Methodological consensus on clinical proton MRS of the brain

Wilson, Martin; Andronesi, Ovidiu; Barker, Peter B; Bartha, Robert; Bizzi, Alberto; Bolan, Patrick J; Brindle, Kevin M; Choi, In-Young; Cudalbu, Cristina; Dydak, Ulrike; Emir, Uzay E; Gonzalez, Ramon G; Gruber, Stephan; Gruetter, Rolf; Gupta, Rakesh K; Heerschap, Arend; Henning, Anke; Hetherington, Hoby P; Huppi, Petra S; Hurd, Ralph E; Kantarci, Kejal; Kauppinen, Risto A; Klomp, Dennis W J; Kreis, Roland; Kruiskamp, Marijn J; Leach, Martin O; Lin, Alexander P; Luijten, Peter R; Marjaska, Magorzata; Maudsley, Andrew A; Meyerhoff, Dieter J; Mountford, Carolyn E; Mullins, Paul G; Murdoch, James B; Nelson, Sarah J; Noeske, Ralph; Öz, Gülın; Pan, Julie W; Peet, Andrew C; Poptani, Harish; Posse, Stefan; Ratai, Eva-Maria; Salibi, Nouha; Scheenen, Tom W J; Smith, Ian C P; Soher, Brian J; Tká, Ivan; Vigneron, Daniel B; Howe, Franklyn A

Magnetic Resonance in Medicine

DOI: 10.1002/mrm.27742

Published: 01/08/2019

Peer reviewed version

Cyswllt i'r cyhoeddiad / Link to publication


Hawliau Cyffredinol / General rights

Copyright and moral rights for the publications made accessible in the public portal are retained by the authors and/or other copyright owners and it is a condition of accessing publications that users recognise and abide by the legal requirements associated with these rights.

- Users may download and print one copy of any publication from the public portal for the purpose of private study or research.
- You may not further distribute the material or use it for any profit-making activity or commercial gain
- You may freely distribute the URL identifying the publication in the public portal

Take down policy

If you believe that this document breaches copyright please contact us providing details, and we will remove access to the work immediately and investigate your claim.

A Methodological Consensus on Clinical Proton MR Spectroscopy of the Brain: Review and Recommendations

Martin Wilson, Centre for Human Brain Health and School of Psychology, University of Birmingham, Birmingham, England;
Ovidiu Andronesi, Martinos Center for Biomedical Imaging, Department of Radiology, Massachusetts General Hospital, Harvard Medical School, Boston, MA 02114, USA;
Peter B. Barker, Department of Radiology, Johns Hopkins University School of Medicine, Baltimore, MD, USA;
Robert Bartha, Robarts Research Institute, University of Western Ontario, London, On, Canada;
Alberto Bizzi, Neuroradiology Unit, Institute Clinico Humanitas IRCCS, Milan, Italy;
Patrick J. Bolan, Center for Magnetic Resonance Research, University of Minnesota, 2021 6th St SE, Minneapolis, MN 55455, USA;
Kevin M. Brindle, Department of Biochemistry, University of Cambridge, Cambridge, England;
In-Young Choi, Department of Neurology, Hoglund Brain Imaging Center, University of Kansas Medical Center, Kansas City, KS 66160, USA;
Cristina Cudalbu, Center for Biomedical Imaging (CIBM), Ecole Polytechnique Federale de Lausanne (EPFL), Lausanne, Switzerland;
Ulrike Dydak, School of Health Sciences, Purdue University, West Lafayette, IN, USA;
Uzay E. Emir, School of Health Sciences, Purdue University, West Lafayette, IN, USA;
Ramon G. Gonzalez, Department of Radiology, Massachusetts General Hospital, Harvard University, Boston, MA, USA;
Stephan Gruber, High Field MR Center, Department of Biomedical imaging and Image-Guided Therapy, Medical University of Vienna, Vienna, Austria;
Rolf Gruetter, Laboratory for Functional and Metabolic Imaging (LIFMET), Center for Biomedical Imaging (CIBM), Ecole Polytechnique Federale de Lausanne (EPFL), Lausanne, Switzerland;
Rakesh K. Gupta, Department of Radiology, Sanjay Gandhi Post Graduate Institute of Medical Sciences, Lucknow, India;
Arend Heerschap, Department of Radiology, Radboud University Medical Center, Nijmegen, the Netherlands;
Anke Henning, Max Planck Institute for Biological Cybernetics, Tuebingen, Germany;
Hoby P. Hetherington, Department of Radiology, University of Pittsburgh, Pittsburgh, PA, USA;
Petra S. Huppi, Department of Pediatrics, University of Geneva, Geneva, Switzerland;
Ralph E. Hurd, Stanford Radiological Sciences Lab, Stanford, CA. USA;
Kejal Kantarci, Department of Radiology, Mayo Clinic, Rochester, MN, USA;
Risto A Kauppinen, School of Experimental Psychology and Clinical Research and Imaging Centre, University of Bristol, Bristol, England;
Dennis W. J. Klomp, University Medical Centre Utrecht, Utrecht, the Netherlands;
Roland Kreis, Departments of Radiology and Biomedical Research, University of Bern, Bern, Switzerland;
Marijn J. Kruiskamp, Philips Healthcare, Best, the Netherlands;
Martin O. Leach, CRUK Cancer Imaging Centre, Institute of Cancer Research and Royal Marsden Hospital, London, England;
Alexander P. Lin, Center for Clinical Spectroscopy, Brigham and Women’s Hospital, Harvard University Medical School, Boston, MA, USA;
Peter R. Luijten, University Medical Centre Utrecht, Utrecht, the Netherlands;
Malgorzata Marjańska, Center for Magnetic Resonance Research, University of Minnesota, 2021 6th St SE, Minneapolis, MN 55455, USA;
Andrew A. Maudsley, Department of Radiology, University of Miami, Miami, FL, USA;
Dieter J. Meyerhoff, DVA Medical Center and Department of Radiology and Biomedical Imaging, University of California San Francisco, San Francisco, CA, USA;
Carolyn E. Mountford, Translational Research Institute, Woolloongabba, Australia;
Paul G. Mullins, Bangor Imaging Unit, School of Psychology, Bangor University, Bangor, Gwynedd, Wales;
James B. Murdoch, Canon Medical Research USA, Inc., Mayfield Village, OH, USA;
Sarah J. Nelson, Department of Radiology and Biomedical Imaging, University of California San Francisco, San Francisco, CA, USA;
Ralph Noeske, GE Healthcare, Berlin, Germany;
Gülin Öz, Center for Magnetic Resonance Research, University of Minnesota, 2021 6th St SE, Minneapolis, MN 55455, USA;
Julie W. Pan, Department of Neurology, University of Pittsburgh, Pittsburgh, PA, USA;
Andrew C. Peet, Institute of Cancer and Genomic Sciences, University of Birmingham, Birmingham, England;
Harish Poptani, Centre for Preclinical Imaging, Institute of Translational Medicine, University of Liverpool, Liverpool, England;
Stefan Posse, Department of Neurology, University of New Mexico, Albuquerque, NM, USA;
Eva-Maria Ratai, Department of Radiology, Massachusetts General Hospital, Harvard Medical School, Boston, MA, USA;
Nouha Salibi, MR R&D, Siemens Healthineers, Malvern, PA, USA;
Tom W. J. Scheenen, Department of Radiology, Radboud University Medical Center, Nijmegen, the Netherlands;
Ian C. P. Smith, Innovative Biodiagnostics, Winnipeg, Canada;
Brian J. Soher, Department of Radiology, Duke University Medical Center, Durham, NC, USA;
Ivan Tkáč, Center for Magnetic Resonance Research, University of Minnesota, 2021 6th Street SE, Minneapolis, MN 55455, USA;
Daniel B. Vigneron, Department of Radiology and Biomedical Imaging, University of California San Francisco, San Francisco, CA, USA;
Franklyn A. Howe, Molecular and Clinical Sciences, St George’s, University of London, London, England.

Correspondence
Martin Wilson, Centre for Human Brain Health, University of Birmingham, Birmingham, B15 2TT, England.
E-mail: wilsonmp@bham.ac.uk

Word count
8930
Abstract
Proton Magnetic Resonance Spectroscopy ($^1$H MRS) provides non-invasive, quantitative metabolite profiles of tissue and has been shown to aid the clinical management of several brain diseases. Whilst most modern clinical MR scanners support MRS capabilities, routine use is largely restricted to specialized centers with good access to MR research support. Widespread adoption has been slow for several reasons, and technical challenges towards obtaining reliable good-quality results have been identified as a contributing factor. Considerable progress has been made by the research community to address many of these challenges, and in this paper a consensus is presented on deficiencies in widely available MRS methodology and improvements that have been validated in the literature and are in routine use at several clinical research institutions. In particular, the degree of chemical shift displacement for the widely used point resolved spectroscopy (PRESS) localization sequence was found to be unacceptably high at 3T, and the use of the semi-adiabatic localization by adiabatic selective refocusing (semi-LASER) sequence is a recommended solution. The incorporation of simulated metabolite basis-sets into analysis routines is highlighted as a recommended method for reliably capturing the full spectral detail available from short echo time (TE) acquisitions. The importance of high quality shimming routines is emphasized and limitations of current hardware discussed. Most recommendations require only software improvements, greatly enhancing the capabilities of clinical MRS on existing hardware. We anticipate the implementation of these recommendations will strengthen current clinical applications and advance progress towards developing and validating new MRS biomarkers for clinical use.

Keywords
MRS; brain; consensus; semi-LASER; shimming; metabolites
Introduction
Proton Magnetic Resonance Spectroscopy ($^1$H MRS) has provided a non-invasive measure of brain metabolites since the late 1980s. Abnormal metabolism is often closely linked to disease processes, therefore MRS may improve clinical diagnosis, treatment effect monitoring and understanding of disease mechanisms (1). However, widespread clinical adoption has been slow, with MRS mainly used in specialized imaging centers.

Increased availability of 3T MR scanners in hospitals presents an important opportunity for MRS since metabolite levels are more reliably measured than at 1.5T due to reduced spectral overlap and an improved signal-to-noise ratio (SNR) (Figure 1) (2). However, additional challenges are associated with 3T MRS compared to 1.5T. Reduced homogeneity of the static magnetic field ($B_0$) at 3T necessitates improved shimming hardware and methodology. Chemical shift displacement (CSD) is also a significant problem, requiring higher $B_1$ radiofrequency (RF) power and/or optimized RF pulse shapes. Recent progress in the research community has addressed these challenges, but a major disparity remains between what is available for research and that for routine clinical applications. In addition, the wide range of MRS sequences, parameters and analysis choices can make the technique particularly difficult for non-expert users.

This paper was written and agreed upon by forty-nine MRS experts, belonging to the International Society for Magnetic Resonance in Medicine (ISMRM) MRS study group with the following aims:

1. To recommend appropriate methodology to improve the quality of future MRS studies and increase MRS standardization in order to facilitate clinical trials and meta-analyses of MRS efficacy.
2. To provide recommendations to vendors regarding the best MRS implementations and practices.
3. To focus the research community on resolving key technical barriers that have delayed wider clinical adoption of MRS.

Following initial discussions, an on-line survey was designed and completed by the group, with the results (supporting information section A) used to guide the further discussions leading to our final consensus and recommendations. We restrict our scope to $^1$H MRS detection of endogenous brain metabolites, using currently available methodology that we believe is ready for incorporation into clinical scanner platforms. We first present a basic introduction to $^1$H MRS methodology designed to give newcomers to $^1$H MRS sufficient background to understand the subsequent consensus sections.

Standard MRS Methodology

Single Voxel Spectroscopy (SVS)
Proton SVS is relatively simple to plan and yields clinically informative results, with robust acquisition procedures available on all commercial systems. SVS is currently the most commonly used method to acquire spectra from a single volume-of-interest (VOI or voxel); generally, a 5-minute acquisition time provides good quality spectra at 3T from tissue volumes.
down to 4 cm³. We briefly outline below the main technical considerations for widely available SVS methodology.

**SVS: acquisition methods**

STEAM (stimulated echo acquisition mode) (3,4) and PRESS (point resolved spectroscopy) (5) are widely used to provide single shot 3D localization from the intersection of three slices. In STEAM, three slice-selective RF pulses each with 90° nutation angles, produce a stimulated echo with typically a shortest echo time of TE≈20 ms. In PRESS, a 90° excitation pulse combined with two refocusing pulses produce a spin echo with a shortest TE of approximately 30 ms. PRESS refocusing pulses are nominally 180°, however some implementations use lower flip angles. Shorter TEs are possible for both STEAM and PRESS, but we restrict this section to commercial implementations. PRESS is more commonly used since its spin echo provides twice the signal compared to the STEAM stimulated echo. Cyclic changes in the phase of the slice-selective RF pulses and the receiver help to suppress outer-volume localization artifacts and those from imbalances in scanner electronics (6,7). Ensuring the last slice selection plane has regions of minimum B₀ inhomogeneity optimizes the overall crusher efficiency (8), particularly for MRS including frontal brain regions (7).

All localization methods based on RF/gradient slice selection, such as PRESS and STEAM, exhibit a localization inaccuracy known as the chemical shift displacement error (CSD). CSD causes a spatial displacement of metabolite resonance localization, where resonances further from the center frequency of the RF pulse are displaced to greater extent (Figure 2). Metabolites with frequency separated J-coupled multiplets, such as lactate, may also experience signal loss due to voxel edges not experiencing all three localization pulses (9). For a given maximum RF amplitude (B₁max), the CSD error worsens with increasing field strength and reduced RF pulse bandwidth. Therefore, CSD is worse for PRESS than STEAM, as conventional 90° pulses have greater bandwidth than conventional 180° pulses for a fixed pulse duration and B₁max. CSD severity and reduction will be described in the Consensus Opinion and Recommendations section.

**SVS: TR and TE considerations**

Proper determination of absolute metabolite concentrations or ratios (e.g. tNAA/tCr) ideally requires long TRs and short TEs to minimize signal loss from T₁ saturation and T₂ relaxation effects, respectively. Short TEs (20-30 ms) also offer improved detection of metabolites with complex J-coupled spectral patterns (e.g. glutamine (Gln), glutamate (Glu) and myo-inositol (mI)) due to reduced dephasing from J-coupling evolution. High quality short-TE MRS allows the quantification of an extended neurochemical profile, including neurotransmitters and antioxidants (10). However, a challenge associated with shorter TE’s is the enhancement of broad short- T₂ signals from high-molecular weight macromolecules (MM) and lipids (11). Appropriate analysis methods are capable of separating metabolite and lipid/MM signals, but poor shimming (see next section) degrades this separation; longer TE acquisitions may be employed to suppress broad MM and lipid signals, and simultaneously refocus weakly J-coupled spins. For instance, a TE of 144 or 288 ms is commonly used to aid the discrimination of lactate from lipids and singlets from overlapping multiplets. Whilst long TRs (>2 s) reduce unwanted T₁ weighting, the metabolite SNR per unit-time is also reduced and therefore a compromise is often required for clinical MRS.
SVS: shimming

In vivo susceptibility-induced magnetic field distortions arise from the presence of air/tissue and tissue/bone interfaces. In brain, field distortions are most apparent in regions close to sinuses such as the prefrontal cortex and temporal lobes, and since susceptibility differences scale with the static field strength, these distortions are stronger at higher fields. A homogenous static magnetic field, \( B_0 \), is essential for MRS, since narrow linewidths provide the spectral resolution critical for observation of multiplet resonances, accurate metabolite quantification and efficient water suppression. In addition to gradient coils, MR systems incorporate shimming coils to compensate for \( B_0 \) inhomogeneity and adjustment of currents flowing through these coils is called shimming. Figure 3 shows the effect of shimming in the frontal cortex. Poor shimming leads to loss of resolution between the singlets from total creatine (tCr at 3.0 ppm) and total choline-containing metabolites (tCho at 3.2 ppm) and to poor spectral separation between total \( N \)-acetylaspartate (tNAA at 2.0 ppm) and Glx (glutamine + glutamate at; \( \sim 2.3 \) and \( \sim 2.75 \) ppm) (Figure 3C). After successful shimming (Fig. 2E), multiplet peaks such as Glx and myo-inositol (\( \sim 3.6 \) ppm) are detected and quantified more reliably (Figure 1).

Vendor-provided shimming routines are currently based on the acquisition of a 3-dimensional \( B_0 \) field map (GRESHIM), \( B_0 \) field mapping along orthogonal projections (FASTMAP and its variants) \( (12–16) \) or along orthogonal planes of the localization VOI. \( B_0 \) field variations calculated from signal phase differences are used to compute the currents needed for each of the available shim coils \( (17–19) \). \( B_0 \) field distortions over typical SVS dimensions are generally compensated using a 1\(^{st} \) order shim (using the linear x, y, and z imaging gradient coils) at 1.5T, whereas the use of 2\(^{nd} \) order shimming elements terms \( (z^2, x^2-y^2, xy, xz, \text{ and } yz) \) is strongly recommended for SVS at 3T \( (20) \).

SVS: water suppression

Detection of millimolar metabolite signals in the presence of 3-4 orders of magnitude higher water (~40 M) signals is challenging due to the spectral baseline interference originating from the tails of the large water resonance. In addition, water “sideband” distortions, originating from various sources including subject movement, as well as mechanical vibration and instability of RF and gradient electronics, produce spurious signals further confounding metabolite signal estimates. Such interferences can be mitigated during acquisition by using a water suppression module prior to the MRS localization module. The most common methods exploit the chemical shift difference between water (4.65 ppm) and the strongest metabolite resonances (4.2 to 0.8 ppm). Common methods include a single chemical shift selective (CHESS) saturation pulse at the frequency of water \( (21) \); water suppression enhanced through \( T_1 \) effects (WET) \( (22) \) and variable pulse power and optimized relaxation delays (VAPOR) \( (23) \) shown in Figure 2. In general, the effectiveness of water suppression is significantly affected by \( B_0 \)-homogeneity as methods require narrow-bandwidth frequency selective pulses to avoid inadvertent suppression of metabolite resonances closest to the water resonance.

SVS: voxel dimensions and placement

In non-focal diffuse brain disease or general physiological studies, the SVS acquisition voxel is preferentially placed far away from air/tissue interfaces, because of shimming difficulties, and
away from the scalp to prevent spurious out-of-volume lipid signals. In a 5-minute acquisition time, an 8 cm³ cuboid volume in parieto-occipital grey matter (GM) or parietal white matter (WM) provides high-quality spectra at 1.5T and down to a 4 cm³ volume at 3T with good $B_0$ homogeneity. In focal diseases or specific anatomical areas of interest, voxels that best fit the targeted anatomy or lesion are commonly used. Contamination by lipid signals occurs for voxels located too close to the scalp, and trial voxel placement in volunteers to assess slice-profile and CSD limitations experimentally is advised. In addition to careful voxel positioning, outer volume suppression (OVS) may be used to suppress signal from unwanted regions in challenging areas. In general, volumes below 4 cm³ do not have an adequate SNR if detailed metabolic profiles are needed to answer clinical questions within a reasonable 5-minute acquisition time, but may be acceptable for restricted analysis such as the tCho / tNAA ratio in brain tumours. Voxel volumes greater than 8 cm³ or prolonged scan times are needed to detect small changes in low SNR metabolites such as Glu, Gln or glutathione.

**SVS: data analysis**

Gross spectral features are amenable to expert visual interpretation (e.g. lipids that indicate a high-grade tumor (24)), however, automated analyses of metabolite signals provide objective measures, such as relative metabolite concentrations or ratios, as biomarkers for clinical decision-making and clinical trials (25). MRS analysis using basis sets of known metabolites, MM and lipid signals parametrically fitted to the data with modelling of baseline and peak line-shape variations, have been particularly successful (example fit shown in Figure 1). Fitting may be performed in the frequency-domain, where baseline distortions may be modelled as smoothly varying spline functions (LCModel (26)) or decomposed using wavelets (MIDAS (27)).

Time-domain analysis may also be used to reduce errors from baseline signals by omitting initial data points from the fit - exploiting the rapid temporal decay of baseline distortions (QUEST (28), TARQUIN (29)). Metabolite basis sets can be experimentally acquired or simulated from known parameters (30–32) and either approach is effective (33,34). The addition of known MM and lipid signals to the basis set results in improved analysis, particularly for short-TE datasets or tumor spectral analyses (35). Cellular-scale and macroscopic $B_0$ inhomogeneities and eddy current effects distort the ideal line-shape (Lorentzian), and correction using the unsuppressed tissue water signal (36) or by its estimation during fitting (26) are commonly used.

**SVS: data quality**

Quality control (QC) requires consideration of 1) SNR; 2) metabolite and unsuppressed water resonance line-widths; 3) residual water signal; 4) line-shape; 5) Cramér-Rao lower bounds (CRLB) of the data-fit; 6) fit quality (relative size of residuals vs. noise standard deviation); and 7) presence of artifacts (spurious signals, baseline distortions, contamination from subcutaneous lipids). The first six can be automatically calculated by spectral analysis software, however the evaluation of artifacts currently necessitates inspection by an experienced reader of MRS (37). Current automatic QC suffers from a lack of evidence-based quality thresholds although general recommendations are available (1,38,39). An accepted numerical quality estimate in relation to model fitting of MRS data is the CRLB: a lower estimate of the error of the concentration measurement as influenced by SNR, linewidth, and mutual signal overlap. A relative CRLB >50% indicates that the estimated value is not significantly different from zero and is therefore often considered not to be reliable. However, it may also indicate that the
estimated value is too low to be reliably measured, which may be clinically significant (40). Influences from artifacts are not included in the CRLB calculation and these are discussed further in the supporting information (section B).

**SVS: absolute quantitation methodology**

The obtained MR signal is proportional to the number of contributing ¹H nuclei in the VOI, but the signal is not calibrated and measurement of absolute metabolite concentration is challenging due to unknown scaling factors such as receiver coil sensitivity and loading. For metabolite ratios, these factors cancel and for this reason metabolite ratios (e.g. tNAA/tCr) are widely reported for clinical use, despite the ambiguity of attributing changes to the metabolite in the numerator or denominator. Additionally, small or inaccurate values for the denominator produce large variance in a metabolite ratio. Semi-quantitative MRS can be obtained from the ratio of each metabolite to the unsuppressed tissue water signal from the same VOI and using an assumed MR visible tissue water content (41). Full concentration quantitation requires correction of water and metabolite signals for relaxation factors, assessment of any tissue partial volume effects (e.g. relative proportions of grey and white matter, cerebrospinal fluid (CSF) or pathological tissue in the VOI), and correction for disease-induced water concentration alterations (42). One challenge with concentration measures is that accurate knowledge of water and metabolite relaxation times are required, which may not be available for pathological tissue. In principle, using TEs shorter than 15 ms minimizes absolute quantitation errors of singlet peaks to less than 10% and reduces the dependence on assumed T₂ relaxation values; however, technical challenges with accurate localization at very short TE make this impractical with currently available commercial MRS sequences.

**Magnetic Resonance Spectroscopic Imaging (MRSI)**

While SVS is appropriate for investigation of a focal lesion, a specific anatomical region, or diffuse brain disease, MRSI is preferred when the location of interest is uncertain or multiple areas need to be evaluated simultaneously, for example when investigating metabolite distributions across heterogeneous lesions due to a tumor. In its most commonly used implementation at 3T, a 16x16 grid of spectra with nominal voxel resolution of 1.5 cm³ may be acquired in approximately 5 min at TR 1500 ms with one average per phase-encoding step and elliptical k-space sampling (see below). Despite the advantage of spatial metabolite information offered by 2D and 3D MRSI, robust acquisitions of good quality data present significant challenges. Practical issues include achieving adequate B₀ homogeneity over a large volume for good spectral resolution and reliable water suppression; good scalp lipid suppression; automated and accurate analysis of large multi-voxel datasets and their presentation for intuitive interpretation. In the following sections, we briefly introduce the technical issues surrounding MRSI acquisition and analysis.

**MRSI: volume localization and phase encoding**

The slice-localization methods described for SVS can also be used to define a larger volume in which 2D and 3D phase-encoding (see below) is performed for MRSI, while excluding contaminating signals from subcutaneous lipids. Within this context, methods such as PRESS or STEAM are known as a pre-localization. Outer volume suppression (OVS) may be used in combination with PRESS or STEAM localization to reduce spurious outer-volume and scalp lipid
signals; this creates a target volume that better conforms to the relevant tissue region and aids optimization of shimming and water suppression (Figure 4). A significant localization problem for MRSI is a large in-plane CSD since weaker gradient strengths are used for larger localization volumes. The displacement per ppm as a percentage of the localization width (%CSD) can be over 10% at 3 T, for example the 8 x 8 cm in-plane localization volumes for tCho and tNAA peaks will be shifted by 2.4 cm relative to each other when using conventional PRESS (Figure 5).

The phase-encoding localization method is based on the incremental adjustment of the strength of an applied magnetic gradient field, and may be applied to MRS acquisitions to create a “spectroscopic” image (43,44). Unlike slice-localization described above, localization by phase-encoding does not exhibit CSD effects or geometrical distortions due to B₀ inhomogeneity, and allows relative metabolite signal frequencies to be preserved independent of spatial position. Although data matrix sizes are small in MRSI compared to MRI, acquisition times become substantial as spatial resolution increases or if 3D data are required (e.g., over 51 min for 3D MRSI when acquiring a full 16 x 16 x 8 matrix with TR 1500 ms). To reduce acquisition times, higher frequency spatial components are sacrificed by limiting the k-space acquisition to a spherical or elliptical region, instead of a rectangular one, which reduces measurement time by 25% for 2D and 50% for 3D MRSI.

MRSI acquisition planning and results are typically displayed as a grid of voxels, where the nominal voxel dimensions in the phase encoded directions are determined by the field of view divided by the number of phase encoding steps. Unlike SVS, where it can be reasonably assumed that most/almost all signal originates from the prescribed voxel location, MRSI “voxels” may have a significant amount of signal contribution from outside the displayed grid boundaries or nominal voxel volumes. The point-spread function (PSF) describes precisely how the signal from the surrounding area contributes to a location on the MRSI grid. For phase encoded MRSI, the PSF is a complex sinc function and therefore the signal within a voxel includes positive and negative contributions that decrease with distance from the voxel center (Figure 6 parts a and c). Highly localized and strong scalp lipid signals can therefore contribute signal to many distant voxels (an effect known as signal “bleed”); causing significant spectral distortions and confounding estimation of lesion lipid levels.

MRSI: acquisition-based suppression of scalp lipid
Scalp lipid suppression may also be achieved with OVS, frequency-selective saturation, and inversion recovery (IR) methods, either in isolation or combined with PRESS or STEAM localization. OVS is typically applied via 8 perpendicular planes to the MRSI excitation arranged in an octagonal pattern around the scalp (45). OVS maintains target region signals, but is cumbersome to plan and has limited lipid suppression efficiency. The inversion recovery (IR) approach with a non-frequency selective adiabatic inversion pulse is relatively insensitive to B₀ inhomogeneity and eliminates the need for careful OVS prescription (46). A drawback of IR lipid suppression, however, is that T₁ relaxation also occurs for metabolites in the IR period, resulting in unwanted metabolite T₂ weighting and signal loss (approximately 25% at 3 T) that is greater the shorter the metabolite T₁. For many neurological disorders, intracerebral lipids are
not of interest, hence global lipid suppression via IR is inconsequential; however, lipid signals in the target volume are important diagnostically for tumors, stroke and lipid metabolism disorders.

MRSI: shimming
Adequate shimming is significantly more difficult for MRSI compared to SVS. The $B_0$ inhomogeneity must be corrected over a larger tissue volume, and $B_0$ inhomogeneity in regions close to the scalp or sinuses may exacerbate artifacts due to PSF effects. Whole brain 3D MRSI with 1st and 2nd order shimming may have up to 35% of voxels with insufficient data quality for analysis at 3 T (47). $B_0$ heterogeneity is so large across the whole brain that the required shim strength for optimal homogeneity increases by an order of magnitude in going from a slice above the ventricles to the temporal lobe (48). Therefore, the need for adequately powered shimming hardware with second-order coils and reliable software shimming algorithms is particularly important for MRSI.

MRSI: parallel imaging
For MRSI over large brain volumes, multi-channel RF coils provide improved sensitivity and enable the use of parallel reconstruction methods, such as sensitivity-encoding (SENSE), to improve spatial resolution or reduce scan times. Potential artifacts from parallel image methods include incorrectly localized signals, due to the imperfect reconstruction, which may not be visually obvious in low spatial resolution MRSI data. This can be notable for incompletely suppressed scalp lipids (49–51), which may be shifted in frequency due to $B_0$ inhomogeneity and aliased to obscure metabolite peaks. Self-calibration parallel imaging methods, such as GeneRalized Autocalibrating Partially Parallel Acquisition (GRAPPA), have also been applied to MRSI acquisition (52). The full impact of parallel imaging methods on MRSI metabolite quantification levels is still under investigation (47,53,54).

MRSI: post-processing
With multi-channel coil acquisitions, corrections for coil dependent signal strength and phase characteristics for each spectrum must be made to combine the data optimally (55–58). An MRSI scan of the unsuppressed water obtained at low spatial resolution can aid zero order phasing, frequency offset and line-shape corrections of the metabolite spectra without a major time penalty. With phased-array coils and SENSE, a water reference signal for metabolite quantitation may be acquired at the same spatial resolution as the metabolite scan, but with a significantly reduced scan duration as an alternative to a low resolution acquisition (59).

Following reconstruction, zero-filling of the k-space data is often used to reduce the apparent voxel size and produce smoother looking metabolite maps. However, this step only amounts to interpolation and does not reduce the effects of the PSF or increase spatial resolution. Spatial filtering is often applied to reduce Gibbs ringing artifacts from scalp lipid and from truncation artifacts associated with reduced k-space acquisitions (60). This also causes widening of the central lobe of the PSF resulting in improved spectral SNR, but reduced spatial resolution.

Figure 6 illustrates the influence of the commonly used Hamming filter on the PSF for typical MRSI acquisition parameters. Following zero-filling and filtering operations performed in k-space, spatial 2D and 3D Fourier transformation is then used to create a multi-voxel array of
free induction decays (FIDs), which can be analyzed with the methods described previously in the section “SVS data analysis”.

MRSI: data analysis and metabolite maps
2D and 3D MRSI produce large amounts of data, and robust automated data-processing to generate metabolite maps is needed for ease of clinical interpretation. Acquisitions at 3 T with phased-array head coils have greater spatial variation in receive sensitivity than studies at 1.5 T with quadrature or birdcage head coil designs, and at 3 T dielectric effects can compromise RF transmission homogeneity. Hence, using a semi-quantitative approach with reference to contralateral brain or a coil reference sample, are problematic. MRSI metabolite concentrations can be calculated if an additional water reference is acquired (42), however, the associated time-penalty may be significant (when using conventional methods) since all phase encoding steps need to be repeated. Metabolite ratio maps can be more robust than maps of “absolute” levels (albeit semi-quantitative if relaxation effects are ignored), as they are less sensitive to tissue partial volume effects with CSF. However, partial volume effects between tissue types such as grey and white matter (61,62) or normal and diseased tissues (63) are important to consider when interpreting the data.

MRSI: cautions and quality control
Automatically generated metabolite maps can be unreliable due to the effects of poor water suppression, lipid contamination and B₀ inhomogeneity, and they are also degraded by subject motion and inaccuracies of spectral fitting. Therefore, quality assessment of individual spectra and their fit is a required step for proper interpretation. However, visual quality assessment is often impractical with MRSI, resulting in a need for automated methods to exclude poor quality data. The relative CRLB derived from fitting each spectrum to a model function is misleading as a numeric estimate of data quality if there is a real absence of specific metabolites; on the other hand, good CRLB values may also arise from fitting bad quality spectra if the noise is underestimated, artifacts are present, or the fitting method has converged to an incorrect solution (local minimum). Alternative quality measures include CRLB values from a fit to a co-located water signal, using confidence limits (64) and linewidths from spectral fitting of metabolites or water, detection of outlying values in the spectrum (65), and use of pattern recognition to classify poor quality spectra (38,66,67). A quality map enables easy interpretation at the time of the clinical read, such as implemented in the MIDAS software (68) shown in Figure 7. Nevertheless, a visual assessment of spectra in key diagnostic locations is still advisable, and a zoomed-in grid of the raw spectra overlaid on the MRI of the abnormal region adds confidence to any interpretation of metabolite maps.

In addition to poor-quality data, visual interpretation of metabolite maps can present difficulties, particularly at the low spatial resolution of MRSI as compared to MRI, for which partial volume effects can be significant. Highly interpolated MRSI data can give a misleading impression of spatial detail, and PSF effects reduce the actual spatial resolution and may create artifactual “hot” or “cold” spots. B₀ inhomogeneity can cause localized signal loss, which may be obvious in temporal-frontal brain regions due to proximity to sinuses but can also occur in areas of blood breakdown products or calcification, and close to surgical entry points (e.g. craniotomy staples).
Finally, maps of metabolite ratios can include exceptionally high values because of a division by a small metabolite value and visual inspection is required to interpret these regions correctly.

Consensus Opinion and Recommendations

The discussion in the following sections is based on the results of an MRS technical consensus survey completed by 35 experts in clinical MRS (supporting information section A) and subsequent discussions among all authors. The survey aimed to determine what could be specifically recommended as best practice using the current standard implementations of $^1$H MRS and to define current limitations of scanner hardware and software. The survey also indicated the practical solutions that have been developed within the research environment. Key areas inhibiting the more widespread clinical use of MRS are: limitations in shimming algorithms, the practicalities of voxel planning and the time penalties associated with these processes, as well as reliable data processing and display; all of these issues are significantly exacerbated for MRSI. We indicate solutions to these issues and also provide guidance for recommended acquisition and data-processing protocols within the context of current scanner capabilities.

SVS acquisition

PRESS localization is the current standard for SVS and commercially available from all scanner manufacturers. We recommend a maximum CSD level of 4% per ppm, where % relates to the spatial displacement as a proportion of the slice selection width. This maximum recommended level of CSD is achieved at 1.5 T but exceeded in some implementations at 3 T due to insufficient bandwidth of conventional RF pulses resulting from a limited maximum transmit field $B_{1}^{\text{max}}$. Increased CSD makes planning and interpretation of SVS more challenging due to different metabolite signals being localized to different volumes. Therefore, methods to reduce CSD, whilst maintaining good SNR, short TE’s and accurate localization, should be employed to achieve the full advantages of clinical MRS at 3 T. One of the first approaches to reduce CSD was the use of Very Selective Saturation (VSS) pulses that have high-bandwidth and can suppress an outer volume signal with minimal CSD. Conventional PRESS is used to excite a larger (approximately 20%) region than required, and VSS pulses redefine a smaller region with minimal CSD (69,70), a method known as OVERPRESS.

More recently, the semi-LASER (71,72) sequence has been developed with a design similar to PRESS - slice selection excitation followed by two slice selective refocusing elements. However, in the case of semi-LASER, a pair of adiabatic refocusing pulses is used in place of each conventional 180° pulse. Higher order hyperbolic secant adiabatic full passage (HS) or gradient-modulated offset-independent adiabaticity (GOIA) pulses may be used for refocusing in semi-LASER, with the latter employing wideband, uniform rate and smooth truncation (WURST) RF and gradient waveform modulation (73,74). The advantage of GOIA pulses compared to HS is a reduced maximum $B_1$ strength required for a given pulse bandwidth, enabling semi-LASER to be used at short TE (~30 ms) on 3T systems with a maximum available $B_1$ strength of 13-15 $\mu$T (75). In addition, GOIA semi-LASER has a reduced RF power deposition compared to HS pulses of the same bandwidth – enabling the sequence to be used with a shorter TR and at higher field strengths within SAR limits. Whilst adiabatic pulse pairs offer improved resilience to $B_1$ inhomogeneity, their primary advantage for SVS at 3 T is increased bandwidth and slice
selection profiles, thereby reducing CSD to acceptable levels. A further advantage of semi-LASER compared to PRESS for the same TE, is longer apparent $T_2$ relaxation times and partially suppressed $J$-coupling evolution, due to the CPMG-like refocusing pulse train, hence the enhanced detection of complex multiplets such as glutamate (72,76).

Overall, we recommend the use of semi-LASER due to its recent validation studies (77–82) and increasing availability on clinical systems as both research work-in-progress sequences and commercial product (Figure 2). Furthermore, in our consensus survey, semi-LASER was ranked as the most likely localization technique to improve clinical MRS. We also recommend the use of PRESS with VSS pulses if only this option is available, but note this option lacks the partially suppressed $J$-coupling evolution and longer apparent $T_2$ relaxation when compared to semi-LASER, and can inadvertently excite or refocus large signals (e.g. scalp lipids) that are difficult to suppress fully.

In general, we recommend the shortest achievable TE for SVS at any field-strength, assuming CSD remains within the acceptable range of 4% per ppm and appropriate analysis methods are used to model the increased amplitude of macromolecular and lipid signals. The advantages of using the shortest possible TE include: 1) improved SNR due to reduced $T_2$ relaxation; 2) more accurate concentration estimation due to a reduced dependence on assumed metabolite $T_2$ values and 3) improved SNR for $J$-coupled metabolite signals due to reduced de-phasing. Other TEs may be appropriate for improved detection of specific metabolites, such as lactate/alanine at TE 144 ms or TE 97ms for 2-hydroxyglutarate with PRESS (83). Longer echo times may also be preferred to improve water and lipid suppression, due to their shorter $T_2$ compared to metabolites, or if optimal long TE biomarkers are targeted, such as $t$Cho/$t$Cr in glial tumor grading (84). A SVS TR of 1.5 s at 1.5 T and 2.0 s at 3T is recommended to provide the maximum SNR per unit time, on average, for the main metabolite signals from $t$Cho, $t$Cr, $t$NAA and lactate. Further justification and discussion may be found in section C of the supporting information.

The recommended number of averages for typical voxel dimensions of 15 x 15 x 15 mm$^3$ and 20 x 20 x 20 mm$^3$ for 1.5 and 3T are given in Table 1. A reduction in voxel dimensions results in a loss of SNR and therefore additional averages are required to attain suitable quality data. Note that signal-to-noise scales with the square-root of the number of averages. Accordingly, an unacceptably long time is required for the 15 mm sided cubic voxels to attain the same SNR as with the 20 mm sided cubic voxels, and therefore the recommended averages represent a compromise that results in an SNR reduction of 40% for the smaller voxel size. However, since shimming is generally improved for smaller voxel sizes, some signal loss is mitigated by narrower linewidths.

In addition to water suppressed data, which are required for metabolite level/concentration estimation, a matched unsuppressed water reference acquisition (acquired with the transmit frequency set to the water resonance) is recommended as part of all SVS clinical protocols. Since the water resonance has high SNR compared to metabolites, we recommend acquiring a single average for the water suppressed data. Where possible, a minimum period of 9 seconds without RF excitation should be ensured before acquisition to essentially eliminate $T_1$ weighting.
for water in normal and pathological tissue. A 9 second delay also ensures the T1-saturation signal loss is less than 10% in cystic regions or CSF. Dummy scans should be used to achieve a steady state in cases where the recommended relaxation period cannot be guaranteed. The water signal may be used to: correct for eddy current line-shape distortions (36); estimate B₀ field homogeneity and frequency offset; evaluate water suppression quality and provide metabolite concentration scaling information for use with short TE acquisitions (41).

**MRSI acquisition**

MRSI acquisitions excite a much larger volume of tissue compared to SVS, and therefore a more stringent maximum CSD of 2% per ppm (across the selected volume) is recommended. Higher CSD levels result in a significant loss of metabolite information around the edges of the MRSI excitation region, making it challenging to obtain the required spatial coverage in the cortex, for example, whilst also avoiding unwanted excitation of scalp lipids. It is recommended that edge voxels are excluded from data analysis. We recommend the use of semi-LASER over PRESS for 3 T MRSI pre-localization, due to its reduced CSD level and additional reasons stated in the SVS acquisition section. Where available, the use of high bandwidth spatially selective saturation bands (85,86) are also recommended to improve conformance between the excitation volume and the region of interest by suppressing scalp lipid regions that are unavoidably exited by the pre-localization scheme. Standard OVS is also recommended for suppressing signals from brain areas with significant B₀ inhomogeneity (e.g., frontal sinuses) to reduce PSF related distortion spread. A narrower point spread function, resulting from acquisition with higher in-plane resolution, is also recognized as an effective strategy for reducing scalp lipid contamination - provided the extra scan time and reduced SNR can be tolerated.

For 2D phase-encoded MRSI we recommend the use of elliptical sampling in k-space and TR’s of 1.5 s at 1.5 T and 2.0 s at 3 T (see supporting information section C). The phase encoding field-of-view should fully encompass the pre-localization volume to avoid aliasing, and the use of a 16x16 imaging matrix with 10 mm in-plane resolution and slice thickness of 15 mm is recommended as a default protocol. Note that the recommended spatial coverage and resolution parameters are intended as a starting point only, and may need to be adjusted to match the disease location, extent and clinical question. For instance, in the case of inadequate pre-localization, the phase encoding field-of-view should be increased to contain regions of spurious signals to avoid aliasing. As with SVS, short TE’s (~30 ms) are recommended for MRSI - due to the greater level of metabolite information, higher SNR and reduced T₂ bias in concentration estimates. However, in areas of greater B₀ inhomogeneity, where water and lipid suppression are less effective, longer TE’s may be necessary to reduce water and lipid signals relative to metabolites. An unsuppressed MRSI water scan should be acquired to aid line-shape correction, phasing, chemical shift referencing and quantitation. Since an additional MRSI water acquisition doubles the effective scan time for MRSI at the same spatial resolution (unlike SVS), a reduced resolution acquisition or parallel imaging (59) approach may be used to mitigate the associated time penalty for acquiring the unsuppressed water MRSI.

Whilst 2D phase-encoded MRSI with pre-localization is currently the most commonly used MRSI technique in the clinical environment, there is strong interest in the development of robust
whole-brain MRSI within a clinically feasible acquisition time. In addition to the obvious advantages for studying diseases that are known to impact multiple brain areas, such as neurodegeneration (87), the ability to plan a MRSI acquisition as easily as standard MRI (i.e., without having to preselect a region of interest) will improve data consistency and acceptance in clinical centers. The use of pre-localization also significantly restricts spatial coverage, in particular making it very difficult to study cortical regions near to the surface of the brain. Using conventional phase encoding, 3D MRSI is only practical for limited brain regions when there is the option of highly efficient outer volume suppression. For instance, the use of VSS pulses (70) or a 3D 8x8x8 MRSI matrix of 15 mm isotropic voxels provides good quality data in 12.8 min for TR 1500 ms and a full k-space acquisition. For whole brain examination, 3D MRSI is prohibitively time consuming, therefore a number of fast acquisition methods have been developed by the research community.

The most commonly used whole brain MRSI acquisition uses the Proton Echo Planar Spectroscopic Imaging (PEPSI) technique (88), where an oscillating readout gradient generates an echo-train, encoding a full plane of k-space for each excitation. The chemical shift and one spatial dimension are simultaneously encoded during the gradient echo readout, and phase encoding is used for the two remaining spatial dimensions. One of the most mature implementations of PEPSI uses a high-resolution acquisition scheme (5.6 mm x 5.6 mm x 10.0 mm) combined with lipid inversion nulling and post-acquisition k-space extrapolation (89) to significantly reduce scalp lipid contamination. In addition, an interleaved water acquisition is used to improve reconstruction in the presence of B₀ inhomogeneity and drift, and to provide a reference signal for metabolite concentration scaling (90). Unlike many other MRSI approaches, this sequence has been implemented and tested on instruments from three manufacturers with encouraging consistency across sites (91). The main advantages of this sequence are greater coverage of the cerebrum and simpler planning. This sequence has been reported to sample on average approximately 70% of the brain volume (92), although with instrumentation dependent differences. Recent 3T studies using PEPSI have been able to achieve an acquisition time of 18 min with a TE of 20 ms, which may be acceptable for clinical trials and some specialized clinical assessments (93).

An alternative approach for whole brain MRSI is to combine medium resolution PEPSI encoding (10 mm isotropic) with OVERPRESS localization and automatically prescribed VSS bands (94). This sequence is relatively fast to acquire (13 min) and does not require lipid inversion nulling, which also suppresses potentially useful lipid signals originating from pathology. In addition, the automated prescription of this sequence is much more practical for clinical use when compared to manually prescribing PRESS volumes in combination with saturation pulses and shim-volumes. This approach has been tested on a clinical cohort of brain tumor patients with encouraging results (94).

Considerable progress towards robust whole-brain MRSI has been made in recent years and a wide range of techniques have been demonstrated to provide high acceleration (95,96). At the time of writing, we were unable to recommend one whole brain MRSI approach over another and identify a need for comparative studies to assess the relative performance of the various approaches.
**SVS and MRSI preprocessing and analysis**

Fully automated analysis methods, which perform phase correction, chemical shift calibration and quantitative analysis of metabolite signals, have been available for 25 years (26) and are strongly recommended for MRS analysis. A choice of commercially licensed (26), free (97) and open-source (29,68,98) software packages have been developed, predominantly by the research community, and are available for off-line analysis. The prevailing trend in the literature is to incorporate a quantum mechanically simulated set of metabolite spectra (basis set) into the fitting routine. Modeled lipid signals and modeled or measured macromolecular signals can be included in the basis set when appropriate (e.g., short TE protocols or investigations of pathologies with hypoxic/necrotic processes such as stroke and tumors). We recommend this approach for clinical and research studies over single peak modelling or spectral integration, since simulated basis sets conveniently incorporate a greater level of prior-knowledge (32,35) into the fitting process, which can be easily adapted to match the acquisition protocol.

Quality assessment must be an integral part of MRS analyses since distorted or poorly fit data leads directly to erroneous results. Metabolite SNR and linewidth estimates are recommended as objective measures of data quality. A number of spectral SNR definitions have been described for MRS. Here, we recommend the signal measure be defined as the height of the largest metabolite data point in the real part of the spectrum minus the fitted baseline at that point; and the noise measure as two times the standard deviation of the spectral data points in a region free from metabolite signals, residual water or other spectral distortions. Whilst other measures of spectral SNR are acceptable, reported values should be accompanied by a definition to avoid ambiguity - for instance the current version of the LCModel (26) analysis package defines the noise as two times the standard deviation of the fit residual. Linewidth estimates (FWHM) are also usually measured, in units of Hz or ppm, from the real part of the phased spectrum and are typically derived from the most prominent singlet metabolite or unsuppressed water resonance. Whilst universal guidelines for acceptable MRS data quality are challenging, due to the heterogeneous metabolite profiles of diseases and wide range of acquisition protocols, spectra with FWHM greater than 0.1 ppm should be regarded as being poor quality. A minimum metabolite SNR of 3 is recommended to confirm the presence of a particular singlet, however greater metabolite SNR values are required for accurate quantification or detection of coupled multiplets. Example spectra with different SNR and FWHM values are shown in Figure 8.

Whilst SNR and linewidth criteria are useful for basic quality assessment, a visual inspection of fitting results remains important since analysis errors and data artifacts may produce unrealistic values. Furthermore, these values do not reliably detect a number of quality issues such as unstable or unrealistic baseline estimates during fitting or out-of-volume lipid contamination. To aid visual assessment we recommend that the analysis output includes plots of the phased spectra with the fit and estimated baseline and residual for retrospective visual assessment of the data (example given in Figure 1 with the optional addition of individual basis signals). For SVS, a quick visual assessment of the data and fit quality can be easily performed by someone with a basic understanding of MRS and common artifacts (37) - and this approach is suitable for the clinical environment. However for MRSI, where hundreds of spectra are typically acquired in
a single scan, visual assessment is often impractical and there is great interest in developing automated quality assessment. One of the first approaches, based on the classification of independent spectral components, was found to provide 87% agreement with expert spectroscopists’ evaluations (67). More recently, methods based on random forest classification of spectral data (66) and features extracted from the time and frequency domain (99) have also been shown to provide high agreement with expert assessment. Whilst these techniques are an important step towards fully automated MRS QC, their applicability across different scanners, experimental protocols and clinical scenarios requires further investigation.

Prior to SVS fitting, eddy-current correction using the unsuppressed water reference is recommended to reduce metabolite signal line shape distortions (36). There is also an increasing trend towards retrospective frequency and phase correction of each average and elimination of motion-corrupted FIDs prior to signal averaging (100,101). Whilst these methods have been successfully applied to scans with high metabolite signal (e.g., normal appearing brain), their suitability for low SNR clinical spectra (e.g., brain tumors) has not been fully investigated and therefore should be used with caution. We also note the promising approach of using “metabolite-cycled” MRS for retrospective correction using the water signal, allowing low SNR spectra to be accurately corrected, whilst maintaining metabolite spectral quality (102).

The analysis pipeline for MRSI includes preprocessing steps to aid interpretation. K-space filtering with a suitable window function, such as Hamming, is recommended for reducing distortions from Gibbs ringing of high intensity signals such as scalp lipids. Fourier interpolation, by k-space zero filling, to twice the acquired dimensions is also recommended to aid display and interpretation of metabolite maps. Optimized reconstruction methods, such as the Papoulis-Gerchberg algorithm, are also recommended to reduce lipid artifacts (89). More recently, a variety of novel reconstruction methods were shown to suppress MRSI lipid signals (103–106) with promising initial results, but validation on clinical data are required before recommending them for general use.

**Shimming and water suppression**

The current generation of commercially available shimming and water suppression algorithms do not provide the reliability and robustness needed for performing MRS/MRSI at all locations in the brain, in addition to being quite time-consuming — these drawbacks are important impediments to the wider clinical use of MRS. Therefore, we encourage the research community and MR vendors to optimize the performance of shimming, water suppression and power calibration steps in both favorable and challenging brain regions.

Both projection (FASTMAP and variants) and volumetric mapping acquisition methods are widely available, and can be effective for $B_0$ shimming, with projection methods generally taking less time. A recent study demonstrated that FASTMAP linewidths were 40% better than a vendor implementation of volumetric mapping for an 8 mL SVS VOI in the posterior cingulate cortex (78). We note that the vendor implementation of volumetric mapping has since been updated and improved, highlighting the challenges of comparing shimming methodology: seemingly minor implementation differences often have a significant impact on the overall performance of a particular strategy.
Whilst volumetric mapping methods are thought to be better suited for large VOIs typical to MRSI, no systematic studies have been published by the time of this writing that test this assertion. In addition to the $B_0$ mapping strategy, the shimming optimization method is crucial for automated and reliable shimming and a number of algorithms have been published and recently compared (107). Improved performance was found for methods incorporating known constraints on the maximum available shim strengths and mitigation for instability, and therefore these types of methods are recommended over simpler unconstrained algorithms.

Adequate hardware is essential for good $B_0$ homogeneity and second-order shim coils are recommended for MRS of acceptable quality, particularly at 3 T. In addition, adequately powered shim amplifiers are important to ensure good quality data in anatomic regions such as the hippocampus and frontal brain regions where there are large susceptibility gradients.

**Data formats and interoperability**

MRS results and data may be exported from the scanner console in the following formats: 1) DICOM MR spectroscopy; 2) DICOM secondary capture and 3) data points stored in a proprietary format. DICOM secondary captures (or “screenshots”) typically store an image of the spectrum and fit, alongside the voxel location and quantitative measures, such as metabolite ratios. For MRSI, metabolite maps are also generally displayed together with relevant MRI scans. Whilst secondary captures are essential for rapid clinical interpretation, they do not allow re-analysis or interactive inspection of the data – which is particularly important for MRSI. For example, following an initial analysis, it may become clear that a brain region has an abnormal metabolite profile consistent with a brain tumor, and therefore a re-analysis is required using a cancer-specific metabolite ratio map with a basis set containing additional metabolites or it may be advisable to re-inspect spectral quality in crucial areas.

The storage of the acquired complex MRS data points, either in the time or frequency domain, is essential to ensure the extraction of maximal information. Historically, each vendor had one or more proprietary formats for exporting raw data files – primarily for off-line analysis with third-party tools such as LCModel (26). However, proprietary formats have the following disadvantages: 1) they require extra time and computing resources to generate and securely archive, particularly when a hospital contains multiple scanners from different vendors. 2) Patient details may not be stored in a way that allows the reliable identification of files, creating problems for comparing scans obtained from multiple sites/scanners and effective de-identification – often a requirement for clinical trials. 3) Additional burden is placed on researchers and third-party software developers to support the file formats, which often change with software updates.

To address these issues, the DICOM standards committee introduced supplement 49 “Enhanced MR Image Storage SOP Class” in 2002, which included the “MR Spectroscopy Information Object Definition” suitable for the storage and transfer of MRS data. This allows MRS data points, and associated localization volume and acquisition information, to be archived to PACS alongside the MRI, via a local network, using the same protocols and infrastructure in place at the vast majority of radiology departments. Therefore, we recommend the use of the
DICOM standard for storage and network transmission of the MRS data points and analysis results from the scanner and strongly encourage MR and PACS vendors to implement the standard. For SVS, the storage of individual averages is recommended to aid retrospective correction of frequency and phase instabilities and the identification/removal of motion corrupted averages. Standard DICOM tags should be used to store all important sequence and localization information, and the use of private tags for this information is strongly discouraged - since private tags cannot be guaranteed to be free from protected health information and are therefore automatically removed as part of the de-identification procedures required for clinical trials.

Whilst the DICOM MRS format is essential for clinical purposes, we also recognize the importance of "raw" data formats. For instance, the data from each coil element needs to be stored separately for researchers to develop and compare reconstruction methods, and the DICOM format may not be suitable for these purposes. For these, reconstruction research orientated applications, formats such as the ISMRM Raw Data Format (ISMRMRD) are recommended for data export (108).

**Reporting**

The ideal for using MRS as a clinical biomarker would be the ability to report tissue metabolite levels fully quantitatively and independent of the scanner type or of the pathology under investigation. In practice, limitations of SNR, the need for reasonable acquisition times, that voxels contain mixed tissue types of variable relaxation times, and the effects of methodological variations due to differences in scanner software and hardware, restrict convergence to the ideal. In addition, there may be reluctance to move too far away from a specific institutional MRS protocol and acquire new data that is no longer comparable to historical studies. Hence, pragmatically, our recommendations balance limited protocol variability with best practices of acquiring high-quality MRS data, avoiding methodology that leads to extreme variation in MRS characteristics. Nevertheless, the appearance of the spectrum may still have characteristics that depend on the acquisition protocol, and the subsequent data processing may lead to further differences, all of which may impact visual or pattern recognition analyses. The most promising clinical applications for MRS have been described (1) but further work is needed for the development of standardized MRS biomarkers that enable rigorous and robust multi-institutional use. Hence it is essential that key details be provided in MRS publications to enable appropriate comparisons between different studies and for meta-analyses to better assess the efficacy of proposed MRS biomarkers.

At the 2016 ISMRM workshop "MR Spectroscopy: From current best practice to latest frontiers" attendees were asked to comment on a set of minimum and recommended requirements for reporting of MRS studies. Level 1 requirements were parameters considered minimum for proper and correct reporting of MRS studies by more than 80% of the attendees. Parameters that had between 40 - 80% support as relevant for inclusion in MRS reporting are ascribed as Level 2. A detailed consensus on MRS reporting is currently being developed for an NMR in Biomedicine Special Issue (expected publication in late 2019), hence only the outline of Level 1 minimum requirements is included here (supporting information section D) in relation to our consensus on $^1$H MRS of the brain.
Summary and Conclusions

Tables 1, 2 and 3 below summarize SVS, 2D MRSI and analysis/interpretation recommendations, respectively. To facilitate greater clinical utility of MRS, we encourage vendors to implement all recommendations as outlined here. We highlight the following three recommendations as being likely to have the greatest importance for improving routine clinical MRS and achieving reliable MRS results:

1. The implementation of a robust semi-LASER protocol to improve the localization of SVS and MRSI at 3 T.
2. The incorporation of simulated metabolite, lipid and macromolecular basis sets in spectral analyses for more robust extraction of the maximum amount of metabolic information available.
3. The use of optimized algorithms to perform time-efficient, robust and high-quality automated shimming (107).

These three recommendations are all software based and they can therefore be implemented on almost all existing clinical scanners, significantly enhancing their MRS capabilities.

Whilst we have intentionally restricted the scope of our recommendations to the most common clinical field strengths of 1.5 and 3 T, the same recommendations are also relevant to ultra-high field (7 T and above), where reducing CSD and implementing robust shimming present significantly greater challenges (109). Furthermore, a greater number of metabolites can be detected reliably at ultra-high field, due to wider chemical shift dispersion; and the use of comprehensive simulated metabolite basis sets is required to capture the full metabolite profile (110).

Consensus on appropriate experimental methodology is an evolving process, and we emphasize that this paper should not represent the final word on the topic. Our intention is to provide an assessment of the current state-of-the-art and recommend improvements to MRS methodology and standardization, with a strong focus on clinical applications. However, variety is essential for fruitful developments of new and alternative methods, yielding the clinical workhorses of the future. An initiative is currently underway to produce a special issue that will expand on this paper with a greater focus on ultra-high field MRS and more novel methods.

In conclusion, a large body of research demonstrates that 1) robust high-quality MRS data may be acquired with the hardware available on current clinical MR systems, and 2) many technical challenges of performing clinically useful SVS and MRSI at 1.5T and 3T can be overcome with software improvements applied to current scanner hardware. In this consensus paper, a series of methodological recommendations have been made to provide a degree of standardization and equivalency of methodology across all scanner platforms and guidelines on the current best-practices for clinical MRS.

Acknowledgements

The following authors drafted specific sections contained within this paper: Robert Bartha, In-Young Choi, Cristina Cudalbu, Anke Henning, Hoby P. Hetherington, Franklyn A. Howe, Kejal
Kantarci, Dennis W. J. Klomp, Roland Kreis, Alexander P. Lin, Malgorzata Marjanska, Andrew A. Maudsley, Paul G. Mullins, Sarah J. Nelson, Gülün Öz, Julie W. Pan, Harish Poptani, Stefan Posse, Eva-Maria Ratai and Martin Wilson. Figures 4 and 5 were provided by Yan Li (University of California San Francisco).

Martin Wilson, Franklyn A. Howe and Gülün Öz played a primary role in editing the contributed text and drafting the final version of the manuscript. Middle author names are listed in alphabetical order.

References


<table>
<thead>
<tr>
<th>Aspect</th>
<th>Consensus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Localization method</td>
<td>Chemical shift displacement less than 4% per ppm. 3 T: semi-LASER with OVS (preferred) or OVERPRESS with VSS. 1.5 T: semi-LASER with OVS (preferred) or PRESS.</td>
</tr>
<tr>
<td>Sequence timings</td>
<td>TE as short as possible (typically 30 ms). Longer TEs may be preferred for lactate (144 or 288ms), at TEs optimized for specific metabolites such as 2HG detection, as well as for enhanced lipid suppression. TR 1.5 s at 1.5 T, 2.0 s at 3 T.</td>
</tr>
<tr>
<td>Typical sequence parameters</td>
<td>128 averages collected from a 15 x 15 x 15 mm³ VOI at 3T. 64 averages from a 20 x 20 x 20 mm³ VOI at 3T. 256 averages collected from a 15 x 15 x 15 mm³ VOI at 1.5T. 128 averages from a 20 x 20 x 20 mm³ VOI at 1.5T. Spectral sampling of 1024 complex data points from 2000 Hz spectral width at 1.5T or 3T.</td>
</tr>
<tr>
<td>Reference acquisition</td>
<td>Recommended in all cases. Collect with the same sequence parameters as the water-suppressed scan, but without water suppression and the transmitter frequency set to the water resonance. A single average should be collected with a pre-acquisition delay time of at least 9s to prevent T1-weighting.</td>
</tr>
<tr>
<td>Shimming hardware</td>
<td>2nd order shim coils with adequately powered amplifiers are recommended at 3T.</td>
</tr>
<tr>
<td>Shimming algorithm</td>
<td>Methods incorporating shim strength limits and instability counter-measures are preferred over unconstrained approaches.</td>
</tr>
</tbody>
</table>

Table 1. SVS acquisition consensus summary.
<table>
<thead>
<tr>
<th>Aspect</th>
<th>Consensus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Pre-localization method</td>
<td>Chemical shift displacement less than 2% per ppm. 3T: semi-LASER with OVS (preferred) or OVERPRESS with VSS. 1.5T: semi-LASER with OVS (preferred) or PRESS with VSS.</td>
</tr>
<tr>
<td>Sequence timings</td>
<td>TE as short as possible (typically 30 ms). Longer TEs may be preferred for lactate detection (144 or 288 ms) and enhanced lipid suppression. TR 1.5 s at 1.5T, 2.0 s at 3T.</td>
</tr>
<tr>
<td>Typical sequence parameters</td>
<td>16x16 matrix with 10 mm in-plane resolution and 15 mm slice thickness. 1 average per phase encoding step. Spectral sampling of 1024 complex data points at 2000 Hz spectral width at 1.5T or 3T.</td>
</tr>
<tr>
<td>k-space sampling and preprocessing</td>
<td>2D phase-encoded Cartesian sampling over an elliptical or circular k-space mask. K-space zero-filling (interpolation) to twice the acquired number of points. Hamming filter. Reduction of subcutaneous lipid contamination (e.g. Papoulis-Gerchberg algorithm).</td>
</tr>
<tr>
<td>Reference acquisition</td>
<td>Should be acquired where possible. Collect with the same sequence parameters as the water-suppressed scan, but without water suppression. Typically, to reduce scan time, a lower resolution scan is acceptable and interpolated post-acquisition to match the metabolite resolution</td>
</tr>
<tr>
<td>Shimming hardware</td>
<td>2nd order shim coils with adequately powered amplifiers are recommended at 1.5T and 3T.</td>
</tr>
<tr>
<td>Shimming algorithm</td>
<td>Methods incorporating shim strength limits and instability counter-measures are preferred over unconstrained approaches.</td>
</tr>
</tbody>
</table>

**Table 2.** 2D-MRSI acquisition and preprocessing consensus summary.
<table>
<thead>
<tr>
<th>Aspect</th>
<th>Consensus</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spectral preprocessing</td>
<td>Time-domain apodization (line-broadening) and zero-filling steps should not be applied before spectral fitting, although may aid visual interpretation. Water reference based eddy-current correction (36) before fitting is recommended where possible.</td>
</tr>
<tr>
<td>Analysis methods</td>
<td>Methods should be fully automated, performing phasing, chemical shift calibration and metabolite amplitude estimation without user intervention. The flexibility to be able to model typical baseline and linewidth variations is an essential requirement; which may be achieved using time or frequency domain approaches.</td>
</tr>
<tr>
<td>Basis set</td>
<td>Methods that incorporate prior knowledge via a basis set are recommended over spectral integration or simple fitting of independent single peaks. Metabolite basis sets simulated from known J-coupling and chemical shift values to match the acquisition protocol are recommended for analysis. Lipid basis signals should also be incorporated for tumor analysis and macromolecule signals for short TE (&lt;80 ms) analyses.</td>
</tr>
<tr>
<td>Quality assessment</td>
<td>Single peak metabolite and, where available, water linewidths should be measured at half height (FWHM), as part of an automated analysis pipeline. Metabolite or water linewidths less than 0.1 ppm are required for accurate analysis, and a metabolite SNR greater than 3 is the minimum criterion for determining the presence of a singlet. Visual assessment of spectral and fit quality is recommended, based on the combined display of the phased spectrum, fit, estimated baseline and fit residual.</td>
</tr>
</tbody>
</table>

*Table 3. Analysis and interpretation consensus summary.*
Figure 1. An adult low grade-glioma brain tumor spectrum acquired at 3T with PRESS SVS localization, 18mm sided cubic voxel, 128 averages, TE=32ms and TR=2s. Parametric fitting was performed with the TARQUIN algorithm (29) using a simulated basis-set of metabolite, lipid and macromolecule signals. Although a greater level of spectral detail is available when compared to 1.5T, particularly for strongly J-coupled metabolites such as Glu and ml, data quality is highly dependent on achieving good shimming. For this example, a metabolite FWHM of 0.03 ppm and SNR of 83 were achieved, where the FWHM was measured from the highest metabolite signal (tCho=GPC+PCh) following baseline subtraction. SNR was calculated as the ratio between the highest, baseline subtracted, metabolite signal intensity and two times the standard deviation of the noise level estimated from a spectral region free from metabolite signals.
Figure 2. Comparison of vendor supplied MRS implementation of PRESS vs. an advanced in-house protocol using semi-LASER (72) and improved water suppression, shimming and data processing on a 3T MRI system. Spectra (TE = 30ms, TR = 5s, 64 transients) were acquired from the same voxel in the cerebellar vermis of a healthy volunteer. A) Water was suppressed by ~150-fold by the CHESS sequence in the PRESS protocol and ~4500-fold by the VAPOR scheme (23) incorporated into the semi-LASER sequence. B) CSD for resonances with a chemical shift difference of 3ppm is 36-39% with the PRESS sequence vs. 6% with the semi-LASER sequence in the two dimensions shown. C) Narrower linewidths (shown for the total creatine signal) are obtained with FASTMAP shimming (12) in conjunction with single shot frequency & phase correction in the semi-LASER protocol vs. product shimming and signal averaging in the PRESS protocol in the absence of motion during scanning. D) Small amount of motion (few degrees in z such that the cerebellar volume-of-interest was still acceptable at the extremes of motion) further degrades linewidths and generates unwanted coherences that are not removed with phase cycling in the PRESS protocol (highlighted by the circle), while spectral quality is unchanged, with an acceptable water residual in the semi-LASER protocol thanks to artifact-free single shots and frequency & phase correction of individual shots.
Figure 3. Effects of shimming on MR spectra obtained from the frontal lobes at 3T using PRESS localization (TR=1.5 s, TE=30 ms). A and B display a 2 x 2 x 2 cm³ voxel in the prefrontal cortex. Spectra illustrate the importance of shim quality for resolving metabolite resonances, with the following linewidths measured from the real part of the phased water resonance: C 23 Hz; D 14 Hz and E 10 Hz.
Figure 4. 3D MRSI (bottom left: TE=144 ms, TR=1.25s; bottom right: TE=35 ms, TR=1.3s) acquired at 3T from a patient with glioblastoma. Both scans had a nominal voxel resolution of 1 cm³, and 4Hz spectral apodization was applied. The PRESS excitation volume (yellow rectangle) and 8 VSS outer suppression bands (shaded purple) are shown in the top panel. Both the PRESS volume and VSS outer suppression bands were automatically prescribed using in-house software (94). Good-quality spectra from the red gridded region (top) of slices close to the center of the PRESS box are highlighted for both long and short echo times (bottom).
Figure 5. Chemical shift artifacts displayed in the 3D mode acquired from a healthy volunteer using a product PRESS sequence at 3T (TE/TR=144/1100ms, matrix=12x12x8, elliptical k-space sampling, nominal voxel resolution = 1cm³; total acquisition time = 11.04min). The thick white border indicates the PRESS excitation region (80x80x40mm) and voxels (highlighted in red) display an abnormal metabolite profile originating from CSD rather than pathology. Note the significantly reduced CSD in the anterior-posterior direction due to the application of the stronger gradient in anterior-posterior direction with the first slice selection pulse (90°).
Figure 6. The 1D point spread function (PSF) for phase-encoded MRSI with 16 points (part a) and with reduced spatial signal spread and resolution when applying a Hamming k-space filter (part b). The PSF of the more commonly used 16x16 circularly sampled 2D phase-encoding scheme for MRSI is illustrated in part c) and the corresponding Hamming filtered PSF in part d). A nominal resolution of 1 cm and 16 cm field-of-view was used for all plots. Due to the PSF and its filtering, the effective voxel size is considerably larger than the nominal voxel size.
Figure 7. Illustration of spectral quality maps generated by the MIDAS software package (68). a) $T_1$-weighted MRI corresponding to the selected MRSI slices. b) The raw tNAA signal intensity map. c) The spectral quality map, showing regions that passed the quality criteria for metabolite linewidth $\leq 0.1$ ppm in white. Regions that failed metabolite linewidth quality criteria, but passed water reference linewidth criteria are shown in mid grey, and regions that failed both linewidth criteria are shown in dark grey. d) The tNAA map with the identified poor-quality voxels set to zero. Note the signal dropout in much of the anterior cingulate cortex related to insufficient shimming close to air-tissue interfaces (sinuses, ear canals). Data were obtained using volumetric (3D) $^1$H MRSI with lipid inversion-nulling at 3 T, TE/TR/TI=17.6/1550/198 ms, 50x50x18 k-space points over 280x280x180 mm$^3$. 

Figure 8. Simulated “normal-brain” spectra for a typical PRESS acquisition at 3T; TE=30ms, 1024 data points and 2000Hz sampling frequency. SNR and FWHM values are given for the largest singlet resonance (tNAA).