Low SAR $^{31}\text{P}$ (multi-echo) spectroscopic imaging using an integrated whole-body transmit coil at 7T

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Phosphorus ($^{31}\text{P}$) MRSI provides opportunities to monitor potential biomarkers. However, current applications of $^{31}\text{P}$ MRS are generally restricted to relatively small volumes as small coils are used. Conventional surface coils require high energy adiabatic RF pulses to achieve flip angle homogeneity, leading to high specific absorption rates (SARs), and occupy space within the MRI bore. A birdcage coil behind the bore cover can potentially reduce the SAR constraints massively by use of conventional amplitude modulated pulses without sacrificing patient space. Here, we demonstrate that the integrated $^{31}\text{P}$ birdcage coil setup with a high power RF amplifier at 7 T allows for low flip angle excitations with short repetition time ($T_R$) for fast 3D chemical shift imaging (CSI) and 3D $T_1$-weighted CSI as well as high flip angle multi-refocusing pulses, enabling multi-echo CSI that can measure metabolite $T_2$, over a large field of view in the body. $B_1^+$ calibration showed a variation of only 30% in maximum $B_1$ in four volunteers. High signal-to-noise ratio (SNR) MRSI was obtained in the gluteal muscle using two fast in vivo 3D spectroscopic imaging protocols, with low and high flip angles, and with multi-echo MRSI without exceeding SAR levels. In addition, full liver MRSI was achieved within SAR constraints. The integrated $^{31}\text{P}$ body coil allowed for fast spectroscopic imaging and successful implementation of the multi-echo method in the body at 7 T. Moreover, no additional enclosing hardware was needed for $^{31}\text{P}$ excitation, paving the way to include larger subjects and more space for receiver arrays. The increase in possible number of RF excitations per scan time, due to the improved $B_1^+$ homogeneity and low SAR, allows SNR to be exchanged for spatial resolution in CSI and/or $T_1$ weighting by simply manipulating $T_R$ and/or flip angle to detect and quantify ratios from different molecular species.

**KEYWORDS**

fast acquisition, in vivo, MRSI, quantification, RF, SAR, X-nuclei
Phosphorus ($^{31}$P) MRSI provides opportunities to monitor tissue metabolism by measuring specific energy metabolites and phospholipid metabolites. Phosphocreatine (PCr), adenosine triphosphate (ATP) (with α-, β- and γ-resonances) and inorganic phosphate (Pi) give insight into cell energy metabolism. Decreased PCr/ATP ratios in the heart can be used as diagnostic indicators in systemic diseases such as Type 2 diabetes and obesity. $^{1–3}$ Pi can be used to calculate tissue pH, as its resonance frequency changes with the acidity of the environment. Phosphomonoesters (PMEs) and phosphodiesters (PDEs) allow assessment of phospholipid metabolism. $^{4–6}$ At ultra-high field (>7 T), the increased signal-to-noise ratio (SNR) and increased spectral resolution facilitate the individual detection of PMEs (phosphocholine (PC), phosphoethanolamine (PE)) and PDEs (glycerophosphocholine (GPC), glycerophosphoethanolamine (GPE)). $^{7}$ Enhanced PME to PDE ratios (PC to GPC, PE to GPE) are indicative of proliferation and are often seen in tumor tissue. $^{5,6,8–13}$ Changes in these ratios during (chemo)therapy are markers of therapy response and take place well before morphological changes can be observed. $^{14–16}$

Still, these potential biomarkers are generally monitored to quantify metabolite concentrations or to investigate ratios between different molecular species, thus requiring solely a metabolite density-weighted signal. From proton MRI it is known that most clinically relevant contrast, when compared with proton density-weighted MRI, is obtained from $T_1$ and $T_2$ weighting. In fact, one study showed that in $^{31}$P spectra the $T_2$ relaxation itself may be used as a marker in breast cancer, and another study reported that intra-mitochondrial and cytosolic Pi can be discriminated based on $T_1$ differences. $^{17,18}$

However, current applications of $^{31}$P MRS are generally restricted to relatively small volumes such as the brain, breast and calf muscle, as small birdcage or conventional surface coils are used. $^{4,19}$ Conventional surface coils require high energy adiabatic RF pulses to achieve flip angle homogeneity, as inhomogeneous excitation leads to discrepancies in spectra over larger fields of view. Consequently, this can lead to high specific absorption rates (SARs), thus limiting the number of consecutive scans, particularly when considering metabolite relaxation parameter quantifications, fast spectroscopy methods or $T_1$- and $T_2$-weighted sequences.

Recent work by van der Kemp et al showed an adiabatic multi-echo spectroscopic imaging (AMESING) sequence, which included voxel-specific $T_2$ quantification of the different metabolites in the acquired spectrum. $^{8,18}$ This allowed $T_2$-weighted SNR enhancement, for an increased metabolite sensitivity, or $T_2$ information per metabolite. In the AMESING sequence, uniform excitation is achieved using adiabatic half pass RF pulses and homogeneous refocusing with adiabatic BIR-4 180° pulses, which require high $B_1^+$ (~100 μT) and relatively long pulse duration (8 ms). These pulses are therefore SAR demanding and consequently restricted to body surface applications.

Moving to larger fields of view in the body is therefore challenging, as greater $B_1$ field discrepancies are present with inhomogeneous excitation. Application of larger surface coils and adiabatic pulses would require even more power, which would limit the acquisition even more. In addition, the long $T_R$ times necessary for sufficient spin relaxation between pulses and for minimization of average SAR result in clinically impractical scan times for a single protocol.

To achieve uniform $B_1^+$ fields as with conventional $^1$H MRI, small X-nuclei RF-birdcage coils for head and extremities allow for diverse pulse sequences and enable numerous contrasts. Indeed even multi-echo acquisitions in the brain at 7 T are possible within SAR guidelines using these plug-and-play devices. $^{19}$

In another recent publication, Lörring et al showed an insertable $^{31}$P birdcage body coil that can produce uniform $B_1$ fields, thus allowing the use of rectangular RF pulse excitations. $^{20}$ This birdcage coil is wide enough to accommodate the human torso, allowing $^{31}$P MRSI of the human body, yet occupies space from the bore limiting inclusion of heavy patients and reduces space for receiver coils. Lörring et al did show the application of low flip Ernst angle excitations, with accompanying short $T_R$, which can result in fairly homogeneous excitation fields over the entire spectral bandwidth for in vivo $^{31}$P MRS at 7 T and acceptable scan times over a larger field of view.

In this work, we demonstrate that the permanent installation of a $^{31}$P body coil behind the covers of the patient tube, ie without sacrificing patient space, while interfacing to a high power RF amplifier, increases its usability. The reduction in SAR with this body coil allows the use of rectangular and even multiple rectangular composite pulses. Applications on the large gluteal muscle and liver are shown, including low flip angle excitation with short $T_R$ for fast 3D CSI and 3D $T_1$-weighted CSI, as well as high flip angle multi-refocusing pulses enabling multi-echo CSI, over a large field of view.

### 2 METHODS

#### 2.1 Coil setup

$^{31}$P-MRSI was performed using an in-house designed birdcage body coil, permanently integrated into a 7 T MRI system (Philips Healthcare, Best, The Netherlands), with a bore diameter of 60 cm for full body coverage. The coil, tuned at 120 MHz, was interfaced to and driven by a 25 kW RF amplifier for a $B_1^+$ field of 15 μT at the center of the bore (Figure 1A). $^{21}$ Two overlapping $^{31}$P receiver coils (10 × 16 cm$^2$) in quadrature mode and
two separate fractionated dipole antennas (30 cm) for proton imaging were used in quadrature transceiver mode. The proton antennas were positioned on the left and right sides of the $^{31}$P receiver coils, as can be seen in Figure 1A and 1B.

### 2.2 In vitro and in vivo setup

In vitro measurements were made on a body-sized phantom created from a barrel (diameter 27 cm, height 38 cm) filled with foam, normal saline and a small sphere (diameter 4 cm) filled with a 1 M Pi solution. The composition ensured loading matched to a human body for both the $^{31}$P coils and $^1$H antennas during measurements. For the in vivo measurements four healthy volunteers, three males and one female, with a body mass index (BMI) range of [21.6–26.5], were imaged in prone position with the gluteal muscles at the isocenter of the MR bore. The $^{31}$P receiver coil and proton imaging setup were placed on the gluteal muscles of the volunteers. One volunteer was imaged in the right decubitus position with the $^{31}$P receiver coils positioned at liver height, between the bed and the volunteer. The study was approved by the UMC Utrecht Medical Ethical Review Board and all volunteers gave written informed consent.

### 2.3 MR data acquisition

First, a proton image for anatomy localization followed by a $B_0$ map for image based $B_0$ shimming were obtained. To make sure that the flip angles were kept similar for all volunteers, a flip angle calibration with the carrier frequency set to PCr (ie set to 0 ppm) was made. The $^{31}$P $B_1^+$ field calibration was done with a non-localized block pulse sequence with a series of increasing flip angles and a $T_R$ of 2500 ms, which included gradient spoiling. The zero-crossing of the signal intensity, marking the actual 180° angle, eg no signal, was used to adjust output power.

Two fast chemical shift protocols using rectangular block pulses with the carrier frequency set to PCr, one with maximized signal intensity for Pi using the Ernst angle for Pi based on a $T_1$ of 4300 ms and another at a higher flip angle to increase $T_1$ weighting, were acquired at an isotropic resolution of 40 mm, matrix size $10 \times 6 \times 6$, 512 acquisition points, bandwidth 8000 Hz, $T_R$ 150 ms, flip angle, $\alpha$, 16° and 40°, number of averages 10 and included elliptical k-space sampling resulting in a scan duration of 7 min 3 s. A multi-echo spectroscopic imaging protocol (MESING), shown in Figure 1C, was used in order to acquire a single free induction decay (FID) by means of a pulse acquire and five full echoes in one k-space step, while k-space data were sampled spherically. The sequence was modified such that the excitation was performed using a rectangular 90° pulse at 15 $\mu$T, followed by a composite refocusing made up of three rectangular RF pulses of equal $B_1^+$ amplitude and flip angles of 57°–57°, 285°–57°, and 57°–57° for a refocusing bandwidth of 2 ppm. The refocusing part of the sequence is repeated five times to obtain five echoes. The carrier frequency of all pulses was set to Pi, and PCr for the in vitro and in vivo experiments respectively. The latter was based on the bandwidth of the refocusing pulses and the in vivo $^{31}$P metabolite with the highest concentration.
(PCr), which results in increased signal intensity favoring SNR. Both the in vitro and in vivo experiments with MESING were performed with an isotropic resolution of 40 mm with the carrier frequency set to P1 and PCr respectively. Other parameters were TR 5000 ms, ΔT 45 ms, bandwidth 7800 Hz, matrix 8 × 8 × 6, 256 acquisition points and a scan duration of 21 min 20 s. The in vitro experiment was used to validate the adapted protocol and applicability over a large field of view in vivo.

Liver spectra were acquired using a 3D 31P CSI protocol with Hamming-weighted acquisition at a 15 mm isotropic nominal resolution. The flip angle of 8° and a TR of 60 ms were chosen for optimal SNR with the Ernst angle for GPE and GPC assuming a T1 of around 6000 ms.23 The carrier frequency was set to PCr and other CSI parameters were T2 0.44 ms, bandwidth 4800 Hz, matrix 12 × 8 × 8, number of sampled averages 80 and 256 acquisition points for a total scan duration of 21 min 48 s.

2.4 | Data processing

All 31P MRSI data were processed in MATLAB 2017 (MathWorks, Natick, MA, USA). Calibrations of the 31P B1+ field from all volunteers were summarized by using the peak intensity of PCr of each flip angle dynamic scan obtained for each volunteer after apodization in the time domain with a Gaussian filter of 30 Hz. Data shown were normalized using the maximum signal intensity of each volunteer and the inter-subject variation was calculated from the variation in the periods of each individual fit per volunteer using

\[ \text{SI} = \frac{\sin(\alpha) \left(1 - e^{-\frac{T_2}{T_1}}\right)}{\left(1 - e^{-\frac{T_2}{T_1}\cos(\alpha)}\right)} \]  

3D multi-echo spectral data were spatially filtered using a 3D Hamming window, followed by inverse Fourier transformation to the spatial domain. FID and echoes were apodized using a 40 Hz Gaussian filter, and in vivo data were zero filled to double the number of samples thereafter. First order phase correction was applied to the FID by circular shifting the first missing data points resulting from the acquisition delay. T2 of the metabolites was calculated by fitting a mono-exponential model using the Levenberg–Marquardt algorithm:

\[ \text{SI} = S_0 e^{-\frac{T_2}{T_1}}. \]  

All other 3D CSI data were averaged, spatially filtered using a 3D Hamming window and transformed to the spatial domain by inverse Fourier transformation. The FIDs were apodized in the time domain with a Gaussian filter of 40 Hz and 24 Hz for the gluteal muscle and liver respectively and zero filled to double the number of samples. Phase corrections were applied manually thereafter.

3 | RESULTS

All subjects fitted well in the 31P whole-body coil setup, as this coil is integrated within the MRI scanner, behind the covers of the bore, providing sufficient space for the dipole transceivers and receive loops as shown in Figure 1A and 1B. The flip angle sweep acquired in four volunteers (different coil load) for sufficient space for the dipole transceivers and receive loops as shown in Figure 1A and 1B. The flip angle sweep acquired in four volunteers (different coil load) for sufficient space for the dipole transceivers and receive loops as shown in Figure 1A and 1B. The flip angle sweep acquired in four volunteers (different coil load) for sufficient space for the dipole transceivers and receive loops as shown in Figure 1A and 1B.

The integrated body coil in combination with the quadrature 31P receive loops showed high SNR 31P MRSI (3.9 for PME to 82 for PCr), as shown in the spectra in Figures 3–6. B0 shimming and partial volume effects were suboptimal over such large field of view, with a measured line width of 0.20 ppm before apodization. T1 weighting is apparent from the relative decrease of PCr and increase of α- and γ-ATP resonances in Figure 3C compared with Figure 3D. In addition, the β-ATP peak is decreased and the PDE peak shows a similar but minor decrease. An increase of the P1 signal is noticed in Figure 3D compared with Figure 3C.

The MESING data were acquired well within SAR limits and with sufficient SNR to allow T2 fitting (Figures 4 and 5). The MESING refocusing 180° composite block pulse used 15 μT and was 4.26 ms long, which compared with the 100 μT adiabatic RF pulse of 8 ms of the AMESING sequence used in the breast by van der Kemp et al resulted in an 83-fold reduced effective B1 integral. The in vitro average T2 of P1 in the phantom using the MESING method was 232 ± 35 ms (Figure 4) and the T2 of PCr from a single volunteer measured 177 ± 35 ms (Figure 5). Base-line roll artifact is visible in the FID in Figures 4C and 5C resulting from equal data processing of FID and echoes by correcting for the bandwidth difference. As the 1H antennas are inherently decoupled from the 31P coils, adequate 1H MR images for localization could be obtained, shown by the proton images in Figures 4B and 5B.
Averaging four local liver voxels from the 3D CSI protocol resulted in sufficient SNR to discriminate PD, PC, GPC, GPE, Pi and ATP resonances (Figure 6). The yellow voxels in Figure 6B indicate the voxels’ origin, and a residual opposite phased PCr peak, denoted by an arrow, is visible in the spectrum.

FIGURE 2  In vivo flip angle series showing the excitation profiles for all four volunteers showing the maximum PCr peak versus each flip angle scale factor. An average inter-subject variation of 30% was calculated. The BMI and gender of each volunteer is denoted in the legend. The $T_2$ relaxation effect is apparent from the asymmetry of the 90° versus 180° pulses and mirrored 360° pulse due to not fully relaxed spins. The zero-crossing of the fit for calculating correct power adjustments is marked by the larger black circle.

FIGURE 3  A, B, two spectral images of the in vivo 3D CSI data sets projected on the MR localizer image with a low flip angle (16°) (a) and a high flip angle (40°) (B) with the same $T_R$ (150 ms). Spectral data are normalized to the maximum signal and the $y$-axes of all voxels are scaled to the maximum signal in the 2D spectral images. C, D, two in vivo $T_1$-weighted spectra of the voxel highlighted by the yellow square in A (C) and the voxel highlighted in B (D), normalized to the noise of each spectrum. Data were acquired using the quadrature mode receiver coil setup in combination with the $^{31}$P body coil, and $T_R$ and flip angles were chosen to introduce $T_1$ weighting. The metabolite peaks of Pi, the PDEs, PCr and $\alpha$, $\beta$- and $\gamma$-ATP are labeled in both individual spectra.

Averaging four local liver voxels from the 3D CSI protocol resulted in sufficient SNR to discriminate PD, PC, GPC, GPE, Pi, and ATP resonances (Figure 6). The yellow voxels in Figure 6B indicate the voxels’ origin, and a residual opposite phased PCr peak, denoted by an arrow, is visible in the spectrum.
**DISCUSSION**

Multiple, SAR demanding, body oriented $^{31}$P MRSI methods were explored successfully using the fully integrated $^{31}$P whole-body coil at 7 T. Power calibrations of the homebuilt birdcage coil in multiple volunteers showed consistent performance with 30% inter-subject variability of coil load. Metabolic information from the gluteus maximus and the full liver was acquired, and the multi-echo CSI method was successfully implemented. Simultaneous use of the $^{31}$P receiver coils with the $^1$H transceiver antennas preserved volunteer comfort, as more freedom was experienced due to the lack of additional enclosing hardware that normally a $^{31}$P transmit coil would require. The $^{31}$P whole-body coil with uniform excitation in the body enabled the use of low demand SAR, conventional rectangular RF pulses instead of the high energy adiabatic RF pulses required with conventional surface coils for achieving homogeneous excitations. This decreases overall SAR, increasing the number of possible RF excitations per scan time to permit reduction of acquisition duration by decreasing $T_R$ or even the use of high flip angles, in $^{31}$P MRSI.

$B_1^+$ field homogeneity was assessed from designs by Lörring as our design is merely scaled to the bore size. The homogeneity of the insert was shown from the use of $B_1$ maps from 3 T proton MRI, which have identical diameter and coil layout to the present $^{31}$P body coil and are tuned to almost the same frequency.

Liver spectra were acquired over a large field of view and with minimal signal contamination by positioning the volunteer in the right decubitus position, weighted $k$-space acquisition and small voxel size. Increasing the number of sample averages regained SNR per voxel. This allowed discrimination of the mono- and di-esters PE, PC and GPC, GPE respectively.

MESING was validated on a phantom with $P_i$, as the average $T_2$ of 232 ms found corresponded to the $T_2$ of the body-sized phantom measured with the conventional AMESING sequence from van der Kemp et al (data not shown).$^8$ The in vivo application of MESING showed an average $T_2$ of PCr in the gluteal muscle of 177 ms ± 35 ms, which is comparable to the reported $T_2$ value of PCr in the calf muscle of around 217 ± 14 ms.$^8,24$ Note that the $T_2$ value measured by Bogner et al.$^{24}$ is an average for seven volunteers, where individual physiological differences between volunteers are averaged out, while our measured value in the gluteal muscle is an average from multiple voxels for just one volunteer, without averaging out possible physiological differences. Another possible cause for a difference in $T_2$ is sub-optimal refocusing pulses caused by imperfect power adjustments; however, the flip angle sweep in Figure 2 shows little variance between subjects, making it less likely to be the source of a lower $T_2$. A difference in physiology of the gluteal muscle compared with the calf muscle could result in a slightly higher chemical exchange rate between PCr and ATP, which leads to a lower $T_2$.$^{18,19,25}$

The $T_2$ relaxation property of the metabolites acquired with MESING provided a higher information density from the $^{31}$P spectra compared with a conventional MRSI experiment. Because metabolite specific MR properties are available, the signal of each individual metabolite of interest
can be corrected for $T_2$ blurring during acquisition, subsequently favoring SNR or used as a new contrast for each metabolite. This increases diagnostic significance and allows for new research in molecular dynamics and tissue environments. It can also be of interest to areas where $B_0$ shimming can be difficult, as the reduced spectral SNR caused by static $B_0$ inhomogeneities could be regained using the MESING method. In

FIGURE 5  A, B, in vivo $T_2$ fits of PCr using the MESING data from a single volunteer in voxels corresponding to the gluteal muscles (A) as shown by the red grid in the $T_1$-weighted localizer image (B). Normalized maximum peak value for the FID and each echo are denoted as open squares and the red dotted lines represent the fit. Echo times applicable to all voxels are shown in the bottom left. Average $T_2$ from all voxels with high muscle tissue content, denoted by the white dots, was $177 \mp 35$ ms. C, spectra of the FID and five echoes for the voxel highlighted by the blue square, acquired using the $^{31}$P dual coil receiver in combination with the $^{31}$P body coil. The frequency scaling shown for the $x$-axis of the FID is equal for all other echoes.

FIGURE 6  A, B, liver spectrum (A) after averaging all liver voxels from the 3D CSI protocol shown in the localizer image (B). Data were acquired with the $^{31}$P whole-body transmit coil in combination with the $^{31}$P receiver coils in quadrature mode. Metabolite peaks of PME, $P_i$, PDE and the three ATP resonances are denoted. The arrow points to the opposite phased PCr resulting from residual signal contamination from the muscles
conventional proton MRI, \( T_2 \) is an important biomarker to discriminate tumor from healthy tissue, aiding in diagnosis and disease prognosis. However, MRI focuses on morphological changes whilst metabolic changes occur prior to any observable structural alterations, creating opportunities for MRSI.\(^{26,27}\) \( T_2 \) contrast in MRSI, however, is still not available in the clinic but may increase insight into diseases when used as a biomarker including relaxation information for each metabolite separately. Though quantification of metabolite concentrations requires no transverse or longitudinal relaxation weighting, it has recently been shown by van der Kemp et al that shortening of the transverse relaxation time of \( P_i \) can be used as a biomarker in breast cancer spectroscopy.\(^{18}\)

In our \( T_1 \)-weighted \(^{31}\)P MRSI focusing on \( P_i \), we choose two \( T_R \) and flip angle combinations, which remained close to and deviated from the optimal Ernst angle condition for cytosolic \( P_i \), allowing for \( T_1 \) weighting with the latter condition. Other metabolites are \( T_1 \) weighted in both situations; however, the weighting is amplified for PCr, PME and PDE, with \( T_1 \) relaxation rates of the order of several seconds (\( \geq 3.1 \) s), whereas the optimal Ernst angle condition is almost met for the \( \beta^+ \) and \( \gamma \)-ATP resonances, with \( T_1 \) relaxation rates of around 1800 ms in the high flip angle experiment.\(^{24}\) As such, SNR remained high and \( T_1 \) contrast fair, as can be seen by the increased peaks of \( \gamma^+ \) and \( \alpha^+ \)-ATP resonances and decrease of PCr. The observed decrease of the \( \beta^+ \)-ATP peak in our measurements can be explained by the decreased excitation bandwidth at higher flip angle.

Figure 3C and 3D shows minor change between the two \( P_i \) peaks with respect to the noticeable decrease of PCr. Theoretically this could suggest an increase in signal contribution from intra-mitochondrial \( P_i \).

The adaptation of the RF pulses to operate at 15 \( \mu \)T rather than 100 \( \mu \)T comes at the cost of a lower pulse bandwidth. The implemented composite refocusing pulse used in MESING has a bandwidth of less than 240 Hz. However, setting the carrier frequency to PCr resulted in higher SNR compared with lower concentration metabolites and allowed validation of the adapted sequence in vivo. The use of multi-band RF pulses may be considered to broaden the bandwidth, or, in analogy with multi-slice TSE, rather than exciting slices sequentially within the \( T_R \), multiple narrow band excitations could be combined to cover the entire spectrum within the same scan time. As RF power deposition with conventional RF pulses is substantially decreased, more alternative pulse sequences, similar to pulses used in proton MRI, can be applied.

In our study we have used a two-channel receiver array, merely to demonstrate the feasibility of body-oriented \(^{31}\)P MRSI. When expanding the receiver array to a total of 16 or 32 elements, as already shown by Valković et al, full body coverage can be obtained.\(^{28}\) Combined information from multiple coils around the body could increase field of depth, as SNR and subsequently sensitivity can be regained by strategic coil combination methods such as whitened singular value decomposition.\(^{29}\) The space requirements for such setup may be comparable to those of conventional clinical MRI receiver arrays, where 16-channel body arrays are being used on a regular basis. The \(^{31}\)P receivers, as demonstrated here, can be merged with the relatively thin dipole antennas as can be seen in Figure 1A and aB, without efficiency losses.\(^{20}\)

While we have shown that a uniform transmit field with highly sensitive reception fields can be achieved with the whole-body coil and merged with a \(^1\)H imaging setup, care must be taken in optimizing scan protocols for motion artifacts and SAR, was successfully implemented for use in the body. The latter technique, though with improvements, allows for further research into new approaches in MRS biomarkers and additional metabolite specific information.

## 5 | CONCLUSION

The homebuilt fully integrated \(^{31}\)P body coil allowed \(^{31}\)P MRS methods to be explored that would have been SAR demanding with surface coils. Without sacrificing bore space, the improved hardware allowed full liver coverage \(^{31}\)P MRSI, and a multi-echo sequence, with inherently lower SAR, was successfully implemented for use in the body. The latter technique, though with improvements, allows for further research into new approaches in MRS biomarkers and additional metabolite specific information.

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