Usefulness of the Delay Alternating with Nutation for Tailored Excitation Pulse with T1-Weighted Sampling Perfection with Application-Optimized Contrasts Using Different Flip Angle Evolution in the Detection of Cerebral Metastases: Comparison with MPRAGE Imaging

D. Kim, Y.J. Heo, H.W. Jeong, J.W. Baek, J.-Y. Han, J.Y. Lee, S.-C. Jin, and H.J. Baek

ABSTRACT

BACKGROUND AND PURPOSE: Contrast-enhanced T1-weighted sampling perfection with application-optimized contrasts by using different flip angle evolution (SPACE) with the delay alternating with nutation for tailored excitation (DANTE) pulse could suppress the blood flow signal and provide a higher contrast-to-noise ratio of enhancing lesion-to-brain parenchyma than the MPRAGE sequence. The purpose of our study was to evaluate the usefulness of SPACE with DANTE compared with MPRAGE for detecting brain metastases.

MATERIALS AND METHODS: Seventy-one patients who underwent contrast-enhanced SPACE with DANTE and MPRAGE sequences and who were suspected of having metastatic lesions were included. Two neuroradiologists determined the number of enhancing lesions, and diagnostic performance was evaluated using figure of merit, sensitivity, positive predictive value, interobserver agreement, and reading time. Contrast-to-noise ratiolesion/parenchyma and contrast-to-noise ratiowhite matter/grey matter were also assessed.

RESULTS: SPACE with DANTE (observer one, 328; observer two, 324) revealed significantly more small (<5 mm) enhancing lesions than MPRAGE (observer one, 175; observer two, 150) (P < 0.001 for observer 1, P = .0001 for observer 2). Furthermore, SPACE with DANTE showed significantly higher figure of merit and sensitivity and shorter reading time than MPRAGE for both observers. The mean contrast-to-noise ratiolesion/parenchyma of SPACE with DANTE (52.3 ± 43.1) was significantly higher than that of MPRAGE (17.5 ± 19.3) (P = .0001), but the mean contrast-to-noise ratiowhite matter/grey matter of SPACE with DANTE (−0.65 ± 1.39) was significantly lower than that of MPRAGE (3.08 ± 1.39) (P ≤ .0001).

CONCLUSIONS: Compared with MPRAGE, SPACE with DANTE significantly improves the detection of brain metastases.

ABBREVIATIONS: CE = contrast-enhanced; CNR = contrast-to-noise ratio; DANTE = delay alternating with nutation for tailored excitation; FOM = figure of merit; JAFROC = jackknife free-response receiver operating characteristic; SPACE = sampling perfection with application-optimized contrasts by using different flip angle evolution
All studies were performed using a 3T MR imaging scanner (Magnetom Skyra; Siemens, Erlangen, Germany) and a 64-channel head coil. After we obtained routine precontrast images using axial fluid-attenuated inversion recovery, T2-weighted, T1-weighted, gradient-echo, 3D SPACE, and 3D MPRAGE images with fat suppression were obtained immediately following intravenous administration of 0.1 mmol/kg body weight of Dotarem (gadoterate meglumine; Guerbet, Aulnay-sous-Bois, France). The scan parameters of postcontrast MR imaging were as follows for SPACE with DANTE imaging: TR, 800 ms; TE, 15 ms; flip angle, variable; fat suppression, Fat-Sat (chemical shift selective suppression); parallel acquisition techniques factor, 2; FOV, 230 × 230; bandwidth, 422 Hz/pixel; matrix, 320 × 320; number of slices, 240; echo spacing, 4.88 ms; voxel size, 0.72 × 0.72 × 0.72 mm; scan time, 5 minutes 54 seconds; DANTE preparation pulse were applied; for MPRAGE imaging: TR, 2200 ms; TE, 3.05 ms; flip angle, 9°; fat suppression, water excitation; parallel acquisition techniques factor, 2; FOV, 230 × 230; bandwidth, 260 Hz/pixel; matrix, 320 × 320; number of slices, 240; echo spacing, 8.8 ms; voxel size, 0.72 × 0.72 × 0.72 mm; flip angle, 9°; scan time, 5 minutes 32 seconds. We obtained sagittal planes covering the whole brain for the SPACE and MPRAGE imaging to reduce the scan time. We performed SPACE with DANTE and MPRAGE sequences in alternative order by random distribution to avoid timing bias, which can increase contrast agent uptake due to the delay after injection. The order of sequences was the following: SPACE with DANTE followed by MPRAGE in 32 patients and MPRAGE followed by SPACE with DANTE in 39 patients.

**Image Analysis**

**Determination of Metastatic Lesions.** Two neuroradiologists, one with 5 years of experience and one with 1 year of experience in neuroimaging, independently evaluated the presence of brain metastases using SPACE with DANTE and MPRAGE imaging with a 4-week interval to minimize any learning bias. One observer evaluated SPACE with DANTE followed by MPRAGE, and another observer evaluated MPRAGE followed by SPACE with DANTE. Both observers reported all enhancing lesions in the brain parenchyma, except for the normal anatomic structures or artifacts. The metastatic lesions were classified into 2 groups by lesion size: large (≥5 mm) and small (≤5 mm). Both observers also reported the level of confidence of metastatic lesions at each location on a rating scale (ranging from a lowest confidence level of 0 to a highest confidence level of 100), and the reading time of each case was recorded.

**Evaluation of Image Quality.** We evaluated and compared the CNR of lesions with normal parenchyma (CNRlesion/parenchyma) and the CNR of white matter with gray matter (CNRwhite matter/gray matter) among the SPACE with DANTE and MPRAGE images. For evaluation of CNRlesion/parenchyma we selected homogeneous, solid enhancing lesions of ≥5 mm and excluded rimlike enhancing lesions due to the difficulty in drawing the ROI. The CNR of enhancing lesions was calculated according to Kammer et al:

$$\text{CNR}_{\text{lesion/parenchyma}} = \frac{SI_{\text{lesion}} - SI_{\text{parenchyma}}}{SD_{\text{parenchyma}}}$$
We calculated the CNR for differentiating the gray and white matter as follows:

\[
\text{CNR}_{\text{white matter/gray matter}} = \frac{\text{SI}_{\text{white matter}} - \text{SI}_{\text{gray matter}}}{\text{SD}_{\text{white matter}}}
\]

Here, SI denotes the mean signal intensity of the ROI, and SD denotes the standard deviation of noise. To the extent possible, we endeavored to ensure identical size and location while drawing the ROI of each sequence, with side-by-side comparison of the two 3D-enhanced sequences and occasional use of zooming. For determining the SI and SD of the parenchyma, the ROIs were placed in the adjacent parenchyma because of inhomogeneous noise distribution in parallel imaging and included both white matter and gray matter. The ROIs of white matter were placed at the genu of the corpus callosum, and the ROIs of gray matter were placed at the head of the normal caudate nucleus. Every ROI of normal parenchyma, white matter, and gray matter measured \(22.73 \text{ mm}^2\). Every ROI of enhancing lesions was placed at the center of the lesion, to the extent possible, by 1 neuroradiologist (one with 5 years of experience), and the area of the ROI was dependent on lesion size, varying between 3.72 and 22.73 \(\text{ mm}^2\).

**Statistical Analysis**

All statistical analyses were performed using SAS, Version 9.4 (SAS Institute, Cary, North Carolina). The variables are presented as number or mean \(\pm\) SD. For evaluation of the diagnostic performance of each observer in each reading session, we used figure of merit (FOM) derived from the jackknife free-response receiver operating characteristic (JAFROC) analysis with method 1 of Chakraborty and Berbaum. A free software JAFROC analysis package is available at http://www.devchakraborty.com. The paired \(t\) test was used to compare the number of lesions, CNR, and reading time between SPACE with DANTE and MPRAGE. The sensitivity and positive predictive value of different MR images were calculated using a 2-way contingency table. \(P\) values \(< 0.05\) were considered statistically significant. Interobserver agreement for each MR image was calculated using \(k\) statistics; \(0–0.20, 0.21–0.40, 0.41–0.60, 0.61–0.80, \) and \(0.81–1.00\) were regarded as slight, fair, moderate, substantial, and almost perfect agreement, respectively, based on the Landis and Koch method.

**RESULTS**

**Diagnostic Performance of the MR Images**

SPACE with DANTE (observer one, 328; observer two, 324) revealed significantly more small (\(<5 \text{ mm}\)) enhancing lesions in the brain parenchyma than MPRAGE (observer one, 175; observer two, 150) \((P = 0.006 \text{ for observer 1, } P \leq 0.001 \text{ for observer 2})\) (Figs 1 and 2 and Table 1). In the detection of lesions of \(>5 \text{ mm}\), SPACE with DANTE (observer one, 186; observer two, 188) also revealed more enhancing lesions than MPRAGE (observer one, 168; observer two, 169), but this difference failed to reach statistical significance. Moreover, SPACE with DANTE showed higher sensitivity than MPRAGE for the detection of brain metastases, regardless of the lesion size (Fig 3). Both SPACE with DANTE and MPRAGE showed almost perfect interobserver agreement \((k = 0.99\) for SPACE with DANTE and 0.98 for MPRAGE for lesions smaller than 5 mm; \(k = 0.99\) for SPACE with DANTE and 0.98 for MPRAGE for lesions of \(>5 \text{ mm}\)) for the detection of brain metastases, regardless of lesion size. The FOM of SPACE with
DANTE was significantly higher than that of MPRAGE for both observers in the detection of lesions smaller than 5 mm \((P = .0017)\) (Table 2). However, the FOM was not significantly different between the 2 sequences in the detection of lesions of \(>5\) mm \((P = .1762)\).

SPACE with DANTE revealed more false-positive findings \((n = 18)\) due to incomplete vessel suppression \((n = 15)\) (Fig 4) and flow-related artifacts \((n = 3)\). On MPRAGE \((n = 10)\), the causes for the false-positive findings were vascular structures \((n = 9)\) and flow-related artifacts \((n = 1)\). However, these findings were not significantly different between the two 3D-enhanced MR images. The average reading time of SPACE with DANTE \((\text{observer one, } 45.4 \pm 31.7\text{ seconds}; \text{observer two, } 53.7 \pm 21.5\text{ seconds})\) was significantly shorter than that of MPRAGE for both observers \((\text{observer one, } 73.0 \pm 54.1\text{ seconds}; \text{observer two, } 72.0 \pm 22.8\text{ seconds})\) \((P \leq .0001)\).

**Evaluation of Image Quality**

A total of 51 patients who exhibited homogeneous, solid enhancing lesions of \(>5\) mm were evaluated. The mean CNR_{lesion/parenchyma} of SPACE with DANTE \((52.3 \pm 43.1)\) was significantly higher than that of MPRAGE \((17.5 \pm 19.3)\) \((P \leq .0001)\) (Table 3 and Fig 5). However, the mean CNR_{white matter/gray matter} of SPACE with DANTE \(\sim 0.65 \pm 1.39)\) was significantly lower than that of MPRAGE \((3.08 \pm 1.39)\) \((P \leq 0.0001)\).

**DISCUSSION**

We compared the diagnostic performance of CE 3D-SPACE with DANTE and MPRAGE for detecting brain metastases. The SPACE with DANTE sequences showed significantly higher sensitivity than the MPRAGE sequences, especially for smaller lesions \((<5\text{ mm})\), and they also showed higher interobserver agreement than the MPRAGE sequences. The SPACE with DANTE sequences showed improved arterial and venous blood suppression compared with SPACE alone.\(^7,13\) Moreover, DANTE can supplement this suppression using the gradient pulse in the phase-encoding direction.\(^13,19\) It also suppresses the signal due to slow blood flow, which is incompletely suppressed by SPACE.\(^13,19\) In addition, SPACE generates intravoxel dephasing and helps maintain the black-blood effect that DANTE cannot sustain during the readout period.\(^13\) These factors may simplify the reading process; thus, the reading time of SPACE

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**Table 1: Comparison of lesion detectability according to lesion size**

| Lesion diameter | SPACE with DANTE | MPRAGE | \(P\) Value |
|-----------------|------------------|--------|------------|
| \(\leq 5\) mm   |                  |        |            |
| Observer 1      | 0.904            | 0.698  |            |
| Observer 2      | 0.861            | 0.702  |            |
| Mean \(\pm SD\) | 0.882 \(\pm 0.023\) | 0.700 \(\pm 0.038\) | .0017     |
| \(>5\) mm       |                  |        |            |
| Observer 1      | 0.957            | 0.921  |            |
| Observer 2      | 0.943            | 0.928  |            |
| Mean \(\pm SD\) | 0.950 \(\pm 0.013\) | 0.925 \(\pm 0.014\) | .1762     |

\(^a\) Data for observers 1 and 2 are mean value of FOM data compared between SPACE with DANTE and MPRAGE.
with DANTE was significantly shorter than that of MPRAGE for both observers.

We measured the CNR of enhancing lesions for both SPACE with DANTE and MPRAGE images using the following formula:
\[
\text{CNR}_{\text{lesion/parenchyma}} = \frac{\text{SI}_{\text{lesion}} - \text{SI}_{\text{parenchyma}}}{\text{SD}_{\text{parenchyma}}}.
\]

Our study showed that the CNR of enhancing lesions to the normal parenchyma of SPACE with DANTE is higher than that of MPRAGE. This finding is in agreement with those of previous studies, which showed lower contrast enhancement of gradient-echo images than of spin-echo images and higher CNR of enhancing lesions using the spin-echo sequence than using the gradient-echo sequence. Previous studies have provided several reasons for the better detection of contrast-enhancing lesions using the spin-echo sequence than using the gradient-echo sequence. Previous studies suggested that the higher CNR of enhancing metastatic lesions to the normal parenchyma of the spin-echo sequence than of the gradient-echo sequence contributes to the higher detectability of metastatic lesions. Our result that the CNR of SPACE with DANTE is higher than that of MPRAGE is consistent with the findings of previous studies. The higher magnetization transfer effect of SPACE than MPRAGE has been suggested as another factor influencing its higher detection accuracy. A variable flip angle, which might be introduced as off-resonance pulses, induces magnetization transfer effects. It preferentially reduces the signal from the brain parenchyma, especially from white matter, rendering enhancing lesions more outstanding. However, SPACE alone is not sufficient for the evaluation of enhancing lesions because residual blood signal can occasionally be mistaken for enhancing lesions.

Therefore, black-blood modules have been used for evaluation of brain metastases. A few studies have evaluated the detectability of brain metastasis using 3D CE MR imaging with the black-blood module, but no study has used the DANTE preparation pulse in the evaluation of brain metastasis. A previous study suggested that DANTE preparation is a promising black-blood module that offers a higher signal-to-noise ratio and allows a shorter acquisition time than other types of black-blood modules, such as double inversion recovery or motion-sensitive driven equilibrium preparation modules. Our findings are in agreement with those of Park et al., who detected significantly more small lesions using the CE 3D black-blood single slab turbo spin-echo sequence than using the CE MPRAGE sequence. They found no significant differences in the detection of larger enhancing lesions (>5 mm), and this finding is in agreement with that of our study.

We could recognize the enhancing lesions more easily using SPACE with DANTE than by using MPRAGE, and the result of our study that SPACE with DANTE required a shorter reading time than MPRAGE for both observers supports this finding. This finding is consistent with those of previous studies that analyzed the reading time.

The CNR of enhancing lesions using SPACE with DANTE (8.62) was higher than that of MPRAGE (3.49).

| CNR (n = 5) | SPACE with DANTE | MPRAGE | P Value |
|------------|------------------|--------|---------|
| Lesion/parenchyma | 52.3 ± 43.1 | 17.5 ± 19.3 | <.0001 |
| White matter/gray matter | -0.65 ± 139 | 3.08 ± 139 | <.0001 |

*Data are presented as means. Values were calculated using paired t tests.
suppression of the blood flow. This finding is consistent with a previous study\textsuperscript{13} reported that CE-SPACE showed more false-positive findings than MPRAGE. The variable flip angle of the SPACE sequence imperfectly suppresses the vascular signal and could lead to misinterpretation of the remaining vessels as a metastatic lesion. Furthermore, turbo spin-echo motion-sensitive driven equilibrium\textsuperscript{11} shows more false-positive lesions than the non-motion-sensitive driven equilibrium sequence due to incomplete blood flow suppression of small peripheral vessels, and these findings hamper interpretation. However, we could easily recognize these structures as false-positive lesions by considering multiplanar reconstruction and MPRAGE images. This method has already been confirmed by a previous study,\textsuperscript{13} which decreased the false-positive rate and preserved diagnostic performance. Kato et al\textsuperscript{8} also reported several false-positive events, but the causes were different from those identified in our study. None of the venous sinuses or choroid plexuses were misdiagnosed as metastasis in our study.

Our study has several limitations. First, this study was retrospective in design. Second, pathologic confirmation of all metastatic lesions was not possible because patients with multiple brain metastases usually do not undergo an operation. Third, we could not include SPACE without a DANTE pulse due to its limited acquisition time. However, a previous study\textsuperscript{13} has already shown the improved suppression of arterial and venous blood using SPACE with DANTE compared with SPACE. Another study\textsuperscript{13} using turbo spin-echo motion-sensitive driven equilibrium showed that it achieves better blood vessel suppression than non-turbo spin-echo motion-sensitive driven equilibrium, with a similar CNR. Nevertheless, further studies comparing SPACE without DANTE and SPACE with DANTE are needed and may support our results. Finally, in the present study, the observers were not blinded to the type of MR images because the differentiation of gray and white matter was evidently different between the 2 sequences.

CONCLUSIONS
Using SPACE with DANTE could improve the diagnostic performance for brain metastases; this approach also has almost perfect interobserver agreement. Compared with MPRAGE, SPACE with DANTE significantly improves the detection of brain metastases, particularly of those of $<$5 mm, without significantly increasing the false-positive rate. This information should be considered in the development of optimal brain tumor imaging protocols.

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