Brain Tumor-Enhancement Visualization and Morphometric Assessment: A Comparison of MPRAGE, SPACE, and VIBE MRI Techniques

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ABSTRACT

BACKGROUND AND PURPOSE: Postgadolinium MR imaging is crucial for brain tumor diagnosis and morphometric assessment. We compared brain tumor enhancement visualization and the “target” object morphometry obtained with the most commonly used 3D MR imaging technique, MPRAGE, with 2 other routinely available techniques: sampling perfection with application-optimized contrasts by using different flip angle evolutions (SPACE) and volumetric interpolated brain examination (VIBE).

MATERIALS AND METHODS: Fifty-four contrast-enhancing tumors (38 gliomas and 16 metastases) were assessed using MPRAGE, VIBE, and SPACE techniques randomly acquired after gadolinium-based contrast agent administration on a 3T scanner. Enhancement conspicuity was assessed quantitatively by calculating the contrast rate and contrast-to-noise ratio, and qualitatively, by consensus visual comparative ratings. The total enhancing tumor volume and between-sequence discrepancy in the margin delineation were assessed on the corresponding 3D target objects contoured with a computer-assisted software for neuronavigation. The Wilcoxon signed rank and Pearson χ² nonparametric tests were used to investigate between-sequence discrepancies in the contrast rate, contrast-to-noise ratio, visual conspicuity ratings, tumor volume, and margin delineation estimates. Differences were also tested for 1D (Response Evaluation Criteria in Solid Tumors) and 2D (Response Assessment in Neuro-Oncology) measurements.

RESULTS: Compared with MPRAGE, both SPACE and VIBE obtained higher contrast rate, contrast-to-noise ratio, and visual conspicuity ratings in both gliomas and metastases (P range, <.001–.007). The between-sequence 3D target object margin discrepancy ranged between 1% and 19.9% of lesion tumor volume. Larger tumor volumes, 1D and 2D measurements were obtained with SPACE (P range, <.01–.007).

CONCLUSIONS: Superior conspicuity for brain tumor enhancement can be achieved using SPACE and VIBE techniques, compared with MPRAGE. Discrepancies were also detected when assessing target object size and morphology, with SPACE providing more accurate estimates.

ABBREVIATIONS: CEL = contrast-enhancing lesion; GBCA = gadolinium-based contrast agent; RANO = Response Assessment in Neuro-Oncology; RECIST = Response Evaluation Criteria in Solid Tumors; SPACE = sampling perfection with application-optimized contrasts using different flip angle evolution; TV = tumor volume; VIBE = volumetric interpolated brain examination

Postcontrast MR imaging plays a fundamental role in brain tumor diagnosis, anatomic delineation, and treatment re-

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due to the inversion recovery preparation pulse. It was recommended for use in brain tumor clinical trials, in both the “Consensus Recommendations for a Standardized Brain Tumor Imaging Protocol in Clinical Trials” and in the modified Response Assessment in Neuro-Oncology (RANO) criteria. However, MPRAGE images exhibit suboptimal postgadolinium enhancement visualization, particularly in lesions with low gadolinium concentration. On the other hand, in previous investigations, 2 alternative techniques available for clinical use, the sampling perfection with application-optimized contrasts using different flip-angle evolution (SPACE sequence; Siemens, Erlangen, Germany) and the volumetric interpolated brain examination (VIBE) demonstrated a higher sensitivity to enhancing brain lesions.

MR Imaging Sequences

For MPRAGE and SPACE, we adapted the sequence default parameters recommended by the vendor, including a sagittally oriented slab with a symmetric k-space acquisition in all encoding directions, resulting in 1-mm³ isotropic voxels. For VIBE, which uses an asymmetric k-space acquisition, we obtained sagittally oriented slabs with a 1 × 1 × 1.27 mm actual voxel size, resulting into a 1-mm isotropic voxel after zero-filling interpolation. To minimize the potential bias related to the time gap between the first and last acquired images after injection, we reduced the scan duration using parallel imaging. As previously proposed, a relatively higher acceleration factor was used for SPACE (X4), to account for the longer acquisition time. Finally, because patients had to be investigated for metastases potentially occurring in nonbrain tissues, fat suppression was also applied to improve contrast enhancement at this level. Scan durations were 1 minute 57 seconds for VIBE, 3 minutes 10 seconds for SPACE, and 3 minutes 46 seconds for MPRAGE. A summary of the main parameters of the sequences is reported in On-line Table 1.

All patients underwent our standard brain tumor MR imaging protocol, including FLAIR, DWI, T2*, T2 TSE, and PWI. For the precontrast T1-weighted images, 1 sequence was randomly chosen among MPRAGE, SPACE, and VIBE. A constant dose of 0.1 mL/Kg of gadobutrol, followed by a saline flush, was administered IV in all examinations. Finally, MPRAGE, SPACE, and VIBE sequences were consecutively performed in a randomized order, starting 5 minutes after contrast injection.

Image Analysis

Quantitative Conspicuity. Two measures, the contrast rate and contrast-to-noise ratio, were calculated with previously used formulas:

Contrast Rate = \[(S_{\text{lesion}} - S_{\text{parenchyma}})/S_{\text{parenchyma}}\] × 100

Contrast-to-Noise Ratio = (S_{\text{lesion}} - S_{\text{parenchyma}})/SD_{\text{parenchyma}}

Where S_{\text{lesion}} and S_{\text{parenchyma}} respectively, represent the CEL and the adjacent parenchyma average signal intensity, and SD_{\text{parenchyma}} represents the standard deviation in S_{\text{parenchyma}}

MATERIALS AND METHODS

This study was approved by the local ethics committee of Canton Ticino after obtaining written informed consent from all patients.

Between April 2015 and May 2017, one hundred ninety-six consecutive adult patients undergoing brain MR imaging for suspected or known tumor or tumor relapse following an operation were prospectively enrolled and examined on a 3T scanner (Magnetom Skyra; Siemens) with a 20-channel head coil. Inclusion criteria were the following: enhancing intra-axial tumors that were candidates for neuronavigation-assisted biopsy, surgical removal, and/or stereotactic radiosurgery treatment based on the literature recommendations and multidisciplinary tumor board consensus. Forty-one patients with intra-axial tumors were reviewed; of these, 3 were excluded for movement artifacts, and 1 for incomplete protocol acquisition. In patients with multiple discrete foci of enhancement separated by nonenhancing parenchyma, these were considered individual lesions. Three additional metastases, which were not detected on MPRAGE (see also “Visual Conspicuity” below), were not included in the subsequent analyses. Fifty-four CELs (38 gliomas in 29 patients, and 16 metastases in 8 patients) were finally included. In 38/38 gliomas and 10/16 metastases, the final diagnosis was based on pathology. The diagnoses for the remaining lesions were based collectively on imaging characteristics, clinical history, evidence of remote (non-brain) malignancy, and MR imaging follow-up. In patients with tumor relapse (16 gliomas), differentiating tumor from treatment-related enhancement was based on perfusion and diffusion imaging, morphology (nodular/expansile versus linear/nonspecific), and sequential MR imaging follow-up.

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previously suggested,\textsuperscript{17,31} because parallel imaging was used, SI and SD for noise calculation were taken in the adjacent parenchyma, instead of background air. By means of iPlan, Version 3.0 (Brainlab, Munich, Germany), ROIs were drawn by an expert neuroradiologist (L.D., with 7 years of experience in neuromaging). For \textit{S\textsubscript{lesion}}, ROIs were drawn distant from lesion edges to avoid partial volume effects (average ROI size = 11.88 mm\textsuperscript{2}). For \textit{S\textsubscript{parenchyma}}, a relatively larger ROI (average size = 74.71 mm\textsuperscript{2}) was placed in the adjacent parenchyma, including the gray and white matter to account for signal intensity differences among the tissues,\textsuperscript{13} and avoiding CSF and vessels.

\textbf{Visual Conspicuity.} As previously noted,\textsuperscript{13,17} it is not possible to perform a visual evaluation blinded to the sequence type because expert readers would easily recognize the distinguishing features of the various sequences. Here, 1 neuroradiologist (E.P.) and 1 neurosurgeon (M.R.) with 10 and 15 years of experience, respectively, were unaware of the clinical data and quantitative results, assessed all 3 sequence types in each patient. The 3 sequences were presented sequentially for each patient, in a random order, using the Brainlab DICOM viewer. They provided a consensus visual score, aimed at comparing the quality of CEL conspicuity among sequences, in a competitive manner and ranking (“best,” “intermediate,” and “worst”), taking into account the following: 1) enhancement conspicuity, 2) clarity of CEL separation from adjacent vessels, and 3) impact of artifacts from paramagnetic deposits from hemorrhage/calcifications. When the overall visual conspicuity of a given lesion was assessed as similar between 2 or among all 3 sequences, the same ranking was assigned to each. Finally, during a separate session, the number of small metastases visible on SPACE and/or VIBE, but not on MPRAGE images, was retrospectively recorded.

\textbf{Volume and Margin Delineation.} Analyses were performed using a semiautomated tool based on a region-growing algorithm (SmartBrush 2.5; Brainlab). This is previously validated software that demonstrated excellent reliability for brain tumor segmentation.\textsuperscript{33,34} All segmentations were performed by a single neuroradiologist (L.D.) to avoid interrater bias.\textsuperscript{34} After image coregistration among all sequences, they were presented independently to the reader. Region-growing 2D segmentations of the CELs were drawn on perpendicular slices and automatically 3D-interpolated by the software. Then, margin refinements were manually performed with the help of the region-growing algorithm.\textsuperscript{33}

Within-sequence reproducibility was performed by repeating segmentations in a randomly chosen subset of 27 (50%) CELs during a subsequent separate session and by calculating the corresponding intraclass correlation coefficients with a 2-way mixed consistency, average-measured approach.\textsuperscript{33,35} Consistent with previous validation studies,\textsuperscript{33} intraclass correlation coefficient values were excellent: 0.998 for MPRAGE, 95% CI, 0.995–0.999; 0.997 for VIBE, 95% CI, 0.993–0.999; and 0.999, 95% CI, 0.997–0.999 for SPACE.

The potential discrepancy occurring between the objects contoured using the different techniques was further assessed concerning lesion margin spatial mismatch. Using the Brainlab object manipulation tool, objects were reciprocally subtracted: 1) \(|\text{CEL\textsubscript{MPRAGE}}| − |\text{CEL\textsubscript{SPACE}}|; 2) |\text{CEL\textsubscript{SPACE}}| − |\text{CEL\textsubscript{MPRAGE}}|; 3) |\text{CEL\textsubscript{MPRAGE}}| − |\text{CEL\textsubscript{VIBE}}|; and 4) |\text{CEL\textsubscript{VIBE}}| − |\text{CEL\textsubscript{MPRAGE}}|. We defined the resulting differences as margin extent discrepancy. This quantified the mismatch volume of the regions where a given sequence exceeded another and vice versa (see the explanatory diagram in Fig 1). The amount of mismatch with respect to the entire CEL was assessed by calculating the ratio (expressed in percentages) between the margin extent discrepancy and the median MPRAGE TV as a reference.

\textbf{ID and 2D Measurements.} In a subsequent session, 3 expert neuroradiologists (L.D., D.D., E.P.), with 7, 8, and 10 years of experience in brain tumor imaging, respectively, each independently measured the longest CEL diameter (1D RECIST\textsuperscript{25}) and the product of the 2 longest perpendicular diameters (2D RANO\textsuperscript{9}), excluding cystic cavities and lesions with poorly defined margins. Only lesions of diameter ≥10 mm were considered measurable. Twenty-nine CELs (22 gliomas, 7 metastases) were evaluated and randomly presented to the readers, while varying sequences and patients. Finally, for each lesion, the 3 readers’ measurements were averaged to perform sequence comparisons.

\textbf{Statistical Analysis.} All computations were performed using the SPSS software package, Version 22.0 (IBM, Armonk, New York). Normal distribution assumptions were checked by Kolmogorov-Smirnov and Shapiro-Wilk tests. Descriptive analyses for quantitative variables used median and interquartile range, while frequencies were used for the qualitative measures. In all CELs, as well as gliomas and metastases separately, differences between pairs of sequences in the contrast rate, contrast-to-noise ratio, and TV were investigated using the nonparametric Wilcoxon test. The visually estimated conspicuity ranking scores among sequences were compared using the \(\chi^2\) test. Bland-Altman diagrams were performed to represent the agreement of TV measures between sequences, and the corresponding repeatability coefficients were calculated.\textsuperscript{36,37} As for 1D and 2D measurements, given the limited sample size, analyses were performed in all CELs only. Interreader reproducibility of 1D and 2D measurements was estimated using the intraclass correlation coefficient.

The FSL General Linear Model (http://fsl.fmrib.ox.ac.uk/fsl/fslwiki/GLM) for repeated measures was used to adjust for the potential confound effect of the acquisition order between compared sequences, with respect to the contrast rate, contrast-to-noise ratio, TV, and 1D and 2D differences on Poisson distribution. The General Linear Model was also applied to test the potential confound of postsurgical and posttreatment tissue changes, which can influence the enhancement intensity at different CEL sites.\textsuperscript{38} This potential confounding factor was addressed through testing the interaction effect of recurrence (as a binomial variable) on TV differences with the General Linear Model. The level of significance was set at \(P < .05\).

\textbf{RESULTS} Of 57 CELs detected by SPACE, 3 small (≤2-mm diameter) metastases (5.3% of all lesions, 15.8% of all metastases) were missed.
on MPRAGE. The remaining 54 CELs were included in the subsequent analyses. Patients’ main demographics and CEL characteristics are presented on On-line Table 2.

Conspicuity Assessment

Both the contrast rate and contrast-to-noise ratio were significantly higher on either SPACE and VIBE images compared with MPRAGE, in all groups. Differences remained significant after adjusting for the acquisition order of the compared sequences (Table 1). For visual assessment, in all CELs, MPRAGE obtained the best ranking in 15/54 (27.8%), intermediate in 26/54 (48.1%), and worst in 13/54 (24.1%). SPACE provided the best CEL visualization in all cases (54/54, 100%); therefore, no statistical comparison was performed with MPRAGE. VIBE obtained a higher frequency of best (24/54, 44.4%) and a lower frequency of worst
ratings (5/54, 9.3%) compared with MPRAGE (P < .001), whereas the frequency of intermediate ratings was not significantly different (25/54, 46.3%). Similar results were obtained by VIBE in the glioma subgroup (P = .001), while no significant differences were found in metastases subgroup. Two illustrative cases, with an example of a tiny metastasis not visible on MPRAGE images, are presented in Fig 2.

**Volume and Margin Delineation Assessment**

Between-sequence TV and margin extent discrepancy estimations are summarized in Table 2. As for the TV, significantly discrepant measurements were obtained between MPRAGE and SPACE in all groups (P range, .001–.007), with SPACE generating larger volumes. The corresponding measure agreement distribution is illustrated by the Bland-Altman plots in On-line Fig 1. The repeatability coefficient was 2.03 cm³. After adjusted for the compared sequence-acquisition order, differences remained significant in all CELs and gliomas and approached significance for metastases (P = .075) (Table 2). No significant interaction effect of recurrence (previously treated lesions) was detected (P = 0.958, P = .332). No discrepancy was found between MPRAGE and VIBE in all groups (P range, .201–.343).

The margin extent discrepancy analysis results are reported in Table 2. As for the |CEL_{MPRAGE} − CEL_{SPACE}| comparison (quantifying the areas where MPRAGE exceeded SPACE margins), the margin extent discrepancy represented 7.4% (all lesions), 6% (gliomas), and 6.1% (metastases) of the TV, whereas for the reciprocal |CEL_{SPACE} − CEL_{MPRAGE}| subtraction (quantifying the areas where SPACE exceeded MPRAGE margins), the margin extent discrepancy was 19.9% (all lesions), 14.2% (gliomas), and 15.2% (metastases). Regarding the |CEL_{MPRAGE} − CEL_{VIBE}| mismatch, the margin extent discrepancy was 11% (all lesions), 7% (gliomas), and 3% (metastases) of the TV, whereas for the |CEL_{VIBE} − CEL_{MPRAGE}| mismatch, it was 11% (all lesions), 7.6% (gliomas), and

![Image of two sample cases showing differences in contrast-enhancing lesion conspicuity between sequences.](image-url)

**Table 2: Morphometric assessment**

<table>
<thead>
<tr>
<th></th>
<th>All CELs (n = 54)</th>
<th>Gliomas (n = 38)</th>
<th>Metastases (n = 16)</th>
</tr>
</thead>
<tbody>
<tr>
<td>TV (median/IQR) (mL)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>MPRAGE</td>
<td>1.36/0.18–10.93</td>
<td>3.02/0.44–28.93</td>
<td>0.33/0.006–2.81</td>
</tr>
<tr>
<td>SPACE</td>
<td>1.78/0.20–11.00</td>
<td>3.5/0.49–29.73</td>
<td>0.39/0.006–3.2</td>
</tr>
<tr>
<td>VIBE</td>
<td>1.62/0.16–10.35</td>
<td>3.39/0.43–27.48</td>
<td>0.40/0.007–2.84</td>
</tr>
<tr>
<td>TV, SPACE vs MPRAGE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P</td>
<td>.001b</td>
<td>.007b</td>
<td>.003b</td>
</tr>
<tr>
<td>(adjusted for sequence acquisition order)c</td>
<td>.034b</td>
<td>.033b</td>
<td>.075</td>
</tr>
<tr>
<td>TV, VIBE vs MPRAGE</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>P</td>
<td>.259</td>
<td>.201</td>
<td>.343</td>
</tr>
<tr>
<td>(adjusted for sequence acquisition order)c</td>
<td>.521</td>
<td>.538</td>
<td>.706</td>
</tr>
</tbody>
</table>

**Note:** MED indicates margin extent discrepancy; IQR, interquartile range.

a Wilcoxon signed rank test.
b Statistically significant differences.
c General Linear Model.
15.2% (metastases). Segmentation examples are provided in Fig 3 and in On-line Fig 2, highlighting sources of mismatch related to differences in the enhancement sensitivity, erroneous margin estimation, and vessel segmentation, as well as susceptibility artifacts.

**1D and 2D Assessment**

The interobserver reproducibility, investigated with intraclass correlation coefficient single measurements, showed good agreement among the 3 readers for both 1D (MPRAGE: intraclass correlation coefficient = 0.69, 95% CI, 0.56–0.79; VIBE: intraclass correlation coefficient = 0.73, 95% CI, 0.61–0.82; and SPACE: intraclass correlation coefficient = 0.74, 95% CI, 0.63–0.83), and 2D (MPRAGE: intraclass correlation coefficient = 0.74, 95% CI, 0.63–0.83; VIBE: intraclass correlation coefficient = 0.81, 95% CI, 0.72–0.88; and SPACE: intraclass correlation coefficient = 0.81, 95% CI, 0.72–0.88). At the between-sequence comparison (On-line Table 3), both measurements were significantly discrepant between MPRAGE and SPACE ($P < .001$ and .001, respectively). Differences were also significant after adjusting for the acquisition order of compared sequences (On-line Table 3). No significant discrepancies with VIBE images were found.

**DISCUSSION**

Contrast enhancement refers to hyperintensity on T1-weighted images following GBCA administration, which is the result of extravasation of gadolinium-containing macromolecules through a leaky BBB. GBCA accumulates within tumor interstitium, resulting in longitudinal relaxation time shortening and consequent T1 signal hyperintensity. Contrast enhancement helps CEL detection, characterization, and pretreatment anatomic assessment and represents a biomarker of potential high-grade tumor within malignant gliomas. A number of factors are known to influence the CEL conspicuity with respect to the background parenchyma: the BBB ultrastructural characteristics, magnetic field strength, GBCA concentration, relaxivity properties, time from injection, and MR imaging technique. Here, we quantitatively and qualitatively show that tumor-enhancement visualization with SPACE and VIBE is superior to that in MPRAGE, the currently recommended standard technique for brain tumor clinical trials. In this study, SPACE and VIBE demonstrated significantly superior visual conspicuity ratings across all the CEL assessments, with SPACE having the best enhancement visualization in 100% of cases and VIBE in 44%, compared with 27.8% for MPRAGE. Conversely, MPRAGE received the worst ranking among sequences in 24.1% of all CELs, compared with only 9.3% for VIBE. On image inspection, we found that subtly enhancing parts of the tumors detected by SPACE and VIBE may be crucially missed by MPRAGE. Accordingly, 3 very small metastases could be detected only by SPACE. These results confirm and extend previous investigations conducted on metastatic lesions. Kato et al and Komada et al documented higher contrast rate and contrast-to-noise ratio estimated by SPACE. In addition, a retrospective study by Kammer et al showed that the use of SPACE allowed increased sensitivity and diagnostic confidence in the detection of small lesions. Previous information about VIBE performance was remarkably scarce: only 1 previous study conducted at 1.5T by Wetzel et al reported higher contrast-to-noise ratio values and visual conspicuity in a heterogeneous group of brain tumors, compared with MPRAGE.

To the best of our knowledge, this is the first study providing information about morphometric discrepancies occurring with brain tumor segmentation, related to sequence-enhancement conspicuity differences. First, we found that discrepant and significantly larger TV estimates were obtained using SPACE com-
pared with MPRAGE, both in all lesions and in gliomas and metastases considered separately. However, the use of VIBE images did not lead to significantly different TV estimates with respect to MPRAGE. Second, consistent with the larger TV estimates, SPACE contours exceeded those created with MPRAGE by 12.8% of the TV in all CELs. Most important, despite providing smaller TV estimates, MPRAGE margins exceeded SPACE margins by 6.2% of the TV, as well. Factors contributing to such a spatial delineation discrepancy included the different techniques sensitive to subtle enhancement, overestimation of tumor borders, erroneous vessel contouring, and/or interference from hemorrhagic/calcium deposit–related susceptibility artifacts (Fig 3 and Online Figs 2 and 3). Although an in vivo reference standard is not available to establish the CEL TV and morphology, the fact that SPACE images obtained higher contrast rate and contrast-to-noise ratio and visual conspicuity ratings from expert readers reasonably supports the higher accuracy of this technique compared with MPRAGE for CEL boundary delineation. Furthermore, because high-flow intracranial arteries or venous sinuses appear dark on TSE images due to the flow void effect, they could be more accurately separated from the adjacent CEL than on fast gradient recalled-echo images, which conversely display enhancing vessels due to the flow-related enhancement effect.

Our findings have several practical implications. First, the use of SPACE and/or VIBE sequences may facilitate the identification of areas with relatively higher grade malignancy within gliomas. This is important for guiding accurate biopsy sampling and consequent correct histologic grading and treatment decision-making. Second, because achieving maximal tumor resection may lead to more favorable patient outcomes and the extent of resection may be increased with image-guided approaches, optimizing target delineation accuracy facilitates precision surgery in individual patients. In the setting of stereotactic radiosurgery planning, using low-contrastibility images could also hinder target-volume margin delineation and result in suboptimal dose delivery to all CEL sites and/or in unnecessary dose delivery to surrounding eloquent brain structures. An exemplary case, demonstrating MPRAGE underestimation of tumor margins adjacent to the optic radiation, is presented in Fig 3.

Third, accurate count of brain metastases for radiosurgery planning is important to decide between stereotactic radiosurgery and whole-brain treatment approaches. This importance is underscored by a previous study that emphasized the use of double-dose GBCA to improve MR imaging lesion-detection sensitivity for treatment planning. In contrast, maximizing image sensitivity to low-GBCA concentrations may play a crucial role whenever dose reduction is deemed appropriate to limit a patient’s gadolinium tissue deposition.

Fourth, the varied postcontrast conspicuity should be taken into account for follow-up examinations when different techniques are used, because this has the potential to confound treatment-response assessment, both in terms of enhancement degree and TV. Finally, our findings lend strength to designing data-acquisition strategies dedicated to MR imaging computer-assisted metastasis detection, as well as multiparametric diagnostic and prognostic assessment of gliomas, which rely on highly accurate and standardized measures of tumor MR imaging features.

The RECIST and RANO criteria, which represent standardized methodologies for the assessment of tumor response to treatment, respectively rely on 1D and 2D CEL measurements. In line with 3D-TV findings, both the 1D and 2D estimates performed in all measurable CELs were larger when using SPACE compared with MPRAGE. However, the limited number of measurable lesions available in this study did not allow a specific assessment of glioma and metastasis tumor types. Nevertheless, if confirmed in larger series, the present results might suggest the use of 3D-TSE techniques rather than MPRAGE to improve diagnostic accuracy in the design of clinical trials and in clinical practice.

Because the main goal of the present study was to test whether SPACE and/or VIBE may represent valid alternatives to MPRAGE, a direct comparison between SPACE and VIBE was not performed. However, we found that contrary to SPACE, VIBE did not lead to significant differences of morphometric assessment. We speculate that these differences are due to varied technique characteristics. For example, in SPACE sequences, the magnetization refocusing is obtained by radiofrequency pulses, which increase image resistance to magnetic field inhomogeneity. Another limitation with VIBE is the use of partial k-space acquisition to increase time efficiency so that the acquired voxel size is larger than the final interpolated size, thus potentially having a blurring effect. In addition, VIBE images show vessel enhancement, which may confound CEL boundary segmentation.

**Limitations and Future Perspectives**

While our sample size is relatively small, thorough selection was performed to represent cases potentially encountered in a routine neurosurgical and/or stereotactic radiosurgery treatment setting. The time gap between image acquisition and contrast agent injection represents a potential bias for the comparison of sequences acquired during the same examination because a larger GBCA amount may accumulate in the tumor within this time. However, obtaining data acquisition at exactly the same time for all test sequences would ideally require repeat contrast injections, resulting in an unnecessary dose increase to patients. We limited this bias by the following: 1) randomizing the sequence-acquisition order, 2) starting image acquisition 5 minutes after injection to allow a plateau phase, 3) reducing scan duration using parallel imaging, and 4) repeating quantitative analyses after adjusting for the acquisition order of compared sequences. Another potential confounding factor included different characteristics of GBCA uptake within posttreatment change. In principle, this factor may bias morphometry assessment in previously treated (relapsing) gliomas. However, all these were carefully selected to differentiate tumor from posttreatment-related change, by using perfusion and diffusion imaging, morphology, and sequential MR imaging follow-up. In addition, we detected no significant effect on the between-sequence volume differences. The potential variability across different magnetic field intensities and vendors providing 3D-TSE T1 (eg, CUBE, GE Healthcare, Milwaukee, Wisconsin; VISTA, Philips Healthcare, Best, the Netherlands) and/or 3D-fast gradient recalled echo volume-interpolated T1 (LAVA; General Electric) techniques was not assessed in this study. Finally, we did not focus on reproducibility assessment across dif-
different software tools, or operators performing segmentations. Further investigations are warranted to extrapolate our results to additional treatment centers using different computer-assisted segmentation approaches and varied operator experience levels.

CONCLUSIONS

SPACE and VIBE postcontrast techniques may provide better visualization of brain tumor enhancement than MP-RAGE. Morphometric discrepancies existed among sequences, with SPACE leading to potentially more accurate contouring estimates.

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