Optimizing flip angles for metabolic rate estimation in hyperpolarized carbon-13 MRI

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Abstract—Hyperpolarized carbon-13 magnetic resonance imaging has enabled the real-time observation of perfusion and metabolism in vivo. These experiments typically aim to distinguish between healthy and diseased tissues based on the rate at which they metabolize an injected substrate. However, existing approaches to optimizing flip angle sequences for these experiments have focused on indirect metrics of the reliability of metabolic rate estimates, such as signal variation and signal-to-noise ratio. In this paper we present an optimization procedure that focuses on maximizing the Fisher information about the metabolic rate. We demonstrate through numerical simulation experiments that flip angles optimized based on the Fisher information lead to lower variance in metabolic rate estimates than previous flip angle sequences. In particular, we demonstrate a 20% decrease in metabolic rate uncertainty when compared with the best competing sequence. We then demonstrate appropriateness of the mathematical model used in the simulation experiments with in vivo experiments in a prostate cancer mouse model. While there is no ground truth against which to compare the parameter estimates generated in the in vivo experiments, we demonstrate that our model used can reproduce consistent parameter estimates for a number of flip angle sequences.

Index Terms—Hyperpolarized carbon-13 magnetic resonance imaging, optimal experiment design, Fisher information, quantitative imaging, parameter mapping

I. INTRODUCTION

HYPERPOLARIZED carbon-13 magnetic resonance imaging (MRI) has enabled the real-time observation of perfusion and metabolism in preclinical and clinical studies [1]–[6]. This technology is made possible by techniques for dynamic nuclear polarization (DNP) that have led to signal-to-noise ratio (SNR) increases of four to five orders of magnitude compared with endogenous signal in dissolved 13C-labelled molecules [7], [8]. Injected [1-13C] pyruvate is frequently used as a substrate in metabolism experiments and its rate of conversion to [1-13C] lactate has been shown to distinguish between healthy and diseased tissues in animal [2], and recently human [11] studies.

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In contrast with conventional MRI, magnetization is a non-renewable resource in hyperpolarized MRI. Conventional imaging relies only on thermal equilibrium polarization, therefore an arbitrary number of acquisitions can be performed if we allow time for the magnetization to return to equilibrium between acquisitions. In contrast, hyperpolarization can only be performed before a 13C-labeled substrate is injected into the body, and once injected the magnetization decays due to $T_1$ relaxation and rapid $T_2$ relaxation following radio frequency (RF) excitation. Thus, the choice of excitation sequence is important for managing the trade-off between present and future measurement quality.

In typical practice a constant flip angle sequence is used for excitation, with typical values ranging from 5–30 degrees. Alternative time-varying acquisition sequences include sequences that attempt to maintain constant observed signal over time [9], maximize the cumulative observed lactate signal over time [10], or saturate the lactate signal in each acquisition [11].

In this paper our goal is to design a time-varying flip angle sequence to achieve maximally reliable quantitative estimates of the metabolic rate that can be compared between tissue regions, across subjects, or over time. To achieve this we develop a statistical model of the observed data as a function of the flip angle sequence and design flip angles to maximize the Fisher information about the metabolic rate parameter.

We begin by presenting a mathematical model of the magnetization dynamics in the observed tissue in Section II. In Section III we introduce a flip angle optimization procedure and present an optimal sequence. Next, we validate this result with computer simulation studies to demonstrate that our optimized sequence yields more reliable metabolic rate estimates than commonly-used flip angle sequences in Section IV. Finally, in Section V we demonstrate the feasibility of this procedure in vivo and demonstrate the well-foundedness of our mathematical model with experiments in a prostate cancer mouse model.

The software and experimental data required to generate the figures in this paper are available at: https://github.com/maidens/TMI-2015

II. MATHEMATICAL MODEL

Linear differential equations are well-established as models of metabolic flux measured using MR spectroscopy [2], [12]. It is shown in [13] that a first-order, two-site model with unidirectional flux of pyruvate to lactate is sufficient to accurately model the appearance of lactate, when observed by
hyperpolarized MR. This work also showed that increasing the fidelity of the model to incorporate bidirectional flux, or transport of lactate outside the cell, did not significantly improve the fit to hyperpolarized MR data. Therefore, we consider a two-dimensional system of ordinary differential equations

$$\frac{dx}{dt}(t) = \begin{bmatrix} -k_{PL} - R_{1P} & 0 \\ k_{PL} & -R_{1L} \end{bmatrix} x(t) + \begin{bmatrix} k_{TRANS} \\ 0 \end{bmatrix} u(t)$$

that models the magnetization dynamics in a tissue with an arterial input function $u(t)$ and uni-directional conversion from the substrate (pyruvate) to a metabolizable product (lactate), which has been commonly applied for hyperpolarized $^{13}$C pyruvate experiments. The state $x_1(t)$ denotes the longitudinal magnetization of pyruvate contained in a particular voxel in the tissue and $x_2(t)$ the longitudinal magnetization of lactate in the same voxel. The rate of metabolism of pyruvate to lactate is denoted $k_{PL}$, the perfusion rate from the arterial input to the tissue is denoted $k_{TRANS}$, and $R_{1P}$ and $R_{1L}$ are lumped parameters that account for $T_1$ decay in the magnetization along with other effects, such as metabolism of pyruvate into products other than lactate as well as flow of magnetization out of the slice. The input to the system $u(t)$ is an unmeasured arterial input function (AIF) resulting from the injection of hyperpolarized [1-$^{13}$C] pyruvate. In an experimental setting an AIF will be estimated based on the data collected, but for the purposes of designing a flip angle sequence, it will be assumed to be of gamma-variate shape

$$u(t) = A_0 (t - t_0)^\gamma e^{-(t-t_0)/\beta}$$

with parameters $t_0, \gamma, \beta, A_0$ given in Table I. We acquire data at $N$ time points separated by intervals of length $T_R$. Each time $t$ an acquisition is made, we must choose a flip angle $\alpha_{k,t}$ for each compound $k$ to be measured. If the magnetization of the $k$-th compound before the acquisition is $x_k$, then this choice of flip angle allows us to measure a signal of magnitude $\sin(\alpha_{k,t}) x_k$, after which $\cos(\alpha_{k,t}) x_k$ magnetization remains for future acquisitions (Fig. 1a). This causes discrete jumps, or resets, in the system state, leading to a hybrid dynamical system (Fig. 1b) (Fig. 15). Since we are only interested in the system’s state at acquisition times, we can avoid technicalities associated with hybrid system modelling by discretizing the system in time and considering a discrete-time dynamical system that simultaneously captures the evolution of the longitudinal magnetization of lactate and the discrete jumps induced by the acquisitions. We define the transition matrices $A_d$ and $B_d$

$$A_d = \exp \left( T_R \begin{bmatrix} -k_{PL} - R_{1P} & 0 \\ k_{PL} & -R_{1L} \end{bmatrix} \right)$$

$$B_d = \begin{bmatrix} -k_{PL} - R_{1P} & 0 \\ k_{PL} & -R_{1L} \end{bmatrix}^{-1} (A_d - I) \begin{bmatrix} k_{TRANS} \\ 0 \end{bmatrix}$$

that correspond to the discretization of $[1]$ assuming a zero-order hold on the input between each acquisition $[15]$. We will construct metabolite maps using magnitude image data, which necessitates a Rician noise model. Using magnitude images allows us to avoid modelling sources of phase in the image which would require additional states and parameters to estimate. It would also be possible to perform the parameter mapping using complex images with Gaussian noise, but this would require revising the model to account for phase, or modifying the image reconstruction to estimate and remove phase from the acquired images.

Accordingly, we model the measurements as independent Rician-distributed random variables $[16]$, which have probability density

$$p_{x,\sigma}(y) = \frac{y}{\sigma^2} \exp \left( -\frac{y^2 + x^2}{2\sigma^2} \right) I_0 \left( \frac{y^2}{2\sigma^2} \right)$$

where $I_\nu$ denotes the modified Bessel function of the first kind of order $\nu$. All together, we have the discrete-time model

$$x_0 = 0$$

$$x_{t+1} = A_d(\theta) \begin{bmatrix} \cos \alpha_{1,t} & 0 \\ 0 & \cos \alpha_{2,t} \end{bmatrix} x_t + B_d(\theta) u_t(\theta)$$

$$\tilde{x}_{k,t} = \sin(\alpha_{k,t}) x_{k,t}$$

$$k = 1, 2$$

$$Y_{k,t} \sim Rice(\tilde{x}_{k,t}, \sigma_k)$$

$$k = 1, 2$$

Simulated trajectories of this model are shown in Fig. 2. The model parameters are

$$\theta = [R_{1P}, R_{1L}, k_{TRANS}, u_0, \ldots, u_{N-1}]$$

and we have the freedom to choose

$$\alpha = \begin{bmatrix} \alpha_{1,1} & \ldots & \alpha_{1,N} \\ \alpha_{2,1} & \ldots & \alpha_{2,N} \end{bmatrix}$$

to generate the best possible estimate of the unknown parameters. The noise parameters $\sigma_k$ for $k = 1, 2$ can be estimated separately from a measurement of the background and are therefore assumed to be known. We fix a sampling interval of $T_R = 2$ seconds, though this could in principle be included as a decision variable.

III. OPTIMAL EXPERIMENT DESIGN

In this section, we present a methodology for designing flip angle sequences to provide maximal information about the metabolic rate parameter $k_{PL}$. This methodology is based on an optimization problem that can be solved to local optimality.
using nonlinear models as it is general, easy to compute, provides good results in practice and can be justified mathematically via asymptotic analysis [18].

In this paper, we will estimate unknown model parameters using the maximum likelihood estimator (MLE) defined as

$$\hat{\theta}_{MLE} \in \arg \max_{\theta \in \mathbb{R}^d} p_0(Y)$$

where \( Y \) is the observed data. Under mild assumptions, the maximum likelihood estimator is asymptotically efficient [19], that is,

$$\sqrt{n}(\hat{\theta}_{MLE} - \theta) \xrightarrow{d} \mathcal{N}(0, I^{-1})$$

as the number \( n \) of independent data sets used to compute the MLE tends to infinity. Here \( \mathcal{N}(0, I^{-1}) \) denotes the multivariate normal distribution with mean zero and covariance \( I^{-1} \) and \( \xrightarrow{d} \) denotes convergence in distribution. Thus, we see that asymptotically the MLE achieves the Cramér-Rao bound. While it is not necessarily the case in general that the MLE based on a single data set, or any finite number of data sets, has covariance equal to \( I^{-1} \), it is often a good approximation. Thus, we attempt to design an experiment to reliably estimate \( \theta \) by maximizing the Fisher information.

The Fisher information is a function of the true value of the parameter vector \( \theta \), which is unknown \textit{a priori}. There are commonly three approaches to overcome this difficulty:

1) Minimax/maximin optimal design: Given a range \( \Theta \) of values for the parameter vector \( \alpha \) to maximize the worst-case information among all potential values of the parameter \( \theta \in \Theta \) (i.e. \( \alpha^* = \arg \max_{\alpha \in A} \min_{\theta \in \Theta} \mu(I(\theta, \alpha)) \)) for some measure \( \mu \) of the size of the information matrix). This approach is advantageous because it provides a guaranteed lower bound on the information gained from an experiment despite parametric uncertainty. But it typically leads to a design \( \alpha^* \) that is optimized for a “corner case” in the parameter space, which may be overly conservative in practice. In addition, minimax/maximin objective functions are non-differentiable and can be difficult to optimize numerically [20, 21].

2) Bayesian optimal design: Given a prior distribution \( p_0(\theta) \) on the space of possible parameter values, maximize the expected information (i.e. \( \alpha^* = \arg \max_{\alpha \in A} \int \mu(I(\theta, \alpha)) p_0(\theta) d\theta \)). This approach handles parametric uncertainty nicely, but can lead to difficulties when it is unclear how to choose an appropriate prior, or when the parameter space is large and hence the computation of the high-dimensional integral is numerically intractable [22].

3) Choose a nominal value \( \theta_0 \) of the parameter vector \( \alpha \) at which to optimize the information (i.e. \( \alpha^* = \arg \max_{\alpha \in A} \mu(I(\alpha, \theta_0)) \)). This approach is conceptually simple and numerically tractable, but may suffer from a lack of robustness to the choice of \( \theta_0 \). In this paper, we take the third approach. We address the potential lack of robustness to the choice of the nominal parameter value \( \theta_0 \) by a systematic numerical study of the robustness to parametric uncertainty in Section IV-C. These
experiments demonstrate that for this particular model, an experiment designed using nominal values of the model parameters performs well across a wide range of values of the true parameter. It is possible that the results presented here could be improved further based on a minimax or Bayesian formulation, but this investigation is beyond the scope of this paper.

B. Computing Fisher information for our model

To find a maximum-likelihood estimate with minimum variance, we choose the sequence \( \alpha \) to maximize the Fisher information matrix at a nominal value of the parameter vector \( \theta \). The nominal parameter values used are given in Table I. The \( T \) relaxation, perfusion and metabolic rate parameters were chosen based on our typical data in a prostate cancer mouse model, and nominal values for the input shape and noise parameters were chosen based on a maximum likelihood fit to an arterial input function measured in a preliminary experiment.

\[
\begin{array}{cccccc}
K_{PL} & K_{NL} & k_{PL} & k_{TRANS} & l_0 & \gamma \\
1.2 & 0.2 & 0.055 & 3.296 & 2.143 & 3.4682 \times 10^{-4} & 1.0411 & 2.3608 \times 10^{-4}
\end{array}
\]

To compute the Fisher information, we use the expression derived in [23] for the \((i, j)\)-th entry of \( \mathcal{I} \):

\[
\mathcal{I}_{ij} = \sum_{t=0}^{N} \sum_{k=1}^{2} \frac{1}{\sigma_k^2} \frac{\partial x_k(t)}{\partial \theta_i} \frac{\partial x_k(t)}{\partial \theta_j} \psi \left( \frac{x_k(t)}{\sigma_k} \right) \tag{5}
\]

where the sensitivities are computed recursively as

\[
\begin{align*}
\frac{\partial x_k(t)}{\partial \theta_i} &= 0 \\
\frac{\partial x_k(t+1)}{\partial \theta_i} &= \frac{\partial x_k(t)}{\partial \theta_i} \begin{bmatrix} \cos \alpha_{1,t} & 0 \\ \sin \alpha_{1,t} & \cos \alpha_{2,t} \end{bmatrix} x_t \\
&\quad + A_d \begin{bmatrix} \cos \alpha_{1,t} & 0 \\ \sin \alpha_{1,t} & \cos \alpha_{2,t} \end{bmatrix} \frac{\partial x_k(t)}{\partial \theta_i} + B_d \frac{\partial u(t)}{\partial \theta_i}
\end{align*}
\]

and \( \psi \) is defined in terms of the integral

\[
\psi(z) = -z^2 + \int_0^\infty \frac{t^2}{2} \exp\left(-\frac{1}{2}(y^2 + z^2)\right) dy.
\]

C. Eliminating nuisance parameters

In practice, we do not necessarily need good estimates of all the unknown parameters in the model. For example, in this paper our primary goal is to estimate the metabolic rate parameter \( k_{PL} \) which is useful for discriminating between cancerous and non-cancerous tissues [5], determining the severity of disease [24], [25] and monitoring response to therapy [2]. Thus we wish to modify our optimality criterion to maximize the sensitivity of the experiments to \( k_{PL} \) while considering the nuisance parameters only insofar as they allow us to estimate the parameters of interest. We do so by partitioning the information matrix as

\[
\mathcal{I} = \begin{bmatrix} I_{11} & I_{12} \\ I_{21} & I_{22} \end{bmatrix}
\]

where the first block corresponds to the parameters of interest and the second block corresponds to the nuisance parameters. The inverse of the Fisher information is given by

\[
\mathcal{I}^{-1} = \begin{bmatrix} (I_{11} - I_{12}I_{22}^{-1}I_{12})^{-1} & -I_{11}I_{12}(I_{22} - I_{22}^{-1}I_{21})^{-1} \\ -I_{22}^{-1}I_{21}(I_{11} - I_{12}I_{22}^{-1}I_{12})^{-1} & (I_{22} - I_{22}^{-1}I_{21})^{-1} \end{bmatrix}
\]

Thus, optimal design for the parameters of interest can be performed by maximizing the Schur complement of \( I_{22} \):

\[
S = I_{11} - I_{12}I_{22}^{-1}I_{21}
\]

which corresponds to minimizing the asymptotic covariance of the marginal distribution of the MLE corresponding to the parameters of interest via (4), or equivalently, minimizing the Cramér-Rao bound on the parameters of interest via (3).

In general, if multiple parameters are of interest then \( S \) will be a matrix and we would be required to choose a suitable scalar criterion for measuring the size of \( S \). The problem of simultaneously estimating \( k_{PL} \) and \( k_{TRANS} \) is considered in [23], where the \( D- \), \( E- \) and \( A- \) optimality criteria are compared. However, in this instance we are considering a single parameter of interest \( k_{PL} \), therefore the Schur complement \( S \) is scalar-valued.

D. Regularization

We desire a smoothly-varying sequence of flip angles for a number of reasons including increasing robustness against model mismatch and interpretability of the resulting sequence of flip angles. We achieve smoothness in the flip angle sequence by adding a regularization term \( \lambda \| \Delta \alpha \|_F \) to the objective function to penalize nonsmooth sequences where the differencing operator \( \Delta \) is defined as

\[
\Delta = \begin{bmatrix} \alpha_{1,1} & \ldots & \alpha_{1,N} \\ \alpha_{2,1} & \ldots & \alpha_{2,N} \end{bmatrix}
\]

and \( \| \cdot \|_F \) denotes the Frobenius norm. The nonnegative parameter \( \lambda \) can be adjusted to achieve the desired degree of smoothness.

Regularization also improves the convexity of the objective function. When the regularization parameter takes the value \( \lambda = 0 \), the objective function possesses multiple local minima, but as \( \lambda \to \infty \) the convex term \( \| \Delta \alpha \|_F \) dominates, which guarantees that any stationary point is a global optimum.

We chose a particular value of \( \lambda \) by comparing the flip angle sequences resulting from the optimization using numerous regularization parameter values. The value \( \lambda = 0.1 \) was found to nicely balance between smoothness and range (between 0° and 90°) of the flip angle sequence.

E. Numerical optimization

To design an optimal flip angle scheme, we must solve the flip angle optimization problem

\[
\begin{align*}
\text{maximize} \quad & S(\theta, \alpha) - \lambda \| \Delta \alpha \|_F \\
\text{subject to} \quad & 0 \leq \alpha_{k,t} \leq 90^\circ \quad k = 1, 2 \quad t = 1, \ldots, N
\end{align*}
\]
for the flip angle sequence $\alpha$ where $\theta$ is fixed to some nominal value for the unknown parameters. The MATLAB Optimization Toolbox [26] provides a derivative-free implementation of the quasi-Newton optimization algorithm of Broyden-Fletcher-Goldfarb-Shanno (BFGS) [27], which is well-suited to finding local optima of this objective function.

F. Results

A solution to the optimization problem [9], initialized at $\alpha_{k,t} = 5^\circ$, is given in Fig. 3. Simulated state and observation trajectories corresponding to this flip angle sequence are shown in Fig. 4.

![Optimized flip angle sequence](image)

**Fig. 3.** Optimized flip angle sequence for estimating the metabolic rate parameter $k_{PL}$ using the nominal parameter values in Table I and a sampling interval $T_R = 2 \text{s}$ between acquisitions.

![Simulated trajectories](image)

(a) states $x_t$

(b) Rician-distributed measurements $Y_t$

**Fig. 4.** Simulated trajectories of the model (3) using the optimized flip angle sequence shown in Fig. 3 and the arterial input function shown in Fig. 2a.

We see that the pyruvate flip angles follow a pattern similar to flip angle sequences designed for other objectives, beginning with small flip angles to preserve magnetization for future acquisitions but increasing toward the end of the sequence [9], [10]. In contrast, the optimized flip angle sequence is much more aggressive with the lactate flip angles at the beginning of the experiment than in other variable flip angle sequences. This provides more reliable information about the leading end of the lactate time series, which contains the most information about the metabolic rate.

For the particular model and regularization parameter values used, the BFGS optimization algorithm converges to the same optimal sequence for a wide range of initializations. To confirm this, we have initialized the search algorithm using three flip angle sequences with angles generated randomly between 0 and 90°. For all three initializations, the algorithm converges to the flip angle sequence shown in Fig. 3. This demonstrates that the flip angle sequence presented is likely a global optimum.

IV. Validation Using Simulated Data

In this section, we demonstrate the advantage of the optimally designed flip angle sequence using computer-simulated data. Working with simulated data allows us to collect a large number of statistically independent data sets and provides us access to a “ground truth” value for the parameter vector. This makes it possible to reliably determine the parameter estimation error that results from noise in the simulated measurements. It is not feasible to acquire such a large number of data sets in vivo, and these would also not include ground truth values. Thus we use simulated data to demonstrate that our optimized flip angle sequence leads to smaller uncertainty in estimates of the metabolic rate parameter $k_{PL}$.

A. Two-step parameter estimation procedure

When fitting the data from in vivo experiments, data from different voxels will correspond to different values of the parameters $k_{TRANS}$, $k_{PL}$, $R_{1P}$ and $R_{1L}$ as these values change with spatial location, but all correspond to the same arterial input $u(t)$. Thus we present a fitting procedure that proceeds in two steps: first we fit a single input function $u(t)$ to the entire data set, then we fix this input function and estimate values of the remaining parameters individually for each of the voxels in the slice.

B. Simulation results and discussion

We wish to compare the reliability of estimates of $k_{PL}$ between data generated using five competing flip angle sequences:

1) a $T_1$-effective sequence [9] that aims to keep the measured signal constant despite repeated RF excitation and magnetization exchange between chemical compounds (Fig. 5a).

2) an RF compensated flip angle sequence [28] that aims to keep the measured signal constant despite repeated RF excitation (Fig. 5b).

3) a constant flip angle sequence of 15°.

4) a sequence that maximizes the total signal-to-noise ratio in the observed signal

\[ SNR_{total} = \sum_{t=0}^{N} \sum_{k=1}^{2} \frac{\tilde{x}_{k,t}}{\sigma_k}. \]

[10] (Fig. 5c), and

5) our flip angle sequence that maximizes the Fisher information about $k_{PL}$ (Fig. 3).

For each of the five flip angle sequences, we simulate $n = 25$ independent data sets from the model (2) using the parameter values given in Table I. We then perform the two-step parameter estimation procedure described in Section IV-A. The resulting parameter estimates are shown in Fig. 6. We see that
for all five flip angle sequences, the parameter estimates congregate near the ground truth value of the model parameters.

To demonstrate that our optimized flip angle sequence provides more accurate estimates of $k_{PL}$ than the competing flip angle sequences, we compare the root mean squared (RMS) estimation error between the sequences. WeRepeat this experiment for various values of the noise parameter $\sigma^2$ ranging from $10^3$ to $10^6$ to demonstrate that the improvement in the estimates is robust to variation in the noise strength. A value of approximately $2 \times 10^4$, in the center of this range, is typical for prostate tumor mouse model experiments. For each value of $\sigma^2$ we compute the RMS error of the $k_{PL}$ and nuisance parameter estimates across the $n = 25$ trajectories and plot these relationships in Figs. 7 and 8 respectively. The average improvement compared with competing sequences, across a range of noise parameter values $\{\sigma_p, p = 1, \ldots, 5\}$, is computed as a percentage

$$100 \times \left( \frac{1}{5} \sum_{p=1}^{5} \left[ 1 - \frac{RMS_{Fisher}(\sigma_p)}{RMS_{competing}(\sigma_p)} \right] \right).$$

These improvement percentages are summarized in Table II.

**TABLE II**

<table>
<thead>
<tr>
<th>Competing sequence</th>
<th>$T_1$ effective</th>
<th>RF compensated</th>
<th>Constant $15^\circ$</th>
<th>total SNR</th>
</tr>
</thead>
<tbody>
<tr>
<td>Improvement achieved</td>
<td>90.6%</td>
<td>23.1%</td>
<td>25.3%</td>
<td>19.8%</td>
</tr>
</tbody>
</table>

Overall, we see that the optimized flip angle sequence provides a more reliable estimate of the parameter interest $k_{PL}$, with a substantial improvement over all four competing flip angle schedules. The magnitude of the improvement varies from a 91% decrease in the estimation error against the $T_1$ effective sequence to a 20% decrease compared against the closest competitor: the maximum total SNR sequence. This improvement comes at the expense of less reliable estimates of some of the nuisance parameters. This highlights the advantage of using optimization-based methods to manage trade-offs in experiment design.
C. Robustness to parametric uncertainty

Based on the simulation experiments described in Section IV-B we have argued that flip angles optimized based on the Fisher information lead to smaller error in estimates of the parameter of interest \( k_{PL} \) when the same parameter values are used for the simulation and flip angle optimization. In this section we dispense with the latter assumption to demonstrate that this improvement is robust to uncertainty in the model parameters. We use the flip angle sequence shown in Fig. 3, which was designed using the specific model parameters given in Table I, along with two competing flip angle sequences to simulate data from models with different values for the parameters \( k_{TRANS}, k_{PL}, R_1P, R_1L \) and \( t_0 \) as well as a factor \( \Delta B_1 \) multiplying the flip angle sequence, used to demonstrate robustness to known \( B_1 \) inhomogeneities.

The ranges of the varying parameters were chosen to represent realistic physiological ranges and to center near the values we typically observe in a prostate tumor (TRAMP) mouse model. In Fig. 9 we plot the ensemble RMS error over \( n=25 \) simulated data sets as a function of the model parameter modified in the simulated data.

We see that the flip angles optimized based on Fisher information lead to better estimates of \( k_{PL} \) across nearly the entire range of model parameters used, despite no longer being optimal for the parameter values used to generate the data. This provides strong evidence that not exactly knowing the parameter values \( a \) priori does not limit the usefulness of our proposed flip angle design methodology.

V. IN VIVO EXPERIMENTS

We now move on to in vivo experiments. In contrast with the in silico experiments, here there is no ground truth value of the model parameters against which to compare our estimates, as the true rates are unknown and may vary between different regions of the tissue. However, the in vivo experiments can be used to show well-foundedness of the model that we have chosen and to demonstrate the feasibility of model-based parameter mapping using our optimized time-varying flip angle sequence. We show that our model can reliably reproduce observed data and achieve consistent parameter estimates across a variety of time-varying flip angle sequences.

A. Experimental setup

To implement this technique in vivo, metabolic data were acquired in a prostate tumor mouse (TRAMP) model using a 3T MRI scanner (MR750, GE Healthcare). Briefly, 24\( \mu \)L aliquots of \([1\text{-}13\text{C}] \) pyruvic acid doped with 15mM Trypti radical (Ox063, GE Healthcare) and 1.5mM Dotarem (Guerbet, France) were inserted into a Hypersense polarizer (Oxford Instruments, Abingdon, England) and polarized for 60 minutes. The sample was then rapidly dissolved with 4.5g of 80mM NaOH/40mM Tris buffer to rapidly thaw and neutralize the sample. Following dissolution, 450\( \mu \)L of 80mM pyruvate was injected via the tail vein over 15 seconds, and data acquisition coincided with the start of injection. Metabolites from a single slice were individually excited with a singleband spectral-spatial RF pulse and encoded with a single-shot symmetric EPI readout [29], with a repetition time of 100ms, a field-of-view of \( 53 \times 53\text{mm} \), a matrix size of \( 16 \times 16 \), an 8mm slice thickness, and a 2 second sampling interval. A \( ^{1}H \) image showing the anatomy contained in the slice in question is given in Fig. 10.

Fig. 10. \( ^{1}H \) image of the slice of interest. A large tumor, outlined in blue, fills a significant portion of the slice. Numbered volumes used to extract trajectories for parameter estimation are outlined in gold.
Datasets were acquired using three time-varying flip angle sequences. The $T_1$-effective and RF-compensated sequences, shown in Figs. 5a and 5b respectively, aim to distribute observed magnetization evenly across acquisitions, leading to roughly constant observed signals over time. The RF-compensated sequence does so by accounting for magnetization loss due to repeated RF excitation, but ignoring exchange between chemical compounds and $T_1$ relaxation \[28\]. In contrast, the $T_1$-effective sequence accounts for exchange and $T_1$ relaxation as well as RF excitation in attempting to achieve a flat time profile \[9\]. We compare these two sequences against our sequence, shown in Fig. 3, that has been optimized with the Fisher information-optimized flip angles, is shown in Fig. 11. Data collected using the optimized flip angles shown in Fig. 3 which corresponds to the spatial response of the RF pulse used experimentally. We assume that at each time step $t$ and for each compound $k$ (i.e. $k = 1$ corresponding to pyruvate, $k = 2$ corresponding to lactate) we can choose a real parameter $\alpha_{k,t}$ such that the flip angle applied at location $z$ is $\theta_{k,t}(z) = \alpha_{k,t}\pi(z)$. To generate a finite-dimensional model of the dynamics, we consider the magnetization dynamics at a discrete set of $z$ coordinates \{$z_1, \ldots, z_N$\}. The magnetization at location $z_i$ in the slice is then governed by the equations

$$x_{i,t+1} = A_d \begin{bmatrix} \cos(\alpha_{1,t}\pi(z_i)) & 0 \\ 0 & \cos(\alpha_{2,t}\pi(z_i)) \end{bmatrix} x_{i,t} + B_d u_t(p)$$

and the total magnetization measured is then assumed to be distributed

$$\bar{x}_t = \sum_{i=1}^{N} \frac{1}{z_{i+1} - z_i} \begin{bmatrix} \sin(\alpha_{1,t}\pi(z_i)) & 0 \\ 0 & \sin(\alpha_{2,t}\pi(z_i)) \end{bmatrix} x_{i,t}$$

$Y_{k,t} \sim \text{Rice}(\bar{x}_{k,t}, \sigma^2)$. This approach accounts for the slice profile effects by modelling the dynamics across the actual slice profile.

![Fig. 12. Comparison between an ideal flip angle profile across the slice and the actual profile for the RF excitation pulse used.](image)

### B. Resulting data

An example of the collected data, from the experiment with the Fisher information-optimized flip angles, is shown in Fig. 11. Experimentally-estimated values for the noise parameters are given in Table III.

![Fig. 11. Data collected using the optimized flip angles shown in Fig. 3. The field-of-view of these images is identical to the $^1$H image in Fig. 10.](image)

<table>
<thead>
<tr>
<th>Table III</th>
<th>Maximum-likelihood estimates of the noise parameter $\sigma_k$ for each of the three data sets collected</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>$T_1$ effective</td>
</tr>
<tr>
<td>$\sigma_k^2$ (pyruvate)</td>
<td>$1.84 \times 10^4$</td>
</tr>
<tr>
<td>$\sigma_k^2$ (lactate)</td>
<td>$2.14 \times 10^4$</td>
</tr>
</tbody>
</table>

### C. Flip angle profile modelling

Due to an imperfect (non-rectangular) slice profile, flip angles applied in practice vary spatially across the slice. This can lead to excess signal coming from regions near the boundary of the slice at later time points in the acquisitions, a phenomenon known as the slice profile effect \[30, 31\]. We have found that it is necessary to account for this effect in order to accurately fit the experimental data.

We consider the actual slice profile $\pi(z)$ shown in Fig. 12 which corresponds to the spatial response of the RF pulse used experimentally. We assume that at each time step $t$ and for each compound $k$ (i.e. $k = 1$ corresponding to pyruvate, $k = 2$ corresponding to lactate) we can choose a real parameter $\alpha_{k,t}$ such that the flip angle applied at location $z$ is $\theta_{k,t}(z) = \alpha_{k,t}\pi(z)$. To generate a finite-dimensional model of the dynamics, we consider the magnetization dynamics at a discrete set of $z$ coordinates \{$z_1, \ldots, z_N$\}. The magnetization at location $z_i$ in the slice is then governed by the equations

$$x_{i,t+1} = A_d \begin{bmatrix} \cos(\alpha_{1,t}\pi(z_i)) & 0 \\ 0 & \cos(\alpha_{2,t}\pi(z_i)) \end{bmatrix} x_{i,t} + B_d u_t(p)$$

and the total magnetization measured is then assumed to be distributed

$$\bar{x}_t = \sum_{i=1}^{N} \frac{1}{z_{i+1} - z_i} \begin{bmatrix} \sin(\alpha_{1,t}\pi(z_i)) & 0 \\ 0 & \sin(\alpha_{2,t}\pi(z_i)) \end{bmatrix} x_{i,t}$$

$Y_{k,t} \sim \text{Rice}(\bar{x}_{k,t}, \sigma^2)$. This approach accounts for the slice profile effects by modelling the dynamics across the actual slice profile.

### D. Parameter estimation

We begin by extracting time evolutions of the measured pyruvate and lactate signal from $n = 9$ voxels in the slice. The chosen volumes from which these signals are extracted are illustrated in Fig. 10. As in Section IV parameter estimation is performed in two steps. First, a single arterial input function and value for the parameter $R_{1L}$ are estimated based on the spatial average of the time series extracted from the tumour region. Second, the estimated input function and $R_{1L}$ value are held fixed while model parameters $k_{\text{TRANS}}$ and $k_{\text{PL}}$ are fit individually to the time series extracted from each of the voxels. To ensure practical identifiability of the model, the parameter $R_{1P}$ is fixed to a value of 0.05 during both steps.
E. In vivo results and discussion

Estimates of the arterial input corresponding to each of the three flip angle sequences are shown in Fig. 13. We see that the estimated inputs are reasonably consistent between the three data sets, but have some variation due to measurement noise in the pyruvate signal. Our optimized sequence yields the most smoothly-varying input function, which suggests that it is likely the most reliable of the three estimated AIFs.

Parameter estimates corresponding to each of the nine voxels are compared between the three flip angle sequences in Fig. 14. Examples of the quality of the fit corresponding to a particular voxel are shown in Fig. 15. We see that the estimated parameter values are consistent between the three flip angle sequences and that our model is able to reliably reproduce the observed data in all three cases. This provides evidence that the model we have used in this paper accurately describes the dynamics of magnetization exchange in vivo, and hence that the decision to use this model for the numerical reliability experiments of Section VI is well-founded.

We also present maps that show the spatial distribution of estimated metabolic and perfusion rates in Fig. 16. We see that the range and spatial distribution of parameter estimates are consistent between acquisition sequences.

VI. CONCLUSION

We have presented a method of generating optimal flip angle sequences for estimating the metabolic rate in a model of pyruvate metabolism. This method uses the Fisher information about the parameter of interest as the objective function that we wish to maximize. We have shown that the resulting flip angle sequence leads to smaller variance in the parameter estimates due to noise in the measured signal. We have demonstrated this in silico where we can explicitly compare the estimated model parameter values against the ground truth value. In this simulation experiment we demonstrated that our flip angle sequence leads to a 20% to 90% decrease in the uncertainty of the estimated metabolic rate, when compared with existing sequences. We also performed in vivo experiments to provide evidence that the model used in the in silico experiments is well-founded and demonstrate the feasibility of metabolic rate estimation and parameter mapping using this novel sequence. Based on the reliability results demonstrated in silico and the in vivo experiments demonstrating the appropriateness of the model used for the in silico experiments we argue that, for experiments that aim to quantitatively compare metabolic rates, optimizing flip angle sequences based on the Fisher information will probably lead to more reliable estimates of the model parameters of interest.

REFERENCES


