The Diagnostic Capacity of Pre-treatment $^{18}$F-FDG PET/CT for Predicting the Extranodular Spread of Lymph Node Metastases in Patients with Oral Squamous Cell Carcinoma

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The aim of this study was to evaluate the ability of pretreatment 90-min $^{18}$F-fluorodeoxyglucose positron emission tomography/computed tomography ($^{18}$F-FDG PET/CT) to predict the extranodular spread of lymph node metastases in oral squamous cell carcinoma. We retrospectively reviewed the cases of 56 patients who underwent pretreatment $^{18}$F-FDG PET/CT and surgery with neck dissection. Maximum standardized uptake value, metabolic tumor volume, and total lesion glycolysis were measured for the 56 primary sites and maximum standardized uptake value was measured for 115 lymph node levels. Extranodular spread was present at 9 lymph node levels in 7 patients. Significant differences were found in metabolic tumor volume and total lesion glycolysis of the primary site, and in lymph node maximum standardized uptake value, between patients with and without extranodular spread ($p<0.05$). Combining primary site total lesion glycolysis and lymph node maximum standardized uptake value at their respective optimal cutoffs, the specificity, and accuracy for predicting extranodular spread were 89%, 92%, and 92%, respectively. Pretreatment $^{18}$F-FDG PET/CT is useful for predicting extranodular spread in patients with oral squamous cell carcinoma. The combined use of primary site total lesion glycolysis and lymph node maximum standardized uptake value showed greater predictive value than either predictor singly.

Key words: $^{18}$F-fluorodeoxyglucose positron emission tomography/computed tomography, extranodular spread, metastasis, oral squamous cell carcinoma

In the treatment of oral squamous cell carcinoma (OSCC), patients with clinically diagnosed cervical lymph node (LN) metastases are subjected to radical neck dissection or modified radical neck dissection. The existence of extranodular spread (ENS) influences the method selection [1].

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88%, respectively, using a maximum standardized uptake (SUVmax) value of 2.25 as the cutoff for LN [4]. However, the detection of cervical LNs with an SUVmax of 2.25 is clinically difficult, because adjacent tissues or substances to which the 18F-FDG are also taken up, such as plaque in the carotid artery, have SUVmax values within the same range [5].

In contrast, measuring the 18F-FDG uptake of the primary site (PS) is relatively easy. Moreover, previous reports have demonstrated the utility of measuring the PS 18F-FDG uptake for predicting LN metastases in patients with primary lung carcinoma and OSCC in whom the 18F-FDG uptake to metastatic LNs was negative [6,7]. Furthermore, several recent papers revealed that metabolic tumor volume (MTV) and total lesion glycolysis (TLG) were more useful prognostic markers compared to SUVmax [7-10].

We hypothesized that the evaluation of the 18F-FDG accumulation at the PS could be another potential predictor of ENS in patients with OSCC. To the best of our knowledge, the correlation between ENS and the 18F-FDG uptake at the PS, and the possibility of diagnosing ENS by evaluating the 18F-FDG accumulation at the PS have not been clarified. Moreover, no report has assessed the utility of 90-min 18F-FDG PET/CT for the prediction of LN metastases in OSCC.

In this retrospective study, we compared four parameters collected by pretreatment with 90-min 18F-FDG PET/CT—SUVmax for PS (SUV-PS), MTV for PS (MTV-PS), TLG for PS (TLG-PS), and SUVmax of LN itself (SUV-LN)—for their ability to predict ENS in OSCC.

**Materials and Methods**

**Patient population.** We retrospectively reviewed the medical records of 56 patients with newly diagnosed OSCC who underwent pretreatment 18F-FDG PET/CT for initial staging between August 2010 and November 2016 (28 males, 28 females; age [mean ± SD], 70.43 ± 13.40; range, 23-92 years). All patients underwent resection of the primary tumor and neck dissection with curative intent, and histological diagnosis of the primary tumor and lymph nodes was performed. Seventeen patients underwent comprehensive neck dissection, and 39 patients underwent selective neck dissection (Level 1, n = 11; Level 2, n = 1; Level 1-2, n = 2; Level 1-3, n = 14; Level 1-4, n = 11). Exclusion criteria were a previous history of malignancy within 5 years, induction chemotherapy, and a serum glucose level of more than 150 mg/dl prior to the radiotracer injection.

Data were obtained from medical records, including clinicopathological variables and treatment. Tumors were staged according to the TNM Classification of Malignant Tumors, 8th edition [11]. The study was approved by the institutional review board of our hospital and the requirement of informed consent was waived (1811-012).

**18F-FDG PET/CT procedures and analysis of PET/CT images.** All 18F-FDG PET/CT examinations were performed using an integrated PET/CT scanner (Biograph 16; Siemens Medical Solution USA, Knoxville, TN, USA) at a diagnostic imaging center adjacent to our institution. After fasting for at least 5 h, the patients received an intravenous injection of 3.7 MBq/kg 18F-FDG. At our institution, PET image acquisition is routinely started 90 min after injection of 18F-FDG, with the patient in a relaxed supine position. First, a total-body low-dose CT scan for the calculation of attenuation correction was performed, using a standardized protocol involving 120 kV, auto mA mode, rotation time of 0.5 sec, pitch of 0.8, section thickness of 3 mm, and scan field from the head to the mid-thigh level. Immediately thereafter, PET imaging consisting of 6-8 bed positions with 2.4 min per position over the same region was performed. The PET images were reconstructed with an ordered-subset expectation maximization iterative reconstruction algorithm.

Integrated, co-registered PET/CT images were obtained using a workstation that enables image fusion and analysis (syngo. via; Siemens Medical Solution USA). For semi-quantitative analyses of 18F-FDG uptake, the images were evaluated by 2 nuclear medicine physicians, by consensus. A volume of interest was drawn semi-automatically over the PS and all measurable dissected cervical LN levels on an axial slice. If 18F-FDG uptake was perceptible in a cervical LN, the SUVmax in the cervical LN was measured to minimize partial-volume effects [12,13]. If multiple nodes in the same LN level were measurable, the highest SUVmax value among them was adopted as the representative SUVmax of the LN level. For calculating MTV and TLG, the threshold values were set at an SUV of 2.5, based on the results of previous studies [8,9,14]. The semi-quantitative values of SUV-PS, MTV-PS, TLG-PS and SUV-LN were calculated automatically.
**Statistical analyses.** All analyses were performed using IBM SPSS Statistics (version 22; IBM Corp., Armonk, NY, USA). The Mann-Whitney U test was used to analyze differences in SUV-PS, MTV-PS, TLG-PS, or SUV-LN between the groups with and without ENS. Receiver operating characteristic (ROC) curve analysis of such semi-quantitative parameters was performed to differentiate the tumors with ENS from those without ENS, and to differentiate the LN levels with ENS from those without ENS. *P* values < 0.05 were considered statistically significant.

**Results**

**Characteristics of patients and oral squamous cell carcinoma.** Table 1 shows the patients’ characteristics. Seven out of 56 patients had 9 LNs with pathologically proven ENS in 115 dissected LN levels. The primary tumor size ranged from 12 to 85 mm, with most lesions being <50 mm in the long axis diameter. Figures 1 and 2 show representative 18F-FDG PET/CT images of OSCC with and without ENS.

**Standardized uptake value, metabolic tumor volume, and total lesion glycolysis.** Table 2 shows the SUV-PS, MTV-PS, TLG-PS, and SUV-LN values in the study groups. The SUV-PS (mean ± SD) values in patients with and without ENS were 16.84 ± 4.95 and 12.54 ± 5.98, the MTV-PS values in patients with and without ENS were 27.05 ± 12.25 ml and 14.09 ± 13.36 ml, and the TLG-PS values in patients with and without ENS were 153.03 ± 68.13 g and 80.85 ± 111.30 g, respectively. The SUV-LN values in patients with and without ENS were 8.18 ± 4.43 and 2.69 ± 2.37. Significant intergroup differences were found in MTV-PS, TLG-PS, and SUV-LN (all *p* < 0.05) but not in SUV-PS (*p* = 0.12).

The areas under the ROC curve (AUCs) for SUV-PS, MTV-PS, TLG-PS, and SUV-LN were 0.684 (*p* = 0.12, 95% CI 0.534-0.834), 0.812 (*p* = 0.008, 95% CI 0.697-0.937), 0.816 (*p* = 0.007, 95% CI 0.676-0.956), and

![Fig. 1](image.png)

An 89-year-old woman with left lower gingival squamous cell carcinoma and level I cervical lymph node metastasis with extranodal spread. Axial computed tomography (CT) (A) shows soft tissue density in the primary site (arrowhead) and an enlarged level I lymph node (arrow). Positron emission tomography (PET) (B), positron emission tomography/computed tomography (PET/CT) (C), and maximum intensity projection (MIP) (D) images show high 18F-FDG uptake of the primary site (SUVmax, MTV, and TLG were 28.54, 25.19 ml, and 178.53 g, respectively) and lymph node (SUVmax was 8.85), which is easy to distinguish from adjacent tissues.
0.932 (p < 0.001, 95% CI 0.882-0.982) with the optimal cut-off values of 13.46, 19.56 ml, 106.26 g, and 3.37, respectively (Fig. 3A, 3B). The sensitivity, specificity, and accuracy for predicting ENS using these cut-off values were 86%, 78%, and 80% for MTV-PS, and 86%, 78%, and 80% for TLG-PS, respectively. The sensitivity, specificity, and accuracy of SUV-LN levels for predicting ENS were 100%, 84%, and 86% using the optimal cut-off value.

**A predictor combining TLG in the primary tumor and SUVmax in lymph nodes.** TLG-PS showed a higher AUC than SUV-PS or MTV-PS, and SUV-LN demonstrated the highest AUC among all parameters. Combining the two criteria—namely, considering a TLG-PS greater than 106.26 g and an SUV-LN greater than 3.37—resulted in sensitivity, specificity, and accuracy of 89%, 92%, and 92%, respectively, for the prediction of ENS.

### Table 2

<table>
<thead>
<tr>
<th>Parameter</th>
<th>ENS group</th>
<th>non-ENS group</th>
<th>p value*</th>
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<tbody>
<tr>
<td>Primary site</td>
<td><strong>SUVmax</strong> 16.84 ± 4.95 (13.16-28.54)</td>
<td>12.54 ± 5.98 (3.38-29.15)</td>
<td>0.12</td>
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<td></td>
<td><strong>MTV (ml)</strong> 27.05 ± 12.25 (11.75-54.57)</td>
<td>14.09 ± 13.36 (0.99-83.12)</td>
<td>0.006</td>
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<td></td>
<td><strong>TLG (g)</strong> 153.03 ± 68.13 (52.34-289.91)</td>
<td>80.85 ± 111.30 (2.89-759.97)</td>
<td>0.005</td>
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<tr>
<td>Lymph node</td>
<td><strong>SUVmax</strong> 8.18 ± 4.43 (3.37-19.5)</td>
<td>2.69 ± 2.37 (1.08-17.02)</td>
<td>&lt;0.001</td>
</tr>
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ENS, extranodular spread; SUVmax, maximum standardized uptake value; MTV, metabolic tumor volume; TLG, total lesion glycolysis. The data are represented as mean ± standard deviation, data in parentheses are the range.

*P value: comparison of semiquantitative values between the ENS and non-ENS groups (Mann-Whitney U-test).
Several previous studies have investigated the utility of SUVmax measured by \( ^{18} \text{F-FDG PET/CT} \) for predicting LN metastases in patients with OSCC \([15-17]\). MTV is a measure of the volume of the tumor displaying \( ^{18} \text{F-FDG} \) uptake, and thus quantifies the overall tumor burden, and is theoretically a better predictor of outcome than the SUVmax \([18]\). TLG represents the metabolic activity throughout the entire tumor above a minimum threshold designed to exclude the background activity \([19]\). Therefore, volume-based parameters such as MTV and TLG may reflect the metabolic burden of the active tumor more accurately than SUVmax and thus could be better surrogate imaging markers of tumor biology \([7, 20, 21]\).

The first finding in the present study was that MTV-PS, TLG-PS, and SUV-LN were useful parameters to predict LNs with ENS in our study population. In a previous study \([4]\) of OSCC patients with 71 dissected LN levels, the SUV-LN in the tumors with ENS was greater than that in those without ENS, and the difference was significant. The ROC curve analysis of SUV-LN in the same study showed an AUC of 0.864 ± 0.045 and an optimal SUVmax cut-off value of 2.25, with 85% sensitivity and 86% accuracy. In our study, we achieved excellent diagnostic capacity by setting the SUV-LN cut-off value at 3.37. In clinical practice, it is relatively easy to detect LNs with an SUVmax greater than 3.37, even when surrounded by adjacent normal tissue with physiological uptake. The difference in diagnostic performance of SUV-LN between our study and the previous study \([4]\) could be due to the difference in the PET acquisition time point. PET image acquisitions were started at 90 min after injection of \( ^{18} \text{F-FDG} \) in the present study and at 60 min after injection in the previous study \([4]\). Another previous research on 2-h dual-time-point \( ^{18} \text{F-FDG PET} \) demonstrated that head and neck malignant tumors had a mean SUV increase of 12% between the 1-h (early) and the 2-h (late) phases, and that the \( ^{18} \text{F-FDG} \) uptake to inflammation and normal tissue had meanwhile decreased \([22]\). Another recent report revealed that the \( ^{18} \text{F-FDG} \) uptake to most normal cervical tissue decreases as time proceeds within 3 h after \( ^{18} \text{F-FDG} \) injection \([23]\). These results suggest that 90-min PET acquisition with SUV-LN measurement is superior to 60-min PET acquisition for the prediction of ENS in OSCC, and that the contrast between metastatic LNs with ENS and surrounding normal tissue might be better on 90-min than 60-min PET images.

This study has several limitations. First, it is inherently limited by its retrospective nature. Second, the study population was relatively small, especially in terms of patients having tumors with ENS. Finally, PET image acquisition was performed only at 90 min and not at 60 min. Multi-institutional trials using a higher performance for predicting ENS than the use of either parameter alone, with 89% sensitivity, 92% specificity, and 92% accuracy. Therefore, these results suggest that measuring TLG-PS in addition to SUV-LN on pretreatment \( ^{18} \text{F-FDG PET/CT} \) could be both feasible and useful to predict LNs with ENS in patients with OSCC.

The second finding of our study is that the optimal cut-off value of SUV-LN was 3.37, with 100% sensitivity, 84% specificity, and 86% accuracy in predicting ENS, and an AUC value of 0.932 ± 0.05. In a previous study \([4]\) of OSCC patients with 71 dissected LN levels, the SUV-LN in the tumors with ENS was greater than that in those without ENS, and the difference was significant. The ROC curve analysis of SUV-LN in the same study showed an AUC of 0.864 ± 0.045 and an optimal SUVmax cut-off value of 2.25, with 85% sensitivity and 86% specificity. In our study, we achieved excellent diagnostic capacity by setting the SUV-LN cut-off value at 3.37. In clinical practice, it is relatively easy to detect LNs with an SUVmax greater than 3.37, even when surrounded by adjacent normal tissue with physiological uptake. The difference in diagnostic performance of SUV-LN between our study and the previous study \([4]\) could be due to the difference in the PET acquisition time point. PET image acquisitions were started at 90 min after injection of \( ^{18} \text{F-FDG} \) in the present study and at 60 min after injection in the previous study \([4]\). Another previous research on 2-h dual-time-point \( ^{18} \text{F-FDG PET} \) demonstrated that head and neck malignant tumors had a mean SUV increase of 12% between the 1-h (early) and the 2-h (late) phases, and that the \( ^{18} \text{F-FDG} \) uptake to inflammation and normal tissue had meanwhile decreased \([22]\). Another recent report revealed that the \( ^{18} \text{F-FDG} \) uptake to most normal cervical tissue decreases as time proceeds within 3 h after \( ^{18} \text{F-FDG} \) injection \([23]\). These results suggest that 90-min PET acquisition with SUV-LN measurement is superior to 60-min PET acquisition for the prediction of ENS in OSCC, and that the contrast between metastatic LNs with ENS and surrounding normal tissue might be better on 90-min than 60-min PET images.

This study has several limitations. First, it is inherently limited by its retrospective nature. Second, the study population was relatively small, especially in terms of patients having tumors with ENS. Finally, PET image acquisition was performed only at 90 min and not at 60 min. Multi-institutional trials using a
larger patient population with 90-min dual-time-point 18F-FDG PET/CT may provide a clearer picture and more confident assessment of the ability of 90-min 18F-FDG PET/CT to predict ENS in patients with OSCC.

In conclusion, the findings of the present study demonstrate that the assessment of 18F-FDG MTV-PS and TLG-PS could be a useful and highly accurate tool for predicting ENS in patients with OSCC. The 90-min 18F-FDG PET/CT is potentially more useful than the 60-min 18F-FDG PET/CT for predicting ENS by evaluating the 18F-FDG accumulation to LNs. Specifically, on 90-min 18F-FDG PET/CT, a TLG-PS cut-off value of 106.26 g, combined with a SUV-LN cut-off value of 3.37, demonstrated excellent diagnostic performance in identifying lymph nodes with ENS in OSCC patients.

References