Detection of Local Recurrence in Patients with Head and Neck Squamous Cell Carcinoma Using Voxel-Based Color Maps of Initial and Final Area under the Curve Values Derived from DCE-MRI

J.Y. Lee, K.L. Cheng, J.H. Lee, Y.J. Choi, H.W. Kim, Y.S. Sung, S.R. Chung, K.H. Ryu, M.S. Chung, S.Y. Kim, S.-W. Lee, and J.H. Baek

ABSTRACT

BACKGROUND AND PURPOSE: Early detection of local recurrence is important to increase the chance of cure because local recurrence is the main cause of treatment failure in head and neck squamous cell carcinoma. We evaluated the added value of voxel-based color maps of dynamic contrast-enhanced MR imaging compared with conventional MR imaging alone for detecting local recurrence of head and neck squamous cell carcinoma.

MATERIALS AND METHODS: We retrospectively enrolled 63 consecutive patients with head and neck squamous cell carcinoma after definitive treatment and posttreatment surveillance MR imaging studies that demonstrated focal enhancement at the primary site. Three independent readers assessed conventional MR imaging and a pair of color maps of initial and final 90-second time-signal intensity areas under the curve from dynamic contrast-enhanced MR imaging. The sensitivities, specificities, and accuracies of both conventional MR imaging alone and combined interpretation of conventional and dynamic contrast-enhanced MR imaging were assessed using the clinicopathologic diagnosis as the criterion standard. \( \kappa \) statistics were used to evaluate interreader agreement.

RESULTS: There were 28 patients with subsequently documented local recurrence and 35 with posttreatment change. Adding dynamic contrast-enhanced MR imaging to conventional MR imaging significantly increased the diagnostic accuracies for detecting local recurrence (48%–54% versus 87%–91%; \( P < .05 \)), with excellent interreader agreement (\( \kappa = 0.8; 95\% CI, 0.67–0.92 \) to \( \kappa = 0.81; 95\% CI, 0.69–0.93 \)). By all 3 readers, the specificities were also significantly improved by adding dynamic contrast-enhanced MR imaging to conventional MR imaging (22%–43% versus 87%–91%; \( P < .001 \)) without sacrificing the sensitivities (68%–82% versus 86%–89%; \( P > .05 \)).

CONCLUSIONS: Adding voxel-based color maps of initial and final 90-second time-signal intensity areas under the curve from dynamic contrast-enhanced MR imaging increases the diagnostic accuracy to detect local recurrence in head and neck squamous cell carcinoma by improving the specificity without sacrificing the sensitivity.

ABBREVIATIONS: CI = confidence interval; CE = contrast-enhanced; HNSCC = head and neck squamous cell carcinoma; DCE = dynamic contrast-enhanced; FAUC\(_{90}^{\text{initial}}\) = initial 90-second time-signal intensity area under the curve; IAUC\(_{90}^{\text{final}}\) = final 90-second time-signal intensity area under the curve; SI = signal intensity; TSI = time-signal intensity

Local tumor recurrence is the main cause (25.0%–54.9%) of treatment failures in head and neck squamous cell carcinoma (HNSCC).\(^1\)–\(^4\) Because distant metastasis occurs less frequently than local recurrence, it is reasonable that posttreatment surveillance should focus on detecting local tumor recurrence to increase the chance of cure.\(^5\) Biopsies can be a dilemma in heavily irradiated tissue because biopsy procedure may increase the risk of superimposed infection, failure to heal, and increased edema.\(^6\) Therefore, diagnostic techniques should provide a better estimate...
of the likelihood of a recurrence and thus enable a more reliable selection of patients who should undergo pathologic examination.

Conventional and advanced imaging techniques, including diffusion-weighted MR imaging and FDG-PET, have shown variable results in distinguishing posttreatment change from recurrent or persistent disease. Dynamic contrast-enhanced (DCE)-MR imaging has emerged as a complementary technique for the posttreatment surveillance of patients with HNSCC and has shown promising results for distinguishing benign lesions from malignancies and predicting future treatment response. A quantitative DCE-MR imaging study by Ishiyama et al demonstrated that posttreatment changes had significantly higher permeability surface area than newly diagnosed or previously treated recurrent tumor. In addition to quantitative analysis, semiquantitative analysis of DCE-MR imaging can also be performed, offering the advantage of easier implementation, because it does not require the acquisition of baseline T1 mapping or arterial input function and does not rely on the assumptions of pharmacokinetic modeling. Regarding semiquantitative analysis, another previous study showed that benign postradiation changes had significantly longer time-to-peak and a lower relative washout ratio than recurrent tumors. A recent study showed that $K_{ep}$ (flux rate constant) and area under the gadolinium concentration-time curve were significantly higher in residual tumor, showing the potential of DCE-MR imaging to distinguish benign posttreatment change from residual HNSCC. Our recent studies using semiquantitative analysis of DCE-MR imaging have shown a significant difference in time-signal intensity (TSI) curve patterns between recurrence and posttreatment change and that histogram analysis of the ratios between the initial and final 90-second area under the TSI curve from DCE-MR imaging improves the sensitivity of MR imaging with 11% more recurrent tumors detected compared with conventional MR imaging without compromising diagnostic specificity. However, there are limitations in using both techniques in clinical practice. Considering that most recurrent tumors are mixed with posttreatment change, TSI curve analysis averaging dynamic characteristics of the individual voxels of the entire volume of interest cannot detect early tumor recurrence or select a proper biopsy site for pathologic confirmation due to loss of anatomic information. Histogram analysis might detect early recurrent tumors mixed with posttreatment change but cannot provide a visual guide to select a proper biopsy site either.

To overcome those limitations, we obtained voxel-based color maps of area under the curve (AUC) values extracted from TSI curves during the initial and final 90 seconds, to assess the voxel-by-voxel difference in signal intensity change according to the time lag. The study aimed to evaluate the feasibility and added value of visual assessment of voxel-based color maps using the initial and final AUC values of TSI curves derived from DCE-MR imaging to detect local tumor recurrence in patients with HNSCC after definitive treatment.

**MATERIALS AND METHODS**

This case-controlled, retrospective cohort was approved by the institutional review board of Asan Medical Center, and acquisition of informed consent from patients was waived due to the retrospective nature of the study.

### Study Patients

A review of the data base of our institution revealed 192 consecutive patients with pathologically confirmed HNSCC and surveillance MR imaging after definitive treatment between March 2014 and May 2015. Definitive treatment included curative resection, an operation and adjuvant radiation therapy, or concurrent chemoradiation. Among these, we found 135 patients with a focal moderate to intensely contrast-enhancing lesion at the primary site on fat-suppressed contrast-enhanced (CE) T1-weighted MR images and, finally, recruited 68 patients with a final diagnosis made by pathology or clinicoradiologic follow-up of >12 months. Subsequently, an independent neuroradiologist (J.Y.L., with 5 years of experience in head and neck imaging) who was not involved in the following image interpretation and data processing checked the image quality of DCE-MR imaging and excluded 5 patients with significant metallic artifacts, leaving 63 patients as a test patient group. Three of 63 patients have been previously reported. This prior article dealt with histogram analysis of DCE-MR imaging to evaluate local tumor recurrence in HNSCC, whereas in the current article, we investigated the feasibility of voxel-based color maps of DCE-MR imaging. Twenty patients were randomly selected as a separate patient group for a training session to distinguish among those with an enhancing lesion at the primary site but not confirmed as either recurrence or posttreatment change, to train readers and to minimize interreader variability (Fig 1).

### MR Imaging Acquisition and Generation of Voxel-Based Color Maps from DCE-MR Imaging

MR imaging examinations were performed with a 3T MR imaging scanner (Magnetom Skyra; Siemens, Erlangen, Germany) using a 64-channel head and neck coil. Our MR imaging protocol for head and neck tumors consists of axial and coronal T1- and T2-weighted turbo spin-echo sequences with DCE-MR images. All of the axial T1- (TR/TE = 790/11 ms) and T2-weighted images (TR/TE = 5470/85 ms) were acquired with an FOV = 190 × 190 mm², matrix size = 448 × 291, and slice thickness = 3 mm without gap. DCE-MR imaging was performed in an axial plane with 3D controlled aliasing in parallel imaging results in higher acceleration (CAIPIRINHA) technique before, during, and after administration of a standard single-bolus administration of 0.1 mmol of gadoterate meglumine (Dotarem; Guerbet, Aulnay-sous-Bois, France) per kilogram of patient body weight dose with a rate of 4 mL/s. Dynamic acquisition was performed with a temporal resolution of 3.2 seconds, and contrast was administered after 11 baseline dynamics (total = 144 dynamics). Axial (TR/TE = 650/12 ms) and coronal (TR/TE = 540/11 ms) CE-T1WIs with fat suppression were obtained after DCE-MR imaging. Detailed imaging parameters for DCE-MR imaging are as follows: slice thickness = 3 mm without gap; 20 slices; z-axis coverage = 60 mm; spatial in-plane resolution = 184 × 160; TR/TE = 6.3/3.1 ms; flip angle = 15°; FOV = 19 cm; total acquisition time = 7 minutes 24 seconds.

The DCE-MR imaging data were transferred to a personal
computer to process perfusion parametric maps with an in-house plug-in program, which was developed for DCE-MR imaging data processing by calculating the trapezoidal integration of normalized signal intensity (SI) using ImageJ software (National Institutes of Health, Bethesda, Maryland). The contrast index was calculated for the voxels on each DCE-MR image using the following formula: Contrast Index = \[ \frac{\text{SI (Postcontrast)} - \text{SI (Precontrast)}}{\text{SI (Precontrast)}} \] The time course of the contrast index was plotted to obtain a TSI curve and the initial and final 90-second AUC TSI values (IAUC90 and FAUC90) for each voxel. IAUC90 was defined as the trapezoidal integration of the normalized TSI curve during the initial 90 seconds from the onset of contrast enhancement in the contrast-enhancing voxel, indicating a measurement of the initial arrival of contrast agent in the tissue of interest after intravenous bolus administration that reflects blood flow, vascular permeability, and the fraction of interstitial space. FAUC90 was defined as the trapezoidal integration of the normalized TSI curve during the last 90 seconds in the same contrast-enhancing voxel, with IAUC90 indicating the amount of contrast agent leakage within the extravascular extracellular space. The IAUC90 and FAUC90 values were then constructed to generate voxel-based color maps, which ranged from blue, green, yellow, to red (Fig 2). The SI of the enhanced tissue was displayed after normalization, relative to the unaffected prevertebral muscles on baseline images. The signal intensity of skeletal muscles was coded blue, and the vascular signal intensity exhibiting maximal enhancement was coded red. The average time for generating a pair of color maps in 1 DCE study was <2 minutes. Details for the postprocessing of the DCE-MR imaging data using the in-house-developed plug-in are described in the On-line Appendix and Online Figs 1 and 2.

Analysis of Conventional and DCE-MR Imaging Examinations

After reviewing axial CE-T1WI of 20 patients in the separate patient group to train readers, a neuroradiologist (K.L.C., with 7 years of experience), who was aware of the location of the primary tumor, constructed a TSI curve of DCE-MR imaging for the focal masslike enhancing lesion at the primary site using a hand-drawn ROI with ImageJ software. The ROIs were double-checked and supervised by another experienced neuroradiologist (J.H.B., with 20 years of experience). All ROIs were free from areas of necrosis or nontumor macrovessels depicted on structural images. The TSI curve was interpreted by the neuroradiologist as progressive increment (type 1), plateau (type 2), or washout (type 3). The same neuroradiologist drew a rectangular ROI on axial CE-T1WIs of the separate patient group used to train readers and the patient group to assess the diagnostic accuracy as a guide for visual assessment of voxel-based color maps by different readers.

Three neuroradiologists (J.H.L., Y.J.C., and H.W.K. with 16, 9, and 5 years of experience in head and neck imaging, respectively) conducted a visual assessment of voxel-based color maps of the

FIG 1. Diagram of study population enrollment.
separate patient group used to train readers to minimize inter-
reader variability in their interpretation. For each patient, they
assessed each pair of IAUC90 and FAUC90 images and then deter-
mined the patterns of the color change as progressive increment
(type 1), plateau (type 2), and washout (type 3). The results were
then correlated with those of the TSI curve-pattern analysis, and
discrepancies were discussed to assess agreement. After the train-
ing session, they performed an independent analysis of the patient
group to assess the diagnostic accuracy blinded to the final diag-
nosis using the same methods. They regarded type 2 (plateau) or
type 3 (washout) as a recurrent tumor and type 1 (progressive
increment) as a posttreatment change.

All 3 neuroradiologists performed a blinded review of conven-
tional MR images of the patient group to test the diagnostic accu-
ry to detect recurrent tumor or posttreatment change, and those results were used to investigate the added value of voxel-based color maps over conventional MR imaging. Image sets included axial T2-weighted images, T1WIs, fat-suppressed CE-T1WIs, and coronal fat-suppressed CE-T1WIs. The interpreta-
tion of conventional MR imaging was performed according to the follow-
ing criteria: Recurrent tumors were defined as a focal mass discriminated by the surrounding tissue, with intermediate to slightly high signal intensity on T2WI, low signal on T1WI, and moderate-to-strong enhancement after gadolinium administra-
tion. Degree of enhancement was defined as “moderate” when enhancement was weaker than mucosal enhancement but greater than skeletal muscles. “Strong enhancement” was defined when the degree of enhancement was similar to or stronger than mucosal enhancement. “Posttreatment change” was defined as a fibrous scar on any focal linear or triangular enhancing lesion with very low signal intensity on T2WIs and low signal intensity on T1WIs, and “posttreatment inflammation,” as a diffuse hyperin-
tense abnormality on T2WIs with strong contrast enhancement. Results of conventional MR imaging analysis for the detection of recurrent tumor were classified into very probable, somewhat probable, somewhat unlikely, and unlikely.

Interpretation of both conventional and DCE-MRIs was conducted with a 1-month interval from the interpretation session for conventional MR imaging alone. To assess the added value of DCE-MR imaging, we accepted the result on DCE-MR imaging with only score 2 or 3 on conventional MR imaging. Score 1 or 4 on conventional MR imaging was ac-
tcepted as a final interpretation result regardless of the pattern on DCE-MR imaging.
The Cohen κ coefficient was used to measure interreader agreement of the results of DCE-MR imaging voxel-based color maps among 3 readers and agreement of the results between the TSI curve pattern and voxel-based color maps in the training session. All statistical analyses were performed using MedCalc for Windows, Version 13.0 (MedCalc Software, Mariakerke, Belgium) and SAS 9.4 software (SAS Institute, Cary, North Carolina). The significance threshold for the differences was set at a P value < .05.

### RESULTS

The patient group to test the diagnostic accuracy included 47 men (mean age, 59 years; range, 25–84 years) and 16 women (mean age, 59 years; range, 33–79 years) with a mean age of 59 years (range, 25–84 years). There were 35 patients (male/female ratio = 29:6) with posttreatment change and 28 (male/female ratio = 18:10) with local tumor recurrence. All cases with local recurrence were located at the area of the primary site. Local tumor recurrence was confirmed by pathologic examination (n = 18, 64%) or size increase during follow-up (n = 10, 36%). Posttreatment change was confirmed by pathologic examination (n = 7, 20%) or a decrease or no growth of the abnormal enhancing lesion without any treatment by at least 12-month follow-up imaging (n = 28, 80%). The median time interval between the end of definitive treatment and the detection of an enhancing lesion in the primary site was 26.3 months (range, 12–60 months) for all patients in the test group, 27.0 months (range, 12–60 months) for the posttreatment-change group, and 24.0 months (range, 10–41 months) for the recurrence group, respectively. All enhancing lesions were detected during routine imaging surveillance in asymptomatic patients. There was no significant difference in demographic data or treatment modalities between the 2 subgroups.

Descriptive statistics regarding the demographic and clinical data obtained in both test and training groups are summarized in Table 1.

Among the separate patient group to train readers, TSI curve patterns were classified into type 1 in 12 (60%), type 2 in 5 (25%), and type 3 in 3 (15%) patients. Voxel-based color maps were classified into type 1 in 11 (55%), type 2 in 6 (30%), and type 3 in 3 (15%) patients by visual assessment. Agreement between the TSI curve pattern and visual assessment of voxel-based color maps was excellent ($\kappa = 0.96$; 95% CI, 0.89–0.98; $P < .001$).

Table 2 shows the detailed results of analysis of conventional MR imaging and the combined approach of conventional MR imaging.
imaging and DCE-MRI for detection of local tumor recurrence after evaluation of the patient group to test diagnostic accuracy. False-positive and false-negative rates from Table 2 were 57%–77% and 18%–32% for conventional MR imaging and 9% and 11%–18% for the combined interpretation of conventional and DCE-MRI by 3 readers, respectively. There was a 48–68 percentage point and 4–14 percentage point reduction in both false-positive and false-negative rates by combined interpretation compared with conventional MR imaging alone. Agreement between the TSI curve pattern and visual assessment of voxel-based color maps was excellent among 3 readers in the patient group to test diagnostic accuracy (κ = 0.86; 95% CI, 0.75–0.96 to κ = 0.87; 95% CI, 0.77–0.97). Interreader agreement for interpretation of voxel-based color maps of DCE-MR imaging was also excellent (κ = 0.80; 95% CI, 0.67–0.92 to κ = 0.81; 95% CI, 0.69–0.93).

Table 3 summarizes diagnostic accuracies of conventional MR imaging alone and combined interpretation of conventional and DCE-MRI. Diagnostic accuracies of conventional MR imaging by all 3 readers were significantly increased from 48%–54% to 87%–91% by adding DCE-MRI to conventional MR imaging in all patients (P < .05). The specificities were also significantly improved by adding DCE-MR imaging to conventional MR imaging (22%–43% versus 87%–91%; P < .001) without sacrificing the sensitivities (68%–82% versus 86%–89%; P > .05). Figures 3–5 are representative cases.

Table 3: Diagnostic accuracies of conventional MRI alone and combined interpretation of conventional and DCE-MRI for detecting local tumor recurrence

| Reader | PPV | NPV | Sensitivity | Specificity | Accuracy |
|--------|-----|-----|-------------|-------------|----------|
|        |     |     |             |             |          |
| Reader 1 |     |     |             |             |          |
| Conventional MRI | 45 (39–51) | 57 (32–80) | 79 (66–90) | 22 (11–32) | 48 (35–61) |
| Conventional and DCE-MRI | 89 (74–96) | 91 (79–97) | 89 (72–98) | 91 (77–98) | 91 (80–96) |
| Reader 2 |     |     |             |             |          |
| Conventional MRI | 49 (39–58) | 63 (38–75) | 68 (54–81) | 43 (25–49) | 54 (41–67) |
| Conventional and DCE-MRI | 82 (67–91) | 91 (79–97) | 88 (70–98) | 87 (71–76) | 87 (77–91) |
| Reader 3 |     |     |             |             |          |
| Conventional MRI | 48 (40–54) | 67 (42–86) | 82 (69–93) | 29 (18–37) | 52 (39–65) |
| Conventional and DCE-MRI | 89 (73–96) | 89 (76–95) | 86 (67–96) | 91 (77–98) | 89 (78–95) |

Note: NPV indicates negative predictive value; PPV, positive predictive value.

Numbers in parentheses are 95% confidence intervals.

Statistical difference between conventional MRI and combined interpretation of conventional and DCE-MRI (P < .05).

FIG 3. A 53-year-old man after hemiglossectomy for squamous cell carcinoma of the oral tongue 12 months previously (T2N0M0; depth of invasion, 10 mm). A and B, T2-weighted and contrast-enhanced fat-suppressed T1-weighted images show an ill-defined enhancing lesion at the operative bed of the oral tongue (arrows). C and D, IAUC90 and FAUC90 images of DCE-MR imaging show contrast washout at the corresponding area. E, The time-signal intensity curve obtained from the enhancing area using a hand-drawn ROI also shows contrast washout. This lesion was confirmed as recurrent tumor on subsequent surgical excision.
**DISCUSSION**

Differentiating residual or recurrent tumor from posttreatment change is a clinically challenging decision, resulting in high false-positive rates (26%–46%) using conventional MR imaging criteria.\(^\text{25,26}\) FDG-PET has also been widely used to detect recurrent tumor due to its low false-negative rates and high negative predictive values for local tumor recurrence.\(^\text{27}\) However, low specificity might be a problem of PET/CT due to posttreatment granulation tissue and inflammatory changes.\(^\text{27}\)

In this study, we demonstrated that visual assessment of voxel-based color maps of DCE-MR imaging is both feasible and technically valid for detecting local tumor recurrence at the primary site in patients with HNSCC during surveillance. When conventional MR imaging showed low specificity for distinguishing recurrence from posttreatment change, there was significant improvement of the specificity and accuracy without sacrificing sensitivity when conventional and DCE-MRIs were interpreted together. Hence, we conclude that voxel-based color maps of DCE-MR imaging might be useful to rule out the probability of local tumor recurrence as a complementary tool to conventional MR imaging during posttreatment surveillance. We believe that this technique has the potential to reduce unnecessary biopsy or imaging studies and to lead to a more targeted diagnostic and therapeutic approach with reduced complications by improving the diagnostic accuracy of conventional MR imaging.

Previous studies on the diagnostic performance of conventional MR imaging for local tumor recurrence showed a wide variation in the diagnostic sensitivity (84%; 95% CI, 72%–92%) and specificity (82%; 95% CI, 72%–89%), according to a recent meta-analysis.\(^\text{28}\) In our study, the specificity of conventional MR imaging was relatively low, which might be attributable to our strict interpretation criteria: We considered any focal enhancement inconsistent with posttreatment scar or inflammatory change as recurrent tumor. In contrast, DCE-MR imaging has the potential of low sensitivity compared with conventional MR imaging due to the relatively low signal-to-noise ratio and the spatial resolution of the color maps and the variable degree of enhancement from posttreatment change. However, our results demonstrated that morphologic information with conventional MR imaging and functional information with color maps of DCE are complementary and might contribute in different ways to improvement of overall diagnostic performance. Therefore, combined interpretation of conventional and DCE-MRIs might be more practical and accurate during posttreatment surveillance of patients with HNSCC.

DCE-MR imaging has been useful in differentiating recurrent tumor from posttreatment change in head and neck cancer as well as in other anatomic areas. Previous studies revealed that enhancement patterns after gadolinium contrast injection differed between malignant and benign lesions: Early enhancement and washout corresponded to recurrence, while a progressive enhancement pattern corresponded to benign lesions, including fibrosis.\(^\text{10,29,30}\) Our study used the same methodology as the previous studies and revealed the feasibility and repeatability of using voxel-based color maps of IAUC\(_{90}\) and FAUC\(_{90}\) values from DCE-MR imaging in differentiating recurrent tumor from post-

---

**FIG 4.** A 40-year-old female patient after wide excision for buccal cancer 36 months previously (T1N0M0). T2-weighted image shows a low signal intensity lesion in the right retroantral region (A) with heterogeneous contrast enhancement on contrast-enhanced fat-suppressed T1-weighted imaging (B) (arrows). IAUC\(_{90}\) (C) and FAUC\(_{90}\) (D) images of DCE-MR imaging show a progressive increment pattern. Time-signal intensity curve obtained at the enhancing portion also shows a progressive increment pattern (E). This lesion was confirmed as fibrosis at excisional biopsy.
treatment change in HNSCC. Increased IAUC90 values in recurrent tumors are a reflection of increased blood flow and contrast agent leakage to the extravascular extracellular space, related to the hypervascularity and neoangiogenesis of malignant tumor. A progressive increase in FAUC90 in posttreatment change is due to increased retention of contrast agent in the extravascular extracellular space due to low cellularity.

On DCE-MR imaging, model-based pharmacokinetic DCE parameters are quantitative, having the potential to measure “true” physiologic mechanisms such as blood flow and permeability. However, model-based DCE parameters is difficult to apply in clinical practice for several reasons, including arterial input function measurement and model fit instability. AUC during a given time is a model-free parameter that describes the uptake of the contrast agent in a tissue of interest. It has advantages over the model-based approach because it does not require arterial input function measurement and does not need complex postprocessing pharmacokinetic modeling. In addition, the model-free technique is more robust, is unlikely to be influenced by scan sequence variation, and is easier and more reproducible compared with the model-based method.

The current study emphasizes the usefulness of visual assessment of voxel-based color maps. Many recurrent tumors occur intermixed with posttreatment changes. Because the TSI curve itself has averaged hemodynamic characteristics of the entirety of voxels in the ROI, the TSI curve has inherent limitations. If there is only a small focal area of recurrent tumor showing washout or a plateau pattern within the entire enhancing lesion, TSI curve analysis may overlook a small focus of viable tumor. To overcome this limitation, histogram-based assessment can be an alternative method because it allows comprehensive evaluation of an entire lesion volume. However, both time and effort to obtain the histogram parameters are required for all patients during daily practice. Voxel-based color maps may compensate for the drawbacks of TSI curve-pattern analysis and histogram analysis. While TSI curve analysis and histogram analysis need an ROI segmentation to extract a TSI curve or AUC-derived parameters, voxel-based color maps can provide simple and visually intuitive images and can help radiologists differentiate recurrent tumor from posttreatment change.

Our study has several limitations. In addition to the retrospective nature, the small sample size may diminish the statistical validity of the results. Future research with a large prospective cohort could further validate the methodology. Second, our study did not investigate any difference in the TSI curve pattern according to the status of human papillomavirus or Epstein-Barr virus, histologic tumor grade, or lesion volume. Given that these factors might affect the TSI curve pattern on DCE-MR imaging, the use of a fixed time interval on the TSI curve for voxel-based color maps can be another limitation. Again, additional investigation...
with a larger sample size may be necessary to verify the preliminary results of the study. Third, our study included patients with various tumor locations, which could present different natural histories. However, we believe that the coherent results of our study reflect the histologic consequences of an operation or chemoradiation that are common (granulation tissue and fibrosis), irrespective of the tumor location. Further studies might be necessary to evaluate any difference in the dynamic features of DCE-MR imaging according to a specific tumor subsite.

**CONCLUSIONS**

We demonstrated that adding visual assessment of voxel-based color maps of IAUC$_{90}$ and FAUC$_{90}$ derived from DCE-MR imaging is both a technically valid and reliable method for detecting local tumor recurrence at the primary site in patients with HNSCC during surveillance. Combined interpretation of both conventional and DCE-MRIs can be useful in ruling out the probability of a locally recurrent tumor by improved diagnostic specificity without sacrificing sensitivity.

Disclosures: Jung Hwan Baek—UNRELATED: Consultancy: RF Medical and StarMed since 2017.

**REFERENCES**

1. Agra IM, Carvalho AL, Ulbrich FS, et al. Prognostic factors in salvage surgery for recurrent oral and oropharyngeal cancer. *Head Neck* 2006;28:107–13 CrossRef Medline
2. Goodwin WJ Jr. Salvage surgery for patients with recurrent squamous cell carcinoma of the upper aerodigestive tract: when do the ends justify the means? *Laryngoscope* 2000;110(3 Pt 2 Suppl 93):1–18 CrossRef Medline
3. Kowalski LP, Bagietto R, Lara JR, et al. Prognostic significance of the distribution of neck node metastasis from oral carcinoma. *Head Neck* 2000;22:207–14 CrossRef Medline
4. Carvalho AL, Magrin J, Kowalski LP. Sites of recurrence in oral and oropharyngeal cancers according to the treatment approach. *Oral Dis* 2003;9:112–18 CrossRef Medline
5. Yom SS, Machtay M, Biel MA, et al. Survival impact of planned restaging and early surgical salvage following definitive chemoradiation for locally advanced squamous cell carcinomas of the oropharynx and hypopharynx. *Ann J Clin Oncol* 2005;28:385–92 CrossRef Medline
6. Bahadur S, Amatyia RC, Kacker SK. The enigma of post-radiation oedema and residual or recurrent carcinoma of the larynx and pyriform fossa. *Laryngol Otol* 1985;99:763–65 CrossRef Medline
7. Cheung PK, Chin RY, Edick GD. Detecting residual/recurrent head neck squamous cell carcinomas using PET or PET/CT: systematic review and meta-analysis. *Otolaryngol Head Neck Surg* 2016;154:421–32 CrossRef Medline
8. Cao Y, Popovtzer A, Li D, et al. Early prediction of outcome in advanced head-and-neck cancer based on tumor blood volume alterations during therapy: a prospective study. *Int J Radiat Oncol Biol Phys* 2008;72:1287–90 CrossRef Medline
9. Chikui T, Kitamoto E, Kawano S, et al. Pharmacokinetic analysis based on dynamic contrast-enhanced MRI for evaluating tumor response to preoperative therapy for oral cancer. *J Magn Reson Imaging* 2011;32:589–97 CrossRef Medline
10. Semiz Oysu A, Ayangolu E, Kodalli N, et al. Dynamic contrast-enhanced MRI in the differentiation of posttreatment fibrosis from recurrent carcinoma of the head and neck. *Clin Imaging* 2005;29:307–12 CrossRef Medline
11. Tomura N, Omachi K, Sakuma I, et al. Dynamic contrast-enhanced magnetic resonance imaging in radiotherapeutic efficacy in the head and neck tumors. *Am J Otolaryngol* 2005;26:163–67 CrossRef Medline
12. Ishiyama M, Richards T, Parvathaneni U, et al. Dynamic contrast-enhanced magnetic resonance imaging in head and neck cancer: differentiation of new H&N cancer, recurrent disease, and benign post-treatment changes. *Clin Imaging* 2015;39:566–70 CrossRef Medline
13. Furukawa M, Parvathaneni U, Maravilla K, et al. Dynamic contrast-enhanced MR perfusion imaging of head and neck tumors at 3 Tesla. *Head Neck* 2013;35:923–29 CrossRef Medline
14. King AD, Chow SK, Yu KH, et al. DCE-MRI for pre-treatment prediction and post-treatment assessment of treatment response in sites of squamous cell carcinoma in the head and neck. *PLoS One* 2015;10:e0144770 CrossRef Medline
15. Choi YJ, Lee JH, Sung YS, et al. Value of dynamic contrast-enhanced MRI to detect local tumor recurrence in primary head and neck cancer patients. *Medicine (Baltimore)* 2016;95:e2698 CrossRef Medline
16. Choi SH, Lee JH, Choi YJ, et al. Detection of local tumor recurrence after definitive treatment of head and neck squamous cell carcinoma: histogram analysis of dynamic contrast-enhanced T1-weighted perfusion MRI. *AJR Am J Roentgenol* 2017;208:42–47 CrossRef Medline
17. Chung WJ, Kim HS, Kim N, et al. Recurrent glomusbloma: optimum area under the curve method derived from dynamic contrast-enhanced T1-weighted perfusion MR imaging. *Radiology* 2013;269:561–68 CrossRef Medline
18. Evelhoch JL, LoRusso PM, He Z, et al. Magnetic resonance imaging measurements of the response of murine and human tumors to the vascular-targeting agent ZD6126. *Clin Cancer Res* 2004;10:3650–57 CrossRef Medline
19. Suh CH, Kim HS, Choi YJ, et al. Prediction of pseudoprogression in patients with glioblastomas using the initial and final area under the curves ratio derived from dynamic contrast-enhanced T1-weighted perfusion MR imaging. *AJR Am J Neuroradiol* 2013;34:2278–86 CrossRef Medline
20. Yabuuchi H, Fukuya T, Tajima T, et al. Salivary gland tumors: diagnostic value of gadolinium-enhanced dynamic MR imaging with histopathologic correlation. *Radiology* 2003;226:345–54 CrossRef Medline
21. Yuan J, Chow SK, Yeung DK, et al. A five-colour colour-coded mapping method for DCE-MRI analysis of head and neck tumours. *Clin Radiol* 2012;67:216–23 CrossRef Medline
22. Ailianou A, Mundada P, De Perrot T, et al. MRI with DWI for the detection of posttreatment head and neck squamous cell carcinoma: why morphologic MRI criteria matter. *AJR Am J Neuroradiol* 2018;39:748–55 CrossRef Medline
23. Becker M, Varoquaux AD, Combscure C, et al. Local recurrence of squamous cell carcinoma of the head and neck after radio(chemo)therapy: diagnostic performance of FDG-PET/MRI with diffusion-weighted sequences. *Eur Radiol* 2018;28:651–63 CrossRef Medline
24. King AD, Keung CK, Yu KH, et al. T2-weighted MR imaging early after chemoradiotherapy to evaluate treatment response in head and neck squamous cell carcinoma. *AJR Am J Neuroradiol* 2013;34:1237–41 CrossRef Medline
25. Gong QY, Zheng GL, Zhu HY. MRI differentiation of recurrent nasopharyngeal carcinoma from postradiation fibrosis. *Comput Med Imaging Graph* 1991;15:423–29 CrossRef Medline
26. Lell M, Baum U, Greess H, et al. Head and neck tumors: imaging recurrent tumor and post-therapeutic changes with CT and MRI. *Eur J Radiol* 2000;33:239–47 CrossRef Medline
27. Mukherji SK, Wolf GT. Evaluation of head and neck squamous cell carcinoma after treatment. *AJR Am J Neuroradiol* 2003;24:1743–46 Medline
28. van der Hoorn A, van Laar PJ, Holtman GA, et al. Diagnostic accuracy of magnetic resonance imaging techniques for treatment response evaluation in patients with head and neck tumours: a systematic review and meta-analysis. *PLoS One* 2017;12:e0177986 CrossRef Medline
29. Dao TH, Rahmouni A, Campana F, et al. Tumor recurrence versus fibrosis in the irradiated breast: differentiation with dynamic gadolinium-enhanced MR imaging. Radiology 1993;187:751–55 CrossRef Medline

30. Kinkel K, Ariche M, Tardivon AA, et al. Differentiation between recurrent tumor and benign conditions after treatment of gynecologic pelvic carcinoma: value of dynamic contrast-enhanced subtraction MR imaging. Radiology 1997;204:55–63 CrossRef Medline

31. Cheng HL. Improved correlation to quantitative DCE-MRI pharmacokinetic parameters using a modified initial area under the uptake curve (mIAUC) approach. J Magn Reson Imaging 2009;30:864–72 CrossRef Medline

32. Galbraith SM, Lodge MA, Taylor NJ, et al. Reproducibility of dynamic contrast-enhanced MRI in human muscle and tumours: comparison of quantitative and semi-quantitative analysis. NMR Biomed 2002;15:132–42 CrossRef Medline