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Metabolite Phantom Correction of the Nonuniform Volume-Selection Profiles in MR Spectroscopic Imaging: Application to Temporal Lobe Epilepsy

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BACKGROUND AND PURPOSE: In MR spectroscopic imaging (MRSI), the volume-selection profiles of metabolites differ from each other. These differences cause variations in metabolite intensities, which are particularly prominent when the hippocampi are evaluated. We hypothesize that the errors arising from these effects cause notable artifact when temporal lobe epilepsy (TLE) is lateralized with MRSI.

METHODS: We examined a metabolite phantom, control subjects, and patients with TLE by using MRSI. We calculated the error arising from the different volume-selection profiles of metabolites *in vitro* and evaluated this correction in the examination of the control subjects and in the lateralization of epilepsy in the patients.

RESULTS: Without a correction, a considerable error in the metabolite content existed, even deep inside the spectroscopic volume of interest. The result was false asymmetry (P < .008) in the hippocampi of control subjects. Among the 11 patients, TLE was correctly lateralized in three only after the correction was made, and in one, TLE was incorrectly lateralized.

CONCLUSION: The volume-selection profiles of *N*-acetylaspartate, choline, and creatine differ enough to cause a significant error, even in the metabolite ratios, when patients with TLE are examined with MRSI. We propose a simple phantom method to correct for this error without a need to modify the pulse sequence.

MR spectroscopic imaging (MRSI) is useful in localizing the focus of seizure in patients with temporal lobe epilepsy (TLE) (1–5) or frontal lobe epilepsy (6, 7). The diseased lobe is identified by assigning an asymmetry percentage that reflects the difference in *N*-acetylaspartate(NAA)/(choline [Cho] + creatine [Cr]) between the temporal lobes. A decreased value of NAA/(Cho + Cr) defines the ipsilateral or diseased side (8).

Spectroscopic imaging with a point-resolved spectroscopy (PRESS) chemical-shift imaging (CSI) sequence (9) combines MRSI with the selection of a

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volume of interest (VOI) with PRESS localization. The VOI is localized with three section-selective pulses (one 90° and two 180° pulses), and the spatial distribution of metabolites is obtained by using phase encoding. Because the VOI is selected by using pulses applied simultaneously with magnetic field gradients, protons of metabolites with different chemical shifts obtain different excitation volumes. These volumes are displaced with respect to the transmitter frequency. The displacement for the signal intensity of a metabolite is proportional to the difference between the resonance frequency of the signal intensity and the transmitter frequency and inversely proportional to the gradient strength. The dimensions of the VOI are determined by using the pulse bandwidth and the gradient strength. Because of radiofrequency power requirements, the bandwidth of the pulse is usually kept constant, and the desired VOI dimension is obtained by varying the gradient strength. This method leads to different volume displacements for VOIs of different sizes. Furthermore, the excitation profiles of section-selective pulses that define the VOI deviate from an ideal rectangular shape (10). This feature is especially pronounced for the 180°

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	Epilepsy Clinical		MRSI Lateralization		
Patient/Age (y)/Sex	Lateralization and Type	MR Imaging	No Correction	Pulse Correction	Comment*
1/9/M	L TLE	L T dysmyelination	None	R	Seizure-free without operation
2/10/F	L TLE	L hippocampal dysgenesis (possible)	None	None	Seizure-free without operation
3/16/F	L TLE, possible R FLE	Normal	L	L	No resection, VNS
4/16/F	L TLE	L MTS	L	L	L TL resection, seizure-free
5/13/F	R/L TLE	Normal	L	L	No operation, monthly seizures
6/17/F	R/L TLE	R/L hamartomas (tuberous sclerosis)	None	R	Daily seizures from R TL, rare from L TL, VNS
7/9/M	R TLE	R hippocampal edema	None	R	No operation, rare seizures
8/12/F	R TLE	R MTS	L	L	R TL resection, seizure-free
9/15/F	R TLE	R hippocampus, small	L	L	R TL resection, seizure-free
10/15/M	R TLE	Sturge-Weber syndrome R TO	R	R	Seizure-free without operation
11/34/F	R TLE	Normal	None	R	R TL resection, seizure-free for 6 mo, postoperative seizures from L TL
12/9/F	Multifocal	R hippocampus, small	None	None	Seizures not from TL
13/14/M	L TOE	Dysgenesis L TO	None	None	None
14/15/M	Undetermined	L hippocampus, abnormal	None	None	Not operation, rare seizures
15/50/F	R TLE or R FLE	Normal	None	None	Not operation, monthly seizures

TABLE 1: Clinical, MR imaging, and MRSI findings in patients with TLE

Note.—FLE = frontal lobe epilepsy, TOE = temporo-occipital epilepsy, MTS = mesial temporal sclerosis, TL = temporal lobe, VNS = vagal nerve stimulation.

* Rare seizures were fewer than one per 6 months.

section-selective pulses of the PRESS sequence. The nonideal excitation profile, together with the volumedisplacement artifact originated from the chemical shift, leads to different volume-selection profiles for the metabolites and therefore distorts the spatial distribution of their signals.

The problem associated with the nonideal volumeselection profiles of metabolites in the PRESS CSI sequence is known, and techniques have been implemented to correct for it or minimize it. These techniques include modification of the pulse sequences (i.e., imaging of the excitation profile and calculating a correction or switching the order of the 90° and 180° pulses) (11-13). To our knowledge, the effect of this error in the lateralization of epileptic foci has not been reported. We hypothesized that the errors arising from these effects cause a significant artifact when TLE is lateralized by using MRSI. To address this problem, we corrected the effects of different volume-selection profiles of metabolites by using a homogenous phantom containing the metabolites studied in vivo. This correction was evaluated in control subjects and in patients with TLE.

Methods

Patients and Control Subjects

Fifteen patients (age range, 9–50 years) with epilepsy (one left handed [patient 15], 14 right handed) and eight healthy, right-handed control subjects (age range, 11–46 years) were examined by using MRSI. The diagnoses of epilepsy were based on detailed seizure histories, clinical findings, and EEG

or video EEG (ictal video EEG recordings 11 patients) and MR imaging results. On the basis of these data, 11 patients were determined to have TLE, one had TLE or frontal lobe epilepsy, one had temporo-occipital epilepsy, one had multifocal epilepsy (with an abnormal right hippocampus), and one had undetermined epilepsy (with abnormal left hippocampus on MR imaging) (Table 1). The study protocol was approved by the Ethical Committee of Helsinki University Central Hospital, and written informed consent was obtained from all patients and parents. The MRSI data were not used for clinical decision making in planning temporal lobectomy. In the epilepsy surgery program, four patients (Table 1) proceeded to temporal lobectomy.

MRSI Study

The measurements were performed on a 1.5-T MR system (Magnetom Vision; Siemens, Erlangen, Germany). A standard PRESS-CSI sequence was used with a TR/TE of 2600/270 and a 16 \times 16-cm FOV with 16 \times 16 phase encoding steps. In the PRESS CSI sequence, the transmitter frequency is set at water resonance. The VOI was carefully placed to symmetrically cover the hippocampi by using T1-weighted localizer images. The VOI dimensions were as follows: thickness = 15 mm, anteroposterior dimension = 60 mm, and right-left dimension = 80 or 100 mm depending on the size of the brain (80 mm) in 11 patients and six control subjects and 100 mm in four patients and two control subjects). Shimming of the VOI and water suppression were manually adjusted after automatic global shimming was done. In the shimming process, the three linear shim gradients were adjusted by maximizing the peak intensity and by minimizing the full width of the water resonance at half height while the shape of the peak was simultaneously preserved.

The data were processed by using a CSI postprocessing program (Luise program, Vision; Siemens). Preprocessing steps included zero filling to 1024 data points, filtering with a



Fig 1. Axial/coronal T1-weighted image at the level of the hippocampi with the MRSI voxels *1*, *2*, and *3* chosen to represent the hippocampus. The 100- and 80-mm-wide VOIs are shown (*white rectangles*).

50% Hamming filter, and Fourier transformation. One author (J.O.J.L.) without knowledge of the participants manually corrected the phase and baseline of the spectra. The signal intensities (i.e., signal intensity areas) of NAA, Cho, and Cr were determined by using Gaussian line-shape fitting. To lateralize the patients' epilepsy, we used an average asymmetry percentage from the three voxels per hippocampus (Fig 1) as follows: $2 \times 100(C - I)/(C + I)$, where C and I indicate contralateral and ipsilateral NAA/(Cho + Cr), respectively. The maximal asymmetry of the controls was used as a threshold to classify the patients.

Phantom Correction

A homogenous spherical phantom (GE Medical Systems, Milwaukee, WI) containing an aqueous solution of NAA, Cho, and Cr was measured by using parameters and VOIs identical to those used for measurements in patient, both without and with water suppression. The result of the unsuppressed measurement was used to examine the combined effect of volume displacement and the nonideal excitation profile of the PRESS CSI sequence in the right-left direction. The profile was calculated by averaging the water intensity of the two middle rows of voxels from the entire FOV. Water-suppressed MRSI was used to determine the error and the corrections for NAA, Cho and Cr. The principle behind the phantom correction was the valid assumption that the metabolites are homogeneously distributed throughout the phantom. Therefore, any systematic errors in the spatial metabolite profile were due to imperfections in the PRESS CSI sequence. To correct for the errors resulting from the sequence, each voxel in the phantom was used to correct the corresponding voxel from the in vivo measurements with the identical VOI. The equation used for correction was the following: $S_{\rm C} = S_{\rm M}(S_{\rm Pmax}/S_{\rm P})$, where C is the corrected signal intensity, M is the measured signal intensity in vivo, P is the corresponding voxel signal intensity in the phantom, and Pmax is the voxel with maximum signal intensity in the phantom The average metabolite asymmetries of the phantom were studied with the percentage $2 \times 100[(R - L)/(R + L)]$, where R and L are the right and left voxels, respectively, that corresponded to the location of the hippocampus.



Fig 2. Phantom profile for water signal intensity with the 180° Hamming-weighted sinc pulse of the PRESS CSI sequence (solid line with black boxes) obtained by selecting the right-left dimension of the 100-mm VOI (vertical lines). Profile shifted by 10 mm (dashed line with white boxes). This displacement approximately corresponds to the chemical-shift difference between water and NAA with the typical excitation bandwidth of 1500 Hz for the section-selective 180° pulse. Images shows the difference between the original profile and the shifted profile spanning the VOI (dotted line with gray circles). The water intensity on the y axis is expressed in institutional units (i.u.), and the x axis corresponds to the 10-mm voxels in the 160-mm FOV.

Evaluation of the Method

To evaluate the phantom correction, we examined the metabolite content of the hippocampi in the control subjects. The content of NAA and Cho + Cr the right and left hippocampus was calculated from representative voxels 1–3. The paired Student t test was used to test for differences in the metabolite content between the right and left hippocampus in the control subjects. The mean and standard deviation of the right-to-left ratio of NAA, Cho + Cr, and NAA/(Cho + Cr) were also calculated.

Results

Phantom Measurements

Figure 2 shows the measured water excitation profile in the right-left direction, a profile artificially displaced by 10 mm, and their difference. This 10-mm displacement was a good approximation of the volume displacement induced by the chemical-shift difference between NAA and water for a 100-mm-wide VOI on our system. Figure 2 demonstrates the problems arising from effects originating from the chemical shift when metabolite intensities with different chemical shifts are compared. The difference was minimal at the center of the VOI but increased with increasing distance from the center.

In the metabolite measurements in the phantom, the voxels on the right side of the VOI had consistently increased intensities for all three metabolites. Table 2 presents the asymmetries of NAA, Cr, Cho, and NAA/(Cho + Cr) for the averaged voxels corresponding to the location of the hippocampus for the *in vivo* measurements with the-80 and 100-mm-wide

 TABLE 2: Intensity asymmetry percentage in the phantom voxels corresponding to the hippocampi

VOI Width (mm)	Cho	Cr	NAA	NAA/ (Cho + Cr)
80 mm	7.9 ± 3.2	9.3 ± 3.4	$\begin{array}{c} 18.1 \pm 1.4 \\ 9.3 \pm 1.3 \end{array}$	9.4 ± 1.5
100 mm	4.0 ± 4.4	7.5 ± 1.9		3.8 ± 1.5

Note.—Intensity asymmetry percentage = $2 \times 100[(R - L)/(R + L)]$. Data are the mean percentage ± standard deviation.

TABLE 3: Means and standard deviations for the right-left ratios of hippocampal metabolites in control subjects before and after correction

Metabolite	Me	an	Standard Deviation	
or Index	Before	After	Before	After
NAA	1.15	0.98	0.07	0.05
Cho + Cr	1.10	1.02	0.09	0.08
NAA/Cho + Cr	1.05	0.97	0.06	0.06

VOIs. NAA had the largest asymmetry and Cho the smallest asymmetry, in accordance with their respective chemical shifts. The error for NAA was 18% for the 80-mm VOI.

In Vivo Measurements

Figure 1 shows the six voxels (three per hippocampus) chosen to represent the hippocampi. The location of the hippocampus near the base of the skull and the paranasal sinuses complicated shimming and water suppression in the measurements. Because of this effect, 50% of the spectra in voxel 1, which was closest to the most inhomogeneous region, had to be discarded from the analysis. In voxels 2 and 3, 90% of the spectra were free from distortion and were included.

Control Subjects

Without the correction, the right hippocampus had NAA (P = .0009) and Cho + Cr (P = .008) intensities significantly higher than those of the left hippocampus. After phantom correction, no significant asymmetry remained (P > .21). When the correction was applied, the right-left ratio approached the value for one for the individual metabolites, as well as for NAA/(Cho + Cr). The standard deviation of the right-left ratio decreased for NAA and Cho + Cr but did not change considerably for NAA/(Cho + Cr) (Table 3).

The asymmetry percentages for NAA/(Cho + Cr) in control subjects were calculated, and the highest was chosen as the threshold level for the evaluation of patients. The threshold levels were 13% and 16% with and without correction, respectively.

Patients

The patients were classified according to their asymmetry percentage for NAA/(Cho + Cr). In pa-

tients whose asymmetry percentage exceeded the threshold level of the control subjects, epilepsy was considered lateralized. With uncorrected MRSI, epilepsy was lateralized to the left in five patients, it was lateralized to the right in one patient, and it was not lateralized in nine. Correction for the effects of the volume-selection profile changed the MRSI lateralization in four patients from not lateralized to lateralized to the right, resulting in five cases lateralized to the left, five to the right, and five not lateralized.

In 11 patients, the classification did not change after the correction. In four patients, the asymmetry increased by 9% for NAA/(Cho + Cr) with the 80-mm VOI (Table 2) from 7-15% to 16-24%, altering the classification from not lateralized to lateralized to the right. In three of these patients, lateralization could be considered accurate after the correction: Two (patients 7 and 11) had converging clinical, video EEG, and MR imaging findings, and one (patient 6) had daily right temporal seizures and monthly left temporal seizures. (Both seizure types were documented on video EEG.) In the fourth patient (patient 1), the corrected lateralization disagreed with MR imaging data; this patient, whose seizures had not been recorded on video EEG, had had left temporal dysmyelination but was seizure-free for several years before MRSI. In one patient with TLE (patient 2) and all four patients with other types of epilepsy (patients 12–15), their condition was not lateralized even after the correction. In three patients (patients 1, 8, and 9), MRSI lateralization disagreed with clinical lateralization. In two (patients 8 and 9), lateralization to the right was confirmed by a good outcome after right temporal lobectomy.

Discussion

Various techniques have been developed to minimize or correct for the effects of the nonideal excitation profiles of volume-selection pulses and for volume-displacement effects with MRSI. The VOIprescribing pulses have been switched to minimize the error in the most important direction. When the less distorted 90° pulse prescribes the VOI in the right-left direction, less error is introduced (11). The transmitter frequency can also be shifted from water at 4.7 ppm to the metabolite region at 2.7 ppm to minimize the chemical shift-originated displacements of the metabolite VOIs, if the MR spectroscopic sequence supports this feature. The excitation profiles of the volume-selection pulses can also be measured and corrected for in postprocessing. The PRESS sequence can be modified to measure the pulse profile with conventional imaging (12, 13). When the profile is known, the errors resulting from the volume displacement of the PRESS VOI can be calculated. However, this calculation requires relatively detailed knowledge of sequence parameters, which is usually not readily available. The proposed phantom method is simple for correcting the problem without the need to resort to technical modifications or to detailed parameter data. For example, no knowledge of the gradients, pulse bandwidths, or transmitter frequency is needed. The method can easily be applied in a clinical environment, and the state of the hippocampi can be evaluated by means of direct left-right comparison of metabolite levels and ratios in a single patient.

Our phantom measurements showed that the metabolite profiles of a homogeneous sample were considerably asymmetric. The voxels on the right side of the phantom that corresponded to the location of the hippocampus had consistently increased metabolite intensities for the 80- and 100-mm VOI. This finding is demonstrated in Figure 2, where the displaced profile corresponding to NAA produced high intensity on the right side of VOI. Also, Cho and Cr had similar effects, as both signals had resonance frequencies lower than that of water. The degree of asymmetry was proportional to the chemical shift of the metabolite and the size of the VOI (Table 2), as predicted in theory. The asymmetries of the 100-mm VOI were small because the voxels corresponding to the hippocampus were deep inside the VOI, as shown in Figure 1. The evaluation of the control subjects showed that the correction improved the standard deviation of the right-left ratio for NAA and Cho + Cr and that the maximal asymmetry decreased.

In healthy control subjects, an even metabolite distribution between the right and left hippocampi can be assumed. This assumption has been shown to be true with single-voxel spectroscopy (14) and with MRSI (15). However, there is speculation about a natural asymmetry of NAA/(Cho + Cr) in the human brain, mainly due to the domination of the left hemisphere in more than 90% of the population. Accordingly, the right-left ratio would be less than 1, as Bernard et al (16) reported. Our original data showed significantly increased signal intensity for NAA and Cho + Cr in the right hippocampus, which contradicts the views of an even metabolite distribution and that of a natural asymmetry. By applying the phantom correction, the significant difference between the right and left hippocampi disappeared, and the average right-left ratio approached 1.

When we classified the patients' condition as lateralized or not lateralized with the threshold obtained from the control subjects, the classification changed in four patients from not lateralized to lateralized with the correction. In all patients whose classification changed, their disease was finally lateralized to the right. This was a result of the original tendency of the sequence to cause overestimation of the metabolites, as well as the NAA/(Cho + Cr) value, in the right hippocampus compared with the left (Fig 2). MRSI lateralization agreed with clinical lateralization in three of these patients (patients 6, 7, and 11) and with the postoperative result in one (patient 11).

In five patients, MRSI result were not lateralized even after the correction. This group included all four patients (patients 12–15) who were clinically determined not to have mesial TLE (multifocal, undetermined, temporal or frontal, and temporo-occipital TLE) and one patient (patient 2) with left hippocam-

pal dysgenesis. In two of the three patients whose MRSI lateralization disagreed with clinical lateralization (patients 8 and 9), the latter was confirmed by a good outcome after resection of the temporal lobe. This result supports the view Suhy et al (17) presented, in that metabolically abnormal hippocampus, as shown on MRSI lateralization, may not always be the primary origin of seizures. Accordingly, the degree of metabolic asymmetry between the hippocampi (i.e., lateralization) would not be a good prediction of the surgical outcome. Rather, the general metabolite level (i.e., the degree of NAA decrease) of both hippocampi would be a more important factor, where better-preserved hippocampal metabolism is predictive of a good outcome independent of asymmetry. Further studies are needed to elucidate the role of MRSI in the preoperative evaluation of patients with TLE.

We did not use segmentation to correct for possible CSF contamination of the data for two reasons. First, because the control subjects did not have hippocampal atrophy, segmentation was not regarded as being necessary, even when individual metabolites were evaluated. Second, in the patient evaluation, only metabolite ratios were used.

Conclusion

We measured and corrected the effects that the nonideal excitation profiles of the volume-selection pulses and the chemical shift-originated volume displacement introduce to the metabolite distribution in MRSI. These distorting effects were significant at the location of the hippocampus. The different classifications of the patients with and without correction indicated that a correction is necessary with MRSI. The use of a phantom correction for MRSI is easy to apply in clinical settings because it allows for direct and accurate left-right comparison of patient data, as well as for indications other than the assessment of TLE. The advantage of the method presented here is mainly its simplicity.

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