Osteoradionecrosis (ORN), often with coexistent osteomyelitis, is a serious and often debilitating complication of radiation therapy for head and neck neoplasms. The mandible is the most common site of ORN due to its tenuous blood supply, though ORN can be seen in almost any bone within a radiated field. The primary factor implicated in the pathogenesis of ORN is the amount of radiation given to the affected bone, with both early (<2 years from radiation) and late onset ORN (>2 years from radiation) seen. There is a wide range of incidence of mandibular ORN reported, ranging from 5% to 22%, with more recent studies showing a decreased incidence, presumably attributable to increasing awareness and to improvements in preventive care and radiation techniques.

Patients are often referred for imaging to evaluate the extent of clinically suspected ORN, and, at the same time, to assess potential tumor recurrence. Multiple previous reports have characterized the imaging findings of ORN, namely soft-tissue thickening and enhancement, cortical bone erosion, trabecular disorganization, and bone fragmentation. All these findings can be seen in tumor recurrence, however, making the imaging differentiation of these 2 entities quite challenging.

We compared the relative frequency of CT and PET/CT findings of ORN and tumor recurrence to find patterns that might allow reliable differentiation of one entity from the other.

**MATERIALS AND METHODS**

**Patient Selection and Image Acquisition**

Our institutional review board approved this study, with a waiver of informed consent. All imaging examinations included in this...
study were performed as standard of care, and the results were retrospectively reviewed.

We searched our enterprise-wide electronic medical record encompassing 20 academic and community hospitals between January 1, 2008, and July 20, 2013, to identify patients in whom ORN was suspected on CT or PET/CT, or in whom a question of ORN versus recurrent tumor was raised. All neck and maxillofacial CT reports (including diagnostic neck CT studies performed as part of PET/CT) were searched by using the keywords “osteoradionecrosis,” “ORN,” “osteoradionecrosis versus tumor recurrence,” “osteoradionecrosis or tumor recurrence,” “osteoradionecrosis versus recurrent tumor,” and “osteoradionecrosis or recurrent tumor.” Patients were excluded if there was not pathologic confirmation of suspected tumor recurrence (1 patient) or if ORN could not be confirmed by either pathologic diagnosis or stability of >2 years on serial imaging (6 patients). A single patient was also excluded due to extensive artifacts from dental amalgam, which precluded accurate measurement of standard uptake value (SUV) values on PET/CT. The average time gap between imaging and pathologic confirmation of tumor recurrence or ORN was 14.3 days (range, 3–26 days) and 18.6 days (2–34 days), respectively. At our institution, all maxillofacial and neck CT examinations, as well as all PET/CT examinations performed for evaluation and/or follow-up of head and neck malignancy, are interpreted by fellowship-trained, Certificate of Added Qualification–certified neuroradiologists as part of a dedicated ear, nose, and throat imaging team. Demographic data collected included age and sex. Clinical data collected included location and type of primary neoplasm; initial imaging findings; follow-up imaging results, including any PET/CT results; time from completion of the last radiation therapy to the detection of the abnormality on imaging; and pathology results from either biopsy or operative intervention.

**CT Examinations**

CT examinations of the neck and maxillofacial bones were performed on 16- or 64-detector multidetector row CT scanners (LightSpeed VCT; GE Healthcare, Milwaukee, Wisconsin). CT acquisitions were obtained from the frontal sinuses to the thoracic inlet for the neck and from the top of the frontal sinuses through the mandible for the maxillofacial bones. Neck CT examinations were performed by using a pitch of 1.0, 2.5-mm collimation, 100 maximal mA, 120 kVp (peak), 21.0-cm FOV, in bone and standard algorithms, with 2.0-mm sagittal and coronal reconstructions. Maxillofacial CT studies were performed by using pitch of 0.5, 1.25-mm collimation, 225 maximal mA, 125 kVp, 19.6-cm FOV, with bone and standard algorithms, with 1.0-mm sagittal and coronal reconstructions.

**PET/CT Examinations**

Patients fasted for at least 6 hours before the PET/CT examination with the exception of water intake. Intravenous access was established for both blood glucose measurement and radiopharmaceutical administration. If serum glucose was >200 mg/dL, the examination was cancelled and rescheduled. Patients received between 10 and 17 mCi of IV [18F] FDG. Following radiopharmaceutical administration, the patients rested quietly during a standard 60-minute uptake period, after which imaging was performed.

The studies were performed on 16- to 64-section PET/CT scanners (Discovery; GE Healthcare). The CT scan parameters were 120–130 kVp, variable/smart milliamperes, and 3.75-mm collimation. CT scanning commenced following a 30-second delay after the administration of IV contrast (125-mL iopamidol, Isovue-370; Bracco Diagnostics, Princeton, New Jersey) and was performed from the top of the skull through the abdomen or pelvis (based on ordering clinician’s preference). Following CT, PET data were acquired by using a 4-minute bed position. The PET acquisition included an on-line delayed coincidence subtraction to correct for random coincidences as well as a dead-time correction. Two different PET/CT scanners were used in this study; one had a bismuth germanium oxide scintillation crystal, and the other had a lutetium oxyorthosilicate crystal. All PET acquisitions were uniform, using a 2D technique. The helical CT scan was reconstructed by filtered back-projection into 512 × 512 pixel images with a section thickness of 2.4 mm to match the PET scan. Images were reconstructed by using ordered-subset-expectation maximization with 2 full iterations of 8 subsets. Rescaled CT images were used to produce attenuation-correction values for the PET emission reconstruction.

**Image Analysis**

**CT Examinations.** All CT examinations were reviewed retrospectively in random order, separately by 2 Certificate of Added Qualification–certified neuroradiologists (L.A., S.F.), blinded to the clinical findings, the original study interpretation, and the final pathologic diagnosis. Reviewers were asked to comment on the presence or absence of the following imaging findings: 1) a solid measurable mass, 2) a cystic mass, 3) interruption of the cortical margin of the affected bone, 4) fragmentation of the affected bone, 5) loss of trabecula of the affected bone, 6) intraosseous gas in the affected bone, and 7) sclerosis of the affected bone. They were instructed that surrounding inflammatory change (ie, amorphous soft-tissue stranding or infiltration) and muscle or tissue thickening (Fig 1A, -B) did not qualify as a solid mass and that a discrete measurable mass (Fig 2) must be present. They were further asked to measure the size of any solid or cystic mass (Fig 2E, -F), and in those cases in which trabecular loss existed, they were asked to classify the pattern of trabecular loss as either permeative or lucent. They were instructed that a permeative pattern indicated multiple, independent focal lucent lesions, which, in sum, did not account for >75% loss of total bone trabecula (Fig 3A). A lucent pattern was defined as >75% loss of total bone trabecula in the area of abnormality (Fig 2C, -D). Radiologists were also asked to record the area or areas of the mandible affected, with choices being either the right or left mandibular condyle, coronoid process, ramus, angle, or body.

**PET Examinations.** In cases in which PET/CT was performed, the radiologists were asked to record both the mean (SUVmean) and maximum (SUVmax) SUV values. All PET studies were analyzed quantitatively with a software platform capable of deformable registration of multimodal images in the sagittal, coronal, and axial planes (Mirada XD3; Mirada Medical, Denver, Colorado). On PET scans,
metabolic volumes were manually delineated at the most metabolically active site of suspected ORN or tumor recurrence.

The SUV was automatically calculated by using the following formula:

$$SUV = \frac{C_{dc}}{d/w},$$

where \(d\) is the injected dose (in becquerels), \(w\) is the patient’s body weight (in grams), and \(C_{dc}\) is the decay-corrected tracer tissue concentration (in becquerels per gram). SUV was measured as \(SUV_{\text{mean}}\) and \(SUV_{\text{max}}\). These parameters were obtained from a 2D region of interest placed onto the axial image containing the most metabolically active area of abnormality (based on visual inspection). The smallest possible region of interest was used that would capture the most metabolically active area of abnormality. \(SUV_{\text{mean}}\) was defined as the average pixel value in the region of interest, while \(SUV_{\text{max}}\) was defined as the highest pixel value in the region of interest.

**Statistical Analysis**

Comparison of the demographic data between the 2 groups was performed by using a Fisher exact test for dichotomous variables and a 2-tailed, unpaired t test for continuous variables. The reproducibility of subjective imaging findings was tested between 2 fellowship-trained neuroradiologists on 50 randomly chosen cases from among patients with ORN and tumor, and a Cohen \(\kappa\) was calculated. Comparison of the prevalence of imaging findings between the 2 groups was performed with the Fisher exact test. Comparison of \(SUV_{\text{mean}}\) and \(SUV_{\text{max}}\) was performed between cases of osteoradionecrosis and tumor recurrence by using a 2-tailed, unpaired t test.

**FIG 1.** Images from 2 different patients with soft-tissue findings of osteoradionecrosis. A, Axial CT scan (soft-tissue window) shows enlargement and thickening of the musculature of the right masticator space (white arrowheads), without evidence of a distinct measurable mass, adjacent to an area of osteoradionecrosis. B, Axial CT scan (soft-tissue window) shows inflammatory thickening of soft tissues (white arrow) adjacent to an area of osteoradionecrosis involving the right mandibular body.

**FIG 2.** Images depicting soft-tissue findings of recurrent tumor. Axial CT scan from a 62-year-old man with recurrent oral cancer shows a large mass (white arrows) adjacent to the left mandibular angle in soft-tissue (A) and bone (B) algorithms. Axial CT scan at a different level shows an area of lucent trabecular loss involving the left mandibular angle (white arrowheads) in soft-tissue (C) and bone (D) algorithms. Axial CT images in an 83-year-old patient with recurrent oral cancer show another finding seen in recurrent tumor, a cystic mass (curved arrows), at 2 different levels (E and F).
Results
Fifty-three neck CTs and 10 maxillofacial CTs were evaluated. Twenty-three patients had a PET/CT performed as well. Seventeen of these studies demonstrated findings of tumor recurrence, while 46 demonstrated findings of ORN. Of the subset of patients with a PET/CT, 14 demonstrated findings of ORN, while 9 demonstrated findings of tumor recurrence. No significant differences were found between the 2 groups with regard to age, sex, or the interval between the end of radiation therapy and the detection of an abnormality on imaging (P = .11, 1.00, and 0.19, respectively). Demographic and clinical data for the 2 groups are summarized in the Table.

Most cases of ORN primarily affected the mandible (43 patients, 93%), with 1 case involving the hard palate, and 2 cases involving the maxilla. Most cases of ORN involving the mandible affected either 1 or 2 subsites of the mandible ipsilateral to the site of primary tumor (31 patients, 72%), with the remaining cases affecting either 3 (6 patients, 14%) or >3 (6 patients, 14%) subsites of the mandible. The preponderance of ORN cases involving the mandible affected the lingual surface (20 patients, 47%), while the remainder affected either the buccal surface (13 patients, 30%) or involved both the lingual and buccal surfaces (10 patients, 23%). None of the patients received hyperbaric oxygen before detection of ORN.

Interpretation of the subjective imaging findings between observers was almost perfect, by using Cohen κ interpreted according to Landis and Koch,15 with κ values as follows: 0.865 (95% CI, 0.683–1.00) for cortical disruption, 0.868 (95% CI, 0.724–1.00) for a pattern of trabecular loss, 0.891 (95% CI, 0.743–1.00) for the presence of a cystic mass, 0.940 (95% CI, 0.823–1.00) for the presence of a solid mass, 0.929 (95% CI, 0.792–1.00) for the presence of bony sclerosis, and 1.000 (95% CI, 1.00–1.00) for the presence of intraosseous gas.

CT Examinations
The CT imaging findings in both groups of patients are summarized in Fig 4. The presence of a measurable solid soft-tissue mass in the tumor-recurrence group was 59% (10 of 17 patients), which is significantly higher than that in the osteoradionecrosis group (1 of 46 patients, 2%, P < .001). Similarly, the presence of a cystic mass was also significantly higher in the recurrence group, 41% (7 of 17 patients), than in the osteoradionecrosis group (3 of 46 patients, 6.5%, P = .025).

Several osseous changes showed a significant difference between the 2 groups as well. A permeative pattern of trabecular loss was significantly higher in the ORN group, 80% (37 of 46 patients), than the tumor-recurrence group (4 of 24 patients, 16.7%, P < .0001). Bone sclerosis was also seen significantly more often in the setting of ORN, 28% (13 of 33 patients), than in tumor recurrence (0 of 17 patients, 0%, P = .013). While interruption of the cortical margin was seen significantly more often in the tumor-recurrence group (17 of 17 patients, 100%) than in the ORN group (33 of 46 patients, 71.7%, P = .013), it was noted to have a rather high prevalence in both groups.

PET Examinations
The SUVmean was significantly higher in cases of tumor recurrence (SUVmean range, 4.6–14.1; mean, 8.0) compared with ORN (SUVmean range, 2.2–7.5; mean, 4.3), with a P value of .0021. Likewise, the SUVmax was significantly higher in cases of tumor recurrence (SUVmax range, 5.7–20.2; mean, 11.3) compared with ORN (SUVmax range, 1.7–9.2; mean, 5.3), with a P value of 16.7%.

Patient demographics and clinical characteristics

| Patients Diagnosed with Osteoradionecrosis | Patients Diagnosed with Tumor Recurrence | P Value |
|------------------------------------------|----------------------------------------|---------|
| No. of patients                          | 46                                     | 17      | NA                  |
| Average age (yr)                         | 62                                     | 69      | .11                 |
| Age range (yr)                           | 33–93                                  | 53–85   | .025                |
| % Men (Nc)                               | 761 (35)                               | 77.5 (13) | 1.00               |
| Primary malignancy site                  | Oral Cavity (n = 27), oropharynx (n = 11), parotid (n = 5), NPC (n = 2), unknown (n = 1) | Oral cavity (n = 15), oropharynx (n = 1), sinonasal (n = 1) |         |
| Primary malignancy histology             | Squamous cell (n = 44), adenoid cystic (n = 2) | Squamous cell (n = 17) |         |
| Mean interval (range) between last RT and detection of abnormality (mo) | 40.56 (6–110) | 48.82 (5–98) | .39               |

Note:—NA indicates not applicable; NPC, nasopharyngeal cancer; RT, radiation therapy.
There was, however, substantial overlap in both SUV_{mean} and SUV_{max} values between the 2 entities (Fig 5A, -B).

In the subset of patients with ORN who underwent histopathologic sampling, 20% (5 of 25 patients), had pathologic evidence of associated osteomyelitis. Unfortunately, only 1 of those patients also underwent a PET/CT examination; therefore, it is difficult to compare the relative SUV values of ORN with and without associated osteomyelitis.

DISCUSSION

The findings of a cystic or solid mass on imaging were significantly more prevalent in patients with tumor recurrence, being relatively
rare in patients with ORN. Conversely, the presence of bony sclerosis was not uncommon in patients with ORN but was never seen in tumor recurrence. Although the SUV values in tumor recurrence were significantly higher than in ORN, the substantial overlap in the SUV values for these 2 entities renders SUV measurements relatively impractical for differentiating ORN from tumor recurrence reliably in the clinical setting.

We attempted to evaluate the utility of various CT findings and PET/CT parameters in an effort to distinguish ORN from recurrent malignancy in the head and neck following irradiation for head and neck cancer. Multiple prior studies have also found the relative unreliability of an elevated SUV in differentiating recurrent tumor from ORN.\(^1,11,12,16-18\) While recurrent tumor does, in general, demonstrate higher SUV\(_{\text{mean}}\) and SUV\(_{\text{max}}\) values compared with ORN, there is significant overlap, which makes differentiating the 2 entities on a case-by-case basis extremely unreliable. This overlap is presumably responsible for previous reports of false-positive results in PET scans attributed to osteoradionecrosis.\(^16\)

Unlike SUV values, there were conventional imaging findings that were rather selective for differentiating ORN from tumor recurrence. In evaluating tumor recurrence, a discrete, associated mass provides a significant diagnostic indication for the presence of recurrent tumor. While ORN may be associated with inflammatory and edematous changes in the surrounding musculature and soft tissues,\(^10,19-21\) an asymmetric, discrete solid mass was seen in more than half of the patients with recurrent malignancy, but it was exceedingly rare in ORN, seen in only 2% of patients with ORN. Similarly, a cystic mass was seen in nearly half of patients with recurrent tumor, but in <10% of patients with ORN. This indicates that the presence of a cystic component in a patient with suspected ORN should not be assumed to be an abscess from associated osteomyelitis but should instead prompt concern for possible recurrent neoplasm.

Osseous findings on CT can also be extremely helpful in differentiating ORN from tumor recurrence. Multiple prior studies have delineated the classic bony CT findings associated with ORN, namely cortical disruption, trabecular loss, bone fragmentation, bony sclerosis, and the presence of intraosseous gas.\(^1,10,11,13,21\) However, we tested the value of these imaging findings for discriminating ORN and tumor recurrence. Traditionally, on the basis of known pathologic findings in ORN, it has been thought that a more permeative pattern of bone loss is seen in ORN as opposed to tumor recurrence. This was thought to be related to the fact that irradiation leads to a relatively hypoxic, hypocellular, and hypovascular substrate with an inconsistent ability to remodel tissue loss secondary to radiation-induced injury\(^22\); this, in turn, would lead to relatively less bony loss than would be seen when tissue is being actively destroyed and replaced by tumor. In our patients, the permeative pattern of bone loss, though significantly more common in ORN, was also seen in a significant minority of patients with tumor recurrence. This may be related to our patient population, which is actively surveyed for head and neck cancer, and PET/CT parameters in an effort to distinguish ORN from tumor recurrence reliably in the clinical setting.

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Perhaps the most useful imaging feature in distinguishing ORN from tumor recurrence was the presence of bony sclerosis (Fig 3B). Seen in nearly one-third of patients with ORN, it was not seen in any of our patients with recurrent tumor. This finding may reflect the tendency of squamous cell carcinoma to induce bony destruction rather than sclerosis in contradistinction to the often chronic nature of ORN that may be associated with bony sclerosis.\(^13,23\) Thus, while most patients with ORN may not have bony sclerosis, the presence of this finding on imaging should prompt a relatively confident diagnosis of ORN rather than tumor recurrence. Similarly, intraosseous gas, which has been suggested to be pathognomonic of superimposed osteomyelitis,\(^24\) was not seen in any of our patients with tumor recurrence. This finding did not reach significance in our study, however, in part due to its overall rarity on imaging for either entity.

These imaging findings are readily incorporated into clinical practice and were highly reproducible among observers in our study. The near-perfect agreement indicated by the very high \(k\) values for each of the imaging findings evaluated in this study suggests that interpretation of these imaging findings will be consistent when implemented clinically.

There were several limitations to our study. We purposefully geared our initial search to evaluate only lesions that were either thought to be ORN on imaging or for which the interpreting radiologist entertained the diagnosis of ORN versus tumor recurrence. While this search results in a selection bias toward cases in which the 2 entities look similar and results in a greater number of patients with ORN, we would argue that these “problem cases” are most relevant to everyday practice and the situation in which specific imaging findings may be the most useful. Additionally, not all patients in our study with ORN had pathologic correlation; in many cases, the clinician and patient preferred a noninvasive method of follow-up. As such, it is possible that undetected tumor recurrence occurred in the bed of ORN, confounding evaluation of imaging findings. However, we believe the chances of recurrent tumor remaining stable on imaging studies during a 2-year period are relatively low. Finally, clinical variables such as radiation dose and time from radiation were not considered in our analysis because we, instead, chose to focus on the imaging picture as it was presented to an interpreting radiologist.

**CONCLUSIONS**

The presence of a discrete solid or cystic mass is associated with tumor recurrence, while bony sclerosis, though not common, was seen exclusively in the setting of ORN in our study. Permeative rather than lucent trabecular loss was most often seen in the setting of ORN, but it may also be seen in a minority of patients with recurrent tumor. Finally, while an elevated SUV does suggest a diagnosis of tumor recurrence, the significant overlap of SUV values in patients with tumor recurrence and ORN renders SUV values relatively impractical for use on an individual clinical basis.
REFERENCES

1. Rabin BM, Meyer JR, Berlin JW, et al. Radiation-induced changes in the central nervous system and head and neck. Radiographics 1996;16:1055–72.
2. Rayatt SS, Mureau MA, Hoffer SO. Osteoradionecrosis of the mandible: etiology, prevention, diagnosis and treatment. Indian J Plast Surg 2007;40:65–71.
3. Bras J, de Jonge HK, van Merkesteyn JP. Osteoradionecrosis of the mandible: pathogenesis. Am J Otolaryngol 1990;11:244–50.
4. Chopra S, Kamdar D, Ugur O, et al. Factors predictive of the severity of osteoradionecrosis of the mandible. Head Neck 2011;33:1600–05.
5. Epstein JB, Wong FL, Stevenson-Moore P. Osteoradionecrosis: clinical experience and a proposal for classification. J Oral Maxillofac Surg 1987;45:104–10.
6. Støre G, Boysen M. Mandibular osteonecrosis: clinical behavior and diagnostic aspects. Clin Otolaryngol Allied Sci 2000;25:378–84.
7. Reuther T, Schuster T, Mende U, et al. Osteoradionecrosis of the jaw as a side effect of radiotherapy of head and neck tumour patients: a report of a thirty year retrospective review. Int J Oral Maxillofac Surg 2003;32:289–95.
8. Jereczek-Fossa BA, Orrechia R. Radiotherapy-induced mandibular bone complications. Cancer Treat Rev 2002;28:65–74.
9. Studer G, Gratz KW, Glanzmann C. Osteoradionecrosis of the mandible in patients treated with different fractionations. Strahlenther Onkol 2004;180:233–40.
10. Chong J, Hinckley L, Ginsberg L. Masticator space abnormalities associated with mandibular osteoradionecrosis: MR and CT findings in five patients. AJNR Am J Neuroradiol 2000;21:175–78.
11. Hamilton JD, Lai SY, Ginsberg LE. Superimposed infection in mandibular osteoradionecrosis: diagnosis and outcome. J Comput Assist Tomogr 2012;36:725–31.
12. Yoo JS, Rosenthal DJ, Mitchell K, et al. Osteoradionecrosis of the hyoid bone: imaging findings. AJNR Am J Neuroradiol 2010;31:761–66.
13. Glastonbury CM, Parker EE, Hoang JK. The postradiation neck: evaluating response to treatment and recognizing complications. AJR Am J Roentgenol 2010;195:W164–71.
14. Wu L, Liu J, Wang C, et al. Osteoradionecrosis of the upper cervical spine after radiation therapy for head and neck cancer: differentiation from recurrent of metastatic disease with MR imaging. Radiology 2012;264:136–45.
15. Landis JR, Koch GG. The measurement of observer agreement for categorical data. Biometrics 1977;33:159–74.
16. Liu S-H, Chang J, Ng SH, et al. False positive fluorine-18 fluorodeoxy-D-glucose positron emission tomography finding caused by osteoradionecrosis in a nasopharyngeal carcinoma patient. Br J Radiol 2004;77:257–60.
17. Krabbé C, Pruim J, Dijkstra P, et al. 18F-FDG PET as a routine post-treatment surveillance tool in oral and oropharyngeal squamous cell carcinoma: a prospective study. Eur J Nucl Med 2009;36:1940–47.
18. Abgral R, Querellou S, Potrad G, et al. Does 18F-FDG PET/CT improve the detection of posttreatment recurrence of head and neck squamous cell carcinoma in patients negative for disease on clinical follow up? Eur J Nucl Med 2009;36:24–29.
19. Ng SH, Liu MH, Ko SF, et al. Post treatment imaging of the nasopharynx. Eur J Radiol 2002;44:82–95.
20. King AD, Griffith JF, Abrigo JM, et al. Osteoradionecrosis of the upper cervical spine: MR imaging following radiotherapy for nasopharyngeal carcinoma. Eur J Radiol 2010;73:629–35.
21. Hermans R, Fossion E, Ioannides C, et al. CT findings in osteoradionecrosis of the mandible. Skeletal Radiol 1996;25:31–36.
22. Marx RE. Osteoradionecrosis: a new concept of its pathophysiology. J Oral Maxillofac Surg 1983;41:283–88.
23. Store G, Larheim TA. Mandibular osteoradionecrosis: a comparison of computed tomography with panoramic radiography. Dentomaxillofac Radiol 1999;28:295–300.
24. Merine D, Fishman EK, Magid D. CT detection of sacral osteomyelitis associated with pelvic abscesses. J Comput Assist Tomogr 1988;12:118–21.