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NOTE

Gamma-aminobutyric acid edited echo-planar spectroscopic imaging (EPSI) with MEGA-sLASER at 7T

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Purpose: For rapid spatial mapping of gamma-aminobutyric acid (GABA) at the increased sensitivity and spectral separation for ultra-high magnetic field strength (7 tesla [T]) an accelerated edited magnetic resonance spectroscopic imaging technique was developed and optimized for the human brain at 7 T.

Methods: A MEGA-sLASER sequence was used for GABA editing and volume selection to maximize editing efficiency and minimize chemical shift displacement errors. To accommodate the high bandwidth requirements at 7 T, a single-shot echo planar readout was used for rapid simultaneous encoding of the temporal dimension and 1 spatial. \( B_0 \) and \( B_1 \) field aspects specific for 7 T were studied together with correction procedures, and feasibility of the EPSI MEGA-sLASER technique was tested in vivo in 5 healthy subjects.

Results: Localized edited spectra could be measured in all subjects giving spatial GABA signal distributions over a central brain region, having 45- to 50-Hz spatial intervoxel \( B_0 \) field variations and up to 30% \( B_1 \) field deviations. MEGA editing was found unaffected by the \( B_0 \) inhomogeneities for the optimized sequence. The correction procedures reduced effects of intervoxel \( B_0 \) inhomogeneities, corrected for spatial editing efficiency variations, and compensated for GABA resonance phase and frequency shifts from subtle motion and acquisition instabilities. The optimized oscillating echo-planar gradient scheme permitted full spectral acquisition at 7 T and exhibited minimal spectral-spatial ghosting effects for the selected brain region.

Conclusion: The EPSI MEGA-sLASER technique was shown to provide time-efficient mapping of regional variations in cerebral GABA in a central volume of interest with spatial \( B_1 \) and \( B_0 \) field variations typical for 7 T.

Keywords
7T, edited MRSI, EPSI, GABA
1 | INTRODUCTION

Gamma-aminobutyric acid (GABA) is the major inhibitory neurotransmitter in the central nervous system and is suggested to be involved in psychiatric and neurological conditions, such as schizophrenia,1 depression,2 epilepsy,3 and Parkinson’s disease4 as well as healthy brain aging.5 Proton magnetic resonance spectroscopy (1H-MRS) can be used to noninvasively assess concentrations of neurometabolites in vivo. Spectra from different metabolites can be separated because each metabolite has its characteristic resonance frequency pattern attributed to the specific chemical environment. Assessment of less-abundant neurochemicals, such as GABA, is hampered by the overlap of spectra from more-abundant metabolites together with limitations in peak linewidths and peak frequency separations.

The J-coupling between the spins of the three CH2 groups of GABA can, however, be utilized to separate the GABA signal. With the Mescher-Garwood (MEGA) J-difference editing technique,6 the J-coupling between the 1.89- and the 3.01-ppm (parts per million) GABA resonances is utilized by a selective pulse applied at the 1.89-ppm resonance to refocus the J-coupling evolution at the 3.01-ppm resonance (ON experiment). The J-evolution is left undisturbed in a second (OFF) experiment, and the 3.01-ppm GABA resonance is isolated after spectrum subtraction.

Further facilitation of the assessment of low-concentration GABA from the 1H-MRS spectrum can be achieved by an increase in magnetic field strength, giving both an increased frequency separation between resonances and an increased sensitivity. Although there is still complete overlap of the GABA resonance with other resonances at 7 tesla (T), cerebral concentrations of GABA have been reported without editing in humans at improved precision compared to lower field strengths using short echo time single-voxel localization techniques.7 The accuracy of unedited measurement of GABA at short echo time is, however, sensitive to experimental conditions.8

The increases in sensitivity and chemical shift dispersion are both favorable for GABA editing, but higher field strengths also imply hampered signal localization from increased chemical shift displacement (CSD) errors. GABA editing using MEGA and semilocalization by adiabatic selective refocusing (MEGA-sLASER) was demonstrated to give efficient spectral editing of the 3.01-ppm resonance with minimal CSD errors at 7 T.9

Studies of regional variations in GABA will benefit greatly from spatial mapping of concentrations using magnetic resonance spectroscopic imaging (MRSI) techniques. MRSI at 7 T requires additional considerations associated with the increased spatial variations in the transmitted radio frequency field (B1) and the permanent magnetic field (B0) that need to be taken into account during data acquisition and postprocessing. Also, the increased radio frequency (RF) power deposition at high field, in combination with high-bandwidth adiabatic pulses, leads to an increased minimum repetition time (TR) as compared to 3 T. However, MRSI can be significantly accelerated by simultaneous encoding of the temporal and spatial dimensions using oscillating readout gradients, as exemplified by, for example, spiral gradient sampling10 and the echo planar spectroscopic imaging (EPSI) technique.11 Utilizing such techniques at 7 T is challenging with respect to gradient performance because of the increased bandwidth required to sample all metabolite signals critically.

The aim of our study was to develop, optimize, and test an accelerated edited MRSI technique for spatial mapping of GABA in the human brain at 7 T. The MEGA-sLASER technique was applied for GABA editing and localization, in combination with a fast noninterleaved EPSI readout to accommodate the high-bandwidth requirements at 7 T. The B0 and B1 field aspects specific for 7 T were studied together with correction procedures, and the feasibility of the EPSI MEGA-sLASER technique was demonstrated in vivo.

2 | METHODS

Measurements were performed on a whole-body 7 T MR scanner (Achieva; Philips, Cleveland, OH) interfaced to a 2-channel volume transmit head coil and 32-channel receiver array (Nova Medical, Inc, Burlington, MA). All measurements were performed in accord with local ethical protocols.

2.1 | Sequence design

A volume-selective MEGA-sLASER sequence9 was combined with an echo-planar gradient waveform during signal readout (Figure 1). Frequency offset corrected inversion FOCI refocusing pulses12,13 (duration = 5 ms, HSn = 2, FOCI factor = 5, B1max = 18 uT) were used in the volume selection giving a CSD error of 2.5% between 1.89 and 3.01 ppm in the refocusing directions. The CSD error along the excitation direction from the asymmetric excitation pulse was 5%. MEGA editing was performed as described elsewhere.9 A crusher gradient configuration minimizing stimulated echo contributions was used. To minimize the effects from spatial B0 variations on the editing performance across the volume selection region (volume of interest; VOI), Gaussian RF pulses of 5-ms duration (full width of 95% maximum of 100 Hz) were used to cover the frequency range of the spatial B0 variation over the VOI. The offset frequency was set to 1.9 ppm in the edit ON acquisitions, and to 7.5 ppm in the edit OFF acquisitions.
An oscillating echo-planar (EP) gradient with plateau sampling was used for simultaneous readout of the temporal dimension and 1 spatial. The optimized gradient waveform had a spectrum bandwidth of 2662 Hz and 62% sampling efficiency (slew rate = 133 T/m/s, gradient amplitude = 9.4 mT/m, ramp time = 70 μs, plateau = 230 μs, pixel bandwidth = 4.3 kHz, train duration = 192 ms, sample points = 512).

Phase encoding was used to cover the second spatial dimension. The edit ON/OFF alternation was placed in the innermost loop of the scan execution followed by a phase-encoding loop and finally an outer averaging loop. This minimizes subtraction errors between the ON and OFF spectra.

2.2 | Reconstruction, data processing, and analysis

The reconstruction and analysis software was developed in-house using Matlab (The MathWorks, Inc, Natick, MA). The calculation steps were applied in the following order: (1) flipping of odd-numbered echoes (readout along $k_x$) and correction for temporal odd-even echo shifts; (2) spatial domain fast Fourier transforms; (3) voxel location-dependent coil weight assessment; (4) coil phase angle assessment and correction; (5) voxel location weighted time-domain signal addition over coils; and (6) voxel location phase angle assessment and alignment. Steps 1 through 6 were first applied to the water reference scan and next to the water-suppressed scan while reapplying the coil weights and phase angles determined from the water reference scan. This was, in turn, followed by: (1) eddy current compensation (ECC) \cite{14,15}; (2) regridding of the data acquired on the non-Cartesian $k_x$-time-domain trajectory to a Cartesian grid; (3) k-space Hamming filtering; and (4) 27- and 13-Hz spectral broadening, pre- and post choline frequency, and phase alignment (FPA), respectively.

For enhanced separation of edited GABA signals in regions with remaining frequency and phase shifts, spectral FPA of the choline peak \cite{9,16} was applied as a final processing step before subtracting the edit ON and OFF sets to generate the final GABA and metabolite spectra. The choline peak FPA was performed by comparing to the average spectrum for the ON and OFF measurement, separately, followed by an alignment of the summed ON and OFF spectra using the choline resonance. $B_1$-mapping \cite{17} was included in the scanning protocol to enable correction of varying editing efficiency (EE). A quantum mechanical simulation (result in Figure 2) of the effect of the $B_1$ field variation on only the MEGA editing pulse and on the EE was performed in VESPA. \cite{18,19}

A third-order polynomial was fitted to the simulated signal amplitudes in the normalized $B_1$ field strength range 60% to 140%, and the resulting EE function was applied to correct the edited spectra normalized to the total creatine signal of the OFF-pulse spectra. The Lorentzian line shape was fitted to the creatine signals (fit error threshold = 20%). The Gaussian line shape (30-Hz fixed linewidth) was fitted to the GABA+ (GABA plus co-edited macromolecules) signals, using a fit error threshold of 30%, to allow fitting of not fully separated GABA+ signals. The fit error is the percentage root mean square difference between data and line-shape model.

2.3 | Sequence tests

The performance of the EPSI MEGA-sLASER sequence was tested in vivo in 5 healthy subjects (2 female, 3 male) of 25 to 45 years of age. The scanning protocol included 2 EPSI MEGA-sLASER sets, a T1-weighted 3D magnetization-prepared rapid gradient echo, and a 3D gradient echo dual TR $B_1$ map.\cite{17} The EPSI MEGA-sLASER imaging plane was a transversal midbrain slice above the ventricles (field of view = 160 × 160 mm², slice thickness = 22 mm, VOI = 80

![Sequence diagram for the echo-planar MEGA-sLASER sequence with adiabatic refocusing (FOCI) and 5-ms Gaussian pulses for the spectral editing (MEGA) followed by prephase (px) and phase-encoding (py) gradients before the spatial and temporal encoding echo-planar readout (EPSI) ]
\[ \times 80 \times 22 \text{ mm}^3, \text{matrix size} = 16 \times 16, B_{\text{max}} = 18 \mu T, \text{pencil beam second-order shimming, in-plane voxel size} = 10 \times 10 \text{ mm}^2 \text{ [effective voxel size} = 3.7 \text{ mL}], \text{TR} = 4281 \text{ ms, echo time} = 74 \text{ ms}. \]

The protocol further consisted of 1 water-suppressed scan (VAPOR [variable pulse power and optimized relaxation delays],\(^{20}\) window = 200 Hz) of 9 min duration with 4 signal averages (number of signals averaged; NSA) and a water reference scan (duration 2 minutes) with NSA = 1 and with the editing pulses turned off.

3 | RESULTS

Spatially localized ON and OFF spectra were measured in all subjects. VOI average and ranges for the spatial distributions (Figure 3, row A) of the creatine signal-to-noise ratio (SNR) of the OFF-pulse measurement are shown in Supporting Information Tables S2, together with the errors in the fit to the creatine and corrected GABA+ signals. Corresponding spatial distributions of the fit errors are shown in Supporting Information Figure 9. VOI ranges for the spatial distributions of the normalized \(B_1\) field strength deviations (Supporting Information Figure 8) are shown in Supporting Information Table S1, together with the maximum of the corresponding spatial distributions of the EE correction factors (Supporting Information Figure 8). The intravoxel \(B_0\) field inhomogeneities, assessed from the linewidth of the 3.01-ppm creatine resonance from the OFF measurements, varied as shown in Supporting Information Table S1, corresponding to the distributions in Figure 4 and Supporting Information Figure 7. Intervoxel \(B_0\) field variations, assessed from the creatine resonance frequency shifts (Figure 4E), were significantly reduced by the ECC and varied in ranges from minimum ± 3 Hz to maximum ± 11 Hz (Supporting Information Table S1).

**FIGURE 2** Simulated \(B_1\) field strength dependence of the signal amplitude of the MEGA edited GABA spin system at 3.01 ppm (sampling bandwidth = 2 kHz, 2048 sampling points). The typical normalized \(B_1\) field strength range 80% to 125% over the VOI is indicated (blue rectangle)

**FIGURE 3** Spatial distributions for the healthy volunteer test scans (no. 1-no. 5) of the OFF-pulse measurement creatine (Cr) signal (row A), uncorrected GABA+ signal (row B), and Cr-normalized GABA+ signal after editing efficiency correction and choline peak frequency and phase alignment (row C)
The impact on the spatial distribution of the measured GABA+ signal from the spectral correction procedures was demonstrated for 1 subject (Figure 5 showing the impact of creatine normalization, EE correction, and choline peak FPA separately). Creatine-normalized GABA+ levels of 0.22 were close to uniformly distributed in a central region of the VOI, corresponding to white matter, with an elevation toward the left and right VOI edges (Figure 5C). This elevation was further enhanced by the EE correction (Figure 5D). The choline FPA gave an increased fraction of voxels with fitting error below the chosen threshold (Figure 5E and Supporting Information Figure S1D). Uncorrected GABA+ signal distributions are shown in Figure 3, row B, for all the 5 scans on healthy subjects. The more pronounced choline peak residuals, disabling a fit to the uncorrected GABA+ signals in central voxels, was present for 1 of the 5 healthy subject scans (Figure 3, row B, no. 1). An increase in GABA+ signal toward the left and right VOI edges were a general characteristic across subjects after Cr normalization (Figure 3, row C). A lower spectral quality was observed in some voxels on the VOI edge columns (Supporting Information Figure S1D). A corrected low GABA+ signal centrally in the VOI was further apparent for 2 subjects (Figure 3, row C, no. 2 and no. 4).

4 | DISCUSSION

An accelerated edited MRSI technique for spatial mapping of GABA+ signal in the human brain at 7 T was developed. The EPSI MEGA-sLASER technique was tested in 5 healthy subjects and studied with respect to effects from spatial variations in $B_0$ and $B_1$ fields specific to 7 T.

4.1 | $B_1$ field strength considerations

Simulation of the edited 3.01-ppm GABA signal shows that with increased deviation from the normalized $B_1$ field, the line shapes deteriorate along with the signal amplitude (Figure 2). The maximum observed deviations from the nominal $B_1$ field strength was $-31\%$ and $-29\%$ (Supporting Information Table S1), occurring at the VOI edge (Figure 3, row C, no. 3 and no. 5), corresponding to EE correction factors of 1.66 and 1.58, respectively (Supporting Information Table S1). The correction can also be performed for this substantial $B_1$ field strength deviation of $-30\%$, but the GABA signal line shapes (Figure 2) suggest that corrections may no longer be possible for $B_1$ field strength deviations beyond $-50\%$. In order to approach full brain coverage, other $B_1$ compensation techniques will be required.
4.2 | B₀ field strength considerations

For the 5 subjects, the typical intervoxel $B₀$ inhomogeneities were spanning ranges of typically 45 to 50 Hz and 6 to 11 Hz before and following (Supporting Information Table S1, Supporting Information Figure 7) the ECC, respectively, with a maximum range of 22 Hz following ECC for subject no. 3. The corresponding intravoxel linewidths were in the range of 5–32 Hz, with a maximum of 46 Hz for subject no. 3 (Supporting Information Table S1, Supporting Information Figure S2). All $B₀$ inhomogeneities were thereby well within the 200- and 100-Hz bandwidths of the water suppression (VAPOR) and MEGA editing pulses, respectively. Water suppression and MEGA editing were therefore considered unaffected by $B₀$ inhomogeneities across the image plane in all subjects.

4.3 | GABA+ signal and spatial distributions

The voxels on the outermost VOI edge columns with less well separated corrected GABA+ signal were typically associated with a low SNR. The volume selection transition region of the FOCI refocusing pulses could, to some extent, have affected the signal on the outermost VOI edges. A major part of the outermost VOI edge voxels were, however, having a well separated GABA+ signal with a quality comparable to the inner VOI voxels. The central regions where corrected low GABA+ signal were observed for 2 subjects (Figure 3, row C, no. 2 and no. 4) could not be identified as to have insufficient spectrum quality from, for example, $B₀$ and $B₁$ field effects or from subject motion, suggesting that they are true reflections of regional variations in GABA+.
4.4 | Motion considerations

In the subtracted spectra, subtle motion during scanning is most likely the main source of degradation in terms of frequency and phase shifts around the 3.01 ppm GABA+ signal (Figure 5D and Supporting Information Figure S1C). Motion artifacts were, to a large extent, corrected by the choline peak FPA (Figure 5E and Supporting Information Figure S1D). Minimization of larger motion artifacts would likely require the technique to be combined with prospective motion correction.\(^{10,21,22}\)

4.5 | Temporal-spatial gradient readout considerations

The oscillating EP gradient was optimized to allow for full spectral acquisition in a single shot at a spatial resolution permitted by the sensitivity at 7 T, and to minimize the influence of motion on the ON/OFF subtraction pairs. Only the central k-space is critically sampled, and spectral-spatial ghosting is thus expected at positions where the signal from off-resonance metabolites varies abruptly, that is, at the border between cerebrospinal fluid (CSF) and gray matter.\(^{23}\) This was not observed to be problematic in the volunteer measurements for the selected slice, but should be more pronounced near more CSF-containing regions. Negligible deviations were observed between the spatiotemporal encoding performed by the optimized EP gradient, compared to a corresponding fully phase encoded version of the sequence (chemical shift imaging [CSI] MEGA-sLASER) in separate phantom verification measurements (data not shown). No significant scanner drift was observed with the optimized EP gradient, although this could lead to significant issues if not accounted for.\(^{24}\)

Other oscillating gradient readout acceleration strategies, such as spiral readouts with high acceleration factors,\(^{4.6}\) have been shown feasible at lower field strength.\(^{10}\) At 7 T, however, these pulses have a less optimal bandwidth and a compromised spectral transition region as compared to the FOCI pulse shapes, which could have a more pronounced effect on the outermost VOI edges. A novel alternative to the VOI selection technique for avoiding lipid signals could be to use specialized hardware in terms of a separate lipid suppression coil\(^{25}\) in combination with the EP readout, following the MEGA pulse set, while excluding the SAR intense refocusing part of the sequence. This would lead to a further significant reduction of the acquisition time.

4.6 | Total acquisition time and specific absorption rate

The EPSI acceleration factor was 16 compared to a fully phase encoded acquisition (CSI), giving a total acquisition time of 9 min with 4 signal averages for acquiring the GABA+ signal on a 16 × 16 grid. The specific absorption rate (SAR) of the RF pulses was the predominant limiting factor determining the minimum TR for the EPSI MEGA-sLASER. Shortening of the TR may be possible if less SAR-restrictive localization schemes are realized. Gradient offset independent adiabaticity pulses exhibit lower power deposition and have successfully been applied at 3 T field strength.\(^{10}\)

5 | CONCLUSION

A MEGA-sLASER sequence was combined with an optimized EP readout for accelerated MRSI while utilizing the high sensitivity at 7 T (EPSI MEGA-sLASER). The technique was fulfilling the high-bandwidth demands for 7 T brain spectroscopy and was demonstrated in 5 healthy subjects to provide time-efficient mapping of regional variations in cerebral GABA+ in a large central VOI with typical spatial B1 and B0 field variations. Effects of motion and acquisition instability were considered and could, to a large extent, be retrospectively corrected. Combining the technique with prospective motion correction techniques may lead to further improvements in cases of pronounced motion. The technique is expected to ultimately benefit studies of neurological and psychiatric disorders and healthy brain maturation and aging.

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**SUPPORTING INFORMATION**

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

**FIGURE S1.** Mosaic spectra view of the spatial distributions of the signals for a healthy volunteer test scan (no1), corresponding to Fig. 4, of the ON-pulse measurement (A), the OFF-pulse measurement (B), subtracted GABA signal (C), and GABA signal after creative normalization, EE correction and choline peak frequency and phase alignment (D). Example VOI sub-region with correctable (D, orange region) remaining choline and creatine residuals in subtracted spectra (C, orange region) is indicated.

**FIGURE S2.** Spatial distributions for the healthy volunteer measurements of the intra-voxel (creatine peak linewidth) and inter-voxel (creatine peak frequency offset) B₀ inhomogeneity, corresponding to the values in Supporting Information Table S1.

**FIGURE S3.** Spatial distributions for the healthy volunteer measurements of the editing efficiency (EE) correction factor and the normalized B₀ field strength, corresponding to the values in Supporting Information Table S1.

**FIGURE S4.** Spatial distributions for the healthy volunteer measurements of the errors in the fit to the creatine and corrected GABA+ signals, corresponding to the values in Supporting Information Table S2.

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