NOTE

Effects of different macromolecular models on reproducibility of FID-MRSI at 7T

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Purpose: A properly characterized macromolecular (MM) contribution is essential for accurate metabolite quantification in FID-MRSI. MM information can be included into the fitting model as a single component or parameterized and included over several individual MM resonances, which adds flexibility when pathologic changes are present but is prone to potential overfitting. This study investigates the effects of different MM models on MRSI reproducibility.

Methods: Clinically feasible, high-resolution FID-MRSI data were collected in ~5 min at 7 Tesla from 10 healthy volunteers and quantified via LCModel (version 6.3) with 3 basis sets, each with a different approach for how the MM signal was handled: averaged measured whole spectrum (full MM), 9 parameterized components (param MM) with soft constraints to avoid overparameterization, or without any MM information included in the fitting prior knowledge. The test–retest reproducibility of MRSI scans was assessed voxel-wise using metabolite coefficients of variation and intraclass correlation coefficients and compared between the basis sets. Correlations of concentration estimates were investigated for the param MM fitting model.

Results: The full MM model provided the most reproducible quantification of total NAA, total Cho, myo-inositol, and glutamate + glutamine ratios to total Cr (coefficients of variations ≤ 8%, intraclass correlation coefficients ≥ 0.76). Using the param MM model resulted in slightly lower reproducibility (up to +3% higher coefficients of variations, up to −0.1 decreased intraclass correlation coefficients). The quantification of the parameterized macromolecules did not affect quantification of the overlapping metabolites.

Conclusion: Clinically feasible FID-MRSI with an experimentally acquired MM spectrum included in prior knowledge provides highly reproducible quantification...
1 | INTRODUCTION

MRSI combines the anatomical and biochemical information of an examined tissue and has been extensively used to detect in vivo metabolite levels in various brain disorders.\(^1\) MRSI benefits especially from ultrahigh field strength (i.e., ≥7 Tesla [T]) as the spectral resolution improves significantly.\(^2\) However, at higher field strengths the T\(_2\) relaxation times are shorter, leading to increased T\(_2\)-related SNR loss for MRSI with long TEs. To maximize the number of detectable signals and overcome some challenges that arise from the ultrahigh field (e.g., chemical shift displacement error), FID-based MRSI with no echo time but with a negligible acquisition delay was proposed and has received increasing attention in the research community.\(^3\)-\(^7\) While the detectability of neurometabolites, such as myo-inositol (mIns), glutamate (Glu), glutamine (Gln), or glutathione (GSH) improves, FID-MRSI is also highly sensitive to the broad background macromolecular (MM) signal. The presence of these strong MM resonances poses a challenge for accurate metabolite quantification.\(^8\),\(^9\) To use FID-MRSI as a reliable tool in a clinical context, it is mandatory to either decrease the MM contribution or include it in the fitting model appropriately while also providing sufficient flexibility for cases in which the MM profile is altered.

One approach is to measure the MM signal separately and subtract it from the metabolite spectrum.\(^10\),\(^11\) Although this is the most direct method with which to identify a subject-specific MM contribution, it is also motion-sensitive, prolongs the acquisition time (particularly in case of high-resolution MRSI), and reduces the SNR, which can eventually decrease the quantification accuracy.

Alternatively, prior knowledge of the MM contribution can be introduced into the fitting process. In case of LCModel, a widely used quantification software, the macromolecules can be approximated by spline functions\(^12\),\(^13\) or experimentally acquired and averaged over a group of healthy volunteers. Studies conducted at 3T have revealed that a measured MM spectrum is superior to a mathematical estimation in terms of MRSI reproducibility because the spline baseline cannot adequately characterize the complicated MM lineshapes.\(^14\) This is even more pronounced at higher field strengths (≥ 7 T) in which the apparent linewidth of the individual MM resonances approaches that of the J-coupled metabolites.\(^15\),\(^16\)

The most common strategy for measuring the MM spectrum in vivo is to use an inversion recovery sequence, thereby exploiting the short T\(_1\) relaxation times of macromolecules in contrast to those of most brain metabolites.\(^8\),\(^17\)-\(^19\) Another approach to capture the macromolecules is to back-extrapolate the metabolite signal and separate it from the MRSI data.\(^20\) Afterward, the measured MM spectrum either can be directly included in the basis set as a single component or parameterized and included as several individual MM components, which adds flexibility when the MM content and composition change, for example, when pathologic changes are present.\(^21\)-\(^23\) However, the inclusion of several MM components increases the degrees of freedom of LCModel analysis, which presumably could lead to inaccurate metabolite estimations and negatively influence the reliability of the quantification. This can be avoided by the application of soft constraints. Both approaches, using a whole MM spectrum\(^17\) or several parameterized MM components,\(^24\) were evaluated for fairly high SNR (thus, long scan times). Although the precision of metabolite quantification was investigated, the effects on reproducibility were not assessed. Yet, the knowledge about variation in repeated measurements is a critical prerequisite for future application of FID-MRSI in (longitudinal) clinical studies.

The aim of the present study was therefore to determine the effect of using different MM models on test–retest reproducibility for data acquired via clinically feasible ~5 min FID-MRSI protocols at 7T.

2 | METHODS

2.1 | Subjects

The study was approved by the local ethics board. Ten healthy volunteers (8 females, 7 males; mean age, 28 ± 4 years) were recruited. Informed, written consent was obtained from all subjects prior to the examination. Each subject was measured twice (i.e., test and retest measurement), preferably on the same day. All subjects were removed from the scanner between the sessions. In 4 subjects, the retest measurements were performed on a different day (up to 7 days) due to restricted scanner availability.
2.2 | Data acquisition

In each subject, test and retest MRI/MRSI scans were performed on a 7T Magnetom MR scanner (Siemens Healthcare, Erlangen, Germany) using a 32-channel receive array coil combined with a transmit volume coil (Nova Medical, Wilmington, MA). In each session 3D T$_1$-weighted MP2RAGE images were acquired to guide the positioning of MRSI slices and to derive the brain tissue type and structural maps. The spectroscopic data were obtained in 2 single-slice FID-MRSI acquisitions in order to test the performance of the sequence over larger brain volume. The first slice was positioned in the transverse plane above the corpus callosum, and the second slice was positioned 12 mm above the first slice. Consistent positioning between test and retest measurement sessions was achieved by using an automatic alignment sequence, reloading the spectroscopic sequence from the test measurement, and confirmed by visual inspection of the grid position. The parameters of the FID-MRSI sequence were as follows: acquisition delay/TR of 1.3 ms/600 ms, FOV of 200 × 200 mm$^2$, matrix size of 64 × 64, nominal voxel volume of 3.4 × 3.4 × 8 mm$^3$, flip angle of 45°, 1024 complex spectral data points, acquisition bandwidth of 6000 Hz, water suppression enhanced through T$_1$ effects (WET), 6-fold 2D-Controlled Aliasing in Parallel Imaging Results in Higher Acceleration (CAIPIRINHA), parallel imaging acceleration, and acquisition time of 5:11 min.

2.3 | Spectroscopic data processing

Brain masks were extracted from T$_1$-weighted images using a brain extraction tool. MRSI data within the brain masks were processed automatically using a script written in MatLab (version R2013a; MathWorks, Inc., Natick, MA) and Bash (version 4.2.25, Free Software Foundation, Boston, MA). The processing included a multichannel spectroscopic data combined by matching image calibration data (MUSICAL) coil combination of the raw data, parallel imaging reconstruction, Hamming filtering, removal of lipid signal via L2-regularization, and fitting of the individual spectra with LCModel (version 6.3; LCModel Inc., Oakville, ON, Canada). For this purpose, 3 different basis sets were used, each consisting of 15 metabolite resonances simulated in NMR Scope (jMRUI 5.0), as well as the following:

1. full MM: A single in vivo MM spectrum obtained in previous study by nulling of brain metabolite signals via double inversion recovery (2 inversion 40 ms Wurst pulses, acquisition delay/TR of 1.3 ms/879 ms, T$_1$ of 570 ms, T$_2$ of 21 ms, flip angle of 55°, matrix size of 32 × 32, nominal voxel volume of 5.6 × 5.6 × 12 mm$^3$, 2048 complex spectral data points, acquisition bandwidth of 6000 Hz, water suppression enhanced through T$_1$ effects [WET]), removing the residual metabolite signals using advanced method for accurate, robust, and efficient spectral fitting (AMARES), as described by Craveiro et al., and averaging over different brain regions and healthy volunteers.
2. param MM: Nine individual MM components extracted from measured in vivo metabolite-nulled spectra, with applied soft constraints (via LCModel parameter CHRATO) to avoid overparameterization of the fitting model (detailed description of the parameterization can be found in Považan et al.).
3. no MM: With no macromolecular information included.

Spectral analysis was performed in a frequency range of 0.2 to 4.2 ppm when using the full MM and the param MM basis set. In case of no MM basis set, the frequency range was reduced to 1.8 to 4.2 ppm to avoid the lipid region and MM peaks below. The LCModel parameter DNTMN, controlling the stiffness of the spline baseline, was set to the default value of 0.15 in all cases. The LCModel control files for the 3 quantification approaches are provided in Supporting Information Text S1.

2.4 | Data evaluation

The spectral and fitting quality of the data was assessed via 3 parameters: Cramér-Rao lower bounds (CRLB) of the metabolites reported by LCModel and SNR and FWHM of the fitted NAA peak. Spectra with CRLB$_{NAA} > 20\%$ or FWHM$_{NAA} > 20$ Hz were excluded from further analysis and display (i.e., <2% of all spectra). The excluded spectra were generally located at the periphery of the brain. Metabolite maps were derived from the LCModel quantification results and displayed by MINC software (version 2.0, McConnell Brain Imaging, Montreal, QC, Canada).

For the evaluation, 5 brain regions were defined: frontal white matter (WM), frontal gray matter (GM), parietal WM, parietal GM, and subcortical WM. As a first step, tissue-type segmentation (GM, WM, and CSF) and structural registration (frontal lobe, parietal lobe, and subcortical WM) were performed on T$_1$-weighted images of both test and retest measurements using automated segmentation tool (FAST) and linear image registration tool (FLIRT) of the FSL package. To match the point-spread function of MRSI and T$_1$-weighted MRI, segmented high-resolution images were Fourier-transformed to k-space, matched to the spatial frequency characteristics of the MRSI data, and converted back to image-space. Only voxels with a minimum of 80% of WM or GM content were used for analysis. The segmented images in MRSI resolution were then combined to create binary masks of the 5 aforementioned regions. Finally, the intersections of binary masks from the test and retest of each subject were used as the final masks.

Statistical analysis and calculations were performed in MatLab (MathWorks) using voxel-wise analysis. The test–retest reproducibility of metabolite ratio levels
was established by an intrasubject coefficient of variation (CV) calculated for each eligible voxel as the SD of the 2 measurements divided by their mean. CVs were grouped to 5 regions according to the predefined binary masks. As a measure of method reliability, the intraclass correlation coefficients (ICC) using an absolute-agreement, 2-way, mixed-effects model were calculated between test and retest. For the param MM basis set, a correlation diagram was derived to investigate whether the concentrations of metabolites and underlying individual MM components were independent of each other. To test for differences in CV, ICC, metabolite ratios, and quantification precision between the 3 basis sets, a nonparametric Friedman test, tailored for comparison of multiple related samples, was used. Subsequently, post hoc analysis by Wilcoxon signed-ranks tests was performed for pairwise comparisons. The differences between brain regions were compared using Kruskal-Wallis tests followed by Mann-Whitney post hoc analysis. Bonferroni correction for multiple testing was applied, and a $P < 0.05$ was considered significant.

3 | RESULTS

3.1 | Spectral quality

Altogether, 7532 pairs of test–retest spectra collected from 10 subjects were evaluated. Each spectrum was quantified in LCMModel by using 3 different basis sets. Of the first MRSI acquisition, 630, 1085, 679, 779, or 1957 spectra were assigned to the frontal GM, frontal WM, parietal GM, parietal WM, or subcortical WM region, respectively. In case of the second MRSI acquisition, 656, 1151, 337, or 258 spectra were assigned to the frontal GM, frontal WM, parietal GM, or parietal WM region, respectively. The subcortical WM of the second MRSI scan was not evaluated because of a small number of assigned spectra. Region-specific spectral quality parameters are displayed in Figure 1A. The overall spectral quality was high, with the lowest median SNR > 20 in the subcortical WM and the highest median FWHM <14 Hz in the frontal GM. The use of different fitting models had no impact on the metrics.

FIGURE 1 (A) Spectral quality (SNR, linewidth) and quantification precision (Cramér-Rao lower bounds) parameters. The circles and error bars show the 25th percentile, median, and the 75th percentile of the total number of analyzed voxels. (B) Representative spectra obtained by FID-MRSI and quantified in LCMModel by 3 different MM prior knowledge models: no MM, with no MM information; full MM, with a single measured MM spectrum; and param MM, with 9 individual MM components included. The fitted baselines have been subtracted from the metabolite and MM spectra. These subtracted baselines and the residua are displayed separately below each LCMModel fit. MM, macromolecular
3.2 | Quantification precision

Sample spectra fitted by different basis sets are displayed in Figure 1B. The CRLBs were consistently low among the 2 MRSI slice positions and brain regions (except for ~5% higher CRLB_Glx in the subcortical WM), as well as between test–retest measurements (median CRLB_NAA ≤ 5%, median CRLB_Cho ≤ 6%, median CRLB_Cr ≤ 7%, median CRLB_mIns ≤ 8%, median CRLB_Glx ≤ 14%). The regional differences in CRLB_Glx (CRLB_GM < CRLB_WM < CRLB_subcorticalWM) were associated with SNR loss (SNR_GM > SNR_WM > SNR_subcorticalWM). We observed small alterations in CRLBs among the basis sets (param MM resulted in a +1% increase in CRLBs) (Figure 1A).

Sample metabolite maps are displayed in Figure 2. The metabolite ratio levels are summarized in Supporting Information Table S1. The levels of tNAA/tCr in GM (by −13%) and in WM (by −8%), as well as mIns/tCr in WM (by −6%) and Glx/tCr in WM (by −5%), were significantly decreased when using full MM in comparison to no MM scheme (all P < 0.001). The differences between full MM and param MM were insignificant, except of ~3% difference in GM of Glx/tCr (P = 0.01).

The correlation analysis revealed that the signal amplitudes of several MM resonances were highly interdependent; however, there were no associations or only very weak correlations between individual parameterized macromolecules and overlapping metabolites (Figure 3).

3.3 | Reproducibility

Figure 4 and Supporting Information Figure S1 summarize the measures of reproducibility for the 4 main metabolite ratios of different fitting approaches. The obtained CVs were relatively consistent and low between the no MM, full MM, and param MM basis sets. The full MM analysis was the most reproducible, with mean CV_tNAA/tCr = 7.0%, mean CV_tCho/tCr = 5.9%, mean CV_mIns/tCr = 7.0%, and mean CV_Glx/tCr = 8.1%. Using the param MM scheme yielded up to a +3% increase in CVs compared to full MM, with the largest differences in CV_Glx/tCr (P < 0.01). The reproducibility of metabolites quantified with lower precision (i.e., higher CRLBs) was similarly reduced by maximum 3% when using the param MM basis set in comparison to the full MM (Supporting Information Figure S2). The mean CVs were <12% for Glu/tCr, <15% for GSH/tCr, and <20% for Glu/tCr and Tau/tCr among all brain regions for full MM basis set. The reproducibility of Gln/tCr was significantly improved when using full MM basis set in comparison to no MM (P < 0.001). Generally, the CVs were comparably low between the brain regions in both MRSI acquisitions, and only CVs_Glx/tCr were slightly higher in the subcortical WM (by ~3%, P < 0.001) and in the WM regions (by ~2%, P < 0.01) than in the GM regions, likely due to the lower signal intensity of Glx in the WM. Similarly, CV_mIns/tCr was increased by ~2% in the subcortical WM (P < 0.01) compared to the other 4 regions.

**FIGURE 2** Metabolite maps of one volunteer obtained by FID-MRSI with 64 × 64 resolution. The quantification of spectra was performed in LCModel using 3 basis sets with different macromolecular prior knowledge included: with no MM information, with a single in vivo MM spectrum (full MM), and with 9 parameterized MM components (param MM) with soft constraints.
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ICC values confirmed a very good agreement between the measurements. Generally, the highest ICC values were in the WM regions when using the full MM basis set (mean ± SD in WM regions, ICC<sup>tCho/tCr</sup> = 0.85 ± 0.07, mean ICC<sup>tNAA/tCr</sup> = 0.80 ± 0.09, ICC<sup>mlns/tCr</sup> = 0.85 ± 0.07, ICC<sup>Glx/Cr</sup> = 0.83 ± 0.08). Overall, the obtained ICCs indicate moderate (0.5 < ICC < 0.75) to good (0.75 < ICC < 0.9) reliability between the test and retest measurements in both MRSI acquisitions. The ICC values were comparably high between the basis sets (mean ± SD, ICC<sub>noMM</sub> = 0.76 ± 0.08, ICC<sub>fullMM</sub> = 0.79 ± 0.09, ICC<sub>paramMM</sub> = 0.74 ± 0.09). The incorporation of MM information into the fitting process either via a single

**Figure 3** Correlation coefficients derived from quantification results of the LCModel analysis of FID-MRSI using the basis set that included 9 individual parameterized MM components (param MM). Each element of the matrix indicates the absolute value of the correlation coefficient between the fitting results of the 2 corresponding metabolites.

**Figure 4** CV values for the voxel-based analysis of the metabolite ratios obtained by different MM prior knowledge included in the basis sets: no MM, with no MM information; full MM, with a single measured MM spectrum; and param MM, with 9 individual MM components with soft constraints. The box-and-whisker-plots represent the minimum, 25th percentile, median, 75th percentile, and maximum number of the analyzed voxels. CV, coefficient of variations.
spectrum or over 9 parameterized MM peaks had no negative effect on the reliability of MRSI; in fact, all ICC_{fullMM} ≥ ICC_{noMM} and only ICC_{paramMM} of Glx/tCr (of both MRSI acquisitions) in the frontal GM or the frontal WM were significantly decreased (P < 0.01) compared to ICC_{noMM} (Figure 5 and Supporting Information Figure S3).

4 | DISCUSSION

This study compares the effect of using 3 different macro-molecular models on the test–retest reproducibility of brain metabolite quantification for 7T FID-MRSI. The comparison was performed between basis sets with no MM information, with a single measured MM spectrum, and with 9 individual MM peaks included in the fitting prior knowledge.

Based on obtained CVs and ICCs, we provide evidence that FID-MRSI can map the spatial distribution of 4 neurotransmitter metabolite ratios with high precision, particularly when using the full MM model (CV ≤ 8%, ICC ≥ 0.76) in only ~5 min. The inclusion of individual MM components into the quantification and using adequate soft constraints resulted in almost the same reproducibility but no systematic absolute error compared to the quantification using full MM, and it also offered the added benefit of full flexibility in case of (un-)expected local changes in the MM profile. We found no strong correlation between individual MM components and overlapping metabolites that would indicate a severe bias due to unintentional overfitting. This indicates that the full MM model may be superior for healthy volunteer studies or those in which no major changes in the MM profile are expected, but the difference in reproducibility is small enough to make the param MM fitting model the preferred choice when the MM composition is expected to change, for example, in pathologies. In such cases, fitting MM resonances individually should help maintain the accurate quantification of metabolites and even macromolecules themselves at the expense of only slightly reduced reproducibility. This could provide additional diagnostic information.

We achieved very good spectral quality and quantification precision, which further support the validity of the FID-MRSI method. The small increase (~1%) in CRLB when param MM basis set was used is probably caused by increased number of degrees of freedom of the model. Several metabolite ratio levels were fairly decreased when incorporating MM prior knowledge into the fitting model. These differences were in good agreement with previously published results in which the tNAA, Glu, mlms, tCr, and tCho signals were decreased by approximately 25%, 15%, 13%, 5%, and 8% in the GM and by 16%, 12%, 12%, 7%, and 11% in the WM when using single measured MM spectrum in the basis set. Thus, the largest changes can be found in the GM of tNAA/tCr. The strong correlations we observed between some of the MM resonances can explain a possible physiologically meaningful relation between these resonances, or these dependencies could be partially attributable to the soft constraints applied. Moreover, MM resonances at 1.22, 1.43, and 1.70 ppm can be affected by the lipid signal removal.

The reproducibility of brain spectroscopic methods have been investigated extensively among different field strengths and acquisition techniques. Previous short TE 7T single-voxel MRS reproducibility studies have differed in their approaches to handle the MM contributions by using either the automatic MM calculation of LCModel, the group-averaged measured MM spectrum (here, full MM) as part of the basis set, the inversion-based suppression of the MM signal, or no MM correction at all. These studies were conducted with STEAM or semi-LASER using a 3 to 27 cm³ voxel volume and TEs ranging from 14 to 72 ms positioned in the anterior cingulate, posterior cingulate, prefrontal, or occipital cortex. The CVs for the same metabolites as reported in our work were below 5% when using the measured MM

![Figure 5](image-url)
spectrum and below 10% when using LCMcldel’s MM calculation or suppression of macromolecules.

To our best knowledge, studies concerning long TE MRS(I) reproducibility were conducted at lower field strengths only. According to Birch et al.\textsuperscript{14} and Inglese et al.,\textsuperscript{45} the use of intermediate/long TE outperformed short TE in terms of reproducibility, except of Glu (or Glx) and mIns, which had better reproducibility when using short(er) TE. The CVs obtained in their works at 3T using PRESS localization (with long or intermediate TE) were comparable to or worse than our CVs from FID-MRSI at 7T.

Birch et al. also compared in their work the influence of experimental and simulated MM models on the MRSI reproducibility at 3T\textsuperscript{14} and concluded that the use of experimental MM basis sets resulted in a slightly better performance than the use of simulated macromolecules. The CV < 6% for tNAA, tCr, and tCho (for TE = 80ms) and the CV < 16% for Glx and mIns (for TE = 35ms) were comparable to or worse than our results from 7T. In contrast, Schaller et al. showed that using a mathematical approximation model for the MM contribution was sufficient at 3T; however, the reproducibility of metabolite quantification through CVs was not assessed.\textsuperscript{12}

Another parameter, ICC, is a commonly used reliability index in test–retest analysis. ICC depends not just on the measurement errors of the method but also on the true heterogeneity in the population. Ideally, the variability in measurements is due to genuine differences between the subjects; then, ICC = 1. However, when ICC = 0, the observed variability is only a measurement error. The slightly lower ICC values of tNAA/tCr compared to those of tCho/tCr or mIns/tCr can be caused by imperfect lipid decontamination, which mostly affects the NAA peak. MRS reproducibility studies performed at 7T lack information about ICC and primarily concentrate on CV. Compared to the spin-echo MRSI sequence at 3T,\textsuperscript{33} we achieved better reliability in the WM (the mean ICC\textsubscript{WM} > 0.8 vs. ICC\textsubscript{WM} > 0.55) and comparable reliability in the GM (mean ICC\textsubscript{GM} > 0.7). Thus, our results show that neither extracranial lipids nor MM contributions significantly reduced reproducibility compared to other reports, although both are enhanced in FID-MRSI, especially when highly accelerated by parallel imaging. A further reduction of lipid artifacts via dedicated lipid removal hardware\textsuperscript{46} or spatial-spectral encoding\textsuperscript{47} is expected to further improve the reproducibility.

The limitations of our study include the absence of absolute metabolite concentrations, which would allow a more direct comparison of our results with previous works. Absolute quantification requires information about water density and relaxation times of the metabolites. Our sequence benefits from negligible T\textsubscript{2}-weighting; however, it is sensitive to incorrect assumptions about T\textsubscript{1} relaxation times due to the relatively short TR of 600 ms. Therefore, we decided to report metabolite levels with respect to tCr, which is a commonly used internal reference peak. Due to the lack of ground truth, only the precision and not the accuracy of the method could be assessed. The LCMcldel’s internal MM quantification was not included in our comparison because of the FID acquisition and the resulting phase problems. Fitting range was reduced for no MM analysis; otherwise, having unfitted peaks in residuum would cause wrong CRLB estimation. The reproducibility was evaluated in 2 axial MRSI slices at different levels, both avoiding the deep brain structures, which are usually strongly affected by spatial B\textsubscript{0} inhomogeneity. Nevertheless, we achieved very good reproducibility in the frontal lobe, which is similarly considered challenging for B\textsubscript{0} shimming. Due to restricted scanner availability, we could not maintain the same interval between the measurements for all the subjects. Some of the repeated measurements were performed on a different day, which could have resulted in biological differences between the datasets, but these should be negligible for young healthy volunteers.\textsuperscript{48}

5  |  CONCLUSION

FID-MRSI with in vivo measured macromolecular contribution included in the fitting prior knowledge provides highly reproducible quantification for common neurometabolites at 7T in only ~5 min. The use of the whole measured MM spectrum provided the highest reproducibility in young healthy volunteers. Parameterization of the MM spectrum yielded only slightly lower reproducibility compared to a single MM component; however, parameterization may be beneficial when the MM profile is expected to be altered due to pathological changes. This makes FID-MRSI a feasible clinical research tool to target brain biochemistry as well as for applications beyond investigations of the common brain metabolites.

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REFERENCES

1. Oz G, Alger JR, Barker PB, et al. Clinical proton MR spectroscopy in central nervous system disorders. Radiology. 2014;270:658–679.
2. Gruber S, Heckova E, Strasser B, et al. Mapping an extended neurochemical profile at 3 and 7 T using accelerated high-resolution
proton magnetic resonance spectroscopic imaging. Invest Radiol. 2017;52:631–639.
3. Bogner W, Gruber S, Trattnig S, Chmelik M. High-resolution mapping of human brain metabolites by free induction decay 1H MRSI at 7 T. *Magn Reson Biomed*. 2012;25:873–882.
4. Henning A, Fuchs A, Murdoch J, Boesiger P. Slice-selective FID acquisition, localized by outer volume suppression (FIDLOVS) for 1H MRSI of the human brain at 7 T with minimal signal loss. *Magn Reson Biomed*. 2009;22:683–696.
5. Hangel G, Strasser B, Považan M, et al. Ultra-high resolution brain metabolite mapping at 7 T by short-TR Hadamard-encoded FID-MRSI. *Neuroimage*. 2018;168:199–210.
6. Chadzynski GL, Bause J, Shajan G, Pohmann R, Scheffler K, Ehses P. Fast and efficient free induction decay MR spectroscopic imaging of the human brain at 9.4 Tesla. *Magn Reson Med*. 2017;78:1281–1295.
7. Nassirpour S, Chang P, Henning A. High and ultra-high resolution metabolite mapping of the human brain using 1H FID MRSI at 9.4T. *Neuroimage*. 2018;168:211–221.
8. Behar KL, Rothman DL, Spencer DD, Petroff OAC. Analysis of macromolecular resonances in 1H NMR spectra of human brain. *Magn Reson Med*. 1994;32:294–302.
9. Cudalbu C, Mlynarik V, Gruetter R. Handling Macromolecule signals in the quantification of the neurochemical profile. *J Alzheimers Dis*. 2012;31:S101–S115.
10. Penner J, Bartha R. Semi-LASER 1H MR spectroscopy at 7 Tesla in human brain: metabolite quantification incorporating subject-specific macromolecule removal. *Magn Reson Med*. 2015;74:4–12.
11. Kassem M, Bartha R. Quantitative proton short-echo-time LASER spectroscopy of normal human white matter and hippocampus at 4 Tesla incorporating macromolecule subtraction. *Magn Reson Med*. 2003;49:918–927.
12. Schaller B, Xin L, Cudalbu C, Gruetter R. Quantification of the neurochemical profile using simulated macromolecule resonances at 3 T. *Magn Reson Biomed*. 2013;26:593–599.
13. Provencher SW. Estimation of metabolite concentrations from localized in vivo proton NMR spectra. *Magn Reson Med*. 1993;30:672–679.
14. Birch R, Peet AC, Dehghani H, Wilson M. Influence of macromolecule baseline on 1H MR spectroscopic imaging reproducibility. *Magn Reson Med*. 2017;77:34–43.
15. Snoussi K, Gillen JS, Horska A, et al. Comparison of brain gray and white matter macromolecule resonances at 3 and 7 Tesla. *Magn Reson Med*. 2015;74:607–613.
16. Otazo R, Mueller B, Ugurbil K, Wald L, Posse S. Signal-to-noise ratio and spectral linewidth improvements between 1.5 and 7 Tesla in proton echo-planar spectroscopic imaging. *Magn Reson Med*. 2006;56:1200–1210.
17. Považan M, Hangel G, Strasser B, et al. Mapping of brain macromolecules and their use for spectral processing of 1H-MRSI data with an ultra-short acquisition delay at 7T. *Neuroimage*. 2015;121:126–135.
18. Giapitzakis I-A, Avdievich N, Henning A. Characterization of macromolecular baseline of human brain using metabolite cycled semi-LASER at 9.4T. *Magn Reson Med*. 2018;80:462–473.
19. Lopez-Kolkovsky AL, Mériaux S, Boumezbeur F. Metabolite and macromolecule T1 and T2 relaxation times in the rat brain in vivo at 17.2T. *Magn Reson Med*. 2016;75:503–514.
20. Lam F, Li Y, Clifford B, Liang Z-P. Macromolecule mapping of the brain using ultrashort-TE acquisition and reference-based metabolite removal. *Magn Reson Med*. 2018;79:2460–2469.
21. Mader I, Seeger U, Weisert R, et al. Proton MR spectroscopy with metabolite-nulling reveals elevated macromolecules in acute multiple sclerosis. *Brain*. 2001;124:953–961.
22. Graham GD, Hwang J-H, Rothman DL, Prichard JW. Spectroscopic assessment of alterations in macromolecule and small-molecule metabolites in human brain after stroke. *Stroke*. 2001;32:2797–2802.
23. Seeger U, Klose U, Mader I, Grodd W, Nägeli T. Parameterized evaluation of macromolecules and lipids in proton MR spectroscopy of brain diseases. *Magn Reson Med*. 2003;49:19–28.
24. Považan M, Strasser B, Hangel G, et al. Simultaneous mapping of metabolites and individual macromolecular components via ultra-short acquisition delay 1H MRSI in the brain at 7T. *Magn Reson Med*. 2018;79:1231–1240.
25. van der Kouwe AJW, Benner T, Fischl B, et al. On-line automatic slice positioning for brain MR imaging. *NeuroImage*. 2005;27:222–230.
26. Strasser B, Považan M, Hangel G, et al. (2 + 1D)-CAIPIRINHA accelerated MR spectroscopic imaging of the brain at 7T. *Magn Reson Med*. 2017;78:429–440.
27. Smith SM. Fast robust automated brain extraction. *Hum Brain Mapp*. 2002;17:143–155.
28. Strasser B, Chmelik M, Robinson SD, et al. Coil combination of multichannel MRSI data at 7 T: MUSICAL. *NMR Biomed*. 2013;26:1796–1805.
29. Bilgic B, Chatnuntawech I, Fan AP, et al. Fast image reconstruction with L2-regularization. *J Magn Reson Imaging*. 2014;40:181–191.
30. Vanhamme L, van den Boogaart A, van Huffel S. Improved method for accurate and efficient quantification of MRS data with use of prior knowledge. *J Magn Reson*. 1997;129:35–43.
31. Craveiro M, Clément-Schatlo V, Marino D, Gruetter R. Cudalbu C. In vivo brain macromolecule signals in healthy and glioblastoma mouse models: 1H magnetic resonance spectroscopy, post-processing and metabolite quantification at 14.1 T. *J Neurochem*. 2014;129:806–815.
32. Jenkinson M, Beckmann CF, Behrens T, Woolrich MW, Smith SM. FSL. *Neuroimage*. 2012;62:782–790.
33. Veenith TV, Mada M, Carter E, et al. Comparison of inter subject variability and reproducibility of whole brain proton spectroscopy. *PLoS ONE*. 2014;9:e115304.
34. Wijtenburg SA, Rowland LM, Edden R, Barker PB. Reproducibility of brain spectroscopy at 7T using conventional localization and spectral editing techniques. *J Magn Reson Imaging*. 2013;38:460–467.
35. Wijtenburg SA, Rowland LM, Oeltschinger G, Barker PB, Workman CI, Smith GS. Reproducibility of brain MRS in older healthy adults at 7T. *Magn Reson Biomed*. 2019;32:e0404.
36. Terpstra M, Cheong I, Lyu T, et al. Test-retest reproducibility of neurochemical profiles with short-echo, single-voxel MR spectroscopy at 3T and 7T. *Magn Reson Med*. 2016;76:1083–1091.
37. Hnilicová P, Považan M, Strasser B, et al. Spatial variability and reproducibility of GABA-edited MEGA-LASER 3D-MRSI in the brain at 3 T. *NMR Biomed*. 2016;29:1656–1665.
38. Hammern T, Stadlbauer A, Tomandl B, et al. Short TE single-voxel 1H-MR spectroscopy of hippocampal structures in healthy adults at 1.5 Tesla—how reproducible are the results? *NMR Biomed*. 2005;18:195–201.
39. Venkatraman TN, Hamer RM, Perkins DO, Song AW, Lieberman JA, Steen RG. Single-voxel 1H PRESS at 4.0 T: precision and variability of measurements in anterior cingulate and hippocampus. *NMR Biomed*. 2006;19:484–491.

40. Maudsley AA, Domenig C, Sheriff S. Reproducibility of serial whole-brain MR spectroscopic imaging. *NMR Biomed*. 2010;23:251–256.

41. Saleh MG, Near J, Alhamud A, Robertson F, van der Kouwe A, Meintjes EM. Reproducibility of macromolecule suppressed GABA measurement using motion and shim navigated MEGA-SPECIAL with LCModel, jMRUI and GANNET. *MAGMA*. 2016;29:863–874.

42. Zhang Y, Taub E, Salibi N, et al. Comparison of reproducibility of single voxel spectroscopy and whole-brain magnetic resonance spectroscopy imaging at 3T. *NMR Biomed*. 2018;31:e3898.

43. Geurts J, Barkhof F, Castelijns JA, Uitdehaag B, Polman CH, Pourwels P. Quantitative 1H-MRS of healthy human cortex, hippocampus, and thalamus: metabolite concentrations, quantification precision, and reproducibility. *J Magn Reson Imaging*. 2004;20:366–371.

44. Prinsen H, de Graaf RA, Mason GF, Pelletier D, Juchem C. Reproducibility measurement of glutathione, GABA, and glutamate: towards in vivo neurochemical profiling of multiple sclerosis with MR spectroscopy at 7T. *J Magn Reson Imaging*. 2017;45:187–198.

45. Inglese M, Spindler M, Babb JS, Sunenshine P, Law M, Gonen O. Field, coil, and echo-time influence on sensitivity and reproducibility of brain proton MR spectroscopy. *Am J Neuroradiol*. 2006;27:684–688.

46. Boer VO, van de Lindt T, Luijten PR, Klomp D. Lipid suppression for brain MRI and MRSI by means of a dedicated crusher coil. *Magn Reson Med*. 2015;73:2062–2068.

47. Hingerl L, Bogner W, Moser P, et al. Density-weighted concentric circle trajectories for high resolution brain magnetic resonance spectroscopic imaging at 7T. *Magn Reson Med*. 2018;79:2874–2885.

48. Maudsley AA, Govind V, Arheart KL. Associations of age, gender and body mass with 1H MR-observed brain metabolites and tissue distributions. *NMR Biomed*. 2012;25:580–593.

**SUPPORTING INFORMATION**

Additional supporting information may be found online in the Supporting Information section at the end of the article.

**FIGURE S1** Coefficient of variations (CV) of the second MRSI slice acquired at the upper level of the brain

**FIGURE S2** Coefficient of variations (CV) of the metabolites quantified with lower quantification precision

**FIGURE S3** ICC of the second MRSI slice acquired at the upper level of the brain

**TABLE S1** Metabolite ratios quantified by LCModel using three different basis sets

**TEXT S1** Control files for LCModel quantification

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