Brain High-Energy Phosphates and Creatine Kinase Synthesis Rate under Graded Isoflurane Anesthesia: An In Vivo $^{31}$P Magnetization Transfer Study at 11.7 Tesla

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**Purpose:** The creatine kinase rate of metabolic adenosine triphosphate (ATP) synthesis is an important metabolic parameter but is challenging to measure in vivo due to limited signal-to-noise ratio and long measurement time.

**Theory and Methods:** This study reports the implementation of an accelerated $^{31}$P Four Angle Saturation Transfer (FAST) method to measure the forward creatine kinase (CK) rate of ATP synthesis. Along with a high-field scanner (11.7 Tesla) and a small sensitive surface coil, the forward CK rate in the rat brain was measured in ~5 min.

**Results:** Under 1.2% isoflurane, the forward CK rate constant and metabolic flux were, respectively, $k_{\text{CK}} = 0.26 \pm 0.02 \text{ s}^{-1}$ and $F_{\text{CK}} = 70.8 \pm 4.6 \mu\text{mol/g/min}$. As a demonstration of utility and sensitivity, measurements were made under graded isoflurane. Under 2.0% isoflurane, $k_{\text{CK}} = 0.16 \pm 0.02 \text{ s}^{-1}$ and $F_{\text{CK}} = 41.0 \pm 4.2 \mu\text{mol/g/min}$, corresponding to a 38% and 42% reduction, respectively, relative to 1.2% isoflurane. By contrast, the ATP and phosphocreatine concentrations were unaltered.

**Conclusion:** This study demonstrated the $^{31}$P FAST measurement of creatine kinase rate of ATP synthesis in rat brain with reasonable temporal resolution. Different isoflurane levels commonly used in animal models significantly alter the CK reaction rate but not ATP and phosphocreatine concentrations.

Key words: rats; high fields; metabolic flux; MRS

**INTRODUCTION**

The majority of the metabolic adenosine triphosphate (ATP) yield in the brain is produced in the mitochondria by oxidative phosphorylation by means of the ATPase pathway (Pi + ADP $\leftrightarrow$ ATP). The adult human brain produces and consumes 5 times its weight (~5.7 kg) in ATP daily, whereas at a given moment the brain contains only approximately 2 g of ATP (1–3). Thus, ATP is rapidly cycled to meet the energetic demands.

Creatine kinase (CK) catalyses the conversion of creatine and consumes ATP to create phosphocreatine (PCr) and adenosine diphosphate (ADP): (ATP + Cr $\leftrightarrow$ PCr + ADP). This CK reaction is reversible and thus ATP can also be generated from PCr and ADP. As such, PCr serves as energy storage for the rapid buffering and regeneration of ATP. Moreover, CK enzymes also facilitate the transfer of the high-energy bond out of the mitochondria to the cytosol in the form of PCr. The high-energy phosphate bond in the cytosol is cycled between the high-energy phosphates (PCr and ATP). The rate of ATP synthesis by means of the CK pathway is 5–6 times higher than the metabolic rate of ATP synthesis by means of the ATPase pathway (4,5). Such high rate of chemical exchange ensures diffusion of the high-energy phosphate bond, so that energy produced at the mitochondria is available at the cell membrane to maintain ionic gradients. Disruption of ATP energy pathways could affect normal cellular function and has been associated with several metabolic diseases (6–8).

$^{31}$P magnetization transfer (MT) by MR offers a unique, noninvasive tool for directly measuring the CK rate of ATP synthesis in vivo. $^{31}$P MT measures the forward creatine kinase rate ($k_{f,\text{CK}}$) by using frequency-selective RF energy to saturate $\gamma$-ATP. Saturating the $\gamma$-ATP resonance results in the attenuation of the PCr amplitude due to chemical exchanges. By measuring the change in signal amplitude of the PCr resonance, the forward creatine kinase rate can be calculated.

$^{31}$P MT experiments to measure CK rate are inherently challenging. $^{31}$P has approximately 1/1000th of the signal-to-noise ratio of the $^1$H$_2$O signal in vivo. The $^{31}$P high-energy phosphates are characterized by long longitudinal magnetization recovery ($T_1$, on the order of seconds) and short transverse magnetization relaxation ($T_2$, on the order of tens of milliseconds). These constraints have prevented widespread use of $^{31}$P MT techniques. Nonetheless, $^{31}$P MT has been used to measure CK rates under different anesthetics, pharmacologic and functional stimulations (4,5,9–11), and in association with stroke (8). The protocols for these applications of $^{31}$P MT ranged from half an hour to ten hours.

This study implemented the accelerated $^{31}$P Four Angle Saturation Transfer (FAST) (12) technique to evaluate the brain high-energy phosphates and the forward creatine kinase synthesis rate under graded isoflurane anesthesia. High field (11.7 Tesla [T]) and a small sensitive surface coil were used to improve $^{31}$P signal sensitivity. BIRP radiofrequency excitation was used to overcome radiofrequency B1 field inhomogeneity.
associated with the use of surface coil. The temporal resolution of the 31P FAST approach was 5 min.

THEORY

31P MT measurements can be used to measure the forward CK rate constant ($k_{f,CK}$ in units s$^{-1}$) of ATP synthesis (PCr $\rightarrow$ ATP). The modified Bloch equation for the MT experiment can be written as (12):

$$M'_{0}/M_{0} = 1 + k_{f} \cdot T_{1}^{\text{int}}$$

where $M'_{0}$ is the magnetization of PCr in the presence of RF saturation, $M_{0}$ is the magnetization of PCr in the absence of RF saturation, and $T_{1}^{\text{int}}$ is the longitudinal relaxation constant for PCr in the presence of saturating RF irradiation of $\gamma$-ATP. In addition, the forward metabolic flux $F_{f,CK}$ (mol/g/min) is related to $k_{f,CK}$ as $60 \times k_{f,CK} \times [\text{PCr}]/1.1$, where [PCr] is the PCr concentration which is usually taken as 5 mM in normal rats (4,13) and the brain tissue density is assumed to be 1.1 g/mL. Absolute measurements of [PCr] using NMR are challenging, requiring a carefully calibrated scheme using an external reference, and often are not necessary. It is valuable to report the metabolic flux, compared with the rate constant only, when altered physiology can be measured in the same animal, thereby observing reliable changes in metabolic concentrations. One caveat, is that the magnetization transfer experiment must be fast relative to the changes of metabolite concentrations, so that the physiological concentrations can be assumed stable over the duration of the measurement.

In vivo 31P MT experiments are generally a variation of the saturation transfer (ST) experiment. However, conventional saturation transfer experiments require acquisition of many data points at long repetition time (TR, requiring lengthy scan times. In this study, we implemented an accelerated magnetization transfer experiment, 31P four angle saturation transfer (FAST) (12), to measure $k_{f,CK}$ in approximately 5 min, instead of hours. The FAST approach uses acquisitions with low flip angles allowing shorter TR. Measuring the parameters $M'_{0}$, $M_{0}$, and $T_{1}^{\text{int}}$ requires only four spectra, acquired with and without saturation at two flip angles: $\alpha$ and $\beta$. The PCr signal from each spectra (M($\alpha$), M($\beta$), M'($\alpha$), and M'($\beta$), where the prime denotes acquisitions with $\gamma$-ATP saturated) are used to calculate $M_{0}$, $M'_{0}$, and $T_{1}^{\text{int}}$ according to:

$$M_{0} = \frac{M(\alpha) \cdot [\cos \beta - \cos \alpha]}{\sin \alpha \cdot [\cos \alpha - 1] - \sin \alpha \cdot [\cos \beta - 1]/R}$$

$$M'_{0} = \frac{M'(\alpha) \cdot [\cos \beta - \cos \alpha]}{\sin \alpha \cdot [\cos \alpha - 1] - \sin \alpha \cdot [\cos \beta - 1]/R}$$

and

$$T_{1}^{\text{int}} = -Tr/ln \left( \frac{\sin \alpha - R\sin \beta}{\cos \sin \alpha - R\cos \beta} \right)$$

where R = M($\alpha$)/M($\beta$) and R' = M'($\alpha$)/M'($\beta$).

Sensitivity for 31P experiments is improved by the use of surface coils. However, using inhomogeneous resonators typically results in a range of flip angle distributions over the sensitive volume. Accurate calculation of the rate constant measured by 31P FAST require very accurate flip angles. Accurate flip angles associated with a surface coil can be achieved using a phase alternated-B1 insensitive rotation (BIRP) (14) acquisition scheme that subtraction averages free induction decay acquired using pairs of four segment B1 insensitive rotation (BIR-4) (15) excitation pulses 180° out of phase. The BIR-4 adiabatic pulses are amplitude and phase modulated (Fig. 1) to produce homogeneous flip angle distributions using inhomogeneous resonators. Commonly, adiabatic pulses are constrained to give 90° or 180° flip angles. However, BIR-4 pulses provide user selected adiabatic plane wave rotation by using abrupt phase discontinuities after the first and third segments. Generally, system limitations implementing the abrupt phase changes result in flip angle errors on the order on 10°. The BIRP, phase alternated, scheme has been shown to average out positive and negative flip-angle errors, whereas, BIR-4 pulses resulted in flip angle errors of $-7^\circ$ to $+9^\circ$, in a direct comparison the nominal flip angle differed from the actual value by only $-1.7^\circ$ to $+1.3^\circ$ when using BIRP (14).

METHODS

Animal Preparation

Animal experiments were performed in accordance with the ARRIVE guidelines on ethics and were approved by the Institutional Animal Care and Use Committee. Male Sprague-Dawley rats (200–300 g; n = 4) were initially anesthetized with 2% isoflurane in air. Animals were secured in a holder with a stereotaxic headset and placed in the magnet. Once in the magnet, isoflurane was reduced to 1.2% for 30 min before beginning data acquisition. Isoflurane was delivered in air at a flow rate was 1–2 L/min using a calibrated vaporizer. Rectal temperature was
Heart rate and blood oxygen saturation level (SaO2) were recorded using a MouseOx system (STARR Life Science, Oakmont, PA) and parameters were maintained within normal physiological ranges.

Shimming and positioning were performed using the 1H frequency. The first 31P data sets were acquired after 30 min of exposure to 1.2% isoflurane. The isoflurane was raised to 2% for 30 min and 31P measurements were repeated. The 31P measurements were repeated at 1.2% and 2.0% isoflurane with 30 min exposures to stabilize before acquiring each of the subsequent data sets.

MR Experiments

MRI was performed on an 11.7T Bruker Biospin Magnet using a custom-made concentric loop 1H/31P (500/202.5 MHz) 2/1.5-cm diameter transceiver surface coil. The 1H (500 MHz) element was used for positioning and shimming before 31P NMR. 31P magnetization transfer data was acquired using the FAST method to determine the creatine kinase rate. Four spectra acquired with $\alpha = 30^{\circ}$ and $\beta = 60^{\circ}$ flip angles with and without $\gamma$-ATP saturation (TR = 1100s, NA = 64, DS = 6). Accurate flip angles throughout the brain were set using BIRP excitation scheme. Narrowband ATP saturation with negligible bleed over was achieved using the BISTRO (16) saturation scheme with eight 50 ms hyperbolic secant RF pulses interleaved with dephasing gradients. Total acquisition time for a $k_{f,CK}$ measurement was ~5 min.

Data Analysis

31P spectra were analyzed using Bruker Topspin software. The chemical shift of the PCr resonance peak was set to zero. Integral values for seven resonance peaks (PME, Pi, PDE, PCr and the three adenosine triphosphate peaks: $\alpha$-, $\beta$-, and $\gamma$-ATP) were acquired in the data set. The spectra baseline and zero and first order phase were corrected. $k_{f,CK}$ is calculated using Eqs. [1–4], using four spectra acquired with $\alpha = 30^{\circ}$ and $\beta = 60^{\circ}$ flip angles with and without $\gamma$-ATP saturation. Data analysis used codes written in Matlab (MathWorks Inc, Natick, MA). A paired two-tailed Student t-test was used to evaluate metabolic biomarkers between isoflurane conditions. Values in text and in graphs are mean ± SEM with $P < 0.05$ considered significant.

RESULTS

The BIR-4 pulses used were 50× the length of optimized square excitation pulses. The increased pulse duration allows $T_2$ dephasing in the rotating frame during excitation. Compared with conventional square pulses, BIR-4 pulses eliminated the short $T_2$ phospholipids that contaminated metabolite signals, thereby improving quantification of signal amplitudes (Fig. 2). The BIRP acquisition scheme with phase alternation scheme averaged out positive and negative flip-angle errors of the BIR-4 acquisition, thereby improving $T_1$ fitting, as demonstrated by measurements of the Pi peak in a dead rat (Fig. 3).

A typical 31P data set consisting of the four spectra used to calculate $k_{f,CK}$ in the FAST method is shown in Figure 4. Spectra were acquired at $60^{\circ}$ and $30^{\circ}$, with and without BISTRO saturation of the $\gamma$-ATP resonance (~2.3 ppm). The pair of spectra acquired without saturation was used to calculate $M_0$ of PCr. The pair of spectra acquired with saturation was used to calculate $M'_0$ and $T_1^{int}$ of PCr. The change in PCr signal was robustly detected, allowing for reproducible measurements of the forward CK rate ($k_{f,CK}$).

The modulation in Pi amplitude, theoretically, also allows for similar calculations of the forward ATPase rate
PCr \equiv * \\

the metabolic flux. This approach has potential applica-

tion for studying neurological disorders with metabolic
dysfunction.

Sauter and Rudin (11) and Du et al (4) have previously
reported $k_{f,CK}$ albeit at much lower temporal resolution.
Sauter and Rudin used a conventional $^{31}$P saturation
transfer method at 4.7T to measure forward CK rate and
high-energy phosphate concentrations under 1–2% halothane,
thiopental sodium and graded bicuculline (0.4 mg/kg and 0.8 mg/kg) and found $k_{f,CK}$ to be $0.25 \pm 0.02$
$s^{-1}$, $0.21 \pm 0.03$ s$^{-1}$, $0.30 \pm 0.04$ s$^{-1}$ and $0.49 \pm 0.04$ s$^{-1}$,
respectively, in normal animals. $k_{f,CK}$ linearly correlated
with electroencephalograph (EEG) activity. The ATP levels
remained constant, while PCr decreased with increased EEG activity. In contrast to expectations, PCr did not increase with decreased EEG activity. These finding demonstrated that $k_{f,CK}$ is a sensitive reliable indicator of changes in metabolic activity, whereas the concentrations of ATP and PCr did not provide consistent useful information.

Du et al (4) used variations of the saturation transfer technique at 9.4T to measure ATP synthesis, including the forward CK and ATP$_{as}$ rates in rats under different depths of anesthesia. The concentrations of the high-energy phosphates, forward ATP$_{as}$ and CK rates and the Spectral Entropy Index of EEG were measured in rats anesthetized using isoflurane (2%), $\alpha$-chloralose, low dose pentobarbital, and high dose pentobarbital (from low to high (isoelectric) anesthetic depth). They found $k_{f,CK}$ to be $0.24 \pm 0.02$ s$^{-1}$, $0.21 \pm 0.03$ s$^{-1}$, $0.21 \pm 0.02$ s$^{-1}$ and $0.19 \pm 0.03$ s$^{-1}$ for animals anesthetized with 2.0% isoflurane, $\alpha$-chloralose, low dose pentobarbital, and high dose pentobarbital, respectively. [PCr] decreased $8 \pm 2\%$ and [Pi] increased $42 \pm 6\%$ in the high dose pentobarbital (isoelectric state) compared with low dose pentobarbital anesthesia. It was concluded that the ATP metabolic rates measured by $^{31}$P MT are more sensitive measures of brain bioenergetics than concentrations of the high-energy phosphates.

Our reported values for the creatine kinase rates under
graded isoflurane anesthesia are in general agreement
with studies by Sauter and Rudin and Du et al (4,11),
although the experimental conditions and type or level
of anesthesia differed. In addition, our results also
showed that after the isoflurane level was returned from
2% to 1.2% for 30 min, the CK rate recovered but did
not reach the prior 1.2% isoflurane level. The CK rate dropped again after another 30
min exposure to 2.0% isoflurane.

**DISCUSSION**

We implemented the accelerated $^{31}$P FAST protocol at
11.7T and measured the concentrations of ATP and PCr,
and the forward CK rate of ATP synthesis with a temporal
resolution of 5 min. The major findings were: (i) the forward creatine kinase rate and the metabolic flux of the rat brain were reliably measured, and (ii) changing isoflurane concentration from 1.2% to 2.0% did not change the PCR and ATP concentrations but significantly decreased the forward creatine kinase synthesis rate and the metabolic flux. This approach has potential applica-
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2% to 1.2% for 30 min, the CK rate recovered but did
not reach the prior 1.2% isoflurane level. The CK rate dropped again after another 30
min exposure to 2.0% isoflurane. These findings suggest that 30 min may not be sufficient for metabolic rate to fully recover and that

<table>
<thead>
<tr>
<th>Isoflurane</th>
<th>PCr, mM</th>
<th>ATP, mM</th>
<th>$k_{f,CK}(s^{-1})$</th>
<th>$F_{f,CK}(\mu mol/g/min)$</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.20%</td>
<td>5</td>
<td>3</td>
<td>0.26±0.02</td>
<td>70.8±4.6</td>
</tr>
<tr>
<td>2.0%</td>
<td>4.75±0.05</td>
<td>2.78±0.13</td>
<td>0.16±0.01*</td>
<td>41.0±4.2*</td>
</tr>
<tr>
<td>1.20%</td>
<td>5.25±0.10</td>
<td>2.87±0.09</td>
<td>0.20±0.01*</td>
<td>58.0±2.6</td>
</tr>
<tr>
<td>2.0%</td>
<td>5.05±0.15</td>
<td>2.91±0.18</td>
<td>0.17±0.02*</td>
<td>45.1±6.8</td>
</tr>
</tbody>
</table>

*Acclimation of 30 min was given after switching isoflurane concentration. Baseline concentration of PCr was assumed to be 5 mM and ATP 3 mM. N = 4, mean ± SEM.

*P < 0.05 with unpaired t-test for comparison with initial 1.2% isoflurane condition.

PCr = phosphocreatine; ATP = adenosine triphosphate; ADP = adenosine diphosphate; Cr = creatine; CK = creatine kinase.
commonly used isoflurane levels can significantly alter cerebral metabolism.

**Alternative Techniques**

Alternatively to $^{31}$P MT, magnetic resonance measurements of cerebral metabolism can be made by $^{13}$C or $^{17}$O or blood oxygen level dependent (BOLD) functional MR techniques (17,18), all measure different aspects of metabolism. $^{13}$C studies use $^{13}$C labeled glucose infusions to measure glucose consumption in the GABAergic tricarboxylic acid cycle (19). $^{17}$O measures cerebral metabolic rate of oxygen. $^{17}$O NMR techniques resemble positron emission measurements of oxygen consumption, both using inhaled $^{17}$O labeled oxygen gas (20). Alternatively, $^{17}$O labeled water ($H_2^{17}$O) may be injected before spectroscopic measurement. It may be of interest to compare different measures of metabolic parameters.

**CONCLUSIONS**

This study implemented and used the $^{31}$P FAST technique at 11.7T to evaluate cerebral high-energy phosphates and creatine kinase synthesis rate under graded isoflurane anesthesia. The advantage of the $^{31}$P FAST technique is that the measurement of creatine kinase synthesis is made practical. A drawback of the $^{31}$P FAST technique is that it is likely less robust than the full saturation recovery MT technique with very long acquisition time. However, the use of high field and small surface coil as well as optimized $^{31}$P FAST acquisition parameters and radiofrequency pulses enable robust measurement of the CK synthesis rate. Further improvement in sensitivity is needed to robustly measure ATPase rate ($k_f$,$\text{ATPase}$). Future studies will incorporate localization by single voxel spectroscopy and chemical shift imaging.

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The authors declare no conflict of interest

**REFERENCES**


