1H and 31P NMR Spectroscopy for the study of brain metabolism at Ultra High Magnetic Field from Rodents to Men

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Spectroscopie RMN du $^1$H et $^{31}$P pour l’Étude du Métabolisme Cérébral à Très Haut Champ Magnétique du Rongeur à l’Homme

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ABSTRACT

$^1$H and $^{31}$P nuclear magnetic resonance spectroscopy allows to detect and to measure \textit{in vivo} and non-invasively the concentrations of biologically relevant compounds associated to metabolic processes such as neurotransmission (glutamate, GABA), neuronal and glial density (N-acetyl-aspartate, myo-inositol) and energetic metabolism (phosphocreatine, ATP) among others. Knowledge of the biochemical profile provides a mean to evaluate the metabolic state of the brain in pathological cases or in evolving physiological conditions, such as aging. Yet, the neural basis of age-related cognitive dysfunction in normal brain aging remains to be elucidated and it has been shown to develop at different rates depending on the structural region.

At ultra-high magnetic fields, magnetic resonance spectroscopy (MRS) benefits from an increased signal-to-noise ratio and a higher chemical shift dispersion, resulting in an increased sensitivity and spectral resolution. To exploit these advantages, $^1$H and $^{31}$P longitudinal studies were carried out \textit{in vivo} at 17.2 Tesla in the aging rat brain to evaluate the progressive metabolic changes within the same individuals from the ages of 1 to up to 22 months of age using two rat cohorts with 1 and 8 months of age at the beginning of the study. For the $^1$H MRS studies, $T_1$ and $T_2$ metabolite relaxation times were measured at each exam in order to control age-related variations and to calculate absolute metabolite concentrations. $^1$H neurochemical profiles from four volumes of interest (VOI) in the brain were studied, revealing a progressive increase in myo-inositol and macromolecule content throughout the brain. In our main VOI composed mostly of cortex but also of corpus callosum and hippocampus, increased levels of choline-containing compounds (tCho) and glutamine were also observed, suggesting a mild neuroinflammation. No changes in NAA were observed in our main VOI, the thalamus or the caudate putamen (striatum). $T_2$ decreases were observed with age for total NAA, tCho and macromolecules. Notably, unexpected effects correlated with the number of NMR exams were observed, the most prominent effect being an increase of the $T_1$ relaxation times of the majority of metabolites.

The second axis of the work done during this thesis was to set up an experimental framework for MR spectroscopic imaging (MRSI) studies at 7 Tesla in the human brain. 2D MRSI pulse sequences were developed for the acquisition of $^{31}$P and $^1$H metabolite maps using either slab selection or STEAM localization, respectively. A WET water suppression scheme was numerically optimized for its application at 7 T. Static $B_1$-shimming configurations were implemented to reduce the inhomogeneity of the excitation field in the volume of interest and to generate outer-volume suppression (OVS) “ring” modes to saturate the signal in the periphery of the head. This approach allows to reduce the energy deposition in comparison to conventional OVS bands. Experiments were done \textit{in vitro} showing their feasibility. The performance of standard OVS bands was also compared to a $B_1$-insensitive train to obliterate signal (BISTRO) scheme \textit{in vivo} using a double-tuned $^1$H/$^{31}$P phased-array coil in a single-channel configuration for transmission. The demonstrated suppression efficacy of BISTRO opens the way for its use as a frequency-selective pre-saturation module for future $^{31}$P magnetization transfer experiments for the study of brain energy metabolism at very high magnetic field.
RESUME

La Spectroscopie RMN (SRMN) du $^1$H et du $^{31}$P permet de détecter et de mesurer in vivo de façon non-invasive la concentration de composés biologiques qui sont pertinents à l’étude des aspects variés du métabolisme cérébral comme la neurotransmission (glutamate, GABA), la densité neuronale (N-acetyl-aspartate) et gliale (myo-inositol) ou le métabolisme énergétique (phosphocreatine, ATP), entre autres. Ainsi, l’analyse des profils biochimiques permet d’étudier longitudinalement l’évolution de la physiologie cérébrale en conditions pathologiques ou normales. Par ailleurs, à ultra-haut champ magnétique la SRMN bénéficie d’une sensibilité et d’une résolution spectrale accrues, maximisant l’information métabolique exploitable.

Au cours de cette thèse, nous nous sommes surtout intéressés à l’étude du vieillissement cérébral normal. Une étude longitudinale en $^1$H et $^{31}$P a été menée in vivo à 17.2 Tesla afin de suivre les altérations métaboliques pendant 14 mois chez deux cohortes de rats Dark Agouti âgés d’un mois et 8 mois au départ de l’étude. Les concentrations ainsi que les temps de relaxation $T_1$ et $T_2$ de plus de 20 métabolites ont été mesurés jusqu’à l’âge de 22 mois. Nous avons notamment observé une augmentation des concentrations de myo-inositol et des macromolécules dans les 4 volumes d’intérêt (VOI) étudiés. Dans le VOI Main, comprenant principalement du cortex mais aussi du corps calleux et de l’hippocampe, ces changements métaboliques ont été accompagnés par une augmentation des niveaux de glutamine et de composés contenant de la choline (tCho). Ces observations sont cohérentes avec une possible neuro-inflammation modéré au cours du vieillissement. Aucun changement du NAA a été observé sur le Main VOI, thalamus et putamen caudé (striatum). Additionnellement, une réduction des temps $T_2$ pour le NAA total, la tCho et les macromolécules a été observée, en accord avec une altération du milieu cellulaire et une accumulation de fer dans les tissus avec l’âge. Etonnamment, nous avons observé un effet corrélé avec le nombre d’examens RMN, qui a été fortement manifesté par une augmentation significative des temps $T_1$ de nombreux métabolites.

Un deuxième axe de travail pendant cette thèse a été la mise en place des outils méthodologiques nécessaires à la réalisation des études par SRMN du $^1$H et du $^{31}$P à 7 Tesla chez l’homme. Des séquences d’imagerie spectroscopique 2D ont été développées pour obtenir des cartes de concentration des métabolites $^{31}$P et $^1$H respectivement par la sélection d’une coupe ou bien d’un voxel par écho-stimulé. Un schéma de suppression d’eau WET a été optimisé pour son application à 7 T. Des modes d’excitation et de saturation du signal extérieur (OVS) en « anneau » ont été implémentés avec la méthode de transmission parallèle pour son application en imagerie spectroscopique $^1$H par l’optimisation des configurations statiques d’excitation ou « shimming-B$_1$ ». Cette approche a permis d’appliquer des champs d’excitation plus homogènes et de réduire le dépôt d’énergie chez le sujet par rapport à l’utilisation des bandes OVS classiques. Des expériences in vitro ont été menées pour démontrer leur faisabilité. Enfin, un module de saturation BISTRO a été implémenté pour l’acquisition in vivo de cartes métaboliques en $^{31}$P. L’efficacité du module BISTRO a été démontrée et ce module peut être adapté pour des expériences $^{31}$P de transfert d’aimantation, ouvrant la voie de l’étude du métabolisme énergétique cérébral chez l’homme à très haut champ magnétique.
GENERAL INTRODUCTION

The study of brain metabolism and its associated biochemistry is of great relevance for the understanding of evolving physiological states such as brain maturation and aging but also for the characterization and monitoring of brain pathologies, namely the neurodegenerative diseases. With the improvement in our healthcare systems over the last decades, the average life expectancy is increasing and the proportion of the aged population (above 60 years old) is expected to double worldwide over the next 50 years.

Aging is the primary non-genetic risk factor associated to the development of neurodegenerative diseases and among them, Alzheimer’s disease (AD) alone is the 6th leading cause of death in the USA. In 2000, dementia affected almost 30% of the aged population in France between 85 and 89 years old. A first step for the comprehension and establishment of early biomarkers in the development of neurodegenerative diseases is therefore the understanding of normal aging. It is therefore crucial to characterize the progressive changes taking place during normal brain aging from a molecular point of view.

Nuclear magnetic resonance spectroscopy (MRS) is a non-invasive technique that allows measuring the concentrations of biologically-relevant molecules involved in major metabolic pathways, with a millimolar sensitivity. These molecules are usually referred to as “metabolites” and are involved in different metabolic functions, such as neurotransmission (glutamate, γ-aminobutyric acid or GABA), neuronal and glial densities (N-acetyl-aspartate, myo-inositol), membrane turnover (choline-containing compounds, NMR-visible proteins) and energy metabolism (Creatine, ATP) among others. Other imaging techniques exist, such as Positron Emission Tomography (PET), which are sensitive enough to detect picomolar concentrations. However, they require radioactive exogenous tracers whereas typical MRS detects endogenous molecules as signal sources.

The signal intensity detected in MR experiments is directly influenced by the strength of the static magnetic field. With the advent of increased magnetic field strengths, the MRS technique can reach new horizons as higher spectral and spatial resolutions can be explored, improving the reliability and specificity of MRS measurements.

But with greater fields come greater challenges, and technical difficulties arise due to the increased inhomogeneity of the radiofrequency (B₁) and static magnetic fields (B₀). The application of radiofrequency (RF) pulses, necessary for the generation of the measured NMR signal, also contribute to the challenges faced at ultra-high fields (UHF) as their required RF energy scales up with the magnetic field intensity. Thus MR scientists must be very attentive to the maximum power output provided and sustained by the hardware but even more to the energy absorbed by the head through the dissipation of the transmitted RF power, potentially leading to temperature elevation above safety limits.
To counteract such limitations at UHF, the introduction of state-of-the-art methods is necessary for the acquisition of robust and high-quality spectra. The technical and safety constrains being less restrictive for animal studies allow the use of a wider panel of techniques to mitigate the consequences of $B_1$ inhomogeneity, namely by using energy-demanding adiabatic RF pulses. The application of such pulses being quite limited for human studies at UHF, an approach based on the efficient use of the available power is preferred. Parallel transmission (pTx) is one of the most promising solutions to tackle $B_1$ field inhomogeneities in the human brain. It makes an efficient use of the available allowed power by adapting the phase and amplitude of each resonating element according to each volunteer and the targeted $B_1$-field. Furthermore, phase interference patterns can be advantageously exploited to remove unwanted signals generated from tissue outside of the skull.

MRS measurements of nuclei other than proton ($^1$H), such as phosphorous ($^{31}$P), are difficult due to their intrinsically low sensitivity. This reduced sensitivity is somewhat compensated at UHF by the raw increase in signal. ATP and phosphocreatine which can be detected using $^{31}$P MRS, are particularly important metabolites due to their key roles in energy metabolism. Therefore, acquiring $^{31}$P metabolic profiles could complement the already rich metabolic information derived from $^1$H neurochemical profiles.
THESIS OVERVIEW

The thesis project was done at the Magnetic Resonance Imaging and Spectroscopy Unit (UNIRS, for its acronym in French), attached to NeuroSpin at the Commissariat à l’Energie Atomique et aux Energies Alternatives (CEA) located at Gif-sur-Yvette, Essonne, France. Several horizontal scanners are currently present at NeuroSpin, including three Bruker MRI scanners at 7, 11.7 and 17.2 T dedicated to rodent studies and also two Siemens MRI scanners at 3 and 7 T dedicated to human studies. The 7 T clinical scanner is also equipped with parallel transmission capabilities, allowing to control 8 individual transmission channels for RF coils built for such modality.

In this context, the presented work focused on the implementation of $^1$H and $^{31}$P MRS sequences at 17.2 T and their application to study longitudinally normal brain aging in the Dark Agouti rat. In parallel, $^1$H and $^{31}$P MRSI pulse sequences were developed for their application in the human brain at 7 T using multi-channel or single-channel volume coil transceivers.

The presented manuscript is divided in three parts:
The first part explains the methodological context of this thesis. Chapter 1 starts by introducing the fundamental concepts of NMR spectroscopy. Afterwards, it provides a brief presentation of the RF pulses and pulse sequences used as well as related aspects such as energy deposition and chemical shift artifacts. The chapter ends by providing a prospective view of the gains and challenges of MRS at UHF. Chapter 2 presents all the practical considerations and technical implementations. It includes a brief description of the 7 T clinical and 17.2 T preclinical scanners, the implementation and in vitro validation of the BISTRO scheme on both systems as well as the numerical optimization of WET water suppression schemes. The methodology related to the $^1$H and $^{31}$P spectral decompositions using LCModel is presented, detailing the macromolecule baseline parameterization used for the analysis of rat brain $^1$H spectra. The absolute metabolite quantification strategy for $^1$H and $^{31}$P data is explained. Finally, practical considerations for the acquisition of water $T_1$ maps at 17.2 T are presented.

In the second part, the work done at 17.2 T using in vivo quantitative $^1$H and $^{31}$P MR spectroscopy is presented. Chapter 3 presents a longitudinal $^1$H MRS study of brain aging in the Dark Agouti rat over a span of 18 months. The $T_1$ and $T_2$ relaxation times measured from one VOI as well as metabolite concentrations from four VOIs in the young rat at 1 month of age are presented. These relaxation times are compared to those measured at lower magnetic fields so as to assess their field-dependency. The evolution of $T_1$ and $T_2$ relaxation times as well as the neurochemical profiles from the 4 VOIs are then presented and examined using a linear regression analysis. Chapter 4 presents a preliminary $^{31}$P quantitative MRS study looking at a subset of our previously studied Dark Agouti rats at ages of 5, 17 and 21 months old. Results from the acquired metabolite profiles and measured $T_1$ times from 8 animals are presented. The limitations and encountered problems, notably due to chemical shift artifacts, are discussed.
The third part presents our MRSI developments and the results obtained on the 7 T scanner. In Chapter 5, the parallel transmission approach is explained along with our methodological framework and the static B₁-shim configurations known as the coil eigenmodes or “rings”. The use of such rings as OVS bands are evaluated and compared to numerically optimized B₁-shim configurations in vitro. Chapter 6 presents the evaluation of ¹H and ³¹P MRI and MRSI sequences for the constitution of a research protocol in the human brain including ¹H and ³¹P metabolite maps acquisitions. The numerically optimized WET scheme is validated in vivo in combination with a 2D CSI-STEAM sequence. Adiabatic and conventional OVS bands configurations are tested and compared in terms of efficiency and energy requirements. The chapter ends showing preliminary in vivo ¹H/³¹P MRSI data.

This thesis ends with a general conclusion summarizing the main results and few perspectives for future work in MR spectroscopy are given.
PART I: METHODOLOGICAL CONTEXT

1. THEORETICAL ASPECTS

1.1. BASIC PRINCIPLES OF NMR

1.1.1. The Zeeman effect

In quantum mechanics, particles such as electrons, protons or neutrons possess an intrinsic angular momentum or spin which may only have discrete values. This angular momentum \( \mathbf{L} \) depends on the spin quantum number \( I \) as:

\[
\mathbf{L} = \hbar \sqrt{I(I+1)} \quad ; \quad \hbar = \left( \frac{\hbar}{2\pi} \right)
\]

(1.1)

where \( \hbar \) is Planck’s constant and \( I \) is integer or half-integer. The spin number \( I \) is unique to each nucleus in its stable ground state. For instance, nuclei possessing an odd mass number have a half-integer spin, such as the hydrogen (\(^1\)H) and phosphorus (\(^31\)P) which both possess a spin of +1/2. The “direction” or state of the angular momentum is determined by the quantum number \( m \) which may have only \( 2I + 1 \) values ranging as \( m = -I, -I+1, \ldots, +I \) and correspond to the basis states of the particle. The component of the angular momentum in the z direction is given by:

\[
\mathbf{L}_z = \hbar m.
\]

(1.2)

The corresponding magnetic moment along the z axis is given by:

\[
\mu_z = \gamma \hbar m.
\]

(1.3)

In an external magnetic field \( \mathbf{B}_0 \) oriented along the z axis, the nucleus acquires a magnetic energy \( E \) given by:

\[
E = -\mu_z B_0 = -\gamma \hbar m B_0.
\]

(1.4)

This Zeeman effect states that, for a particle with a spin \( I = 1/2 \), the energy difference between its two possible quantum states is:

\[
\Delta E = \gamma \hbar B_0.
\]

(1.5)

The energy required to transition from one energy state to the other corresponds to a photon or electromagnetic wave with an energy \( \Delta E \) and a frequency \( \nu_0 \) proportional to the magnetic field \( B_0 \) (figure 1.1A).
This resonance frequency is called the Larmor frequency and is given by:

$$\nu_0 = \left(\frac{\gamma}{2\pi}\right)B_0$$  \hspace{1cm} (1.6)

The basic principle of NMR is to apply a transverse radiofrequency (RF) wave at the Larmor frequency (usually in the 10 to $10^3$ MHz range) to excite nuclear spins and then detect the electromagnetic wave emitted when the spins return to their lower energy state (figure 1.1B). Notably, equation (1.5) shows that increasing $B_0$ intensity leads to more energy being involved in the quantum state transitions.

Figure 1.1. The Zeeman Effect and the NMR phenomenon.
A. Upon the application of an external magnetic field $B_0$, quantum energy states for a spin $I (= \frac{1}{2})$ split depending on its quantum number $m$ ($\frac{1}{2}$ or $-\frac{1}{2}$). The energy difference between the two spin states is proportional to the magnetic field strength as well as the Larmor frequency of the radiofrequency wave used to excite the spins. B. The number of spins with a preference for the low energy state increases with the magnetic field, leading to higher polarization in the macroscopic sample.

1.1.2. Boltzmann distribution

When considering a macroscopic sample with a large number of spins in an external magnetic field, the methods of Statistical Physics can be used to calculate its macroscopic magnetization. For the case of a population of spins of $I = 1/2$, only two spin states are allowed: $m = +1/2$ ($\mu$ parallel to $B_0$) or $m = -1/2$ ($\mu$ antiparallel to $B_0$), respectively referred to as the $\alpha$ and $\beta$ spin states. Under thermal equilibrium conditions, the two corresponding energy levels are populated according to the Boltzmann probability distribution:

$$N_\alpha, N_\beta = \frac{\exp(\pm \gamma hB_0/2k_BT)}{\exp(-\gamma hB_0/2k_BT)+\exp(\gamma hB_0/2k_BT)}$$ \hspace{1cm} (1.7)

where $k_B$ is the Boltzmann equilibrium constant, $T$ is the absolute temperature and $N_\alpha$ and $N_\beta$ the mean spin populations in the $\alpha$ and $\beta$ spin states. The population difference ratio is given by:
\[
\frac{N_\alpha}{N_\beta} = \exp\left(\frac{\gamma \hbar B_0}{k_B T}\right) \tag{1.8}
\]

Since at room temperature the thermal energy \(k_B T\) is five orders of magnitude larger than \(\gamma \hbar B_0\), the population ratio can be approximated to:

\[
\frac{N_\alpha}{N_\beta} \sim 1 + \left(\frac{\gamma \hbar B_0}{k_B T}\right) \tag{1.9}
\]

which dictates that for \(^1\text{H}\) nuclei at 7 Tesla, the population difference between both energy levels or polarization, corresponds to 0.00464 % of the total spin population while at 17 T it increases to 0.0113 %. Due to their lower gyromagnetic ratio and their lesser natural abundance, nuclei other than \(^1\text{H}\) are more difficult to detect and have lower Larmor frequencies than \(^1\text{H}\). Table 1.1 summarizes the NMR characteristics of the nuclei studied in vivo (Bernstein, et al., 2004 p. 960; de Graaf, 2007 p. 9).

<table>
<thead>
<tr>
<th>Isotope</th>
<th>Spin</th>
<th>Natural abundance (%)</th>
<th>Gyromagnetic ratio (\gamma/2\pi) (MHz/T)</th>
<th>NMR frequency in MHz</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>3.0 T</td>
</tr>
<tr>
<td>(^1\text{H})</td>
<td>1/2</td>
<td>99.985</td>
<td>42.58</td>
<td>127.728</td>
</tr>
<tr>
<td>(^2\text{H})</td>
<td>1</td>
<td>0.015</td>
<td>6.54</td>
<td>19.608</td>
</tr>
<tr>
<td>(^3\text{He})</td>
<td>1/2</td>
<td>1.4x10(^{-4})</td>
<td>32.43</td>
<td>97.302</td>
</tr>
<tr>
<td>(^7\text{Li})</td>
<td>3/2</td>
<td>92.580</td>
<td>16.55</td>
<td>49.638</td>
</tr>
<tr>
<td>(^13\text{C})</td>
<td>1/2</td>
<td>1.108</td>
<td>10.70</td>
<td>32.112</td>
</tr>
<tr>
<td>(^14\text{N})</td>
<td>1</td>
<td>99.630</td>
<td>3.08</td>
<td>9.231</td>
</tr>
<tr>
<td>(^15\text{N})</td>
<td>1/2</td>
<td>0.370</td>
<td>-4.32</td>
<td>-12.948</td>
</tr>
<tr>
<td>(^17\text{O})</td>
<td>5/2</td>
<td>0.037</td>
<td>5.77</td>
<td>17.316</td>
</tr>
<tr>
<td>(^19\text{F})</td>
<td>1/2</td>
<td>100</td>
<td>40.05</td>
<td>120.156</td>
</tr>
<tr>
<td>(^23\text{Na})</td>
<td>3/2</td>
<td>100</td>
<td>11.26</td>
<td>33.786</td>
</tr>
<tr>
<td>(^31\text{P})</td>
<td>1/2</td>
<td>100</td>
<td>17.24</td>
<td>51.705</td>
</tr>
<tr>
<td>(^39\text{K})</td>
<td>3/2</td>
<td>93.10</td>
<td>1.99</td>
<td>5.964</td>
</tr>
<tr>
<td>(^\infty\text{Xe})</td>
<td>1/2</td>
<td>26.44</td>
<td>11.78</td>
<td>35.331</td>
</tr>
</tbody>
</table>

*Table 1.1. NMR properties of most nuclei used for in vivo NMR studies.*

### 1.1.3. Macroscopic Magnetization

All individual magnetic moments in a sample add up to a net macroscopic magnetic moment \(M\). Although the spins are not parallel to the main magnetic field \(B_0\) and have rotational components on the xy-plane perpendicular to the axis of the static magnetic field (referred as the transverse plane), there is no net transverse component of \(M\) as the phase of the spins is randomly distributed. Nevertheless, there will be a net longitudinal magnetization component \(M_0\) along the
direction of $B_0$ proportional to the population difference $\overline{N}_\alpha - \overline{N}_\beta$ (figure 1.2). From eq. (1.3), the expression for $M_0$ at thermal equilibrium can be found:

$$M_0 = \sum_{i=1}^{N} \mu_i = \overline{N}_\alpha \mu_z + \overline{N}_\beta \mu_z = \gamma \left( \frac{\hbar}{2} \right) (\overline{N}_\alpha - \overline{N}_\beta)$$  

(1.10)

where $N$ is the total spin population. At room temperature, one can derive $M_0$ using eq. (1.9):

$$M_0 = \frac{N \gamma \frac{\hbar^2}{4k_B T}}{4k_B T} B_0$$  

(1.11)

This expression reveals the importance of the static magnetic field as well as the gyromagnetic ratio on the magnitude of the macroscopic magnetization. Nevertheless, the interest of the NMR scientist is not the maximum available signal itself but rather the available signal-to-noise ratio (SNR) that is achievable in a given time interval. Other intrinsic factors such as noise levels, sample volume and signal loses due to relaxation time effects also play a role on the determination of the sensitivity.

**Figure 1.2.** The net magnetization $M$ vector in an external magnetic field.

**A.** A single spin precesses around the external static magnetic field $B_0$. The spin magnetic moment $\mu$ has quantized transverse and longitudinal components. **B.** The sum of all spin magnetic moments in either the $\alpha$ or $\beta$ states amount to the macroscopic magnetization vector $M$. The lack of phase coherence in the transverse plane leaves only a net longitudinal component $M_0$.

### 1.1.4. Excitation and the Bloch Equations

At the thermal equilibrium state, the net magnetization vector $M_0$ is aligned along the direction of the main magnetic field $B_0$ and there is no precession motion measurable in the transversal plane. This precession is needed in order to generate a detectable electromotive force (emf) in an electric circuit through induction (Faraday’s law). Thus, a second transversal magnetic field ($B_1$) is applied to “excite” the spins and rotate the magnetization towards the transverse plane. In modern impulsonal NMR, $B_1$ is applied as a short RF pulse at the Larmor frequency (1.6).
Following this RF pulse, the transverse magnetization \( M_{xy} \) will precess around \( \mathbf{B}_0 \) at the Larmor frequency, generating an emf in the receiver coil adjacent to the sample.

In practice, an induction coil is used to generate the RF excitation field. For amplitude-modulated RF pulses (discussed in the following section), their duration and intensity will determine the nutation or “flip angle” experienced by the net longitudinal magnetization. Accordingly, the transverse magnetization \( M_{xy} \) reaches its maximum value \( (M_{xy} = M_0) \) for a 90° flip angle (figure 1.3).

The evolution of the macroscopic magnetization \( \mathbf{M} \) under the effects of an external magnetic field \( \mathbf{B}_{ext} \) composed of both \( \mathbf{B}_0 \) and \( \mathbf{B}_1 \) fields is derived from the following equation:

\[
\frac{\partial \mathbf{M}}{\partial t} = \mathbf{M} \times \gamma \mathbf{B}_{ext} = \mathbf{M} \times \gamma (\mathbf{B}_0 + \mathbf{B}_1)
\] (1.12)

![Diagram](image)

**Figure 1.3. Effect of an RF pulse \( \mathbf{B}_1 \) on the macroscopic magnetization \( \mathbf{M} \).**

At thermal equilibrium, the net magnetization \( \mathbf{M} (= M_0) \) is parallel to the main magnetic field \( \mathbf{B}_0 \). In order to generate a detectable NMR signal, a time-varying transversal RF pulse \( \mathbf{B}_1(t) \) is applied tilting \( \mathbf{M} \) about its axis. By calibrating \( \mathbf{B}_1 \) properly, \( \mathbf{M} \) is rotated by 90° resulting in a maximum amplitude for \( M_{xy} \). The diagram is shown in the rotating frame of reference at the Larmor frequency.

Taking into account the return to thermal equilibrium, equation (1.12) can be expanded to yield the Bloch equations:

\[
\frac{\partial M_x(t)}{\partial t} = \gamma \left[ M_y(t)B_0 - M_z(t)B_{1y} \right] - \frac{M_x(t)}{T_2}
\] (1.13)

\[
\frac{\partial M_y(t)}{\partial t} = \gamma \left[ M_z(t)B_{1x} - M_x(t)B_0 \right] - \frac{M_y(t)}{T_2}
\] (1.14)

\[
\frac{\partial M_z(t)}{\partial t} = \gamma \left[ M_x(t)B_{1y} - M_y(t)B_{1x} \right] - \frac{M_z(t) - M_0}{T_1}
\] (1.15)

The relaxation times \( T_1 \) and \( T_2 \) describe the mono-exponential recovery of \( M_z \) and decay of \( M_{xy} \) as shown in figure 1.4.
Figure 1.4. Effects of $T_1$ and $T_2$ relaxation on the longitudinal $M_z$ and transverse $M_{xy}$ components.

A. After RF excitation, the transverse magnetization $M_{xy}$ decreases exponentially with time corresponding to the dephasing of the spins in the sample. At $t = T_2$, $M_{xy}$ has been reduced to 37% of its initial value. B. After RF excitation, the longitudinal magnetization $M_z$ increases exponentially reaching 63% of its maximal value of $M_0$ at $t = T_1$ and $M_0$ at $t>5T_1$. The $T_1$ relaxation process corresponds to the spin population’s energy surplus being transferred to its surrounding. In biological tissue, $T_1$ is usually an order of magnitude longer than $T_2$.

1.1.5. Longitudinal relaxation time $T_1$

The spin-lattice or $T_1$ relaxation process takes place following a perturbation of the net magnetization $M$ and determines the speed at which the longitudinal magnetization $M_z$ returns to its thermal equilibrium value $M_0$. $T_1$ relaxation originates from a constant growth rate of $M_z$ due to interactions between the excited spins and the lattice, which acts as a *reservoir* that stocks the energy of the excited spin allowing it to return to its minimal energy state. The relaxation process is done through dissipation of energy into the nearby atoms as thermal energy. The process of energy transfer from the excited spins to the lattice does not induce any increase of temperature in the sample as the proton spin energy is minor compared to the thermal energy scale $k_B T$ *in vivo*.

From equation (1.15), the time-dependency of $M_z$ can be derived following an excitation pulse:

$$\frac{\partial M_z(t)}{\partial t} = \frac{M_0 - M_z(t)}{T_1}$$ (1.16)

Solving this partial differential equation provides the expression of the $M_z(t)$ at time $t$:

$$M_z(t) = M_z(0) e^{-t/T_1} + M_0 (1 - e^{-t/T_1})$$ (1.17)

where $M_z(0)$ is the longitudinal magnetization just prior to the 90° excitation pulse.

When the experiment is sequentially repeated with a repetition time $TR$ and no residual transverse magnetization remains at the beginning of each repetition, then a *steady-state* magnetization is achieved after a few repetitions:
where \( \alpha \) is the excitation flip angle (FA). Experimentally, the residual transverse magnetization can be eliminated by applying dephasing gradients or by using a TR longer than \( 5^*T_2^* \). One can notice that the steady-state magnetization is highest at long repetition times (TR > \( 5^*T_1 \)). However, to optimize the SNR per unit time, one should rather use a short TR (such as TR = \( T_1 \)) so as to increase the number of signal averages. For a given TR and \( T_1 \), the optimal excitation angle or Ernst angle is given by the expression:

\[
\alpha_{\text{Ernst}} = \arccos(e^{-\text{TR}/T_1})
\]  

(1.19)

The use of short TRs and corresponding Ernst angle can be an efficient way to accumulate signal in a given time, such as in NMR studies of low sensitivity nuclei like \(^{31}\)P. However, it can be problematic for the quantification of the observed NMR signals especially if they possess distinct \( T_1 \) relaxation times. Alternatively, long TRs can be used if differential \( T_1 \)-weighting wants to be avoided, in particular when the \( T_1 \) relaxation times are unknown.

1.1.6. \( T_1 \) estimation

Several methods are available for the determination of \( T_1 \) relaxation times:

a) The progressive saturation techniques use the dependency of the steady-state magnetization to the repetition time and flip angle. These methods are suited for low-sensitivity nuclei where long scans are needed to achieve an acceptable SNR or whenever the available scanning time is reduced (Freeman, et al., 1971; Hofmann, et al., 2001; Deoni, et al., 2003; Haacke, et al., 1999 pp. 637-653).

b) The Look-Locker method where several data points with varying \( T_1 \)-weighting are acquired within a single TR using a single inversion pulse and a train of small FA. This method is time-efficient and allows estimating an apparent \( T_1 \) relaxation time. Yet it is not suited for NMR spectroscopy experiments as it requires high SNR (Deichmann, et al., 1992; Schmitt, et al., 2004; Rooney, et al., 2007).

c) The single inversion-recovery (SIR) method consists in acquiring \( T_1 \)-weighted data sets for which the delay (inversion time = TI) between the inversion pulse and the excitation is varied while keeping a fixed recovery time (= TR - TI) to prevent partial saturation effects (de Graaf, et al., 2006; Vold, et al., 1968; Lu, et al., 2014).

The SIR method (figure 1.5A) is considered the “gold-standard” for measuring \( T_1 \) relaxation times (Cudalbu, et al., 2009; Wright, et al., 2008). After the inversion pulse, the longitudinal magnetization evolves accordingly to eq. (1.17) as:

\[
M_z(t) = -M_0 e^{-t/T_1} + M_0(1 - e^{-t/T_1}) = M_0(1 - 2e^{-t/T_1}), \quad 0 < t < TI
\]

(1.20)
As depicted in figure 1.5B, the return to the thermal equilibrium $M_0$ will depend on the $T_1$ values of the spin populations in the sample.

In practice, the MRS measurement will consist of numerous repetitions and the steady-state magnetization must be considered instead. For a $90^\circ$ excitation, the expression is the following:

$$M_z = M_0(1 + e^{-TR/T_1} - 2e^{-TI/T_1})$$  \hspace{1cm} (1.21)

**Figure 1.5. Magnetization inversion-recovery method for $T_1$ measurements.**

**A.** Chronogram of the SIR pulse sequence. An inversion pulse is applied at the beginning of the experiment and an excitation pulse is applied after a delay called the inversion time. The recovery time, corresponding to the delay between the excitation pulse and the next inversion pulse, allows $M_z$ to reach a steady-state magnetization. **B.** After inversion of $M_0$, the longitudinal magnetization $M_z$ recovers at a rate determined by the $T_1$ value. By varying the inversion time, better contrasts can be observed between spin populations with different $T_1$ times. Measuring $M_z$ at different TIs allows estimating the $T_1$ value through equation (1.17).

### 1.1.7. Transverse relaxation times $T_2$ and $T_2^*$

The spin-spin or $T_2$ relaxation process takes place in an excited spin population where the net transverse magnetization decreases in magnitude through dephasing of the individual nuclear magnetic moments. A qualitative explanation of the relaxation process is that spins experience local magnetic fields which are a combination of the main magnetic field $B_0$ and small local magnetic field changes created by the neighboring molecules. The variation of the local fields leads to fluctuations of the individual precessional frequencies of the spins, which in turn tend to present a different individual phase. This induces a loss in the phase coherence of the excited spin population. The consequence on the macroscopic $M_{xy}$ component is a reduction in magnitude, the so-called $T_2$ decay.

From equations (1.13) and (1.14), the time-dependency of $M_{xy}$ can be derived (for $B_1 = 0$):

$$\frac{\partial M_{xy}(t)}{\partial t} = -\frac{M_{xy}(t)}{T_2}$$  \hspace{1cm} (1.22)
which leads to:

\[ M_{xy}(t) = M_{xy}(0)e^{-t/T_2} \]  

(1.23)

where \( M_{xy}(0) \) is the transverse magnetization just after the excitation pulse.

In practice, there are additional factors to the dephasing of the transverse magnetization, in particular non-random, local magnetic field inhomogeneities \( \Delta B_0(r) \). These local inhomogeneities are most prominent at the interfaces between media with strong magnetic susceptibility differences, such as brain tissue and air. This apparent relaxation time \( T_2^* \) is shorter than the intrinsic \( T_2 \) relaxation time. Nevertheless, this additional non-random dephasing can be compensated by “refocusing” the individual spins using a spin-echo (SE) or Hahn refocusing pulse sequence (figure 1.6A). If \( \Delta B_0(r) \) is time-independent, the acquired phase of the spins before refocusing at the position \( r \) is:

\[ \phi(r) = \gamma \Delta B_0(r) \cdot TE/2 \]  

(1.24)

where TE/2 is the delay between the excitation and refocusing pulses. As illustrated in figure 1.6B, the application of the 180° refocusing pulse about the xy-plane will cause the individual spins to rotate in the inverse sense prior to the refocusing pulse. Therefore, the phase acquired during the first TE/2 time lapse due the non-random field inhomogeneities will be compensated after the second TE/2 delay. A complete refocalization of the transverse magnetization will occur, forming an echo (figure 1.6B). The TE delay is called the echo-time. The SE sequence is commonly used to estimate \( T_2 \) relaxation times (figure 1.7).

The measurement of intrinsic \( T_2 \) relaxation times can be further improved by using a Carr-Purcell-Meiboom-Gill (CPMG) refocusing pulse train, which is more effective at refocusing diffusion-related dephasing than the Hahn spin-echo sequence. The CPMG pulse train consists of an excitation pulse at time \( t = 0 \) and a train of refocusing pulses at time \( t = (2n+1)\tau \), where the phase of the 90° RF pulse is shifted by 90° with respect to the \( n \) inversion pulses and \( \tau \) is the middle time-point between the excitation pulse and the first 180° pulse. This phase-cycling approach compensates for imperfections in the refocusing pulse’s flip angles, which otherwise would induce a cumulative error with the number of 180° pulses (Bernstein, et al., 2004 pp. 420-422; Meiboom, et al., 1958).
PART I: THEORETICAL CONTEXT

A. The spin-echo pulse sequence consists in a 90° excitation pulse followed by a 180° refocusing pulse applied at TE/2. The NMR signal is refocused at TE and its intensity is $T_2$-weighted. B. After excitation, the NMR signal decreases rapidly due to $T_2^*$ relaxation. However, the application of a 180° refocusing pulse leads to the inversion of the spin dephasing in the xy-plane. The phase acquired due to non-random field inhomogeneities before refocusing is cancelled out and a “spin-echo” is formed.

Figure 1.6. The Spin-echo experiment.

Figure 1.7. Measurement of $T_2$ relaxation times.

After excitation, the transverse magnetization $M_{xy}$ decays exponentially according to equation (1.23). By employing a Hahn SE sequence (or better yet, a CPMG refocusing pulse train), $T_2^*$ effects (and diffusion-related signal loss) are compensated and values closer to the intrinsic $T_2$ can be estimated.
1.1.8. NMR Spectrum and the Discrete Fourier Transform

As it was described in section 1.1.4, when the transverse magnetization $M_{xy}$ is non-null, a detectable emf oscillating at the Larmor frequency is induced on the received coil positioned in the transverse plane. Following the $T_2^*$ decay of the transverse magnetization, a time-varying complex signal designated as the free induction decay (FID) will be acquired by the NMR spectrometer. This FID can be examined in the frequency domain by applying a complex Fourier transform, resulting in a NMR “spectrum”. In practice, the number of sampled complex values of the FID is finite and the calculation of the spectrum is done instead by applying a discrete Fourier transform (DFT), replacing the theoretical integral by a finite summation. When the number of registered time-points is a power of 2 (such as $2048 = 2^{11}$) then a fast Fourier transform (FFT) can be applied (figure 1.8) which is computationally more efficient than the DFT (Cooley, et al., 1965; Brigham, 1988).

Once phased, the real and imaginary components of the spectrum correspond to Lorentzian absorption and dispersion lineshapes. For the analysis of the NMR spectrum, the better resolved absorption spectrum is analyzed allowing for the measure of the resonance frequency $\nu$, the peak intensity, area and linewidth at half maximum (FWHM or $\Delta\nu_{1/2} = (\pi T_2^*)^{-1}$). If needed, the spectrum can be analyzed in magnitude but the lineshape is broader due to the dispersive component.

![Figure 1.8. Free induction decay and NMR spectrum](image)

The duality between the measured FID in the time domain and its corresponding spectrum in the frequency domain is established by the application of a discrete Fourier transform. The real and imaginary components of the FID and spectrum are shown in blue and red, respectively.
PART I: THEORETICAL CONTEXT

1.1.9. Chemical Shift

Nuclei are surrounded by local magnetic fields, in particular the magnetic field created by the movement of electrons in the external magnetic field $B_0$. This induced magnetic field opposes $B_0$ (accordingly to Lenz’s law) and its intensity depends on the density of the electronic cloud around the considered nucleus. Thus, the nuclear spins experience a magnetic “shielding” whose characteristics depend on the molecule the nucleus belongs to or on the molecules in its immediate vicinity. As a consequence, the Larmor frequency will be shifted by a factor $\sigma$ known as the chemical shift expressed in parts per million (ppm):

$$\omega_0 = 2\pi \nu_0 = \gamma (B_0 - \sigma)$$

(1.25)

A positive $\sigma$ will correspond to a shielding (higher electronic density) and a negative one will correspond to an anti-shielding (lower electronic density) and it may vary due to changes in the chemical environment such as pH or temperature. The definition of the chemical shift of a group of equivalent nuclei is:

$$\sigma = \frac{\nu_0 - \nu_{ref}}{\nu_{ref}}$$

(1.26)

where $\nu_{ref}$ is the resonance frequency of a reference compound. The standard reference compounds used for system and coil calibrations is tetra-methyl-silane (TMS) for $^1$H MRS and phenyl-phosphonic acid (PPA) or hexa-methyl-phosphorous-triamide (HMPT) for $^{31}$P MRS.

1.1.10. J-Coupling

When two nuclei are involved in a covalent bound, the nuclear spins interact indirectly through their shared electrons. This scalar, spin-spin or J-coupling results in a splitting of their respective resonances into hyperfine structures. Scalar coupling originates from intramolecular coupling mechanisms, in contrast with dipole-dipole interactions which take place between different molecules.

In its most simple case, two bounded atoms A and X with a large gap in resonance frequencies are considered (figure 1.9A). By the Pauli Exclusion Principle, the interaction between the electrons inside the covalent bond is forced to be antiparallel. In the case where the AX spin system is in the high-energy ($\beta\beta$) or low energy states ($\alpha\alpha$), one of the nuclei (say A) will be in a parallel spin state to that of its electron, which is less favorable than an antiparallel state. The energy level will therefore increase by an amount of $J_{AX}/4$ where $J_{AX}$ corresponds to the single-bond scalar coupling constant between A and X. For the intermediate states ($a\beta$ and $\beta a$), the electrons can be in antiparallel state with the nuclei spins which is energetically favorable, decreasing the energy levels of these states by $J_{AX}/4$. In the case of heteronuclear coupling such as $A = ^1H$ and $X = ^{13}C$, the energy required to excite the $^1H$ spin corresponds to the transitions $a\beta \rightarrow \beta\beta$ or $\alpha\alpha \rightarrow \beta\alpha$, 

25
which are $\gamma_H (h/2\pi) B_0 - \hbar J_{HC}/2$ and $\gamma_H (h/2\pi) B_0 + \hbar J_{HC}/2$, respectively. The associated NMR spectrum consists in a pair of resonances centered on the Larmor frequency with a $J_{HC}$ frequency gap (figure 1.9B). In the case of a strongly coupled system AB in which the difference in resonance frequencies is small ($|v_A - v_B| \approx J_{AB}$), there are still 4 energy level transitions but the intensities of the corresponding NMR resonances are distorted by second order effects. Notably, J-coupling constants are independent of the magnetic field intensity.

Figure 1.9. Energy levels splitting due to J-coupling and NMR spectrum of a weakly-coupled AX spin system. A. Two atoms interacting through a covalent bond are linked through an electron bond. The energy levels of the spins A and X are influenced by the anti-parallel pairing of the electrons due to the Pauli Exclusion Principle. This scalar coupling $J_{AX}$ leads to 4 different possible energy states. B. This leads to 4 possible single spin transitions ($E_2-E_1$ and $E_4-E_3$ for A and $E_2-E_1$ and $E_4-E_3$ for X). If it is a weakly-coupled spin system ($|v_A - v_X| \gg J_{AX}$), the NMR spectrum consists in 2 doublets of resonances centered on the Larmor frequencies of A and X each with a $J_{HC}$ frequency gap.
1.2. MRS Sequences

1.2.1. Radiofrequency Pulses

1.2.1.1. Amplitude Modulated Pulses

As explained in section 1.1.5, the longitudinal magnetization is projected towards the transverse plane by applying a secondary transverse magnetic field \( B_1 \) oscillating at or close to the Larmor frequency of the spin population of interest. The \( B_1 \) “excitation” field is created by a surface or volume transmit coil. The resolution of the Bloch equations (eq 1.13-1.15) during the application of the RF “pulse” \( B_1(t) \) allows for the calculation of the magnetization \( M(t) \). To do so, it is convenient to describe the NMR experiment in a frame rotating about \( B_0 \) at the angular carrier frequency \( \omega \) as well as an effective magnetic field \( B_{\text{eff}} \):

\[
B_{\text{eff}} = \sqrt{B_1^2 + \left( \frac{\Delta \omega}{\gamma} \right)^2}; \quad \Delta \omega = \omega_0 - \omega
\]

(1.27)

with \( \omega_0 \) being the angular Larmor frequency.

On-resonance \((B_1 \gg \frac{\Delta \omega}{\gamma})\), the magnetization rotates about \( B_1 \) as shown in figure 1.3 and the FA \( \alpha \) is given by:

\[
\alpha = \gamma B_1 T
\]

(1.28)

with \( T \) being the RF pulse duration. However, off-resonance effects are present when the frequency offset \( \Delta \omega \) is comparable to \( \gamma B_1 \). In MRS, this is usually the case since the spectral bandwidth is several kHz wide.

By incorporating \( B_{\text{eff}} \) into the Bloch equations, frequency-dependent profiles of the three magnetization components \( M_x, M_y \) and \( M_z \) can be calculated. A simpler method to estimate the frequency profile of the RF pulse is to apply a Fourier transformation (FT) in the time domain. Yet, the FT being a linear operation, it must be taken into consideration that the said frequency profile is an approximation only valid for small nutation angles \((\alpha<30^\circ)\) where the sinus function is almost linear \((\sin(\alpha) \approx \alpha)\).

Another important factor for the characterization of RF pulses is the Time-Bandwidth or \( R \) factor. The \( R \) factor corresponds to the product of the pulse length \( T \) and the pulse bandwidth \( \Delta \omega \). The latest is measured as the full width at half maximum (FWHM) of the frequency profile of the resulting magnetization after a 90° or 180° FA for excitation and inversion/refocusing pulses, respectively. For a given RF pulse modulation, the \( R \) factor is constant and is useful for determining the pulse duration needed to achieve a target bandwidth:

\[
T = \frac{R}{\Delta \omega}
\]

(1.29)
In the case of amplitude modulated (AM) pulses, the amplitude integral $\kappa$ is also a pulse-specific constant that represents the ratio between the area of the normalized pulse ($B_{1_{\text{max}}} = 1$) and a normalized square pulse of the same duration:

$$\kappa = \sqrt{\left(\sum_{i=1}^{N} V_i,\text{Real}\right)^2 + \left(\sum_{i=1}^{N} V_i,\text{Imag}\right)^2 / N}$$ (1.30)

where $V$ is the (discrete) vector representation of the pulse envelope defined by $N$ complex points. The amplitude integral $\kappa$ is useful for calculating the required $B_1$ for a particular pulse and a desired flip angle, provided that a reference value is available from a reference pulse (usually a 1 ms long square pulse). The mathematical expression is directly derived from eq. (1.28):

$$\alpha = \gamma \cdot B_1 \cdot T \cdot \kappa$$ (1.31)

The choice of a RF pulse will largely depend on the experimental demands and constrains, such as spectral bandwidth and desired sharpness (transition bandwidth), available RF power, gradient capabilities and Specific Absorption Rate (SAR) limitations. The R and $\kappa$ factors for the most common AM pulses used during this thesis are listed in table 1.2.

The transition bandwidth $T_{bw}$ is the frequency gap for which the condition $0.05 \leq M_{xy}/M_0 \leq 0.95$ holds for a 90° excitation pulse and it should be considered when sharp excitation profiles are required. Sinc pulses with a high number of lobes have good $T_{bw}$ while Gaussian pulses have rather degraded ones but demand a smaller $B_{1_{\text{max}}}$. Hermitian shaped pulses show a good compromise between $T_{bw}$, R value and peak $B_1$.

<table>
<thead>
<tr>
<th>Pulse Shape</th>
<th>Time-Bandwidth R</th>
<th>Amplitude Integral $\kappa$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gaussian</td>
<td>2.17</td>
<td>0.5530</td>
</tr>
<tr>
<td>Sinc 3-lobes (FLASH,Siemens)</td>
<td>3.35</td>
<td>0.3504</td>
</tr>
<tr>
<td>Sinc 3-lobes (CSI,Siemens)</td>
<td>3.84</td>
<td>0.2800</td>
</tr>
<tr>
<td>Sinc 7-lobes</td>
<td>8.92</td>
<td>0.1137</td>
</tr>
<tr>
<td>SLR (VERSE)$^a$</td>
<td>15.00</td>
<td>0.1049</td>
</tr>
<tr>
<td>Square</td>
<td>1.37</td>
<td>1.0000</td>
</tr>
<tr>
<td>Hermitian</td>
<td>5.43</td>
<td>0.1794</td>
</tr>
</tbody>
</table>

Table 1.2. Time-Bandwidth product and amplitude integrals of common AM pulses.

$^a$ (Amadon, et al., 2010)

1.2.1.2. Adiabatic Pulses

In most MRS studies, surface coils are used due to their high sensitivity and elevated peak power. Nevertheless, they generate a quite inhomogeneous $B_1$ excitation field, leading to artifacts when a deep structure is studied or when the volume-of-interest (VOI) extends in depth with respect to
the plane of the transmitter’s inductive loop(s). Although it is possible to solve the $B_1$ inhomogeneity problem by using a transmit-only volume coil in conjunction with a receive-only surface coil, the maximum $B_1$ power needed to apply high-bandwidth RF pulses can be compromised (aside from the increasingly complicated set-up and possible coupling effects). An elegant and reliable solution consists in using RF pulses for which the FA will be less sensitive to the amplitude of the applied $B_1$. Adiabatic pulses are frequency- and amplitude-modulated pulses that generate a uniform nutation angle even in an inhomogeneous $B_1$ field provided that the RF amplitude is above a certain threshold so that it stays in the adiabatic regime.

A qualitative illustration of the application of an adiabatic pulse is as follows. The frequency modulation on the applied $B_1$ field has an effect equivalent to modulating the off-resonance component of $B_{\text{eff}}$ as shown in equation (1.27), where $\Delta\omega$ is substituted by the term $\Delta\omega(t) = \omega(t) - \omega_0$:

$$B_{\text{eff}}(t) = \sqrt{B_1^2(t) + \left(\frac{\Delta\omega(t)}{\gamma}\right)^2} \quad (1.32)$$

Through the simultaneous modulation of the RF field intensity and carrier frequency, the effective $B_{\text{eff}}(t)$ field changes in orientation over time. If the orientation (with respect to the reference frame rotating at the Larmor frequency) given by:

$$\alpha(t) = \arctan\left(\frac{\Delta\omega(t)}{\gamma B_1(t)}\right) \quad (1.33)$$

changes slowly enough such that it complies with the adiabatic condition:

$$\left|\frac{d\alpha(t)}{\gamma dt}\right| \ll |B_{\text{eff}}(t)| \quad (1.34)$$

then the magnetization that was parallel ($M_z$) and perpendicular ($M_{xy}$) to $B_{\text{eff}}$ at the beginning of the RF pulse will keep their orientation to $B_{\text{eff}}(t)$ for the duration of the pulse. If this condition is not fulfilled, there will be a loss of coherence among the longitudinal magnetization vectors at different frequencies according to $\Delta\omega(t)$ and $B_1$.

Adiabatic pulses can be easily calibrated by keeping a constant pulse length and gradually increasing the transmitter power up to a threshold at which the recorded signal intensity remains constant. The most commonly used adiabatic pulses are the hyperbolic secant (HS) adiabatic half-passage (AHP) $90^\circ$ excitation pulse and the adiabatic full-passage (AFP) $180^\circ$ inversion pulse (Silver, et al., 1984; Baum, et al., 1985), whose amplitude and frequency modulation functions $F_1(t)$ and $F_2(t)$ are:

$$B_1(t) = B_{1\text{max}} \ast F_1(t) = B_{1\text{max}} \ast \text{sech}\left[\beta \tau(t)\right] \quad (1.35)$$

$$\Delta\omega(t) = \omega_c + \Delta\omega_{\text{max}} \ast F_2(t) = \omega_c + \Delta\omega_{\text{max}} \ast \frac{\text{tanh}\left[\beta \varepsilon(t)\right]}{\text{tanh}[\beta]} \quad (1.36)$$

where $\tau = (1-2t/T_p)$, $T_p$ is the pulse length, $\beta$ is a truncation factor close to 5.3 so that $\text{sech}(\beta) = .01$, $\omega_c$ is the central frequency and $t$ varies from 0 to $T_p/2$ for an AHF pulse and from 0 to $T_p$ for
an AFP pulse. For AFP pulses, the R product is determined by the maximum frequency modulation $\Delta \omega_{\text{max}}$, as the inversion bandwidth equals $2 \times \Delta \omega_{\text{max}}$. For AHP pulses, the effective excitation bandwidth is $B_1$ dependent as the angle $\alpha(t)$ for off-resonance excitation varies with $\Delta \omega$ and $B_1$ as shown by eq. (1.40). It is worth noticing that AHP and AFP pulses are not plane rotations as the existing transverse magnetization $M_{xy}$ at the start of the pulse rotates around the effective field $B_{\text{eff}}$ acquiring a linear phase as a function of the RF intensity and the frequency offset, effectively dephasing during the pulse. However, if an AFP pulse has been applied, running a second AFP just afterwards inverts the effective field $B_{\text{eff}}$ canceling the acquired linear phase by adding the same opposite phase to the magnetization vector. This method of adiabatic refocusing is employed by the LASER localization sequence as it will be discussed later on this chapter.

When the power requirements are too elevated for the spectrometer or the SAR levels are too high, other modulation functions can be used to reduce the maximum required RF amplitude $B_{1\text{max}}$ while degrading moderately the transition bandwidth of the pulse profile. Modulating function pairs $F_1$ and $F_2$ may be used as long as they fulfill the Offset-Independent Adiabaticity (OIA) condition:

$$\left(\frac{\gamma B_{1\text{max}}F_1(t_\Omega)}{\Delta \omega_{\text{max}}F_2(t_\Omega)}\right)^2 \gg 1$$ (1.37)

at time $t_\Omega$ for an on-resonance frequency isochromat $\Omega$. As it was shown by Tannus and Garwood (Tannús, et al., 1997), the peak power of OIA pulses is determined by the AM function while the average power is determined by the inversion bandwidth (for a fixed pulse duration $T_p$). Among the modulating functions that fulfill the OIA conditions, the HSn pulse family was used in this thesis to limit the required $B_{1\text{max}}$. Their AM and FM functions are given by the formulas:

$$F_1(t) = \text{sech}\left(\beta \tau^n(t)\right)$$ (1.38)

$$F_2(t) = \int \text{sech}^2\left(\beta \tau^n(t)\right)$$ (1.39)

The use of HS8 ($n = 8$) pulses with respect to the standard HS AFP shows a dramatic improvement in the required peak power, as there is a 51% reduction in $B_{1\text{max}}$ while the average applied power changes by less than 1% ($T_p = 2\text{ms}$, Bandwidth ($\Omega$) = 50 kHz). Nevertheless, $T_{bw}$ is degraded by 5%.

1.2.1.3. Energy deposition

Along with the magnetic field $B_1$, an associated electrical field $E$ is created which induces electrical currents in the sample. These induced electrical currents are dissipated into heat which may become harmful if the ensuing increase in temperature reaches a hazardous level, particularly in tissues with low perfusion. The local temperature elevation at a position $r$ depends on the conductivity $\sigma(r)$ and density $\rho(r)$ of the tissue as well as the duration $T$ of the local
electric field $E(r,t)$. The level of energy deposition is determined through the specific absorption rate (SAR) and is given in watts per kilogram.

The IEC has determined safety guidelines (International Electrotechnical Commission, March 2010) based on global and local SAR limits over periods of integration of 6 minutes and of 10 seconds. Global SAR corresponds to the energy deposition on the entire head and local SAR on the energy deposition over the mass of any closed 10-grams volume. The local SAR is limited to a maximum of 20 W/kg over a period of 10 seconds and 10 W/kg over a 6-min window. The global SAR is limited to a maximum of 9.6 W/kg over a period of 10 seconds and 3.2 W/kg over a 6-min window. The local SAR can be calculated using the following expression:

$$SAR_{local}(r) = \frac{\sigma(r)}{2\rho(r)} \frac{1}{T} \int_0^T \|E(r,t)\|^2 dt$$  \hspace{1cm} (1.40)

The E-field depends on the transmitter coil and quality factor as well as the characteristics and positioning of the sample within the coil. Global energy deposition can be calculated based on the quality factor of the loaded and unloaded coil, but this information does not provide information about the eventuality of local “hot-spots” (Haacke, et al., 1999 pp. 858-860). From the SAR mathematical expression, it can be observed that:

1) the power deposition scales quadratically with the intensity of the electric field,
2) for a given flip angle, SAR values can be reduced by increasing the duration of the pulse (at the cost of decreasing its spectral bandwidth) and
3) longer TRs may help overcome SAR limitations at the cost of longer acquisition times.

1.2.2. SVS localization

Spatial localization of the NMR signal is an essential aspect of in vivo MRS. In order to acquire physiologically relevant and good quality spectra, a proper localization sequence must be used to select the NMR signal from within the VOI and avoid spurious signal from outside. Indeed, localization errors introduces unwanted signal that biases the validity of a spectrum for physiological or clinical research purposes. Furthermore, experimental adjustments such as the $B_0$ and $B_1$ field homogenization or “shimming” and RF pulses calibration are spatially-dependent. Therefore, a good spatial localization will guarantee that the NMR signal is originating from a region where experimental conditions have been optimized hence improving the quality of the spectra. It will also confine the NMR experiment to the VOI and minimize partial volume effects. Using high-bandwidth RF pulses for spatial selection is an efficient way to limit the mismatch between the volumes localized for spins resonating at different frequencies. This artifact is detailed in section 1.2.4 and is known as the chemical shift displacement artifact (CSDA).

Single volume spectroscopy (SVS) sequences that were most relevant to this work are briefly described below.
1.2.2.1. FID with Outer Volume Suppression localization

The Pulse-Acquire or FID sequence is the simplest signal acquisition method as it consists of a non-selective excitation RF pulse prior to the acquisition of the FID. A VOI can be selected using a set of Outer-Volume Suppression (OVS) bands applied prior to the excitation pulse (figure 1.10). The OVS scheme saturates the NMR signal from a slab next to the VOI by projecting $M_z$ into the transverse plane and subsequently dephasing $M_{xy}$ using strong $B_0$ gradients (Shungu, et al., 1993). The dephasing of the signal may not be completely satisfactory (notably in clinical systems) as it requires important spoiling (Hurley, et al., 2012). For this reason RF-spoiling is also used. RF-spoiling consists in using adiabatic or frequency-modulated (FM) RF pulses such as truncated HS inversion pulses to achieve an incoherent 90° excitation which greatly reduces the net magnetization vector.

The advantages of a FID-OVS sequence with respect to other SVS sequences, especially for X-nuclei MRS, are threefold:

1) As the signal is acquired shortly after the non-selective excitation, the differential $T_2^*$-weighting between metabolites is reduced. This is particularly interesting for studying NMR species with short $T_2^*$ relaxation times, such as phosphorous metabolites.

2) It presents a low sensitivity to $B_1$ field inhomogeneity as it does not require any inversion or refocusing pulses for localization. In addition, the choice of the flip angle is not critical, which is useful during experiments where the calibration of the RF power is uncertain such as in X-nuclei MRS experiments using a surface coil.

3) It provides a high degree of freedom in the determination of the VOI geometry, giving the possibility to acquire spectra from the entire brain while limiting the artifacts due to chemical shift.

Furthermore, when large VOI are studied and the $B_1$ field is moderately inhomogeneous over the sample, recurring to adiabatic pulses such as $B_1$-independent rotation phase cycled (BIRP) and variable angle adiabatic plane rotation (BIR-4) pulses help achieving an arbitrary homogeneous flip angle over the entire VOI (Bottomley, et al., 1994; Staewen, et al., 1990).
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Figure 1.10. FID acquisition pulse sequence with OVS localization. The FID sequence achieves spatial localization by eliminating the undesired volume contributions through an outer-volume suppression module consisting of the application of n separated suppression bands prior to the non-localized excitation of the VOI.

1.2.2.2. Stimulated Echo Acquisition Mode (STEAM)

STEAM is a robust MRS localization technique consisting of three slice-selective 90° RF pulses (figure 1.11) (Frahm, et al., 1987). Due to the nature of the pulses, it generates a total of 5 spin echos and 3 FIDs, where the time interval between the first two pulses is TE/2 and the delay between the last two is the mixing time or TM (Hahn, 1950). Usually, only the stimulated echo at t = TE+TM is of interest as it corresponds to the signal generated exclusively from the VOI. Other NMR signals are destroyed by applying crusher gradients. Due to the application of a dephasing gradient during TM, only half of the total available magnetization can be refocused. In spite of this, STEAM is a very popular MRS sequence due to its versatility and robustness since it remains effective even when the calibration of the 90° RF pulses is not optimal. Also the TE can be considerably reduced (down to few ms) by employing asymmetric or inherently refocused 90° RF pulses (Geen, et al., 1991; Tkàc, et al., 1999).
Figure 1.10. STEAM localization pulse sequence.
The STEAM pulse sequence consists of 3 $90^\circ$ excitation pulses applied on the rotational axis. After the second $90^\circ$ pulse, the magnetization of the xy-plane is rotated into the xz-axis. Application of strong dephasing gradients ($G_2$) during the mixing time removes any transverse component, leaving only a net coherent magnetization on the longitudinal axis. The second $G_1$ spoiler allows dephasing the unwanted FID signal from the third excitation pulse, and compensates for the first $G_1$ spoiler allowing the formation of the stimulated echo. The second $G_1$ spoilers also contain the refocusing gradients. To prevent the apparition of unwanted FIDs or spin echoes, $G_1$ must be different from $\pm G_2$ or $\pm G_2/2$ (de Graaf, 2007).

1.2.2.3. Point resolved spectroscopy (PRESS)

The PRESS pulse sequence is a commonly used localization method consisting of a slice-selective excitation and two slice-selective refocusing pulses (Bottomley, 1987). The slice-selective pulses are applied in orthogonal planes and as such the magnetization outside of the VOI is either not excited or not refocused. To completely eliminate residual signal from outside the VOI, spoiler gradients pairs surrounding the $180^\circ$ refocusing pulses are applied. The refocusing pulses correspond to two Hahn spin-echoes which generate a first spin-echo at time $2t_1$, where $t_1$ is the time between the excitation pulse and the first refocusing pulse, and a second spin-echo at time $TE = 2t_1 + 2t_2$, where $t_2$ is the time between the first spin-echo and the second refocusing pulse. PRESS is a robust pulse sequence which, unlike STEAM, measures the full available magnetization in the VOI. However, the use of PRESS at high fields is challenging due to the increased $B_1$ inhomogeneity, which degrades the efficacy of the two refocusing pulses. Furthermore, the elevated pulse bandwidths needed to reduce chemical shift displacement artefacts (which will be discussed in section 1.2.4) are difficult to achieve using amplitude-modulated RF pulses due to their elevated peak power requirements.
1.2.2.4. Localization by adiabatic selective refocusing (LASER)

The LASER sequence is a localization sequence that allows for a sharp localization profile and is highly insensitive to $B_1$-inhomogeneities. It consists of a non-selective 90° RF pulse, typically an AHF, followed by three pairs of slice-selective refocusing AFP pulses (figure 1.12), each of which delineates an orthogonal slice. Each AFP pulse is surrounded by identical crusher gradients to destroy the signal outside of the slice being selected (Garwood, et al., 2011). By employing adiabatic pulses with an elevated R factor, sharp selection profiles can be achieved while reducing chemical shift artifacts.

A major advantage of LASER compared to other SVS pulse sequences is that its train of 180° refocusing pulses acts as a CPMG pulse train, reducing anti-phase coherence from J-coupling and apparent $T_2$-shortening effects from diffusion and chemical exchange (Carr, et al., 1954; Allerhand, et al., 1966). Under the CPMG regime, the dephasing effects of spin-spin interactions are therefore reduced, increasing the apparent $T_2$ relaxation time. The benefits of the CPMG regime are not the same for all spin systems, as it has been experimentally observed at 9.4 T (Deelchand, et al., 2015). Spin pairs AB satisfying the condition:

$$\sqrt{\Delta \omega_{AB}^2 + J_{AB}^2 \cdot T_{E_{CPMG}}} \ll 1$$ (1.41)

have a greater enhancement of their signal detection capabilities, as is the case for strongly coupled systems (AB) such as glutamate and glutamine up to moderate echo times (~45 ms at 4T, ~10 ms at 17.2 T) (Allerhand, et al., 1966; Hennig, et al., 1997).

Nevertheless, LASER presents two main drawbacks. Firstly, the elevated energy deposition due to the use of three pairs of adiabatic refocusing pulses and secondly, the difficulty to reach relatively short TEs, although it is less sensitive to apparent $T_2$ relaxation effects due to the CPMG pulse train as explained before.
Figure 1.11. LASER localization pulse sequence

The LASER pulse sequence consists in a non-localized 90° pulse followed by three pairs of slice-selective adiabatic 180° pulses. The crusher gradients surrounding each refocusing pulse dephase the signal outside of the selected slice. The total TE is the sum of the 3 independent echo-times. To improve its insensitivity to B$_1$ field inhomogeneities, an adiabatic 90° pulse can be used such as an AHP or a BIR-4 pulse.

1.2.3. Chemical Shift Imaging

One drawback of SVS is that the metabolic information originates from a parallelepipeded VOI usually encompassing different tissue types. The technique of MRS Imaging (MRSI) or Chemical Shift Imaging (CSI) allows for the simultaneous acquisition of different volumes (voxels), enabling the study of distinct tissue types under the same physiological and experimental conditions. Additionally, tissue-specific profiles can be extracted if a high resolution acquisition matrix is employed or when large pathological features are present (such as tumors).

Any CSI pulse sequence consists in combining a SVS localization sequence with a 2D or 3D spatial encoding scheme. In order to preserve the spectroscopic information, only phase encoding (PE) steps are used (figure 1.13).

The phase acquired by the NMR signal for any PE step along an arbitrary axis x is given by the expression:

$$\phi(x) = \gamma * x * G_x * t$$  \hspace{1cm} (1.42)

where $G_x$ is the amplitude of the gradient and t its duration. If there is no phase encoding (i.e $G_x = 0$), a regular NMR spectrum is acquired. Else, the acquired NMR signal is the sum of phase shifted resonances accordingly to their spatial position x. The spectrum of the entire volume $F_{VOI}$ corresponds to the FT of the individual spectra $F_{Ele}$. The spatial distribution across the PE gradient is given by:

$$F_{VOI}(G_x, \omega) = \int_{-\infty}^{+\infty} F_{Ele}(x, \omega) e^{i \gamma G_x x} dx = \int_{-\infty}^{+\infty} F_{Ele}(x, \omega) e^{i k_x x} dx = F_{VOI}(k_x, \omega)$$  \hspace{1cm} (1.43)
Applying the inverse FT gives the single voxel spectrum at position $x$:  

$$ F_{Ele}(x, \omega) = \int_{-\infty}^{+\infty} F_{VOI}(k_x, \omega) e^{-ik_x x} dx $$

In the case of a 3D acquisition, a single voxel spectrum will be obtained by applying the spectral FT to the sum of the acquired data (which corresponds to $F_{VOI}$) and then applying a spatial 3D FT to the matrix of PE spectra.

Figure 1.12. 2D CSI-SE pulse sequence
This typical 2D MRSI sequence consists in a slice-selective spin-echo sequence for which the phase of each echo is encoded using phase-encoding gradients in the frequency and phase directions.

1.2.3.1. The Point-Spread Function and Apodization Methods

The spatial resolution of a voxel in a CSI is primarily determined by the number of PE steps $N_P$ over the field of view (FOV), modulated by a convolution function called the point-spread function (PSF). The PSF expresses the fact that a limited phase-encoded sampling generates a signal “bleeding” effect after data reconstruction with the Fourier transform. For each voxel, some signal is distributed over adjacent voxels leading to a degraded spatial resolution and partial volume effects (Posse, et al., 2012). For a regular sampling scheme, the PSF corresponds to a cardinal sinus and it is determined by $N_P$ and the FOV ($= 1/ \Delta k_x$):

$$ PSF = \Delta k_x \sin(\pi N_P \Delta k_x x) / \pi N_P \Delta k_x x $$

Due to the limited number of PE steps in MRSI, the effects are significant and the PSF may be improved by using apodization methods either as a weighted sampling scheme or as a post-processing step.
Apodization of the PSF as a post-processing method limits the “bleeding” effects into distant voxels but degrades the voxel resolution by widening the PSF and removes high-frequency k-space voxels contributions (figure 1.14A). An apodization function showing a good compromise between ripple reduction and widening of the effective voxel size is the Hann (or “Hanning”) filter (figure 1.14.B):

\[
W(k) = \alpha + \beta \cos\left(\frac{\pi k}{k_{\text{max}}}\right), \quad -k_{\text{max}} \leq k \leq k_{\text{max}}
\]

where \(k_{\text{max}}\) is the largest sampled position in the k-space and \(\alpha = \beta = 0.5\). For \(\alpha = 0.53836\) and \(\beta = 0.46164\) (also known as Hamming filter), the largest pair of sidelobes are minimized, further reducing ripples (Ernst, et al., 1987; Harris, 1978).

Another filtering method is the apodization of the PSF by acquiring a weighted k-space, where a higher number of acquisitions are repeated for PE steps closer to the center of the k-space (figure 1.15). The effect of such weighted acquisition is an increase in SNR at the expense of the spatial resolution. The density of repetitions per PE step can be estimated by distributing the allotted number of repetitions to fit the shape of the apodizing function.

![A. (Left). 2D unfiltered PSF for a 25x25 matrix. Ripples appear several voxels away from the center, causing bleeding effects. (Right). 1D unfiltered PSF (black). A regular sampling scheme of the k-space where no voxel weighting is present (red). B. (Left) Filtering the k-space using a Hanning function reduces the bleeding effects at the cost of a degraded spatial resolution. (Right) The Hanning-weighted sampling scheme (red) has smoothed and enlarged the resulting PSF (black).](image)

Figure 1.13. Point-Spread Function and apodization of the CSI k-space.

A. (Left). 2D unfiltered PSF for a 25x25 matrix. Ripples appear several voxels away from the center, causing bleeding effects. (Right). 1D unfiltered PSF (black). A regular sampling scheme of the k-space where no voxel weighting is present (red). B. (Left) Filtering the k-space using a Hanning function reduces the bleeding effects at the cost of a degraded spatial resolution. (Right) The Hanning-weighted sampling scheme (red) has smoothed and enlarged the resulting PSF (black).
1.2.4. Chemical Shift Artifact and outer-volume contamination

In a localized acquisition sequence, slice-selection is achieved by applying a gradient $G_x$ during the RF pulse:

$$G_x = \frac{\Delta \omega}{\gamma x_{sl}}$$  \hspace{1cm} (1.47)

where $\Delta \omega$ is the RF pulse bandwidth and $x_{sl}$ the slice thickness. As given by equation (1.40), the difference in chemical shift between different spin populations will cause them to resonate at different frequencies. For a spin population resonating at $\omega_m$ and a RF carrier frequency $\omega$, there is a spatial displacement of the excited volume known as the chemical shift displacement artifact (CSDA). The chemical shift displacement $\Delta x$ is given by the expression:

$$\Delta x = \frac{\omega - \omega_m}{\gamma G_x} = \frac{\omega - \omega_m}{\Delta \omega} x_{sl}$$  \hspace{1cm} (1.48)

To appreciate the practical consequences of the CSDA, one can consider the following $^1$H MRS experiment at 17.2 T: a carrier frequency at 2.7 ppm, an RF pulse with a spectral bandwidth of 20 kHz and a 10 mm slice thickness. In this situation, the water signal (~4.7 ppm) originates from a slab displaced by ~0.73 mm, while the signal from the macromolecule and lipid resonances at 0.9 ppm comes from a slab at +0.66 mm from the intended slab (figure 1.16A). If extended to a 3D volume selection, then only 64% of the signal of the recorded spectra (from 4.7 to 0.9 ppm) will originate from the same location.
A. The chemical shift displacement artifact (CSDA) is proportional to the frequency difference between the carrier and spin resonance frequencies and inversely proportional to the spectral bandwidth of the pulse used for slice-selection. By choosing a carrier frequency at the center of the spectrum (close to the Cr(CH₃) resonance at 3.02 ppm) one can minimize the spatial mismatch between resonances. B. Spectra acquired with a carrier frequency centered at 3 ppm using a "H 2D CSI-SE sequence (slice thickness = 6 mm, matrix size 25 x 25, circular k-space sampling) with hermit excitation pulses with spectral widths of 9 kHz (black spectrum) and 5.4 kHz (blue spectrum) at 17.2 T. The spectrum shown in blue displays a more pronounced CSDA as the chemical shift displacement artefact increases.

Large CSDA may also degrade the quality of the spectra through outer-volume contamination, notably by unintended excitation of the extra-cranial lipids (figure 1.16B). CSDA is reduced through the use of large bandwidths relative to the spectral bandwidth. If it is not possible, a smaller VOI can be defined avoiding outer-volume contamination from undesired structures at the cost of reduced SNR.

1.3. IN VIVO NMR SPECTROSCOPY AT HIGH MAGNETIC FIELDS

1.3.1. Benefits

As shown in eq. (1.11), the magnetization $M_0$ is proportional to $\gamma^2B_0$. Due to the principle of reciprocity, the induced emf in the reception coil vary proportionally to $\omega_0*M_0$. Therefore, the NMR signal $S$ is proportional to:

$$S(\gamma, B_0) = \gamma^3 B_0^2 = \gamma (\omega_0)^2$$  \hspace{1cm} (1.49)

when the sources of noise are ignored. Noise is nevertheless present due to the resistances of the coil, hardware and sample. It has been shown that the effective resistance $R_{eff}$ is frequency-dependent. At low frequencies, the coil and electronics resistances dominate over the sample resistance whereas at high frequencies (above 0.5 T for $^1$H) the sample resistance dominates.
(Hoult, et al., 1976). The thermal noise in the system $\sigma_{\text{Thermal}}$ depends on the sample resistance $R_{\text{eff}}$ as given by:

$$\sigma_{\text{Thermal}} \propto \sqrt{4kT \cdot R_{\text{eff}} \cdot \omega_{\text{Rec}}} \quad (1.50)$$

where $\omega_{\text{Rec}}$ is the detecting system receiver bandwidth (Haacke, et al., 1999). When only white noise is present, the SNR is:

$$\text{SNR}(\gamma, B_0) = \frac{S(\gamma, B_0)}{\sigma_{\text{Thermal}}} \propto \frac{\gamma (\gamma B_0)^2}{\gamma B_0 \sqrt{4kT \cdot \omega_{\text{Rec}}}} = \frac{\gamma^2 B_0}{\sqrt{4kT \cdot \omega_{\text{Rec}}}} \quad (1.51)$$

A linear gain in SNR can be obtained by increasing the magnetic field strength (Radpath, 1998), when the changes in relaxation times, receiver bandwidth or coil sensitivity are ignored. Furthermore, one can notice that there is a greater sensitivity with larger gyromagnetic ratios.

Another important aspect for MRS at higher magnetic fields is the greater spectral resolution. While the chemical shifts are $B_0$-independent, the frequency difference between spins increases linearly with $B_0$. Provided that $B_0$ field inhomogeneities are properly accounted for, the overlap between peaks is $a priori$ reduced as they are spread over a larger frequency range, as it can be appreciated in $^1\text{H}$ MRS brain spectra (figure 1.17). This is particularly notorious for glutamate and glutamine, which are hardly distinguishable below 7 T (Gruetter, et al., 1998). Also, $J$-coupling constants being $B_0$-independent, some hyperfine structures $in vivo$ becomes less manifest, simplifying the spectrum and facilitating its spectral decomposition.
Figure 1.16. Improvement of MRS spectra with $B_0$. Increasing the magnetic field strength improves SNR, spectral resolution and simplifies J-coupling patterns. Spectra acquired in the rat brain in vivo at increasing magnetic fields are shown. A LASER sequence was used for the acquisition of the spectra at 11.7 T (TE = 25 ms, TR = 3s) and at 17.2 T (TE = 16.5 ms, TR = 5s). The spectrum at 7 T was measured with STEAM (TE = 3, TR = 1.5s). No filtering was applied.
1.3.2. Challenges

Despite the benefits of ultra-high magnetic fields (UHF), moving towards higher magnetic fields is not straightforward as several difficulties arise in vivo. Notably, the gain in SNR described by equation (1.51) does not consider signal loses due to faster $T_2$ decay. Also, the eventually longer $T_1$ relaxation times lead to reduced steady-state magnetization unless the total scan time is prolonged. Fortunately, for some nuclei such as $^{31}$P, the $T_1$ relaxation times of metabolites decreases with $B_0$, improving the normalized SNR at UHF (de Graaf, et al., 2006; Lu, et al., 2014).

Another important aspect is the increase of magnetic susceptibility effects with the external magnetic field, accentuated near tissue interfaces. The increased $B_0$ inhomogeneities have an important impact in MRS because the spatially-varying magnetic field creates local changes of the Larmor frequency for chemically-identical spin populations, no longer adding coherently at a single frequency but spreading instead over different frequencies, losing phase coherence and forming a broader non-Lorentzian resonance shape. Water suppression becomes less effective due to a broadened water resonance. This can be countered by increasing the spectral bandwidth of the frequency-selective water saturation pulses but the detection of the resonances close to 4.7 ppm can be compromised by the saturation pulses and baseline distortions. The use of more intense shimming coils or of additional higher order shimming coils alleviates static field inhomogeneities in small to moderate brain regions at UHF and the use of dynamic shimming methods further improves the quality of multi-slice CSI acquisitions at 7 T (Boer, et al., 2012).

As the magnetic field increases, so does the energy required to excite the spins from the $\beta$ to the $\alpha$ energy state [eq. (1.5)]. The required $B_1$ scales linearly with $B_0$ and the power required for its production increases approximately linearly as well (Vaughan, et al., 2001). In order to keep the CSDA within acceptable levels, higher spectral bandwidths are also required, further exacerbating the need for stronger excitation field intensities.

This creates two main challenges: First, the peak power demands at UHF can quickly reach the hardware limits when elevated spectral bandwidths are demanded. Secondly, global power deposition $P$ increases dramatically as it shows a quadratic increase with both the $B_1$ intensity and the RF frequency $\omega_0$:

$$P = \frac{4}{15} \pi \sigma \omega_0^2 B_1^2 R^5$$  

(1.52)

as calculated for the ideal case of a homogeneous spherical phantom of radius $R$ and conductivity $\sigma$ (Hoult, et al., 2000). Yet, human research protocols at 7 T are typically constrained by SAR restrictions rather than hardware limitations, such as the RF power amplifier.

Finally, the $^1$H resonance frequency at 7 T is such that the wavelength shortens to ~12 cm, reaching a similar length to the human head dimensions (Yang, et al., 2002; Vaughan, et al., 2001). The generated interference patterns further complicate the $B_1$ and E fields, reducing $B_1$ field inhomogeneity and exacerbating the apparition of local SAR “hot spots” [equation (1.40)].
To tackle the aforementioned $B_1$-related complications, numerous methods exist to reduce $B_1$ power demands, such as using variable-rate selective excitation (VERSE) amplitude or adiabatic pulses (Conolly, et al., 1988; Conolly, et al., 1991) or numerically optimized pulses such as phase-modulated SLR pulses (Shinnar, 1994), GOIA HS(n) or wideband, uniform rate and smooth truncation (WURST) frequency-modulated pulses (Tannús, et al., 1997; Andronesi, et al., 2010).

$B_1$ field inhomogeneity can be compensated in a subject-dependent manner with parallel transmission (pTx) methods employing static or dynamic $B_1$ shimming (Emir, et al., 2012; Boer, et al., 2012) while reducing power deposition (Emir, et al., 2012; Boer, et al., 2012; Avdievich, et al., 2009). The increased degree of freedom of pTx transmitters allow to employ circularly polarized (CP) eigenmode “rings” to eliminate the signal of extracranial lipids with reduced energy deposition compared to standard OVS bands (Hetherington, et al., 2010). Other methods such as double-row transmit array coils (Shajan, et al., 2014) or passive RF shimming using dielectric pads (O’Brien, et al., 2014) can also contribute to $B_1$ homogeneity and global SAR reduction.
2. PRACTICAL ASPECTS

2.1. NMR SPECTROMETERS AND ADJUSTMENT PROCEDURES

The work presented in this thesis was done on a 17.2 T preclinical Biospec Bruker scanner and on a 7 T clinical Magnetom Siemens scanner at NeuroSpin, a research center focused on the development of high-field MR imaging and spectroscopy and its application in neurosciences and translational research. Some practical and methodological aspects of this work are presented here.

2.1.1. Biospec Bruker 17.2 T Preclinical Scanner

The preclinical experiments were performed on a horizontal 17.16 T horizontal magnet (Biospec, Bruker BioSpin, Ettlingen, Germany) with an internal bore (without the gradient coil) of 25 cm and a magnetic drift of 5.4 Hz per hour. It is equipped with an actively shielded gradient system capable of delivering gradients up to 1 T/m on each axis with slew rates of up to 9000 T/m/s, allowing to switch from 5% to 95% of the maximum gradient strength within 100 µs and gradient linearity of 3% over 3.5 cm. The gradient bore diameter is 8.5 cm. The shim coils enable to correct for B₀ inhomogeneities up to the second order spherical harmonics. The spectrometer is controlled using Paravision 5.1. Details about the RF coils used for the experiments are provided in chapter 3 and 4.

2.1.2. Siemens Magnetom 7 T Scanner

Experiments in humans were performed on a Siemens Magnetom 7 T scanner (Siemens Medical System, Erlangen, Germany), equipped with an 8-channel Tx-array and a AC84 head gradient set (max. strength 80 mT/m, slew rate 400 T/m/s). A whole-body gradient (max. strength 45 mT/m, slew rate 200 T/m/s) is also present but its use has been currently set to a static B₀ shimming mode to dephase signal originating from the shoulder regions in imaging experiments. It is equipped with a set of shim coils adjusting up to the second order spherical harmonics. However, the actual shimming was done using the more efficient shim coils attached to the AC84 head gradient set. The system was built around an unshielded 90-cm-diameter-bore superconducting magnet (Magnex Scientific, Oxford, England). It is enclosed in a 300 ton steel room which acts as a Faraday cage and passive shielding, minimizing the leaking of external RF noise. The control of the scanner is carried out using the SyngoMR VB17 interface. The development platform is done in IDEA. The RF coils used for the work presented in this thesis will be presented in chapters 5 and 6.
2.1.3. $B_1$ calibration procedures

2.1.3.1. Experiments at 17.2 T

Due to the strong $B_1$ inhomogeneity when using surface coil, RF power calibration for $^1$H experiments was checked manually by optimizing the transmitter power while keeping constant all other acquisition parameters using a STEAM sequence centered on the VOI. The power difference (in decibels) between the system’s reference value and the calibrated power (for which the STEAM signal was maximized) was added manually to set the correct power of all other RF pulses. For $^{31}$P measurements, this calibration was a lengthier process since the PCr signal used is several orders of magnitude less intense than water. Therefore, signal averaging was required. In this case, the calibration was done with a FID-OVS sequence, the OVS being positioned between the RF coil and the VOI. The RF power for the adiabatic OVS bands had been determined beforehand and was kept constant for all experiments.

2.1.3.2. Experiments at 7 T

$B_1$ calibration for $^1$H experiments consisted in determining the reference voltage by measuring the $B_1$ field. The $B_1$ mapping methods used on this work were a modified version of the Actual Flip angle Imaging (AFI) sequence (Yarnykh, 2007; Amadon, et al., 2008; Boulant, et al., 2010) for single-channel transmit coils and the XFL sequence (Amadon, et al., 2010; Brunner, et al., 2008; Amadon, et al., 2012; Fautz, et al., 2008) for multi-channel transmit coils. The AFI sequence, which is considered the gold standard for $B_1$ mapping, determines the FA based on the ratio of two amplitude images $I_1$ and $I_2$ acquired with identical RF pulses, both applied within the same TR. The XFL sequence consists qualitatively, in the application of a sharp slice-selective saturation followed by a sharp excitation within the slice. The “saturated” images are then compared to a reference slice with no saturation to estimate the $B_1$ maps for each excitation channel. Further details about the AFI and XFL sequences are provided in chapter 5 and 6.

For $^{31}$P experiments, the reference voltage was derived from the reference voltage estimated for the $^1$H channel. In vitro reference values were acquired using STEAM with long repetition times ($TR = 40$ s) to reduce $T_1$-weighting. The relative differences in vitro between the $^1$H and $^{31}$P reference values corresponded to a factor of 1.57 at the center of the $^1$H/$^{31}$P coil. This ratio was assumed to be constant for any coil loadings.

2.1.4. $B_0$ shimming procedures

A major challenge for MRS at UHF is the greater inhomogeneity of the static magnetic field $B_0$. The frequency shifts and line-broadening that occur due to $B_0$ inhomogeneity can be one order of magnitude larger than the intrinsic linewidths leading to a degradation of the spectral resolution.
Therefore, $B_0$ inside the spectroscopic voxel must be as homogeneous as possible. To do so, $B_0$ shimming is performed using the gradient coils for first order spherical harmonic shimming and a dedicated set of shim coils for higher order shimming.

Among the $B_0$ shimming methods, the most common for MRI studies consists in mapping the $B_0$ field (Jezeard, et al., 1995) by acquiring two images using a fast-gradient recalled echo sequence where a phase evolution is allowed on the second image by increasing the TE. The $B_0$ field map is then obtained by solving:

$$\Delta \phi = \arctan \left( \frac{R_1 I_2 - R_2 I_1}{R_1 R_2 + I_1 I_2} \right)$$  \hspace{1cm} (2.1)

where $R_i$ and $I_i$ are the real and imaginary parts of the first ($i = 1$) and second ($i = 2$) acquisitions. This first method was used for the initial $B_0$ shimming prior to the acquisition of anatomical images. Since the determination of the phase difference between the two images is independent of the $B_1$ field homogeneity, $B_0$ mapping was also used for shimming on large volumes in the rat brain where strong $B_1$ inhomogeneity were expected.

The second method is the FASTMAP or the FAST(EST)MAP automatic shimming procedures (Gruetter, 1993; Gruetter, et al., 2000). The principle of the FASTMAP family method consists in measuring six orthogonal linear projections across the volume of interest, which can then be used to determine the required first- and second- order shim terms. The process is done iteratively and three to four iterations are usually sufficient to converge towards a stable shim configuration. For preclinical experiments, a Bruker’s implementation of FASTMAP was available and the FAST(EST)MAP sequence used on the Siemens 7 T scanner was provided by Dr. Malgorzata Marjanska (CMRR, University of Minnesota, USA). In general, using FASTMAP on small volumes provided better shimming results than the field map method.

Since both previous methods rely on calibrations of the different shim coils to calculate the best theoretical shim configuration, a third $B_0$ shimming procedure was applied to experiments done at 17.2 T. Using a STEAM sequence for localization, first-order shims intensities were iteratively optimized around the initial shim configuration so as to simply maximize the signal. This automatic shimming step was about 1 minute long for a TR = 3 s and was particularly useful to correct first-order shims for non-cubic volumes.

2.2. METHODOLOGICAL DEVELOPMENTS

2.2.1. Localization optimization

For preclinical studies, single voxel spectroscopy pulse sequences were primarily used and existing Bruker’s methods were either modified or new methods were programmed for Paravision 5.1. For experiments on the 7 T magnet, pulse sequences were edited using IDEA.
The FID-OVS localization was modified by replacing classical suppression bands with a $B_1$-insensitive train to obliterate signal (BISTRO) OVS scheme in order to improve the OVS effectiveness in regions presenting strong $B_1$ inhomogeneity.

The LASER method was implemented based on the existing source code of the PRESS method. In order to minimize both the chemical shift displacement artifact (CSDA) and the echo-time, frequency-modulated refocusing pulses with a high time-bandwidth product of 20 were chosen, resulting in a spatial displacement of 14.6% between 4.7 to 0.7 ppm at 17.2 T [eq. (1.48)] when 1-ms long RF pulses were applied. A minimal TE of 16.5 ms was achieved by applying short but intense spoiler gradients. The power requirements for different pulse waveforms were evaluated experimentally. For most SVS experiments, hermit and AFP HS8 pulses were used.

The main challenges of in vivo $^1$H MRSI at UHF are the $B_0$ and $B_1$ inhomogeneities together with the SAR limits which can be easily reached in the human head. For this reason, low energy-demanding and robust CSI sequences are needed. The two pulse sequences that were used in this work were the CSI-STEAM sequence, where a voxel is selected using STEAM, and the 2D CSI-FID where a single slab is selected. For the CSI-FID, OVS bands are required to suppress unwanted signal from the exterior of the brain. Similarly to the FIDLOVS sequence (Henning, et al., 2009), two variants of the CSI-FID sequence were developed with different OVS modules. One consisted in replacing the standard OVS sequence by a BISTRO scheme (presented below) and its application is shown in $^{31}$P MRSI in chapter 6. The second one was developed using static $B_1$-shimming methods to generate a “ring-like” excitation covering the external part of the brain at a low SAR cost. The theoretical background and its applications are presented in chapter 5.

### 2.2.1.1. $B_1$-insensitive train to obliterate signal (BISTRO) bands

The localization with the FID-OVS sequence was achieved by applying outer-volume-suppression bands enclosing the volume of interest, as explained in section 1.2.2.1. Using frequency-modulated pulses provides a good way to spoil unwanted signal through the combination of RF spoiling and strong crusher gradients. Nevertheless, $B_1$-sensitivity may still be an issue in very inhomogeneous $B_1$-fields: the adiabatic regime may not be reached due to insufficient RF power, resulting in a correct slice and bandwidth selection but with an excitation flip angles below 90°, leaving a portion of the $M_z$ magnetization unaffected. To confront this problem, a more effective suppression scheme was implemented, known as BISTRO for $B_1$-insensitive train to obliterate signal (Luo, et al., 2001). The BISTRO OVS scheme consists of a train of $n$ AFP pulses whose amplitudes $B_{1i}(t)$ are modulated as it would be for an adiabatic amplitude-modulating function:

$$B_{1i}(t) = B_{1\text{max}} \times F_{1i}(t) \times \text{sech}\left(\beta \left(1 - \frac{i-1}{n-1}\right)\right)$$

(2.2)

where $F_1$ is the amplitude-modulating function of individual $n$ pulses and $B_{1\text{max}}$ the maximal RF intensity of the last pulse.
Frequency-modulated RF pulses such as OIA are well suited to be used in the BISTRO pulse train as their $M_{xy}$ shape and width profiles obtained in sub-adiabatic conditions are very similar to those obtained under the adiabatic regime. The RF pulse train allows to achieve a homogeneous and effective RF spoiling in regions with strong $B_1$ inhomogeneity by gradually increasing the amplitudes of the individual pulses ($B_{1i}$). When multiple BISTRO bands are used, the spoiling efficiency can be further improved by interleaving the RF pulse trains, allowing $M_{xy}$ for each band to be spoiled longer by all the slice selection and crusher gradients.

A BISTRO module was implemented in Paravision 5.1 with $n=8$ and combined to the pre-existent FID-OVS localization sequence. Similarly, for the acquisition of $^{31}$P MRSI data in the human brain, BISTRO bands were added in a CSI-FID sequence with a maximum of 6 interleaved bands ($n=6$, figure 2.1).

The efficacy of the BISTRO bands was evaluated in vitro on both systems using either a FLASH sequence (at 17.16 T) or a 2D CSI-FID sequence (at 7 T). The BISTRO bands were compared to single-pulse OVS bands (figure 2.2). Measurement on the FLASH images revealed that the BISTRO scheme was more effective when low RF powers were applied (13.3 % increased suppression efficacy). Both the standard and the BISTRO OVS schemes were equally effective (less than 1% difference) when both were applied with elevated RF power (figure 2.2A). CSI measurements on the clinical 7 T scanner showed similar results, as illustrated by figure 2.2B (14.8 % increased suppression efficacy).
Figure 2.2. *In vitro* validation of the BISTRO bands at 17.2 and 7 Tesla

A. FLASH images acquired at 17.16 T using either 4 standard, single-pulse OVS bands or 4 BISTRO bands. Low- and high-power settings (12 dB difference) were used to evaluate the efficacy of the OVS bands. Contrast was enhanced for visualization. When using elevated power, no difference was noted between both OVS schemes. A 3-ms sech pulse was used for saturation (6.6 kHz bandwidth).

B. 2D CSI water map (matrix size: 16 x 16; FOV = 220 x 220 mm²) showing the relative water intensity across the phantom with and without BISTRO bands. Two water spectra identically scaled from the same ROI (white circle) are shown. An 8-ms HS8 pulse was used for saturation (2.5 kHz bandwidth).

2.2.2. Water suppression optimization

In this thesis, the water suppression (WS) was performed with a numerically optimized “water suppression enhanced through T₁ effects” (WET) method consisting of 3 or 4 frequency-selective Gaussian excitation pulses (Ogg, et al., 1994). This choice was done over other WS methods such as variable pulse powers and optimized relaxation delays or VAPOR (Tkác, et al., 1999), because of the reduced module duration and smaller energy deposition while still providing more flexibility than other common WS schemes such as chemical shift selective (CHESS) suppression (Haase, et al., 1985).

Since a greater number of pulses leads to a more efficient and B₁-insensitive WET scheme, 4 pulses were considered for the MRSI studies at 7 T instead of the 3 equally distant RF pulses found in the default Siemens implementation of WET.

Inter-pulse delays and nutation angles were optimized using equation (1.17). The spoiling was assumed to be ideal. The influence of each set of parameters on the suppression efficacy was evaluated for realistic ranges of T₁ values and B₁ distributions (Wyss, et al., 2013; Wright, et al., 2008). The suppression efficiency was defined as the sum of the residual signals for each T₁ and B₁ value combination under the WET parameterization being tested. The optimization algorithm constrains were the minimum and maximum inter-pulse delay. The minimum inter-pulse delay was set to the duration of the frequency-selective Gaussian pulses. The parameter corresponding
to the maximum allowed delay between successive pulses had a major impact on the suppression efficiency (figure 2.3).

For our experimental set-up at 7 T, the optimal solution for an allowed maximum delay of 380 ms was chosen. The parameters of the default WET scheme provided by the constructor in the CSI sequence as well as the numerically-optimized parameters are summarized in table 2.1. The theoretical performance of the solution is shown in figure 2.4. Assuming B\textsubscript{1} field variations of 10% from the nominal value and T\textsubscript{1} relaxation times ranging from 1000 to 5000 ms, the solution showed an improvement of a factor of 28 over the default parameterization provided by the constructor (figure 2.5). The increased suppression efficacy is experimentally demonstrated \textit{in vivo} in chapter 6.

For preclinical experiments at 17.2 T, \textsuperscript{1}H MRS studies on the rat brain were focused on small VOIs using a surface transceiver coil. Therefore a moderate B\textsubscript{1} field inhomogeneity was considered (20 % variation) for the WET optimization (figure 2.6). Additionally, single-inversion recovery experiments were carried out and for the purpose of reducing the minimum available TI, a short water-suppression scheme was needed. Therefore, only 3 pulses were considered for the WET module and 2 distinct set of parameters were calculated (table 2.1). As illustrated in figure 2.7, both WET modules were expected to be equally effective. Figure 3.5 shows spectra acquired with the two WET parameter sets. Figure 3.9 shows spectra acquired from different brain regions with the longer 269-ms-long WET module.

![Figure 2.3. WET efficiency as a function of the maximum allowed pulse delay using 4 pulses.](image)

Chosen Max. Delay: 380 ms

The efficiency of water suppression was evaluated by summing the residual M\textsubscript{z} signal for T\textsubscript{1} values ranging from 1000 to 5000 ms and B\textsubscript{1} field variations of up to 10% from the nominal value. The ranges of T\textsubscript{1} times were considered from previously reported values (Wyss, et al., 2013; Wright, et al., 2008) and on the observed B\textsubscript{1} inhomogeneity of 10% of our coil on typical VOI sizes used for in vivo MRSI acquisitions at 7 T. Since the solution was dependent on the initial conditions, 250 different initial conditions were evaluated for each point.
Figure 2.4. WET performance using 4 optimized flip angles and delays for experiments at 7 T. The ratio $M_z/M_0$ was used to express the efficiency of the water suppression for $T_1$ values ranging from 1000 to 5000 ms and $B_1$ field with up to 40% variations from the nominal value. The mean suppression factor over a 20% $B_1$ variation at the displayed $T_1$ ranges was 73163.

Figure 2.5. Water suppression efficiency as a function of the $B_1$ inhomogeneity considered at 7 T. The WS efficiency for a $T_1 = 1850$ ms with the optimized 4-pulse parameter set (blue) and the standard 3-pulse WET module proposed by the constructor in the CSI-FID sequence (red). The optimized WET module showed an improved efficiency by a factor of 28 for $B_1$ variations of 20% and $T_1$ ranges from 1000 to 5000 ms.
Figure 2.6. WET performance using optimized flip angles and delays for experiments at 17.2 T
At 17.2 T, small VOIs were studied and for this reason, $B_1$ inhomogeneity was considered to be mild. This assumption permitted to calculate WET parameters which were more efficient in relatively homogeneous $B_1$ fields (up to 20% variation). The results showed small water suppression efficiency changes with respect to $T_1$.

Figure 2.7. Water suppression efficiency as a function of the $B_1$ inhomogeneity considered at 17.2 T
The WET WS efficiency for a $T_1 = 2000$ ms with a module duration of 259 ms (blue) and 105 ms (red). Over the range of $T_1$ values from 1400 to 5000 ms and assuming 20% of $B_1$ variation, the shorter WET module was 28% more efficient according to numerical simulations.
<table>
<thead>
<tr>
<th>Experimental Setup</th>
<th>Total duration (ms)</th>
<th>Delay 1 (ms)</th>
<th>Delay 2 (ms)</th>
<th>Delay 3 (ms)</th>
<th>Delay 4 (ms)</th>
<th>Angle 1 (deg)</th>
<th>Angle 2 (deg)</th>
<th>Angle 3 (deg)</th>
<th>Angle 4 (deg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>7 T †</td>
<td>487</td>
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<td>56</td>
<td>30</td>
<td>15</td>
<td>100.4</td>
<td>82.2</td>
<td>87.1</td>
<td>116.6</td>
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<td>150</td>
<td>60</td>
<td>40</td>
<td>-</td>
<td>90.1</td>
<td>75.1</td>
<td>117.3</td>
<td>-</td>
</tr>
<tr>
<td>17.2 T §</td>
<td>105</td>
<td>35</td>
<td>35</td>
<td>25</td>
<td>-</td>
<td>84.5</td>
<td>79.5</td>
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<td>-</td>
</tr>
<tr>
<td>7 T ¶</td>
<td>188</td>
<td>60</td>
<td>60</td>
<td>60</td>
<td>-</td>
<td>89.2</td>
<td>83.4</td>
<td>160.8</td>
<td>-</td>
</tr>
</tbody>
</table>

Table 2.1. Parameters for the WET schemes used at 7 T and 17.2 T. The total duration includes RF pulses and spoiler gradient durations. The optimization algorithm was run with specific constrains for each of the three optimized schemes.

† Assumed B_1 inhomogeneity of 10 %, T_1 ranges from 1 to 5 s, maximum inter-pulse delay of 380 ms.
‡ Assumed B_1 inhomogeneity of 20 %, T_1 ranges from 1.4 to 5 s, maximum inter-pulse delay of 150 ms.
§ Assumed B_1 inhomogeneity of 20 %, T_1 ranges from 1.4 to 5 s, maximum inter-pulse delay of 35 ms.
¶ Default parameters of the WET scheme provided by the constructor for the CSI-FID pulse sequence.

### 2.3. Macromolecule Baseline Parameterization

At short echo-times, *in vivo* spectra acquired in the brain not only show contributions from mobile low-molecular-weight molecules such as metabolites but also present broad signal contributions from high-molecular-weight lipids and proteins (Behar, et al., 1994). Previous studies in humans and rodents have assigned some of these resonances to methyl and methylene resonances of amino acids such as leucine, isoleucine, valine, threonine, alanine, lysine, arginine, glutamate, glutamine and α-methylene protons (Behar, et al., 1993). The macromolecule (MM) signal originates mostly from their mobile chains, whereas the bulk of the macromolecular protons, having a restricted mobility and very short T_2 values are difficult to observe directly. MMs have different physical properties than metabolites: they possess shorter T_1 and T_2 relaxation times and 10 to 20 times shorter apparent diffusion coefficients (de Graaf, et al., 2006; Kreis, et al., 2005; Pfeuffer, et al., 2000). These properties are used for the acquisition of “metabolite-nulled” spectra or to reduce MM contributions in ¹H spectra.

#### 2.3.1. Macromolecule baseline fit

Macromolecule NMR signals are an important aspect to consider in short-TE ¹H MRS studies as they largely overlap with low concentrated metabolites, introducing systematic quantification errors if they are not properly accounted for. Furthermore, their own, proper quantification may provide useful clinical information as it has been shown in different pathologies such as in stroke, in the presence of tumors or in multiple sclerosis (Kaminogo, et al., 2001; Saunders, et al., 1997; Narayana, et al., 1992; Davie, et al., 1994).

Several methods have been proposed to fit the MM baseline (Cudalbu, et al., 2012). One common approach is to acquire a “metabolite-nulled” spectrum using either the inversion-recovery (IR) or the progressive saturation recovery methods. The MM spectrum is then included as a basis
element for the spectral decomposition or simply subtracted from the short-TE spectrum to obtain a macromolecule-free neurochemical profile. The approach adopted in this thesis was to parameterize the MM baseline using LCModel based on experimental metabolite-nulled spectra (Seeger, et al., 2003). This technique can be easily adapted for the analysis of spectra with strong T\(_1\)- or T\(_2\)-weighting and permits the reliable quantification of separate MM resonance groups or components.

2.3.2. Metabolite-nulled spectrum using DIR

For our preclinical experiments at 17.2 T, a Double Inversion Recovery (DIR) module (TI\(_1\)/TI\(_2\) = 2600/600 ms) was combined with a LASER localization to acquire metabolite-nulled spectra as shown in figure 2.8. Two adiabatic AFP HS8 inversion pulses (2 ms, 10 kHz bandwidth) were placed prior to the localization sequence. The DIR module was chosen over a SIR module due to the increased T\(_1\)-weighting introduced by the second inversion pulse. The inversion delays TI\(_1\) and TI\(_2\) were adjusted to achieve minimal metabolite residual contributions.

![Figure 2.8. Metabolite-nulled spectra at varying echo times at 17.2 T.](image)

Metabolite-nulled spectra acquired from the rat brain using LASER (128 avg, 1800 ms TR) with T\(_1\)-weighting introduced with a DIR module (TI\(_1\)/TI\(_2\) = 2600/600 ms) at echo times varying from 16.5 to 80 ms. Due to their short T\(_2\), MM resonance intensities rapidly decay with increasing TEs. Metabolites with short T\(_1\) relaxation times showed residual signal, notably from Cr(CH\(_2\)) and tCho. Their contributions can be clearly observed at the longer TEs.
2.4. SPECTRAL DECOMPOSITION

The analysis of $^{1}$H and $^{31}$P spectra was done using LCModel 6.2 (Provencher, 1993). This software uses a basis set of predefined spectra to calculate a linear combination of its components so as to fit an experimental MR spectrum in the spectral domain (de Graaf, et al., 1990). The concentrations of the fitted metabolites (expressed in mM) are calculated with respect to a reference signal which can be either an external water reference spectrum or a metabolite found in the spectrum (usually tCr or NAA). This is done by comparing the integral of each individual metabolite with that of the reference compound. However, the integral for each metabolite or functional group is weighted by $T_1$ and $T_2$ relaxation effects that may not correspond to those in the compound of reference, resulting in an over- or underestimation of their concentration. These “apparent” concentrations can be corrected to yield “absolute” concentrations, which will be further discussed in section 2.5.

Other methods and software packages exist for the analysis of spectra, the majority in the time domain: AMARES (Vanhamme, et al., 1997), MRUI (Naressi, et al., 2001), VARPRO (van der Veen, et al., 1988) and others (Slotboom, et al., 1998). The LCModel software has, nevertheless, the advantage of including into the spectral analysis experiment-dependent factors which are difficult to model, such as Eddy current and spatial profiles, which can be corrected using a water reference signal. It is also well suited for handling J-coupled resonances as the spectral patterns at different echo-times can be directly provided with measured or simulated spectra.

2.4.1. Basis set generation using the spin density formalism

The first step is the constitution of the basis set of metabolite spectra with which the in vivo data will be fitted. Although it is possible to use in vitro spectra acquired experimentally using the same MRS sequence as for the in vivo experiment, this approach is sensitive to experimental biases such as the temperature and pH conditions (Kaiser, et al., 2010). For this work, the basis sets were constituted of simulated spectra generated using the spin density formalism. This approach calculates the evolution of the initial density matrix by the application of energy operators corresponding to the RF pulses, applied magnetic field gradients and delays that constitute the experimental pulse sequence (Mulkern, et al., 1994). The spin density matrixes were defined using published chemical shifts and J-coupling values for $^{1}$H (Govindaraju, et al., 2000; Near, et al., 2012) and $^{31}$P (Jung, et al., 1997; Jensen, et al., 2002) metabolite resonances. The practical generation of the basis sets was done in Matlab (The MathWorks, MA, USA) using spin density simulation software developed by R.A de Graaf (SpinWizard, Magnetic Resonance Research Center, Yale School of Medicine). The metabolites that were included in the basis sets for the $^{1}$H and $^{31}$P studies in the aging rat brain are given in section 3.2.5 and 4.1.4.1.
Simulated \(^1\)H spectra of \(N\)-acetyl aspartate, phosphocreatine, glutamate, glutamine, myo-inositol and taurine were compared with \textit{in vitro} acquisitions done at room temperature at varying TE (16.5, 24, 35, 50, 60, 80, 100, 120, 140, 160, 180 and 200 ms). Concomitantly, the simulated basis-sets were calibrated so as to obtain the correct concentrations when using an unsuppressed water spectrum as an internal reference of concentration.

2.4.2. LCModel Parameterization

LCModel is based on a constrained non-linear optimization algorithm that minimizes the difference between a linear combination of model spectra and the spectrum being fitted. In order to achieve results with minimal residual noise, several parameters can be adjusted (frequency shifts, linewidths, zero- and first-order phases, etc). The control parameters also allow setting soft constraints to improve the consistency and the quality of the fits. Among the most relevant ones are the predefined concentration ratios, initial phase parameters and the expected variability ranges. This set of parameters is particularly important for the quantification of low-concentrated metabolites such as GSH and GABA and for inversion-recovery experiments where metabolites are close to zero-crossing (figure 2.9). Up to 18 metabolite ratios were used for the proper fitting of \(^1\)H spectra and 2 for the \(^{31}\)P spectra.

It should be noted that the metabolite ratios were determined based on typical metabolite concentrations for healthy rats (de Graaf, 2007 p. 51). Only soft ratio constraints were used in LCModel due to the wide range of concentration values reported in the reference. Additionally, the neurochemical profile varies with the rat strain, age, pathological state or brain region (Hong, et al., 2011; Tkac, et al., 2003; Ross, et al., 2010; de Graaf, et al., 2011). The analysis of brain extracts through biochemical measurements such as high-resolution liquid-phase NMR or high-resolution magic angle spinning (HR-MAS) NMR permits to determine with high precision the concentration of metabolites and therefore to improve the ratios used in LCModel. Nevertheless, caution should be taken as immobile or NMR-invisible metabolite pools revealed in liquid-state NMR, such as glutamate (Kauppinen, et al., 1994), could bias the expected concentration values observed in \textit{in vivo} MRS. Similarly, handling and storage conditions of the sample as well as the mechanical stress from spinning in MAS NMR have shown to cause significant increases in metabolites (Esteve, et al., 2014) such as the release of NMR-invisible bound creatine (Opstad, et al., 2008), alanine, glucose and lactate (Opstad, et al., 2008), increased levels of acetate and aspartate due to NAA degradation (Cheng, et al., 1997) and loss of metabolites of up to 30 to 50% due to washing of the thawed sample (Bourne, et al., 2003).
LCModel fit of a T$_1$-weighted $^1$H MRS spectrum acquired in the rat brain in vivo at 17.2 T using SIR-LASER (TR/TE/TI = 5000/16.5/1000 ms, 128 avg). A priori knowledge of the sign of metabolite and macromolecule contributions and fitting soft constraints were essential for achieving meaningful and consistent fitting results in low SNR spectra. The shown spectrum is phased so as to exhibit a positive tCr(CH$_3$) contribution.

2.4.3. Macromolecule Baseline Parameterization

For short-TE $^1$H MRS, it is critical to correctly fit the macromolecule baseline. Although LCModel offers to fit the slow modulation of the MM baseline using a cubic spline function, the results may not be satisfactory and it is not suited for fitting heavily T$_1$-weighted spectra (figure 2.9). The approach adopted for this thesis was to use the LCModel built-in Gaussian functions to account for the MM signal. To this end, a high-SNR MM spectrum was obtained from averaging the metabolite-nulled spectra acquired from the brains of 6 healthy rats using a DIR-LASER sequence (TR/TE/TI$_1$/TI$_2$ = 5000/16.5/2400/600 ms, 128 averages, 2048 complex points, further details in section 3.2.4). To fit this MM spectrum, more than 40 Gaussian-shaped functions were considered as well as the residual resonances of tCr(CH$_3$). Other metabolites with short or very long T$_1$ were later added into the fit: tCho, tNAA(CH$_2$) and Tau (figure 2.10). Strict metabolite concentration ratios were considered between them based on their estimated T$_1$ relaxation times (figure 2.10). The MM parametrization was confronted to series of T$_1$- and T$_2$-weighted spectra acquired in the rat brain (n = 6) for validation, as shown in figures 3.5 and 3.6. The retained MM baseline parameterization consisted of 32 individual elements grouped into 4 MM components.
(figure 2.11 and table 2.2). The definition of 4 groups of MM components based on their resonance frequencies was necessary to limit the degrees of freedom for their fit.

Figure 2.10. Residual metabolite magnetization calculated for metabolite-nulled spectra at 17.2 T. Residual signal after the application of a double-inversion recovery module (TI1/TI2 = 2600/600 ms) as a function of the T1 relaxation time with a recovery time of 1800 ms. The metabolites considered to have an important residual contribution for the parameterization of the MM baseline were tCr(CH2), tCho, NAA(CH2) and Tau. T2-weighting was ignored. The considered T1 values were: Cr(CH3): 1753 ms, tCr(CH2): 1203 ms, tCho: 1408 ms, NAA(CH2): 1527 ms and Tau: 2211 ms.

Figure 2.11. MM baseline parameterization at 17.2 T in the rat brain. Average macromolecule spectrum (black) and its LCM offense (red). The macromolecule baseline was fitted by 4 non-overlapping groups of MM resonances. The CRLB values were 1% for MM1, MM2 and MM3 and 2% for MM4. No filtering was applied and all spectral elements were scaled identically. MM definitions and relative amplitudes are shown in Table 2.2. Due to their short or long T1 relaxation times, residual signals from tCr(CH2), tCho, tNAA(CH2) and Tau were also considered for the fit and their contributions were subtracted.
Table 2.2. Definition of the MM components used for the rat brain at 17.2 T

Definition of the 4 MM components used for fitting the macromolecule and lipid baseline in LCModel. The chemical shift, linewidth and amplitude parameters of the Gaussian basis functions were derived from the mean metabolite-nulled spectrum shown in figure 2.11. The apparent signal integrals are normalized with respect to MM1.

Figure 2.12. LCModel fit of metabolite-nulled spectra at varying echo times.

T2-weighted metabolite-nulled spectra acquired in vivo at 17.2 T from a single animal (black, left) using LASER (TR/TI1/TI2 = 1800/2600/600 ms, 128 avg). Their LCModel fit (red, left) was done using the macromolecule parameterization proposed in Table 2.2 and their corresponding residuals are also shown (right). Due to their short or long T1 relaxation times, residual signals from tCr(CH2), NAA(CH2), tCho and Tau were also considered in the fit. No filtering was applied and all spectral elements were scaled identically. At TE of 50 and 60 ms, J-coupled MM residuals were observed at 2.2, 3.05, 3.7 and possibly at 4.33 ppm. Residual signal was also found at 1.45 ppm, probably due to a difference in T2 relaxation times among the macromolecule resonances which constitute our MM1 component.
The definition of separate MM components and MM ratios permitted to properly fit T₁- and T₂-weighted metabolite spectra (see chapter 3). No MM J-coupling effects were considered for fitting the MM baseline at TE>16.5 ms (Behar, et al., 1994; Henry, et al., 2001). Nevertheless, only minor residuals contributions were observed at TE of 50 and 60 ms when the MM parameterization was tested on two metabolite-nulled data sets at varying TEs (figure 2.12).

2.4.4. Error estimation

To measure the level of uncertainty of a spectral decomposition using LCModel, the so-called Cramer-Rao Lower Bounds (CRLB) are considered. The CRLBs provide a lower bound for the minimum possible error on the parameter estimation given by LCModel. It is defined as the inverse of the covariance matrix of the parametric model used to describe the experiment, taking into account the experimental noise. From this definition, it follows that the CRLBs are only an approximation of the “real” error estimation as the parametric model is not a perfect model of the real data and as such the omission of important fitting parameters may provide unrealistic CRLBs. The inclusion of parameterized MM components may thus highly reduce the CRLBs of overlapping metabolites. In practice, the CRLBs are used as an index of the reliability of the LCModel analysis since lower SNR data exhibit higher CRLBs. It is generally accepted that metabolites displaying CRLBs < 10% are determined with sufficient precision whereas CRLBs between 20 to 30% should be considered with caution. In this thesis, metabolite concentration results were discarded when their CRLBs were above 25%.

2.5. ABSOLUTE QUANTIFICATION

NMR spectra provide information that can be used to derive the absolute concentrations of several chemical compounds. Although calculating relative metabolite ratios provide pertinent biological information, it can be ambiguous for pathologies where multiple variations occur (or are expected to occur) simultaneously. For instance, total creatine, a commonly used internal reference of concentration, has been observed to change in the normal aging brain or in diseases such as in tumor and stroke (Howe, et al., 2003; Kinoshita, et al., 1997). Calculating absolute concentrations becomes a necessary approach when tissue metabolism is altered in an unpredictable manner or when apparent metabolite concentrations alterations are due to concomitant changes in metabolite relaxation times.

2.5.1. Quantification strategy for ¹H MRS

In this work, the method of absolute quantification was to use the water resonance as an internal reference of signal. It has been extensively used in the past (Kaiser, et al., 2005; Kreis, et al.,
1993) and it consists in comparing the metabolite signal to the water signal from the same VOI assuming a known water content. For the correct estimation of the tissue water concentration in the VOI, tissue segmentation is required since different tissue types are known to have different water concentrations (Lentner, 1981).

Among the caveats to consider when water is used as a reference of concentration is that tissue water content varies with age. Diffusion tensor imaging (DTI) studies have shown that the newborn brain has higher water contents than the adult brain in both gray matter (GM) and white matter (WM) tissues (Neil, et al., 1998). In the aging brain, shrinkage of the brain volume has been observed (Raz, et al., 2005) and alterations of the WM microstructure were found with DTI (Pfefferbaum, et al., 2000; Coutu, et al., 2014), although neuronal density is preserved (Terry, et al., 1987; Mrak, et al., 1997). Variations in water content have a direct impact on metabolite quantification and their monitoring is an important aspect during longitudinal studies. For the particular case of the rat, the adult brain size is achieved until the 60th post-natal day (Dobbing, et al., 1971) and myelination is achieved at 90th post-natal day (Juraska, et al., 2004) although it has been observed to continue in the cortex after 3 month (Mengler, et al., 2014). This indicates that important WM and GM water content variations occur during the first 3 months of age.

While the first developmental stages of the rat brain are rather well characterized (Watson, et al., 2006), this is not the case for the elderly rat. The mean water content in the mouse brain has been observed to decrease during the second year of age from 79% to 75% (Duarte, et al., 2014), providing indicative values for the evolution of water content in the rat brain but a thoroughly study is required to determine the water content in the aging rat brain.

In addition to the water content of each compartments, correction for relaxation effects have to be considered to achieve absolute quantification. For our preclinical studies, tissue segmentation was based on T1 maps which were also used to estimate the T1 relaxation times of water in GM, WM and cerebrospinal fluid (CSF). Segmentation was done based on a rat brain atlas (Paxino, et al., 1998). T1 relaxation times of “pure” GM, WM and CSF were expected to correspond to those of the cingulate cortex (GM), corpus callosum and optical nerve (WM) and the lateral ventricles (CSF). GM and WM quantification was achieved by first determining the mean T1 value from a manually-drawn VOI and then calculating the proportions of “pure” GM and WM required to reproduce it, where the GM and WM reference values were taken from the same T1 map. Pixels showing an elevated T1 value (above 2400 ms) were considered to correspond to CSF and were excluded from the calculations.

CSF contributions need to be considered to account for partial volume effects, which may otherwise have a “diluting” effect on the tissular concentrations in the brain. Proper discrimination of the CSF water signal can be achieved by acquiring water spectra of the VOI with varying TEs and applying a double-exponential fit on the decaying water signal (Kreis, et al., 1993; Gasparovic, et al., 2006). This method also allowed to determine the T2 relaxation time of brain tissue water.
With these elements, the apparent water concentration in the VOI can be calculated as:

\[
[W]_{VOI} = [W]_{GM} R_{GM} + [W]_{WM} R_{WM} + [W]_{CSF} C_{CSF}
\]  (2.3)

where \([W]_{GM}\), \([W]_{WM}\) and \([W]_{CSF}\) are the GM, WM and CSF water concentrations; \(R_{GM}\), \(R_{WM}\) and \(R_{CSF}\) their respective T1 and T2 weightings given by:

\[
R_C = \exp\left(-\frac{TE}{T_{2,C}}\right) \left(1 - \exp\left(-\frac{TR}{T_{1,C}}\right)\right)
\]  (2.4)

where \(T_{2,C}\) and \(T_{1,C}\) are the compound or tissue T2 and T1 relaxation times.

Similarly to other studies (Christiansen, et al., 1993 pp. p. 466-472; de Graaf, et al., 2006; Kaiser, et al., 2005), the absolute metabolite concentrations \([\text{met}]\) were then calculated as follows:

\[
[\text{met}] = \frac{S_m}{S_W} \frac{[W]_{VOI}}{[(1 - f_{CSF}) R_m]}
\]  (2.5)

where \(S_m\) and \(S_W\) correspond to the metabolite and water measured signal intensities and \(R_m\) was determined for each metabolite using their estimated T1 and T2 relaxation times obtained from measured T1- and T2-weighted spectra.

### 2.5.2. Quantification strategy for \(31^P\) MRS

The absolute quantification strategy in \(31^P\) studies consisted in comparing \(31^P\) MRS signal to the \(1^H\) water signal. The water spectrum has to be acquired from the same VOI in order to preserve an identical spatial dependence. Similarly to eq. (2.5), the absolute metabolite concentration \([\text{met}]\) can be determined using:

\[
[\text{met}] = \frac{S_m}{S_W} \frac{[W]_{VOI}}{[(1 - f_{CSF}) C_{PH} R_m]}
\]  (2.6)

where \(C_{PH}\) is a scaling factor accounting for the difference in \(1^H\) and \(31^P\) sensitivities of the dual coil used (Buchli, et al., 1994; Bottomley, et al., 1996). \(C_{PH}\) was estimated from in vitro measurements directly after each in vivo experiment using identical coil and VOI positioning as for the in vivo MRS acquisitions and assuming the same sensitivity and B1 profiles for both \(31^P\) and \(1^H\) RF coils. Since the acquisition delay of the FID-OVS localization sequence was short, the signal loss due to T2 relaxation was considered to be minor and therefore T2-weighting was not accounted for. For each \(31^P\) spectrum analyzed in LCModel, the parameter FCALIB was set to the specific value \([W]_{VOI} C_{PH}/S_W\) since the \(1^H\) raw water spectrum was not considered by the software as a proper reference scan.
2.6. WATER $T_1$ MAPPING

Water $T_1$ mapping was important for the purpose of tissue segmentation and for the determination of the apparent water concentration in tissues and in our VOIs. The methods used for $T_1$ estimation are presented in section 1.1.6.

2.6.1. MRI sequence and the Look-Locker method

Among the different $T_1$ mapping methods, the Look-Locker method (Deichmann, et al., 1999; Look, et al., 1968) was used based on the acquisition of 3D inversion-recovery fast gradient echo (IR-FGE) images. Unlike the standard inversion-recovery method where a 90° excitation is applied after the inversion delay TI, the Look-Locker method applies successive small FA excitation pulses allowing to sample the longitudinal recovery at numerous inversion times within a single TR$_1$ (figure 2.13). The final $T_1$ maps are generated by fitting the IR-FGE data as a function of TI using a 3-parameter model (Marty, et al., 2013) implemented in C and Matlab (The MathWorks, MA, USA). The theoretical framework derives from Deichmann et al. (Deichmann, et al., 1992). First, an apparent $T_1^*$ relaxation time is determined from the acquired data by adjusting the 3-parameter model:

$$M(t) = A - B \cdot \exp \left(-\frac{t}{T_1^*}\right)$$  \hspace{1cm} (2.7)

where $A = M_0^* = M_0 T_1^*/T_1$; $B = M_0 + M_0^* = M_0 (1 + T_1^*/T_1)$ and $M(t)$ is the measured signal at $t = TI$. The $T_1$ relaxation time can then be estimated as:

$$T_1 = T_1^* \left(\frac{B}{A} - 1\right)$$  \hspace{1cm} (2.8)

It should be noted that the choice of the flip angle $\alpha$ is critical for the correct estimation of $T_1$. Although the precedent equations do not require the knowledge of $\alpha$ for the calculation of $T_1$, it has a direct impact on the rate at which the steady-state magnetization is achieved. Using an elevated flip angle therefore reduces the quality of the estimation as only the first points for each TR$_1$ provide information allowing to estimate and differentiate compounds with different $T_1$ times (figure 2.14). Choosing short flip angles provide more reliable results at the expense of a reduced SNR. For our experiments, nominal flip angles of 5° degrees were targeted.
At $t = 0$, an inversion pulse is applied. After a first $T_1$ delay, a train of $N$ small excitation pulse $\alpha$ is applied every $TR_2$ leading to the measure of $k$-space line after an effective inversion time $TI_i = TI_1 + (i-1)*TR_2$. The pulse sequence is repeated every $TR_1$ until all $k$-space lines are covered.

Figure 2.14. $T_1^*$ dependence on the flip angle for two different relaxation times. The choice of the flip angle has a direct impact on the measured $T_1^*$ relaxation time. Choosing elevated flip angles reduce the contrast between different $T_1$ times and cause shorter $T_1$ values to be estimated. Illustration from: (Deichmann, et al., 1992).

2.6.2. Sequence evaluation at 17.2 T

One limitation of the IR-FGE pulse sequence is that it requires a homogeneous inversion pulse over the entire VOI. Achieving an inversion may become challenging when using a surface coil due to the strong $B_1$ field inhomogeneity profile. To tackle this problem, an AFP inversion pulse (HS8, 4 ms, 5 kHz bandwidth) was used. To test the robustness of this method at 17.2 T, the
results acquired on 6 rats using a transceiver surface coil were compared to data acquired using a volume coil in an equivalent set of 6 rats (similar weight and age). The sequence parameters and T$_1$ map reconstruction are detailed in sections 3.2.3 and 3.2.6. Two other sequences, the IR-FISP (Schmitt, et al., 2004) and the RARE (Hennig, et al., 1986), were also tested with the volume coil.

The results obtained from the volume coil were highly reproducible while the IR-FGE maps acquired with the surface coil were discouraging as they showed unrealistically low values ranging from 1200 to 1500 ms for WM and GM in regions close to the surface coil and increasingly high values as the imaging slice was further away from the coil (up to 1850 ms for GM, ~1500 ms for WM). Based on figure 2.14: this behavior is assumed to be the consequence of a high excitation FA close to the coil which was gradually reduced for deeper structures leading to an increasing T$_1^*$ contrast and higher T$_1^*$ values. Measuring T$_1$ maps using a surface coil with the Look-Locker method was therefore abandoned.

When comparing T$_1$-mapping data obtained using the volume coil, consistent results were found between the IR-FGE and IR-FISP sequences, while the RARE sequence showed more elevated values. This overestimation of T$_1$ relaxation times with RARE could be due to the application of an inhomogeneous refocusing pulse over the VOI, which was performed by an amplitude-modulated pulse (hermit modulation), contrary to the adiabatic inversion pulse used for the other two pulse sequences (HS modulation). The results are summarized in Table 2.3 and two T$_1$ maps from two rats are shown in figure 2.15. Interestingly, a constant factor of 1.4 was found between the measured T$_1$ times determined with the RARE sequence and the IR-FISP or IR-FGE sequences.

<table>
<thead>
<tr>
<th>Sequence</th>
<th>VOI (ms)</th>
<th>GM (ms)</th>
<th>WM (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>IR-FGE</td>
<td>1839 ± 54</td>
<td>1877 ± 77</td>
<td>1670 ± 34</td>
</tr>
<tr>
<td>IR-FISP</td>
<td>1850 ± 62</td>
<td>1925 ± 11</td>
<td>1669 ± 37</td>
</tr>
<tr>
<td>RARE</td>
<td>2602 ± 60</td>
<td>2724 ± 121</td>
<td>2369 ± 47</td>
</tr>
</tbody>
</table>

Table 2.3. T$_1$ relaxation times measured using 3 different T$_1$-mapping methods using a volume coil.

The volume-of-interest (5x5x2 mm$^3$, figure 2.5) consisted mostly of cortex (~60%) and had contributions from the hippocampus and the corpus callosum.
Figure 2.15. T1 maps acquired at 17.2 T using the IR-FGE and IR-FISP sequences. T1 maps reconstructed from two healthy rats. The data sets were acquired using an IR-FGE (TE/TR1/TR2 = 3.02/10/10000 ms, 60 TIs ranging from 90 to 9530 ms, 6 segments, 125x125x1000 µm resolution) and a IR-FISP (TE/TR1/TR2 = 4/8.4/10000 ms, 40 TIs ranging from 150 to 5064 ms, 6 segments, 250x250x1000 µm resolution). The images were cropped to focus on the brain area covered by our spectroscopic VOIs. Average T1 value from the purple VOI is given in table 2.3. The T1 maps are scaled identically.
PART II: PRECLINICAL STUDIES

3. QUANTITATIVE STUDY OF METABOLIC ALTERATIONS DURING AGING IN THE RAT BRAIN

MRS is a technique that allows measuring non-invasively the concentration of brain metabolites \textit{in vivo} and is very well suited for longitudinal studies. As previously discussed in chapter 1, MRS benefits from increased SNR and spectral dispersion at higher magnetic fields. Working at UHF is therefore propitious for metabolite quantification studies.

The work presented in this chapter was done at 17.2 T using quantitative \textit{in vivo} \textsuperscript{1}H MRS with the objective to follow the changes of the neurochemical profile during normal aging in the rat brain. In this longitudinal study, data was gathered over an age span of 18 months in healthy Dark-Agouti rats with the purpose of assessing this strain as a practical rodent model of brain aging.

3.1. INTRODUCTION TO NORMAL BRAIN AGING

The process of aging is a complex process associated with a progressive decline of biological, cognitive and psychomotor functions whose rate is partially determined by the genetic background but it is also influenced by nutrition, lifestyle and environmental factors (Busse, 1987). Changes observed affecting the central nervous system (CNS) include a decline in cognitive performance, as it was observed for 38\% of people aged between 60 to 78 years old (Kolvisisto, et al., 1995), affecting episodic memory processing, working and spatial memory as well as a declining processing speed (Hedden, et al., 2004; Woodruff-Pak, et al., 1988). Elderly people also show a slowing of motor movements and loss of fine motor control (Mattay, et al., 2002). Whether these alterations are due to neuronal loss and cell damage (Pakkenberg, et al., 1997) or to an age-related reduction in synaptic efficacy (reduced neurotransmitter supply, reuptake and receptor abundancy) (Morrison, et al., 2012) is in debate (Mattay, et al., 2002; Yeoman, et al., 2012).

Aging is the primary risk factor for stroke (Duijn, et al., 1992; Moreno-Torres, et al., 2005) and neurodegenerative disorders such as AD (Heun, et al., 1997) Parkinson’s disease (Ellis, et al., 1997) or other types of dementia (Kantarci, et al., 2004; Charles, et al., 1994; Kantarci, et al., 2006). Common features of neurodegenerative diseases are elevated levels of reactive oxygen species (ROS) and an accumulation of misfolded proteins in the CNS (Hung, et al., 2010). It has been suggested that the elevated levels of ROS are not attributed to an increased oxidative stress but rather to a functional decline of the antioxidant defense and repair mechanisms (Andersen, 2004). Neuroinflammation, which involves the activation of glial cells (mainly microglia) and an eventual infiltration of peripheral immune cells through the blood-brain barrier, has been
observed in a majority of neurodegenerative diseases as well as in the normal aging brain. However, it is not clear if there is a causal relation between chronic neuroinflammation, which creates neurotoxic and neurotrophic factors (Power, et al., 2001), and the onset of the disease (Garden, et al., 2006; Witte, et al., 2010).

With the improvement of health care over the last century, the average life expectancy has increased from 68.3 years in 1950 to 77.5 in 2003 and the proportion of elder people is expected to double worldwide over the next 50 years, leading to a predictable larger rate of neurodegenerative diseases (Hoyert, et al., 2005). In particular, the prevalence of AD nearly doubles with every five years of age and for instance, in 2000 dementia affected almost 30% of those aged 85 to 89 in France and it is the 6th leading cause of death in the USA. To date, only treatments for the symptoms are available but no cure has been found and the morbidity costs associated to aging are a primary challenge to health and social care systems. The associated expenses due to hospitalization or custodial care were evaluated as 1% of the world gross income in 2010 (Joseph, et al., 2009) and are expected to increase in the following decades (Comas-Herrera, et al., 2007; Alzheimer's Association, 2013).

With the advent of MRS and MRI, disease progression can be followed non-invasively over time (Parsons, et al., 2000; Herminghaus, et al., 2003). However, changes in brain metabolism due to healthy aging may become a confounding effect with respect to disease-related alterations. Yet, it is not well established which are the changes of the major metabolites during healthy aging in the human brain as results found in the literature vary among brain regions and in some cases contradictory trends have also been reported, notably for NAA in the frontal cortex (Haga, et al., 2009; Reyngoudt, et al., 2012).

### 3.1.1. Neurobiology of normal aging in the human brain

Normal brain aging in humans has been associated with a wide range of landmark alterations detected with several imaging techniques. A loss of neurons and brain volume reduction has been associated with age although neuronal density remains intact (Terry, et al., 1987; Mrak, et al., 1997; Raz, et al., 2005). Changes in white matter have also been observed in the elderly and it has been associated with altered functional connectivity (Almkvist, et al., 1992; O'Sullivan, et al., 2001; Esposito, et al., 1999). Reduction of fractional anisotropy and diffusional kurtoses has revealed changes in white matter microstructure with age (Pfefferbaum, et al., 2000; Coutu, et al., 2014). Decreases in cerebral blood flow have also been found in the frontal and parietal cortices (Loessner, et al., 1995) using positron emission tomography (PET). At the cellular level, the progressive decline in the CNS has been associated to the accumulation of damages due to oxidative stress in the form of ROS, impaired cellular energy metabolism and DNA damage repair, telomere erosion and accumulation of damaged proteins and organelles (Harman, 1956; Mattson, et al., 2006; Floyd, et al., 2002; Sahin, et al., 2010).
Among the probable causes are the oxidative stress generated by excessive levels of mitochondrial free radicals and the accumulation of protein aggregates due to dysfunctional mitochondria and reduced cellular autophagic capacity (Rubinsztein, et al., 2011; Moràn, et al., 2012). Microglia act as the active immune cells in the CNS and upon activation due to insults in the brain they release inflammatory mediators (including ROS and nitric oxide) to further recruit microglia and to a lesser extent activate astrocytes, which in turn may allow the entry of leukocytes by disrupting the blood-brain barrier (Garden, et al., 2006). Although neuroinflammation has a protective role, chronic neuroinflammation can become neurotoxic since fully activated microglial cells release a variety of compounds (ROS, nitric oxide, tumor necrosis factor-α,…) in quantities which can in turn damage neurons, oligodendrocytes or extracellular matrix structures (Block, et al., 2007; Power, et al., 2001; Sawada, et al., 2008). Chronic neuroinflammation has thus been increasingly recognized as a process underlying neurodegeneration (Witte, et al., 2010; Sawada, et al., 2008). Yet, it is not clear whether neuroinflammation in the elderly is onset by aging effects alone (Suridjan, et al., 2014). Furthermore, several studies have shown that the initial microglial activation is induced by the expression of potent pro-inflammatory factors such as IFN-γ, whose expression is increased in the brain of aged animals. However, their source has not been identified (Lynch, 2009) although glia-neuron cross-talk seems to have a central role and high IFN-γ levels impede cell renewal (Walter, et al., 2009). Therefore, detecting the onset of neuroinflammation and its progression may be of particular interest to better understand the process of aging.

### 3.1.2. NMR Spectroscopy of aging

#### 3.1.2.1. Clinical studies

Numerous MRS studies have looked at the aging human brain (Reyngoudt, et al., 2012; Haga, et al., 2009) presenting, in general, an increase of tCho, Ins and tCr with age. Choline-containing compounds and myo-inositol are, respectively, biomarkers of membrane turn-over and of the glial compartment (Jenden, 1979 pp. 13-24; Brand, et al., 1993). Their increased levels points at an astroglial activation and reactive gliosis, as observed during neuroinflammation (Chang, et al., 2013).

On the contrary, NAA changes with age showed less concluding results as it was observed to increase, show no change or decrease even within the same studied brain region. NAA is localized and synthetized mostly in neurons (Moffett, et al., 2006; Baslow, 2007 p. 418; Nordengen, et al., 2015; Urenjak, et al., 1993) and plays several roles in in the developing and adult brain. Notably, it is involved in the myelination of postnatal brain and myelin turn-over in adults (Chakraborthy, et al., 2001; Namboodiri, et al., 2006). NAA is also a precursor for NAAG and together they have act as neurotransmitters for the axon-glial signaling of neurons with oligodendrocytes and astrocytes, respectively (Baslow, 2010). A biochemical coupling between NAA synthesis and energy metabolism in neural mitochondria (Clark, 1998) further suggests
NAA as a biomarker for neuronal health and viability (Moffett, et al., 2007), although elevated levels has been found in Canavan’s disease (Wittsack, et al., 1996). The conflicting observations reported for NAA in the aging brain could be obscured by the numerous functions and varying concentrations of NAA across the brain (Moffett, et al., 2006; Moffett, et al., 1995). The process of aging has also been found to vary depending on the brain region, where glutamate changes have been observed in the striatum but not in the cerebellum or the pons (Zahr, et al., 2013). Differences have also been shown even within similar brain structures such as in the corona radiata and the mesial motor cortex (Kaiser, et al., 2005) or the posterior cingulate cortex and the hippocampus (Reyngoudt, et al., 2012). Nevertheless, a large variability is present among the age-related variations reported in the MRS literature due to the use of different subject inclusion criteria, regions of interests and quantification methods.

3.1.2.2. Preclinical studies and the Dark-Agouti rat model

An important challenge for the study of human brain aging is the inter-subject variability and inhomogeneous biological background among the elderly population, making it difficult to determine a “control” group for which “healthy” aging could be acknowledged. Furthermore, longitudinal studies for a fixed human cohort aiming at following the metabolic changes from the beginning of adulthood (~20 years old) up to old age (~80 years old) are not feasible. To overcome these limitations, aged rodents and in particular rats, provide a popular model to examine the neurobiological basis of aging. Rats have a lifespan of 3 years (Smith, et al., 2010) and they have been studied extensively during the postnatal brain development period and also as aging models in behavioral and physiological studies (Yeoman, et al., 2012; Watson, et al., 2006). There are several neurological processes that are common for humans and rats which permit to establish a chronological correspondence between the brain maturation timelines of both species. Among the shared traits, humans and rats have a relatively immature CNS at birth and extensive postnatal neurological development takes place afterwards (Watson, et al., 2006). Based on the number of synapses with respect to the adult brain, the appearance of neurological electrical activity characteristic of the active and quiet sleep and the development of the glutamate decarboxylase and choline acetyltransferase enzymes, it has been proposed (Romijn, et al., 1991) that the approximate equivalent between a human newborn brain is that of a rat pup at 12 to 13 postnatal days (PND). Mature, aerobic metabolism is reached at 1 year of age in humans (Bentourkia, et al., 1998) and at PND 21 in rats (Booth, et al., 1980). Adult brain size and weight in the human is achieved at ages of 9 and 11 years old, respectively; whereas in the rat it happens after puberty at 60 PND (Dobbing, et al., 1971). The developmental scheme is similar across mammalian species although relative brain volumes and mass vary, notably for the neocortex and visual system which are larger in the human while the olfactory system is much larger in rodents (Rice, et al., 2000), which in addition, it also presents
neurogenesis throughout the life of the rat (Bayer, et al., 1982; Bayer, 1982; Altman, et al., 1990). In both species, the prefrontal cortex is the latest structure to reach maturity, achieving complete synaptogenesis and myelination at approximately 17 to 25 years of age in humans (Giedd, et al., 1999; Sowell, et al., 2001a; Sowell, et al., 2001b) and at 90 PND in rats (Juraska, et al., 2004). In the context of the aging rat, a decline in learning as well as motor and cerebellar function has been found (Villarreal, et al., 2004; Weiss, et al., 1991; Rapp, et al., 1987; Barnes, 1979; van der Staay, et al., 1993; Luu, et al., 2008). Studies done in the cerebellum also showed a reduction in the total number of synapses and Purkinje cells in 26-months old Sprague-Dawley rats (Rogers, et al., 1984; Chen, et al., 1999). In the hippocampus, impairments in synaptic plasticity including deficits in the induction and maintenance of long-term potentiation and long-term depression observed during aging may have a detrimental effect on the encoding and preservation of memories (Rosenzweig, et al., 2003; Thibault, et al., 2007). Neuroinflammation during aging was also studied in the hippocampus based in immunohistochemical methods, showing an activation of astrocytes and microglia in 22-months old Wistar rats accompanied by higher levels of neuronal debris and a reduced number of neurons with respect to 3 months old rats in the CA1 region (Cerbai, et al., 2012).

Previous MR spectroscopy studies have been done looking at the early developmental stages in the rat brain (Tkac, et al., 2003) and comparisons have also been done between young (5 months old) and elderly (24 months old) Brown-Norway rats but these studies were focused on the mitochondrial oxidative cycle under dietary caloric restrictions (Lin, et al., 2014). Recently, a long longitudinal study done in mice, covering the ages of 3 to 24 was done (Duarte, et al., 2014). However, no corrections for relaxation times were applied.

The Dark-Agouti (DA) rat strain was chosen for this longitudinal aging study because of its moderate and stable weight during adulthood (~350 g) without dietary restrictions which is a practical requirement for in vivo MRS experiments due to the size of our hardware (rat holder and gradient bore). To avoid confounding physiological changes due to estrogens (Li, et al., 2005), only male rats were included in this study. Among others, DA rats have been previously used for the study of MS-like neuroinflammation in the spinal cord using MRI and PET (Abourbeh, et al., 2012; Birdsall Abrams, et al., 2007). To our knowledge no MRS study was done to date using this rat strain.

3.1.3. Objectives

The main goal of this study was to investigate the metabolic alterations occurring during aging in the DA rat brain and to validate this strain as an aging model. $^1$H MRS data were collected from rats up to 22 months old and from four different brain regions. In order to achieve an absolute quantification of a maximum number of metabolites, $T_1$ and $T_2$ relaxation times of metabolites and tissue water were measured. Although numerous studies of relaxation times have been completed in human and animal models using MRI (Bottomley, et al., 1984; Wright, et al., 2008)
or MRS (Ganji, et al., 2012; Li, et al., 2012; de Graaf, et al., 2006; Cudalbu, et al., 2009), no such studies had been previously done at 17.2 T \textit{in vivo}. Therefore, $T_1$ and $T_2$ relaxation times of singlet and J-coupled metabolites and 4 groups of macromolecular resonances were measured, at different time points, as relaxation times have been observed to change during evolving pathological (Ongür, et al., 2010; Lei, et al., 2003) or physiological states such as aging (Kreis, et al., 1993; Marjanska, et al., 2013; Bottomley, et al., 1984). The relaxation times at 17.2 T were compared with experimental values reported at lower magnetic fields (Cudalbu, et al., 2009; de Graaf, et al., 2006; Xin, et al., 2008; Xin, et al., 2013) to evaluate their magnetic field dependency \textit{in vivo}.

3.2. MATERIALS AND METHODS

3.2.1. Management of the DA rats and scheduling of the MRS acquisition

\textit{In vivo} experiments were performed on 18 DA male rats. Twelve rats were received at the age of 4 weeks (220 ± 20 g) and 6 other rats were acquired at 8 months of age (326 ± 28 g) (Janvier Labs, Le Genest-Saint-Isle, France). A first cohort of young rats (C0, $n = 6$) was dedicated to the measurement of water $T_1$ relaxation times using $T_1$ mapping and a volume coil. No subsequent $T_1$ mapping experiments were done at older ages due to the small diameter of the volume coil which could cause suffocation of older rats. MRS data were acquired from the second cohort of young rats (C1, $n = 6$) and the cohort of 8-month old rats (C2, $n = 6$). $^1$H MRS data was acquired as shown in table 3.1. Due to time limitations, $T_1$ and $T_2$ relaxation measurements were not measured for the last time-points of each cohort C1 and C2 (12 and 18 months, respectively). As illustrated by figure 3.1, the weight of the C1 and C2 rats were regularly monitored. Weight perturbations were observed after each MRS session, probably due to stress and anesthesia after-effects. Metabolic profiles (and metabolite-nulled spectra) were acquired from 4 different VOIs. Relaxation times were acquired from a single region. Further details are given in the data acquisition section.
### Table 3.1. Experimental design for $^1$H MRS acquisition

<table>
<thead>
<tr>
<th>Cohort</th>
<th>Measurement</th>
<th>Age (Month)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C0</td>
<td>T$_1$ Maps</td>
<td>1 - - - - - -</td>
</tr>
<tr>
<td>C1</td>
<td>Metabolic Profiles</td>
<td>1 4 7 - - 12 - -</td>
</tr>
<tr>
<td></td>
<td>Relaxation Times</td>
<td>1 4 7 - - - - -</td>
</tr>
<tr>
<td>C2</td>
<td>Metabolic Profiles</td>
<td>- - - 8 11 - 14 18</td>
</tr>
<tr>
<td></td>
<td>Relaxation Times</td>
<td>- - - 8 11 - 14 -</td>
</tr>
</tbody>
</table>

Columns indicate the age at which MR measurements took place for each rat cohort.

### Figure 3.18. Weight monitoring during the longitudinal MRS aging study.

The weight of both C1 and C2 cohorts steadily increased over time, never exceeding 380 g. Small weight losses were occasionally observed after an exam. The weight loss observed for C2 between weeks 50 to 52 was caused by two rats which suffered severe dehydration (40% weight loss) due to their water bottle being plugged. MRS data from these rats was acquired at week 50 and both animals regained their normal weight within four weeks.

#### 3.2.2. Experimental set-up

Animals were anesthetized during the experiments with 1.0-1.5% isoflurane in pure O$_2$. Respiration rate was monitored (40–60 breaths/min) and body temperature was held constant (37.5 ± 0.5° C) using a warm-water circuit for the whole duration of the experiment. The head was stereotaxically restrained by a bite bar and ear pins. The study protocol was approved by the Committee on the Ethics of Animal Experiments of the Commissariat à l’Energie Atomique (CETEA, Permit Number: ID 12-058).

The study was performed on a horizontal 17.2 T MRI scanner (Biospec, Bruker BioSpin, Ettlingen, Germany) equipped with an actively shielded gradient system capable of delivering gradients up to 1 T/m. MRS data acquisitions were performed using a 20-mm-diameter single-
loop surface coil as transceiver while water T\textsubscript{1} maps were acquired with a Bruker 38-mm-diameter birdcage volume coil (figure 3.2).

![Figure 3.2. 17.2 T Bruker Biospec scanner and surface and volume coils used during the \textsuperscript{1}H MRS study.](image)

3.2.3. MRI data acquisition

Water T\textsubscript{1} maps were obtained using a 3D IR-FGE sequence (TE/TR\textsubscript{1}/TR\textsubscript{2} = 3.02/10/10000 ms, 6 segments, flip angle = 5°, resolution: 125x125x1000 µm\textsuperscript{3}, matrix size: 256x96x14) with 60 inversion times (90 to 9530 ms) using a non-adiabatic hermit excitation pulse (1 ms, 5.4 kHz bandwidth) and an HS8 AFP inversion pulse (4 ms, 5 kHz bandwidth). For comparison, the T\textsubscript{1} relaxation time of physiological water (Proamp, Aguettant) was also measured at 37 °C.

For positioning the spectroscopic voxel, reference images were acquired using an axial rapid acquisition with relaxation enhancement (RARE) sequence covering the entire brain (TE/TR = 20/3000 ms, slice thickness: 0.5 mm, 24 slices, in-plane resolution: 180 x 180 µm\textsuperscript{2}).

3.2.4. MRS data acquisition

\textsuperscript{1}H spectra were acquired using a LASER sequence (TE/TR = 16.5/5000 ms, 128 averages, 2048 complex points) consisting of a non-selective excitation pulse (Hermit, 0.4 ms duration, 13.5 kHz bandwidth) followed by 3 pairs of slice-selective HS8 AFP inversion pulses (hyperbolic secant modulation, n = 8, 1 ms duration, 20 kHz bandwidth) (Garwood, et al., 2011). The resulting chemical shift displacement artifact was a spatial shift of 13% between water and lipids at 1.3 ppm.
Four different volumes of interest (VOI) were explored, covering the thalamus (Thal), cingulate cortex (CC), caudate putamen (CP) and a region referred to as the “main” region (Main) composed mostly of cerebral cortex with minor contributions from the corpus callosum and hippocampus. VOIs sizes and positioning are given in figure 3.3.

First- and second-order shims were adjusted to optimize local $B_0$ field homogeneity. For our Main region, FASTMAP was used on a cubic volume (27 $\mu$L) centered on the VOI followed by the application of the Bruker MAPSHIM routine on the VOI leading to typical water linewidths of $23 \pm 3$ Hz. For the remaining 3 regions only FASTMAP was used with typical water linewidths of $23 \pm 2$ (Thal), $20 \pm 2$ (CP) and $21 \pm 3$ (CC). Further $B_0$-shimming details are given in section 2.1.3. Non-suppressed water spectra were acquired for water referencing (Kreis, et al., 1993) and eddy current correction.

$T_1$-weighting was introduced by incorporating a single IR module consisting of a non-selective HS8 AFP inversion pulse (2 ms, 10 kHz bandwidth) followed by an inversion delay (TI) prior to the LASER localization scheme. In total, 9 $T_1$-weighted IR spectra were acquired ($TI = 109, 264, 500, 750, 1000, 1250, 1500, 2000, 3000$ ms). To conserve the same steady-state magnetization, all 9 IR spectra were acquired with a recovery time of 5000 ms.

$T_2$-weighted spectra were acquired by increasing the echo-time from 16.5 to 200 ms leading to 12 different TEs ($16.5, 24, 35, 50, 60, 80, 100, 120, 140, 160, 180$ and $200$ ms). The delays were equally and symmetrically split on the 3 pairs of AFP pulses with the exception of the TE of 16.5 ms. $T_1$ and $T_2$-weighted spectra were only acquired in the Main VOI.

To parameterize the MM baseline, metabolite-nulled spectra were acquired for each VOI using a double inversion recovery (DIR) module (HS8, $TI_1/TI_2 = 2600/600$ ms) incorporated prior to the LASER sequence ($TE/TR = 16.5/5000$ ms, 128 averages, 2048 complex points).

Water suppression was achieved using numerically optimized WET schemes as presented in section 2.2.2.

To check for signal stability, localized water spectra were acquired prior to each MRS acquisition. A small decrease of the water signal was observed during the first hour of the experiment affecting the sensitivity up to 1.5%. The average water frequency drift was 5.4 Hz per hour.
3.2.5. MRS data analysis

Spectral preprocessing consisted in zero-filling to 4096 data points, Fourier transformation and zero-order phase correction. After removal of the residual water signal using the HLSVD algorithm (Pijnappel, et al., 1992) implemented in the jMRUI software (available online at http://www.mrui.uab.es/mrui), spectra were analyzed using LCModel 6.2.

For each echo-time, a basis-set of metabolite spectra was simulated as described in section 2.4.1. For the LCModel analysis, the IR spectra were phased such that the tCr resonance at 3.03 ppm was positive. Consequently, the basis-set used for analyzing T1-weighted data included inverted spectra to account for the metabolites presenting an inverted signal with respect to the methyl resonance of tCr.

The following 21 metabolites were considered at all echo-times: alanine (Ala), aspartate (Asp), ascorbate (Asc), creatine (Cr), Ethanolamine (EA), γ-amino-butyric acid (GABA), glucose (Glc), glutamate (Glu), glutamine (Gln), glycine (Gly), glutathione (GSH), glycerophosphoryl-choline (GPC), phosphoryl-choline (PCh), myo-inositol (Ins), lactate (Lac), N-acetyl-aspartate (NAA), N-acetyl-aspartyl-glutamate (NAAG), phosphocreatine (PCr), phosphoryl-ethanolamine (PE), serine (Ser) and taurine (Tau). Due to their differences in T1 values, the methyl (CH3 at 3.03
ppm) and methylene (CH\textsubscript{2} at 3.93 and 3.91 ppm) moieties of Cr and PCr were separately accounted for at all echo times. Likewise, the three NAA moieties (CH\textsubscript{3} at 2.01 ppm, CH\textsubscript{2} at 2.49 and 2.67 ppm and CH at 4.38 ppm) were also fitted independently. The sums Cr+PCr, PCh+GPC and NAA(CH\textsubscript{3})+NAAG were addressed as tCr, tCho and tNAA respectively. Acetate and scyllitolositol were not considered due to their low concentration in the brain (below 0.2 and 0.1 mM concentrations, respectively), as measured with high-resolution liquid-phase \textsuperscript{1}H NMR from brain extracts (de Graaf, et al., 2011).

To estimate T\textsubscript{1} and T\textsubscript{2} relaxation times, 2-parameter fits consisting of mono-exponential functions were used to fit the equilibrium magnetization and T\textsubscript{1} or T\textsubscript{2} relaxation times respectively using a non-weighted Levenberg-Marquardt algorithm. Both fits were applied individually on each animal data set. Cramer-Rao Lower Bounds (CRLB) were used as error estimator (section 2.4.4) and all data points presenting values higher than 25% were discarded. The coefficient of determination (R\textsuperscript{2}) was used to evaluate the quality of the fits. Metabolite resonance intensities obtained from LCModel were converted to absolute concentrations using water signal as an internal reference of concentration (section 2.5.1).

3.2.6. MRI data analysis and segmentation

T\textsubscript{1} maps were generated based on the IR-FGE data acquired from one-month old rats (section 2.6, figure 2.15) and tissue segmentation was done manually (section 2.5.1). For the determination of the T\textsubscript{1} values of each VOI, outlier pixels showing elevated values (above > 2300 ms) were considered to correspond to CSF and were thus removed from the VOI. The thalamus regions contained the largest number of CSF pixels, which accounted for 5% of the total thalamus VOI. Table 3.2 summarizes the T\textsubscript{1} measurement results. WM contributions were estimated as follows: 14 ± 10 % (Main), 60 ± 14 % (Thal), 34 ± 12 % (CP) and none for the CC. The assumed water content of GM and WM were 82% and 73% respectively (Lentner, 1981). The water content of GM, WM and CSF as well as the GM and WM fractional contributions for each region were assumed to be constant at all ages.

<table>
<thead>
<tr>
<th>Region</th>
<th>T\textsubscript{1} (ms)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Main VOI</td>
<td>1839 ± 54</td>
</tr>
<tr>
<td>Gray Matter</td>
<td>1877 ± 77</td>
</tr>
<tr>
<td>White Matter</td>
<td>1670 ± 34</td>
</tr>
<tr>
<td>Thalamus</td>
<td>1610 ± 55</td>
</tr>
<tr>
<td>Caudate Putamen</td>
<td>1732 ± 52</td>
</tr>
<tr>
<td>Cingulate Cortex</td>
<td>1835 ± 45</td>
</tr>
<tr>
<td>Cerebrospinal Fluid</td>
<td>2842 ± 318</td>
</tr>
<tr>
<td>Physiological Water (37°)</td>
<td>2966 ± 60</td>
</tr>
</tbody>
</table>

Table 3.2. T\textsubscript{1} relaxation times measured in the rat brain at 17.2 T using an IR-FGE sequence
3.2.7. Macromolecule baseline parameterization

The MM baseline parameterization was estimated based on a mean metabolite-nulled spectrum obtained from spectra acquired from the first scan of the C1 cohort. The method and the results are detailed in section 2.4.3.

3.2.8. Statistical analysis

A one-way ANOVA was used to evaluate the metabolic differences between the four VOIs for the C1 cohort at 1 month of age. The effect of age and the number of experienced examinations was evaluated for the metabolite profiles of the 4 VOIs and the relaxation times using a linear regression model. Linear-regression with mixed effects analysis considering age and the number of experienced examinations as simultaneous predictors were used to determine the contribution of each variable to the metabolite concentrations and relaxation times changes. To be able to join the two C1 and C2 cohorts within a single statistical analysis, the “rat” variable was considered as a random effect. The model corresponded to:

\[ Y = \beta_0 + \beta_1 \times \text{Age} + \beta_2 \times \text{Exams} + e \]  

(3.1)

where \( e \) is the residual error, \( Y \) is the observed data corrected for rat effects. The age and number of exams were considered to be independent factors. The analysis was done using R, version 3.1.2 (R Core Team, 2014) with linear mixed-effects model package lme4, version 1.1-7 (Bates, et al., 2014). No corrections were applied to reduce type I errors (such as the Bonferroni correction). Statistically significant threshold was set as \( P < 0.05 \).

3.3. RESULTS: qMRS IN YOUNG RATS

Figure 3.4 shows a representative spectrum and its LCModel fit, the individual metabolites contributions as well as the MM fit, baseline and residual signal. Overall, the measured data showed high SNR and good quality as no artifacts were observed. The parameterized MM fit correctly accounted for the broad resonances resulting in a rectilinear baseline on the fitted spectra, despite the \( T_1 \) or \( T_2 \) weighting introduced by the TE or TI parameters. The water suppression scheme was quite effective resulting in a water residual of similar magnitude as the NAA(CH₃) resonance and no baseline distortions due to eddy currents were observed. Nevertheless, the baseline between 4.5 and 4.3 ppm presented minor distortions, which seldom affected the quantification of the MM4 group and the NAA(CH) moiety resonating at 4.38 ppm.
Figure 3.4. Spectral decomposition of a $^1$H spectrum at 17.2 T.
LCModel spectral decomposition of a representative spectrum acquired from the Main VOI at TE = 16.5 ms. 21 different metabolites were considered in the basis-set in addition to the 4 MM components, however EA was seldom detected. The simulated MM baseline, the residual signal and regularized spline baseline are shown. No filtering was applied and all spectral elements shown are scaled identically. $^a$The CH$_3$ and the CH$_2$ moieties of tCr were considered separately in the basis-sets. $^b$The CH$_3$, CH$_2$ and CH moieties of NAA were considered separately in the basis sets.
3.3.1. T₁ and T₂ relaxation in young rats

Figure 3.5 shows a set of T₁-weighted spectra (left) and their LCModel fit (right). The fits were in good agreement at all inversion times. CRLBs lower than 5% were consistently found for the most prominent resonances (Gln, Glu, Ins, NAA(CH₃), NAA(CH₂), tCr(CH₃), tCr(CH₂), MMs) while GPC, PCh, GABA, Asp, Glc and Tau showed values from 6% to 10%. Overall, the CRLBs were fairly stable at all TI with the exception of the inversion times between 0.75 and 1.25 s where the metabolites were zero-crossing, resulting in CRLB above 20%.

Figure 3.5. T₁-weighted spectra acquired in the rat brain at 17.2 T
Set of T₁-weighted spectra (TE/TR = 16.5/5000 ms) from a 1-month old rat obtained with the IR method (left) and their LCModel fit (right) with inversion times ranging from 109 (bottom) to 3000 ms and a non-inverted spectrum (top). Water removal was performed using HLSVD. No filtering was applied. The most notable features were the macromolecules being the first to invert at TI = 500 ms, followed by tCr(CH₂) at 750 ms. Most metabolites were inverted between 1000 and 1250 ms. Tau and the NAA(CH) moiety were the last to invert at 1500 ms.

Figure 3.6 shows a set of T₂-weighted spectra and their corresponding LCModel fits. Figure 3.7 shows the normalized apparent concentration as a function of TE for the singlet tCr(CH₃), the major J-coupled metabolites (GABA, Glu, Gln, Ins, NAA(CH₂), Tau) and the MM1 group from a single rat. As demonstrated by both the quality of the LCModel fits at all TE (fig. 3.6) and the R² values obtained for the determination of the T₂ times (fig. 3.7), the J-modulation of most metabolites was adequately accounted for by our simulated model spectra. In particular, one can appreciate the accurate adjustment of the NAA aspartate moiety at echo-times 35 to 80 ms, the
strongly coupled spin system reaching a minimal signal intensity at 35 ms and its phase being clearly inverted (for the resonance at 2.67 ppm) at TE = 50 and 60 ms.

Figure 3.6. T₂-weighted spectra acquired in the rat brain at 17.2 T
Set of T₂-weighted spectra (left) and their corresponding LCModel fit (right) from a 1-month old rat. TE varied from 16.5 to 200 ms. Water removal was performed using HLSVD. No filtering was applied.

Figure 3.7. Normalized apparent concentrations with respect to echo time.
The normalized apparent concentrations (red) is shown as a function of TE for 6 J-coupled metabolites, tCr(CH₃) and MM1 from a single animal. Only data points with CRLB below 25% are shown. For each linear regression (black), the slope is equal to the estimated T₂ relaxation time. The corresponding T₂ values and coefficients of determination ($R^2$) were: GABA: 80 ms, 0.964; Glu: 96 ms, 0.931; Gln: 46 ms, 0.961; MM1: 18 ms, 0.999; Ins: 170 ms, 0.898; NAA(CH₂): 173 ms, 0.939; Tau: 124 ms, 0.966; tCr(CH₃): 138 ms, 0.979.
Likewise, the J-modulation of Ins was properly accounted for, notably the phase-inverted 3.52 ppm resonance at TE = 80 and 100 ms. At TE values 50 to 80 ms, strong J-modulation led to a major signal attenuation of Glu and Tau resonances which are visibly refocused for TE = 100-140 ms due to the J-induced rephasing of the signal intensity (TE ~ 1/J).

The CRLBs reported by LCModel were below 5% for tCr(CH$_3$), tCho and the three moieties of NAA even at long echo times. Ins, Glu, Tau and Gln showed values below 5% at short echo times (< 60 ms) but were considerably increased due to the signal loss induced by J-coupling effects, notably at TE of 60, 80 and 140 ms. The MM resonance groups also had low CRLB values (<5%) up to TE = 60 ms and reached 10% at TE = 100 ms. Above this TE, the macromolecule signals were no longer observed and so they were removed from the LCModel analysis. Overall, the high spectral resolution and signal-to-noise ratio accessible at 17.2 T allowed us to properly analyze 18 distinct metabolites with the current methods up to TE = 50 with a maximum CRLB of 22.2%.

The estimated T$_1$ and T$_2$ relaxation times for young adult rats are shown in Table 3.3 with their corresponding coefficients of determination ($R^2$). Only relaxation times with a mean $R^2$ value higher than 0.80 are shown. Figure 3.8 shows the distribution of T$_1$ and T$_2$ times and their standard deviations in decreasing order. The mean T$_1$ and T$_2$ metabolite relaxation times were 1721 ± 237 ms and 148 ± 53 ms respectively, which were very close to the relaxation times of tCr(CH$_3$).

3.3.2. Neurochemical profile in young rats

Following the quantification method described in section 2.5.1, the concentrations of 21 brain metabolites as well as the apparent $^1$H concentration of the 4 macromolecule resonances groups were determined from the 16 ms neurochemical profiles.

The absolute concentrations from the Main VOI are summarized in Table 3.3. Based on the bi-exponential fit of the T$_2$-weighted unsuppressed water spectra, the fractional CSF content for each rat was determined and was corrected individually for each animal, giving a mean $f_{CSF} = 4 \pm 2 \%$. The mean T$_2$ time of brain tissue was 24 ± 2 ms whereas the T$_2$ value of CSF was 410 ± 490 ms. Although the precision of the CSF T$_2$ time was poor, the TE of our neurochemical profile was short (16.5 ms) and therefore only negligible errors were expected to be introduced in the quantification. In average, the apparent water concentration in the MAIN VOI was 22.5 ± 0.5 mol/L.

Since T$_2$-weighted water spectra were not acquired for the Thal, CC and CP regions, their CSF contributions were estimated from the volume fraction in the T$_1$ maps that presented T$_1$ values above 2300 ms (section 3.3.6). The found values showed fractional CSF content of 5 % for Thal, 0.5 % for CC and no CSF content for CP. The calculated relaxation times were applied for the relaxation corrections of the 4 VOIs. The apparent water concentration for Thal, CP and CC VOIs were 21.9 ± 0.3, 21.4 ± 0.2 and 22.2 ± 0.1 mol/L respectively. When a T$_1$ or T$_2$ time was
not determined with sufficient precision ($R^2<0.80$) for a given metabolite, the average $T_1$ or $T_2$ metabolite relaxation time was used instead. Macromolecule MM proton concentrations were determined by referencing to the area under the curve of $tCr(CH_3)$ for each individual spectrum.

Figure 3.8. $T_1$ and $T_2$ relaxation times distribution in the young rat brain at 17.2 T Metabolite and macromolecule (A) $T_1$ and (B) $T_2$ relaxation times obtained in vivo in the rat brain at 17.2 T with coefficients of determination higher than 0.80. Mean values from six 1 month-old rats are shown in decreasing order along with their standard deviation. The mean $T_1$ and $T_2$ relaxation times of metabolites were $1721 \pm 237$ ms and $148 \pm 53$ ms respectively, closely matching those of $tCr(CH_3)$. 
### Table 3.3. Summary of absolute concentrations and relaxation times in young rats at 17.2 T

Metabolites and macromolecules relaxation times and absolute concentrations estimated at 17.2 T from 1-month old rats *in vivo* (mean ± SD) acquired from the Main VOI. To evaluate the quality of the T<sub>1</sub> and T<sub>2</sub> fits, the coefficient of determination (R<sup>2</sup>) are presented. Concentrations (mean ± SD) from the 16 ms neurochemical profiles were calculated using equation (2.6). The mean CRLBs are shown. Macromolecule proton concentrations were referenced to the area under the curve of Cr. The number of times each metabolite was detected out of 12 scans is shown.

<table>
<thead>
<tr>
<th>Metabolites</th>
<th>T&lt;sub&gt;1&lt;/sub&gt; (ms)</th>
<th>R&lt;sup&gt;2&lt;/sup&gt;</th>
<th>T&lt;sub&gt;2&lt;/sub&gt; (ms)</th>
<th>R&lt;sup&gt;2&lt;/sup&gt;</th>
<th>Concentration (mmol/L)</th>
<th>CRLB</th>
<th>Measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ala&lt;sup&gt;a&lt;/sup&gt;</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1.0 ± 0.2</td>
<td>14</td>
<td>12</td>
</tr>
<tr>
<td>Asp&lt;sup&gt;b&lt;/sup&gt;</td>
<td>2142 ± 170</td>
<td>0.897 ± 0.040</td>
<td>-</td>
<td>-</td>
<td>2.2 ± 0.4</td>
<td>8</td>
<td>12</td>
</tr>
<tr>
<td>Asp</td>
<td>1653 ± 48</td>
<td>0.978 ± 0.012</td>
<td>77 ± 14</td>
<td>0.906 ± 0.038</td>
<td>2.9 ± 0.3</td>
<td>6</td>
<td>12</td>
</tr>
<tr>
<td>Cr&lt;sup&gt;c&lt;/sup&gt;</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.1 ± 0.4</td>
<td>4</td>
<td>12</td>
</tr>
<tr>
<td>EA</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>0.6 ± 0.1</td>
<td>17</td>
<td>2</td>
</tr>
<tr>
<td>GABA</td>
<td>1643 ± 36</td>
<td>0.971 ± 0.017</td>
<td>78 ± 20</td>
<td>0.960 ± 0.012</td>
<td>2.0 ± 0.3</td>
<td>6</td>
<td>12</td>
</tr>
<tr>
<td>Glc</td>
<td>1780 ± 104</td>
<td>0.956 ± 0.035</td>
<td>170 ± 26</td>
<td>0.872 ± 0.044</td>
<td>2.2 ± 0.6</td>
<td>6</td>
<td>12</td>
</tr>
<tr>
<td>Gln</td>
<td>1799 ± 61</td>
<td>0.987 ± 0.008</td>
<td>58 ± 10</td>
<td>0.945 ± 0.022</td>
<td>4.6 ± 0.4</td>
<td>3</td>
<td>12</td>
</tr>
<tr>
<td>Glu</td>
<td>1699 ± 48</td>
<td>0.993 ± 0.004</td>
<td>79 ± 23</td>
<td>0.929 ± 0.030</td>
<td>9.6 ± 0.9</td>
<td>2</td>
<td>12</td>
</tr>
<tr>
<td>Gly&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1943 ± 165</td>
<td>0.883 ± 0.075</td>
<td>-</td>
<td>-</td>
<td>0.2 ± 0.04</td>
<td>16</td>
<td>6</td>
</tr>
<tr>
<td>GPC</td>
<td>1412 ± 75</td>
<td>0.953 ± 0.033</td>
<td>220 ± 32</td>
<td>0.930 ± 0.029</td>
<td>0.6 ± 0.1</td>
<td>5</td>
<td>12</td>
</tr>
<tr>
<td>GSH</td>
<td>1658 ± 341</td>
<td>0.905 ± 0.059</td>
<td>105 ± 39</td>
<td>0.839 ± 0.066</td>
<td>1.0 ± 0.1</td>
<td>10</td>
<td>12</td>
</tr>
<tr>
<td>Ins</td>
<td>1608 ± 34</td>
<td>0.995 ± 0.092</td>
<td>170 ± 15</td>
<td>0.948 ± 0.025</td>
<td>5.9 ± 0.6</td>
<td>2</td>
<td>12</td>
</tr>
<tr>
<td>Lac</td>
<td>1766 ± 184</td>
<td>0.966 ± 0.026</td>
<td>121 ± 26</td>
<td>0.827 ± 0.092</td>
<td>1.6 ± 0.3</td>
<td>6</td>
<td>12</td>
</tr>
<tr>
<td>NAA(CH&lt;sub&gt;3&lt;/sub&gt;)</td>
<td>1810 ± 64</td>
<td>0.993 ± 0.004</td>
<td>226 ± 26</td>
<td>0.938 ± 0.042</td>
<td>9.2 ± 0.7</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>NAA(CH&lt;sub&gt;2&lt;/sub&gt;)</td>
<td>1527 ± 48</td>
<td>0.981 ± 0.009</td>
<td>172 ± 14</td>
<td>0.939 ± 0.022</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>NAA(CH)</td>
<td>1967 ± 188</td>
<td>0.885 ± 0.053</td>
<td>182 ± 22</td>
<td>0.876 ± 0.043</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>NAAG</td>
<td>1510 ± 55</td>
<td>0.939 ± 0.034</td>
<td>-</td>
<td>-</td>
<td>0.8 ± 0.1</td>
<td>7</td>
<td>12</td>
</tr>
<tr>
<td>PCh</td>
<td>1413 ± 71</td>
<td>0.950 ± 0.033</td>
<td>204 ± 31</td>
<td>0.930 ± 0.027</td>
<td>0.3 ± 0.1</td>
<td>7</td>
<td>12</td>
</tr>
<tr>
<td>PCr&lt;sup&gt;c&lt;/sup&gt;</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>4.9 ± 0.3</td>
<td>3</td>
<td>12</td>
</tr>
<tr>
<td>PE&lt;sup&gt;b&lt;/sup&gt;</td>
<td>1868 ± 109</td>
<td>0.975 ± 0.007</td>
<td>-</td>
<td>-</td>
<td>1.7 ± 0.2</td>
<td>9</td>
<td>12</td>
</tr>
<tr>
<td>Ser&lt;sup&gt;a&lt;/sup&gt;</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>-</td>
<td>1.0 ± 0.2</td>
<td>20</td>
<td>5</td>
</tr>
<tr>
<td>Tau</td>
<td>2211 ± 94</td>
<td>0.990 ± 0.003</td>
<td>118 ± 5</td>
<td>0.950 ± 0.017</td>
<td>5.5 ± 0.5</td>
<td>3</td>
<td>12</td>
</tr>
<tr>
<td>tCho</td>
<td>1408 ± 67</td>
<td>0.954 ± 0.033</td>
<td>212 ± 21</td>
<td>0.919 ± 0.049</td>
<td>0.9 ± 0.2</td>
<td>4</td>
<td>12</td>
</tr>
<tr>
<td>tCr(CH&lt;sub&gt;3&lt;/sub&gt;)</td>
<td>1753 ± 60</td>
<td>0.995 ± 0.002</td>
<td>141 ± 7</td>
<td>0.981 ± 0.009</td>
<td>8.9 ± 0.6</td>
<td>1</td>
<td>12</td>
</tr>
<tr>
<td>tCr(CH&lt;sub&gt;2&lt;/sub&gt;)</td>
<td>1203 ± 45</td>
<td>0.946 ± 0.010</td>
<td>132 ± 10</td>
<td>0.927 ± 0.023</td>
<td>-</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>tNAA&lt;sup&gt;c&lt;/sup&gt;</td>
<td>1786 ± 55</td>
<td>0.997 ± 0.002</td>
<td>218 ± 17</td>
<td>0.935 ± 0.025</td>
<td>9.9 ± 0.7</td>
<td>1</td>
<td>12</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>MM groups</th>
<th>T&lt;sub&gt;1&lt;/sub&gt; (ms)</th>
<th>R&lt;sup&gt;2&lt;/sup&gt;</th>
<th>T&lt;sub&gt;2&lt;/sub&gt; (ms)</th>
<th>R&lt;sup&gt;2&lt;/sup&gt;</th>
<th>Concentration (mmol/L)</th>
<th>CRLB</th>
<th>Measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>MM1</td>
<td>748 ± 14</td>
<td>0.962 ± 0.013</td>
<td>20 ± 2</td>
<td>0.994 ± 0.003</td>
<td>170.1 ± 14.1</td>
<td>2</td>
<td>12</td>
</tr>
<tr>
<td>MM2</td>
<td>652 ± 41</td>
<td>0.977 ± 0.020</td>
<td>28 ± 3</td>
<td>0.980 ± 0.008</td>
<td>114.4 ± 8.6</td>
<td>3</td>
<td>12</td>
</tr>
<tr>
<td>MM3</td>
<td>574 ± 22</td>
<td>0.971 ± 0.012</td>
<td>55 ± 5</td>
<td>0.931 ± 0.055</td>
<td>26.6 ± 2.8</td>
<td>5</td>
<td>12</td>
</tr>
<tr>
<td>MM4</td>
<td>792 ± 39</td>
<td>0.948 ± 0.021</td>
<td>43 ± 6</td>
<td>0.959 ± 0.013</td>
<td>82.5 ± 5.8</td>
<td>4</td>
<td>12</td>
</tr>
</tbody>
</table>

<sup>a</sup> Mean metabolite T<sub>1</sub> and T<sub>2</sub> were considered for quantification.

<sup>b</sup> Mean T<sub>2</sub> was considered for quantification.

<sup>c</sup> T<sub>1</sub> and T<sub>2</sub> of the methyl moiety were considered for quantification.
3.3.3. Regional variability

Measured spectra and their LCModel fits from the 4 studied regions are shown in Figure 3.9. The metabolite concentrations of the C1 cohort at the first time-point of acquisition are shown in Table 3.4. Statistical differences were determined using a one-way ANOVA test. The results for the main metabolites are shown in figure 3.10.

Numerous metabolite concentration differences were found between the Main VOI and the Thal, CP and CC regions. Among these variations, one can notice that tCho and Ins were more concentrated in the thalamus, which is consistent with the elevated WM percentage measured from the $T_1$ maps. On the contrary, Ins, tNAA and the Glu/Gln ratio were lowest in the striatum (CP) while it presented the highest Tau levels. The CC region exhibited the highest Glu level and PCr/Cr ratio. The Main VOI showed the highest tNAA and tCr levels although their ratio was slightly above average. Macromolecule concentrations were found to be similar in the four regions although higher concentrations were found in the CC, mainly for the MM3 component ($p = 0.003$). Lactate levels were highest in the thalamus and lowest in the CC.

Figure 3.9. Neurochemical profiles from the 4 VOIs acquired from a single rat. Spectra (red) and the corresponding LCModel fits (black) are shown for the Main, thalamus, caudate putamen and cingulate cortex VOIs (figure 3.3). Minor baseline distortions were seldom observed beyond 4.2 ppm for the CP and CC spectra.
### Table 3.4. Metabolite concentrations in the 4 studied regions from the C1 rats at 1 month of age.

Metabolite and macromolecule concentrations (mM, mean ± SD) of the 4 VOIs were calculated using the $T_1$ and $T_2$ relaxation times estimated from the Main region. The same procedure was applied as for the values shown in Table 3.3 for the Main VOI concentrations.

<table>
<thead>
<tr>
<th>Metabolite</th>
<th>Main</th>
<th>Thalamus</th>
<th>Caudate P.</th>
<th>Cingulate C</th>
<th>p-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ala</td>
<td>1.0 ± 0.2</td>
<td>0.7 ± 0.2</td>
<td>1.2 ± 0.3</td>
<td>0.9 ± 0.2</td>
<td>0.001</td>
</tr>
<tr>
<td>Asc</td>
<td>2.2 ± 0.4</td>
<td>1.4 ± 0.3</td>
<td>2.5 ± 1.5</td>
<td>1.9 ± 0.4</td>
<td>0.003</td>
</tr>
<tr>
<td>Asp</td>
<td>2.9 ± 0.3</td>
<td>3.1 ± 0.4</td>
<td>2.3 ± 0.2</td>
<td>2.9 ± 0.5</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Cr/Pc</td>
<td>0.8 ± 0.1</td>
<td>0.8 ± 0.1</td>
<td>0.7 ± 0.1</td>
<td>0.6 ± 0.1</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>GABA</td>
<td>2.0 ± 0.3</td>
<td>2.9 ± 0.9</td>
<td>2.2 ± 0.3</td>
<td>2.0 ± 0.3</td>
<td>0.043</td>
</tr>
<tr>
<td>Glc</td>
<td>2.2 ± 0.6</td>
<td>2.0 ± 0.3</td>
<td>1.9 ± 0.4</td>
<td>2.1 ± 0.4</td>
<td>0.212</td>
</tr>
<tr>
<td>Glu</td>
<td>4.6 ± 0.4</td>
<td>10.4 ± 1.1</td>
<td>9.7 ± 0.7</td>
<td>11.2 ± 1.1</td>
<td>0.009</td>
</tr>
<tr>
<td>Gln</td>
<td>9.6 ± 0.9</td>
<td>5.0 ± 0.6</td>
<td>6.1 ± 0.6</td>
<td>6.2 ± 1.1</td>
<td>0.001</td>
</tr>
<tr>
<td>Gly</td>
<td>0.2 ± 0.04</td>
<td>0.3 ± 0.1</td>
<td>0.4 ± 0.1</td>
<td>0.4 ± 0.1</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Glu/Gln</td>
<td>2.1 ± 0.3</td>
<td>2.1 ± 0.3</td>
<td>1.6 ± 0.2</td>
<td>1.8 ± 0.3</td>
<td>0.001</td>
</tr>
<tr>
<td>GSH</td>
<td>1.0 ± 0.1</td>
<td>1.3 ± 0.2</td>
<td>1.4 ± 0.4</td>
<td>1.6 ± 0.3</td>
<td>0.001</td>
</tr>
<tr>
<td>Ins</td>
<td>5.9 ± 0.6</td>
<td>6.5 ± 0.9</td>
<td>4.6 ± 0.3</td>
<td>5.7 ± 0.2</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Lac</td>
<td>1.6 ± 0.3</td>
<td>1.9 ± 0.3</td>
<td>1.3 ± 0.3</td>
<td>0.9 ± 0.2</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>PE</td>
<td>1.7 ± 0.2</td>
<td>1.6 ± 0.3</td>
<td>1.6 ± 0.2</td>
<td>1.6 ± 0.1</td>
<td>0.141</td>
</tr>
<tr>
<td>Tau</td>
<td>5.5 ± 0.5</td>
<td>3.1 ± 0.9</td>
<td>9.5 ± 0.7</td>
<td>6.6 ± 0.5</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>tCho</td>
<td>0.9 ± 0.2</td>
<td>1.6 ± 0.3</td>
<td>1.1 ± 0.1</td>
<td>1.0 ± 0.2</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>tCr</td>
<td>8.9 ± 0.6</td>
<td>8.3 ± 0.9</td>
<td>7.9 ± 0.3</td>
<td>7.7 ± 0.5</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>tNAA</td>
<td>9.9 ± 0.7</td>
<td>10.1 ± 1.5</td>
<td>7.8 ± 0.4</td>
<td>9.0 ± 0.4</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>tCr/tNAA</td>
<td>0.90 ± 0.04</td>
<td>0.83 ± 0.07</td>
<td>1.01 ± 0.07</td>
<td>0.86 ± 0.06</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>MM1</td>
<td>170 ± 14</td>
<td>176 ± 17</td>
<td>171 ± 6</td>
<td>182 ± 19</td>
<td>0.141</td>
</tr>
<tr>
<td>MM2</td>
<td>114 ± 9</td>
<td>119 ± 12</td>
<td>124 ± 9</td>
<td>131 ± 12</td>
<td>0.013</td>
</tr>
<tr>
<td>MM3</td>
<td>27 ± 3</td>
<td>30 ± 3</td>
<td>31 ± 3</td>
<td>33 ± 4</td>
<td>0.003</td>
</tr>
<tr>
<td>MM4</td>
<td>82 ± 6</td>
<td>81 ± 12</td>
<td>84 ± 26</td>
<td>98 ± 14</td>
<td>0.027</td>
</tr>
</tbody>
</table>

$^a$ $T_1$ and $T_2$ of the methyl moiety were considered for quantification.

$^b$ Concentrations are expressed as $^1$H mM.
Figure 3.10. Regional variability between the 4 VOIs in 1 month-old rats. (Top) Metabolites concentrations were compared using a one-way ANOVA (Table 3.4). (Left) MM concentrations were rather stable between regions. The same T₁ and T₂ relaxation time corrections were applied for all VOIs. Statistical significance: (*) p<0.05, (**) p<0.005, (***) p<0.0005.

3.4. RESULTS: QMRS IN AGING RATS

The results obtained from the C1 and C2 cohorts at increasing ages (Table 3.1) are presented. In general, the quality of the data and the CRLBs did not change over time.

3.4.1. T₁ and T₂ relaxation in aging rats

Over the span of 14 months, relatively few T₁ and T₂ statistically significant changes with age were observed in the Main region. Figures 3.11 and 3.12 illustrate the evolution of relaxation times of a few major metabolites and brain water, showing age-related changes. Table 3.5 summarizes the results from the linear regression analysis presenting the relative changes attributed either to aging (in % per month) or to the number of scans experienced by the rats (in % per exam).
3.4.1.1. Changes with age

Except for a slight $T_1$ increase for GABA, there were no $T_1$ changes correlated with age. Similarly, only minor changes in the $T_2$ decay times were correlated with age (tCho, NAA(CH$_3$) and MM1).

3.4.1.2. Changes with the number of scans

On the contrary, the $T_1$ relaxation times of many metabolite functional groups (Asp, GABA, Gln, Glu, NAA(CH$_3$), NAAG, tCr(CH$_3$) and MM2) were shown to increase with the number of scans at an approximate rate of 2-3% per exam. Asc, Glc and Tau also seemed to increase but they were just above significance levels ($p < 0.06$). The methylene moieties of NAA and tCr did not seem to change. The observed $T_1$ changes due to the number of exams displayed a very reproducible pattern, as it can be observed in figure 3.11 for GABA, NAA(CH$_3$) or tCr(CH$_3$).

Diminutions of the $T_2$ relaxation time were also found for MM3, Tau and tCr(CH$_2$) at a rate of approximately 5-7% per exam. Brain tissue water relaxation times did not change significantly for any of the two predictors.
Figure 3.11. T₁ relaxation times variations from 1 to 14 months of age. Boxplots of T₁ relaxation times at different ages are shown for the C1 (blue) and C2 (red) cohorts for 8 metabolites and brain tissue water. Measurements were done on the Main VOI. Significant changes are indicated for each metabolite and are listed in Table 3.5. 6 individual measurements were acquired at each time point. (*) p < 0.05, (**) p < 0.005 and (***) p < 0.0005.
Figure 3.12. $T_2$ relaxation times variations from 1 to 14 months of age. Boxplots of $T_2$ relaxation times at different ages are shown for the C1 (blue) and C2 (red) cohorts for 5 metabolites moieties and brain tissue water. Measurements were done on the Main VOI. Significant changes are indicated for each metabolite and are listed in Table 3.5. 6 individual measurements were acquired at each time point. (*) $p < 0.05$ and (**) $p < 0.005$. 
<table>
<thead>
<tr>
<th></th>
<th>T1 Relaxation Time</th>
<th></th>
<th>T2 Relaxation Time</th>
<th></th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Age</td>
<td>No. Exams</td>
<td>Age</td>
<td>No. Exams</td>
</tr>
<tr>
<td></td>
<td>Change (%)</td>
<td>p-value</td>
<td>Change (%)</td>
<td>p-value</td>
</tr>
<tr>
<td>Asc</td>
<td>0.16</td>
<td>0.754</td>
<td>-4.21</td>
<td>0.057</td>
</tr>
<tr>
<td>Asp</td>
<td>-0.28</td>
<td>0.108</td>
<td>3.08</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>GABA</td>
<td>0.69</td>
<td>0.017</td>
<td>3.68</td>
<td>0.006</td>
</tr>
<tr>
<td>Glc</td>
<td>-0.29</td>
<td>0.571</td>
<td>4.18</td>
<td>0.051</td>
</tr>
<tr>
<td>Gln</td>
<td>-0.57</td>
<td>0.024</td>
<td>3.34</td>
<td>0.001</td>
</tr>
<tr>
<td>Glu</td>
<td>0.06</td>
<td>0.603</td>
<td>2.59</td>
<td>&lt; 0.001</td>
</tr>
<tr>
<td>Gly</td>
<td>1.45</td>
<td>0.387</td>
<td>4.51</td>
<td>0.645</td>
</tr>
<tr>
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<td>0.190</td>
<td>1.42</td>
<td>0.142</td>
</tr>
<tr>
<td>GSH</td>
<td>0.53</td>
<td>0.329</td>
<td>-0.97</td>
<td>0.701</td>
</tr>
<tr>
<td>Ins</td>
<td>-0.12</td>
<td>0.357</td>
<td>1.37</td>
<td>0.019</td>
</tr>
<tr>
<td>Lac</td>
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<td>0.810</td>
</tr>
<tr>
<td>NAA(CH₃)</td>
<td>-0.08</td>
<td>0.647</td>
<td>2.88</td>
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<tr>
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<tr>
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<tr>
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<tr>
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<tr>
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<tr>
<td>tCr(CH₃)</td>
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<tr>
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</tr>
<tr>
<td>MM2</td>
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<tr>
<td>MM3</td>
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<tr>
<td>MM4</td>
<td>0.29</td>
<td>0.167</td>
<td>1.05</td>
<td>0.332</td>
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Table 3.5. Relaxation times variations with age and number of exams.
Results from the linear regression model analysis for T₁ and T₂ relaxation times using age and the number of exams as predictors. The observed change is shown along with its associated p-value. Age-related changes per month were calculated as \( \beta_1 \times 100/\beta_0 \) and changes per exam as \( \beta_2 \times 100/\beta_0 \) according to equation (3.1). Measurements were done on the Main VOI.
3.4.2. Neurochemical profile in aging rats

The same quantification method was used for the determination of the neurochemical profile in the Main VOI at each time point for both cohorts as described in section 3.2.6 and 3.3.2. For each cohort, the first three time-points were corrected using their measured relaxation times and their fractional CSF contributions (Table 3.2). Since the relaxation times could not be measured for the fourth time-point, the values measured on the third time-point were considered.

![Boxplots of absolute concentration variations at different ages](image)

**Figure 3.13.** Concentration variations in the Main region observed during the study. Boxplots of absolute concentration variations at different ages are shown for the C1 (blue) and C2 (red) cohorts for 6 metabolites measured from the Main VOI. Significant changes are indicated for each metabolite and are listed in Table 3.6. The number of measurements at each age were 12, 13, 18 and 8 for C1 and 13, 13, 16 and 6 for C2. (*) \( p < 0.05 \) and (**) \( p < 0.005 \).

3.4.2.1. Changes with age

Significant metabolic alterations in metabolite concentrations were associated with aging in the Main VOI. As shown in figure 3.13, increases of the concentrations of Gln, Ins, MM1, tCho and
tNAA were found (from 0.5 to 2 % per month). The [Glu]/[Gln] ratio was also found to decrease with age (-1.8 %, p = 0.025).

3.4.2.2. Changes with the number of exams

As for the relaxation times, numerous changes were also associated with the number of exams (Table 3.6): increased concentrations of tCho, Asc, MM2 and MM3 were found along with decreases for GABA, Gln, tNAA and MM1 concentrations. The analysis results did not show any significant change of tCr.

3.4.3. Regional variability

Quantification of the neurochemical profiles from the thalamus, CP and CC regions was done as shown in section 3.3.3. The linear regression results, with age and the number of exams as estimators, are shown in Table 3.6. The evolution of the metabolites concentrations showing major changes as well as the [Cr]/[PCr] and [Glu]/[Gln] ratios, for the 4 studied regions, are shown in figure 3.14.

3.4.3.1. Changes with age

Some consistent alterations were found across all regions to be correlated with age: a decreased Glu/Gln ratio (from -1.2 to -1.8%), increased Ins concentrations in the thalamus (2%) and the striatum (0.7% but p = 0.06), and increased macromolecular contributions in the thalamus (from 1.1 to 1.7%) and the cingulate cortex (from 1.2 to 1.6%). In addition, an increase of the Cr/PCr ratio with age was observed for the cingulate cortex (3%, p = 0.02).

3.4.3.2. Changes with the number of exams

Across the 3 VOIs, several concentrations changes were found to be correlated with the number of exams for Asc, Glu, PCr, PE, NAA, NAAG and MM4. However, only increases in Asc and Glu were observed in at least two different VOIs (Main+Thal for Asc and Main+CC for Glu). A smaller number of significant changes were observed for the CP (2) and Thal (4) VOIs compared to the CC (16) and Main (11) regions (which notably, have higher GM contributions than the CP and Thal regions).
Figure 3.14. Metabolic changes in the aging rat brain in the four VOIs. Mean concentrations are presented along with their standard deviations as error bars for the different time-points for the C1 (green colors, ages 1, 4, 7, 12) and C2 (blue colors, ages 8, 11, 14, 18) cohorts. Identical corrections for $T_1$- and $T_2$-weighting were applied to the concentrations of the four VOIs at each time-point. *MM1 is expressed in units of $^1$H mmol/L.
<table>
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<th>Region</th>
<th>Change (%)</th>
<th>P. Value</th>
<th>Change (%)</th>
<th>P. Value</th>
<th>Change (%)</th>
<th>P. Value</th>
<th>Change (%)</th>
<th>P. Value</th>
<th>Change (%)</th>
<th>P. Value</th>
<th>Change (%)</th>
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<td>0.586</td>
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<td>0.64</td>
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<td>0.791</td>
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<td>0.014</td>
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<td>0.008</td>
<td>0.61</td>
<td>0.796</td>
<td>0.46</td>
<td>0.355</td>
<td>3.77</td>
<td>0.103</td>
</tr>
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<td>0.202</td>
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<td>0.63</td>
<td>0.403</td>
<td>-3.75</td>
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</table>

Table 3.6. Metabolite concentration variations with age and number of exams as predictors.
Results from the linear regression model analysis for the absolute metabolite concentrations using age and the number of exams as predictors. Each region was analyzed independently. The observed change is shown along with its associated P. value. Age-related changes per month were calculated as $\beta_1 \times 100/\beta_0$ and changes per exam as $\beta_2 \times 100/\beta_0$ according to equation (3.1).
3.5. DISCUSSION

3.5.1. qMRS of the young rat brain

In this first study, we present the first $^1$H NMR spectra at 17.2 Tesla in the rat brain in vivo. The $T_1$ and $T_2$ relaxation times were determined for 22 and 18 functional groups respectively at different ages, as well as the macromolecule baseline divided into 4 resonance groups. Good quality data and spectral decomposition were achieved due to the increased spectral dispersion and sensitivity at UHF. In addition, the adopted parameterization of the macromolecule baseline (table 2.2) suitably accounted for the macromolecule and lipid contributions and allowed to account for differences in $T_1$ and $T_2$ relaxation times between the four MM components.

3.5.1.1. Challenges of $^1$H MRS at 17.2 T

The difficulties of UHF MRS are increased chemical shift displacement and a higher sensitivity to magnetic field susceptibility effects. Motivated by the need to limit the required $B_1$-field intensity while using large-bandwidth slice-selective adiabatic pulses, HS8 refocusing pulses with a 20kHz bandwidth were used at the cost of a larger transition bandwidth (duration = 1 ms, $B_1$ = 3.2 kHz, 6.3 kHz transition bandwidth) compared to other hyperbolic-secant pulses such as a HS1 with the same bandwidth-length product (duration = 1 ms, $B_1$ = 6.2 kHz, 3.3 kHz transition bandwidth). The use of FASTMAP combined to the manufacturer’s MAPSHIM procedure allowed us to obtain a mean water spectral linewidth of 0.031 ± 0.004 ppm in the Main region, which is comparable to values reported at 11.7 T (0.032 ppm) with similar VOI positioning and dimensions (de Graaf, et al., 2006).

3.5.1.2. MM parametrization method

The MM parameterization method used provided excellent results for fitting the macromolecule and lipid signals as illustrated by figures 3.4, 3.5, 3.6 and table 3.3. A similar approach has been previously proposed (Hong, et al., 2011) and was successfully applied for the analysis of ultra-short-echo-time neurochemical profiles of the rat brain in vivo at 16.4 T. Contrary to their methods, our MM parameterization allowed us to use a regularized spline baseline with a knot spacing of 1.0 leading to a flat baseline, even for the $T_1$-weighted spectra. Additionally, separately accounting for the MM resonances was particularly appealing for this study as it allowed us to estimate for the first time the $T_2$ relaxation times of macromolecule and lipids groups outside of the 0.5 to 2.0 ppm region of the $^1$H spectrum (MM2 to MM4) and their $T_1$ relaxation times with an IR technique. Indeed, MM baseline measurements using a saturation
recovery technique have been shown at 1.5 T (Hofmann, et al., 2001) although their precise values were not provided.

3.5.1.3. $T_1$ and $T_2$ relaxation times

The distribution of $T_1$ relaxation times found in the young DA rat at 17.2 T is shown in figure 3.8. Overall, the $T_1$ values are similar for the majority of metabolites. The mean metabolite $T_1$ relaxation time (1720 ms) was close to that of tCr(CH$_3$) (1750 ms) and Glu (1700 ms). The resonances of Tau, Asc, Gly and NAA(CH) present the longest $T_1$ values while NAA(CH$_2$), PCh, GPC, NAAG and tCr(CH$_2$) have the shortest ones. The $T_1$ values of the MM groups were found within a short range from 570 ± 20 (MM3) to 800 ± 40 ms (MM4). The relatively narrow spread of longitudinal relaxation times shows that a similar $T_1$-weighting will be experienced at shorter repetition times. For instance, if TR were 2000 ms, the equilibrium magnetization of tCr would be 72% of the fully relaxed magnetization and only marginally less for Tau with 65%.

$T_2$ relaxation times of singlets and J-coupled resonances were reliably estimated for 18 metabolites moieties and 4 groups of MM resonances ($R^2 > 0.80$). They are shown in descending order in figure 3.8. Similar to the $T_1$ measurements, the mean metabolite $T_2$ relaxation time (148 ms) was close to that of tCr(CH$_3$) (141 ms). The resonances of NAA(CH$_3$), GPC and PCh exhibited the longer $T_2$ values, whereas the shortest metabolite $T_2$s were found for the J-modulated resonances of Gln, Asp and GABA. As expected, macromolecules $T_2$ times were the shortest, ranging from 20 (MM1) to 55 ms (MM3).

3.5.1.4. Magnetic field strength dependency of $T_1$ and $T_2$ relaxation times

Previous $T_1$ measurements using the IR technique have been reported at lower magnetic fields using either the LASER sequence at 4.0, 9.4 and 11.7 T (de Graaf, et al., 2006) or the SPECIAL sequence at 9.4 and 14.1 T (Cudalbu, et al., 2009). De Graaf et al considered the 7 most prominent metabolites moieties (tCr(CH$_3$), tCr(CH$_2$), tCho, NAA(CH$_3$), Glu, Ins and Tau) and the macromolecules resonances at 1.7, 1.4, 1.1 and 0.9 ppm. Using AMARES, Cudalbu et al also estimated the $T_1$ values of these metabolites with an IR technique. De Graaf et al concluded on the increase of the $T_1$ relaxation times with the magnetic field strength accordingly to the Bloembergen-Purcell-Pound (BPP) theory of dipolar relaxation (Bloembergen, et al., 1948). However, by comparing our results at 17.2 T with those reported at 11.7 T, a mild increase of $T_1$ was observed only for NAA(CH$_3$), Glu and MM1 while no clear increase was observed for the other metabolites due to the larger standard deviations for these measurements. Likewise, Cudalbu et al found a statistically significant $T_1$ difference for NAA(CH$_3$) when increasing from 9.4 T to 14.1 T but the remaining measured metabolites were found to be the same within experimental error. To further examine the dependence of the spin-lattice relaxation times with the magnetic field strength, the following power law was considered:
to describe the evolution of $T_1$ with the magnetic field $B_0$ (Bottomley, et al., 1984). To assess the agreement between our values at 17.2 T and those reported by de Graaf et al, the empirical parameters $\alpha$ and $\beta$ were estimated for NAA(CH$_3$), tCr(CH$_3$), tCr(CH$_2$), tCho and MM1 using only their reported values at 4.0, 9.4 and 11.7 T (Table 3.7). Glu and Ins were not considered due to the elevated experimental error at 9.4 T and 11.7 T. The resulting extrapolation curves (Figure 3.15A, dashed lines) for NAA(CH$_3$) and the tCr moieties were all within error margins of our measured $T_1$ values at 17.2 T. For tCho, the larger discrepancies observed should be linked to its less precise quantification due to its spectral overlap with Tau. Including our results in the estimation of $\alpha$ and $\beta$ had only a visible impact on the extrapolation results for tCho and MM1 (Figure 3.15A, continuous line). From these simulated curves, the expected increases for $T_1$ from 11.7 T to 17.2 T are the largest for NAA (79 ms) and MM1 (132 ms) while they are relatively minor for tCr(CH$_2$), tCr(CH$_3$) and tCho (with 57, 41 and 45 ms respectively). Such differences are difficult to detect as the experimentally-induced errors are of a similar magnitude or larger (Table 3.7). This could explain both our results and the lack of remarkable $T_1$ increases above 9.4 T observed by Cudalbu et al, with the exception of NAA(CH$_3$) that shows an statistically significant increase. Nevertheless, methodological differences between the three studies could also thwart the comparison. Notably, it has been shown (de Graaf, et al., 1999; de Graaf, et al., 2014; Shemesh, et al., 2013) that the water suppression scheme and the spectrally-selective inversion pulses may have a non-negligible effect on metabolite intensity modulation and apparent longitudinal relaxation rates through (i) magnetic transfer effects between mobile and immobile proton pools or (ii) chemical exchange and cross-relaxation mechanisms between water and non-exchangeable protons.

Previous studies have measured the $T_2$ relaxation times of metabolites in the human (Michaeli, et al., 2002) and rat (de Graaf, et al., 2006; Deelchand, et al., 2015) brains using the LASER sequence. The reported $T_2$ values of NAA(CH$_3$), tCr(CH$_3$), tCr(CH$_2$), tCho and MM1 are summarized in table 3.7. The data demonstrate a consistent metabolite $T_2$ decrease and a narrowing of $T_2$ times with the increasing magnetic field. To further analyze these results ranging from 4.0 T to 17.2 T, the empirical expression:

$$T_2(B_0) = \gamma \exp\left(-\frac{B_0}{\delta}\right)$$

was used to describe the $B_0$ dependency of $T_2$ values. The $\gamma$ and $\delta$ coefficients were fitted for the 4 metabolites and MM1 using all the values shown in table 3.7. The resulting extrapolation curves and coefficients are shown in figure 3.15B and table 3.7. These simulations predict a narrowing of the $T_2$ distribution and a further decrease of the $T_2$s at higher magnetic fields. The resonances of NAA(CH$_3$), tCr(CH$_3$) and tCho exhibit the strongest dependency to the magnetic field intensity. Interestingly, the $T_2$ weighting parameter $\delta$ is very similar for NAA(CH$_3$), tCr(CH$_3$) and MM1.
The simulation therefore implies that a similar relative decrease is experienced by the three resonances and the main difference comes from the term $\gamma$ which could be correlated to the intrinsic rotational correlation time and the effectiveness of the pulse sequence to refocus diffusion-related dephasing. The largest $\delta$ value was found for tCr(CH$_2$) which showed no apparent dependence to the magnetic field strength. With the exception of tCr(CH$_2$), these results are in contradiction with the BPP theory (Bloembergen, et al., 1948) which predicts field-independent $T_2$ relaxation times for a large range of rotation correlation times. But our data is rather in agreement with the field-dependency of $T_2$ proposed by Michaeli et al originated from increased microscopic susceptibility gradients at higher fields, inducing a more important dynamic dephasing through diffusion.

$T_2$ values obtained at 9.4 T and 14.1 T (Xin, et al., 2008; Xin, et al., 2013) using a Hahn spin-echo also showed a decrease of metabolite $T_2$s with $B_0$, notably for J-coupled metabolites Glu, Ins.

<table>
<thead>
<tr>
<th>Metabolite</th>
<th>4.0 T</th>
<th>7.0 T</th>
<th>9.4 T</th>
<th>9.4 T</th>
<th>11.7 T</th>
<th>17.2 T</th>
<th>$\gamma$</th>
<th>$\delta$</th>
</tr>
</thead>
<tbody>
<tr>
<td>tCr(CH$_3$)</td>
<td>233 ± 6</td>
<td>221 ± 19</td>
<td>171 ± 4</td>
<td>170 ± 12</td>
<td>159 ± 7</td>
<td>141 ± 7</td>
<td>277.65</td>
<td>22.4</td>
</tr>
<tr>
<td>tCr(CH$_2$)</td>
<td>141 ± 11</td>
<td>-</td>
<td>128 ± 6</td>
<td>146 ± 16</td>
<td>125 ± 7</td>
<td>132 ± 10</td>
<td>142.87</td>
<td>168.9</td>
</tr>
<tr>
<td>tCho</td>
<td>570 ± 29</td>
<td>-</td>
<td>441 ± 36</td>
<td>445 ± 67</td>
<td>366 ± 73</td>
<td>212 ± 21</td>
<td>768.46</td>
<td>15.4</td>
</tr>
<tr>
<td>NAA(CH$_3$)</td>
<td>392 ± 15</td>
<td>341 ± 25</td>
<td>294 ± 5</td>
<td>321 ± 30</td>
<td>285 ± 26</td>
<td>226 ± 26</td>
<td>459.65</td>
<td>23.9</td>
</tr>
<tr>
<td>MM1</td>
<td>27a ± 2</td>
<td>-</td>
<td>28a ± 2</td>
<td>-</td>
<td>25a ± 2</td>
<td>20 ± 2</td>
<td>42.50</td>
<td>22.6</td>
</tr>
</tbody>
</table>

Table 3.7. Metabolite relaxation times at increasing magnetic fields

Metabolites and macromolecules $T_1$ and $T_2$ relaxation times measured at different magnetic fields in the brain in vivo (mean ± SD). Rat brain values at 4.0, 9.4, 11.7 and 14.1 T and human brain values at 7.0 T are taken from references (de Graaf, et al., 2006; Michaeli, et al., 2002; Cudalbu, et al., 2009; Deelchand, et al., 2015). The localization sequence for each study is indicated. For the $T_1$ studies at 9.4 and 14.1 T only the results obtained with the IR technique were considered. Reported MM1 values were calculated from references as the mean relaxation times of the MM resonances from 0.9 to 1.8 ppm when available.

$^a$ SD is the mean value of the individual SD reported for the different MM components in (de Graaf, et al., 2006).

$^b$ Values were estimated from 3.2 and 3.4 from (Cudalbu, et al., 2009).

$^c$ Values were obtained using the progressive saturation recovery technique (Cudalbu, et al., 2009).
and Tau. Yet, these values were not compared to those obtained using Carr-Purcell refocusing pulses, which are closer to the intrinsic $T_2$ value due to a reduction of the diffusion and proton-exchange contributions of the apparent $T_2$ (Michaeli, et al., 2002) and also for the refocusing of cross-relaxation or interference effects due to dipole-dipole interactions in scalar-coupled metabolites (Deelchand, et al., 2015).

Figure 3.15. $T_1$ and $T_2$ times estimations at higher magnetic fields. Calculated (A) $T_1$ and (B) $T_2$ relaxation times up to 25 T using the functions (3.2) and (3.3) based on in vivo data measured with LASER at different fields. LASER $T_1$ values at 4.0, 9.4 and 11.7 T were fitted and the corresponding curves were plotted as dashed lines. The continuous lines were obtained from fitting all the LASER $T_1$ values in table 3.7. Standard deviations are shown as reported in their respective study. The coefficients for the continuous curves are summarized in table 3.7.

3.5.1.5.  *In vivo* $^1$H neurochemical profiles

This study presents the first in vivo $^1$H neurochemical profiles acquired at 17.2 T (figure 3.4), demonstrating the possibility to detect and quantify up to 23 metabolite moieties. The absolute concentrations found here for young DA rats were in good agreement with other studies, most of
them using ultra-short-echo-time acquisitions (Hong, et al., 2011; Choi, et al., 2009; Pfeuffer, et al., 1999; Mans, et al., 1994; Mlynàrik, et al., 2008). Correcting for T1- and T2-weighting resulted in substantial increases of concentrations, ranging from 11 % (GPC) to 42 % (Gln) for metabolites and up to a 130 % (MM1) increase for the MM components. Our Gln concentration was higher than those reported in other MRS studies but are in good agreement with values from in vitro rat brain extracts (Mans, et al., 1994). This difference could be originated from the different handling of the macromolecule baseline among studies. Contrary to the study of Hong et al, we were not able to detect acetate when its model spectrum was added to the LCModel basis-set. However, we were able to detect other low-concentration metabolites such as alanine (CRLB = 14 %), glycine (CRLB = 16 %) and serine (CRLB = 20 %). While Ser is seldom detected and quantified in vivo (Choi, et al., 2009), its addition to the basis-set was necessary to properly account for the NMR signal at 3.84 ppm. A more specific approach such as difference editing would be necessary to validate its detection at 17.2 T.

3.5.1.6. Regional variability

Neurochemical profiles from 4 different VOIs (Thal, CP, CC and Main) were acquired in this study, showing comparable quality despite the differences in voxel size and distance to the coil. Metabolic differences between the VOIs (figure 3.10) were expected (Tkac, et al., 2003; Duarte, et al., 2014) and they may be linked to the presence of tissue-specific neural cell types (Urenjak, et al., 1993) and organization based on their GM and WM compositions (Banerjee, et al., 2012; Wang, et al., 1998; Bhattacharyya, et al., 2011). Even if numerous 1H MRS studies have measured the neurochemical profile in the rat brain from various brain region, the large variability in methods doesn’t allow us to definitely confirm our observations. If we consider the metabolic data obtained from voxels located in the hippocampus, the striatum and the cortex by Tkac et al. (Tkac, et al., 2003) from 1 month-old Sprague-Dawley rats, most relative differences in concentrations between our Main (encompassing both cortex and a some hippocampus) and CP VOIs are in agreement. Only our lower Ins concentration in the Main region seems to be in contradiction with their study. However, this may be explained by the contributions of the hippocampus and WM to our Main VOI. Metabolic variations between the main and thalamus region were mostly in agreement with spectroscopic data presented by Just et al. (Just, et al., 2014), with the exception of Glu, Asp and Ala which were less concentrated in their thalamic VOI compared to their cortical VOI. Again, this could be due to larger WM contributions in our Main VOI. The possibility that small discrepancies in metabolite concentrations are due to the different rat strains used in these studies cannot be overruled.
3.5.2. qMRS of the aging rat brain

Two remarkable features of the normal aging brain are progressive mitochondrial dysfunction (Federico, et al., 2012) and chronic neuroinflammation (Czeh, et al., 2011). Both processes promote the production of ROS, which are detrimental to neurons. There is growing evidence that these factors play major roles in age-related neurodegenerative diseases (Witte, et al., 2010; Karbowski, et al., 2012; Moràn, et al., 2012). The accumulation of molecular insults from oxidative stress leads to a persistent perturbation of the various mitochondrial functions. Progressively, neuronal oxidative energy synthesis is impaired and glial cells, in particular microglial cells, are activated. In time, the aging brain’s capabilities for efficient energy production and DNA repair diminish; the prolonged glial activation becomes neurotoxic and the elevated IFN-γ levels reduce or impede cellular regeneration (section 3.1.1). This chain of events eventually leads to an impaired cellular homeostasis and neurotransmission and ultimately to loss of cognitive function.

The main goal of this study was to follow the metabolic changes happening in the early stages of normal aging in the aging DA rat. By following in parallel 2 cohorts of healthy rats, an age span of 18 months could be covered leading to the observation of multiple metabolic alterations using quantitative in vivo $^1$H MRS.

3.5.2.1. Metabolic alterations with age

Based on the results from the linear regression analysis (table 3.6 and figure 3.15), the most robust metabolic changes observed over the 4 regions were increases of Ins and MM concentrations and a decrease in the [Glu]/[Gln] ratio. The concentrations appeared to vary from 1 to 2 % per month and are consistent with a progressive glial activation (Jenden, 1979 pp. 13-24; Brand, et al., 1993). These observations are in agreement with previous aging studies (Reyngoudt, et al., 2012; Haga, et al., 2009). The increase in MM content is also in agreement with an increased membrane turn-over and a possible accumulation of protein aggregates or cellular debris. No changes were observed for NAA on any of the 4 VOIs. This is in agreement with previous human studies (Reyngoudt, et al., 2012; Haga, et al., 2009) although increases or decreases of NAA with age have also been reported in humans (Brooks, et al., 2001; Fukuzako, et al., 1997; Urrila, et al., 2004). Our results are in agreement with work done in the aging mouse (Duarte, et al., 2014) as they did not find NAA changes in the cortex and found a decrease in the striatum (CP) only at 24 months of age.

The observed decreases of the Glu/Gln ratio can be attributed to an increase in Gln concentrations accompanied with a decrease in Glu concentrations. Even if both metabolic changes were not always statistically significant for all the VOIs, these changes may indicate the beginning of previously observed shifts with age between Glu synthesis in neurons and its recycling into Gln in glial cells as it has been shown in $^{13}$C MRS studies (Rothman, et al., 2011). Indeed, $^{13}$C MRS
allows to probe metabolic pathways such as glycolysis, the tricarboxylic (TCA) cycle or the glutamate-glutamine cycle using $^{13}$C-labeled energy substrates and previous $^{13}$C MRS studies in aging mice (Patel et al., 2012; Patel et al., 2014) and humans (Boumezbeur et al., 2010) have shown an accelerated glial TCA cycle while the neuronal TCA and glutamate-glutamine cycles slow down. Yet, the decrease of neuronal TCA in the study done by Boumezbeur et al. was highly correlated with concomitant decreases in Glu and tNAA.

The CC region presented a decrease in PCr/Cr indicating a reduction in energy metabolism. This was accompanied by decreasing levels of tNAA and Glu, although neither NAA (P = 0.051) nor NAAG showed significant changes with age. There was no evidence of glial activation as Gln and Ins levels were unchanged. These changes could suggest a mild decrease in neuronal metabolic activity with age, which has been reported in AD studies (Salehi et al., 1999) where the decrease in neuronal metabolic activity was correlated with the genetic background (polymorphism of apolipoprotein E) and cell characteristics (cell profile area and size of the Golgi apparatus). Furthermore, frailness in the posterior cingulate region has also been observed in AD patients (Zimny et al., 2011; Nestor et al., 2003). Clearly, the application of supplementary techniques is needed for the validation of this hypothesis, notably with dynamic $^{13}$C MRS using $^{13}$C-labeled glucose and histology. Furthermore, the presence of confounding factors in the analysis of the data, notably the effect of the number of exams, invites for caution on the interpretation of the data based solely on the applied statistical model.

Our results are rather consistent with the aging mouse study done by Duarte et al. as they did not find NAA and PCr/Cr decreases in the cortex (closely matching our Main VOI) but they did observe significant changes in the striatum (CC). Their Ins and tCho also were observed to increase with age. The Glu/Gln ratio decreased in their three explored regions in the mouse as well as the four regions in the DA rat, making it a potentially interesting biomarker for future longitudinal aging studies.

### 3.5.2.2. Relaxation times alterations with age

In addition to metabolic changes, modest shortenings of the T$_2$ relaxation times were observed for tNAA (-1%, P = 0.02) tCho (-1.5%, P = 0.05) and MM1 (-0.7%, P = 0.03). Overall, a tendency towards a reduction of the T$_2$ times was observed, which could indicate a mild change in the cellular environment with age, such as increases in iron content or protein aggregates, which is in agreement to the observed increase of macromolecule content and previous work done in humans (Marjanska et al., 2013).

Concerning the evolution of the T$_1$ relaxation times, our analysis revealed only two statistically significant and very small changes associated with age for GABA (+0.7%, P = 0.02) and Gln (-0.6%, P = 0.02).
3.5.2.3. Alterations with the number of exams

The number of exams experienced by the rats was found to be an important and unexpected contributing factor to the evolution of the neurochemical profile during this study. Both the Main and the CC regions presented numerous changes linked to the number of scans. Since these two regions had the highest GM/WM ratios, a possible sensitivity of GM tissue could be speculated by the physiological stresses suffered during the experiments. However, the causes underlying the physiological alterations during the MR experimental sessions have yet to be investigated. Nonetheless, the interaction with the aging process was accounted for by using a linear-regression with mixed effects analysis with the number of exams as one of the regressors.

The most prominent observation was an increase of the T\textsubscript{1} relaxation times in the Main region for Asp, GABA, Gln, Glu, NAA(CH\textsubscript{3}), NAA(CH), NAAG, tCr and MM2 at a rate of 2 to 3% per exam. These increased T\textsubscript{1} times could indicate a possible cell swelling or another alteration of the intracellular matrix leading to an enhanced mobility of the metabolites. Transient changes such as an increase in water content or brain temperature have also been associated to T\textsubscript{1} increases (Barbier, et al., 2005; Fatouros, et al., 1991; Lin, et al., 1997) but such reasons do not seem pertinent since the T\textsubscript{1} measured for brain tissue water did not change throughout the study. In addition to these T\textsubscript{1} increases, the T\textsubscript{2} times of Tau, tCr(CH\textsubscript{2}) and MM3 decreased at rates of 4 to 7% per exam.

The observed alterations associated to the number of scans are difficult to interpret since, to our knowledge, this is the first time that such effects have been observed.

One possible explanation could be the direct effect of the static magnetic field (SMF), as it has been shown in rats to have a reduced neuronal density in the hippocampus after being exposed to small magnetic fields (up to 1 μT) during gestation (Whissell, et al., 2009). Oxidative stress has been shown to decrease at 1.5 T in healthy volunteers (Sirmatel, et al., 2007), however mice exposed at 3 and 4.7 T showed increased metallothionein (a ROS by-product) and lipid peroxidation in the liver (Satoh, et al., 1996; Watanabe, et al., 1997). Additionally, even small SMF from 3 to 110 mT have shown increased effects of tumor-treating drugs in mice exposed to SMF for 35 mins to 4 hours a day (Tofani, et al., 2003; Gray, et al., 2000). In vitro tests at 7 T have revealed tumor growth inhibition through the effect of SMF alone and increased tumor cell apoptosis at 1 T (Ghibelli, et al., 2006; Raylmann, et al., 1996). These changes are nevertheless cell-line dependent and it has been concluded that the overall effects of SMF do have an effect on free radical metabolism, but the impairments on normal cell-growth and toxicity are minor or non-existent (Ghodbane, et al., 2013). The issue of B\textsubscript{0} effects on health is still in debate (Richard, 2005) and it is not clear whether there would be a persistent effect on the adult brain and it is yet to determine the effects of sporadic but prolonged exposure (~6-8 h per experiment) at 17.2 T.

Another plausible cause could be the intense time-varying B\textsubscript{1} fields required to achieve refocusing using short AFP pulses at 17.2 T. Since cooling in the brain is achieved mainly through perfusion (which is reduced by the anesthesia) and the experimental sessions for each rat...
consisted of at least 5 continuous scanning hours using LASER, there could be temperature increases especially close to the coil. Thus, these temperature elevations would be more intense in the cortical areas (Main and CC) compared to sub-cortical area (CP and Thal). To validate this hypothesis, thermometry measurements could provide relevant information about the temperature evolution in the rat brain non-invasively. This technique relies on measuring the local shifts of the Larmor frequency caused by local changes in temperature (Hindman, 1966). To evaluate tissue and cellular damage, temperature models have been used to simulate the induced damage on the brain using microwave probes (Sherar, et al., 2000; Skinner, et al., 1998). These models provide a framework to evaluate the potential damaged caused by the applied B1 field and they have been used to predict the extent of tissue damage based on parameters of cell death, microvascular blood flow stasis and protein coagulation (Sherar, et al., 2000). Yet, perfusion effects are not integrated in the Arrhenius equation despite that it plays a critical role in the cooling of the brain during long exams.

Histological validation could be achieved by looking at morphological changes in cells such as increases in cytoplasm and shrinking of the cellular nucleus; signs of necrosis and blood coagulation and microglial activation. To avoid a possible confounding effect due to other factors such as handling stress, SMF and anesthesia, a control group should be needed where an identical experimental protocol would be applied on rats of the same age with RF pulses turned off and sacrifices made either immediately after experiment or one week after, as it has been shown in the rabbit brain that damages may become visible several days after the experiment took place, notably coagulative necrosis (Matsumi, et al., 1994).

Another cause could be the cumulative effect of the isoflurane anesthesia, which has been associated to lasting alterations on cerebrovascular regulation when exams were done under hypoxia conditions (Wegener, et al., 2008). Contrary to other halogens, it has also been shown that isoflurane is hepatotoxic exclusively under hypoxia and reduced blood-flow conditions in the rat (Van Dyke, 1982; Schieble, et al., 1988; Harper, et al., 1982). Although the rats in our study breathed pure oxygen during the experiments and lactate levels were stable over their whole duration, the long duration of the experiment had most likely weakened the animal. It has been shown that fasting rats in hypoxia have suffered of liver necrosis (Van Dyke, 1982) and that hepatotoxicity levels of isoflurane may dependent on the genetic background, as it has been shown for halothane (Gourlay, et al., 1981). Therefore, the possibility of liver injury should not be discarded and verification tests would be needed to verify the condition of the liver, notably by measuring transaminase concentration levels in the blood (De Ritis, et al., 1955).

A last possibility could be that these observations are related to the stress experienced by the rats during or in-between MRS sessions. Indeed, it has been shown that depressed patients exhibit increased glutamate (Sanacora, et al., 2012) and decreased GABA (Sanacora, et al., 1999; Hasler, et al., 2007). Both neurotransmitters alterations are rather consistent with our observations. However, the MRS literature in that matter is rather inconsistent depending on the method and the conditions studied. For example, in a recent ex vivo MRS study of the effect of acute restraint
stress (Drouet, et al., 2015), it was shown that both GABA and Glu levels were increased in rats being restrained for 1 hour.

3.5.2.4. Impact of the metabolite relaxation times on absolute quantification

For this study, T₁ and T₂ metabolite relaxation times were measured at most time-points on the Main region and were assumed to be identical for the 3 other VOIs. Measuring the relaxation times with the presented methods was a rather time-consuming process as the majority of the experimental session was dedicated to the acquisition of T₁- and T₂-weighted spectra. For this reason, measurements of metabolite relaxation times are rarely done in MRS studies, neglecting the effects of differential T₁- and T₂-weighting amidst metabolites. Therefore, it is interesting to evaluate the impact of metabolite relaxation time corrections in our study by looking on how different our observations would have been without such correction.

As a matter of fact, most statically significant metabolic alterations were very similar (not detailed). The differences were the following: Glu/Gln changes were no longer observed for Thal and CC and the MM changes in the CC were no longer significant; instead NAA was now increasing with age in CC. Also, in the Main region, Gln and MM1 decreases with the number of exams were no longer statically significant.

These relatively few differences demonstrate that considering relaxation times for the purpose of investigating metabolic alterations is relevant when data are acquired using short TR or long TE. For this study, long TR (5s) and rather short TE (16.5 ms and CPMG refocusing) were used. As a consequence, the quantification of species with strong T₂-weighting such as Glu, Gln and the macromolecules (Table 3.3) were the only ones impacted.

3.5.2.5. Alternative statistical analysis

Our MRS data were examined using a linear-regression with mixed effects analysis assuming that the age and the number of exams were independent variables and considered the data of both cohorts. Yet, a more conservative approach could have been considered by omitting the β₂*Exams term in equation (3.1), leaving age as the only regressor and analyzing each cohort separately. The results of such analysis are presented in tables 3.8 and 3.9, respectively summarizing the relaxation times and concentration results.

Several interesting observations can be done. First, the main changes observed simultaneously on the VOIs with the mixed effect analysis, namely the decrease in Glu/Gln ratio and the increase of Ins and MMs, are preserved although they are present in different “stages”. The first stage, corresponding to the young C1 cohort, only presents an increase in Ins, Gln and tCr but stable Glu levels, suggesting an onset of glial activation. For the second stage, corresponding to the older C2 cohort, Ins continues to increase along with MM content and it is accompanied by a
reduction of the PCr/Cr ratio, suggesting a possible decline of mitochondrial function and the accumulation of protein residues.

A second observation is that numerous metabolites show a $T_1$ increase for C1 and C2. In addition, metabolites presenting $T_1$ increases in C1 are also presenting a $T_1$ increase in C2, as it can be clearly observed in figure 3.8. Yet, the relaxation times of brain water tissue still do not change.

3.5.2.6. Stability of the MM parametrization

The initial MM parameterization was done based on data from 1 month-old rats. In order to check that the macromolecule baseline was correctly fitted at any ages, mean metabolite-nulled spectra were acquired at each age and were fitted. As illustrated by figure 3.16, the MM parameterization allowed for the proper fit of the metabolite-nulled spectra at all ages.

![Figure 3.16. Mean metabolite-nulled spectra and fits in the rat brain from 1 to 18 months of age. Mean MM spectra (black) and LCModel fit (red) after removal of residual metabolite contributions (as shown in figure 2.12). Besides the MM spectrum at 1 month of age, each spectrum was obtained by adding automatically-phased individual spectra weighted by their measured noise levels. Mean spectra are scaled with respect to the resonance at 0.9 ppm.](image)

3.5.2.7. Methodological limitations

In this study, several limitations have to be considered. First, the water content for GM and WM were considered to be constant at all ages (Lentner, 1981). Yet, the GM water content has been found to vary significantly in humans (Chang, et al., 1996), in particular for subjects above 50
years old. In mice, total brain water content is stable for the first year of life but it shows a global decrease from 79 % to 75 % water content during the second year (Duarte, et al., 2014). Assuming similar changes for the GM in the rat brain, a 2% decrease of water content between 12 to 18 months could be expected leading to an over-estimation of 2% of the reported metabolite concentrations for the latest time-point.

Secondly, the T\textsubscript{1} maps acquired from the initial time point at the age of 1 month were used to determine the T\textsubscript{1} relaxation times of water in GM and WM and also to determine the GM, WM and CSF compositions of the VOIs. As discussed in section 2.5.1, increases in brain volume and myelination in the rat brain take place until maturation is achieved at 3 months of age (Juraska, et al., 2004; Dobbing, et al., 1971) and myelination has even been shown to continue at least up to 6 months of age (Mengler, et al., 2014). It is therefore likely that water content estimation errors are introduced due to biases in the determination of GM and WM contributions as well as relaxation time changes over the duration of the study. On the other hand, the T\textsubscript{1} and T\textsubscript{2} relaxation times measured from the brain tissue water in our Main VOI employing MRS did not show significant variations, suggesting that age-related changes of GM and WM relaxation times could be negligible. Also, a relatively long repetition time (5 s) was used for the acquisition of water reference spectra, reducing the sensitivity to small variations in T\textsubscript{1} relaxation times. A clear limitation of this study remains as the GM and WM fractions used for the water content calculation were fixed for all ages to those found in the juvenile, immature brain.

The calculated CSF fractional volume values for the thalamus, CP and CC regions were estimated based on the number of pixels presenting elevated T\textsubscript{1} values in the T\textsubscript{1} maps, which could lead to estimation errors due to the relative coarse resolution of the maps for such kind of application. Nevertheless, these errors are not expected to alter our results as minor CSF fractional volumes were found for the CC and CP regions.

Due to time-constraints, metabolites relaxation times could not be measured in all rats for the last time-points. Therefore, the relaxation times measured in the precedent exam were considered for the correction of the metabolites’ T\textsubscript{1} and T\textsubscript{2} relaxation weightings. Based on the result of the linear regression, the associated quantification errors are expected to be at most 2.5 ± 1.9 % for metabolites and 5.6 ± 2.1 % for macromolecules.
<table>
<thead>
<tr>
<th>Metabolite</th>
<th>T1 Relaxation Time</th>
<th>T2 Relaxation Time</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Change (%)</td>
<td>P. Value</td>
</tr>
<tr>
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</tr>
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</tr>
<tr>
<td>Gly</td>
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</tr>
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<tr>
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</tr>
<tr>
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</tr>
<tr>
<td>Lac</td>
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<td>0.042</td>
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<tr>
<td>MM1</td>
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<td>0.148</td>
</tr>
<tr>
<td>MM2</td>
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</tr>
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<td>MM3</td>
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<tr>
<td>MM4</td>
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</tr>
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<td>PE</td>
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<td>0.119</td>
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<td>Tau</td>
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<tr>
<td>tCho</td>
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<td>0.563</td>
</tr>
<tr>
<td>tCr(CH₃)</td>
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</tr>
<tr>
<td>tNAA(CH₃)</td>
<td>-0.21</td>
<td>0.444</td>
</tr>
</tbody>
</table>

Table 3.8. Metabolite relaxation time variations with age as the only predictor.

Results from the linear regression model analysis for relaxation times using age as the only predictor and considering “rat” as random effect. The observed changes for each cohort are shown along with their associated P. value. Age-related changes per month were calculated as $\beta_1 * 100 / \beta_0$, following the nomenclature of equation (3.1).
### Table 3.9. Metabolite concentration variations of individual cohorts with respect to age as sole predictor

Results from a linear regression model analysis for the absolute metabolite concentrations using age as the only predictor and “rat” as a random effect. Each region was analyzed independently. The observed changes for each cohort are shown along with their associated P. values. Age-related changes per month were calculated as $\beta_1 \times 100/\beta_0$, following the nomenclature of equation (3.1).
3.6. CONCLUSION

The main goal of this study was to develop quantitative $^1$H MR spectroscopy at 17.2 Tesla and apply it to evaluate the Dark Agouti rat as a murine model of normal brain aging. $T_1$ and $T_2$ relaxation times were measured in the rat brain in vivo for the first time at 17.2 T for the purpose of absolute quantification using $^1$H MRS. In summary, the $T_1$ relaxation times of 22 metabolite functional groups and the $T_2$ relaxation times of 18 singlets and J-coupled metabolites were measured, as well as the relaxation times of 4 macromolecule resonance groups covering the macromolecule and lipids baseline from 0.5 to 4.5 ppm. Our data are in good agreement with the field-dependency of $T_1$ and $T_2$ relaxation times of brain metabolites as observed at lower magnetic fields. Absolute concentration measurements corrected for relaxation effects were also in good agreement with previously published values.

To our knowledge, this is the first longitudinal study of the aging brain using MRS covering a life-span of 18 months in the rat. Our results are consistent with a mild age-related glial activation which has been reported previously for the aging human, mouse and rat brains using either $^1$H, $^{13}$C MRS or histological methods (Cerbai, et al., 2012; Chang, et al., 1996; Boumezbeur, et al., 2010; Duarte, et al., 2014). Contrary to our expectations, no clear signs of neurodegeneration or mitochondrial dysfunction were observed. These findings suggest that at 18 months of age the aging process in the Dark Agouti rats has only begun. The observed regional differences hint at differential rates of aging process between cortical and sub-cortical brain area.

Unexpectedly, various alterations were correlated with the number of NMR examinations. Even if no clear explanation can be presented, these observations should be considered cautiously for future longitudinal MRS studies at UHF. These external effects to the process of aging complicate the validation of the DA rat as a model of aging. Nevertheless, the hallmarks of brain aging reported in humans were indeed observed, validating a priori its use for future aging studies.
4. Preliminary $^{31}$P MRS Data in the Aging Rat Brain

Phosphorus MRS allows to study of in vivo brain energy metabolism by directly measuring key high-energy phosphate compounds such as adenosine-triphosphate (ATP) and phosphocreatine (PCr). Moreover, there is a growing interest in the study of changes in phosphomonoesters (PMEs) and phosphodiesters (PDEs) levels in neurodegenerative or psychiatric afflictions (Mecheri, et al., 1997; Wijnen, et al., 2010). Similarly to the work presented in chapter 3, the main goal of this work was to develop quantitative in vivo $^{31}$P MRS data at 17.2 T. $^{31}$P Spectra were acquired using a FID-OVS sequence using BISTRO bands for localization (Luo, et al., 2001) from aging Dark Agouti rats. Spectra decomposition was done in the frequency domain using LCModel (Provencher, 1993; Deelchand, et al., 2012) and quantification was performed using brain tissue water as a reference of concentration as proposed by Bottomley et al. (Bottomley, et al., 1996). As an internal reference of concentration, $^1$H water signal was acquired from the same volume using a similar localization scheme. Furthermore, the $T_1$ relaxation times of phosphorylated metabolites were measured using the single inversion recovery method. The estimated $T_1$ times were compared to the extrapolated $T_1$ values as predicted by (Lu, et al., 2014).

4.1. Materials and Methods

4.1.1. Experimental set-up

In vivo experiments were performed on the previously studied cohorts of Dark Agouti rats (Janvier Labs, Le Genest-Saint-Isle, France): C0: 6 young rats (278 ± 13 g, 5 months old); C1: 6 middle-aged rats (333 ± 37 g, 17 months old); and C2: 3 elderly rats (360 ± 37 g, 21 months old). Rats were anesthetized during the experiments with 1.0-1.5% isoflurane in pure O$_2$. Respiration rate was monitored (40–60 breaths/min) and body temperature was held constant (37.5 ± 0.5° C) using a warm-water circuit for the whole duration of the experiment. The head was stereotaxically restrained by a bite bar and ear pins. The study protocol was approved by the Committee on the Ethics of Animal Experiments of the Commissariat à l’Energie Atomique (CETEA, Permit Number: ID 12-058).

The study was performed on a horizontal 17.2 T MRI scanner (Biospec, Bruker BioSpin, Ettlingen, Germany) equipped with an actively shielded gradient system capable of delivering gradients up to 1 T/m. MRS data acquisitions were performed using a $^1$H/$^{31}$P dual-resonance coil consisting of a 20-mm single-loop $^{31}$P surface coil and a butterfly $^1$H geometrically decoupled surface coil (RAPID Biomedical GmbH, Rimpar, Germany) operating respectively at 295.587 MHz and 730.195 MHz. A dedicated home-made holder was designed and built to fit properly.
the $^1$H/$^{31}$P RF coil with respect to the animal head pins (figure 4.1). Since the tuning and matching of both transceivers was fixed, the reflected power for each channel was checked and noted using the “wobble” function in Paravision 5.1 (figure 4.2) at the beginning of each experiment in order to account for loading and transmission differences between experiments.

Figure 4.1. Dual $^1$H/$^{31}$P coil and a dedicated home-made holder used for $^{31}$P MRS experiments.

Figure 4.2. Tuning and matching of the $^{31}$P transceiver using the wobble function in Paravision 5.1. In this example, the reflected power at the Larmor frequency is 13.4 %. Such values were used for quantification.

4.1.2. MRI data acquisition

For positioning, reference images were acquired using an axial rapid acquisition with relaxation enhancement (RARE) sequence covering the entire brain (TE/TR = 20/3000 ms, slice thickness: 0.5 mm, 24 slices, in-plane resolution: 180x180 µm²).
4.1.3. MRS data acquisition

First- and second-order shims were adjusted by acquiring a B\(_0\) field map and using Bruker’s MAPSHIM routine on a large VOI (4.5 x 7 x 7 mm\(^3\)) centered on the brain. A subsequent first-order shim correction using Paravision automated procedure was applied on the same VOI, as explained in section 2.1.3. The average water linewidth measured on the VOI was 28 ± 2 Hz.

![Axial View](image)

![Coronal View](image)

![Sagittal View](image)

Figure 4.3. \(^{31}\)P MRS VOI definition using BISTRO bands

8 manually-placed OVS BISTRO bands were used to saturate signal contributions from outside of the rat brain. Positioning was done based on \(T_2\)-weighted anatomical images (axial) and scout \(T_2^*\)-weighted images (coronal, sagittal).

\(^{31}\)P MR Spectra were acquired with a pulse-acquisition sequence (100 µs square pulse, 12.8 kHz excitation spectral width, 1024 points, 15 kHz receiver bandwidth, TR = 2 s, 906 averages). The VOI was defined manually by a set of 8 BISTRO bands (HS8 adiabatic pulses: 3 ms duration, 6.6 kHz, from 2 to 6 mm thick) using a hyperbolic secant envelope of 8 pulses (Luo, et al., 2001; Silver, et al., 1984). As a result, the selected VOI encompassed most of the brain (figure 4.3). The RF pulses carrier frequency was set to the PCr resonance frequency. The resulting chemical shift
artifact for the OVS bands caused a large spatial displacement of 30% between PCr and PE (6.78 ppm) and of -34% between PCr and ATPa (-7.35 ppm).

From a subset of animals consisting of 3 old, 3 middle-aged and 2 young rats, 31P T1-weighted spectra were acquired using the same localization method (TR = 2s, 320 averages). T1-weighting was introduced by incorporating a non-selective HS8 AFP inversion pulse (3 ms, 6.6 kHz) prior to the FID-OVS localization scheme. In total, 10 T1-weighted IR spectra were acquired (TI = 4, 176, 250, 500, 750, 1000, 1250, 1500, 1750 and 2000 ms). To conserve the same steady-state magnetization, all 10 IR spectra were acquired with a recovery time of 2 s.

A non-suppressed water spectrum was acquired from the same VOI and FID-OVS sequence (200 µs square pulse, 1024 points, TR = 10 s, 8 averages) for the purpose of water referencing (Bottomley, et al., 1996). Additional water spectra were acquired with increasing acquisition delays (0.11, 10, 20, 30, 40, 50, 80 and 100 ms) in order to determine CSF contributions based on T2* relaxation times differences, similarly to the method considering T2 decay as described in section 2.5.1.

To estimate the scaling factor CPH between 1H and 31P channels (section 2.5.2), in vitro measurements using a phosphoric acid sample (50 mM concentration, diluted monopotassium phosphate, pH = 7.4, room temperature) were performed immediately after each experimental session using the same coil and VOI positioning.

In order to evaluate the impact of the CSDA on our 31P data, 31P MR spectra (604 averages) were acquired from two rats with a carrier frequency shifted by -7.5, -5, -2.5, 0, +2.5, +5 and +7.5 ppm with respect to PCr. Moreover, two 31P MR spectra (604 averages) were also acquired with and without OVS bands.

4.1.4. MRS data analysis

4.1.4.1. LCMModel parameterization

Pre-processing steps consisted in zero-filling to 2048 data points, Fourier transformation and zero- and first-order phase correction. Spectra were analyzed using LCMModel 6.2.

The basis set was simulated as described in section 2.4. Published chemical shift values and the homonuclear J-coupling constants were considered (Jung, et al., 1997; Jensen, et al., 2002; Lu, et al., 2013).

For the LCMModel analysis, the IR spectra were phased such that the PCr resonance was positive. Consequently, the basis-set used for analyzing T1-weighted data included inverted spectra for the ATP resonances due to their shorter T1 relaxation times compared to PCr.

The LCMModel basis consisted of a total of 12 metabolites as shown in Table 4.1 along with their chemical shifts and allowed frequency shifts (SDSH). Due to their differences in T1 and T2 values (Lei, et al., 2003; Remy, et al., 1987), the three separate resonances of ATP were separately accounted for. Due to the magnitude of the chemical shift artifact, the ATPβ resonance was rarely
observed and was not considered for the LCModel analysis. For the same reason, the analysis window was limited to the [10, -10] ppm range. Adenosine-monophosphate (AMP) was not included because its concentration was expected to be close to noise levels (513 µmol/L) (Pollesello, et al., 1995).

LCModel was originally implemented solely for the analysis of $^1$H spectra and therefore hidden control parameters needed to be adjusted so that X-nuclei spectra could be handled (Henry, et al., 2003; Deelchand, et al., 2012). In addition to the declaration of standard parameters such as the operating frequency, the number of points and dwell time (HZPPPM, NUNFIL, DELTAT), the following parameters were modified:

- **XSTEP**: Increment between the numbers of the displayed x-axis. It was set to 1 to allow LCModel to display the results.
- **DKNTMN**: The knot spacing for the spline baseline was set to 2, allowing to obtain a baseline which could account for large first-order phase variations without affecting the quantification of metabolites and greatly increasing the calculation speed.
- **DESDSH**: Default allowed frequency shifts between the in vivo data and the basis set elements. It was set to 0.01 (ppm) instead of the standard 0.004 value. Specific frequency shifts were set for certain metabolites (Table 4.1) using the (N-, CH-, AL-) SDSH commands.
- **DESDT2**: The variation of the expected spectral linewidths due to the increased $1/T_2$ at higher magnetic fields. Its value was set to 12. The associated EXT2 parameter was not modified. The improvement of the LCModel fit, notably for PCr and ATPγ, can be observed in figure 4.4.
- **FCALIB**: The calibration factor was used to scale the measured $^{31}$P signal to absolute concentration units, by adjusting it with respect to the $C_{PH}^[W]/S_W$ term as shown in eq. (2.6).
<table>
<thead>
<tr>
<th>Metabolite</th>
<th>PPM</th>
<th>SDSH</th>
</tr>
</thead>
<tbody>
<tr>
<td>Adenosine Triphosphate</td>
<td>ATP</td>
<td>α</td>
</tr>
<tr>
<td></td>
<td></td>
<td>β</td>
</tr>
<tr>
<td></td>
<td></td>
<td>γ</td>
</tr>
<tr>
<td>Fructose-6-phosphate</td>
<td>F6P</td>
<td></td>
</tr>
<tr>
<td>Glucose-1-phosphate</td>
<td>Glc1P</td>
<td></td>
</tr>
<tr>
<td>Glucose-6-phosphate</td>
<td>Glc6P</td>
<td></td>
</tr>
<tr>
<td>Glycerol-1-phosphate</td>
<td>Gly1P</td>
<td></td>
</tr>
<tr>
<td>Glycerol-3-phosphorylcholine</td>
<td>GPC</td>
<td></td>
</tr>
<tr>
<td>Glycerol-3-phosphorylethanolamine</td>
<td>GPE</td>
<td></td>
</tr>
<tr>
<td>Inorganic phosphate</td>
<td>Pi</td>
<td></td>
</tr>
<tr>
<td>Phosphocreatine</td>
<td>PCr</td>
<td></td>
</tr>
<tr>
<td>Phosphorylcholine</td>
<td>PCh</td>
<td></td>
</tr>
<tr>
<td>Phosphorylethanolamine</td>
<td>PE</td>
<td></td>
</tr>
<tr>
<td>NAD+NADP</td>
<td>NADH</td>
<td></td>
</tr>
</tbody>
</table>

Table 4.1 Metabolites included in the LCModel basis for the analysis of our $^{31}$P spectra. Chemical shifts values are relative to the PCr resonance. The SDSH parameters used in LCModel determines the allowed frequency shift for a given metabolite. Although GPC, GPE and Pi were attributed with elevated SDSH values, resonance frequency shifts were only observed for Pi.

Figure 4.4. LCModel $^{31}$P fits using two parameter sets. The residual noise (red) was used to evaluate the quality of the fit. Using a constrained parameterization consisting of the default value of DESDT2 and SDDEGP = 0.05, strong residual signal was observed at the base of the PCr and ATP, resonances and the Pi and PE were often incorrectly fitted. Setting DESDT2 = 12 and SDDEGP = 4 largely reduced the residual noise.
4.1.4.2. Water referencing

The FCALIB variable for each experiment was determined by the $C_{PH}$ conversion factor. The precise formula for the calculation of $C_{PH}$ based on in vitro measurements was the following:

$$C_{PH} = \frac{\overline{S_{W}}}{\overline{S_{P}}} \cdot \frac{[P_v]}{[W_v]} = \frac{S_{W_v}/Load_{W_v}}{S_{P_v}/Load_{P_v}} \cdot \frac{[P_v]}{[W_v]}$$  \hspace{1cm} (4.1)

where $[P_v]$ and $[W_v]$ were the phantom apparent water and phosphorous concentrations; $\overline{S_{W}}, \overline{S_{P}}$ were the measured phantom water and Pi signals compensated for differences in matching conditions using the parameters $Load_{W_v}$ and $Load_{P_v}$, obtained prior to the MRS sequence using the “wobble” function of Paravison 5.1 as shown in figure 4.2. The FCALIB was then derived for each in vivo experiment from the following formula:

$$\text{FCALIB} = C_{PH} \cdot \frac{[W]}{S_{W}} \cdot \frac{Load_{W}}{Load_{P}}$$  \hspace{1cm} (4.2)

where $[W]$ were the assumed, apparent brain water content, $S_{W}$ the water signal, $Load_{W}$ and $Load_{P}$ the reflected power values recorded just before launching the MRS sequence. An additional constant factor ($= 464673$) was used for all LCModel analysis to account for the calibration of the simulated basis-set, the number of averages and receiver gain differences.

4.1.4.3. $T_1$ relaxation times estimation

To estimate the $T_1$ relaxation times, a 3-parameter fit consisting of a mono-exponential function was used to fit the equilibrium magnetization and the $T_1$ relaxation times of each metabolite using a non-weighted Levenberg-Marquardt algorithm and leaving the actual excitation angle as a free parameter. $T_1$ estimation was done separately for each animal. Cramer-Rao Lower Bounds (CRLB) were used as error estimator (section 2.4.4) and all data points presenting values higher than 25% were discarded. The coefficient of determination ($R^2$) was used to evaluate the quality of the fits.

4.1.4.4. Statistical analysis

A linear-regression analysis considering “Rat” as a random effect and “Age” as a predictor was used to assess the statistically significant changes of metabolite concentrations and $T_1$ times with age. The analysis was done using R, version 3.1.2 (R Core Team, 2014). No corrections were applied to reduce type I errors. Statistically significant level was set as $P < 0.05$. 

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4.2. RESULTS

4.2.1. Variability of $C_{PH}$

The $C_{PH}$ conversion factor was calculated based on the apparent water and inorganic phosphate concentrations *in vitro* (equation 4.1). $T_1$ relaxation times of water and phosphorous in the 50 mM sample were measured with the same method that was used *in vivo*. The measured water $T_1$ was 2.766 s, in agreement with the value found for injectable water (2.966 s) using LASER as shown in chapter 3. The measured phosphorous $T_1$ time was 5.224 s. After each scanning session, *in vitro* measures were done using the 50 mM phantom to determine $C_{PH}$ considering the exact same VOI geometry and position while trying to keep the same positioning of the coil relative to the sample. Since the RF power calibration for the $^{31}$P channel was done by maximizing the measured signal (section 2.1.1.1) using a TR of 2s, equations (1.18) and (1.19) were used to estimate the Ernst angle and steady-state magnetization and determine an apparent $[P_v]$ concentration. The mean $C_{PH}$ was 0.00238 ± 0.00036, similar to the variation of 0.0018 ± 0.0003 found in humans (Bottomley, Atalar et Weiss 1996).

4.2.2. Assessment of the CSDA-related attenuation of $^{31}$P metabolites

Even if the $^{31}$P spins were excited using a non-selective RF pulse, each BISTRO band was saturating a different slab in space for each metabolite of interest leading to an effective different volume of acquisition for each metabolite. The impact of this chemical shift artifact was experimentally assessed *in vivo* by acquiring identically parameterized spectra with a shifted carrier frequency. By modeling the variation of the PCr resonance according to the frequency shift, one could estimate the attenuation of the different metabolites in the $^{31}$P MRS data. This approach is only phenomenological and assumes a homogenous distribution of the $^{31}$P metabolite in the brain.

Figure 4.5 (left) shows a set of PCr resonances for one rat, positioned according to the frequency shift used for their acquisition. Figure 4.5 (right) shows the PCr maximal intensity and areas plotted with respect to the frequency shift after normalization to the “true”, on-resonance values. The PCr area variation with respect to the frequency offset (figure 4.5, right, black line) could be approximated by a linear function of the frequency shift in ppm (figure 4.5, right, blue line):

$$A(\Delta f) = 1 + 0.044*\Delta f$$  \hspace{1cm} (4.3)

Since LCModel metabolite concentrations are calculated from the measured area for each metabolite, the linear approximation $A(\Delta f)$ of the CSDA-related attenuation profile was used to correct for the differences in localization and sensitivity for each metabolite.
Figure 4.5. CSDA-related attenuation observed in vivo for the PCr signal. PCr resonances (left) and normalized intensities and area (right) are plotted as a function of the carrier frequency shift. The shown spectra and values correspond to the LCModel fitted values for PCr. Values shown in the right figure are from two rats. The observed PCr area can be approximated as a linear function of the frequency shift (blue line, R² = 0.988).

4.2.3. $^{31}$P neurochemical profile

Figure 4.6 shows a representative spectrum and its LCModel fit, residual noise, baseline and the individual metabolite contributions. The ATP α and γ resonances showed distinct linewidths and intensities, notably due to differences in relaxation times but also to the magnitude of the CSDA at -2.48 and -7.5 ppm. The spline baseline generated by LCModel (DKNTMN = 2) permitted to obtain a flat spectrum without effecting the quantification of any metabolite. The metabolites that were systematically detected with CRLBs below 10% were GPC, PE, PCh, PCr, Pi and the ATP resonances. GPE was also systematically detected but the average CRLB was 17%. The increase of the SDT2 parameter greatly reduced the residual noise levels (figure 4.4) for PCr and ATP but there was an overall increase in the CRLB values of metabolites by 1 to 2%.
Figure 4.6. Spectral decomposition of a $^{31}$P spectrum acquired in the rat brain at 17.2 T. LCModel spectral decomposition of a typical spectrum acquired from the rat brain using BISTRO bands for localization (figure 4.3). 12 different metabolites were considered in the basis-set although F6P was seldom detected. No filtering was applied and all spectral elements shown are scaled identically. *The $\alpha$ and $\gamma$ ATP resonances were considered separately in the basis sets.

4.2.4. T$_1$ relaxation times

Figure 4.7 shows a set of T$_1$-weighted spectra (left) and their LCModel fit (right). Due to their shorter T$_1$ times, the ATP resonances were inverted with respect to the remaining metabolites already at TI = 176 ms. At TI = 750 ms, the remaining metabolites were zero-crossing. The CRLBs of PCr ranged from 1 to 11% over the set of T$_1$-weighted spectra. Other metabolites that systematically showed CRLB below 25% were GPC, Glc6P, PCh and the ATP moieties. The elevated noise levels for the T$_1$-weighted spectra made it difficult to accurately fit metabolites with low concentrations.

The mean T$_1$ relaxation times estimated from the 8 rats are shown in table 4.2 with their corresponding mean coefficients of determination ($R^2$). Only T$_1$ fits presenting a $R^2 > 0.6$ were accepted. The T$_1$ times of PCr found at different ages were: $1110 \pm 250$ (21 months old, $n = 3$), $850 \pm 135$ (17 months old, $n = 3$) and $1110 \pm 170$ ms (5 months old, $n = 2$).
Figure 4.7. $^{31}$P $T_1$-weighted spectra acquired in the rat brain at 17.2 T
Set of $T_1$-weighted spectra (left) and their LCModel fit (right) acquired from the rat brain with the IR method with inversion times ranging from 3 to 2000 ms. A recovery time of 2 s was kept for all spectra. For displaying purposes, a Gaussian smoothing of 8 Hz was applied on the data in the frequency domain. The most notable features was the inversion of the ATP resonances already at $TI = 176$ ms followed by the inversion of the remaining metabolites between 500 and 750 ms.

<table>
<thead>
<tr>
<th>Metabolite</th>
<th>$T_1$ (ms)</th>
<th>$R^2$</th>
<th>Measures</th>
</tr>
</thead>
<tbody>
<tr>
<td>GPC</td>
<td>1080 ± 210</td>
<td>0.78 ± 0.13</td>
<td>5</td>
</tr>
<tr>
<td>GPE</td>
<td>1170</td>
<td>0.64</td>
<td>1</td>
</tr>
<tr>
<td>Glc6P</td>
<td>1030 ± 185</td>
<td>0.79 ± 0.12</td>
<td>7</td>
</tr>
<tr>
<td>PE</td>
<td>1000 ± 275</td>
<td>0.84 ± 0.1</td>
<td>3</td>
</tr>
<tr>
<td>PCr</td>
<td>1010 ± 210</td>
<td>0.95 ± 0.02</td>
<td>8</td>
</tr>
<tr>
<td>NADH</td>
<td>1800</td>
<td>0.69</td>
<td>1</td>
</tr>
<tr>
<td>Pi</td>
<td>1110</td>
<td>0.86</td>
<td>1</td>
</tr>
<tr>
<td>PCh</td>
<td>1160 ± 140</td>
<td>0.87 ± 0.09</td>
<td>3</td>
</tr>
<tr>
<td>ATP$_α$</td>
<td>280 ± 70</td>
<td>0.72 ± 0.10</td>
<td>3</td>
</tr>
<tr>
<td>ATP$_γ$</td>
<td>390 ± 100</td>
<td>0.77 ± 0.04</td>
<td>5</td>
</tr>
</tbody>
</table>

Table 4.2. $T_1$ relaxation times of $^{31}$P metabolites measured in the rat brain in vivo at 17.2 T.
Data from rats of all ages were pooled. LCModel results presenting CRLB > 25 % were discarded.
*Only results showing a coefficient of determination higher than 0.6 were considered.

4.2.5. Metabolite concentrations

Applying the quantification method described in section 2.5.2, the concentrations of 12 metabolites were calculated. Based on the bi-exponential fit of the $T_2^*$-weighted unsuppressed water spectra, the fractional CSF content was estimated as merely 1.7 ± 0.6 % of the total measured water signal ($R^2 = 0.999$) and it was not observed to change with age. A mean brain tissue water content of 79.55 % was assumed, based on published values (Lin, et al., 2000). The $T_1$ value of the gray matter shown in table 3.1 was used to correct for $T_1$-weighting on the reference water spectrum, although it had only a minor impact (below 2%) due to the long TR (= 10 s) employed for its acquisition. The water apparent concentration was 43.9 M. Corrections for $T_2$-weighting were omitted due to the short acquisition delays used. Metabolites were corrected
for $T_1$-weighting using the values shown in table 4.2. For Gly1P, F6P and Glc1P the $T_1$ value of PCr was used. $T_1$-weighting was corrected by assuming a FA of $82^\circ$, corresponding to the Ernst’s angle of the mean PCr $T_1$ value (1010 ms) with TR = 2 s.

Table 4.3 summarizes the metabolite concentrations for each of the three rat cohorts and their pooled results provided by LCModel and their values after correction for $T_1$ relaxation effects and CSDA-attenuation. The number of measurements, CRLBs and values found in the literature are also shown.

<table>
<thead>
<tr>
<th>Metabolite</th>
<th>5 Months</th>
<th>17 Months</th>
<th>21 Months</th>
<th>All</th>
<th>Corrected for $T_1$ and CSDA</th>
<th>Measures$^a$</th>
<th>CRLB$^b$</th>
<th>Literature</th>
</tr>
</thead>
<tbody>
<tr>
<td>Gly1P</td>
<td>0.33 ± 0.14</td>
<td>0.30 ± 0.04</td>
<td>-</td>
<td>0.32 ± 0.11</td>
<td>0.29 ± 0.10</td>
<td>6</td>
<td>20</td>
<td></td>
</tr>
<tr>
<td>GPC</td>
<td>0.74 ± 0.31</td>
<td>0.73 ± 0.23</td>
<td>0.51 ± 0.14</td>
<td>0.70 ± 0.26</td>
<td>0.73 ± 0.27</td>
<td>18</td>
<td>7</td>
<td>0.6-1.05-1.14$^H$</td>
</tr>
<tr>
<td>GPE</td>
<td>0.26 ± 0.10</td>
<td>0.28 ± 0.11</td>
<td>0.19 ± 0.02</td>
<td>0.25 ± 0.10</td>
<td>0.27 ± 0.10</td>
<td>17</td>
<td>17</td>
<td>0.74-0.83$^H$</td>
</tr>
<tr>
<td>F6P</td>
<td>0.29 ± 0.16</td>
<td>-</td>
<td>-</td>
<td>0.29 ± 0.16</td>
<td>0.26 ± 0.15</td>
<td>3</td>
<td>22</td>
<td>0.012, 0.050</td>
</tr>
<tr>
<td>Glc1P</td>
<td>0.19 ± 0.05</td>
<td>0.25 ± 0.15</td>
<td>0.22</td>
<td>0.23 ± 0.12</td>
<td>0.23 ± 0.12</td>
<td>9</td>
<td>20</td>
<td>0.02</td>
</tr>
<tr>
<td>Glc6P</td>
<td>0.06 ± 0.12</td>
<td>0.06 ± 0.10</td>
<td>-</td>
<td>0.05 ± 0.10</td>
<td>0.20 ± 0.06</td>
<td>4</td>
<td>21</td>
<td>0.09, 0.19, 0.8$^H$</td>
</tr>
<tr>
<td>PE</td>
<td>0.97 ± 0.39</td>
<td>1.09 ± 0.30</td>
<td>0.69 ± 0.06</td>
<td>0.97 ± 0.34</td>
<td>0.87 ± 0.30</td>
<td>18</td>
<td>8</td>
<td>1.7, 1.0-20, 0.5-1$^H$</td>
</tr>
<tr>
<td>PCr</td>
<td>4.87 ± 1.90</td>
<td>4.29 ± 1.17</td>
<td>3.06 ± 1.18</td>
<td>4.34 ± 1.60</td>
<td>5.08 ± 1.87</td>
<td>18</td>
<td>2</td>
<td>4.2-4.5, 2.4-5$^H$</td>
</tr>
<tr>
<td>NADH</td>
<td>0.13 ± 0.05</td>
<td>0.12 ± 0.06</td>
<td>0.08 ± 0.02</td>
<td>0.11 ± 0.06</td>
<td>0.26 ± 0.13</td>
<td>11</td>
<td>18</td>
<td>0.36$^H$</td>
</tr>
<tr>
<td>Pi</td>
<td>0.73 ± 0.33</td>
<td>0.71 ± 0.23</td>
<td>0.53 ± 0.10</td>
<td>0.69 ± 0.27</td>
<td>0.68 ± 0.26</td>
<td>18</td>
<td>9</td>
<td>0.94$^H$, 1.63-2.8</td>
</tr>
<tr>
<td>PCh</td>
<td>0.44 ± 0.19</td>
<td>0.42 ± 0.14</td>
<td>0.26 ± 0.07</td>
<td>0.40 ± 0.16</td>
<td>0.38 ± 0.16</td>
<td>18</td>
<td>9</td>
<td>0.3, 0.46$^H$</td>
</tr>
<tr>
<td>ATP$_a$</td>
<td>0.54 ± 0.24</td>
<td>0.72 ± 0.21</td>
<td>0.38 ± 0.24</td>
<td>0.59 ± 0.25</td>
<td>0.90 ± 0.38</td>
<td>18</td>
<td>11</td>
<td>-</td>
</tr>
<tr>
<td>ATP$_r$</td>
<td>1.21 ± 0.57</td>
<td>1.19 ± 0.39</td>
<td>0.73 ± 0.27</td>
<td>1.12 ± 0.48</td>
<td>1.30 ± 0.56</td>
<td>18</td>
<td>7</td>
<td>2.3-3.3, 2.19-3.8$^H$</td>
</tr>
<tr>
<td>Pi/PCr</td>
<td>0.15</td>
<td>0.16</td>
<td>0.18</td>
<td>0.16</td>
<td>0.14</td>
<td>-</td>
<td>-</td>
<td>0.27-0.53, 0.32</td>
</tr>
<tr>
<td>ATP$_r$/PCr</td>
<td>0.24</td>
<td>0.27</td>
<td>0.25</td>
<td>0.25</td>
<td>0.26</td>
<td>-</td>
<td>-</td>
<td>0.45-0.71,0.51-1.27$^H$</td>
</tr>
</tbody>
</table>

Table 4.3. $^{31}$P MRS metabolite concentrations and ratios at 17.2 T in the rat brain in vivo. Metabolite concentrations (mM) are shown for the separate rat cohorts at ages 5, 17 and 21 months old and their pooled data. Also listed are the concentrations after $T_1$ and CSDA corrections [eq. (4.3)]. $T_1$ relaxation times shown in table 4.2 were used for $T_1$-weighting correction and PCr values were used for Gly1P, F6P and Glc1P. CSDA correction was applied using the ppm values shown in table 4.1. Metabolite ratios were calculated for each individual spectrum before averaging.

$^a$Occurrence of a metabolite with CRLB below 25 among 18 spectra.

$^b$Mean CRLB for the considered occurrences.

$^H$Values reported for human brain studies (Zhu, et al., 2012; Hetherington, et al., 2001; Zhu, et al., 2015; Jensen, et al., 2002; Blümli, et al., 1999). Other values were reported from rodent studies (Plaschke, et al., 1993; Mans, et al., 1994; Erekinska, et al., 1989; Mlynárik, et al., 2012; Smart, et al., 1994). The underlined values are those measured here using $^1$H MRS (Table 3.3).
4.3. **DISCUSSION**

In this preliminary study, we present the first $^{31}$P NMR data acquired in the rat brain *in vivo* at 17.2 T. 12 metabolites were quantified using water as an internal reference of concentration and the T1 relaxation times of 10 metabolite moieties were measured ($R^2 >0.6$). Localization was achieved employing BISTRO bands.

4.3.1. **Signal localization and CSDA-attenuation**

In order to appreciate the effectiveness of our localization using BISTRO bands, two spectra were acquired from the same rat with and without OVS bands. Figure 4.8 compares the FID spectrum and the LCModel fit of the FID-OVS spectrum corrected for the CSDA-attenuation. The ratio between the PCr signal areas was 12.4. Also, a 5.2 difference was found between the PCr maximum values. As can be appreciated from the slender linewidth of PCr in figure 4.8, the application of OVS drastically reduced the contributions outside of the VOI, where $B_0$ shimming had taken place. Due to the strong difference in PCr areas, $^1$H FLASH images were acquired *in vitro* using the identical BISTRO module as used for the FID-OVS scan. The scan parameters and images are shown in figure 4.9. A difference between the nominal cubic volume and the actual imaged volume using HS8 pulses was found, with a factor of 3.2. This factor was reduced to 1.4 when Bruker’s HS (3 ms, 6.6 kHz) pulse was applied, demonstrating the impact of the transition bandwidth of the HS8 pulses used for the VOI selection. These results partially explain difference between the PCr maximum values of the localized and non-localized spectra (figure 4.8, top).

This effect had a major impact on the SNR of all $^{31}$P MRS acquisitions and should be taken into account for future work, by modifying the declared bandwidth of the HS8 pulses to account for the transition bandwidth. However, using more selective adiabatic RF pulse envelopes than the HS8 pulses with the same bandwidth might be limited by our broad-band amplifier power output since we already had to work at up to 2 dB of attenuation from the maximal power.
Figure 4.8. Effect of OVS localization on $^{31}$P spectra in the rat brain. OVS-localized (black) and non-localized (red) $^{31}$P spectra acquired from the same rat (604 averages). (Top) The raw spectra showed a 5.21 and a 12.4 fold difference between the PCr maximum height and area, respectively. (Bottom) The fit results from LCModel are shown after normalization by the PCr height and corrected for the frequency-dependent CSDA-attenuation [eq. (4.3)]. A ratio $[\text{ATP}/\text{PCr}]$ of 0.29 and 0.39 was found for the localized and the non-localized acquisitions, respectively. Baseline correction was applied on both spectra. The ATP resonance was not fitted in this example and to correctly fit the non-localized spectrum, the DEEXT2 parameter was set to 12.

Figure 4.9. Transition bandwidth effects of HS8 and Paravision’s HS pulses
Relative sizes of the nominal volume selected by the BISTRO bands and the actual imaged volume using a (A) HS8 pulse and a (B) paravision’s HS pulse designed for saturation on a FLASH imaging sequence (TR/TE = 1200/6 ms, FOV = 40x40 mm$^2$, OVS band thickness = 8 mm). The purple bands represent the nominal sizes and positioning of the suppression bands (8-mm thick) used for this experiment. The ratios of the selected ROI and the nominal area are 0.46 and 0.81 for the HS8 and HS pulses, respectively.
To understand the attenuation profile shown in figure 4.5, we must consider the RF coil sensitivity profile, the homogeneity of the $B_0$ field and the volume of distribution of PCr. If we assume that the PCr signal originates from the brain only and that our “on-resonance” VOI includes most of it (figure 4.3), the spatial mismatch should lead to an attenuation being a function of the cube of the frequency shift. For “off-resonance” volumes, the VOI is spatially-shifted and present stronger $B_0$ field irregularities and consequently stronger $T_2^*$ attenuation and larger linewidths. The spatial displacement at off-resonance values of $\pm 7.5$ ppm for a 4-mm thick OVS band is 1.36 mm. This shift is nevertheless comparable to the spatial mismatch due to the transition bandwidth of our 3-ms long HS8 pulses (2.08 kHz transition bandwidth, 1.26 mm). As a consequence, the volume suppressed by the BISTRO bands at off-resonance values were still reducing extracranial and muscle contributions. Therefore, the CSDA-attenuation profile should be dominated by the inhomogeneous sensitivity of the $^{31}$P coil. From figure 4.5, the VOI seems to move towards the RF coil for positive frequency shifts whereas for negative shifts, the signal is originating from a deeper VOI. In this regards, our experimental set-up and the estimated attenuation profile was probably very sensitive to the distance between the coil, the center of our VOI and the positioning of the top and inclined OVS bands (figure 4.3). It is therefore possible that part of the variability observed for our metabolites concentrations and our low [ATP] concentrations is related to the simplistic model of our CSDA-attenuation profile.

The CSDA also had a major impact on the detection of the ATPγ and ATPα resonances, reducing their intensity by 11% and 33%, as estimated from equation (4.3). The $A(\Delta f)$ function was used to correct for the bulk effect of CSDA but residual errors are expected since the OVS positioning varied slightly between rats. For future work, it will be critical to reduce the CSDA, notably by reducing the thickness of the OVS, reducing the number of OVS bands to cover only critical regions or by using shorter HS8 pulses to increase the gradient strength. This solution may indeed provide better results because the $B_1$ field intensity is stronger close to the coil, allowing for shorter pulses and the coil sensitivity is also higher, making it the dominant noise contributor and therefore the most important region where signal suppression is required.

To further investigate if our low ATP concentrations are due to an inexact attenuation profile, the $[\text{ATP}_\gamma]/[\text{PCr}]$ ratio for the non-localized spectrum was calculated. At 0.39 without and 0.35 with correction for $T_1$-weighting, both values are indeed higher that the $[\text{ATP}_\gamma]/[\text{PCr}]$ ratio measured from the OVS-localized spectrum (0.26). Although these values remain lower than those found in the literature (Table 4.3), strong signal contributions were originated from outside the localized volume, presumably from muscle and extracranial lipids. Since the muscle $[\text{ATP}]/[\text{PCr}]$ ratio is 0.22 (Bottomley, et al., 1996) and the FOV of the $^{31}$P was possibly quite large (20-mm-diameter loop), it is possible that our non-localized spectra PCr and ATP values were strongly altered by signal contributions from the scalp and jaw muscles.
4.3.2. Spectral decomposition using LCModel

Using LCModel software for the analysis of $^{31}$P spectra proved to be a simple and reliable approach and besides the recently published work of (Deelchand, et al., 2015), no other LCModel applications in $^{31}$P have been reported. LCModel managed to readily account for strong baseline distortions (notably for $^{31}$P spectra acquired with no OVS) and provides a method for automated and operator-independent analysis. The starting point for the initial first-order phase was found to be a critical parameter for correctly fitting lowly concentrated metabolites. The initial phasing parameters were found to be constant for a given pulse sequence and ADC filter. The main drawback of using LCModel for the analysis of $^{31}$P spectra is its relatively cumbersome handling for frequency shifts, which are relatively common in $^{31}$P spectra due to variations in the chemical environment, such as pH. Setting metabolite-specific SDSH values may solve this problem but only when individual, non-overlapping resonances are affected. For ATP, this may not be a solution as the three resonances may not necessarily show the same chemical shift change (notably in phantom studies) and so the individual resonances must be accounted for by dividing the three ATP resonances into separate elements in the basis-set or using the LCModel built-in Gaussian functions.

To our knowledge, heteronuclear J-coupling constants were not available for the metabolites included table 4.1. Recently, Deelchand et al. has included these values (Deelchand, et al., 2015) and for future work they should be included in the simulation of the spectra in the basis set.

4.3.3. $T_1$ relaxation times

$T_1$ relaxation times of PCr, ATP$_\alpha$ and ATP$_\gamma$ resonances were estimated to be $1010 \pm 210$, $280 \pm 70$ and $390 \pm 100$ ms respectively. From the excellent study by Lu et al. (Lu, et al., 2014) looking at the relaxation times of $^{31}$P metabolites from 4 to 16.4 T, one can calculate the $T_1$ values for these resonances at 17.16 T based on an extrapolation formula similar to eq. (3.2). The corresponding $T_1$ times are respectively 1430, 530 and 800 ms, much longer than our results. These differences could be attributed to our lower SNR, to the shorter range of TI explored (0.003 to 2 s vs 0.01 to 20 s) or the shorter recovery-time used for our experiment (2 s vs 16 s). In order to increase the SNR of our $T_1$-weighted spectra, it could be envisaged to limit the number of OVS bands, retaining only the bands used to suppress the jaw muscles, which have strong PCr and ATP contributions (Bottomley, et al., 1996). The complete removal of the OVS bands could also be considered at the cost of reduced specificity assuming similar $T_1$ relaxation times for $^{31}$P metabolites in the brain and muscles. The strong baseline distortions could be easily managed by LCModel. The main drawback would be the increased linewidths, although at 17.2 T they only have a moderate impact in differentiating the individual metabolites, as it is shown in the raw, non-localized spectrum in figure 4.8.
4.3.4. Metabolite quantification

In this study, water was used as an internal reference of concentration. For this, a calibration factor $C_{PH}$ was calculated for each experiment. The variability that was observed (15%) was congruent with values found in similar studies (16%, (Bottomley, et al., 1996)). Two other common methods used for $^{31}$P referencing use either an external homonuclear reference close to the head or a phantom with similar brain conductive properties, geometry and coil loading with those of the volunteer. It has been shown that the internal water reference is two times less accurate (11% mean error) and reproducible (15% SD of error) than the other two methods (Buchli, et al., 1994). Indeed, a high variability was observed for all metabolites concentrations (mean variation of 34%), although SNR limitations had an important impact on the detection of metabolites with low concentrations. The theoretical framework of this method assumes an identical sensitivity profile for both channels, an assumption which may contribute to the variability of the quantification.

Despite this limitation, the concentration values that were found in this work were in agreement with published results (Table 4.3). However, the concentrations for GPC, GPE, Glc1P and ATP were found to be underestimated. In particular, our $[\text{ATP}_\gamma]/[\text{PCr}]$ ratio of 0.26 was lower than values reported for the dog (0.45), mouse (0.71) and human (0.51 to 1.27) brain (Hetherington, et al., 2001; Jensen, et al., 2002; Mlynárik, et al., 2012; Erecinska, et al., 1989). $T_1$-weighting correction was applied to all metabolites based on the calculated $T_1$ times (table 4.2). The PCr and ATP$_\gamma$ corrections corresponded to an increase in signal of 15 and 2 % from the uncorrected values, respectively. If instead, the predicted $T_1$ times at 17.16 T from Lu et al were used, the PCr and ATP$_\gamma$ concentrations would increase by a further 13 and 9 %, decreasing the $ATP_\gamma/\text{PCr}$ ratio by 5%. Our low ATP/PCr ratio could be associated to an increased $T_2^*$- differential weighting of ATP with respect to PCr, as their $T_2$ differences have been shown to be quite large (by ~ 20 fold) already at low magnetic fields (Brooks, et al., 1986) and for which no correction was applied.

The CSF fractional volume was found to be $1.72 \pm 0.59 \%$. Due to the large transition bandwidth of the HS8 OVS bands, it is most likely that this CSF contribution correspond to the ventricular CSF volume as the OVS were placed tangentially to the encephalon. In comparison, the CSF fractional volume in the human brain has been reported at 11%, most of it being extra-ventricular (Matsumae, et al., 1996). In the 2 and 7 months-old rat brain, the ventricular CSF fractional volume was respectively estimated as 1.06 and 1.22 % of the total brain volume (Tajima, et al., 1993; Sahin, et al., 2001; Chiu, et al., 2012), in agreement with our results (1.72 %). Furthermore, our measured CSF fraction was not observed to increase with age. This is in agreement with measurements done in rats from 3 to 20 months old, where the ventricular volume remained negligible compared to the brain volume (~2000 µL at 6 months of age (Tajima, et al., 1993)) increasing from 7 to 34 µL but the total CSF increased from 275 to 375 µL (Chiu, et al., 2012).
4.3.5. Comparison to $^1$H MRS data

The metabolites concentrations measured with $^{31}$P MRS, corrected for T$_1$-weighting and compensated for CS DA-attenuation, were in good agreement with the metabolite concentrations found in the Main region using $^1$H MRS. The GPC, PCr and PCh concentrations showed relative differences of 22, 4 and 27 % respectively, which are within experimental error. The PE concentration measured in $^{31}$P (0.87 mM) was half of that found in the $^1$H studies (1.7 mM), presenting similar CRLBs in both nuclei (8 % and 9 % for the $^{31}$P and $^1$H analysis, respectively). Interestingly, both metabolite concentrations are consistent with their various reported concentrations either in $^1$H or $^{31}$P MRS studies (Blüml, et al., 1999; Smart, et al., 1994). The relative difference may originate from the difference between both VOIs but also from the inaccuracy of some of our correction factors (T$_1$-weighting, CS DA attenuation).

4.3.6. Changes with age

Considering the small size of each cohort, it was difficult to determine statistically significant changes. Age-related metabolite concentration or T$_1$ relaxation changes could not be observed using $^{31}$P MRS and only statistically significant increases of [PE]/[PCr] with age were found. [GPC]/[PCr] showed an increase but it was above the significance threshold (P = 0.055). The PCr concentrations were observed to decrease with age but it was not significant either (P = 0.15).

The absence of age-related changes should be considered with caution, as measurement errors introduced by the CS DA, the quantification method intrinsic variations and the reduced number of elderly rats (n = 3) compromises our evaluations. Furthermore, no age-specific T$_1$ corrections were applied due to the reduced number of T$_1$ measurements and this may also play an important role as the results found in the $^1$H studies revealed strong T$_1$ variations due to age and the number of examinations.

4.4. CONCLUSION

The present work points out the main challenges for achieving $^{31}$P MRS quantification in the rat brain, exposing the shortcomings of our localization scheme using HS8 pulses for OVS. Despite these drawbacks, future $^{31}$P MRS measurements at 17.2 T look promising and several solutions can be adopted to tackle the observed problems, notably by shortening the thickness of our OVS and considering the transition bandwidth for positioning of the OVS bands.

Up to 12 metabolites were quantified using LCModel and the T$_1$ relaxation times of 10 metabolite moieties were estimated. The metabolite concentrations of GPC, PCr and PCh were in agreement with those found with $^1$H MRS using LASER. The ATP concentration and the
[ATP]/[PCr] ratio was lower than expected. Based on our low ATP concentrations, it might be interesting to account for T₂-weighting in the future. Preliminary results in the aging DA rat brain were also presented.

In the future, the presented approach could be readily used for other experimental setups. Notably, the implementation of the BISTRO scheme opens the path for saturation transfer experiments, in particular for the measurement of the rate of ATP synthesis, as it has been done in the mouse model of AD at 14.1 T (Mlynàrik, et al., 2012).
PART III: MRSI STUDIES AT 7 TESLA

5. APPLICATION OF PARALLEL TRANSMISSION TO $^1$H MRSI AT 7 TESLA

At high magnetic field, $^1$H MR Spectroscopy benefits from an increase in sensitivity and spectral resolution by an intrinsic increase of the Signal-to-Noise Ratio (SNR) and a higher chemical shift dispersion. Yet, at 7 Tesla and higher, the wavelength of radiofrequency (RF) pulses is shortened to the size of the human head, creating interference patterns exalting $B_1$ field inhomogeneity (Yang, et al., 2002; Van de Moortele, et al., 2005). In consequence, sub-optimal results are obtained in MRSI and MRS studies due to the application of inhomogeneous flip angles over the volume of interest. Furthermore, at elevated magnetic fields, the increased energy deposition (Section 1.2.1.3) generated by the RF pulses limits the available power and number of RF pulses per unit of time (Avdievich, et al., 2009).

Parallel transmission (pTx) is one of the most promising and elegant solutions to tackle the problem of $B_1$ inhomogeneity through the use of independently-driven coil elements, providing a great flexibility to achieve adjustable excitation patterns (Katscher, et al., 2003; Zhu, 2004). However, careful monitoring of the apparition of highly localized energy deposition regions in the brain or “hot spots” is needed due to the complexity of the applied E-fields.

During this thesis, pTx methods were tested to tailor localized excitation fields and less energy-demanding OVS using different static $B_1$-shimming configurations within a single pulse sequence under conservative SAR constraints. Energy deposition levels followed the IEC guidelines for humans (International Electrotechnical Commission, March 2010) and were monitored using an in-house simulation software (Cloos, 2012).

5.1. OBJECTIVES

The primary goals of the presented work were to take advantage of the increased degrees of freedom offered by the pTx mode (i) to improve the excitation field homogeneity for a 2D, pulse-acquire chemical shift imaging (CSI) sequence (detailed in section 2.2.1) and (ii) to reduce extracranial lipid contributions with effective, low power demanding outer-volume suppression (OVS) pulses. To this end, static $B_1$-shim configurations corresponding to the circularly polarized (CP) eigenmodes or “rings” were used as OVS. Employing OVS rings make it possible to suppress signal originated from the periphery of the brain within a single RF pulse while preventing chemical shift artifacts since no selection gradients are used for such class of OVS (Hetherington, et al., 2010). The ring modes efficacy and robustness were compared to a numerically optimized OVS mode, whose calculation was based on evaluating a cost function in
two volumes of interest (VOI). The efficacy of these methods was evaluated in vitro. All the sequences, parameterizations and SAR limits were set under in vivo constrains.

5.2. STATIC B\textsubscript{1} SHIMMING

Static B\textsubscript{1} shimming in the context of parallel transmission consists in providing fixed amplitude and phase values for each individual element of a multi-transmit coil during RF transmission, resulting in a specific excitation pattern that can improve both RF homogeneity and efficiency (Mao, et al., 2006; Ibrahim, et al., 2007). The B\textsubscript{1} field generated at any point r in space when using n independent resonators is given by:

\[ B_{1}(r) = \sum_{i=1}^{n} A_i B_i(r) e^{-i(\phi_i(r)+\phi_0i)} \]  

(5.1)

where the amplitude B\textsubscript{i}(r) and phase \( \phi_i(r) \) are spatial functions determined by the coil geometry and loading conditions and \( A_i \) and \( \phi_0i \) are amplitude and phase parameters defined by the user. The set of the n parameters \( A_i \) and \( \phi_0i \) constitute a B\textsubscript{1} shimming configuration. Since the loading conditions and efficiency of each resonator changes from subject to subject and the phase relationships between the coils are spatially dependent, the calculation of the B\textsubscript{1} configuration varies with the coil, subject and brain region and must therefore be calculated for every experiment. In practice, \( B_i(r) \) and \( \phi_i(r) \) are determined for each resonator through B\textsubscript{1} mapping and \( A_i \) and \( \phi_0i \) are typically calculated as to reduce B\textsubscript{1} field inhomogeneity inside a specific VOI. The default B\textsubscript{1}-shim configuration of the coil corresponds to the pseudo-CP mode, which has full power amplitude (\( A_i = 1 \)) for all channels and there is a relative phase difference of the coil elements of 45°, the first coil starting with a 0° phase.

5.2.1. OVS ring modes

At 7 T, SAR constrains limit the available B\textsubscript{1} field required for high-bandwidth RF pulses, which are required for good spatial localization such as for the application of OVS bands. Furthermore, the numerous OVS bands required to correctly suppress signal originated from extracranial lipids rapidly becomes prohibitive in terms of SAR limitations. To tackle this problem, a different and less-SAR demanding approach can be adopted, consisting of the use of static B\textsubscript{1}-shim configurations aiming at exciting only the rim of the head in a single pulse without the use of selection gradients (figure 5.1). One of these methods consists in using the CP eigenmode “rings” obtained by altering the relative phase of the N resonating elements in a multi-transmit coil, each eigenmode being orthogonal to the other N-1 modes (Alagappan, et al., 2007; Hetherington, et al., 2010). The determination of the individual phase of each coil element \( j \) for an eigenmode \( m \) is given by the expression:
\[
\phi_{0,m,j} = -\phi_{0,0,j}(\text{VOI}_{\text{Inner}}) + (j - 1) \times \frac{2\pi}{n} \times m, \quad m = 1, 2, \ldots, n - 1
\]  

(5.2)

where \(\phi_{0,0,j}(\text{VOI}_{\text{Inner}})\) corresponds to the phase maximizing the constructive interference of the \(n\) channels inside the volume \(\text{VOI}_{\text{Inner}}\), typically chosen in the center of the coil’s field of view (FOV) as seen in figure 5.2C. The amplitudes of each channel are adjusted for the CP mode \((n = 0)\) using a least-squares algorithm to match a target \(B_1\) profile and the resulting \(A_i\) values are then used for the 7 other eigenmodes. The 8 eigenmodes or “ring modes” (hereafter referred to as ring mode \(M_i, n = i\)) for an 8 channel transmit/receive coil are shown in figure 5.2A. The ring mode \(M_2\) presents the excitation pattern that is most suitable for the spoiling of extracranial lipids. In comparison, the \(M_1\) excitation pattern is not selective enough while the \(M_3\) and \(M_4\) eigenmodes cause unwanted excitation inside the VOI.

![Multiple OVS Bands and Single OVS Ring](image)

Figure 5.1. Extracranial lipid signal suppression through spatially selective saturation. **Left.** Several OVS bands are required to cover the majority of the head skin. **Right.** A single OVS “ring” obtained with a specific \(B_1\) shim configuration results in a reduction of the required number of pulses while covering the entire brain.

### 5.2.2. Numerically optimized \(L_0\) and \(L_R\) modes

In parallel to the development of the ring modes, numerically-optimized \(B_1\) shim configurations were also used (figure 5.2B). For their numerical optimization, the 8 amplitudes and phase offsets were simultaneously allowed to vary freely. The calculations were done using the cost functions shown below:

\[
C(a_1, \ldots, a_8, \phi_1, \ldots, \phi_8, r) = \sum_{i=1}^{8} a_i B_i(r) e^{-i(\phi_i)}
\]  

(5.3)

\[
L_0 = \min_{V} \frac{\sum_{r \in \text{VOI}_{\text{Outer}}} |C(V,r)|}{\sum_{r \in \text{VOI}_{\text{Inner}}} |C(V,r)|^2}
\]  

(5.4)

\[
L_R = \min_{V} \frac{\sum_{r \in \text{VOI}_{\text{Inner}}} |C(V,r)|}{\sum_{r \in \text{VOI}_{\text{Outer}}} |C(V,r)|}
\]  

(5.5)
where $\mathbf{B}_i$ is the complex value of the $\mathbf{B}_1$ field measured for channel $i$.

The $L_0$ shim configuration was meant for homogenous excitation within the VOI. The purpose of the $L_R$ shim configuration was to be used as an OVS ring by minimizing the excitation in the inner volume and maximizing the peripheral excitation. The inner and outer volumes were defined manually (as ellipses) using reference images and a fixed ratio between their major axis of 0.6 was used (figure 5.2C). The $\mathbf{B}_1$ maps had an identical FOV as the images used for reference.

Figure 5.2. $\mathbf{B}_1$ shim configurations of an 8-channel multi-transmit coil in a water phantom

A. The circularly polarized (CP) mode where the mean phase at the center of the coil is zero. The eigenmodes or rings were generated by adding a phase shift between adjacent coil elements of $n*2\pi/8$, $n=1,\ldots,7$. B. Numerically optimized $\mathbf{B}_1$ shim for the excitation in the center of the coil ($L_0$) and for peripheral excitation ($L_R$). C. Outer and Inner volumes used for the calculation of both the ring modes (inner volume) or the numerically optimized $L_0$ and $L_R$ shim configurations (inner and outer volumes). During this work, the ratio of the inner volume was fixed to 0.6 of that of the outer VOI.

5.3. $\mathbf{B}_1$ MAPPING

Equation (5.1) shows that the $\mathbf{B}_1$ profile depends on the individual 8 resonators of the multi-transmit coil. In order to calculate the spatially-dependent $\mathbf{B}_1(r)$ and $\phi_i(r)$ profiles of each channel, a fast and reliable $\mathbf{B}_1$ mapping sequence is necessary. To this end, numerous techniques have been proposed, the Actual Flip angle Imaging (AFI) sequence being among the most popular (Yarnykh, 2007; Amadon, et al., 2008; Boulant, et al., 2010). However, the AFI sequence is time-consuming for pTx applications, limiting its use on clinical exams.

The XFL sequence (Amadon, et al., 2010; Amadon, et al., 2012; Fautz, et al., 2008) is a 2D multi-slice, magnetization-prepared turbo-FLASH pulse sequence. It employs a sharp slice-selective saturation SLR VERSE’d pulse (table 1.2) to pre-saturate the magnetization in a slice and its saturation efficacy depends on the spatially-dependent FA of the transmitting channel. The RF saturation is immediately followed by a gradient spoiler and a slice-selective excitation pulse using a FLASH readout. To increase SNR, the excitation pulse is applied by all the 8
channels with a linear combination specific to the channel whose $B_1$ profile is being measured. The application of the 8 channels for excitation is known as the interferometric method (Brunner, et al., 2008) and requires a post-processing step to retrieve the individual $B_1$ amplitude maps and requires an additional reference image with no saturation. Non-saturated FLASH images are also acquired to determine the individual phase maps.

With the XFL sequence, the $B_1$ maps of the 8 channels were acquired in less than 5 minutes with excellent correlation and identical resolution as the AFI sequence. Figure 5.3 shows typical $B_1$ profiles of the 8 resonators measured in a water phantom.

Figure 5.3. $B_1$ profiles of the 8 individual coils acquired in a water phantom using XFL. The presented $B_1$ maps were calculated at every experiment. The images show an axial slice rotated 90° clockwise.

5.4. MATERIALS AND METHODS

5.4.1. Experimental set-up

Experiments were performed on a Siemens Magnetom 7 Tesla MRI scanner (Siemens Medical System, Erlangen, Germany), equipped with an 8-channel Tx-array set and an AC84 head gradient coil (max. strength 80 mT/m, slew rate 400 T/m/s). A home-made 8-channel transmit-receive array head coil was used consisting of 8 strip-line dipoles distributed every 42.5° on a cylindrical surface of a 27.6 cm diameter (figure 5.4). All dipoles were tuned ideally at 297.18 MHz corresponding to the proton Larmor frequency at 7 T and matched identically to a 50 Ohm line impedance. The coil was built according to the norm ISO 13485 and certified by Bureau Veritas (France). A water spherical phantom (diameter 16 cm, salted with 4 g/L) matching the conductivity of the human brain was used for this study. Both the local and global average RF power deposition limits for the human subject were monitored in real time to ensure compliance to the SAR guidelines (International Electrotechnical Commission, March 2010).
5.4.2. MRI data acquisition

For all experimental sessions, the 8 individual channel \( B_1 \) profiles were obtained using a XFL sequence for fast \( B_1 \)-mapping (resolution: 5 mm isotropic, 64 x 40 matrix, 32 slices, 2.5 mm slice thickness with a 2.5 mm gap between slices) using the interferometric method (Amadon, et al., 2010; Amadon, et al., 2012).

\( B_0 \) mapping was performed using a fast 2D 3-echoes spoiled gradient echo sequence based on the AFI sequence (\( TR_1/TR_2 = 40/200 \), \( TE_1/TE_2/TE_3 = 1/2/3 \) ms, 5 mm isotropic resolution, 5 mm slice thickness) and it was used to correct for off-resonance errors for the \( B_1 \) profiles (Amadon, et al., 2008). A better resolved AFI sequence (2.5 mm isotropic) was also acquired for the generation of masks for \( B_1 \)-shimming calculations.

FASTESTMAP was used for iterative \( B_0 \) shimming on a 60 x 60 x 30 mm\(^3\) volume positioned at the center of the CSI slice.

Validation of the RF power calibration was done using the modified AFI sequence. To this end, the power required to achieve several flip angles (15°, 30°, 45°, 60°, 75° and 90°) in a determined VOI was calculated based on the acquired XFL \( B_1 \) maps and equation (1.31). The AFI sequence was run with the calculated RF power and the experimental FA was compared to the targeted FA.

All the FA validation tests were done using the pseudo-CP \( B_1 \)-shim configuration described in section 5.2.
For validation of the B\textsubscript{1}-shim configurations used for the CSI experiments, FLASH images were acquired (1 average, TR = 450 ms, TE = 4.2 ms, 1.2 mm isotropic resolution, 20 mm slice-thickness, 15° FA, 1.28 ms long sinc pulse). The WS and OVS preparation modules (figure 5.5) were added to this FLASH sequence with the same timing and RF characteristics as in the CSI sequence. Independent static B\textsubscript{1}-shim configurations could be set for the excitation, OVS and WS pulses. 5 FLASH images were acquired with the following B\textsubscript{1}-shim configurations:

- pseudo-CP, CP or L\textsubscript{0} mode for the excitation, no OVS;
- pseudo-CP mode for the excitation and either M\textsubscript{2} or L\textsubscript{R} mode for the OVS;
- L\textsubscript{0} mode for the excitation and L\textsubscript{R} mode for the OVS.

The required B\textsubscript{1} intensities were calculated for each configuration based on the values shown in table 1.2. Since the goal of the experiments was to test the efficacy of the OVS module using a water phantom, the OVS carrier frequency was centered on the water resonance frequency and the WS module was deactivated for the experiments shown in this chapter.

![Chronogram showing the WS and OVS modules incorporated prior to the FLASH and 2D CSI sequences. The WS (blue), OVS (dark red) and excitation (gray) RF pulses were managed independently and each could use a different static B\textsubscript{1}-shim configuration. The spacing between consecutive pulses were 401, 355, 172, 57, 49, 26, 18 and 10 ms. For clarity, the WS and OVS modules are displayed on two separate lines. Spoiler gradients are schematically presented in yellow and slice-selection gradient in orange.](image)

5.4.3. MRI data analysis

The reconstruction of the AFI sequences was done using in-house reconstruction routines and the masks were generated using the brain extraction tools available in the FSL software package (Smith, 2002). Matlab (The MathWorks, MA, USA) was used for the analysis of the imaging data, for retrieving the B\textsubscript{1} maps out of the interferometry data and for the calculation of the ring modes. The numerical optimization of the L\textsubscript{0} and L\textsubscript{R} shim configurations [equ.(5.4) and (5.5)]
was done using a non-linear unconstrained minimization algorithm (*fminunc* matlab function). The \(B_0\) map was used for phase correction of the non-saturated XFL reference image.

### 5.4.4. MRSI data acquisition

A 2D pulse-acquire CSI sequence was used (5 averages, 16 x 16 matrix, FOV 240 x 240 x 20 mm\(^3\), TR = 1500 ms, TE = 1.26 ms, FA = 90°, 4 preparation scans, elliptical encoding, 1.28 ms sinc pulse duration). The WS and OVS preparation modules shown in figure 5.5 were incorporated. CSI data were acquired using the following \(B_1\)-shim configurations:

- CP or \(L_0\) mode for the excitation, no OVS;
- CP mode for the excitation and \(M_2\) mode for the OVS;
- \(L_0\) mode for the excitation and \(L_R\) mode for the OVS.

As for the FLASH measurements, the OVS carrier frequency was centered on the water resonance frequency and the WS module was deactivated.

### 5.4.5. MRSI data analysis

A 50 Hz Hamming filter was applied on the reconstructed CSI data (32 x 32 matrix). Visualization of the water CSI maps was done using Syngo’s spectroscopy card.

### 5.4.6. SAR evaluation

SAR evaluation was done using the “CEASAR” software (figure 5.6) developed in-house (Cloos, et al., 2010), based on a conservative approach for the calculation of the allowed power limits. To this end, the model assumed constructive RF interference at all points in the brain. The 10 s and 6 min time averaged power limits were restricted to the more conservative guidelines of the 6-min SAR limits. An average light head of 5 kg was assumed, in order to provide subject-independent power limits. Complete absorption of the incident power by the head was also assumed.

At the beginning of the experiment, an initial set of time-averaged power limits was fixed for each individual channel (1.2 W time-averaged power limit over 10 s). For every pulse sequence, power settings and TR, the global and local 10-gram SAR limits were calculated using four pre-simulated datasets with different anatomies and positions inside the coil (Aarkid, East Lothian, Scotland and Virtual Family (Christ, et al., 2010)). The coil structure was simulated with HFSS (ANSYS, Canonsburg, PA, USA). The results from the worst-case scenario were considered and if necessary more appropriate time-average power limits were set for each channel. The setting of individual power limits reduced the power constrains when using 8 separate transmit channels.
while enforcing SAR safety. When SAR limits were exceeded, the software proposed either an increase in TR or a reduction of the applied voltage for SAR re-evaluation. Without CEASAR’s validation, the sequence could not be launched. On average, a safety margin of a factor of 5 was found between the true absorbed energy and the simulated result provided by CEASAR (Boulant, et al., 2011).

It should be noted that, the power emitted by the RF amplifiers is measured by the constructor through the use of directional couplers and time-averaged power meters (TALES system). If at any time the power limits are exceeded, the acquisition is terminated. All CEASAR-validated protocols comply to such limits and only non-validated protocols may trigger a sudden termination of the sequence (Cloos, 2012).
Figure 5.6. SAR evaluation tool CEASAR. Screenshots of the CEASAR software used for evaluation of SAR limits and individual coil time-averaged power limits for peak, 10s and 6 mins. Features are indicated with arrows and labels. A. Main screen shows a protocol in compliance with the SAR guidelines. The three columns correspond to the peak, 10 s and 6 mins time-averaged power limits for each of the 8 independent channels. As explained in the text, these limits are evaluated with respect to the most conservative guideline of 6-min limits. B. Display of the individual pulses to be applied on each channel, showing the RF-waveform intensities and lengths.
5.5. RESULTS

5.5.1. B₁-mapping and RF Calibration

Figure 5.3 shows typical B₁ profiles obtained during the experiments. To validate the RF power calibration method, the required power needed to achieve a precise excitation FA for the VOI shown in figure 5.7 was estimated. The FA distributions in the VOI are shown in table 5.1. On average, the deviation between the measured and target angles was 5.9 % ± 1.6 %. Thus, the linearity of the RF coil was confirmed for the range of used voltages (19 to 111 V, for a maximum power of 190 V).

![VOI used for RF power calibration tests.](image)

<table>
<thead>
<tr>
<th>Target (deg)</th>
<th>Measured (deg)</th>
<th>Deviation</th>
</tr>
</thead>
<tbody>
<tr>
<td>15</td>
<td>16.2 ± 1.5</td>
<td>8.2 %</td>
</tr>
<tr>
<td>30</td>
<td>31.6 ± 2.6</td>
<td>5.5 %</td>
</tr>
<tr>
<td>45</td>
<td>47.3 ± 3.9</td>
<td>5.2 %</td>
</tr>
<tr>
<td>60</td>
<td>63.4 ± 4.8</td>
<td>5.7 %</td>
</tr>
<tr>
<td>75</td>
<td>78.7 ± 6.0</td>
<td>5.1 %</td>
</tr>
<tr>
<td>90</td>
<td>92.9 ± 5.6</td>
<td>3.2 %</td>
</tr>
</tbody>
</table>

Table 5.1. Validation of the RF calibration pipeline.
Flip Angle measurements from a 150 cm³ VOI using the modified AFI sequence. The RF power was manually determined based on the measured XFL B₁ maps, aiming for a target FA (shown in the first column). The measured FA distributions as well as the deviation from their target angle are shown.
5.5.2. Pseudo-CP, CP vs $L_0$ excitation modes

Figure 5.8 shows two sets of experiments done with slightly different set-ups (using two phantoms with the same composition but slightly different volumes). The $B_1$ profiles for each $B_1$-shim configuration are shown along with the resulting FLASH images. The excitation homogeneity was evaluated by measuring the mean FA in the inner VOI from the FA maps (table 5.2) and the signal intensity from the FLASH images (table 5.3). Figure 5.9 displays the $B_1$ field profiles along the line and column crossing the center of the phantom for each $B_1$ configuration used.

For a similar nominal power setting, the $L_0$ mode is 40% more efficient and half as inhomogeneous within the inner VOI than the CP mode (Table 5.2). Consistently, figures 5.9A and 5.9B show less distorted profiles for the $L_0$ mode (black) than for the CP mode (red).

In term of signal intensity (Table 5.3), the $L_0$ mode led to a 29% more homogeneous signal at the center of the phantom (Purple ROI, figure 5.10). Also, the $L_0$ mode didn’t excite as much the periphery of the phantom as the CP mode.

Compared to the CP and the $L_0$ modes, the pseudo-CP mode led to more signal overall but not as homogenous as the other two “focalized” $B_1$ excitation modes in both VOIs.

5.5.3. $M_2$ ring vs $L_R$ modes

The efficacy of the $M_2$ and $L_R$ OVS ring modes were compared using the same pseudo-CP mode for excitation and pulse sequence (figure 5.5) with the WS turned off.

Compared to $M_2$, the $L_R$ mode led to a 78 % less intense excitation and a 65% more homogeneous FA distribution in the inner VOI while retaining 90% of the $B_1$ intensity in the outer VOI (table 5.2). Nevertheless, the $L_R$ mode had one disadvantage, which was the FA variability over the outer VOI, leaving a star-like pattern of residual signal (clearly visible in figure 5.8 and 5.9).

Among the CP-eigenmodes, $M_2$ is the most suitable for a use as an OVS ring mode thanks to its strong and homogenous excitation at the periphery of the phantom (figure 5.2, table 5.2). However, it also presents a moderate degree of excitation within the inner VOI.

The impact of the $M_2$ mode in the center of the phantom was confirmed by a 12% increased reduction in signal intensity compared to the $L_R$ mode. A small ROI (yellow, figure 5.10) placed in the periphery of the phantom was used to estimate the maximum efficacy of the OVS modes which was found similar for both OVS modes (88% vs 91%).
Figure 5.8. Separate phantom validations of the explored $B_1$ shim modes. $B_1$ profiles and their corresponding FLASH images determined in two in vitro experiments. To facilitate the comparison, an identical grayscale range was used for FLASH CP and $L_0$ images and OVS ring $M_2$ and $L_R$ images. The same pseudo-CP $B_1$-shim configuration was used for excitation when the two OVS ring modes were evaluated.
Figure 5.9. $B_1$ profiles along the central column and line crossing at the center of the phantom. Profiles along the central line (A, C) and column (B, D) of the $B_1$ maps shown for phantom 1 in figure 5.8. A and B show the CP and $L_0$ distributions. C and D show the distributions for $L_R$ and the $M_2$, $M_3$ and $M_4$ ring modes (respective phase shifts of $\pi/2$, $3\pi/4$ and $\pi$).

<table>
<thead>
<tr>
<th>$B_1$ shimming</th>
<th>VOI$_{Inner}$ (A.U.)</th>
<th>$\Delta B_1$ (%)</th>
<th>VOI$_{Outer}$ (A.U.)</th>
<th>$\Delta B_1$ (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CP</td>
<td>71 ± 18</td>
<td>25</td>
<td>89 ± 21</td>
<td>24</td>
</tr>
<tr>
<td>$L_0$</td>
<td>100 ± 9</td>
<td>9</td>
<td>81 ± 24</td>
<td>30</td>
</tr>
<tr>
<td>$M_1$</td>
<td>80 ± 24</td>
<td>30</td>
<td>58 ± 24</td>
<td>41</td>
</tr>
<tr>
<td>$M_2$</td>
<td>68 ± 31</td>
<td>46</td>
<td>92 ± 14</td>
<td>15</td>
</tr>
<tr>
<td>$M_3$</td>
<td>45 ± 25</td>
<td>55</td>
<td>79 ± 26</td>
<td>33</td>
</tr>
<tr>
<td>$M_4$</td>
<td>46 ± 20</td>
<td>43</td>
<td>69 ± 30</td>
<td>43</td>
</tr>
<tr>
<td>$L_R$</td>
<td>15 ± 11</td>
<td>73</td>
<td>83 ± 27</td>
<td>32</td>
</tr>
</tbody>
</table>

Table 5.2. Regional $B_1$ distributions of the $B_1$-shim configurations.

Mean $B_1$ values (A.U.) in the inner and outer volumes defined in figure 5.2C. The $B_1$ profiles were calculated assuming an identical power input. FLASH images acquired with the CP, $L_0$, $M_2$ and $L_R$ modes are shown in figure 5.8.
Figure 5.10. Definitions of the inner (purple) and outer (yellow) ROIs used for the measurements presented below in table 5.3.

<table>
<thead>
<tr>
<th>Excitation</th>
<th>OVS</th>
<th>Inner ROI</th>
<th>Outer ROI</th>
<th>Voltage(^{a}) (Exc / OVS)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pseudo-CP</td>
<td>-</td>
<td>295 ± 82</td>
<td>218 ± 16</td>
<td>17 / -</td>
</tr>
<tr>
<td>CP</td>
<td>-</td>
<td>231 ± 65</td>
<td>160 ± 9</td>
<td>14 / -</td>
</tr>
<tr>
<td>(L_{0})</td>
<td>-</td>
<td>226 ± 46</td>
<td>74 ± 10</td>
<td>20 / -</td>
</tr>
<tr>
<td>Pseudo-CP (M_{2})</td>
<td></td>
<td>262 ± 82</td>
<td>25 ± 8</td>
<td>17 / 10</td>
</tr>
<tr>
<td>Pseudo-CP (L_{R})</td>
<td></td>
<td>303 ± 64</td>
<td>19 ± 13</td>
<td>17 / 19</td>
</tr>
</tbody>
</table>

Table 5.3. Impact of the tested \(B_{1}\)-shim configurations on the signal intensity. OVS ring modes were tested using the same pseudo-CP mode for excitation. The ROIs are shown in figure 5.10.

\(^{a}\) Applied voltage when \(A_{1} = 1\) [eq. (5.1)].

5.5.4. Application to CSI

As shown in figure 5.11, a good match was found between the FLASH and CSI images acquired simultaneously when using different \(B_{1}\)-shimming configurations. Based on this observation, one can expect the explored \(B_{1}\)-shim modes to display the same properties when applied to CSI acquisitions. Notably, the numerically-optimized \(L_{R}\) still provided good results even for VOIs close to the rim of the phantom.
Figure 5.11. Simultaneous CSI and FLASH profiles with different $B_1$-shim configurations. The display of the CSI maps was obtained by fitting the water resonance in each voxel using the spectroscopy routines available in Syngo. For comparison, all FLASH images were scaled identically. The $B_1$-shim configurations were: A) CP mode, B) $L_0$ mode, C) CP excitation with OVS ring $M_2$ and D) $L_0$ excitation with OVS ring $L_R$. E and F are the VOIs used for calculating the $B_1$-shim configurations shown in (A, C) and (B, D), respectively. The dark bands traversing each CSI images are found on the reference image (localizer) to which the CSI image is co-registered in Syngo.

5.5.5. SAR evaluation

Local and global SAR values were estimated for the different CSI acquisitions using “CEASAR” as a percentage of the maximum time-averaged power limits of 10 W/Kg and 3.2 W/Kg, respectively. Results were summarized in Table 5.4. The numerically optimized $L_0$ and $L_R$ modes showed higher SAR levels than the CP and $M_2$ modes. Since phase information was ignored by “CEASAR”, only constructive interference was assumed. Therefore, SAR differences are only due to the different amplitude coefficients $A_i$ and nominal voltages used.

OVS and WS individual pulses introduced only a mild increase in SAR. This was expected as they were relatively long RF pulses (5.12 and 6.6 ms respectively) and they were applied using a Gaussian waveform, which is the second most RF-efficient pulse considered within the framework of this thesis (amplitude integral of 0.55, table 1.2). Using sharper amplitude-modulated pulses such as the hermit pulse for excitation and OVS rings lead to an increase in the local and global SAR levels by a factor of 2, as shown in table 5.4.
Table 5.4. SAR measurements with several B\textsubscript{1} shimming combinations

<table>
<thead>
<tr>
<th>Excitation</th>
<th>OVS\textsuperscript{a}</th>
<th>WS\textsuperscript{a}</th>
<th>Local SAR(%)</th>
<th>Global SAR(%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CP</td>
<td>-</td>
<td>-</td>
<td>2.89</td>
<td>1.45</td>
</tr>
<tr>
<td>L\textsubscript{0}</td>
<td>-</td>
<td>-</td>
<td>3.46</td>
<td>1.80</td>
</tr>
<tr>
<td>M\textsubscript{2}\textsuperscript{b}</td>
<td>-</td>
<td>-</td>
<td>0.34</td>
<td>0.17</td>
</tr>
<tr>
<td>L\textsubscript{R}\textsuperscript{b}</td>
<td>-</td>
<td>-</td>
<td>0.73</td>
<td>0.29</td>
</tr>
<tr>
<td>CP</td>
<td>M\textsubscript{2}</td>
<td>CP</td>
<td>5.41</td>
<td>2.69</td>
</tr>
<tr>
<td>CP\textsubscript{Herm}</td>
<td>M\textsubscript{2}\textsubscript{Herm}</td>
<td>CP</td>
<td>11.22</td>
<td>5.61</td>
</tr>
<tr>
<td>L\textsubscript{0}</td>
<td>L\textsubscript{R}</td>
<td>L\textsubscript{0}</td>
<td>7.70</td>
<td>3.64</td>
</tr>
</tbody>
</table>

Excitation pulses had a 3-lobes Sinc RF envelope. OVS and WS pulses had a Gaussian waveform. Pulse durations for the 4 OVS and WS pulses were 5.12 and 6.6 ms, respectively. Maximum SAR values correspond to 10 W/kg and 3.2 W/kg.

\textsuperscript{a} OVS and WS pulses were applied with a 90° FA.

\textsuperscript{b} These pulses were applied using a Gaussian RF envelope with a 5.12 ms pulse duration (i.e., as an OVS pulse).

\textsuperscript{Herm} Pulses were applied using a Hermit RF waveform.

5.6. DISCUSSION

5.6.1. B\textsubscript{1}-inhomogeneity

B\textsubscript{1} field inhomogeneity is a major problem for NMR at 7 T which can be tackled by using parallel transmission methods. Several methods have been proposed, notably for MR imaging applications, where spatial inhomogeneity is highly reduced with tailored RF pulses such as k\textsubscript{T}-points or spokes (Cloos, et al., 2012; Katscher, et al., 2003; Zhu, 2004; Zelinski, et al., 2008). These methods define the excitation field by modulating the RF contributions at different positions of the k-space, using the coil gradients between RF pulses for positioning in the k-space. Although RF-tailoring methods achieve good spatial localization, their use is limited in MR spectroscopy due to the strong off-resonance effects and narrow-band nature (Setsompop, et al., 2007). Setsompop et al. have managed to apply wideband spokes for CSI applications, but B\textsubscript{1}+ mitigation was achieved only for a 600 Hz bandwidth (Setsompop, et al., 2009).

On the other hand, pTx has been widely introduced into MRS and MRSI studies through static or dynamic B\textsubscript{1}-shimming (Boer, et al., 2012; Zhu, et al., 2013; Emir, et al., 2012; Pan, et al., 2010; Marjanska, et al., 2012). Static B\textsubscript{1}-shimming and the use of multi-transmit array coils improves the FA distribution homogeneity and can help reducing the energy deposition by increasing the efficiency at which the individual coil elements are used (Avdievich, et al., 2009; Cloos, et al., 2012). This is particularly important at high fields, where strong B\textsubscript{1} field inhomogeneities are present and SAR limits are quickly reached due to the increased power demands at higher magnetic fields [equation (1.52)].
The method explored here took advantage of the coil eigenmodes to generate low power-demanding OVS rings. This approach is an interesting answer for high-resolution CSI-FID studies where short TRs are needed and the chemical shift artifacts have to be minimized (Henning, et al., 2009). Furthermore, the individual power amplitudes of each resonator are determined for each subject and experimental session during the static $B_1$-shimming calculation. This allows a more efficiently use of the available power with respect to the SAR limitations.

5.6.2. Chemical shift artifact and SAR limits

The presented $M_2$ and $L_R$ ring modes showed a good efficacy while showing reasonable SAR demands. The absence of a slice-selection gradient abolished the need of large-bandwidth RF pulses, whose use is problematic due to SAR limitations. Covering the extra-cranial space is usually done using 6 to 8 individual OVS bands and OVS schemes require the application of 2 to 3 separate pulses for a good efficiency. Assuming 3 blocks of 6 OVS bands, a hermit waveform with a moderate bandwidth of 1.33 kHz and $FA = 90^\circ$, the SAR introduced by the OVS module alone would be 36.7% (local) and 18.4% (global) using our current experimental setup ($TR = 1.5$ s) and SAR evaluation method. In contrast, local and global SAR values for the $M_2$ and $L_R$ ring modes were below 3% and 1.2%, respectively. These SAR values were calculated considering a common reference power of 110 V. However, the reference voltage for human head is closer to 150 V (based on values from other studies in the research group). Using this more realistic voltage would lead to a 35% increase of the local and global SAR values of 63.9% and 31.86%, respectively. Although such set of OVS would still be accepted for a moderate TR of 1.5 s, the chemical shift artifact from lipids (~0.9 ppm) to the $tCr(CH_3)$ moiety (3.03 ppm) would correspond to a vast 48% spatial shift.

5.6.3. In vitro validation of static $B_1$-shimming excitation and OVS ring modes

The work presented in this chapter was meant as the first step towards the application of static $B_1$-shimming excitation and OVS ring modes for $^1H$ CSI-FID acquisitions in the human brain. The acquisition and processing steps were identical to those needed for in vivo studies and the presented SAR values were evaluated for simulated human heads. The CP mode and $L_0$ configurations showed similar results in the acquired images. Yet the $B_1$ profile of the $L_0$ configuration showed an improved excitation homogeneity (30%) within a small VOI while reducing the excitation at the periphery. Being the result of a numerical optimization, it is also likely that its performance will be higher than the $M_0$ mode for a larger ROIs used in the brain. Therefore, the $L_0$ configuration would be the first candidate for future in vivo applications. The disadvantages of the OVS ring modes over conventional OVS bands is that excitation inside the VOI cannot be completely nulled, leading to a partial saturation of the metabolites resonances.
(as shown by the *in vitro* measurements in table 5.3) and that the excitation profiles may vary between subjects. The numerical optimization used for the L_R mode limited both inconveniences compared to the M_2 mode. The experimental results were in agreement with the simulations as shown in figure 5.8. Nevertheless, the B_1 profiles for the M_2 mode showed mild differences between experiments, probably due to experimental variations particularly in coil loading and manual definition of the ROIs used for power calibration. The numerically optimized L_0 and L_R modes showed less inconsistency as illustrated by the B_1 profiles among the different experiments.

The L_R configuration had the disadvantage of presenting small gaps between each pair of resonators, giving a star-like profile. This irregular pattern could lead to a probable contamination by extra-cranial lipids *in vivo* and as such its validation is required. However, similar patterns have been used *in vivo* previously providing good results (Hetherington, et al., 2010; Boer, et al., 2012).

The ratio between the outer and inner VOI radii used for calculation of the B_1 shims (figure 5.2C) was fixed at 0.6. Other studies have used a ratio of 0.8 (Hetherington, et al., 2010). The similarity of our rings compared to those obtained by Hetherington et al. *in vitro* suggests that the ratio used here could be used for human studies.

In this study, the timing of our OVS module was not thoughtfully optimized as it was not the primary goal of the work. Nevertheless, the timings used here allowed appreciating the relative efficacy differences of each OVS ring modes *in vitro*. Similarly to the work done in section 2.2.2 for water suppression, such optimization could be done later using the range of T_1 values for lipid contributions at 7 T and the experimental FA distribution.

5.6.4. Possible improvements

The disadvantages that the OVS ring modes present over conventional OVS bands is that signal inside the VOI is partially suppressed (as shown by the *in vitro* measurements in table 5.3) and the spatial suppression profile may vary between subjects. The L_R mode is expected to reduce this inner excitation at the cost of some gaps at the periphery. Such gaps might be removed by altering the cost function shown in equation (5.5) by adding an additional weighting function penalizing the variance within the external VOI of adjustment. Combining the L_R and M_2 ring in the 4 OVS pulses by alternating the B_1-shim configurations may also be an alternative to remove the signal gaps observed in the L_R mode while reducing the unwanted signal suppression in the brain.

A more elegant solution would be the use of k_1-points or spokes for spatial selection of the extra-cranial tissue while selectively exciting only the lipids resonance frequencies. This approach requires a completely different implementation and could not be developed within the duration of this PhD thesis.
5.7. CONCLUSION

In this chapter, a parallel transmission approach was adopted to evaluate efficient, low-energy demanding OVS pulses at 7 T. An acquisition, data processing and SAR evaluation pipeline was validated in vitro. The use of static B1 shimming methods improved the FA homogeneity within a medium-size VOI at the center of the sample. The two methods for B1 shimming calculation, based on the coil eigenmode rings and the numerically optimized L0 and L_R configurations, showed encouraging results. The L_0 mode resulted in a slightly better performance than the CP mode in terms of FA homogeneity. The ring mode M_2 showed good suppression efficiency but also affected the VOI of interest. The L_R mode showed a consistent pattern between experiments, leaving untouched the inner VOI but leaving small unsuppressed gaps in the external VOI. The presented method has very low SAR demands and is a starting point for latter developments at 7 T and 11.7 T were SAR limitations will have an even greater impact on the parameterization of MRS pulse sequences. Due to numerous technical issues related to the prototype multi-transmit coil, the upgrade of our pTx interface (Tim Tx Step 2) and other regulatory pitfalls, this work could not go further during this PhD thesis.
6. **H AND **P MRSI IN THE HUMAN BRAIN AT 7 T

MRS studies in the human brain at 7 T benefits from higher SNR and increased spectral resolution, permitting in particular, the reliable and individual detection of Glu and Gln based on their respective H4 resonances at 2.35 and 2.45 ppm. Yet, the technical difficulties that arise at higher fields, namely the increased B1 inhomogeneity and increased power requirements, limit the design of clinical protocols and complicate the use of MRS sequences due to SAR constrains. Indeed, it has been shown that when using single transmit volume coils at 7 T, there is a 50 % variation in B1 homogeneity across axial brain slices and the required power to achieve similar B1 intensities is also increased by ~4-fold compared to that at 4 T (Vaughan, et al., 2001; Avdievich, et al., 2009). Also, increasing the longitudinal dimensions of head volume coils do not provide a substantial increase in longitudinal brain coverage (Nabetani, et al., 2006). To compensate for these technical difficulties, a double-tuned 1H/31P 16-element transceiver phased array (8 for each channel) with multi-transmit capabilities and elevated B1\textsuperscript{+} performance (B1 generated per unit of power) was acquired from Resonance Research Inc. (Billerica, MA, USA). Taking advantage of its dual-tune capabilities, our objective was to acquire preliminary 1H and 31P CSI data in healthy volunteers with the goal of defining a human research protocol at 7 T.

In this chapter, an optimized WET water suppression module (section 2.2.2) and a BISTRO OVS module (section 2.2.1.1) were evaluated at 7 T. To this end, a 2D CSI sequence using STEAM for localization (CSI-ST) was used for the acquisition of 1H MRS data and comparison of the WET modules. 31P metabolic maps were also acquired using a 2D CSI-FID sequence (section 2.2.1), employing either standard or BISTRO bands.

6.1. MATERIALS AND METHODS

6.1.1. Experimental set-up

Experiments were performed on a Siemens Magnetom 7 Tesla MRI scanner (Siemens Medical System, Erlangen, Germany) with a AC84 head gradient set (max. strength 80 mT/m, slew rate 400 T/m/s). For this study, a double-tuned 1H/31P 16-element transceiver phased array (8 for each channel) with multi-transmit capabilities was used (figure 6.1), operating at 297.182 MHz and 120.301 MHz at the 1H and 31P frequencies, respectively. The center of the B1\textsuperscript{+} FOV differed by less than 1 cm between the two nuclei. The coil was built using tight-fitted, inductively decoupled resonators which were manually tuned and matched for the load of a human head, increasing transmit and reception efficiency when similar “head” sizes are scanned. Further technical details can be found in the paper of its inventor N. Avdievich (Avdievich, 2011 p. Sect. 3.4).
Experiments were performed on a total of 6 young healthy volunteers. This preliminary study was approved by our institutional review board and informed consent was obtained from all volunteers.

![Double-tuned $^{31}$P/$^1$H transceiver 8-channel phased array.](image)

**Figure 6.1.** Double-tuned $^{31}$P/$^1$H transceiver 8-channel phased array. The array contains two concentric layers of evenly-spaced rectangular surface coils, the $^{31}$P surface coils corresponding to the inner layer. The array consisted of two detachable parts, the bottom section having 10 surface coils (5 per nucleus) and the top part with 6 (3 per nucleus). Although different tops were available to adjust for the volunteer head size, only one was used for this study leading to a 23 cm major axis.

6.1.2. MRI data acquisition

$T_2$-weighted GRE scout images were acquired (2 average, $TR = 8.6$ ms, $TE = 4$ ms, 0.96 mm isotropic resolution, 240x240 mm$^2$ FOV, 20 mm slice-thickness, 3° FA, 1.28 ms long sinc pulse) for positioning of the 2D CSI slice and anatomical reference.

$B_1$ maps were acquired using a 3D modified AFI sequence covering the entire brain (Yarnykh, 2007) with spoiling improvements (Nehrke, 2009) and $B_0$ field inhomogeneity corrections (Amadon, et al., 2008; Boulant, et al., 2010). The sequence parameters were: $TR_1/TR_2 = 20/100$ ms; $TE_1/TE_2/TE_3/TE_4 = 3/3/4.5/7$ ms; 4-mm isotropic resolution with a 64x64x36 matrix and acceleration factor of 3. The sequence duration was 2m 53s.

Two 3D GRE images with flip angles of 5° and 20° were acquired ($TR/TE = 14/3$ ms, 1-mm isotropic resolution, sequence duration 4m 38s) from the same volume to estimate water $T_1$ maps.
6.1.3. MRI data analysis

A dedicated ICE reconstruction pipeline calculated the $B_1$ maps and “reference voltage” map (voltage required to achieve a 180° FA for a 1-ms long square pulse, according to the Syngo MR B17A IDEA User’s Guide). The reference voltage at the center of the CSI slice was then used as the default value for the rest of the experiment.

$T_1$ maps were calculated using the $B_1$ map generated from the AFI sequence and the two GRE acquisitions based on a 2-point version of the DESPOT1 method (Li, et al., 2006; Deoni, et al., 2003), where the FA is no longer estimated but is provided by our $B_1$ map.

6.1.4. MRSI data acquisition

For $^1$H MRSI data acquisition, a 2D modified CSI-ST sequence was used (section 2.2.1) : TR/TE/TM = 2000/9/10, 2048 complex points, 6kHz bandwidth, 2 averages, 19x19 encoding steps with elliptical acquisition weighting, 180x180 mm² FOV, 90x90 mm² VOI, 20 mm slice thickness,. The carrier frequency was shifted by -2 ppm relatively to the water resonance to minimize CSDA for most metabolites. The duration of the acquisition was 10m 52s. The WET water suppression module consisted of 4 frequency selective RF pulses (Gaussian pulse, 135 Hz bandwidth) with optimized flip angles and timings as shown in table 2.1. For comparison, $^1$H CSI-ST data were also acquired using the default water suppression scheme.

$^{31}$P MRSI data were acquired with a 2D CSI-FID sequence (section 2.1.3.1). OVS bands were used for the removal of signal coming from the skin and muscle (notably the temporalis muscle, used for mastication). The tested OVS modules were: a single BISTRO band ($n = 6$, HS8, 5.12 ms pulse length, 4.2 kHz bandwidth, total duration 48.75 ms), 1 or 2 OVS bands with a single adiabatic RF pulse (HS8, 5.12 ms pulse length, 4.2 kHz bandwidth) and 6 bands with a single conventional RF pulse (Hermit, 4.096 ms, 1.33 kHz bandwidth). All OVS bands had a thickness of 37 mm. A $^{31}$P CSI-FID sequence was acquired without OVS. The acquisition parameters were the following: TR = 2000, acquisition delay of 1.06 ms, 2048 complex points, 6kHz bandwidth, 5 averages, 19x19 encoding steps with elliptical acquisition weighting, 220x220 mm² FOV, 30 mm slice thickness, 18m 18s total scan duration. A flip angle of 56° was used, maximizing the available signal per unit time for PCr assuming a $T_1$ relaxation time of 3.54 s in humans at 7 T (Lu, et al., 2014). The $^{31}$P carrier frequency was shifted by +600 Hz so that the PCr resonance would be at 0 ppm.

$^1$H and $^{31}$P MRSI data were acquired from axial slices positioned above the lateral ventricles where the $B_1^+$ efficacy of the $^1$H coil was maximal (figure 6.2).

For $^{31}$P RF power calibration, reference voltages for both $^1$H and $^{31}$P channels were estimated at the center of the coil in vitro (spherical water phantom with phosphoric acid at 40 mM concentration, 16 cm diameter, 4 g of salt per L) by acquiring localized spectra using a STEAM
sequence and varying voltages (TR = 40 s, 8 averages) (section 2.1.3.2). For in vivo experiments, the ratio between reference voltages was assumed to be preserved and provided us with an a priori value for the reference voltage for $^{31}$P experiments. This reference voltage was revised by acquiring additional $^{31}$P STEAM spectra using varying voltages for one volunteer (TR = 2 s, 32 averages).

To determine the $B_1^+$ profile of the $^{31}$P phased array, a methodology similar to the AFI sequence was explored, where the signal ratio between two images with different TRs is used to derive the flip angle. To minimize the error, $r (= \text{TR}_1/\text{TR}_2)$ should be between 1/8 and 1/2 (Chmelik, et al., 2014). Thus, two $^{31}$P CSI maps were acquired using a TR$_1$ = 2s and TR$_2$ = 0.4 s, corresponding to a ratio $r$ of 1/5. No OVS bands were applied and the resulting acquisition times were 18m18s and 3m40s, respectively.

FAST(EST)MAP (Gruetter, et al., 2000) was used for $B_0$-shimming. The in-plane shim volume corresponded to a large area (~80 x 80 mm$^2$) covering the majority of the CSI VOI, excluding the skull and extracranial tissue. The thickness of the shimming volume was 40 mm, to prevent steep $B_0$ transitions adjacent to the MRSI slices.

6.1.5. MRSI data analysis

Visualization of the residual water signal in $^1$H CSI data and PCR in $^{31}$P CSI acquisitions was done using Syngo’s spectroscopy card. CSI data was processed using CSIAPO software (Le Fur, et al., 2010) developed on the IDL platform (Interactive Data Language, Research System, Inc., Boulder, CO, USA). Post-processing steps for the CSI data consisted in applying a Hanning spatial filtering and applying zero and first-order phase correction. An exponential filtering of 3 Hz was applied to all $^{31}$P CSI data in the time domain. Individual voxel spectra from CSI maps were retrieved using CSIAPO.

6.2. RESULTS

6.2.1. $B_1$-mapping and RF calibration

6.2.1.1. $^1$H channel

On average, the $^1$H reference voltage measured using the AFI sequence in vivo was 288 ± 21 V at the center of the brain. Typical $B_1$ maps within the CSI slice are shown in figure 6.2.
6.2.1.2. \( ^{31}\text{P} \) channel

*In vitro* RF calibration experiments using a STEAM sequence led to estimating a ratio between \( ^{1}\text{H} \) and \( ^{31}\text{P} \) reference voltages of 1.57. Since the \( ^{1}\text{H} \) reference voltage was approximately 288 V, the reference voltage for the \( ^{31}\text{P} \) channel was on average 452 V. For one volunteer, a higher \( ^{31}\text{P} \) reference voltage was found at 500 V. Therefore, the presented power referencing method underestimated the reference voltage at the center of the brain.

Estimation of the \( \text{B}_1^+ \) profile based on evaluating the signal ratio of two identically acquired scans could not be carried out as the SNR of the short TR acquisition was too low, effectively confounding the PCR signal to noise over the majority of the brain (figure 6.4).
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6.2.2. $B_0$-shimming

The water linewidths obtained from the large VOI used for FASTESTMAP was $18 \pm 2$ Hz. However, PCr linewidths of the central voxel from $^1$H CSI STEAM acquisitions showed an average width of $12 \pm 2$ Hz.

6.2.3. Water suppression efficacy

Figure 6.4 displays the residual water signal for two $^1$H CSI datasets acquired from two different volunteers using either the default water suppression scheme (6.4A) or the numerically optimized water suppression module (6.4B). Spectra corresponding to the same positioning relative to the VOI are shown. The suppression efficiency was estimated for the 5 shown regions by calculating the ratio between the residual water resonance and NAA(CH$_3$) heights. Ratios of $218 \pm 150$ and
25 ± 27 were found respectively for the default and numerically optimized WET schemes. This corresponds to an average improvement in suppression efficacy by a factor of 8.8.

Figure 6.4. WET water suppression efficacy evaluated in vivo at 7 T
A. Residual water distribution and (C) $^1$H spectra using the constructor’s WET module. B. Residual water distribution and (D) metabolite spectra using the optimized WET module shown in table 2.1. The spectra shown in each column were scaled identically and corrected for zero-order phase. Spectra in column (C) were corrected for first-order phase. Water distributions displays (A, B) were obtained by fitting the water resonance using Syngo’s spectroscopy card. The shown circles correspond to the point spread function (PSF) contour at 64% of the maximal PSF intensity. One can notice the 20% CSDA due to the 2 ppm shift between the water resonance and the carrier frequency.

6.2.4. Water $T_1$ mapping

For each volunteer, brain water $T_1$ maps were calculated based on the 2-points DESPOT1 method (figure 6.5A). Based on the histogram of $T_1$ values across the brain (figure 6.5B), distribution of
T₁ values in GM and WM were estimated: $T₁ = 1954 \pm 400$ for GM and $1314 \pm 294$ ms for WM. These values are in agreement with the literature (Wright, et al., 2008). Although there was a substantial inter-subject variation of the median T₁ relaxation times, the difference between GM and WM T₁ values was rather constant ($640 \pm 120$ ms), allowing the use of these maps for brain segmentation.

Figure 6.5. 3D T₁ maps established using the 2-point DESPOT1 method at 7 T in a normal brain.
A. The axial slice corresponds to the plane of the CSI VOI. B. Histogram of the T₁ values over the entire brain. The T₁ median values for WM and GM in this volunteer were 1050 and 1650 ms respectively.

6.2.5. OVS performance evaluation

Figure 6.6 displays a $^{31}$P map of the PCr signal acquired with a single BISTRO band using AFP HS8 pulses, positioned where the $B₁^+$ profile was observed to be most variable (possibly due to changes in coil loading between subjects). Based on our preclinical $^{31}$P experiments, the transition bandwidth was properly accounted for in the definition of the used HS8 pulses. As observed from the PCr map and the $^{31}$P spectra, the BISTRO band was effective in reducing the metabolite signals to noise levels. Yet, the elevated power requirements (SAR limits = 100%) did not allow the application of more than a BISTRO band for a TR of 2.3 s.
Figure 6.6. Single BISTRO band efficacy for $^{31}$P signal suppression in the human head at 7 T
A. PCr distribution in the brain under the application of a BISTRO band using HS8 RF pulses. B. Spectra corresponding to the regions marked in A. The spectrum #1 shows no residual signal above noise levels. Spectra are scaled identically and zero- and first-order phase corrections were applied. A 3 Hz exponential filtering was applied. The overlaid circles correspond to the point spread function (PSF) contour at 64% of the maximal PSF intensity.

Figure 6.7. Efficacy of a single adiabatic OVS band.
A. PCr signal distribution in the brain with an OVS band applied on one side of skull. The OVS band consisted of a single HS8 saturation pulse. B. Spectra from the regions marked in the PCr map. A clear reduction can be observed between the spectrum #1 and the other two spectra, showing a good suppression efficacy. All spectra are scaled identically. Zero- and first-order phase corrections were applied. A 3 Hz exponential filtering was applied. The overlaid circles correspond to the point spread function (PSF) contour at 64% of the maximal PSF intensity.
Figure 6.7 shows a map of the PCr signal and the efficiency of a single OVS band using a single adiabatic HS8 pulse. The OVS band was applied on only one side of the head to evaluate the suppression efficacy. A reduction of the PCr signal (+0.5 to -0.5 ppm) by a factor of ~8 was found between spectrum #1 and the non-suppressed spectrum #3. In another subject, 2 adiabatic OVS bands were applied (figure 6.8A) and compared to the application of 6 conventional OVS bands (figure 6.8B). Even if the second configuration led to a better definition of the VOI, the conventional OVS bands demonstrated a less consistent suppression efficacy than the adiabatic ones as it can be clearly observed from the comparison of spectra #3 and #5 in both columns. Both configurations were at 100% SAR limits for a TR = 2.3s.

Figure 6.8. Efficacy of conventional and adiabatic OVS bands for in vivo $^{31}$P MRSI at 7 T. PCr concentration maps in the same volunteer for two OVS configurations consisting in (A) 2 adiabatic OVS bands (5.1 ms HS8 pulse) and (B) 6 conventional OVS bands (4.1 ms hermit pulse). (C, D) Spectra from the 5 shown regions are displayed. For each column, the spectra are scaled identically and are corrected for zero- and first-order phase. A 3 Hz exponential filtering was applied. The overlaid circles correspond to the point spread function (PSF) contour at 64% of the maximal PSF intensity.
6.2.6. $^{31}$P Metabolite signal maps

$^{31}$P metabolite signal maps obtained using CSIAPO are shown in figure 6.9C along with spectra corresponding to voxels containing mostly either WM (#1) or GM (#2). Three additional spectra are shown corresponding to spectra averaged over the frontal (#3) and occipital cortex (#4) and the right superiori corona radiata (#5) (Wakana, et al., 2004). Consistently, decreased signal intensity was observed from the frontal area, probably due to lower coil sensitivity and effective FA compared to the back of the head where the coil elements were closer.

![Image of $^{31}$P Metabolite signal maps](image)

Figure 6.9. $^{31}$P Metabolic signal maps acquired *in vivo* in the human brain at 7 T.
A. Reference image showing the 2 adiabatic OVS bands and the different ROIs. B Spectra from GM (#1) and WM voxels (#2) and averaged spectra from the frontal (#3) and the occipital cortex (#4) and from the right superiori corona radiate (#5). All spectra were scaled identically. C. $^{31}$P metabolite signal maps co-registered with the anatomical image. All spectra were corrected for zero- and first-order phase and a 3 Hz exponential filter was applied. The overlaid circles correspond to the point spread function (PSF) contour at 64% of the maximal PSF intensity.

6.3. **DISCUSSION**

In this chapter, several technical aspects of MRSI at UHF were evaluated *in vivo* in order to orient future developments and constitute a $^1$H/$^{31}$P MRSI clinical protocol at 7 T.
6.3.1. *In vivo* validation of our clinical set-up

The good performance and dual-tune capabilities of our RF coil permitted the acquisition of $^1$H and $^{31}$P metabolite maps within a total scan time of 29 minutes with a moderate resolution (2.07 and 2.52 cm diameter of the PSF at 64% maximal intensity before Hanning filtering) at 7 T. As illustrated by figure 6.2, the $^1$H coil had a relatively inhomogeneous and narrow profile in the Z direction. This made positioning of the volunteer a critical aspect of our set-up. When the subject was placed too deep into the coil, the center of the $B_1^+$ field was shifted towards the lateral ventricles, resulting in a higher reference voltage for our CSI slice and an eventual TR lengthening to comply with SAR limitations. Placing the VOI too close to the ventricles jeopardized the convergence of FAST(EST)MAP toward a stable $B_0$-shim configuration and degraded the efficacy of the water suppression module, globally deteriorating the quality of the data.

For $^{31}$P CSI acquisitions, the frontal brain region showed a high inter-subject variability with respect to the detected signal. This is probably due to differences in coil loading between the head used for the preliminary tuning and matching of the coil and the actual volunteer. A solution could be to use the smaller top (leading to a 22 cm major axis) in the future to improve the sensitivity in the frontal area when volunteers with smaller heads are scanned. Another possibility would be to tune and match the three $^{31}$P resonators of the top part of the coil on each individual so as to restore the efficacy in the frontal area, at the cost of 2 or 3 minutes of experiment time.

6.3.2. $^1$H MRSI and WS efficiency

Based on the data acquired on our 6 volunteers, the efficiency of our WET scheme was evaluated. The optimized WET showed an indicative improvement of the water suppression efficacy in the brain by a factor of 8.8 with respect to the standard WET configuration. The efficacy of the optimized WET scheme is compatible with spectral decomposition software such as LCModel and application of an HLSVD water removal routine could further improve the results.

In this work, 2D $^1$H CSI maps were acquired in order to provide fair conditions for validation of our methods and also because of the reduced FOV of the coil in the Z direction. However, for 3D $^1$H CSI, the presented water suppression scheme may not provide optimal results. Other schemes, such as VAPOR, should provide better results due to the increased number of frequency selective pulses that are applied. Nevertheless, an increased power deposition would be accompanied and most importantly, a longer minimum TR would be required. A different approach, based on the acquisition of $^1$H metabolite data using CSI without the application of water suppression has been proposed (Le Fur, et al., 2014). Based on the modulus method (Serrai, et al., 2002), good quality
metabolite data has been obtained and it could provide an alternative or complementary solution for 3D $^1$H CSI acquisitions at UHF (Le Fur, et al., 2015).

6.3.3. $^{31}$P MRSI and OVS efficiency

Despite its efficiency, the energy requirement for the adiabatic BISTRO scheme was so elevated that only one was allowed for a TR of 2.4 s. The alternative single-pulse adiabatic OVS bands were efficient in spoiling the signal contribution from the temporalis muscles. However, two bands are not enough to properly delimit the brain VOI. Therefore, conventional OVS bands using AM pulse (or pTx OVS rings) need to be considered. Yet, the observed efficacy of the conventional OVS bands was not satisfactory in our experiments. This inefficiency was probably due to RF calibration errors and $B_1$ field inhomogeneity along the OVS bands. Measuring the correct $^{31}$P $B_1$ profile over the brain would allow to properly adjust for the applied power of the OVS bands.

Another approach would be to increase the number of pulse per OVS bands. As in a BISTRO or WET module, a scheme could be devised to reduce the sensitivity to $B_1$ inhomogeneity. However, the current configuration was already at maximum SAR limits, indicating that the pulse bandwidth would have to be reduced. Since the bandwidth of evaluated hermit pulses (1.33 kHz for a 4.1 ms duration) is already small (CSDA displacement of 26 % from PCr to ATPγ (-2.48 ppm) at 7 T), further decreasing the bandwidth while keeping a similar absolute CSDA displacement would demand the use of very thin OVS bands, losing in coverage. Therefore, restricting to a lesser number of bands seems as the only alternative if amplitude modulated pulses are to be retained. Furthermore, the major source of spurious signal observed in this study was originated from the temporalis muscles, which could be covered by two effective adiabatic suppression OVS bands.

Although the adiabatic BISTRO module cannot be considered as an option for OVS bands due to SAR limitations, it could instead be used for frequency-selective saturation of the ATP resonances so as to perform $^{31}$P magnetization-transfer MRSI experiments. Characterization of the BISTRO train efficacy as a function of train length and AFP modulating functions would nevertheless be required (Lei, et al., 2003).

6.3.4. Towards $^1$H and $^{31}$P qMRSI at UHF

In this chapter it is shown that it is possible to implement a complete analysis pipeline of the $^1$H and $^{31}$P MRSI data. As illustrated by figures 6.3 and 6.9, a good SNR and spectral resolution was obtained. For $^1$H MRSI, the total acquisition of the $B_1$ and water $T_1$ maps was 12 minutes.

Calculating the $B_1^+$ map of the $^{31}$P array was attempted based on the method of the signal ratio initially proposed by (Yarnykh, 2007). Contrary to the work done by (Chmelik, et al., 2014), we
did not manage to estimate the $B_1^+$ profile because the short TR acquisition had very poor SNR despite the fact that an almost identical TR was used (0.43 s). However, their flip angle was higher (90°) and the matrix size was smaller (16x16). This suggests that the power may have been underestimated and that noise sources such as elevated receiver spectral bandwidth should be avoided (for this particular scan), notably because the majority of the signal will originate from the PCr and ATP$\gamma$ resonances. Tests on a phantom using a similar $T_1$ relaxation time and concentrations as those found in vivo will be necessary for validation of the method. Post-processing of the data using LCModel can be easily adapted to analyze both the $^1$H and $^{31}$P spectra from the metabolite maps. To this end, CSIAPO will prove particularly useful once its export function in LCModel format (.RAW) will be validated.

6.4. CONCLUSION

In this pilot study, low energy demanding $^1$H and $^{31}$P CSI pulse sequences were used for the acquisition of preliminary $^1$H and $^{31}$P MRS data. Our optimized WET water suppression scheme exhibited a higher efficacy with respect to the WET module provided by the constructor. Among the different OVS bands evaluated, the single-pulse adiabatic OVS bands were found to be the most effective. However, the number of these OVS bands was drastically restricted by the SAR limitations. The alternative OVS scheme based on AM pulses needs further development, notably a correct, independent RF power calibration to properly determine their suppression efficacy.
**GENERAL CONCLUSION**

*Methodological Advances*

The main objective of this PhD thesis was to develop acquisition methods that could allow us to exploit the benefits that UHF provides for MR spectroscopy while compensating for the disadvantages linked to higher magnetic fields, notably the elevated $B_1$ and $B_0$-field inhomogeneities.

To tackle the important $B_1$-field inhomogeneities expected from using a surface coil at 17.2 T, a completely $B_1$-insensitive acquisition method was required. The LASER sequence was chosen due to its relative $B_1$-insensitivity thanks to the use of 6 AFP pulses but also because of the elevated excitation bandwidth that could be achieved with AFP pulses (20 kHz for a 1ms long HS8 RF pulses). Without such wide bandwidths, CSDA effects at 17.2 T could have jeopardized the quality of the spectra. To further improve our $^1$H spectroscopic data, a short, numerically optimized WET scheme was implemented. The resulting acquisition method was successfully applied in our longitudinal study of aging in the DA rat.

Different guidelines were followed for the $^{31}$P study in the rat brain, since the main technical objective was to achieve an identical localization between the $^1$H and $^{31}$P VOIs, allowing to do water referencing. Taking into account that the $^1$H/$^{31}$P dual tune coil was composed of two surface coils and that our targeted VOI was most of the rat brain, accounting for $B_1$ inhomogeneity was critical. The BISTRO module has been used in previous MRS studies as an effective method to localize $^{31}$P signal in presence of strong $B_1$ inhomogeneities. It was therefore implemented leading to a FID-BISTRO localization sequence.

The work done on the Siemens 7 T scanner was a first step towards the development of $^1$H and $^{31}$P MRSI protocols at NeuroSpin. The technical difficulties changed from the preclinical environment since SAR limitations stand as a major obstacle for employing energy-demanding pulses (AFP, SLR pulses). Moreover, strong $B_1$ inhomogeneities appear as a result of wavelength interference patterns, originated from the comparable size of the $^1$H wavelength in the brain dielectric compared to the human head. A 2D pulse-acquire CSI sequence with OVS localization was chosen to limit CSDA along one spatial direction while keeping a short acquisition time. This approach is interesting for quantitative $^1$H and $^{31}$P measurements as it minimizes $T_2^*$-weighting. For these reasons, the main work at 7 T was focused on developing and testing different OVS schemes.

The multi-transmit (pTx) approach offers a greater freedom than the conventional, single-channel transmission mode. It allows tailoring the $B_{1+}$ field in a specific VOI by adjusting the individual coil amplitudes and phases but it can also be used to generate “ring-like” excitation patterns which are well-suited for removal of extracranial tissue signal within a single pulse, reducing SAR. Another advantage of such mode is that it does not require a slice-selection gradient,
preventing CSDA effects. This technique was implemented and tested *in vitro* with promising results for $^1$H CSI-OVS acquisitions with a reduced SAR cost. Unfortunately, technical issues and some regulatory difficulties prevented their application *in vivo* during this thesis. Therefore, BISTRO and classical OVS bands were implemented based on the experience acquired at 17.2 T, combined with a $^{31}$P 2D CSI-FID sequence and evaluated *in vivo*. The BISTRO band proved to be inadequate due to its large SAR levels. The single-pulse adiabatic OVS bands showed good efficiency for a relatively large bandwidth of 4.2 kHz, reducing CSDA artifacts. The conventional OVS bands were not as effective and exhibited larger CSDA artifacts. But their low efficacy could be linked to an incorrect RF calibration at the rim of the head. If accounted for properly using an independent RF calibration of each band, a total of 6 conventional OVS bands could be placed simultaneously (for a TR of 2 s), leading to more appropriate delimitation of our VOI compared to the adiabatic OVS. In addition to OVS scheme, an improved WET scheme was evaluated for *in vivo* $^1$H MRSI in combination with a 2D CSI-ST sequence leading to satisfactory water suppression.

**Main Scientific Contributions**

The main scientific contributions of this PhD thesis emerges from the longitudinal study of normal brain aging in DA rats using quantitative $^1$H MRS. Brain aging is a central issue in today’s society as the aged population (above 60 years) is expected to double in proportion in the following decades and numerous neurodegenerative diseases display aging as the primary risk factor. Since it is unrealistic to plan for a longitudinal study in humans over decades, animal models provide an interesting alternative for the characterization of the normal physiological changes during aging. For this reason, we focused on studying the aging process in the rat brain at 17.2 T and chose the Dark Agouti rat strain as a practical model due to its rather stable weight (below 380g) without dietary restriction. To our knowledge, only one other longitudinal study was published recently by Duarte et al. (Duarte, et al., 2014) investigating normal brain aging in the 3 to 24 month-old mouse.

$T_1$ and $T_2$ relaxation times, including $J$-coupled resonances and macromolecules (MM), were measured from a large VOI along the $^1$H neurochemical profiles. Regional variability was examined also by acquiring spectra from 4 brain regions (thalamus, cingulate cortex and striatum). Through the careful parameterization of the MM baseline, the robustness of our spectral decomposition was improved while allowing the assessment of 4 groups of MM resonances from 0.5 to 4.5 ppm.

Due to time constrains, it was not possible to follow a single cohort over the full span of a rat lifetime. For this reason, two distinct cohorts of 1 and 8 month-old rats were followed in parallel and their spectroscopic data were analyzed together through a linear regression model accounting for individual rat variations as random effects. The results were consistent with a mild gliosis and neuroinflammatory response, confirming the observations found in the aging human, mouse and
rat brains using either $^1$H, $^{13}$C MRS or histological methods (Cerbai, et al., 2012; Chang, et al., 1996; Boumezbeur, et al., 2010; Duarte, et al., 2014). A regional variability of the age-related changes was observed, confirming that the aging process takes place at different rates on the cortical and sub-cortical areas.

One remarkable finding was that the number of scans had a confounding effect in our analysis. Such effect was unexpected but was discerned through the clear and reproducible patterns observed for our T$_1$ measurements between the two cohorts. To the best of our knowledge, this is the first time such observation has been reported. Similar influence of the number of exams was not reported by Duarte et al as they had only one cohort possibly leading to the confusion between the effects of aging and of the number of scans, if there was any in their study. Several hypotheses were presented to explain our observations but without a specific study, this cumulative effect of the number of exams remains open to discussion.

Quantitative $^1$H MRS data acquired from 1 month-old rats was published in Magnetic Resonance in Medicine. Notably, our T$_1$ and T$_2$ relaxation times at 17.2 T were compared to relaxation times measured at lower magnetic fields so as to assess their field-dependency. In addition, some preliminary data from our aging study were presented as poster or oral presentations at the International Society for Magnetic Resonance in Medicine (ISMRM) congress in 2014.

Preliminary $^{31}$P MRS data were acquired at 17.2 T from a subset of aging DA rats, covering the ages of 5, 17 and 21 months. These experiments allowed the evaluation of the quantification method proposed by Bottomley et al. (Bottomley, et al., 1996) based on the water signal as a heteronuclear reference of concentration. They also permitted to pinpoint some errors made in the parametrization of our adiabatic OVS bands. Applying the necessary corrections should help in improving the localization of $^{31}$P MRS data and the achievable SNR. Nevertheless, 12 phosphorylated metabolites were quantified and for 10 of them their T$_1$ relaxation times were also measured at least once. The GPC, PCr and PCh concentrations found using $^1$H or $^{31}$P MRS were in agreement. The PE concentrations differed but both the $^1$H and $^{31}$P values were consistent with the literature. LCModel was used for the analysis of the $^{31}$P spectra, allowing for an automated and operator-independent spectral decomposition. Recently, the use of LCModel for the analysis of $^{31}$P spectra was published and the results were compared with AMARES (Deelchand, et al., 2015). The work presented here was done independently and some improvements to our LCModel analysis could be made based on this recent paper, notably the simulation of model spectra with different linewidths, as observed in vivo. These $^{31}$P MRS results were accepted for a poster presentation at the 23$^{th}$ ISMRM congress in Toronto.

**Perspectives**

The unexpected cumulative effect of the number of exams observed during our aging study is intriguing and would deserve to be investigated further. However the large number of potential cofactors (B$_0$ and B$_1$ intensities, anesthesia, scan duration, stress) would make such study quite
difficult to design, especially considering that our clearest observation was an increase of the $T_1$ relaxation times for most metabolites. At least, one should be cautious about such unexpected effects in future longitudinal MRS studies at UHF. For instance, some relevant biological parameters could be checked, notably blood pressure and cortisol levels in order to monitor stress levels.

For the $^{31}$P quantitative MRS studies, the water referencing method was used due to its simplicity and adaptability to other research models. Unfortunately, this method provided with rather variable results as previously reported (Buchli, et al., 1994). Therefore, it may be convenient to adopt a less variable quantification method such as an external reference of signal close to the VOI. A very interesting approach would be the use of an electronic reference such as the ERETIC method (Ding, et al., 2012). However, its introduction requires hardware modifications and is therefore a solution that may require substantial effort to be implemented.

The implemented BISTRO module shown in this work was used exclusively as OVS bands and notably at 7 T, SAR limitations allowed the application of a single BISTRO band. However, the implementation of the BISTRO scheme opens the path for saturation transfer experiments, notably for the measurements of the equilibrium constant of the creatine kinase enzyme, as it has been done in the Alzheimer mouse model at 14.1 T or in healthy humans at 7 T (Mlynárik, et al., 2012; Lei, et al., 2003).

Furthermore, continuous work is necessary for the application of OVS bands for $^{31}$P MRSI human studies at 7 T. Although the present solution of using 2 OVS bands using an AFP pulse correctly eliminates the main extracranial signal from the temporalis muscles, this approach may no longer be enough for 3D MRSI acquisitions since at least a third OVS band will be required, notably for the suppression of the plane covering the eyes and mouth. Since $B_1$ inhomogeneity is expected to be quite elevated for such band, a mixed approach for signal saturation could be proposed, employing an adiabatic OVS band for the eyes while using conventional OVS bands, correctly calibrated, for the temporalis muscles.

The increases in $B_0$ and $B_1$ inhomogeneity at higher fields and the associated increase in power requirements are a major challenge for MRSI in the human brain particularly for signal localization. At the moment, our $^1$H/$^{31}$P dual coil was only used as a single-channel transmit coil for both nuclei, restricting its potential since SAR reduction and increased FA homogeneity can be obtained by applying static $B_1$-shimmings (Avdievich, 2011). Enabling it for pTx applications and validating the associated safety regulations could therefore improve substantially our $^1$H MRSI protocol.

There is also work to be done on the aspect of data analysis notably for the establishment of an automatic pipeline for spectral analysis and visualization of MRSI data. This is planned to be done in collaboration with Dr. Yann LeFur (Centre de Résonance Magnétique Biologique et Médicale - UMR 7339) and his software C SIPAO (Le Fur, et al., 2010), which will soon allow the exportation of pre-processed MRSI data in LCModel format.
Overall, this Ph.D. project demonstrated the benefits and challenges of MRS at UHF. The first longitudinal study in the aging rat brain was done, indicating the onset of the physiological changes expected in the aging brain. Yet, a confounding effect was detected and $^{31}$P results concerning the changes with age were limited. Although few technical improvements remain necessary, several tools were developed during this PhD thesis allowing the acquisition of high quality, preclinical and clinical $^1$H and $^{31}$P MRS data at ultra-high magnetic fields for the exploration of brain metabolism. This should clear the path for future translational MRS studies from rodents to men.
LIST OF PUBLICATIONS

Peer-reviewed journal

A.L. Lopez Kolkovsky, S. Mériaux, F. Boumezbeur. Metabolite and Macromolecule T₁ and T₂ Relaxation Times in the Rat Brain in vivo at 17.2 T. DOI: 10.1002/mrm.25602

Conference Proceedings


# Abbreviations and Symbols

<table>
<thead>
<tr>
<th>Abbreviation</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ace</td>
<td>Acetate</td>
</tr>
<tr>
<td>AD</td>
<td>Alzheimer’s disease</td>
</tr>
<tr>
<td>ADP</td>
<td>Adenosine-diphosphate</td>
</tr>
<tr>
<td>AFI</td>
<td>Actual flip angle imaging</td>
</tr>
<tr>
<td>AFP</td>
<td>Adiabatic full passage</td>
</tr>
<tr>
<td>AHP</td>
<td>Adiabatic half passage</td>
</tr>
<tr>
<td>Ala</td>
<td>Alanine</td>
</tr>
<tr>
<td>AMARES</td>
<td>Advanced Method for Accurate, Robust and Efficient Spectral fitting.</td>
</tr>
<tr>
<td>AMP</td>
<td>Adenosine-monophosphate</td>
</tr>
<tr>
<td>Asc</td>
<td>Ascorbic acid</td>
</tr>
<tr>
<td>Asp</td>
<td>Aspartate</td>
</tr>
<tr>
<td>ATP</td>
<td>Adenosine-triphosphate</td>
</tr>
<tr>
<td>Avg</td>
<td>Number of averages</td>
</tr>
<tr>
<td>BIR</td>
<td>$B_1$-insensitive rotation</td>
</tr>
<tr>
<td>BISTRO</td>
<td>$B_1$-insensitive train to obliterate signal</td>
</tr>
<tr>
<td>BPP</td>
<td>Bloembergen-Purcell-Pound</td>
</tr>
<tr>
<td>CHESS</td>
<td>Chemical shift selective</td>
</tr>
<tr>
<td>Cho</td>
<td>Choline-containing compounds</td>
</tr>
<tr>
<td>CNS</td>
<td>Central nervous system</td>
</tr>
<tr>
<td>CPMG</td>
<td>Carr-Purcell-Meiboom-Gill</td>
</tr>
<tr>
<td>Cr</td>
<td>Creatine</td>
</tr>
<tr>
<td>CRLB</td>
<td>Cramer-Rao lower bound</td>
</tr>
<tr>
<td>CSDA</td>
<td>Chemical shift displacement artifact</td>
</tr>
<tr>
<td>CSF</td>
<td>Cerebrospinal fluid</td>
</tr>
<tr>
<td>CSI</td>
<td>Chemical shift imaging</td>
</tr>
<tr>
<td>CYCLOPS</td>
<td>Cyclically ordered phase sequence</td>
</tr>
<tr>
<td>1D</td>
<td>One-dimensional</td>
</tr>
<tr>
<td>2D</td>
<td>Two-dimensional</td>
</tr>
<tr>
<td>3D</td>
<td>Three-dimensional</td>
</tr>
<tr>
<td>DA</td>
<td>Dark Agouti (rat strain)</td>
</tr>
<tr>
<td>dB</td>
<td>Decibel</td>
</tr>
<tr>
<td>DFT</td>
<td>Discrete Fourier transform</td>
</tr>
<tr>
<td>DIR</td>
<td>Double Inversion m</td>
</tr>
<tr>
<td>EA</td>
<td>Ethanolamine</td>
</tr>
<tr>
<td>FA</td>
<td>Excitation flip angle</td>
</tr>
<tr>
<td>F6P</td>
<td>Fructose-6-phosphate</td>
</tr>
<tr>
<td>FFT</td>
<td>Fast Fourier transform</td>
</tr>
<tr>
<td>FID</td>
<td>Free induction decay</td>
</tr>
<tr>
<td>FLASH</td>
<td>Fast low-angle shot</td>
</tr>
<tr>
<td>FOV</td>
<td>Field of view</td>
</tr>
</tbody>
</table>
ABBREVIATIONS & SYMBOLS

FT  Fourier Transformation
FWHM Frequency width at half maximum
GABA  γ-Aminobutyric acid
GE  Gradient echo
Glc  Glucose
Glc1P  Glucose-1-phosphate
Glc6P  Glucose-6-phosphate
Gln  Glutamine
Glu  Glutamate
Glx  Glutamine and glutamate
Gly  Glycine
Gly1P  Glycerol-1-phosphate
GM  Grey matter
GPC  Glycerophosphorylcholine
GPE  Glycerophosphorylethanolamine
GSH  Glutathione
HF  High (magnetic) field
HLSVD Hankel Lanczos singular value decomposition
HMPT  Hexamethylphosphoroustriamide
HS  Hyperbolic secant modulated AFP RF pulse
HS8  Hyperbolic secant modulated AFP RF pulse, n = 8
IEC  International Electrotechnical Commission
IR  Inversion recovery
Lac  Lactate
LASER  Localization by adiabatic selective refocusing
MAS  Magic angle spinning
mI  Myo-inositol
MRI  Magnetic resonance imaging
MRS  Magnetic resonance spectroscopy
MRSI  Magnetic resonance spectroscopy imaging
MRUI  Magnetic resonance user interface method
MT  Magnetization transfer
NAA  N-Acetyl aspartate
NAAG  N-Acetyl aspartyl glutamate
NAD(H)  Nicotinamide adenine dinucleotide oxidized (reduced)
NADP(H)  Nicotinamide adenine dinucleotide phosphate oxidized (reduced)
NMR  Nuclear magnetic resonance
OVS  Outer volume suppression
PCh  Phosphocholine
PCr  Phosphocreatine
PDE  Phosphodiesters
PE  Phosphorylethanolamine
Pi  Inorganic phosphate
PND  Postnatal day
PPA  Phenylphosphonic acid
PPM  Parts per million
PRESS  Point resolved spectroscopy
PSF  Point spread function
R_{eff}  Effective resistance equal to the sum of the coil, electronics and sample resistances
ROI  Region of interest
ROS  Reactive oxygen species
RF  Radiofrequency
SAR  Specific absorption rate
SE  Spin echo
SI  Spectroscopic imaging
sI  Scylo-inositol
SIR  Single Inversion Recovery
SMF  Static magnetic field
SNR  Signal-to-noise ratio
STEAM  Stimulated echo acquisition mode
SV  Single voxel (or volume)
SVS  Single Volume Spectroscopy
TALES  Transmit Antenna LEvel Sensor
Tau  Taurine
tCho  Total choline
tCr  Total creatine
TMS  Tetramethylsilane
tNAA  Total NAA
UHF  Ultra high (magnetic) field
VAPOR  Variable pulse powers and optimized relaxation delays
VARPRO  Variable Projection method
VERSE  Variable rate selective excitation
VOI  Volume of interest
WM  White matter
WURST  Wideband, uniform rate and smooth truncation
A.U.  Arbitrary Units
B_0  External static magnetic field
B_1  Magnetic radiofrequency field of the transmitter coil
C_{PH}  Proton-Phosphorous sensitivity scaling factor of a dual coil
E  Energy
E  Electrical field
e  Particle charge
emf  Electromotive force
f_{CSF}, f_{GM}, f_{WM}  Fractional CSF, GM and WM volumes
G  Gradient strength
h  Planck’s constant (6.626208 x 10^{-34} J s)
I  Spin quantum number
<table>
<thead>
<tr>
<th>Symbol</th>
<th>Description</th>
</tr>
</thead>
<tbody>
<tr>
<td>J</td>
<td>Spin-spin or scalar coupling constant (in Hz)</td>
</tr>
<tr>
<td>$k_B$</td>
<td>Boltzmann equilibrium constant ($1.38066 \times 10^{-23} \text{ J K}^{-1}$)</td>
</tr>
<tr>
<td>$L$</td>
<td>Angular momentum</td>
</tr>
<tr>
<td>$m$</td>
<td>Mass</td>
</tr>
<tr>
<td>$m$</td>
<td>Magnetic quantum number</td>
</tr>
<tr>
<td>$M$</td>
<td>Macroscopic magnetization</td>
</tr>
<tr>
<td>$M_i$</td>
<td>Eigenmode of phase shift $i*2\pi/8$</td>
</tr>
<tr>
<td>met</td>
<td>Metabolite (concentration)</td>
</tr>
<tr>
<td>$M_{x,y,z}$</td>
<td>Orthogonal Cartesian components of the macroscopic magnetization</td>
</tr>
<tr>
<td>$M_0$</td>
<td>Macroscopic equilibrium magnetization</td>
</tr>
<tr>
<td>$P$</td>
<td>Power deposition</td>
</tr>
<tr>
<td>$r$</td>
<td>Distance</td>
</tr>
<tr>
<td>$R$</td>
<td>Product of the bandwidth at FWHM by the pulse duration</td>
</tr>
<tr>
<td>$R_m$</td>
<td>Weighting factor due to $T_1$ and $T_2$ relaxation of species $m$</td>
</tr>
<tr>
<td>$R^2$</td>
<td>Coefficient of determination</td>
</tr>
<tr>
<td>$S$</td>
<td>Measured NMR signal</td>
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<tr>
<td>$S_m$</td>
<td>Measured metabolite signal</td>
</tr>
<tr>
<td>$S_w$</td>
<td>Measured water signal</td>
</tr>
<tr>
<td>$t$</td>
<td>Time</td>
</tr>
<tr>
<td>$T$</td>
<td>Temperature (in K°)</td>
</tr>
<tr>
<td>$T$</td>
<td>Torque</td>
</tr>
<tr>
<td>$T_p$</td>
<td>RF pulse length</td>
</tr>
<tr>
<td>$T_1$</td>
<td>Longitudinal relaxation time constant</td>
</tr>
<tr>
<td>$T_1^*$</td>
<td>Observed longitudinal relaxation time constant</td>
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<tr>
<td>$T_2$</td>
<td>Transverse relaxation time constant</td>
</tr>
<tr>
<td>$T_2^*$</td>
<td>Apparent transverse relaxation time constant</td>
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<tr>
<td>$T_{acq}$</td>
<td>Acquisition time</td>
</tr>
<tr>
<td>$T_{bw}$</td>
<td>Transition bandwidth</td>
</tr>
<tr>
<td>TE</td>
<td>Echo time</td>
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<tr>
<td>TI</td>
<td>Inversion time</td>
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<tr>
<td>$T_{I1}$</td>
<td>First inversion recovery time</td>
</tr>
<tr>
<td>$T_{I2}$</td>
<td>Second inversion recovery time</td>
</tr>
<tr>
<td>TR</td>
<td>Repetition time</td>
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<tr>
<td>$\hat{T}R$</td>
<td>Recovery time</td>
</tr>
<tr>
<td>$v$</td>
<td>Velocity</td>
</tr>
<tr>
<td>$W(k)$</td>
<td>Spatial frequency weighting function</td>
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<tr>
<td>$W$</td>
<td>Apparent water concentration</td>
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<tr>
<td>$\alpha$</td>
<td>Nutation angle</td>
</tr>
<tr>
<td>$\gamma$</td>
<td>Gyromagnetic ratio</td>
</tr>
<tr>
<td>$\delta$</td>
<td>Chemical shift (in ppm)</td>
</tr>
<tr>
<td>$\Delta B_0$</td>
<td>Magnetic field shift</td>
</tr>
<tr>
<td>$\Delta \omega$</td>
<td>Frequency offset</td>
</tr>
<tr>
<td>$\omega_{\text{max}}$</td>
<td>Spectral bandwidth of an RF pulse</td>
</tr>
<tr>
<td>Symbol</td>
<td>Definition</td>
</tr>
<tr>
<td>--------</td>
<td>-----------------------------------------------</td>
</tr>
<tr>
<td>$\omega_{\text{rec}}$</td>
<td>Receiver bandwidth of the detecting system</td>
</tr>
<tr>
<td>$\theta$</td>
<td>Nutation angle</td>
</tr>
<tr>
<td>$\kappa$</td>
<td>Amplitude integral of an amplitude-modulated pulse</td>
</tr>
<tr>
<td>$\mu$</td>
<td>Magnetic moment</td>
</tr>
<tr>
<td>$\mu_z$</td>
<td>Magnetic moment component in the z direction</td>
</tr>
<tr>
<td>$\sigma$</td>
<td>Magnetic shielding constant</td>
</tr>
<tr>
<td>$\sigma_{\text{thermal}}$</td>
<td>Thermal noise</td>
</tr>
<tr>
<td>$\nu_0$</td>
<td>Larmor frequency</td>
</tr>
<tr>
<td>$\nu_{\text{ref}}$</td>
<td>Reference frequency</td>
</tr>
<tr>
<td>$\phi$</td>
<td>Phase</td>
</tr>
<tr>
<td>$\phi_0$</td>
<td>Zero-order (constant) phase</td>
</tr>
<tr>
<td>$\phi_1$</td>
<td>First-order (linear) phase</td>
</tr>
<tr>
<td>$\Omega$</td>
<td>Frequency Isochromat</td>
</tr>
<tr>
<td>$\omega_0$</td>
<td>Larmor frequency</td>
</tr>
<tr>
<td>$[]$</td>
<td>Molar concentration</td>
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