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Rapid MR Imaging of the Pediatric Brain Using the Fast Spin-Echo Technique

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Purpose: To evaluate diagnostic reliability and to establish optimal scanning techniques of a recently developed Fast Spin-echo MR pulse sequence that allows rapid proton density-weighted and T2-weighted imaging. **Methods:** We compared lesion conspicuity and signal intensity measurements on Fast Spin-echo and conventional spin-echo sequences in 81 patients ranging from 1 week to 25 years in age on a 1.5-T MR unit. A total of 28 Fast Spin-echo dual-echo images (14 slice locations) were obtained in 2:08 minutes with a 256×128 matrix or in 3:12 minutes with a 256×192 matrix at a TR of 2000 msec and two excitations. **Results:** Lesion conspicuity and characterization on Fast Spin-echo images compared favorably with conventional spin-echo images in our series when pseudo-TEs of 15 and 90 msec were employed for proton density-weighted and T2-weighted images, respectively. Fast Spin-echo images yielded diagnostic information in four nonsedated patients whose conventional spin-echo images were either degraded by motion or unobtainable. Fat signal remained bright on T2-weighted Fast Spin-echo images. Magnetic-susceptibility effects were slightly reduced with Fast Spin-echo but did not pose any diagnostic problem in our series. **Conclusion:** Diagnostically reliable rapid dual-echo brain images can be obtained with Fast Spin-echo sequences.

Index terms: Magnetic resonance, technique; Brain, magnetic resonance; Pediatric neuroradiology

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Magnetic resonance (MR) imaging has proven to be the study of choice for many neuroradiologic applications since its introduction into clinical practice. The popularity of the conventional spin-echo (CSE) sequence is attributed to its simple contrast behavior and relative insensitivity to magnetic-field inhomogeneity and magnetic-susceptibility effects (1).

One drawback of the CSE sequence is a relatively long acquisition time. A typical dual-echo CSE sequence employing 2000/30, 80/2 (TR/TE/excitations) and a 256×128 (frequency \times

phase) matrix requires nearly 9 minutes to complete. This long imaging time contributes to motion artifact and patient discomfort. The long imaging time is especially problematic in pediatric neuroimaging because of the increased need for sedation. In addition, higher resolution studies using larger matrices have been impractical with CSE techniques because of time constraints. For example, a CSE study (2000/30, 80/2) would take more than 13 minutes for a 256×192 matrix and more than 18 minutes for a 256×256 matrix.

The potential benefits of faster sequences have stimulated research in the development of MR sequences that reduce acquisition times without sacrificing the image quality provided by CSE techniques. Several methods have been implemented (2, 3), but poor image quality has hindered widespread clinical use.

Fast Spin-echo (FSE) generates dual-echo images in a fraction of the time required for CSE. FSE is based on the Rapid Acquisition with Relaxation Enhancement (RARE) sequences described by Hennig and others (4-6) but modified with a specific phase encode-reordering algorithm

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to provide flexible T2 contrast manipulation (7). The algorithm is based on the principle that tagging early echoes with small phase-encoding gradients minimizes T2-weighting, while tagging late echoes with small phase-encoding gradients maximizes T2-weighting (6). Adjusting the echoes that are tagged with small phase-encode gradients is the means by which pseudo-echo times (pTEs) ranging from the earliest echo time to the last echo time within a given echo train can be chosen.

The purpose of our study was to evaluate the diagnostic reliability of the dual-echo FSE brain images in comparison to CSE images and to establish optimal scanning techniques for FSE neuroimaging in pediatric and adolescent patients.

Materials and Methods

In the standard dual-echo CSE technique, a 90° excitation radiofrequency (RF) pulse is followed by a 180° RF refocusing pulse with echo sampling between 15 and 30 msec to produce a proton density-weighted (PDW) image. Then, another 180° pulse, with echo sampling usually between 70 and 100 msec, is performed to acquire data for the more heavily T2-weighted (T2W) image. The CSE sequence is depicted in Fig. 1A. Note that the sequence must be repeated for each individual phase-encoding step.

In FSE, a 90° pulse is followed by a series of 180° pulses that are closely spaced (15 msec in our study), resulting in the acquisition of eight echoes as opposed to the two

echoes obtained with the CSE technique (Fig. 1B). The first four echoes of the eight-echo train are used to collect four k-space (8) data lines for the PDW image, while the next four are used to gather four k-space data lines for the T2W image. Thus, FSE can theoretically reduce the time required to gather the full number of phase-encoding steps by a factor of four. Low phase-encoding gradient echoes give rise to the strongest signals and so dominate the contrast (6). Therefore, the extent of the T2-weighting for an FSE image is controlled by the choice of which echoes are encoded with the lowest phase-encoding gradients. This allows the operator to select an effective or pseudo-echo time (pTE), which is analogous to the echo time (TE) of CSE techniques (7).

FSE sequences were implemented on a 1.5-T Signa system (General Electric, Milwaukee, WI) equipped with actively shielded gradient coils. All studies were performed with the quadrature head coil designed to reduce RF power deposition and improve signal-to-noise ratio. The study was approved by the Committee on Clinical Investigation at the authors' institution.

The FSE dual-echo brain images were acquired in addition to conventional CSE images in 81 patients ranging in age from 1 week to 25 years. The composition of the study group is listed in Table 1. Two of the nonsedated patients had excessive motion artifact on initial T1-weighted sagittal CSE images. Only FSE axial images were obtained in these cases.

All CSE images were obtained in the axial plane with a TR of 2000 msec and TEs of 30 and 80 msec for PDW and T2W images, respectively. Other CSE imaging parameters included a 256×128 matrix, 2 excitations, 5-mm slice thickness, 2.5-mm interslice gap, a 20-cm field of

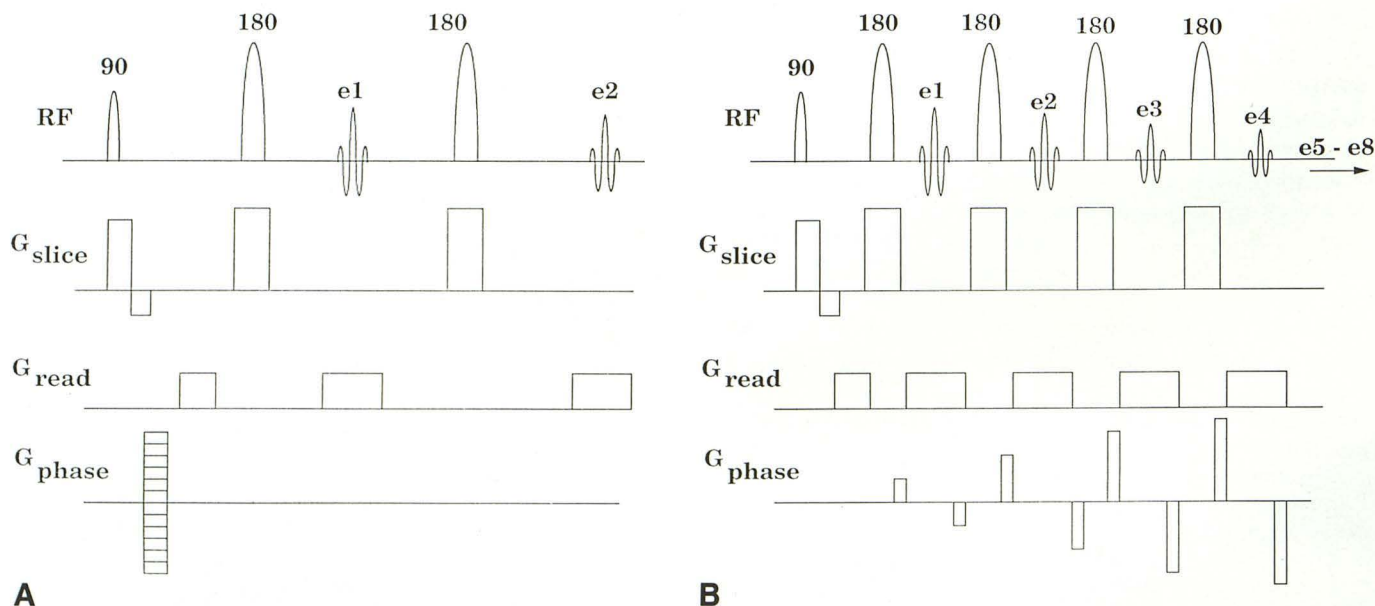


Fig. 1. A, CSE pulse sequence; the top line indicates the RF transmit pulses and the acquired echoes. The next three lines indicate the application of slice, read, and phase encode gradients (e = echo sampling).

B, FSE pulse sequence; lines are as in A. Note the collection of a large number of echoes (only four shown) and how distinct phase-encode gradients are applied to each echo and then "unwound" following echo collection.

TABLE 1: Composition of the study group (N = 81)

Diagnosis	No.
Neoplasm	
Astrocytoma	12
Oligodendroglioma	2
Germinoma	2
Ganglioglioma	1
Medulloblastoma	1
Meningioma	1
Other	1
Vascular lesions	
Cavernous hemangioma	3
Ischemia/infarction	3
Cortical hemosiderosis	2
Hemorrhagic contusion	2
White matter disease	
Encephalomalacia	2
Postoperative/postradiation white matter changes	2
Nonspecific white matter disease	1
Demyelinating white matter disease	1
Delayed myelination	1
Others	
Congenital hydrocephalus	3
Arachnoid cyst	3
Tuberous sclerosis	2
Mesial temporal/hippocampal sclerosis	2
Atrophy	2
Mega cisterna magna	1
Porencephaly	1
Subdural hematoma	1
No abnormality seen	30

view, and 18 slice locations. The acquisition time was 8:56 minutes.

The FSE images were also obtained with a TR of 2000 msec and 2 excitations. In the early phase of the study, PDW and T2W images were obtained with pTEs of 30 and 75 msec in 24 and 10 patients, respectively. In the remaining subjects, pTEs of 15 and 90 msec were used for PDW and T2W images. Matrix size was either 256×128 (35 patients), 256×192 (41 patients), or 256×256 (five patients). Acquisition times were 2:08, 3:12, and 4:16 minutes, respectively. Slice thickness, interslice gap, and field of view were identical to those for the CSE sequences. However, because of the longer echo trains in FSE, only 14 slice locations were available using the TR of 2000 msec. FSE images were acquired in either the axial or coronal plane. Coronal images were often obtained to facilitate diagnosis or surgical planning.

The FSE images were compared with CSE images for lesion detectability and characterization (ie, number, size, and signal behavior) by a group of three neuroradiologists in an unblinded fashion. In addition to the visual comparison of signal characteristics, signal intensities of white and gray matter, cerebrospinal fluid (CSF), and fat, as well as the

background noise along the phase-encoding direction, were measured in six selected sets of FSE and CSE images utilizing region-of-interest (ROI) analyses by one of the authors. For each ROI, absolute signal intensity measurement, which is independent of window level or width settings, was obtained from ROIs containing between 100 and 300 pixels. The signal intensity measurements were made from studies in which both FSE and CSE images were obtained in the axial plane. Only the noise in the direction of the phase-encoding gradient was measured since it is greater in magnitude than the noise in the readout direction (7) and is considered a more valid measurement when "... assessing the impact of various techniques on image quality" (9). Contrast-to-noise ratios (CNR) were calculated for gray matter versus white matter, CSF versus white matter, CSF versus gray matter, and fat versus gray matter using the equation

$$\text{CNR} = (S_1 - S_2)/N_p \quad (\text{A})$$

where S_1 and S_2 are the mean signal intensities of the ROIs within the two tissue types being compared and N_p is the signal intensity from an ROI along the phase-encoding direction.

A phantom study was performed with a homogeneous water phantom in order to demonstrate the effect of matrix size on Gibbs artifact in FSE imaging.

Results

All the lesions detected on CSE images were seen on FSE images when pTEs of 15 and 90 msec were employed. Of the 24 patients whose FSE PDW images were obtained with pTEs of 30 msec, two findings (cystic ganglioglioma and mega cisterna magna) were inconspicuous on PDW images (although detected on the T2W images) because free water and CSF were nearly isointense to the adjacent gray matter (Fig. 2).

No additional lesions were demonstrated by FSE sequences. Measurements of the maximal dimensions of 10 randomly chosen lesions, ranging between 8 and 45 mm, revealed a strong correlation between FSE and CSE images (Fig. 3). FSE images yielded diagnostic information in two patients whose CSE images were degraded by motion artifact. In two agitated patients, one with cavernous hemangiomas and the other with tuberous sclerosis, diagnostically useful images were obtained with only FSE images in the axial plane (Fig. 4).

The mean signal intensities reported in Table 2 compare the FSE sequence of choice (TR 2000/pTE 15, 90; 2 excitations, 256×192 matrix) with the standard CSE brain examination used in our series (TR 2000/TE 30, 80; 2 excitations, 256×128 matrix). Mean signal intensi-

Fig. 2. Poor CSF/free water versus gray matter contrast in a FSE PDW image using a pTE of 30 msec; 7-year-old boy with a cystic ganglioglioma (scan time 3:12 min).

A, Axial CSE (2000/30) image demonstrates a low-intensity mass in the right posterior temporal lobe (scan time of 8:56 min).

B, Axial FSE (2000/pTE 30) image shows the lesion to be isointense and indistinguishable from adjacent gray matter, although the distinction was made on the T2W image (data not shown).

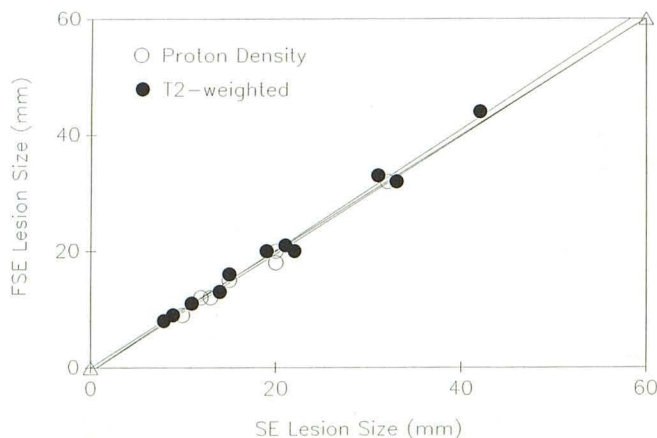
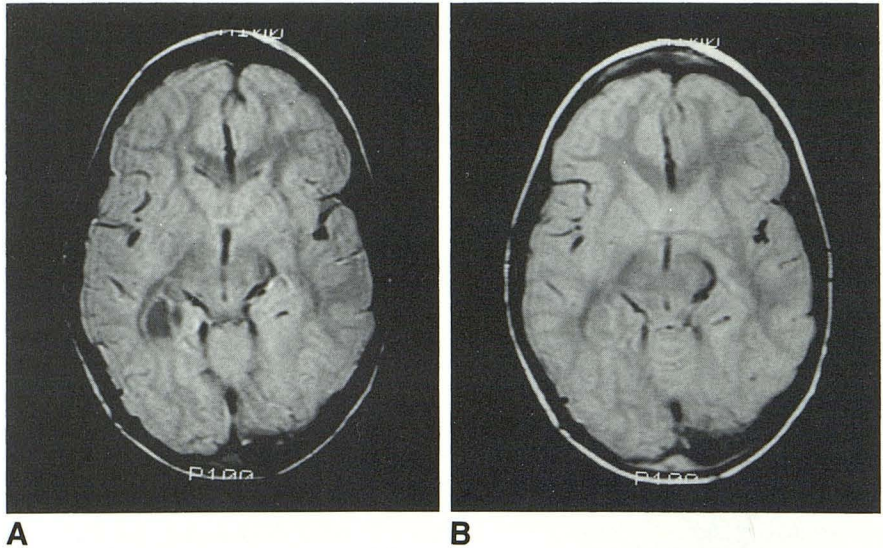


Fig. 3. CSE versus FSE correlation plot for lesion size. Open circles are measurements on PDW images; Closed circles are measurements on T2W images. The diagonal line closed by open triangles represents a perfect correlation, while the two other lines shown are linear fits for the PDW and T2W data. The correlation coefficients are all above 0.99

ties for white and gray matter and CSF were comparable to those in the CSE technique on the PDW images. However, in the T2 FSE images, signal intensities from these tissues were slightly reduced compared to their CSE counterparts. Scalp fat signal was much higher with FSE on both PDW and T2W sequences than with CSE (Table 2). The CNR characteristics of FSE compared favorably with those of the CSE sequence, though fat versus gray matter CNRs were significantly different (Table 3). Figure 5 presents exemplary FSE and CSE PDW and T2W images filmed using the same window level and width to illustrate the signal intensity behavior and the CNRs between the two techniques.

The phantom study demonstrated significant Gibbs artifact on the FSE image when 256×128

matrix was used. This artifact was reduced when matrix sizes of 256×192 or 256×256 were used (Fig. 6).

Discussion

The CSE technique has been the standard pulse sequence in neuroimaging since MR was introduced to diagnostic radiology. Newer MR pulse sequences that require shorter imaging times but retain time-tested spin-echo contrast would be desirable for clinical imaging.

FSE is a recently developed rapid MR pulse sequence that utilizes the Rapid Acquisition with Relaxation Enhancement (RARE) sequences described by Hennig and others (4–6), but modified with a specific phase encode-reordering algorithm (7).

Our study shows that FSE images can provide rapid dual-echo long TR brain images, with image quality matching that of CSE images. Lesion conspicuity and characterization on FSE images in 51 patients with intracranial abnormalities correlated well with the CSE sequence in our series.

FSE, in its current development, can shorten spin-echo acquisition times up to a factor of four compared to CSE images acquired with identical scan parameters by acquiring four phase-encode values for each TR. The time saving afforded by FSE can be utilized in several ways. One can keep FSE imaging parameters (TR, matrix, and number of excitations) the same as CSE and increase the speed of image acquisition. The speed of FSE can be traded off for increased spatial resolution by using larger image matrices, while still maintaining short imaging times. Alter-

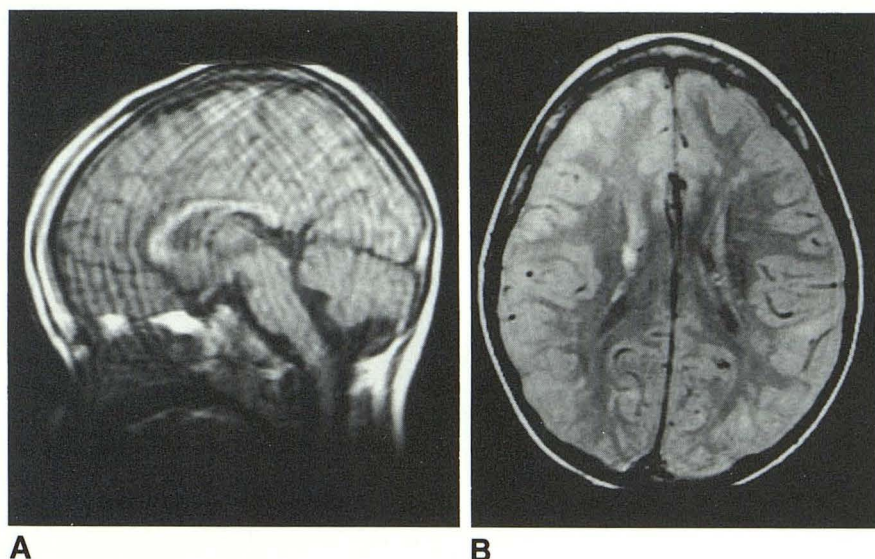


Fig. 4. Six-year-old nonsedated boy with tuberous sclerosis.

A, T1-weighted sagittal CSE image (600/15) is severely degraded by motion artifact (scan time 3:53 min).

B, FSE axial image (2000/15/2) shows minimal motion artifact allowing visualization of the periventricular hyperintense lesions (scan time 3:12 min). The patient cooperated better for the faster FSE acquisition. The longer CSE sequence would have most likely resulted in severe motion artifact.

TABLE 2: Mean signal intensities and standard deviations obtained from six patient studies performed in the axial plane using CSE and FSE

Pulse Sequence	GM	WM	Fat	CSF	N _p
PD-weighted					
CSE, TE 30	673 ± 33	586 ± 28	696 ± 33	553 ± 39	12 ± 1.7
FSE, pTE 15	672 ± 73	592 ± 65	1117 ± 72	507 ± 39	15 ± 4.7
T2-weighted					
CSE, TE 80	372 ± 30	275 ± 9	250 ± 19	507 ± 28	10 ± 2.7
FSE, pTE 90	329 ± 40	257 ± 33	546 ± 12	523 ± 38	14 ± 2.9

Note.—Measurements were made from axial brain images obtained with a TR of 2000 msec and 2 excitations. Matrix size for CSE images was 256 × 128 while FSE image matrix was 256 × 192. PD = proton density, GM = gray matter, WM = white matter, and N_p is the noise signal intensity in ROIs along the phase encoding direction.

TABLE 3: Contrast-to-noise ratios of conventional spin-echo and Fast Spin-echo images as obtained from the data in Table 2

Pulse Sequence	GM-WM	CSF-GM	CSF-WM	Fat-GM
	N _p	N _p	N _p	N _p
PD-weighted				
CSE, TE 30	7.3 (6.0)	-10.0 (-8.2)	-2.8 (-2.3)	1.9 (1.6)
FSE, pTE 15	5.3	-11.0	-5.7	29.7
T2-weighted				
CSE, TE 80	9.7 (7.9)	13.5 (11.0)	23.2 (18.9)	-12.2 (-11.4)
FSE, pTE 90	5.1	13.9	19.0	15.5

Note.—Numbers in parentheses are the CNRs anticipated if the CSE sequences had been acquired with the same matrix size (256 × 192) as the FSE sequences. The correction factor is (128/192)^{1/2} (see Ref. 9). For abbreviations, see Table 2.

natively, one may acquire additional imaging sequences or planes without significantly prolonging examination times. Nongap thin-slice imaging of the entire brain is also possible with FSE in less time than with CSE.

In general, FSE images obtained with pTEs of 15 and 90 msec produced tissue contrast similar

to that of the 30- and 80-msec CSE images, respectively (Table 3). Fat tissues remain bright on FSE T2W images as opposed to the signal loss seen with T2W CSE images (Table 2). This effect is believed to be due to the lengthening of T2 that occurs with short echo spacings, leading to decreased contributions from lipid protons to the T2 decay process due to spin-spin splittings (7, 10). Although no diagnostic problem was encountered in our series, difficulty may arise, for example, in differentiating extracellular methemoglobin from fat. Standard fat suppression schemes can be applied to FSE and the method even allows a novel fat-suppression method that can be implemented without decreasing the number of slices available for each TR (11). A pTE of 30 msec tended to produce PDW images with too much T2-weighting, resulting in unacceptably poor contrast between CSF and gray matter with the 2000-msec TR. This explains why two findings, mega cisterna magna and cystic gangli-

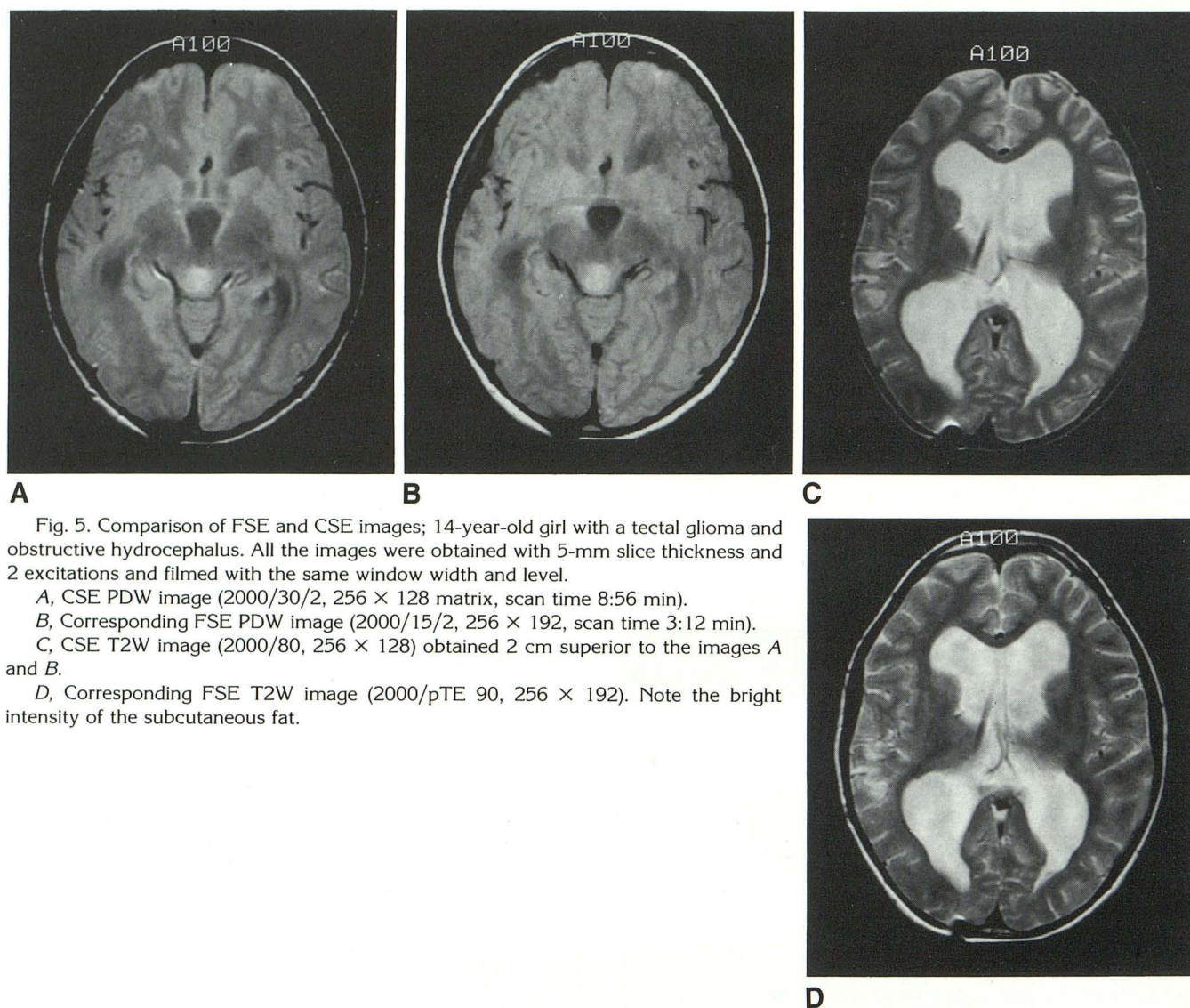


Fig. 5. Comparison of FSE and CSE images; 14-year-old girl with a tectal glioma and obstructive hydrocephalus. All the images were obtained with 5-mm slice thickness and 2 excitations and filmed with the same window width and level.

A, CSE PDW image (2000/30/2, 256 × 128 matrix, scan time 8:56 min).

B, Corresponding FSE PDW image (2000/15/2, 256 × 192, scan time 3:12 min).

C, CSE T2W image (2000/80, 256 × 128) obtained 2 cm superior to the images A and B.

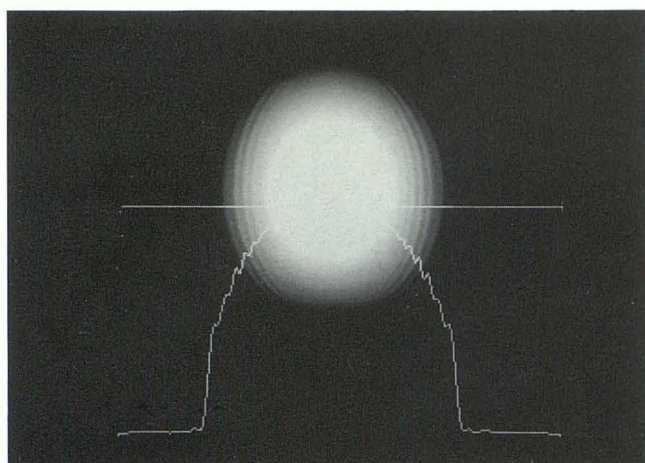
D, Corresponding FSE T2W image (2000/pTE 90, 256 × 192). Note the bright intensity of the subcutaneous fat.

oglioma, were not visualized on FSE with a pTE of 30 msec (Fig. 2), although both pathologies were well demonstrated on the late pTE FSE images. The main disadvantage of the 15-msec pTE FSE images is increased blurring of images compared to 30-msec pTE images. The blurring effect at the shorter pTE is due to using later, more T2-attenuated, echoes to obtain high-frequency phase-encoding information (4, 6, 7).

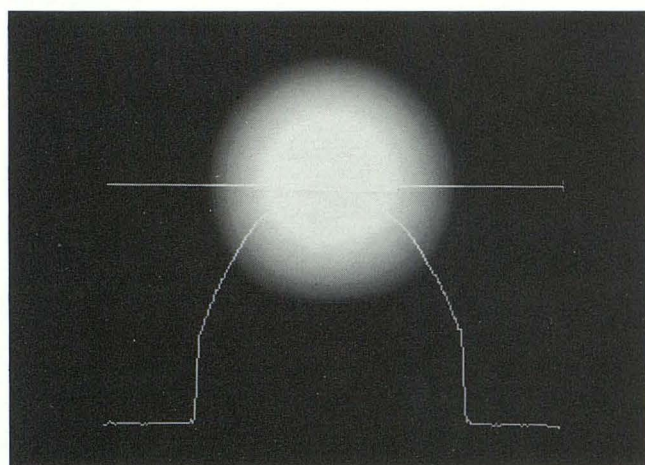
The Gibbs or truncation artifact is a well-known effect causing concentric or curvilinear lines adjacent to edges (12). This artifact is increased with FSE sequences and is related to T2 decay and concomitant discontinuities in the k-space trajectories encountered during a full FSE acquisition (7, 13). The authors preferred increasing matrix size from 256 × 128 to 256 × 192 for brain FSE imaging. This not only allowed better

spatial resolution but also reduced Gibbs artifact (Fig. 6). The dual-echo FSE acquisition time with a 256 × 192 matrix is roughly 1/3 the time required for the CSE sequence obtained with a 256 × 128 matrix (3:12 min vs 8:56 min).

For a given TR, fewer slices are possible with FSE than with CSE because of the longer echo trains used for FSE (120 msec vs 80 msec). In our series with a TR of 2000 msec, the FSE technique provided 14 slice locations as opposed to 18 with the CSE sequence. This limitation is rarely a problem in imaging of the infant or child brain, but may present a dilemma in imaging older children or adults by forcing a decision in favor of a fast scan time, but fewer slices. This problem can be overcome in several ways. The echo spacing in the FSE echo train may be shortened even further to allow more slices for a



A



B

Fig. 6. Phantom study; a homogeneous water phantom was imaged with the FSE sequence (2000/90/2) with a 256×128 (A) and a 256×192 matrix (B). The profile line at the bottom corresponds to signal intensities along the horizontal line through the center of the phantom parallel to phase-encoding axis. The Gibbs artifact is significantly reduced with a 256×192 matrix. There was further reduction of the Gibbs artifact when a matrix of 256×256 was used (data not shown).

given TR without increased scan time. For example, if the echo train is shortened to a 12-msec spacing as opposed to a 15-msec used in this study, the number of slices increases from 14 to 18. However, the decreased echo spacing requires smaller echo-acquisition windows, with subsequent increased bandwidth and decreased signal-to-noise ratio. Another method of obtaining more slice locations is to increase TR. A TR of 2500 msec will allow 18 slice locations with an eight-echo FSE. Even though this strategy increases the scan time by 25% over that obtained with a TR of 2000 msec, it further reduces the T1

influence, which may be a desirable feature. However, increased TR also results in decreased CNRs for CSF versus gray matter contrast on PDW images, which may pose a diagnostic problem. The FSE sequence with a TR of 2500 msec was not evaluated in this study. A third method for increasing the number of slices without increasing TR is to use a six-echo FSE sequence that utilizes the first three echoes for the PDW image and the last three for the T2W image. The number of available slices in a 2000-msec TR becomes 18, but the matrix size is restricted to 256×192 , and the scan time becomes 4:12 minutes (2000/pTE 15, 80/2 excitations). We are currently evaluating this sequence.

The use of many closely spaced refocusing pulses can also lead to an effective lengthening of T2 that reduces magnetic-susceptibility contributions to signal loss (7, 14). In fact, the reduced susceptibility effect was helpful in demonstrating a region of the brain distorted by a metallic artifact from dental braces, not an uncommon situation in pediatric patients (Fig. 7). Reduced susceptibility effects may also result in sharper delineation of the interfaces between air and soft tissues or bone and soft tissues on FSE images, although this was not evaluated in our study. The susceptibility signal loss associated with products of hemorrhage is reduced with FSE (Fig. 8), although not significantly (15).

The number of excitations and interslice gaps were arbitrarily chosen to be 2 and 50%, respectively, for FSE imaging in order to be comparable to the CSE image techniques employed at the authors' institution. These parameters can be varied in FSE sequence with similar changes in signal-to-noise ratios as in CSE sequence. Likewise, there is no limitation in choice of TR, pTE, slice thickness, or slice planes (including oblique) with FSE. While flow compensation techniques have been available for CSE, similar methods are not currently available for FSE. Although not a major problem at present, flow artifacts have been observed with FSE. We have not quantitatively evaluated their contribution to image degradation in this study.

Increased RF power deposition is a feature of FSE. In the brain MR studies performed with the CSE, estimated specific absorption rates (SARs) ranged from 0.01 to 0.1 W/kg. In general, the SAR values for FSE are increased by a factor of 3 to 4 as compared to CSE because of the increased number of 180° pulses per TR interval.

Fig. 7. FSE images displaying reduced magnetic-susceptibility effects as compared with CSE images.

A, CSE image (2000/80/2, scan time 8:56 min) demonstrates severe signal dropout and distortion from the dental brace.

B, FSE image (2000/90/2, scan time 3:12 min) acquired at the same slice location shows much less artifact with better visualization of the orbits and the temporal lobes.

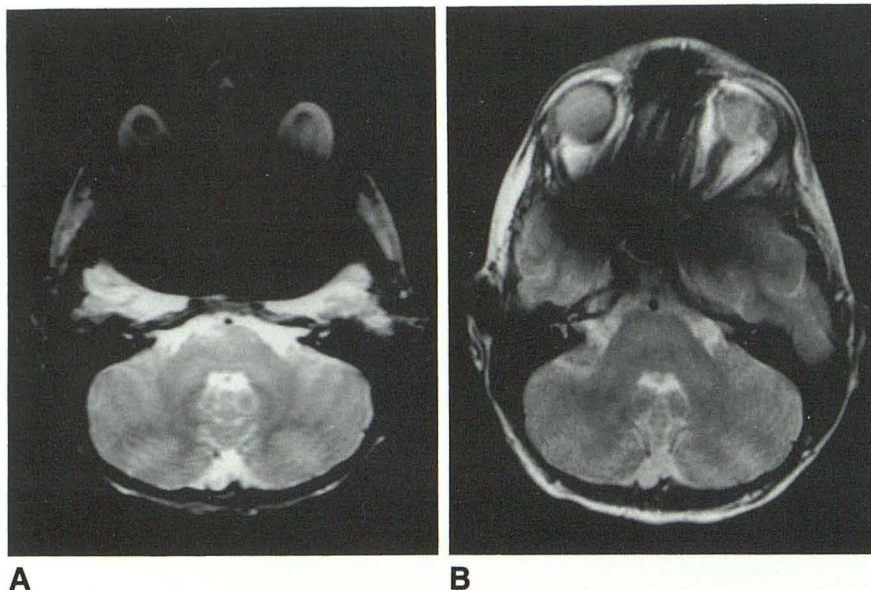
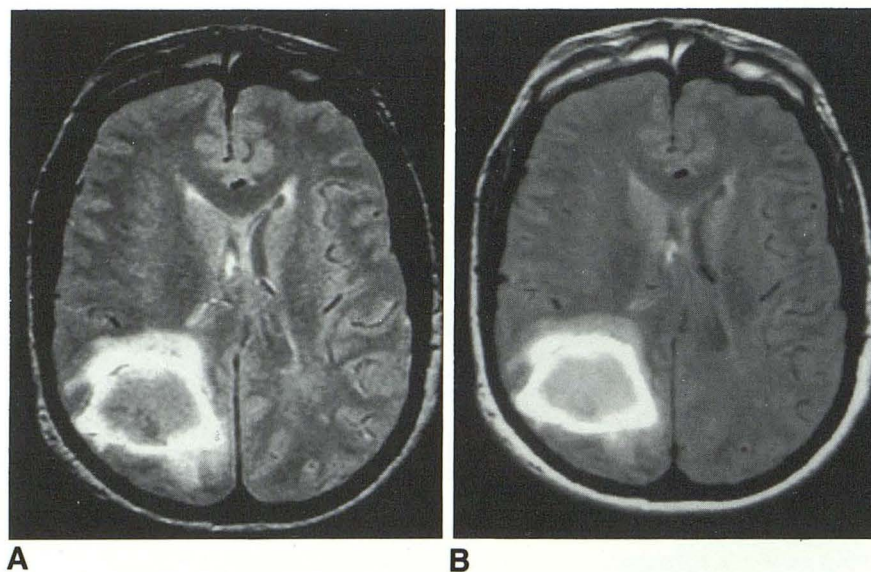


Fig. 8. Example of intracranial hemorrhage on CSE and FSE.

A, CSE PDW image (2000/30/2, scan time 8:56 min) demonstrating large right parieto-occipital junction hemorrhage. Note the susceptibility-induced signal loss from blood byproducts in the central region of the lesion.

B, FSE PDW image of the same slice location (2000/15/2, scan time 3:12 min) demonstrating slightly less signal loss in the same region.



This implies that for the head studies reported here, SAR values were under the 0.4 W/kg limit espoused by both the United States and Great Britain and well under the 1 W/kg limit suggested by the Federal Republic of Germany (16).

In conclusion, we have demonstrated that diagnostically reliable dual-echo long TR brain MR images can be obtained in a pediatric population using a newly developed FSE sequence in approximately $\frac{1}{4}$ to $\frac{1}{3}$ of the CSE acquisition time employing comparable TR and number of excitations without compromising lesion detectability. Pseudo-TEs of 15 and 90 msec appear to provide PDW and T2W signal contrasts very similar to those of CSE. FSE proved helpful in

reducing motion artifact, especially in marginally cooperative patients and patients of borderline age for sedation. Fat remains bright on T2W FSE images, but can be suppressed. A matrix size of 256×192 or larger reduces Gibbs artifact inherent in FSE to an acceptable level. Susceptibility effects were slightly reduced with FSE, but did not pose a diagnostic problem in our series.

References

1. Edelman RR, Kleefield J, Wentz KU, Atkinson DJ. Basic principles of magnetic resonance imaging. In: Edelman RR, Hesselink JR, eds. *Clinical magnetic resonance imaging*. Philadelphia: Saunders, 1990:23-25
2. Haake EM, Tkach JA. Fast MR imaging: techniques and applications.

- AJR 1990;155:951-964
3. Cohen MS, Weisskoff RM. Ultra-fast imaging. *Magn Reson Imaging* 1991;9:1-37
 4. Hennig J, Naureth A, Friedburg H. RARE imaging: a fast imaging method for clinical MR. *Magn Reson Med* 1986;3:823-833
 5. Henning J, Friedburg H. Clinical applications and methodological developments of the RARE technique. *Magn Reson Imaging* 1988;6:391-395
 6. Mulkern RV, Wong STS, Winalski C, Jolesz FA. Contrast manipulation and artifact assessment of 2D and 3D RARE sequences. *Magn Reson Imaging* 1990;8:557-566
 7. Melki PS, Mulkern RV, Panych LP, Jolesz FA. Comparing the FAISE method with conventional dual echo sequences. *J Magn Reson Imaging* 1991;1:319-326
 8. Twieg DB. The k-space trajectory formulation of the NMR imaging process with applications in analysis and synthesis of imaging methods. *Med Phys* 1983;10:610-621
 9. Kaufman L, Kramer DM, Crooks LE, Ortendahl DA. Measuring signal-to-noise ratios in MR imaging. *Radiology* 1989;173:265-267
 10. Allerhand A. Analysis of Carr-Purcell spin-echo NMR experiments on multiple-spin systems. I. The effect of homonuclear coupling. *J Chem Phys* 1966;44:1-9
 11. Higuchi N, Hiramatsu K, Mulkern RV. A novel "powerless" method for fat suppression in RARE sequences. Book of Abstracts. Annual Meeting of the Society of Magnetic Resonance in Medicine, 1991. San Francisco: SMRM
 12. Wood ML, Henkelman RM. Truncation artifacts in magnetic resonance imaging. *Magn Reson Med* 1985;2:517-526
 13. Mulkern RV, Melki PS, Jakab P, Higuchi N, Jolesz FA. Phase encode order and its effect on contrast and artifact in single-shot RARE sequences. *Med Phys* 1991;18:1032-1037
 14. Bendel P. Spin-echo attenuation by diffusion in non-uniform field gradients. *J Magn Reson* 1990;86:509-515
 15. Jones KM, Mulkern RV, Mantello MT, et al. Evaluation of brain hemorrhage: comparison of Fast Spin-echo and conventional dual spin-echo images. *Radiology* 1992;182:53-58
 16. Persson BR, Stahlberg F. *Health and safety of clinical NMR examinations*. 1st ed. Boca Raton, FL: CRC Press, 1989;108-112