18F-FDG PET/CT Metabolic Tumor Volume and Total Lesion Glycolysis Predict Outcome in Oropharyngeal Squamous Cell Carcinoma

Remy Lim1, Anne Eaton2, Nancy Y. Lee3, Jeremy Setton3, Nisha Ohri3, Shyam Rao3, Richard Wong4, Matthew Fury5, and Heiko Schöder1

1Molecular Imaging and Therapy Service, Department of Radiology, Memorial Sloan-Kettering Cancer Center, New York, New York; 2Department of Biostatistics, Memorial Sloan-Kettering Cancer Center, New York, New York; 3Department of Radiation Oncology, Memorial Sloan-Kettering Cancer Center, New York, New York; 4Head and Neck Service, Department of Surgery, Memorial Sloan-Kettering Cancer Center, New York, New York; and 5Head and Neck Oncology Service, Department of Medicine, Memorial Sloan-Kettering Cancer Center, New York, New York

Treatment of oropharyngeal squamous cell carcinoma with chemoradiotherapy can now accomplish excellent locoregional disease control, but patient overall survival (OS) remains limited by development of distant metastases (DM). We investigated the prognostic value of staging 18F-FDG PET/CT, beyond clinical risk factors, for predicting DM and OS in 176 patients after definitive chemoradiotherapy. Methods: The PET parameters maximum standardized uptake value (SUVmax), metabolic tumor volume (MTV), and total lesion glycolysis (TLG) were recorded. Univariate Cox regression was used to examine the prognostic value of these variables and clinical prognosticators for local treatment failure (LTF), OS, and DM. Multivariate analysis examined the effect of SUVmax, TLG, and MTV in the presence of other covariates. Kaplan–Meier curves were used to evaluate prognostic values of PET/CT parameters. Results: Primary tumors were distributed across all stages. Most patients underwent chemoradiotherapy only, and 11 also underwent tonsillectomy. On univariate analysis, primary tumor MTV was predictive of LTF (P = 0.005, hazard ratio [HR] = 2.4 for a doubling of MTV), DM and OS (P < 0.001 for both, HR = 1.9 and 1.8, respectively). The primary tumor TLG was associated with DM and OS (P < 0.001, HR = 1.6 and 1.7, respectively, for a doubling of TLG). The primary tumor SUVmax was associated with death (P = 0.029, HR = 1.1 for a 1-unit increase in standardized uptake value) but had no relationship with LTF or DM. In multivariate analysis, TLG and MTV remained associated with death after correcting for T stage (P = 0.0125 and 0.0324, respectively) whereas no relationship was seen between standardized uptake value and death after adjusting for T stage (P = 0.158). Conclusion: Parameters capturing the volume of 18F-FDG–positive disease (MTV or TLG) provide important prognostic information in oropharyngeal squamous cell carcinoma treated with chemoradiotherapy and should be considered for risk stratification in this disease.
tion than SUVmax and whether these indices have incremental prognostic value beyond widely accepted clinical factors for DM and death.

**MATERIALS AND METHODS**

**Patient Population and Treatment Regimens**

This retrospective analysis was approved by our Institutional Review Board; patient consent was not required. Our analysis was restricted to patients with 18F-FDG PET/CT results (rather than PET-only results) because combined PET/CT improves the anatomic localization of abnormalities and accuracy of scan interpretation (12). To ensure relative uniformity in technique and quality of scans, those patients having undergone 18F-FDG PET/CT elsewhere before referral to our institution were excluded. Patients with DM at presentation noted on conventional imaging or staging 18F-FDG PET/CT were also excluded. All patients were uniformly treated with IMRT to 70 Gy (simultaneous integrated boost), and most received concurrent chemotherapy. Single-agent cisplatin was the primary choice of chemotherapeutic agent (2–3 cycles of a 100 mg/m² dose of cisplatin on days 1, 22, and 43). Some individuals were treated with other regimens (Table 1). After completion of therapy, patients were assessed by clinical examination and PET/CT at approximately 10–12 wk. Thereafter, clinical examinations and CT or MRI of the neck were performed every 2–3 mo for the first 2 y and every 4–6 mo thereafter (3). The occurrences of local treatment failure (LTF), DM, and death were recorded from the medical records.

**18F-FDG PET/CT**

In preparation for PET/CT, patients fasted for 6 h, but water intake was encouraged. After a blood glucose level of less than 200 mg/dL (median, 95 mg/dL; mean, 99 mg/dL; range, 55–198 mg/dL) was confirmed, 18F-FDG (425.5–573.5 MBq [11.5–15.5 mCi]) was injected intravenously, followed by a 72 ± 17 min uptake period, during which patients drank diluted oral contrast. Low-dose CT (120–140 kV, 80 mA) and PET scans were then obtained for the torso (3 min/bed position, thoracic inlet to upper thigh) with the arms up, followed by dedicated images of the head and neck (5 min/bed position) with the arms down. For radiotherapy planning scans, intravenous contrast was also administered. All images were reviewed on a workstation integrated with a PACS (Volume Viewer 2, AW Suite, version 2.0; GE Healthcare) that allowed multiplanar reformating of images.

**18F-FDG PET Image Analysis**

Volumetric regions of interest were placed over areas of abnormal 18F-FDG uptake. Initially, SUVmax (maximum voxel intensity within the volumetric region) was recorded for the primary tumor and neck lymph nodes. Region-of-interest borders were set by manual adjustment in 3 planes to conform to nodal levels in the neck and to exclude adjacent 18F-FDG–avid structures. Nodal disease was recorded for each neck level (13). A threshold of 42% of the maximum signal intensity (14) was used to delineate the metabolic tumor volume (MTV). Total lesion glycolysis (TLG) was calculated as the product of lesion mean standardized uptake value (SUV) and MTV (15).

**Statistical Analysis**

Missing uptake data were imputed for 3 patients with no measurable 18F-FDG uptake in primary tumors (2 with tonsillectomy before PET/CT and 1 with unknown primary) using cold deck imputation with the mid-point method (16). Eighteen individuals without nodal disease on 18F-FDG PET/CT had values for nodal SUVmax, MTV, or TLG imputed analogously. For analysis of lymph nodes, we used the highest SUVmax (respectively, the highest MTV or TLG) in any neck level. In the cases with bilateral neck disease, only the highest SUVmax (respectively, the highest MTV or TLG) in the neck level was used. MTV and TLG were found to be right-skewed and were log-transformed to better fit a normal distribution. Descriptive statistics for PET/CT variables are based on the preimputation data. Outcomes recorded included time to death, DM, and LTF measured from the start of radiotherapy. Follow-up for LTF was censored at the time of DM or neck recurrence. We examined the association between SUVmax, MTV, and TLG for primary tumor and metastatic lymph nodes and clinical variables (demographics, location of primary tumor, T and N stages, nodal spread beyond level 2) and patient outcome using univariate Cox proportional hazards regression. The effect of tobacco...
exposure was studied using both smoking status (current, former, and never) and pack-years (less than 20 and 20 or more; the latter is not included in final models because it was not significant on its own and did not alter the effect of smoking status). Lymph node PET parameters were not examined as predictors of LTF because an association seemed implausible. We further explored the relationship between PET/CT variables, DM, and overall survival (OS) using Kaplan–Meier curves stratified by quartiles of SUVmax, MTV, and TLG and the log-rank test.

The small number of events in this dataset limited the number of factors we could adjust for in multivariate modes. Because T stage is a strong prognostic variable and may hold some of the same information as the PET parameters, we built OS models with T stage and each PET parameter that satisfy the widely accepted criterion of requiring 10–15 events per variable in a multivariate model. To assess the ability of different models to discriminate between high- and low-risk individuals, the concordance probability estimate (CPE) method was used to estimate the concordance probability for various models (17). The optimal cutoff for primary tumor MTV as a predictor of survival was chosen using the maximally selected log-rank statistic (18). A 2-sided P value of less than 0.05 was considered significant. All analyses were performed in SAS 9.2 (SAS Institute Inc.) and R 2.11.1 (The R Foundation for Statistical Computing).

RESULTS
Patients and Treatment
An institutional database showed that 442 patients with OPSCC underwent chemoradiotherapy between September 1998 and May 2009. Of these, 266 patients were not considered eligible because 18F-FDG imaging had been performed as PET-only (rather than as PET/CT), or PET/CT had been performed at outside institutions. All of the remaining 176 patients underwent initial-staging 18F-FDG PET/CT and therapy at our institution in 2002 or later and form the basis for this study (Table 1). Eleven patients underwent tonsillectomy in addition to chemoradiotherapy (3 of them before PET/CT; in 2 cases, the subsequent PET/CT scan still showed abnormal uptake in the tonsillar fossa). Twenty-five patients underwent neck dissection after chemoradiotherapy for suspected residual disease; only 5 showed minimal residual disease on histopathology. The mean time interval between PET/CT and the start of therapy was 3 wk (median, 2.4 wk).

18F-FDG PET/CT Findings
The median (±SD) primary tumor SUVmax, MTV, and TLG were 13.6 ± 5.5 g/mL (range, 3.4–28.2 g/mL), 9.7 ± 12.9 cm³ (range, 1.2–113.0 cm³), and 81.4 ± 191.8 g (range, 5.4–1,964.4 g), respectively. The median (±SD) SUV, MTV, and TLG for all metastatic lymph nodes were 8.3 ± 4.6 g/mL (range, 2.4–33.3 g/mL), 5.0 ± 8.4 cm³ (range, 0.5–60.8 cm³), and 23.1 ± 70.9 g (range, 2.7–442.7 g), respectively. PET parameters for the various neck node levels are shown in Supplemental Table 1 (supplemental materials are available online only at http://jnm.snmjournals.org). TLG and MTV were highly correlated, and TLG and SUV were moderately correlated. Correlation between MTV and SUV was weak (Supplemental Table 2).

Patient Outcome
There were 137 survivors during the study period, of whom 125 (91%) were followed for more than a year and 99 (72%) were followed for more than 2 y. The median follow-up period was 41.0 mo (range, 6.5–80.9 mo) for surviving patients. Over the course of the study period, local failure occurred in 13 of 176 patients. Two of these 13 local failures were censored for analysis because of earlier regional or distant recurrences. The estimated 2-y local failure rate was 7.6% (95% confidence interval [CI], 4.2%–13.4%). DM occurred in 32 of 176 patients, and the estimated 2-y DM rate was 17.9% (95% CI, 12.7%–24.8%). In total, 39 patients died. Twenty-five patients had a documented DM at the time of death, 3 patients died of a second primary malignancy, 2 died from progressive locoregional disease, and the remainder died from cardiovascular, neurologic, or infectious diseases; the cause of death remained unclear in 3 patients. The estimated OS at 2 y was 83.9% (95% CI, 77.2%–88.8%).

Univariate Analysis
Univariate Cox analysis investigated the association between clinical parameters; 18F-FDG PET/CT variables; and LTF, OS, and DM. These results are shown in Table 2. Among clinical parameters, advanced T stage was associated with local failure (P = 0.014) and poor OS (P < 0.001), whereas advanced N stage was associated with poor OS (P = 0.032) and an increased hazard of DM (P = 0.014). In addition, age was associated with LTF. Smoking status was not associated with LTF, DM, or OS (all P > 0.20). Primary tumor MTV and TLG were found to be strong predictors of DM and OS (P < 0.001). A doubling of MTV resulted in a 1.8-fold increase in the hazard of death and a 1.9-fold increase in the hazard of DM. Similarly, a doubling of TLG resulted in a 1.7-fold increase in the hazard of death and a 1.6-fold increase in the hazard of DM. MTV and TLG were also associated with LTF, with hazard ratios (HRs) of 2.4 and 1.9, respectively, for doubling. Primary tumor SUVmax was associated with death (an HR of 1.1 for a 1-unit increase in SUVmax) but did not have an association with DM or LTF. A spread of metastatic disease beyond neck level 2 (defined as 18F-FDG–positive lymph node on PET/CT) resulted in a 2.2-fold increased hazard of death and a 2.5-fold increased hazard of DM.

We next analyzed the relationship between PET/CT variables and DM and OS stratified by quartiles of each variable; Kaplan–Meier curves are shown in Figures 1 and 2 for MTV and TLG and in Supplemental Figure 1 for SUVmax. Results from the log-rank test comparing quartile groups were similar to those observed with Cox models. There was a weak trend toward worse survival with increasing SUVmax (log-rank P = 0.085); TLG quartile was associated with OS (P = 0.018); and MTV quartile was associated with both DM (P = 0.007) and OS (P = 0.027). Because patients in the highest quartile of MTV
had notably worse survival than patients in the other 3 groups, we considered using MTV to define 2 groups rather than 4. To find the optimal MTV for separating patients with high hazard of death, we used a maximally rather than 4. To find the optimal MTV for separating groups, we considered using MTV to define 2 groups had notably worse survival than patients in the other 3

<table>
<thead>
<tr>
<th>Variable</th>
<th>n</th>
<th>OS HR</th>
<th>OS HR 95% CI</th>
<th>OS P</th>
<th>DM HR</th>
<th>DM HR 95% CI</th>
<th>DM P</th>
<th>LTF HR</th>
<th>LTF HR 95% CI</th>
<th>LTF P</th>
</tr>
</thead>
<tbody>
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<td>Age &gt; 70 y</td>
<td>176</td>
<td>2.24</td>
<td>0.99–5.08</td>
<td>0.054</td>
<td>1.29</td>
<td>0.45–3.69</td>
<td>0.632</td>
<td>5.42</td>
<td>1.58–18.62</td>
<td>0.007</td>
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<tr>
<td>T stage</td>
<td>176</td>
<td>&lt;0.001</td>
<td>1.00–0.86</td>
<td>0.16</td>
<td>1.06</td>
<td>0.96–1.17</td>
<td>0.263</td>
<td>0.21</td>
<td>0.03–1.76</td>
<td>0.014</td>
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<td>1 vs. 4</td>
<td>0.09</td>
<td>0.01–0.65</td>
<td>0</td>
<td>Not applicable, no events</td>
<td>1.54</td>
<td>0.113</td>
<td>1.89</td>
<td>0.641</td>
<td>0.21</td>
<td>0.36–1.54</td>
</tr>
<tr>
<td>2/3 vs. 4</td>
<td>0.32</td>
<td>0.17–0.62</td>
<td>0.42</td>
<td>0.21–0.86</td>
<td>0.62</td>
<td>0.64</td>
<td>1.08</td>
<td>0.11</td>
<td>0.36–1.54</td>
<td>0.113</td>
</tr>
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<td>Primary tumor SUV</td>
<td>176</td>
<td>1.06</td>
<td>1.01–1.12</td>
<td>0.029</td>
<td>1.03</td>
<td>0.97–1.09</td>
<td>0.406</td>
<td>1.06</td>
<td>0.96–1.17</td>
<td>0.263</td>
</tr>
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<td>Primary tumor MTV*</td>
<td>176</td>
<td>1.77</td>
<td>1.31–2.40</td>
<td>&lt;0.001</td>
<td>1.89</td>
<td>1.35–2.65</td>
<td>&lt;0.001</td>
<td>2.38</td>
<td>1.30–4.36</td>
<td>0.005</td>
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<td>Primary tumor TLG*</td>
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<td>1.66</td>
<td>1.29–2.14</td>
<td>&lt;0.001</td>
<td>1.56</td>
<td>1.20–2.04</td>
<td>&lt;0.001</td>
<td>1.90</td>
<td>1.17–3.11</td>
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<td>SUVmax in nodes</td>
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<td>1.00–1.11</td>
<td>0.033</td>
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<td>Maximum MTV in nodes*</td>
<td>176</td>
<td>1.05</td>
<td>0.86–1.28</td>
<td>0.641</td>
<td>1.21</td>
<td>0.96–1.54</td>
<td>0.113</td>
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<td>Maximum TLG in nodes*</td>
<td>176</td>
<td>1.11</td>
<td>0.94–1.32</td>
<td>0.217</td>
<td>1.25</td>
<td>1.03–1.52</td>
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<td>Level of spread in lymph nodes†</td>
<td>176</td>
<td>2.21</td>
<td>1.16–4.19</td>
<td>0.015</td>
<td>2.47</td>
<td>1.21–5.05</td>
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<td>Smoking status</td>
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<td>0.3102</td>
<td>0.9747</td>
<td>1.09</td>
<td>0.406</td>
<td>1.06</td>
<td>0.96–1.17</td>
<td>0.263</td>
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<tr>
<td>Former vs. never</td>
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<td>0.43–2.06</td>
<td>1.59</td>
<td>0.63–4.04</td>
<td>1.15</td>
<td>0.29–4.62</td>
<td>0.9747</td>
<td></td>
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<td></td>
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<tr>
<td>Current vs. never</td>
<td>1.78</td>
<td>0.77–4.13</td>
<td>2.24</td>
<td>0.80–6.29</td>
<td>1.02</td>
<td>0.17–6.11</td>
<td>0.9747</td>
<td></td>
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<td></td>
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</table>

*HR represents doubling of covariate value.
†Greater than level 2 vs. levels 1, 2, or no spread.
95% CI = 95% confidence interval.

DISCUSSION

In the largest series to date, we show that 18F-FDG PET/CT indices of metabolic tumor burden (MTV, TLG) show a strong correlation with the development of DM or death in patients with OPSCC treated with IMRT and concurrent chemotherapy.

Prior studies evaluating the prognostic value of 18F-FDG PET in head and neck cancer mostly applied SUVmax, as the biomarker of choice, because it is commonly used in clinical practice; many of these studies noted a prognostic value of SUVmax (8, 19–21). For instance, in a study of 120 patients, primary tumor SUVmax correlated strongly with poor local control and disease-free survival (8). However, in some other studies, SUVmax was not associated with short-term outcomes or disease-free survival (9) and did not predict recurrence (22). A major confounding factor in all of these studies was the inclusion of tumors from various head and neck sites, with different tumor biology; differing
etiology, risk factors, and clinical presentation; and different overall prognosis (23). Moreover, the location of the primary tumor may mandate different treatment regimens. All of these factors may potentially skew the data analysis. Therefore, our study focused on a relatively large population of patients with primary OPSCC treated with IMRT and concurrent chemotherapy.

Our findings suggest that TLG may provide better prognostic information than SUVmax. This finding appears plausible because TLG considers the volume of metabolically active disease (in addition to the level of tumor glucose utilization). Although this is concordant with surgical data showing worse outcome with increasing primary tumor size and burden of nodal metastases, our comprehensive analysis circumvents the obvious limitations inherent to simple clinical size assessment or size measurements by imaging. First, traditionally T stage for oropharyngeal carcinoma takes into account only the greatest single dimension of the primary tumor (T1–T3) or infiltration of surrounding tissue (T4). In addition, neither clinical examination nor simple structural assessment can differentiate reliably between viable tumor, edema, and necrosis. In contrast, our data are based on the combined measurement of metabolically active tumor burden and glucose utilization in these lesions. Of note, studies in oral cavity and larynx cancer have shown that tumor volume derived from 18F-FDG PET/CT data correlates better with gross pathology volume than volumes measured on CT or MRI (24,25).

In our study, the estimated concordance probability for TLG and survival time was higher than the estimated concordance probability for SUVmax (0.69 vs. 0.59), and TLG similarly performed better for the DM outcome. In addition, in OS models adjusted for T stage, TLG yielded a higher estimated concordance probability than SUV. Although the differences did not reach significance, the notion that TLG appears to be a better prognostic marker than SUV alone is also supported by survival curves, which

![Figure 1](image1.png)  
**FIGURE 1.** Kaplan–Meier curves for PET/CT variable TLG for developing DM or death are stratified by quartiles of each variable. mets = metastases.

![Figure 2](image2.png)  
**FIGURE 2.** Kaplan–Meier curves for PET/CT variable MTV for developing DM or death are stratified by quartiles of each variable. mets = metastases.
show stronger separation of patient groups when stratified by TLG quartile than by SUV quartile (Figs. 1 and 2).

We found that an MTV greater than \(19.7\, \text{cm}^3\) places patients at high risk of death. Some other studies suggested MTV as a prognostic parameter. Interestingly, the median MTV varies from \(9.7\, \text{cm}^3\) (this study) to \(11.2\, \text{cm}^3\) (the study by La et al. \((26)\)). 23.5 \(\text{cm}^3\) (the study by Seol et al. \((27)\)), and 39 \(\text{cm}^3\) (the study by Chung et al. \((9)\)). In a study of 85 patients with pharyngeal squamous cell carcinoma (53% oropharynx, 26% nasopharynx), an MTV increase of \(17.4\, \text{cm}^3\) (the difference between the 75th and 25th centile) was prognostic for recurrence (composite of locoregional and distant disease) and death \((26)\). In another study of 59 patients (64% larynx, 22% oropharynx, and 14% hypopharynx), an MTV of greater than \(9.3\, \text{cm}^3\) was associated with increased risk of recurrence and death (composite of progression free and OS) \((27)\). Finally, in a study of 82 patients (77% nasopharynx, 16% oropharynx), patients with greater than median MTV had a worse disease-free survival \((9)\). There are 2 possible explanations for the wide variation in median MTV. First, there appears to be a general trend toward lower MTV with increasing proportion of patients with OPSCC. For example, the study reporting a median MTV of 39 \(\text{cm}^3\) \((9)\) included a large proportion of patients with nasopharynx cancers, which may be larger in size at detection than OPSCC by virtue of their location. Second, differences may also occur because of differences in threshold methods used for delineating MTV. We selected a threshold of 42\% for volume definition, based on prior phantom studies \((14)\), whereas other investigators used 50\% of maximum intensity \((26)\) or a predefined absolute SUV number \((28)\). Regardless of the chosen threshold, however, all studies showed MTV as an indicator of poor outcome.

Advanced cervical nodal stage \((29)\) and level of neck node involvement \((30,31)\) are recognized risk factors for the development of DM. In our study, using PET/CT-based staging, we confirmed these earlier data from surgical series. Nearly 90\% of our cohort demonstrated nodal involvement by PET, concurring with historic studies showing nodal metastases in neck dissection specimens in up to 84\% of patients with OPSCC, mostly in levels 2–4 \((30)\). We found that the hazard of DM and death increased significantly with higher N stage and more than doubled when \(^{18}\text{F}-\text{FDG}\)-positive nodes were seen beyond neck level 2, confirming advanced level of nodal involvement as an adverse prognostic factor \((30–33)\).

Our study has some limitations. Although histopathology was available for all primary tumors, we did not have histologic confirmation for all lymph nodes but surmised that all \(^{18}\text{F}-\text{FDG}\)-positive nodes in typical distribution for OPSCC were indeed metastatic. We excluded a substantial proportion of patients whose pretreatment PET/CT was performed elsewhere because of the potential wide variability in scan techniques and quality. However, all patients who underwent PET/CT for staging and subsequent chemoradiotherapy at our institution since 2002 were included; there was no selection bias. Patients were imaged on several state-of-the-art PET/CT systems, which show a less than 1\% error in SUV in large, homogeneous \(^{18}\text{F}-\text{FDG}\) phantoms. We also used comparable reconstruction parameters (subsets, number of iterations, and filters) for all scanners. We therefore believe that fluctuations in SUV derived from different scanners are well within the statistical error of SUV calculated from repeated measurements on the same scanner. The low rate of LTF limited our evaluation; however, we do not believe that LTF is an important driving force of survival, because most patients appear to die from DM. Additionally, because of the limited number of deaths observed, we were unable to perform risk stratification for each T stage separately. In

![Overall survival by MTV](image)

**FIGURE 3.** Kaplan–Meier curve illustrating prognostic value of primary tumor MTV for OS. This optimal MTV for identifying patients at high hazard of death is \(19.7\, \text{cm}^3\). RT = radiotherapy.

### TABLE 3
Association Between PET Variables and OS Adjusted for T Stage

<table>
<thead>
<tr>
<th>Variable</th>
<th>HR</th>
<th>HR 95% CI</th>
<th>(P)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Model 1—SUV</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>T stage</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 vs. 4</td>
<td>0.11</td>
<td>0.01–0.89</td>
<td>0.0013</td>
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<tr>
<td>2/3 vs. 4</td>
<td>0.33</td>
<td>0.17–0.63</td>
<td></td>
</tr>
<tr>
<td>Primary tumor SUV</td>
<td>1.04</td>
<td>0.98–1.10</td>
<td>0.1577</td>
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<tr>
<td><strong>Model 2—MTV</strong></td>
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<td></td>
<td></td>
</tr>
<tr>
<td>T stage</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1 vs. 4</td>
<td>0.17</td>
<td>0.02–1.43</td>
<td>0.0485</td>
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<tr>
<td>2/3 vs. 4</td>
<td>0.45</td>
<td>0.22–0.91</td>
<td></td>
</tr>
<tr>
<td>Primary tumor MTV*</td>
<td>1.45</td>
<td>1.03–2.03</td>
<td>0.0324</td>
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<td><strong>Model 3—TLG</strong></td>
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<tr>
<td>T stage</td>
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<tr>
<td>1 vs. 4</td>
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<td>2/3 vs. 4</td>
<td>0.45</td>
<td>0.22–0.91</td>
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<td>Primary tumor TLG*</td>
<td>1.43</td>
<td>1.08–1.89</td>
<td>0.0125</td>
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*HR represents doubling of covariate value.
95% CI = 95% confidence interval.
OS models adjusted for T stage, TLG was the PET parameter that provided the best discrimination. However, when stratifying by quartiles of risk based on the model, the second and third quartiles did not differ significantly with regard to survival (Fig. 4). Therefore, whereas the model successfully identified a group of patients with a high risk of death and a group with low risk, it showed less discriminatory power among patients with intermediate risk, possibly related to sample size. A small number of patients in our study had a short follow-up period; however, because we used survival methods to account for follow-up time and the largest part of the cohort (72%) was followed beyond the 2-y high-risk window, these should not affect our results. Human papilloma virus (HPV) infection status may influence prognosis in OPSCC (34). Because HPV status was not routinely assessed in our institution during the study period, this factor could not be entered in the multivariate analysis. Instead, we applied smoking status as a surrogate marker because it appears inversely related to HPV tumor status (34). Of note, smoking status was not a significant parameter in multivariate analysis. Quantitative PET measurements depend on technical parameters of data acquisition and processing, including the segmentation algorithm and threshold chosen for MTV and TLG. Our choice of a 42% threshold was based on prior phantom and modeling studies (14,35). Although this threshold may not be optimal for all lesion sizes and target-to-background ratios, we believe that our results are valid: applying a nonoptimum threshold for a group of lesions with a given size range may result in uncertainty of the molecular target volume but would not be expected to cause a systematic reshuffling of data between the 4 groups (based on the lesion size) defined in the study. Finding an optimum PET threshold and method for data segmentation is the subject of ongoing research. This uncertainty does not undermine the concept that MTV has prognostic value. However, although we found a worse outcome for patients with a primary MTV greater than 19.7 cm³, this specific number and the chosen cut points for TLG require validation in future studies and may vary between institutions depending on the local PET protocol. Nevertheless, the general association of high MTV and TLG with poor outcome would still hold true. Finally, future studies will have to compare the prognostic value of MTV and TLG with that of 3-dimensional tumor volumetric measurements from CT or MRI. Although these data are currently not available, we expect that MTV and TLG will retain their prognostic significance, because they are not simple measurements of tumor size (albeit in 3 dimensions) but instead are based on glucose metabolism, an important hallmark of the malignant phenotype.

CONCLUSION

Metabolically active tumor volume, captured by PET indices of MTV or TLG, is an important prognostic factor in OPSCC, provides incremental prognostic information beyond clinical T stage, and should be considered for risk stratification in these patients.

DISCLOSURE STATEMENT

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REFERENCES


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18F-FDG PET/CT Metabolic Tumor Volume and Total Lesion Glycolysis Predict Outcome in Oropharyngeal Squamous Cell Carcinoma

Remy Lim, Anne Eaton, Nancy Y. Lee, Jeremy Setton, Nisha Ohri, Shyam Rao, Richard Wong, Matthew Fury and Heiko Schöder

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