and CBF (Shen et al., 2004b). Basal $T_2^*$-weighted signal intensities, magnitude and percent signal changes due to oxygen challenges, and their times to peaks were analyzed for the three tissue types. These data were also evaluated as a function of binned ADC and/or CBF on a pixel-by-pixel basis. Moreover, ischemia-induced and OC-induced $T_2^*$ changes were analyzed to evaluate their potential effects on CBF and $T_2^*$-weighted signals in stroke animals.

2. Results

Fig. 1 shows the ADC, CBF, OC $T_2^*$-weighted percent-change maps at 30 min, and $T_2^*$-weighted MRI at 24 h after permanent MCAO from one animal and all animals. ADC and CBF were reduced in the stroke hemisphere as expected. ISODATA – iterative self-organizing data analysis – used clustering analysis and classified pixels into normal, mismatch and core clusters. The ischemic core pixels showed a mixture of low positive and negative percent changes. Mismatch cluster showed regions of markedly higher percent changes compared to the homologous region in the contralateral hemisphere. The exaggerated OC responses are most consistent and apparent in the primary somatosensory cortex (solid long arrows) and less apparent in brain structures along the midline (dash short arrows). The area with exaggerated OC response was larger than the ISODATA mismatch area which was hyperintense on $T_2^*$-weighted MRI at 24 h. In the normal hemisphere, there were some exaggerated OC responses which were likely from large vessels as indicated by their shapes. These vessels could be readily identified.

The OC response time courses from all animals showed that the mismatch OC response was about twice that of the normal OC response, and the core cluster showed negligible OC response (Fig. 2A). Time to peak of the mismatch was significantly delayed compared to normal tissue. Group-averaged OC percent changes, TTP, ADC and CBF values for the three types of tissues were quantified (Fig. 2B). Group-averaged OC percent changes were significantly different among all three tissue types ($P<0.05$). Mismatch OC% change was 1.5 times higher than normal OC% changes and the core OC% change was 3.5 times smaller than normal. Mismatch TTP was about 2.5 times longer than normal TTP.

In addition, basal $T_2^*$-weighted signal, OC-induced magnitude changes and OC-induced percent changes were analyzed as a function of ADC and CBF at 30 min after stroke (Fig. 3). The vertical solid lines and shaded bars indicate, respectively, normal values of ADC and CBF and their standard deviations obtained from this study. Two vertical dot lines and the shaded bars indicate, respectively, the viability thresholds and their standard deviations as reported in (Shen et al., 2003). $T_2^*$-weighted signal intensities of normal ADC and CBF pixels were generally high (Fig. 3A). There were sharp transitions in basal $T_2^*$-weighted signal intensities between the two vertical lines. For ADC and CBF below the viability thresholds, $T_2^*$-weighted signal intensities was low and relatively independent of ADC and CBF. Fig. 3B and C show that the OC-induced changes in pixels with ADC and CBF below their viability thresholds were markedly reduced. By contrast, pixels with ADC and CBF between normal and viability thresholds showed markedly higher changes than those of normal tissue.

The OC responses were also analyzed on the combined ADC-CBF contour plot with color-code scale indicating baseline $T_2^*$-weighted signal intensities, OC magnitude changes, and OC percent changes (Fig. 4, 30 min after MCAO). Baseline $T_2^*$-weighted signal intensities of the mismatch were lower than normal, and those of the ischemic core were lower than mismatch. The OC magnitude and percent changes of the mismatch were markedly higher than normal, whereas those of the core were negligible. There were some distinct differences between OC magnitude- and percent-change contour plots in that the peak of the percent-change responses were localized to the mismatch zone on the contourplot whereas the peak of the magnitude-change responses were also located at relatively higher CBF values. In the OC magnitude change plot, the largest changes centered about 0 ml/g/min and 0.55 $\times 10^{-3}$ mm$^3$/s as well as 0.3 ml/g/min and 0.65 $\times 10^{-3}$ mm$^3$/s. By contrast, in the OC% change plot, the largest changes centered about 0 ml/g/min and 0.55–0.65 $\times 10^{-3}$ mm$^3$/s. Note that a few negative CBF pixels in Fig. 4 were due to subtraction errors in the continuous ASL technique for CBF close to zero (Shen et al., 2003).

The effects by ischemia and OC on $T_1$ and CBF changes were analyzed for the 30-min time point after stroke (Fig. 5). Basal $T_1$ increased slightly in the ischemic core ($8.2\times 2\%, P<0.05$). OC decreased $T_1$ significantly in the normal hemisphere (1.8$\pm$ 0.8%, $P<0.05$), but not in ischemic core ($-0.2\pm 0.2\%, P>0.05$) and mismatch ($-0.6\pm 1.0\%, P>0.05$). OC decreased CBF significantly in the normal hemisphere (0.15$\pm$ 0.07 ml/gram/min, 16$\pm$8%, $P<0.05$), but had no significant effect on the core and mismatch tissue. Although OC responses on sham rats were not measured,
our previous data on normal animals subjected to OC challenges were not statistically different from the normal hemisphere of stroke rats herein (data not shown).

3. Discussion

This study further characterized the effects of transient oxygen challenge on $T_2^*$-weighted signal intensities in permanent stroke rats. The major findings were:

i) The ischemic core cluster, derived automatically from perfusion and diffusion data, showed no significant response, whereas the mismatch cluster showed markedly higher percent changes relative to normal tissue in the acute phase. The exaggerated OC responses are more apparent in the primary somatosensory cortex than other brain structures at risk in this stroke model. ii) Many of the mismatch pixels showed some exaggerated OC responses which became hyperintense on $T_2^*$-weighted MRI at 24 h. iii) Basal $T_2^*$-weighted signal intensities on the perfusion–diffusion contourplot were high in the normal cluster and low in the core cluster, with a sharp transition in the mismatch cluster. iv) OC-induced changes on the perfusion–diffusion contourplot dropped as perfusion and diffusion values fell below their respective viability thresholds. v) Basal $T_1$ increased slightly in the ischemic core. OC decreased $T_1$ significantly in the normal hemisphere, indicative of hyperoxia-induced vasoconstriction or increased dissolved oxygen in the plasma, but OC had no significant $T_1$ effect in the ischemic core and mismatch pixels. vi) OC decreased CBF.

Fig. 1 – (A) ADC, CBF, ISODATA and oxygen-challenge percent change maps at 30 min after MCAO and $T_2^*$-weighted MRI at 24 h after MCAO from (A) one animal and (B) all 8 animals. Scale bar unit for ADC is $10^{-3}$ mm$^2$/s and for CBF is ml/gram/min. In the ISODATA maps, normal is shown as orange, mismatch as green, and core as cyan. Excluded CSF and corpus callosum are shown as black. Arrows indicate regions of exaggerated OC responses.
significantly in the normal hemisphere, but had no significant CBF effect on the mismatch and the ischemic core pixels.

The present study corroborated Santosh’s findings (Santosh et al., 2008) in the following ways. In normal tissue, the increased oxygen delivery during OC resulted in increased $T_2^*$-weighted signal intensities because capillary and venous oxygenation still have some room for increased oxygen saturation. In the ischemic core, there was no blood-flow and no oxygen delivery and thus OC $T_2^*$-weighted change was negligible. In the mismatch, some tissue was still metabolically active but with restricted blood flow, and thus oxygen extraction fraction and deoxyhemoglobin concentration in the capillaries and veins were higher than normal. Upon OC, $T_2^*$-weighted signal magnitude increased in the mismatch was higher than in normal tissue. The exaggerated OC% changes were likely because of the smaller denominator. Blood-gas $O_2$ saturation was ~90% under air and 100% under oxygen inhalation. Thus, the $T_2^*$-weighted signal changes were unlikely to come from the arterial side; rather they came predominantly from the venous side.

The present study also extended Santosh’s study in several ways. First, it probed the OC response in the normal, mismatch and ischemic core clusters that were objectively defined. Second,
basal $T_2$-weighted signal intensities and magnitude change of OC and the response time to peak were analyzed as a function of ADC or CBF on a pixel-by-pixel basis as well as the ADC-CBF contour plots. Results were interpreted with respect to the ADC and CBF viability thresholds. Third, the potential effects on $T_2^*$-weighted signal responses by ischemia and OC were analyzed. Our data had higher spatial resolution which allowed improved separation of different tissue types. These results are discussed below.

3.1. OC responses in ischemic core, normal and mismatch

The OC% map showed that many of the ISODATA-defined mismatch pixels had exaggerated OC responses compared to normal pixels (Fig. 1), whereas the ischemic core had no significant OC responses. We avoided ROI analysis because it could unintentionally bias as there were considerable heterogeneous $T_2$-weighted signal changes across the brain. A case in point is that even within the normal hemisphere, there were small areas that had high or low OC responses (Fig. 1) due to large vessels or white matter, respectively. The advantage of the cluster analysis is that these clusters include all pixels within the three defined classes, which improved certainty and demonstrated that most mismatch pixels exhibited exaggerated OC responses. In some animals, the area with exaggerated OC response was larger than the mismatch area, suggesting that some tissue with reduced ADC were at risk and thus potentially salvageable. ISODATA analysis showed that pixels with exaggerated OC responses became hyperintense on $T_2$-weighted MRI in permanent stroke.

The exaggerated OC response was often found in the primary somatosensory cortex. The anterior communicating artery offers collateral flow and thus the primary somatosensory cortex is among the few structures that have a better chance of being salvaged by reperfusion. This is consistent with the variable and higher than normal CBF and ADC in this region compared

Fig. 3 – (A) Group-averaged profiles of baseline $T_2$-weighted signal intensity, (B) magnitude changes in $T_2$-weighted intensity by oxygen challenge, and (C) percent changes by oxygen challenge as function of ADC and CBF at 30 min after stroke. The vertical solid lines indicate normal ADC and CBF with shaded bars indicating their standard deviations from this study. Two vertical dot lines indicate the viability thresholds with shaded bars indicating their standard deviations as reported in [Shen et al., 2003].
to many subcortical regions. Moreover, unlike the hypoperfused regions in the subcortical regions which have embedded large vessels, the OC response in primary somatosensory was not contaminated by large vessels, which made the high OC response in primary somatosensory region readily visible. It is interesting to note that the TTP of OC response was longer in mismatch tissue. The longer TTP of OC response in the mismatch tissue was due to delayed delivery of oxygen in a manner similar to TTP in dynamic contrast susceptibility CBF measurement. As such the TTP of OC response could potentially be used to identify at risk tissue.

Some of the scattered core pixels had moderately high OC response (Fig. 2B), likely due to some residual oxygen delivery and/or minor contamination by non-core pixels. There were no significant differences in baseline CBF or ADC between pixels with and without OC response in the core.

There are some confounds in this study. Although ISODATA can include analysis of all pixels within the three defined classes,
its disadvantage is that there could be scattered misclassified pixels. Another confound is that some pixels in the normal hemisphere showed exaggerated OC responses, likely from large-vessel contamination. Similarly high OC response patterns were also seen in normal rats. By doing pixel-by-pixel normalization using the contralateral hemisphere response as control, exaggerated OC responses could also be detected. Not all mismatch pixels showed exaggerated OC responses. This could be because the mismatch and exaggerated OC responses depicted different physiological status. Further studies are needed to compare the tissue fate of the mismatch and exaggerated OC responses (i.e., to evaluate the tissue fate with reperfusion).

3.2. OC responses as a function of ADC and CBF

There was also considerable heterogeneity of ADC and CBF within each tissue cluster. We thus analyzed the basal $T_2^*$-weighted signal intensities, magnitude and percent change by OC as a function of ADC alone and CBF alone, and with respect to viability thresholds (Shen et al., 2003). For pixels below ADC and CBF viability threshold (Fig. 3), basal $T_2^*$-weighted signal intensities were low and constant, whereas OC% changes drop from high to low rapidly with decreasing ADC or CBF. The likely explanation is that, at sufficiently low ADC and CBF, the occlusion essentially occluded the vessel and tissue oxygenation is anoxic without oxygen gradient under basal (air) conditions. By contrast, an oxygen gradient was likely present in pixels with mildly reduced ADC which were likely close to patent vessels. Reduced $T_2^*$-weighted signals have been observed to vary among infarct core, penumbra and normal tissue (Kavec et al., 2001; Kavec et al., 2004). However, sensitivity is low precluding practical application of basal $T_2^*$-weighted MRI to identify salvageable and non-salvageable tissue.

For pixels with ADC and CBF above the viability thresholds but below normal values, basal $T_2^*$-weighted signal intensities decreased progressively with decreasing ADC or CBF, whereas the OC changes were higher than normal pixels. The ADC and CBF values of tissue with enhanced OC responses were above the ADC and CBF viability thresholds (Meng et al., 2004; Shen et al., 2003), supporting the notion that pixels with exaggerated OC responses were at risk tissue and thus potentially salvageable. These observations further corroborated the significances of viability thresholds established previously under similar experimental conditions (Meng et al., 2004; Shen et al., 2003).
Note that baseline $T_2^*$-weighted intensities at the high ends of the ADC and CBF values continued to rise slightly instead of reaching a plateau. The high ADC is likely due to contamination by cerebrospinal fluid. The high CBF is likely due to contamination by large vessels and high vascular density. OC-induced responses however showed a plateau (Fig. 3B and C). These high ADC and CBF pixels could be readily distinguished on maps.

Although the information from OC responses as a function of ADC or CBF alone were informative, it was incomplete and was even misleading because these plots did not take into account the CBF status for each ADC bin and vice versa. A case in point is that the peak of the contour plots (Fig. 4B) is localized at 0 ml/g/min and 0.55×10⁻³ mm²/s whereas peaks appeared at 0.6 ml/g/min and 0.6×10⁻³ mm²/s (Fig. 3B). This discrepancy was the result of mixing normal and low CBF pixels in Fig. 3B, shifting the OC response peaks. Thus, the OC responses on the CBF–ADC contour plots should be more accurate because they accounted for both tissue ADC and CBF properties.

Careful inspection of the ADC-CBF contour plots showed there were some distinct differences in OC magnitude and percent changes. The peaks of the exaggerated OC changes tended to be at lower CBF in the % contour plot, likely because of the division by smaller basal $T_2^*$-weighted signal intensities. This raises the question whether magnitude changes should be used instead. While the OC percent changes may be biased by the low basal $T_2^*$-weighted signal, it may be more sensitive. Additional studies are needed to determine which analysis method will provide more accurate prediction of final infarct volume (Huang et al., 2011). With improved MRI sensitivity, the ability of basal $T_1^*$ or $T_2^*$-weighted MRI to identify salvageable and non-salvageable tissue may be warrant.

3.3. $T_1$ effects on OC responses

In this study, the TR (1 s with 60° flip angle) was not infinitely long and thus, there could be $T_1$-weighting in the $T_2^*$-weighted signals. There are two potential $T_1$ effects on OC-induced $T_2^*$-weighted signal changes: i) $T_1$ induced by ischemia, and ii) $T_1$ induced by dissolved molecular oxygen during OC. In the acute phase (up to 3 h) during which vasogenic edema in the permanent MCAO is likely minimal, the ischemia-induced $T_1$ change is likely small, as indeed the case herein. In the chronic stroke phase, during which edema could be significant (Tanaka et al., 2011), longer $T_1$ due to edema effect in cerebral ischemia would result in a reduced OC response.

Oxygen inhalation increased dissolved paramagnetic molecular oxygen decreases $T_1$ in normal tissue. Oxygen delivery and consumption might differ among the three different tissue types, the $T_1$ effect by dissolved oxygen could vary among the tissue types. In the core, this was not an issue because there was no oxygen delivery and OC-induced % changes were low. In the normal tissue, dissolved oxygen due to OC decreased $T_1$, which exaggerated $T_2^*$-weighted OC response for short TR. In the mismatch tissue, OC increases dissolved oxygen somewhat (but less than normal) and, thus, augmented $T_2^*$-weighted signal for short TR but to a less extent than in normal tissue. That said, with a 60° flip angle and 1 s TR, these effects were likely small for all tissue types in the permanent MCAO model in the acute phase.

In normal tissue, OC decreased CBF by 16±8% due to hyperoxia-induced vasoconstriction, consistent with the ~10% hyperoxia-induced CBF reduction reported previously (Kety and Schmidt, 1948; Sicard and Duong, 2005). In the mismatch, OC increased instead of decreased CBF likely because the vessels were either damaged and did not experience significant hyperoxia. The improved oxygenation by OC could also improve perfusion (Henninger et al., 2007; Shin et al., 2007). Dissolved O2 may change blood $T_1$ and the brain $T_1^*$, which could affect CBF calculation. This effect on CBF calculation, although unknown, is likely small. Some studies have shown prolonged oxygen inhalation in cerebral ischemia to be beneficial (Henninger et al., 2007; Shin et al., 2007) while other studies have shown it to be harmful (Michalski et al., 2010; Padma et al., 2011). The beneficial effects include reversal of ADC lesion, prolongation of the mismatch status, extending the therapeutic window. The harmful effects include increased generation of reactive oxygen species. The brief (2 min) OC in our study is unlikely to significantly alter ischemic tissue fate. A short OC duration can be used in future studies to avoid potential unwanted effects on ischemic tissue fate. It is also worth mentioning that it would be of interest to compare OC findings with spontaneous breathing as it is clinically more relevant. Unlike ventilated condition, respiration rate, blood pressure and heart rate could change (Duong et al., 2001) which could affect the OC responses.

In summary, this study presents further characterization of the effects of transient oxygen challenge on $T_2^*$-weighted signal intensities in stroke rats as function of ADC and CBF on a pixel-by-pixel basis. The key finding is that many of the perfusion–diffusion mismatch shows exaggerated oxygen-challenge responses. Future studies will evaluate the effects of treatment (such as reperfusion) on the exaggerated OC responses, quantitatively define the OC-response viability threshold, and employ predictive model to determine if exaggerated OC responses improves prediction accuracy.

4. Experimental procedure

4.1. Animal preparation

Animal experiments were performed with approval of the Institutional Animal Care and Use Committee, UT Health Science Center San Antonio. Eight male Sprague Dawley rats (250–300 g) were anesthetized with 2% isoflurane in air during surgery. Permanent focal brain ischemia of the right hemisphere was induced using the intraluminal middle cerebral artery occlusion (MCAO) method (Shen et al., 2003). Animals were mechanically ventilated and maintained at ~1.2% isoflurane in air (unless during OC) during MRI. End-tidal CO₂ was monitored via a Surgivet capnometer (Smith Medical, Waukesha, WI, USA). Noninvasive end-tidal CO₂ values have been calibrated previously against invasive blood-gas samplings under identical settings (Sicard et al., 2003). The rectal temperature was maintained at 37.0±0.5 °C. The heart rate and blood oxygen saturation level were monitored using a MouseOx system (STARR Life Science Corp., Oakmont, PA, USA). All recorded physiological parameters were maintained within normal physiological ranges unless otherwise perturbed by OC.
Quantitative CBF and ADC were acquired every 30 min post-occlusion up to 180 min. Transient oxygen challenge (ventilated with oxygen) was performed at about 45 min post-occlusion. T2*-weighted MRI was performed during which the animals breathed 1 min air, 2 min O2, 5 min air, and 2 min O2 and 2 min air. To evaluate the possible effects of ischemia and O2 on T1 and CBF which could indirectly affect T2*-weighted signal changes during OC, quantitative T1 and CBF were measured during air and during oxygen inhalation (6 min). These additional T1 and CBF measurements were performed on some animals (N=5). All rats were imaged again at 24-hr post-occlusion and T2 maps were acquired to confirm infarct volume.

4.2. MRI experiments

MRI experiments were performed on a 7-T/40-cm magnet, a Biospec Bruker console (Billerica, MA), and a 40-G/cm gradient insert (ID=12 cm, 120-μs rise time). A surface coil (2.3-cm ID) was used for brain imaging and a neck coil (Duong et al., 2000) for perfusion labeling. Coil-to-coil electromagnetic interaction was actively decoupled.

ADC: Averaged ADC was obtained by averaging three ADC maps with diffusion-sensitive gradients separately applied along the x, y or z direction. Single-shot, spin-echo echo-planar images (EPI) were acquired (Shen et al., 2005) with matrix=96×96 with partial Fluorier (5/8) acquisition and reconstructed to 128×128, FOV=2.56×2.56 cm, seven 1.5-mm slices, TR=3 s (90° flip angle), TE=37 ms, b=4 and 1200 s/mm^2, λ=17.53 ms, δ=5.6 ms, and 16 averages.

CBF: CBF was measured using the continuous arterial spin-labeling (cASL) technique with single-shot, gradient-echo, echo-planar-imaging (EPI) acquisition as described previously (Shen et al., 2005). Continuous arterial spin labeling employed a 2.7-s square radiofrequency pulse to the labeling coil. Acquisition parameters were matrix=96×96 with partial Fluorier (5/8) acquisition and reconstructed to 128×128, FOV=2.56×2.56 cm, seven 1.5-mm slices, TR=3 s (90° flip angle), and TE=10.2 ms.

T2*-weighted MRI of oxygen challenge: OC fMRI was acquired using T2*-weighted gradient-echo EPI, matrix=96×96 (reconstructed to 128×128), FOV=2.56×2.56 cm, seven 1.5-mm slices, TR=1 s, TE=26 ms, 60° flip angle. OC experiment paradigm was: 1 min OFF, 2 min ON, 5 min OFF, 2 min ON and 1 min OFF, and 720 repetitions in total.

T1: T1-weighted images were acquired using single-shot inversion-recovery gradient-echo EPI sequence with 6 different inversion delay times (0.025, 0.5, 1, 2, 4 and 8 s), matrix=96×96 (reconstructed to 128×128), FOV=2.56×2.56 cm, TR=12 s (90° flip angle), and 4 signal averages.

T1: T1-weighted images were acquired using fast spin-echo pulse sequence with two effective echo times (50 and 80 ms), TR=2 s (90° flip angle), matrix=128×128, FOV=2.56×2.56 cm, echo train length 8, and 8 averages.

4.3. Data analysis

Data analysis used codes written in Matlab (MathWorks Inc., Natick, MA, USA) and the STIMULATE (University of Minnesota) software. Data were reported as mean±SD, with P<0.05 (t-test) taken to be statistically significant. ADC, CBF and T2 maps were calculated as described previously (Shen et al., 2003; Shen et al., 2004a). OC response percent change maps with respect to basal (air) conditions were calculated. T1 maps with intensity in unit of seconds were calculated pixel-by-pixel by fitting the model of $S_i = S_0 - 2AE^{-t/T_{1i}}$ where $Si$ is the signal intensity obtained with inversion delay time $Ti$.

Cerebrospinal fluid and the corpus callosum were excluded from analysis via ISODATA method (Shen et al., 2004b). Three tissue types (normal, perfusion-diffusion mismatch and ischemic core) were characterized by using auto-clustering ISODATA method (Shen et al., 2004b) based on ADC and CBF data. Images among different animals were co-registered using custom-designed semi-automatic co-registration software (Liu et al., 2004; Schmidt et al., 2006; Shen and Duong, 2008; Shen et al., 2005) to generate group-averaged maps. Percent change, time to peak (TTP), ADC and CBF values were analyzed for different tissue types for individual animals, followed by group averaging. Time to peak was defined as the time from one standard deviation above the mean of baseline to 90% of the mean peak value.

Profiles of baseline T2*-weighted signal intensity, OC signal magnitude change, and OC% change were plotted as function of ADC or CBF. Baseline T2*-weighted signal intensity, signal intensity change by OC and OC% changes were also plotted on ADC–CBF contour plots. The grid size was 0.02×10^-3 mm^2/sec for ADC or 0.12 mL/gram/min for CBF, respectively. Data were interpreted with respect to viability thresholds (Shen et al., 2003).

T1 and CBF changes induced by oxygen were analyzed for the ISODATA determined ischemic core, mismatch and compared with the homologous regions in the contralateral hemisphere.

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References


