Magnetic resonance signals from different nuclei can be excited or received at the same time, rendering simultaneous or rapidly interleaved multi-nuclear acquisitions feasible. The advantages are a reduction of total scan time compared to sequential multi-nuclear acquisitions or that additional information from heteronuclear data is obtained at the same time and anatomical position. Information content can be qualitatively increased by delivering a more comprehensive MR-based picture of a transient state (such as an exercise bout). Also, combining non-proton MR acquisitions with $^1$H information (e.g., dynamic shim updates and motion correction) can be used to improve data quality during long scans and benefits image co-registration. This work reviews the literature on interleaved and simultaneous multi-nuclear MRI and MRS in vivo. Prominent use cases for this methodology in clinical and research applications are brain and muscle, but studies have also been carried out in other targets, including the lung, knee, breast and heart. Simultaneous multi-nuclear measurements in the liver and kidney have also been performed, but exclusively in rodents. In this review, a consistent nomenclature is proposed, to help clarify the terminology used for this principle throughout the literature on in-vivo MR. An overview covers the basic principles, the technical requirements on the MR scanner and the implementations realised either by MR system vendors or research groups, from the early days until today. Considerations regarding the multi-tuned RF coils required and heteronuclear polarisation interactions are briefly discussed, and fields for future in-vivo applications for interleaved multi-nuclear MR pulse sequences are identified.

**KEYWORDS**
interleaved, MRI, MRS, multi-nuclear, simultaneous, X-nucleus

---

**Abbreviations:**
- ASL, arterial spin labelling
- ATP, adenosine triphosphate
- BOLD, blood oxygenation level dependent
- CA, contrast agent
- CBF, cerebral blood flow
- CMRO$_2$, cerebral metabolic rate of oxygen consumption
- dMb, deoxymyoglobin
- DQF, double quantum filtered
- G6P, glucose-6-phosphate
- GRE, gradient echo
- HP, hyperpolarized
- Lac, lactate
- Mb, myoglobin
- MRSI, magnetic resonance spectroscopic imaging
- nOe, nuclear Overhauser effect (or enhancement)
- PCr, phosphocreatine
- PD, proton density
- Pi, inorganic phosphate
- PO$_2$, oxygen partial pressure
- SAR, specific absorption rate
- SNR, signal-to-noise ratio
- SVS, single-voxel spectroscopy
- T/R switch, transmit/receive switch
- UTE, ultra-short T$_E$
- ZTE, zero echo time

This is an open access article under the terms of the Creative Commons Attribution License, which permits use, distribution and reproduction in any medium, provided the original work is properly cited.

© 2022 The Authors. NMR in Biomedicine published by John Wiley & Sons Ltd.

DOI: 10.1002/nbm.4735

https://doi.org/10.1002/nbm.4735
1 | INTRODUCTION

MRI has become a major diagnostic tool in medical routine, and the main application modality of magnetic resonance in vivo. Beyond this, MRS has been established as a clinical research tool for brain disorders. The vast majority of these NMR applications performed today are based on exciting the magnetic moment of hydrogen (1H) nuclei. However, the application of NMR is not limited to 1H, as other physiologically relevant nuclei (e.g., 2H, 3He, 13C, 17O, 19F, 23Na, 31P or 129Xe) give rise to an NMR signal, but are less abundant and intrinsically less sensitive than 1H.

Such non-proton or ‘X-nucleus’ studies can provide complementary information not available from 1H NMR. For example, phosphorus-31 (31P) MRS has been employed as a tool to investigate intracellular pH and metabolism in vivo since its early days and continuously throughout, particularly in skeletal muscle. It has proven useful to study metabolites in liver, heart and brain as well as bone mineralization and oncology. Carbon-13 (13C) can provide information about the metabolism of glucose and glycogen in vivo. Deuterium-2 (2H) is also suited to evaluate glucose metabolism and as a tracer for cell tracking, monitoring of fluorinated drugs and as an alternative to hyperpolarized (HP) helium-3 (3He) and xenon-129 (129Xe) gases in functional lung imaging and ventilation studies or oxygen-17 (17O) to image and quantify the metabolic rate of oxygen consumption and as a tracer of cerebral blood flow (CBF). The viability of healthy and tumorous tissue can be studied with sodium-23 (23Na) imaging and spectroscopy, which is also a valuable tool for the diagnosis and research of kidney and cartilage defects.

Often 1H and X-nuclear MR data from the same subject are required, for instance to correlate high-resolution anatomic 1H images with metabolic information from X-nuclear MR or to confront different types of functional information based on different nuclei. Acquiring these datasets sequentially has several disadvantages. Most obviously, the acquisition time adds to the (costly) total scan time, with negative bearing on the subject’s comfort and cooperation. However, also comparison of datasets acquired during transient stimuli is hampered with sequential acquisitions because the stimulation and response may not be strictly reproducible as such; additionally, repeated stimulation may have undesired effects (e.g., fatigue or habituation). Data requiring an identical anatomical position, such as 1H, 3He or 129Xe images during ventilation studies, may also be challenging to obtain over separate breath-holds. Furthermore, with sequential acquisitions, X-nuclear MR cannot benefit from real-time adjustments derived from 1H MRI, such as navigators or dynamic shim updates.

Fortunately, the Larmor frequencies of the pertinent nuclei are at least several hundred kilohertz apart at clinically relevant field strengths and it is therefore possible to independently excite and receive signals from different nuclei at the same time. This allows for multi-nuclear acquisitions in a single scan by collecting data of each nucleus either truly simultaneously or in rapidly interleaved acquisitions.

The feasibility of the approach, which can help overcome the disadvantages of sequential measurements, was demonstrated by Thulborn et al. as early as 1981 and was then employed in several pioneering works in humans. The potential of reducing total measurement time was demonstrated in various studies and the possibility to obtain multiple datasets in a single measurement has been exploited by acquiring complementary data from transient states that are problematic to repeat precisely—for example, in exercising muscles during hypcapnia or in ventilation studies.

NMR sensitivity increases with magnetic field strength, motivating the trend towards higher B0 fields. The field strength of 3 T is becoming the standard for clinical scanners, while 7 T and above are becoming more widespread for research systems. This development is particularly interesting for non-proton MR, as X-nuclear MR examinations are now feasible in clinically relevant scan times, providing more specific data at higher temporal or spatial resolution than at lower fields. Consequently, the increasing availability of high-field MR scanners has renewed the interest in non-proton MR in general, a key prerequisite for simultaneous and interleaved multi-nuclear MR. The off-the-shelf hardware support of interleaved multi-nuclear measurements in modern clinical scanners has also contributed to the latest increase of interleaved applications.

In this review, a consistent terminology is proposed, in line with the literature on interleaved and simultaneous multi-nuclear MR in vivo. Technical obstacles and solutions are discussed, as well as the main applications and their advantages over conventional, sequential acquisition. Dual-tuned RF coils, necessary for multi-nuclear measurements, and potential heteronuclear interactions, such as nuclear Overhauser enhancement (nOe), are briefly discussed. Some perspectives for clinical applications using interleaved measurements are indicated to conclude this review.

2 | TERMINOLOGY

The topic of this review is the acquisition of datasets from different nuclei simultaneously or in close succession within a pulse sequence, in vivo. In agreement with the literature in this field, we suggest some consistent definitions:

- **Multi-nuclear**: describes acquisitions with more than one type of NMR-visible nucleus. This commonly refers to 1H and another nucleus, but combinations without 1H (References) or with three to four different nuclei (have been realized.
- **Non-proton X-nucleus**: designates MR measurements with any nucleus other than 1H.
Interleaved: a multi-nuclear measurement is considered ‘interleaved’ when different datasets are acquired sequentially within a short time, typically within the repetition time $T_R$ of a pulse sequence. The criterion is that the signal of only one type of nucleus is received at a time (Figure 1). The sequence elements (i.e., RF and gradient pulses) for the different nuclei are played out either consecutively without mutual overlap (Figure 1A) or interspersed before data are sampled, still consecutively for each dataset (Figure 1B). This latter variant has been termed ‘synchronous’ acquisition.\(^\text{58}\)

Simultaneous: multi-nuclear datasets can be acquired by receiving NMR signals of different nuclei truly simultaneously, that is, ADC sampling of signals with different resonance frequencies at the same time (Figure 1C).

It is worth stressing that the criterion for multi-nuclear interleaved or simultaneous acquisition lies in the reception of the NMR signal and not in the RF transmission for different nuclei.

The term ‘interleaved’ is also used outside the context of multi-nuclear MR: for example, for \(^1\)H imaging with different contrasts,\(^\text{49}\) parameters or slice positions\(^\text{50,51}\); combining MRS sequences sensitive to different metabolites, voxel positions\(^\text{52–54}\) or with added editing pulses\(^\text{32,55}\) or merging imaging and spectroscopy sequences into a single experiment.\(^\text{56–58}\) While such ‘interleaved’ techniques are not per se the topic of this review, they can be and have been combined with multi-nuclear interleaved measurements.\(^\text{32,34,55,59,60}\)

Finally, the terms ‘interleaved’ and ‘simultaneous’ have sometimes been used in the literature to describe measurements with different nuclei that were actually performed in consecutive scans\(^\text{61,62}\) and not even necessarily in the same scan session.

Other terminology has been used, for example, occasionally ‘time shared’\(^\text{63}\) for ‘interleaving’, or ‘parallel’\(^\text{64}\) for ‘simultaneous’. The latter is common terminology for high-resolution NMR in liquids and solids, but is uncommon with in vivo literature (where it would conflict with, e.g., ‘parallel imaging’). Interleaved sub-variants have also been defined for diverse polarization transfer and indirect detection methods, but these have not been applied in vivo.\(^\text{64}\)

### 3 | BENEFITS OF INTERLEAVED AND SIMULTANEOUS MULTI-NUCLEAR MR

#### 3.1 | Scan time reduction

The most obvious advantage of multi-nuclear interleaving is a reduction of the total scan time. Interleaved and synchronous measurements reduce the total duration by using the idle period of the first dataset acquisition for a second dataset acquisition. The waiting times present in $T_1$, $T_2$, or diffusion-weighted imaging, during the post-labelling delay of arterial spin labelling (ASL) measurements or simply to allow for longitudinal magnetization recovery are typical examples of idle periods suitable for secondary nucleus acquisitions. Also, different rates (i.e., different $T_R$ values) or MR signal recording times (i.e., the ADC sampling durations) can be implemented, taking into account the different relaxation times of nuclei or echo train lengths to optimize signal-to-noise ratio (SNR), while still reducing the total acquisition time.\(^\text{28,31,65–68}\) Simultaneous multi-nuclear acquisitions can further reduce the total sequence duration by overlapping the ADC recordings for the two nuclei, and may be particularly useful in measurements where little to no delay time is used, such as \(^1\)H and \(^23\)Na gradient echo (GRE) and ultra-short $T_E$(UTE) MRI.\(^\text{59,70}\)
Reducing the total acquisition time by interleaving has been achieved in the brain, \textsuperscript{30,65,70,71} knee, \textsuperscript{31,48} and breast. \textsuperscript{69} In a recent work, \textsuperscript{70} simultaneous \textsuperscript{23}Na and \textsuperscript{1}H radial imaging was used to acquire \textsuperscript{1}H T\textsubscript{1}, T\textsubscript{2}, proton density (PD) and B\textsubscript{1} maps using MR fingerprinting and \textsuperscript{23}Na density images, at 7 T in the brain (Figure 2). 

Reducing the total acquisition time by interleaving has been achieved in the brain, \textsuperscript{30,65,70,71} knee, \textsuperscript{31,48} and breast. \textsuperscript{69} In a recent work, \textsuperscript{70} simultaneous \textsuperscript{23}Na and \textsuperscript{1}H radial imaging was used to acquire \textsuperscript{1}H T\textsubscript{1}, T\textsubscript{2}, proton density (PD) and B\textsubscript{1} maps using MR fingerprinting and \textsuperscript{23}Na density images, at 7 T in the brain (Figure 2).

3.2 | Multiparametric information

Dynamic studies greatly benefit from acquiring multiple datasets simultaneously. Through interleaving, a single transient test can generate complementary information that can be readily combined to extract multiparametric biological variables that would be challenging to calculate otherwise.\textsuperscript{37,72} During an MR examination, the physiological response to a dynamic stimulus or precise lung inflation state may be difficult to reproduce. In certain cases the test might even be impossible to repeat in the same examination, notably following the injection of contrast agent (CA) or with patients showing slow or compromised physiologic recovery. Other examples of functional paradigms outside the brain include exercise bouts, muscle ischaemia or the administration of tracers, drugs or enriched substrates to study their biodistribution, metabolism or pharmacokinetics. Furthermore, measuring multiple MR parameters during stimulation can reveal alterations within the probed concomitant biological processes that may otherwise not manifest in a basal state\textsuperscript{73–75} and show the temporal relationships between them.\textsuperscript{32,55,76} The technique is particularly interesting in pathologies where compensatory biological adaptations could be masking a failing physiological variable, misleading the clinical diagnosis. For instance, patient cases with abnormally low mitochondrial adenosine triphosphate (ATP) production have been characterized using a multiparametric sequence interleaving \textsuperscript{1}H imaging and spectroscopy with \textsuperscript{31}P MRS,\textsuperscript{73} with mitochondrial diabetes clearly distinguished.
from peripheral arterial disease by the normal perfusion and myoglobin (Mb) resaturation profiles (Figure 3). Other conditions can also be evaluated, such as the impact of ageing, physical training, nutritional supplementation, drugs and so forth.

3.3 | Including dynamic $^1$H-based adjustments and navigators

The quality of X-nuclear MR data can potentially be improved by including dynamic adjustments derived from $^1$H MR. Examples are MR navigators, which can be used for prospective correction of respiratory and rigid bulk motion. Feedback-based motion tracking and correction, $B_0$ shimming and frequency correction can increase the robustness of measurements and provide the means for real-time quality control by rejecting or repeating data acquisitions compromised by motion. This is particularly useful for pulse sequences where artefacts are difficult to detect or to correct (as in magnetic resonance spectroscopic imaging, MRSI), during exercise paradigms in the magnet where movement-induced artefacts and $B_0$ variations are common or to alleviate examinations with patients experiencing difficulty in lying still. Fast $^1$H imaging can also be used for retrospective motion correction. Alternative non-MR motion correction methods track rigid-body movements only and require additional hardware. An example is a cardiac MRS study in humans, demonstrating $^1$H MR based volume tracking for compensation of respiratory motion to avoid contamination from chest wall and liver $^{31}$P MRS. Results from nine healthy volunteers measured at 1.5 T showed an average increase in fitting accuracy and signal amplitude with respect to the reference data. More recently, at 7 T, retrospective motion correction was applied to $^{23}$Na MRI of the human brain using interleaved $^1$H 3D navigator images, increasing the consistency between consecutive scans and improving the robustness of image quality against motion. Motion correction has further been exploited in rodents, for X-nucleus imaging of lung, heart and kidney.

3.4 | nOe, polarization transfer and $^1$H decoupling

Signal enhancement of low-sensitivity nuclei, such as $^{13}$C, $^{15}$N, $^{19}$F or $^{31}$P, can be achieved by exploiting the heteronuclear spin–spin or dipolar coupling interactions with $^1$H nuclei by means of nOe, polarization transfer or $^1$H decoupling. Although these methods can be applied without simultaneous or interleaved multi-nuclear signal reception and are therefore not per se the core topic of this review, they are closely...
related and can be combined. NOe has been frequently observed with interleaved sequences, as it can be induced by the pulses for the $^1$H acquisition, even without addition of dedicated NOe pulses. Polarization transfer requires deliberate adjustment of flip angles and echo time, taking scalar coupling constants into account. Heteronuclear decoupling is achieved by transmitting on one Larmor frequency while receiving on the other, and hence still allows for interleaving but conflicts with simultaneous acquisition.

The NOe originates from the dipole interactions with the saturated $^1$H nuclei, with the effective enhancement value depending on numerous experimental aspects including magnetic field strength, $^1$H irradiation intensity, biological tissue type and physiological state.\textsuperscript{96,97} The application of $^1$H decoupling, typically achieved using a WALTZ scheme\textsuperscript{101} during the X-nuclei read-out, collapses the split peaks of coupled resonances into singlets, greatly improving the sensitivity and simplifying spectral fitting. Decoupling pulses generate NOe by themselves but additional irradiation can be applied to achieve full NOe. During $^1$H polarization transfer, broadband RF pulses with appropriate phases and flip angles are played out simultaneously for both nuclei, enhancing the heteronuclear $J$-coupled resonances while removing uncoupled ones. The X-nuclei spectrum is thus simplified and the baseline is flattened. The sequence timings are chosen based on the $J$-coupling constant of the resonance of interest.

Unfortunately, the $^1$H irradiation needed in these techniques will increase energy deposition and unavoidably impact the $^1$H equilibrium magnetization. Though this might not be a concern in interleaved sequences where the $^3$H signal is solely used for motion or frequency corrections, it could be a limiting factor when $^1$H SNR is critical (such as MRS) or lead to bias in acquisitions employing magnetization-preparation modules (such as ASL or long-$T_1$-mapping MRI). Conversely, unintended heteronuclear NOe can be generated by the application of on-resonance RF-intensive $^1$H pulses, which is typically the case of magnetization-preparation modules or in sequences employing adiabatic pulses.\textsuperscript{60,71,102} Furthermore, a larger signal enhancement will result if the complete volume of interest of the X-nucleus measurement is irradiated.\textsuperscript{40}

While NOe can increase SNR and repeatability\textsuperscript{103} and provide biological information by itself,\textsuperscript{104} its magnitude will depend on the experimental setting but also on the tissue type and potentially on the pathological state.\textsuperscript{97} Therefore, similar NOe values between studies can be assumed only if the sequence parameters and experimental conditions are largely conserved. Acquiring reference data for NOe characterization will come at a cost in additional acquisition time, which will depend on the available X-nucleus SNR and encoding scheme. Estimating NOe in preparatory measurements is recommended,\textsuperscript{6,96} and will allow calculation of ‘NOe-free’ metabolite concentrations and ratios.\textsuperscript{60}

### 4.1 Basic principles

Simultaneous multi-nuclear acquisition offers true synchronicity of signal recording, at the cost of higher technical demands than interleaved acquisitions, and it causes dependences between acquisition parameters. While RF excitation and reception is independent between nuclei (setting aside heteronuclear polarization transfer, $^1$H decoupling and potentially NOe, as discussed above), magnetic field gradients always act on transverse magnetization and higher-order spin coherences of all spin systems. The gradient trajectory being identical during simultaneous acquisitions leads to different fields of view for nuclei with different gyromagnetic ratios, which can be corrected for by $k$-space regridding,\textsuperscript{67,70} as a gradient-linearity correction term to the MR system\textsuperscript{94} (when the gyromagnetic ratios are close, such as $^1$H and $^{19}$F) or, in principle, by setting the readout bandwidth per nucleus, via different ADC dwell times of separate receivers or by using sufficient oversampling. Similarly, simultaneous slice-selective excitation or refocusing results in identical imaging slab orientations while the slice profiles and thicknesses are controllable via the RF pulse shape and bandwidth, which can be set individually for each nucleus. The alternative approach of interleaving multi-nuclear acquisition relaxes the timing constraints to beyond the data acquisition duration (typically fractions of seconds) or repetition times. Interleaving can offer more flexibility with respect to field of view geometries, matrix sizes and repetition times, and even different types of acquisition scheme can be used: for example, combining $^1$H imaging and X-nucleus spectroscopy.\textsuperscript{32,34,72}

### 4.2 Requirements on the MR scanner

Several prerequisites on the MR scanner’s hardware and software have to be met for multi-nuclear interleaved or simultaneous acquisitions. The system must be able to transmit RF pulses at multiple resonance frequencies within one pulse sequence, either in rapid succession or simultaneously. This requires an RF transmit and receive system (including power amplifiers, multiple-tuned RF coils, interfaces and the signal acquisition chain from preamplifiers to sampling hardware) that can operate at different Larmor frequencies and allows for rapid switching between nuclei within a pulse sequence. Finally, the pulse sequence and data processing pipeline (e.g., inline image reconstruction systems) have to be implemented so as to drive the RF pulses and to record and store the NMR signal at the required frequencies.

The challenge is that many systems, even when ready for measuring X-nuclear data, are designed to acquire data of only a single nucleus within a pulse sequence. Most (clinical) MRI systems today are equipped with a power amplifier that can transmit RF within a narrow frequency
band at the scanner’s $^3$H frequency, sometimes wide enough for alternative $^{19}$F excitation. X-nucleus excitation is usually achieved with an additional broadband amplifier, often with lower peak power and usually lower maximum output frequency than the $^3$H amplifier. The MR system’s synthesizer frequency is mixed into the RF waveforms and then fed to the respective power amplifier. Transmission at two Larmor frequencies in one pulse sequence is fairly standard with X-nucleus capable MR systems, for heteronuclear polarization transfer, nOe or indirect detection. Monitoring the specific absorption rate (SAR) is mandatory on human MRI systems and must therefore be readily implemented by the manufacturer, also for multi-nuclear RF transmit. Therefore, no additional risk arises from using this capability for simultaneous or interleaved acquisitions. However, there is room for improvement in MR system and coil vendors’ SAR management, which may often be too conservative because local SAR differs between $^1$H and X-nuclei, and flip angle measurements are challenging with lower sensitivity. Unfortunately, systems capable of multi-nuclear transmit cannot necessarily receive signals from different nuclei in one scan and may require hardware modifications in addition to the adaptations of pulse sequences and reconstruction pipelines. Handling the timing within the sequence, increased complexity (e.g., when parametrizing the protocol), additional data reconstruction steps and higher total SAR demand may also constitute additional challenges, depending on the application.

4.3 RF coils

RF coils are used to apply RF pulses and to receive the MR signals. On human systems with field strengths of up to 3 T, $^1$H transmit is commonly achieved with a body coil installed in the magnet bore, and X-nucleus transmit is nearly always done with dedicated coils. Some systems disable the ($^1$H transmit) body coil when a local transmit coil is plugged in, making dedicated dual-frequency (X-nucleus and $^3$H) local transmit coils obligatory for $^3$H and X-nucleus RF measurements in the same examination. Body coils are not standard on ultra-high-field systems (although a $^{31}$P whole-body coil has been presented at 7 T, Reference $^{106,107}$), and dedicated coils are generally used for all nuclei.

In principle, simultaneous and interleaved measurements are not limited by the RF coil itself, as long as it comprises channels for both Larmor frequencies. It may be necessary to adapt coil-related software parameters, to allow the pulse sequence to activate the required transmit/receive (T/R) switches and preamplifiers at the necessary times. Further precautions should be taken, for example, to deal with transmission on one frequency while the preamplifier is active for the other, or to guarantee that this is avoided, to prevent hardware damage.

A practical difficulty of interleaved multi-nuclear applications is the increased complexity (and cost) of dual-tuned coils. Highly optimized $^1$H coils with a high channel count deliver maximum performance (high SNR, low mutual decoupling of elements and optimal placement for parallel imaging), but generally are proton-only coils. Dual-tuned coils are typically optimized for X-nucleus sensitivity, with $^3$H elements designed for scout imaging and $B_0$ field mapping. They typically have a lower $^3$H channel count and inferior performance than coils optimized for $^3$H MR only, $^{108-111}$ which may limit the potential of interleaved and simultaneous multi-nuclear applications. To improve the overall dual-tuned coil performance and to allow for acquisition of high-quality $^1$H data, innovative and organ-specific coil designs have been developed. $^{37,60,112-122}$

An in-depth discussion on the trade-offs for single- and multi-structure dual-tuned RF coils designs (focused on the brain but applicable to other anatomical targets) can be found elsewhere. $^{123}$

4.4 Implemented MR system solutions for simultaneous and interleaved multi-nuclear acquisitions

Simultaneous or interleaved signal reception has been realized in various ways by vendors and—in the early times of in vivo MR and later in cases where this was not possible on clinical MR scanners—by different research groups. The receiver of most MRI systems is based on the superheterodyne principle, that is, the signal is converted to an intermediate frequency of the order of a few megahertz, in one or several stages. The intermediate frequencies may or may not be different for different nuclei, according to the implementation by the manufacturer of the MR system. $^{125}$ Hence, simultaneous or interleaved multi-nuclear acquisitions may be possible straightforwardly (from the user perspective) or may necessitate hardware modifications.

An overview of the published implementation strategies for interleaved and simultaneous multi-nuclear MR is given in Table 1, which is structured into three categories: (1) Early experimental MR systems built by commercial vendors or by the research groups, (2) MR imagers designed for clinical routine that require hardware modifications and (3) commercial MR scanners, on which this is possible without or with only minimal hardware modifications (e.g., rerouting cables) by the user.

The early works on multi-nuclear interleaved and simultaneous measurements, particularly during the 1980s, profited from the research systems’ relative openness of the hardware and software, that is, those systems (described in terms of ‘spectrometer and data processing system’ rather than ‘MR scanner’) required—and allowed—low-level access to the hardware for operation. Solutions were to add spectrometers, $^{27,126}$ switches to alternate transmitters and receivers $^{63}$ or, e.g., ‘simply changing the synthesizer frequency under computer control’ (see Schnall et al., $^{128}$). Several groups had designed custom-built MR systems, foreseeing such capabilities. $^{28,43}$
Starting in the mid-1990s, multi-nuclear interleaved and simultaneous measurements were performed on large-bore human MRI scanners, but because the capability was not implemented by the manufacturers this required custom hardware adaptations. Solutions involved auxiliary spectrometers or even an additional full RF transmit chain. An alternative approach is to shift either the local oscillator frequency of the superheterodyne receiver or the frequency of the received NMR signal itself. That is, to receive a second NMR signal, either the frequency of the local oscillator signal provided to the mixing stage in the receiver cassette is appropriately set, or the NMR signal’s frequency is shifted using a mixer before being routed to the receiver. In both cases the resulting frequency at the digitization stage is what the system expects for acquisition of the default nucleus. This period saw declining publication activity in this field, which may well be a consequence of the technical and administrative difficulties arising from modifying the hardware of systems designed and certified for clinical applications.

Since 1999 and until today, vendors of pre-clinical and human research systems have been offering hardware solutions allowing for interleaved or simultaneous multi-nuclear MR. On clinical systems this became again possible without modifying the hardware about 10 years later, followed by a resurgence in publication activity involving human subjects after 2010. The vendor-specific solutions (e.g., on the Bruker Avance, Siemens VD and upwards, and Philips Achieva platforms) generally involve one or several constant (i.e., independent of the nucleus) intermediate frequencies during signal reception in a superheterodyne receiver. Today, direct digitization of the NMR signal is implemented in the most recent hardware generations (e.g., Philips dStream technology), which in principle allows for acquisition of the signals of multiple nuclei at a time. Throughout all periods, simultaneous and interleaved techniques were used, though the majority of publications (about three in four) report on the latter approach.

### 5 | CLINICAL AND RESEARCH APPLICATIONS

Interleaved and simultaneous acquisitions of multi-nuclear MRI and MRS have been applied in clinical studies and research applications in human and animal studies. Table 2 gives an overview of these applications.
5.1 Skeletal muscle

The vast majority of interleaved multi-nuclear papers so far published are reports of studies performed in skeletal muscle. The explanation is two-fold: first, skeletal muscle is the organ that experiences by far the fastest and greatest physiological and metabolic adaptations upon activation, and only interleaved acquisitions are capable of monitoring multiple physiological variables quasi-simultaneously, which is necessary to study their interactions with sufficient temporal resolution. Second, limb investigation is much less constraining in terms of spatial localization, which simplifies coil setup and sequence design.

The first interleaved $^1$H/$^31$P MR study of human skeletal muscle investigated the effect of hypoxia during an incremental knee-extension exercise, monitoring in parallel intramyocytic oxygen partial pressure (PO$_2$) calculated from the deoxymyoglobin (dMb) desaturation level, the high-energy phosphates and intracellular pH in the quadriceps.

The main contribution of the study to exercise physiology however was through integration of dMb-derived intramyocytic PO$_2$ with invasive determination of blood flow, arterial and venous PO$_2$ to determine for the first time the O$_2$ diffusional conductance at intermediate muscle O$_2$ consumption ($V_{O_2}$).

Also during an incremental knee-extension protocol with a very similar $^1$H/$^31$P setting (see Figure 4), electrically stimulated muscle contractions were compared with voluntary contractions.

| Organ | Species | Sequences | Type | References |
|-------|---------|-----------|------|------------|
| Muscle | Human   | $^1$H MRI + $^31$P MRS | int  | 34,77,141,144 |
|        |         | $^1$H MRI + $^1$H MRS + $^31$P MRS | int  | 32,60,72,74,76,146 |
|        |         | $^1$H MRS + $^31$P MRS | int  | 33,35,55,127,138-140 |
|        | Mouse   | $^1$H MRI + $^31$P MRS | int  | 59,136,137 |
|        | Rabbit  | $^1$H MRS + $^31$P MRS | sim  | 27 |
| Brain  | Human   | $^1$H MRS + $^31$P MRS | int/sim | 30,35,65,71 |
|        |         | $^1$H MRI + $^2$H MRS | int/sim | 28,68,70,92 |
|        | Cat     | $^1$H MRS + $^31$P MRS (+ $^23$Na MRI (+ $^{19}$F MRI)) | int | 46,47,128 |
|        | Rat     | $^1$H MRS or $^1$H MRI + $^31$P MRS or HP $^{13}$C MRS | int/sim | 63,142,148,149 |
| Lung   | Human   | $^1$H MRI + $^3$He (+ $^{129}$Xe MRI) | int  | 36,37 |
|        | Rat     | $^1$H MRI + $^{19}$F MRI | int  | 95 |
| Knee   | Human   | $^1$H MRI + $^{23}$Na MRI | int/sim | 31,48,134 |
|        | Rabbit  | $^1$H MRI + $^{19}$F MRI | sim  | 132 |
| Liver  | Rat     | $^1$H MRS + $^31$P MRS | sim  | 44 |
|        |         | $^1$H MRI + HP $^{13}$C MRS | sim  | 142 |
| Kidney | Mouse   | $^1$H MRI + HP $^{13}$C MRS | sim  | 67 |
| Breast | Human   | $^1$H MRI + $^{23}$Na MRI | syn  | 69 |
| Heart  | Human   | $^{31}$P MRS + $^1$H pencil navigators | int  | 90,91 |
| Whole body | Mouse | $^{19}$F MRI + $^1$H MRI motion correction | sim  | 94 |
|        | Rabbit  | $^{19}$F MRI + $^1$H MRI motion correction | sim  | 93 |
synthesis rate was lower in the patients during the insulin-dependent phase (second to fifth hour of recovery). In contrast, no statistically significant difference of the mean G6P concentration was found, despite being systematically lower in the control group. This technique was also used in a separate study, performed in healthy subjects, showing an increase in G6P concentration and glycogen synthesis during the first 15 min after heavy exercise but a reduced glycogen resynthesis rate for several hours in muscle with high glycogen concentration, suggesting an inhibiting feedback mechanism of glycogen in its resynthesis. A major step forward occurred with the addition of a perfusion imaging module to the $^1$H/$^3$P non-localized MRS sequence. In the first studies, skeletal muscle perfusion was measured with an MR version of venous occlusion

**Figure 4** Mean ± SE curve obtained from interleaved $^1$H/$^3$P MRS acquisitions during the different force steps (10, 20, 30, 40 and 50% of quadriceps maximal isometric voluntary torque) and the subsequent recoveries (R) of the voluntary contraction exercise session for PCr (A), [P]/[PCr] ratio (B), pH (C) and dMb (D). Figure reproduced with permission from Reference 140.
plethysmography, which was rapidly replaced by a more efficient pulsed ASL variant (SATIR). A spin echo blood oxygenation level dependent (BOLD) signal reflecting capillary oxygenation was also possible to obtain with SATIR. This sequence was used to investigate a number of conditions, as described in the following paragraphs.

In the field of exercise physiology, differences in skeletal muscle energy metabolism and perfusion control were documented between endurance and sprint athletes. Evidence was collected linking Mb concentration and energy metabolism efficacy. On the assumption that arterial O₂ content and mitochondrial oxidative coupling are normal, it was shown that multi-nuclear interleaving during the recovery phase of a plantar flexion bout could provide O₂ supply, uptake and consumption rates in the calf from ASL perfusion values, Mb resaturation and creatine rephosphorylation rates, respectively. By gathering these elements within the same physiological stress, the oxygen extraction rate could be calculated.

In relation with aging, it was demonstrated that in healthy elderly subjects the perfusion response to aerobic exercise was somewhat reduced as compared with young adults, but no difference in maximum mitochondrial ATP production was observed. However, during the exercise bout itself, adenosine diphosphate control of oxidative phosphorylation appeared to be slightly but significantly impaired. In a subsequent study, acute administration of an antioxidant cocktail was shown to improve both perfusion and mitochondrial ATP production during exercise recovery in the elderly subjects only.

The interleaved sequence was also able to reveal previously unidentified pathological mechanisms in Type 3 glycogen storage disease. In addition to a defective debranching enzyme activity, the patients have abnormal muscle perfusion response to moderate exercise. Combined analysis of dMb, BOLD, perfusion and PCr curves during exercise recovery concluded a role of perfusion in the lower ATP production, on top of the enzyme deficiency, that might contribute to the phenotype shift of the disease from childhood to adulthood. More anecdotally, the aetiology of abnormally low mitochondrial ATP production in mitochondrial diabetes and peripheral artery disease patients was characterized using this sequence. More recently, this interleaved sequence was implemented on a 3 T clinical scanner, without the need of any hardware modifications from the user side, other than employing a dual-tuned T/31P RF coil. The repeatability of the multiparametric acquisition during an ischaemia-hyperaemia paradigm and a plantar flexion exercise bout was assessed, while taking into account visit- and subject-specific nOe effects. In both paradigms, negative correlations were found between T₂* and pH during recovery and, at the end of exercise, the PCr depletion correlated with the percentage of Mb desaturation.

A sequence performing 1H T₂* mapping and adiabatic pulse-acquire 31P MRS was used on peripheral artery disease patients, finding a negative correlation between PCr recovery rate and the BOLD amplitude during hyperaemia. The sensitivity available with a 7 T human scanner was invested into improving spatial information of interleaved measurements by implementing, for the first time, multi-slice pulsed ASL in combination with multiple 31P semi-LASER voxels placeable at arbitrary positions. Two 31P spectra were acquired from the gastrocnemius muscle every 6 s (Figure 5A), while perfusion and T₂ contrast were measured in 10 slices (Figure 5B). The study showed that metabolic activity, which was recently found to vary significantly along a single muscle, was tightly coupled to haemodynamic changes measured during the same exercise: the significantly higher end-exercise PCr depletion, stronger pH drop and slightly elevated PCr recovery times were positively correlated with perfusion and T₂ changes measured in the gastrocnemius (Figure 5C).

Studying acid-base metabolism and glycolytic control requires concurrent quantification of 31P MRS-visible high-energy metabolites and lactate (Lac), which has 1H resonances that overlap with much stronger lipid resonances in muscle. A sequence interleaving non-localized 1H and 31P spectroscopy with 1H double quantum filtered (DQF) MRS for Lac detection was implemented at 3 T. Repeatability and feasibility were demonstrated in the tibialis anterior muscle during ischaemic dorsi-flexion exercise in healthy subjects. After ordering effects dominating the appearance of Lac resonances in anisotropic muscle tissue had been discovered, 1H and 31P STEAM-localized spectroscopy was interleaved with localized 1H DQF, taking muscle orientation into account. In this work, absolute quantification of Lac and phosphorylated metabolites was achieved in situ, during and following ischaemic plantar flexion exercise. Despite the complexities of Lac quantification in the presence of lipid signals and in anisotropic tissue, the method showed excellent agreement of estimated [Lac] with values quantified ex vivo. Consistent results from interleaved direct pH and Lac measurements and from indirect analysis of proton handling confirmed assumptions on cytosolic buffer capacity in vivo. The concept of dynamic investigations using interleaved multi-nuclear measurements was also adapted for small-animal skeletal-muscle applications at 4 T.

Myostatin inhibition causes an increase in muscle mass, but compromised force production has been reported in isolated mstn(−/−) muscle. Exerting the interleaved dynamic protocol on mstn(−/−) mice revealed a reduced oxidative mitochondrial capacity, a reduced BOLD contrast (indicating a possible decrease in oxygen extraction) and a prolonged hyperaemia response with respect to wild-type mice. Additionally, an increased proportion of Type Iib fibre and an unaltered capillary density were observed with histology, leading to the conclusion that the mstn(−/−) model has a non-pathologic shift towards glycolytic metabolism.

The effect of electropermeabilization was evaluated on muscle function using an empty plasmid 15 d after electroporation, considered as the end of the regenerative phase. Interleaved measurements showed altered perfusion and bioenergetics in electropermeabilized mice, whereas histological findings demonstrated a decreased number of Type Iib fibre but increased capillary density and number of Type I and Ila fibres. Although a decrease in 10% of cross-sectional muscle area was found, the specific muscle force did not change.
(A) Time-resolved multivoxel localised $^{31}$P MRS
gastrocnemius medialis

PCr time series and fit

recovery

exercise

rest

frequency / ppm

(B) Multislice $^1$H pASL and $T_2^*$-weighted MRI

perfusion

(60 – 300 s post exercise)

$T_2^*$ weighted signal increase

(post exercise, peak)

post ex. perfusion w.
signal / a.u.

slices

dist. prox.

slices

dist. prox.

post-exercise perfusion-weighted signal / a.u.

end-exercise PCr depletion / %

peak $T_2^*$-weighted signal increase / a.u.

end-exercise pH

$^{31}$P MRS
During the past decade, interleaving of $^1$H, $^3$He and $^{129}$Xe imaging has been developed for lung studies. The different diffusivity and solubility properties of $^3$He and $^{129}$Xe MR provide complementary information on ventilation, perfusion and lung microstructure, while $^1$H MRI provides anatomical and functional data. Interleaving enhances the complementarity of these methods by acquiring the datasets within the same physiological state, reducing spatial mismatches caused by variations in lung inflation or diaphragm position, and shortens the required breath-hold duration. Furthermore, acquiring $^1$H MRI anatomical data simultaneously with $^3$He or $^{129}$Xe images would allow their co-registration with anatomical CT images, the clinical gold standard in diseases, such as emphysema and cystic fibrosis, and in lung radiotherapy.

Wild et al. performed interleaved $^1$H and HP $^3$He MRI in vivo at 3 T using the scanner’s $^1$H quadrature body coil and a linear $^3$He Helmholtz coil, each coil actively detuning while the other one was active. GRE images of both nuclei were acquired during a 15 s breath-hold, in healthy subjects and in a patient afflicted by lung cancer and chronic obstructive pulmonary disease. When the $^1$H and $^3$He images were acquired in separate breath-holds, the ventilation volume overlap between repeated breath-holds was 87.4% and 86.7% for a volunteer and the patient, respectively. In the patient, despite the effort to replicate the breath-hold manoeuvre, misregistration was always visible. The authors noted that by interleaving the measurement of individual phase encoding lines of the $^1$H and $^3$He images (5 ms gap), motion misregistration errors were further reduced by limiting the effect of cardiac pulsatility. This work was later extended to include HP $^{129}$Xe imaging using a setup of electrically isolated RF coils comprising a flexible $^{129}$Xe quadrature vest transceiver inside an elliptical $^3$He birdcage coil nested inside the $^1$H body coil (Figure 6). The $^3$He and $^{129}$Xe coils’ tunings were verified while nested and with the load of a volunteer. By taking advantage of the different diffusivities of $^{129}$Xe and $^3$He, dual-gas imaging could be used to enhance detection of partial obstructions in the same inflation state while the ventilation volumes would be provided by the $^1$H anatomic images.

A triple-tuned RF coil with improved $^1$H reception was later created by the same group for 1.5 T use, although interleaved acquisitions were limited by the requirement of the new coil to manually activate the T/R switch of the nuclei.

Studies employing $^1$H and $^{19}$F MRI simultaneously with retrospective motion correction were performed in rabbits, mice and rats. The 6% larger field of view of $^1$H images, originating from the gyromagnetic ratio differences, was compensated during image post-processing. Lowering voxel resolution for increased SNR on $^{19}$F images was obtained by applying a spherical weighting to the image k-space, reducing its radius.
5.3 | Brain

Reductions of scan time and improvements of SNR per unit time have been achieved with interleaved acquisitions in human brain, in particular when interleaving 2D 1H MRSI with 3D 31P MRSI. SNR per unit time increased by 12% and 80% for the 31P and 1H datasets, respectively, for the same duration (50 min) compared with non-interleaved, serial acquisitions. 31P MR fingerprinting and 23Na MRI has also been carried out at 7 T (Figure 2). At 4 T, fluid-attenuated inversion recovery (FLAIR) images (14 slices) were acquired over 7 min while interleaving with 2H MRSI (13 × 9 × 11 matrix, spherical encoding, 2 averages), 60 min after an oral intake of [6,6'-2H2]-glucose.

Non-localized interleaved 1H and 31P MRS has been performed in the hypoxic cat brain and the ischaemic mouse brain to monitor concentration changes of ATP, PCr, P, and Lac as well as intracellular pH changes. In the mouse, PCr and ATP had completely depleted 10 min after the arterial occlusion. At the end of the 30 min ischaemia, Lac concentration had increased 10-fold (15.8 ± 2.5 μ mol/g) and pH had decreased from 7.14 ± 0.01 to 6.32 ± 0.10. About 1 h after reperfusion, metabolite concentrations had returned to baseline levels. A linear regression analysis showed a strong correlation (−0.97) between intracellular pH and [Lac].

Interleaved 1H PRESS (alternating between 30 ms and 136 ms T1 for Lac detection) and 31P slab-selective pulse-acquire spectroscopy measurements were made at 4 T in five awake humans during hypocapnia. During the 20 min-long hyperventilation period, a (modest) maximum increase in pH (0.047) occurred at the 14th minute, maximum Lac accumulation was reached 1 min later and only minor PCr (−3.4%) and P, (+6.4%) changes were observed. At the end of the 20 min recovery period, the partial pressure of carbon dioxide, pH and P, had not recovered to pre-hypocapnia values. The modest changes observed during hyperventilation, contrasting with studies carried out in anaesthetized animals, suggested an adaptive response of human brain to hyperventilation or a deregulation of cerebrovasculature under anaesthesia.

Taylor et al explored the ischaemic rat brain with an interleaved 1H PRESS/31P FID sequence. The interleaving measurements revealed that PCr responses to occlusion were quite similar between subjects, as previously thought, but the Lac responses showed higher inter-individual variability. By comparing the Δ([Lac]/[PCr]) ratio, two rat subgroups could be differentiated. The authors hypothesized that the low Δ([Lac]/[PCr]) value was an indicator of a reduced metabolic reserve of glucose and glycogen. Again, a strong correlation was found between [Lac] and pH (−0.85) during the 12 min ischaemia.

A study carried out in cats with interleaved 17O and 19F MRSI aimed to evaluate the cerebral metabolic rate of oxygen consumption (CMRO2) in a 0.8 cm3 voxel in the parietal cortex. While breathing a gas mixture of 17O2 and CHF3, CBF was measured with the inert CHF3 tracer while CMRO2 was estimated from the H217O concentration (above the natural abundance value) in the voxel and the measured CBF using a single-compartment model. From the seven studied animals, a wide variation of CBF and CMRO2 was observed, but a good correlation between CBF and CMRO2 was also found.

The study of metabolite kinetics using HP 13C in an organ is affected by non-specific signal contributions arising from vascular and extracellular compartments. To circumvent this difficulty, an injection of HP 13C-labeled pyruvate, administered intravenously, followed by a gadolinium-based CA was performed on the rat to isolate the signal from the intracellular compartment in the brain and liver. The pyruvate and Lac dynamics were then evaluated using a two-compartment model. Performing simultaneous 1H and 13C MRI allowed inclusion of the T1 variability (from the CA concentration) as an additional model parameter.

In humans, retrospective motion correction of 23Na MRI was recently performed using 1H 3D navigator image volumes. Navigator data had a temporal resolution of 6 s and matched the spatial resolution of the 23Na data. Both 23Na data consistency between consecutive scans and image quality were improved.

5.4 | Kidney

Simultaneous spectral–spatial Cartesian 1H and spiral HP 13C MRI acquisitions were used to track pyruvate and Lac dynamics in the kidney during free breathing. Every 5 s, two 1H images of water and fat, a 1D 13C spectrum used to measure the relative frequency of pyruvate and two 13C images of Lac and pyruvate were acquired. The 1H images were used to retrospectively compensate for motion during region-of-interest (ROI) positioning and to discard motion-corrupted images.

5.5 | Heart

Cardiac studies in humans have employed interleaved acquisitions for 1H image navigation to compensate for respiratory motion in 31P spectra. The effectiveness has been demonstrated using pencil-beam shaped 1H excitations at 1.5 T and has also been implemented with image-based navigators at 7 T. In rabbits, motion correction was implemented by simultaneous 1H and 19F MRI for imaging of the heart.
5.6 | Knee

In a study in the knee,\textsuperscript{33} the acquisition time of four different 3D datasets including $^{23}$Na images with or without the contribution of long-$T_2$ components, a $^1$H $T_2^*$ map and a three-point Dixon (GRE multi-$T_E$ acquisition) was halved, resulting in a total scan time of 23 min and 25 s when acquiring in interleaved mode. Another study\textsuperscript{48} also halved the acquisition time by simultaneously acquiring 3D UTE radial $^1$H and $^{23}$Na images.

5.7 | Breast

In breast imaging, simultaneous $^1$H and $^{23}$Na MRI reduced the acquisition time by half at 3 T.\textsuperscript{69}

6 | PERSPECTIVES FOR CLINICAL APPLICATIONS

The added value of multi-nuclear interleaving in combination with decreasing technical difficulties for its implementation, notably in the clinical setting, are incentives for the MR community to further invest in this methodology. The design and benefits of interleaved and simultaneous pulse sequences will nevertheless depend on the application, organ of interest and availability of a dual-tuned coil that fulfils the required sensitivity and spatial coverage.

| Target | Application | Gain | Multi-nuclear methods |
|--------|-------------|------|-----------------------|
| Oncology | Improved tumour characterization and monitoring | Reduced acquisition time | $^1$H + $^{31}$P MRS |
| Brain | Metabolic profiling and quantification | Reduced acquisition time | $^1$H + $^{31}$P MRS/MRSI |
| | Richer examination in bipolar disorder (Li concentration, membrane turnover, pH and Mg$^{2+}$) | Reduced acquisition time | $^7$Li + $^{31}$P MRSI |
| | Motion correction for long 3D $^7$Li MRSI acquisitions in bipolar disorder | Improved data quality | $^7$Li MRSI + $^1$H navigator |
| Muscle | Faster $^1$H and $^{31}$P examinations in neuromuscular diseases | Reduced acquisition time | $^1$H MRI + $^{31}$P MRS/MRSI |
| | Discrimination of alkaline P$\_r$ resonances in dystrophic muscle | Reduced acquisition time | $^1$H + $^{31}$P MRS |
| | Blood flow and energy metabolism evaluation in individual muscles | Multiparametric information | $^1$H MR + localized $^{31}$P MRS |
| | Simultaneous measurement of IMCL, glycogen and G6P synthesis and storage following exercise | Multiparametric information | $^{13}$C + $^{31}$P MRS, $^{13}$C + $^1$H MRS |
| Heart | Motion correction of localized $^{31}$P MRS | Improved spectral quality | Localized $^{31}$P MRS + $^1$H navigators |
| | Measurement of metabolic biomarkers (CK reaction, [PCr], [ATP]) | Reduced acquisition time | $^1$H MR + localized $^{31}$P MRS |
| Lung | Evaluate gas uptake and transfer times with anatomical or perfusion information | Multiparametric information | $^{129}$Xe MR + $^1$H MRI |
| | Continuous ventilation imaging in normoxia conditions with anatomical or perfusion information | Multiparametric information | $^{19}$F + $^1$H MRI |
| Liver | Combined fat fraction, IHCL and $^{31}$P MRS measurements in NAFLD and NASH | Reduced acquisition time | $^1$H MR + $^{31}$P MRS |
| | Motion correction from breathing in $^{31}$P MRSI acquisitions | Improved data quality | $^1$H navigator + $^{31}$P MRSI |
| Cartilage | Inclusion of $^{23}$Na imaging for improved detection of osteoarthritis and cartilage repair monitoring | Reduced acquisition time | $^1$H + $^{23}$Na MRI |
| Bone | Complementary quantitative mineral bone content and bone matrix density values for improved diagnosis | Reduced acquisition time | $^1$H UTE or ZTE + $^{31}$P ZTE |

Abbreviations: IMCL, intramyocellular lipid; CK, creatine kinase; IHCL, intrahepatocellular lipid; NAFLD, non-alcoholic fatty liver disease; NASH, non-alcoholic steatohepatitis.
Based on studies using classical ‘sequential’ sequences, a few multi-nuclear applications are briefly discussed below and are summarized in Table 3. Furthermore, in the specific case where the individual datasets acquired during multi-nuclear interleaving present very different VOIs, volume-specific $B_0$-shimming configurations could greatly improve data quality.\textsuperscript{161–163}

### 6.1 | MRS in brain and oncology

MRS has demonstrated its usefulness in classifying mass lesions and tumours and in monitoring their therapeutic treatment.\textsuperscript{12} The complementary information brought by $^1$H MRS alone during an MRI examination increased the rate of correct diagnoses by 15.4%.\textsuperscript{164} $^{31}$P MRS has also been used, to a lesser extent, for tumour classification and monitoring.\textsuperscript{12,165–168} Interleaving $^{31}$P MRS with $^1$H MRS could prove a useful application in oncology and in numerous brain studies by extracting complementary metabolic information within clinically feasible time constraints.\textsuperscript{29,30,71}

In bipolar patients under lithium administration, interleaving $^7$Li and $^{31}$P MRSI could simultaneously provide $^7$Li levels in tissue and information on free Mg\textsuperscript{2+}, pH and cell membrane anomalies.\textsuperscript{169,170} Moreover, long X-nucleus acquisitions, such as $^7$Li 3D MRSI (46 min, Reference\textsuperscript{170}), could benefit from $^1$H navigators for movement correction.\textsuperscript{81}

### 6.2 | Muscle

Evaluations in neuromuscular diseases typically include fat infiltration, muscle water $T_2$, lean mass and muscle cross-sectional area measurements using $^1$H MRI. $^{31}$P MRS is also included in mitochondrial myopathies, congenital lipodystrophy, muscular dystrophies and fibromyalgia.\textsuperscript{171,172} Cellular membrane damage and ‘leakiness’ in dystrophic muscle has also been evaluated by comparing $^1$H- and $^{31}$P-based pH values.\textsuperscript{173} Interleaving $^1$H MRI (or MRS) and $^{31}$P MRS could provide in these diseases a reduction in acquisition time. Glycogen detection by $^{13}$C MRS could also be combined with $^{31}$P or $^1$H MRS after a physical effort to simultaneously evaluate glucose transportation and phosphorylation, glycogen synthesis and the changes of lipids and glucose storage and utilization with respect to exercise and diet.\textsuperscript{66,129,174}

During a transient state, such as exercise, dynamic acquisitions interleaving fast multiparametric imaging schemes such as vPIVOT or SAGE\textsuperscript{160,175} with localized $^{31}$P single-voxel spectroscopy (SVS)\textsuperscript{33,34,176} could provide a more detailed evaluation of energy metabolism and oxygen consumption in individual muscles.\textsuperscript{72,177} Translation of DQF Lac MRS to ultra-high field\textsuperscript{178} has the potential to further increase sensitivity of interleaved $^1$H/$^{31}$P measurements\textsuperscript{55} to study acid–base metabolism and glycolytic control. By reducing the temporal resolution, fast MRSI modalities could replace the localized SVS module.\textsuperscript{57,179}

### 6.3 | Lung

Dynamic $^{129}$Xe MRS yields information on the surface-to-volume ratio and gas transfer times, while $^{129}$Xe MRI explores ventilation, regional gas uptake\textsuperscript{180} and alveolar-capillary exchange.\textsuperscript{181} $^{19}$F MRI has also been used for ventilation imaging under normoxic conditions at high temporal resolutions.\textsuperscript{182–184} The value of such $^{129}$Xe or $^{19}$F MR datasets could be enriched by interleaving them with anatomical\textsuperscript{185} or perfusion\textsuperscript{186} information by $^1$H MRI during single breath-hold or continuous ventilation.

### 6.4 | Heart

Localized $^{31}$P MRS provides relevant biomarkers (CK reaction, [PCr], [ATP]) in cardiomyopathies, diabetes, heart failure, aortic stenosis after valve replacement and during exercise paradigms.\textsuperscript{187,188} Interleaving $^{31}$P MRS with $^1$H MRI sequences is a viable clinical option as the individual datasets have similar acquisition lengths (\~10 min for gated $^{31}$P MRSI at 1.5 or 3 T\textsuperscript{189,190}). Other than examination length reduction, interleaving could enable navigators during $^{31}$P MRS, potentially increasing data quality and repeatability.

### 6.5 | Liver

$^{31}$P MRS(I) has been used to assess regenerative activity\textsuperscript{191,192} and to evaluate graft function following transplantation.\textsuperscript{193} Localized $^{31}$P MRS could benefit from MR navigators by reducing the impact of breathing, whereas interleaving $^{31}$P MRSI with $^1$H MR, for monitoring intrahepatocellular lipids or fat-fraction values, could reduce total scan time. These tools could constitute interesting clinical applications in prevalent diseases such as non-alcoholic fatty liver disease and non-alcoholic steatohepatitis.\textsuperscript{194}
6.6 | Cartilage

Changes in sodium concentration, an indirect measure of glycosamine sulfate proteoglycan (GAG) content, is evidence of early osteoarthritis and correlates with cartilage repair. In this context, $^{23}$Na imaging could be interleaved with $^1$H MRI for morphological or comparative information (UTE $T_2$, gagCEST$^{197,199}$) at a reduced total examination time and without requiring CAs.

6.7 | Bone

$^1$H UTE and zero echo time (ZTE) imaging provides information on the density and mechanical properties of bone matrix whereas high-resolution MRI has been used for microarchitecture imaging in trabecular bone, albeit at clinically long acquisition times. $^{31}$P MRI can provide mineral content information at the cost of long acquisition times (~ 20 to 37 min$^{9,10,201}$). The repetition times used in bone $^{31}$P ZTE MRI (~ 150 ms) could be used for interleaved $^1$H ZTE and high-resolution acquisitions for a reduced examination length. Combining bone matrix density and mineral content information could allow differentiation of osteoporosis from demineralizing disorders, a necessity for accurate diagnosis, intervention and monitoring of clinical responses to treatment.

7 | CONCLUSION

Interleaved and simultaneous multi-nuclear MR acquisition protocols have a wide range of applications, from reduction of total acquisition time to improved X-nucleus data quality by adding $^1$H-derived dynamic adjustments to multiparametric acquisitions within a single dynamic experiment, granting insights that are difficult or impossible to obtain by other means. While some early and experimental systems allowed for such measurements relatively straightforwardly, this became increasingly difficult on clinical MRI scanners in the past and required specific hardware modifications. Fortunately, the latest generation of MR systems of major vendors removed this hardware limitation, enabling interleaved or simultaneous multi-nuclear acquisition provided that the system supports X-nucleus measurements and a dedicated dual-tuned coil is available. The dual-tuned RF coil plays an important role, and multiple designs exist for optimal performance in a specific organ and application. The significant added value of interleaving for clinical applications and research, accompanied by the decreasing technical difficulties for its implementation, are major incentives to further invest in and standardize interleaved and simultaneous multi-nuclear acquisitions. While most results have been obtained in muscle, promising non-proton MR applications are abundant throughout different organs, particularly due to increasing sensitivity of MR systems, and new multi-nuclear MR applications can be envisaged to increase the value of clinical MR.

ACKNOWLEDGEMENTS

We thank Urs Sturzenegger for information on Philips systems and Guido Kudielka for information on GE systems. Support from the Austrian Science Fund FWF http://doi.org/10.13039/501100002428 (Grant P 35305-B) and from the French National Research Agency ANR (Grant 14-CE23-0005) is gratefully acknowledged.

DATA AVAILABILITY STATEMENT

Data sharing is not applicable to this article as no new data were created or analyzed in this study.

ORCID

Alfredo L. Lopez Kolkovsky https://orcid.org/0000-0002-9375-1465
Benjamin Marty https://orcid.org/0000-0002-4983-647X
Martin Meyerspeer https://orcid.org/0000-0002-0295-8218

REFERENCES

1. Oz G, Alger JR, Barker PB, et al. Clinical proton MR spectroscopy in central nervous system disorders. Radiology. 2014;270(3):658-679. doi:10.1148/radiol.13130531
2. Moon RB, Richards JH. Determination of intracellular pH by 31P magnetic resonance. J Biol Chem. 1973;248(20):7276-7278. doi:10.1016/S0021-9258(19)43389-9
3. Ackerman JJ, Grove TH, Wong GG, Gadian DG, Radda GK. Mapping of metabolites in whole animals by 31P NMR using surface coils. Nature. 1980;283:167-170. doi:10.1038/283167a0
4. Chance B, Eleff S, Leigh JS. Noninvasive, nondestructive approaches to cell bioenergetics. Proc Natl Acad Sci U S A. 1980;77(12):7430-7434. doi:10.1073/pnas.77.12.7430
5. Valković L, Chmelik M, Kršík M. In-vivo 31P-MRS of skeletal muscle and liver: a way for non-invasive assessment of their metabolism. Anal Biochem. 2017;529:193-215. doi: 10.1016/j.ab.2017.01.018

6. Meyerspeier M, Boesch C, Cameron D, et al. 31P magnetic resonance spectroscopy in skeletal muscle: experts' consensus recommendations. NMR Biomed. 2020;2020:e424. doi: 10.1002/nbm.4246

7. Purvis LAB, Clarke WT, Valković L, et al. Phosphodiester content measured in human liver by in vivo 31P MR spectroscopy at 7 tesla. Magn Reson Med. 2017;78(6):2095-2105. doi: 10.1002/mrm.26635

8. Henning A. Proton and multinuclear magnetic resonance spectroscopy in the human brain at ultra-high field strength: a review. NeuroImage. 2018;168:181-198. doi: 10.1016/j.neuroimage.2017.07.017

9. Wu, Reese TG, Cao H, et al. Bone mineral imaged in vivo by 31P solid state MRI of human wrists. J Magn Reson Imaging. 2011;34(3):623-633. doi: 10.1002/jmri.22637

10. Zhao X, Song HK, Seifert AC, Li C, Wehrli FW. Feasibility of assessing bone matrix and mineral properties in vivo by combined solid-state 1H and 31P MRI. PLoS ONE. 2017;12(3):e0192186. doi: 10.1371/journal.pone.0192186

11. Jagannathan NR, Sharma U. Breast tissue metabolism by magnetic resonance spectroscopy. Metabolites. 2017;7(2). doi: http://www.ncbi.nlm.nih.gov/pubmed/28590405

12. Payne GS. Clinical applications of in vivo magnetic resonance spectroscopy in oncology. Phys Med Biol. 2018;63(21):21TR02. doi: 10.1088/1361-6560/aae61e

13. Rothman DL, Magnusson I, Katzrobert LD, Shulman G, Shulman GI. Quantitation of hepatic glycogenolysis and gluconeogenesis in fasting humans with 13C NMR. Science. 1991;254(5031):573-576. doi: 10.1126/science.1948033

14. Gonen O, Hu J, Murphy

15. Gonen O, Murphy

16. Lee SW, Hilal SK, Cho ZH. A multinuclear magnetic resonance imaging technique for non-invasive assessment of their metabolism. J Magn Reson Imaging. 2010;257(1):175-184. doi: 10.1002/jmri.25002

17. Madelin G, Regatte RR. Biomedical applications of sodium MRI in vivo. Magn Reson Imaging. 2010;28(12):1751-1764. doi: 10.1016/j.mri.2010.02.017

18. Thulborn KR, Soffe NF. NMR spectroscopy of skeletal muscle and liver: a way for non-invasive assessment of their metabolism. Anal Biochem. 2017;529:193-215. doi: 10.1016/j.ab.2017.01.018

19. Mateescu GD, Ye A, Flask CA, Erokwu B, Duerk JL. In vivo assessment of oxygen utilization via deuterium magnetic resonance. Magn Reson Imaging. 2010;28(12):1751-1764. doi: 10.1016/j.mri.2010.02.017

20. Kreis F, Wright AJ, Hesse F, Fala M, En Hu D, Brindle KM. Measuring tumor glycolytic flux in vivo by using fast deuterium MRI. J Cereb Blood Flow Metab. 2017;37(11):3518-3530. doi: 10.1177/0271267817706444

21. De Feyter HM, Behar KL, Corbin ZA, et al. Deuterium magnetic resonance imaging (DMI) for MRI-based 3d mapping of metabolism in vivo. Sci Adv. 2018;4(8): eaat7314. doi: 10.1126/sciadv.aat7314

22. Ruiz-Cabello J, Barnett BP, Bottomley PA, Bulte JWM. Fluorine 19F NMR and MRI in biomedicine. NMR Biomed. 2011;24(2):114-129. doi: 10.1002/nbm.1570

23. Halawesheh AF, Moon RE, Foster WM, et al. Perfluoropropane gas as a magnetic resonance imaging contrast agent in humans. Chest. 2013;144(4):1300-1310. doi: 10.1378/chest.12-2597

24. Kruger SJ, Nagle SK, Couch MJ, Ohno Y, Albert M, Fain SB. Functional imaging of the lungs with gases. J Magn Reson Imaging. 2016;43(2):289-296. doi: 10.1002/jmri.25002

25. Zhu KH, Zhang Y, Wiesener HM, Uguribil K, Chen W. In vivo measurement of CBF using 17O NMR signal of metabolically produced H217O as a perfusion tracer. Magn Reson Med. 2013;70(2):309-314. doi: 10.1002/mrm.24469

26. Zhu KH, Chen W. In vivo 17O NMR imaging—quantitative assessment of regional oxygen consumption and perfusion rates in living brain. Anal Biochem. 2017;529(4):171-178. doi: 10.1016/j.ab.2016.08.026

27. Madelin G, Regatte RR. Biomedical applications of sodium MRI in vivo. J Magn Reson Imaging. 2013;38(3):511-529. doi: 10.1002/jmri.24168

28. Trattnig S, Welch GH, Juras V, et al. 23Na MR imaging at 7 T after knee matrix associated autologous chondrocyte transplantation preliminary results. Radiology. 2010;257(1):175-184. doi: 10.1148/radiol.2571081505

29. Thulborn KR, Soffe NF, Radda GK. Simultaneous in vivo measurement of oxygen utilization and high-energy phosphate metabolism in rabbit skeletal muscle by multinuclear 1H and 31P NMR. J Magn Reson. 1981;45(2):362-366. doi: 10.1016/0022-2364(81)90136-0

30. Lee SW, Hilal SK, Cho ZH. A multinuclear magnetic resonance imaging technique—simultaneous proton and sodium imaging. Magn Reson Imaging. 1986;4(4):343-350. doi: 10.1016/0730-725X(86)91044-1

31. Gonen O, Murphy-Boesch J, Srinivasan R, et al. Simultaneous and interleaved multinuclear chemical-shift imaging, a method for concurrent, localized spectroscopy. J Magn Reson Biomed. 1994;9(4):198-204. doi: 10.1002/jmri.19941050

32. Gonen O, Hu J, Murphy-Boesch J, Stoyanova R, Brown TR. Dual interleaved 1H and proton-decoupled-31P in vivo chemical shift imaging of human brain. Magn Reson Med. 1994;32(1):104-109. doi: 10.1002/mrm.1910320114

33. de Bruin PW, Koken P, Versluis MJ, et al. Time-efficient interleaved human 23Na and 1H data acquisition at 7 T. NMR Biomed. 2015;28(10):1228-1235. doi: 10.1002/nbm.3368

34. Brillault-Salvat C, Giacomini E, Wary C, et al. An interleaved heteronuclear NMR–NMR approach to non-invasive investigation of exercising human skeletal muscle. Cell Mol Biol. 1997;43(5):751-762.

35. Lopez Kolkovsky AL, Reynghoudt H, Giacomini E, Niess F, Meyerspeier M, Carlier PG. pH evaluation in the exercising muscle by interleaved localized NMR spectroscopy of carnosine and inorganic phosphate. In: 34th ENSMRB annual scientific meeting; 2017; Barcelona, Spain.956.

36. Niess F, Schmid Al, Bogner W, et al. Interleaved 31P MRS/1H ASL for analysis of metabolic and functional heterogeneity along human lower leg muscles at 7T. Magn Reson Med. 2020;83(6):1909-1919. doi: 10.1002/mrm.28088

37. Friedman SD, Jensen JE, Frederick BB, Artru A, Renshaw PF, Dager SR. Brain changes to hypocapnia using rapidly interleaved phosphorus–proton magnetic resonance spectroscopy at 4 T. J Cereb Blood Flow Metab. 2007;27(3):646-653. doi: 10.1089/jcbfm.20060083

38. Wild JM, Ajraoui S, Deppe MH, et al. Synchronous acquisition of hyperpolarised 3He and 1H MR images of the lungs—maximising mutual anatomical and functional information. NMR Biomed. 2011;24(2):130-134. doi: 10.1002/nbm.1565

39. Wild JM, Marshall H, Xu X, et al. Simultaneous imaging of lung structure and function with triple-nuclear hybrid MR imaging. Radiology. 2013;267(1):251-255. doi: 10.1148/radiol.12121153
199. Wehrli FW. Structural and functional assessment of trabecular and cortical bone by micro magnetic resonance imaging. *J Magn Reson Imaging*. 2007; 25(2):390-409. doi:10.1002/jmri.20807

200. Chang G, Boone S, Martel D, et al. MRI assessment of bone structure and microarchitecture. *J Magn Reson Imaging*. 2017;46(2):323-337. doi:10.1002/jmri.25647

201. Robson MD, Gatehouse PD, Bydder GM, Neubauer S. Human imaging of phosphorus in cortical and trabecular bone in vivo. *Magn Reson Med*. 2004; 51(5):888-892. doi:10.1002/mrm.20055

**How to cite this article:** Lopez Kolkovsky AL, Carlier PG, Marty B, Meyerspe M. Interleaved and simultaneous multi-nuclear magnetic resonance in vivo. Review of principles, applications and potential. *NMR in Biomedicine*. 2022;35(10):e4735. doi:10.1002/nbm.4735