Pre-Therapeutic Total Lesion Glycolysis on \([^{18}F]FDG\)-PET Enables Prognostication of 2-Year Progression-Free Survival in MALT Lymphoma Patients Treated with CD20-Antibody-Based Immunotherapy

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Abstract

Purpose: Standardized uptake values (SUV), total metabolic tumor volumes (TMTV), and total lesion glycolysis (TLG) based on positron emission tomography with 2-deoxy-2-[\(^{18}\)F]fluoro-D-glucose ([\(^{18}\)F]FDG/positron emission tomography (PET) are established outcome predictors in FDG-avid lymphomas. We therefore investigated whether these biomarkers also have prognostic value in extranodal marginal zone B cell lymphoma of the mucosa-associated lymphoid tissue (MALT lymphoma), with a focus on patients treated with anti-CD20 antibody-based immunotherapy.

Procedures: Pre-therapeutic \([^{18}\)F]FDG/PET scans of 61 treatment-naïve MALT lymphoma patients, including 35 scheduled for anti-CD20 antibody-based immunotherapy, were included in this retrospective study. SUVmean, SUVmax, TMTV, and TLG were measured and tested for 2-year progression-free survival (PFS) prognostication, using Cox regression analyses. Receiver operating characteristic curves were used to determine optimal cutoffs for prognostic \([^{18}\)F]FDG/PET parameters, and Kaplan–Meier estimates with log rank tests were performed.

Results: After 2 years, progression had occurred in 12/61 patients (CD20-antibody group 6/35). TLG emerged as the only significant prognostic factor for 2-year PFS in the multivariate analyses with forward selection, both in entire cohort (hazard ratio HR, 1.001; 95 % CI, 1.001–1.002; \(P<0.0001\)) and in the CD20-antibody group (HR, 1.001; 95 % CI, 1.001–1.002; \(P=0.001\)). However, in the entire population, where 8/26 patients with a TLG > 90 (30.8 %) vs. 4/35 patients with a TLG ≤ 90 (11.4 %) showed...
Introduction

Outcome prognostication and identification of risk groups play an important role in the management of patients with malignant lymphoma. An increasing number of recent studies suggest that, for patients with classical Hodgkin, diffuse large B cell (DLBCL), follicular, as well as T cell lymphomas, quantitative parameters derived from pre-therapeutic positron emission tomography with 2-deoxy-2-[18F]fluoro-D-glucose ([18F]FDG)/positron emission tomography (PET) — such as maximum standardized uptake value (SUV max), total metabolic tumor volume (TMTV), or total lesion glycolysis (TLG) — are prognostic for clinical outcomes, and may therefore improve risk stratification [1–5].

For extranodal marginal zone B cell lymphoma of the mucosa-associated lymphoid tissue (MALT lymphoma), only three studies have, so far, evaluated quantitative pre-therapeutic [18F]FDG/PET-derived parameters for outcome prognostication [6–8], probably because of the variable FDG avidity of this lymphoma subtype, due to which [18F]FDG/PET is not recommended for imaging of MALT lymphomas by the International Conference on Malignant Lymphoma (ICML) [9]. Notably, none of the these studies — including the single, very recently published study that included an evaluation of TMTV and TLG, and was unable to establish either parameter as a prognostic factor for overall survival (OS) or progression-free survival (PFS) [6] — focused on a specific treatment group, contrary to studies in other lymphoma subtypes. Instead, therapeutic approaches in the investigated MALT lymphoma populations ranged from antibiotics and radiation therapy to surgery and chemotherapy — a heterogeneity that may have influenced the results of the respective studies. Apart from antibiotics, CD20-antibody-based immunotherapy is the most commonly used systemic treatment for MALT lymphoma, and has been shown to carry significant anti-tumor activity in untreated as well as relapsed MALT lymphoma [10].

In the present study, it was therefore our aim to determine whether pre-therapeutic [18F]FDG/PET-derived quantitative parameters, i.e., SUV, TMTV, and TLG, carry prognostic information, with regard to 2-year PFS, in (1) MALT lymphoma patients treated with CD20-antibody-based immunotherapy, as compared to (2) a general population of MALT lymphoma patients receiving different types of treatment.

Materials and Methods

Patients and Design

Patients with histologically proven, treatment-naïve MALT lymphoma, who had undergone [18F]FDG/PET/CT (x-ray computed tomography) or [18F]FDG/PET/MRI (magnetic resonance imaging) for routine pre-therapeutic staging at the local tertiary care center between 2008 and 2015, had shown at least some degree of [18F]FDG avidity (equivalent to Deauville score of ≥2 [11]), and for whom a follow-up over a period of ≥2 years after diagnosis/imaging, or until documented progression, was available, were included in our retrospective study. The study was performed in accordance with the ethical standards as laid down in the 1964 Declaration of Helsinki and its later amendments. Ethics committee approval was obtained; for this type of study (i.e., with a retrospective design), formal consent is not required. The histological diagnosis of MALT lymphoma was made by a reference pathologist who analyzed tissue samples (obtained by biopsy or surgery) according to the 2016 revision of the World Health Organization classification of lymphoid neoplasms. According to our institutional standard of care, the decision to perform [18F]FDG/PET was independent of age, performance status, laboratory findings, disease stage or localization, or treatment strategy.

Imaging Protocols

For both [18F]FDG/PET/CT and [18F]FDG/PET/MRI, PET was performed 60 min after intravenous administration of a target dose of 3 MBq/kg (minimum injection dose, 200 MBq) of [18F]FDG. All patients had fasted for at least 5 h before imaging; the upper blood glucose level limit was 150 mg/dl.

Whole-body [18F]FDG/PET/CT was performed using a 64-row, multi-detector hybrid scanner (Biograph TruePoint 64; Siemens, Erlangen, Germany) with an axial field-of-view (FOV) of 216 mm, and a PET sensitivity of 7.6 cps/kBq. PET was performed with 3 min per bed position, four iterations and 21 subsets, a 5-mm slice thickness, and a 168 × 168 matrix, using the point spread function (PSF)-based reconstruction algorithm TrueX. Contrast-enhanced venous-phase CT was used for attenuation correction, and
used to assist with lesion delineation. CT or DWI components of PET/CT or PET/MRI could be
surrounding tissues (e.g., with Deauville scores 2 or 3), the CT or DWI components of PET/CT or PET/MRI could be
used to assist with lesion delineation.

Post-reconstruction harmonization of SUVs, using the previously described ComBat method [13], was performed
to correct for technical differences between the PET/CT and PET/MRI scanners that are known to affect SUVs [14].
Finally, TLG (product of TMTV and SUVmean) [15] was calculated.

Statistical Analysis

Descriptive statistics for demographic, clinical, and imaging parameters were calculated. Independent sample $t$ tests were
used to test for significant differences of the imaging parameters between gastric and non-gastric MALT lymphomas. Univariate Cox regression analyses were performed to model the influence of demographic, clinical, and imaging parameters on the PFS. For all parameters that demonstrated statistical significance in the univariate analysis, a multivariate Cox regression analysis with forward selection (based on the likelihood ratio) was performed.

Receiver operating characteristic (ROC) curves were constructed, and used to determine optimal cutoffs for those
parameters on the PFS. For all parameters that demonstrated statistical significance in the univariate analysis were calculated, using Kaplan–Meier estimates, and log rank tests were used for group comparisons.

The above evaluation was performed for the entire patient cohort (regardless of the type of treatment received) on the one hand, and then also separately for the subgroup of patients that received CD20-antibody-based immunotherapy. All statistical tests were performed using IBM SPSS 23.0 (Armonk, NY, USA).

Results

Sixty-one consecutive patients (37 women and 24 men; mean age, 66.5 ± 12.6 years) met our criteria for participation in the
study. Fifty-two patients had undergone pre-therapeutic $[^{18}F]$FDG/PET using contrast-enhanced PET/CT, and nine patients using PET/MRI. Low $[^{18}F]$FDG uptake (≤ mediastinum; equivalent to Deauville 2) was observed in 12 patients (18.7 %); uptake higher than that of the mediastinum, but not exceeding that of the liver (equivalent to Deauville 3) was observed in 11 patients (17.2 %); and uptake greater than that of the liver (equivalent to Deauville 4 or 5) was observed in the remaining 41 patients (64.1 %). Involved anatomic regions/ organs are listed in Table 1.

Thirty-five patients received CD20-antibody-based immuno
therapy (rituximab, 27; ofatumumab, eight); seven received clarithromycin (one in combination with lenalidomide); five exclusively underwent helicobacter pylori eradication; four were treated with radiotherapy; two underwent surgery; and one patient each was treated with chlorambucil, Zevalin, R-CHOP (rituximab, cyclophosphamide, doxorubicin, vincristine, and prednisone), R-BENDA (rituximab, bendamustine), or R-2CD (rituximab, chlorambine). In three patients, a watch-and-wait strategy was used.

Entire Cohort (All Treatments)

After 2 years, progression had occurred in 12/61 patients (19.67 %), including three deaths. Descriptive data for
demographic, clinical, and imaging parameters and results of the univariate Cox regression analyses are provided in
Table 2. Gastric ($n=15$) and non-gastric ($n=46$) MALT lymphomas did not differ with regard to SUVmean (mean,
3.79 ± 1.56 vs. 3.68 ± 1.85; P = 0.84), SUVmax (mean, 7.62 ± 5.62 vs. 7.22 ± 4.17; P = 0.77), TMTV (mean, 88.17 ± 129.56 vs. 72.30 ± 200.59; P = 0.78), or TLG (mean, 472.22 ± 1009.59 vs. 224.79 ± 512.03; P = 0.22).

None of the demographic or clinical parameters, but all [18F]FDG/PET-derived parameters except SUVmean were significant at univariate analysis. In the multivariate analysis, only TLG (hazard ratio HR, 1.001; 95 % CI, 1.001–1.002; P < 0.0001), but not SUVmax (P = 0.85) or TMTV (P = 0.27), retained the significant association with PFS.

The ROC analysis revealed an optimal TLG cutoff value of 90 for separation of patients who achieved, and those who did not achieve, 2-year PFS; the AUC was 0.68 (95 % CI, 0.50–0.87; P = 0.05) (Fig. 2). Based on this threshold, 8/26 patients with a TLG > 90 (30.8 %) vs. 4/35 patients with a TLG ≤ 90 (11.4 %) showed progression within the 2-year observation period (HR, 0.35; 95 % CI, 0.10–1.15; P = 0.069) (Fig. 3).

**CD20-Antibody Group**

After 2 years, progression in this treatment group had occurred in 6/35 patients (17.1 %), including one death. Descriptive data for demographic, clinical, and imaging parameters and results of the univariate Cox regression analyses are provided in Table 3. Similar to the entire cohort, gastric (n = 7) and non-gastric (n = 28) MALT lymphomas did not differ with regard to SUVmean (mean, 4.02 ± 1.98 vs. 3.89 ± 1.57; P = 0.53), SUVmax (mean, 9.21 ± 7.92 vs. 7.42 ± 3.99; P = 0.40), TMTV (mean, 124.32 ± 176.11 vs. 61.68 ± 111.59; P = 0.25), or TLG (mean, 762.30 ± 1447.90 vs. 228.93 ± 411.30; P = 0.37).

TMTV (HR, 1.006; 95 % CI, 1.002–1.01; P < 0.0001) and TLG (HR, 1.002; 95 % CI, 1.001–1.003; P < 0.0001), but not SUVmean (P = 0.53) or SUVmax (P = 0.091), were prognostic for 2-year PFS at univariate analysis; whereas in the multivariate analysis, only TLG (HR, 1.001; 95 % CI, 1.001–1.002; P = 0.001), but not TMTV (P = 0.71), retained its significant association with PFS.

Similar to findings in entire cohort, the ROC analysis revealed an optimal TLG cutoff value of 90 for separation of patients who achieved, and those who did not achieve, the 2 year PFS; the AUC was 0.68 (95 % CI, 0.72–1.00; P =
Table 2. Demographic, clinical, and imaging data, and results of univariate Cox regression analyses for the 61 MALT lymphoma patients

| Characteristics                  | Frequency (%) or mean (± SD) | Univariate analysis of PFS |
|----------------------------------|-----------------------------|---------------------------|
|                                  |                             | HR (95 % CI)              | P value   |
| Age ≥ 70 years                   | 30 (49.2 %)                 | 1.038 (0.34–3.22)         | 0.94      |
| Ann Arbor stage                  |                             |                           |           |
| I                                | 36 (59.0 %)                 | 1                         |           |
| II                               | 10 (16.4 %)                 | 0.45 (0.10–2.01)          | 0.30      |
| III                              | 2 (3.3 %)                   | 1.44 (0.29–7.14)          | 0.66      |
| IV                               | 13 (21.3 %)                 | 5.10 (0.84–31.12)         | 0.077     |
| Plasmacytic differentiation      | 15 (24.6 %)                 | 1.14 (0.31–4.20)          | 0.85      |
| Elevated LDH                    | 5 (8.2 %)                   | 0.38 (0.08–1.73)          | 0.21      |
| Elevated β2 microglobulin       | 22 (36.1 %)                 | 0.81 (0.25–2.66)          | 0.73      |
| Missing                          | 6 (9.8 %)                   | 24.44 (0.12–49,496.68)    | 0.20      |
| ECOG ≥ 1                        | 7 (11.5 %)                  | 1.13 (1.004–1.23)         | 0.043     |
| SUVmax                           | 7.32 ± 4.52                 | 1.16 (0.86–1.56)          | 0.34      |
| SUVmean                          | 3.71 ± 1.77                 | 1.002 (1.001–1.004)       | 0.002     |
| TMTV (ml)                        | 76.20 ± 184.78              | 1.001 (1.001–1.002)       | < 0.0001  |
| TLG                              | 258.63 ± 667.84             |                           |           |

0.006) (Fig. 2). Based on this threshold, 6/16 patients with a TLG > 90 (37.5 %) vs. 0/19 patients with a TLG ≤ 90 (0.0 %) showed progression within the 2-year observation period (HR, 0.010; 95 % CI, 0.0001–8.068; P = 0.003) (Fig. 3).

Non-CD20-Antibody Group (All Other Treatments)

After 2 years, progression in this treatment group had occurred in 6/26 patients (23.1 %), including two deaths. Gastric (n = 8) and non-gastric (n = 18) MALT lymphomas did not differ with regard to SUVmean (mean, 3.59 ± 1.19 vs. 3.35 ± 2.23; P = 0.73), SUVmax (mean, 6.22 ± 2.14 vs. 6.91 ± 4.55; P = 0.60), TMTV (mean, 56.54 ± 67.36 vs. 88.81 ± 293.70; P = 0.66), or TLG (mean, 218.41 ± 289.55 vs. 218.34 ± 652.12; P = 1.0).

Only Ann Arbor stage (P = 0.039), but none of the PET-based parameters, and also none of the other clinical and laboratory parameters showed a significant association with PFS at univariate analysis, and therefore, no multivariate analyses were performed.

Discussion

There are two main conclusions that can be drawn from the results of our study. First, TLG may be a useful prognostic factor for 2-year PFS in patients treated with CD20-antibody-based immunotherapy, but not necessarily in an unselected population of MALT lymphoma patients receiving different types of treatment; here, contrary to the CD20-antibody cohort, TLG did not reach statistical significance for separation of groups at risk for progression (Fig. 3). Second, in accordance with previous findings in MALT lymphoma and DLBCL, TLG seems to be slightly superior to TMTV as a prognostic biomarker, and clearly superior to the clinical standard PET parameter, the SUVmax, in MALT lymphoma patients [3, 4, 6]. The latter may be explained by the variable degree of [18F]FDG avidity in this lymphoma subtype [9]—both metabolic tumor volume and the degree of glucose metabolism, which the TLG combines in a single measure, appear to carry prognostic information here.

Contrary to other lymphoma subtypes, there is currently no universally accepted standard treatment for MALT lymphoma [16]. For instance, limited disease may be treated with radiation therapy or surgery, and in gastric MALT lymphoma, eradication of H. pylori may even be sufficient. The randomized, multicenter IELSG-19 trial suggested that PFS and EFS (event-free survival) were comparable between MALT lymphoma patients that received CD20-antibody-based immunotherapy with rituximab (monotherapy) and those that received standard cytostatic chemotherapy with chlorambucil [17]. Unlike rituximab, the fully humanized, second-generation CD20-antibody ofatumumab can bind both the small and large extracellular loop of CD20 [18]; it has already been approved by the FDA (Food and Drug Administration) for the treatment of chronic lymphocytic leukemia, and has shown promise in different lymphoma subtypes, although head-to-head comparisons with rituximab in randomized controlled trials are presently still lacking [19]. In our study, TLG demonstrated a potential utility as a prognostic biomarker in the CD20-antibody group: the dichotomized TLG enabled the identification of a “low-risk population” in which not a single patient showed progression within the 2-year observation period, suggesting a very favorable response to this type of treatment (Fig. 2). We hypothesize that, in such a low-risk population, the use of longer time intervals between follow-up examinations may be an option.

The literature on the use of quantitative [18F]FDG/PET parameters in MALT lymphoma is scarce, and in part, contradictory. Qi, et al. recently reported, in a series of 123 patients with FDG-avid disease, that the SUV was an independent prognostic factor for 5-year OS, but not PFS [7]—the latter finding being partly in accordance with our
results. The patients included in this study had received different types of treatment: in Ann Arbor's stages I and II, they were predominantly treated with radiotherapy, with only 8% receiving immunotherapy not further specified; in stages III and IV, 47% received systemic treatment not further specified, and 36% underwent active surveillance. Contrary to these results, Albano et al. very recently reported, based on a cohort of 94 patients with [18F]FDG-avid MALT lymphoma, a significant association of TMTV and TLG, but not SUV, with PFS, albeit only in the univariate, but not the multivariate analysis; no imaging parameter was associated with OS [6]. Their cohort also consisted of a mixed population of MALT lymphoma patients treated with antibiotic therapy, surgery, radiotherapy, chemotherapy, or a combination thereof. The fact that none (Albano et al.), or only a small percentage (Qi, et al.), of the patients were treated with immunotherapy, limits the comparability between these study and our own. Nevertheless, the TLG cutoff of 84 in the study by Albano et al. was similar to that in our study (TLG > 90), and then again, in our own mixed population of 61 patients that received different types of treatment, TLG was also unable to separate groups at risk for progression, despite a trend in that direction.

We used PFS instead of overall survival (OS) as the clinical endpoint of our study, mainly because MALT lymphoma is a less frequently seen, although not uncommon, lymphoma subtype, with quite a good prognosis, compared to other non-Hodgkin lymphoma subtypes, such as mantle cell lymphoma. As a consequence, with PFS, smaller patient populations, as well as shorter observation periods, may be used to obtain a sufficiently large patient sample with the outcome of interest (i.e.,

![ROC curves for the four [18F]FDG-PET-derived quantitative parameters SUVmean, SUVmax, TMTV, and TLG with regard to separation of MALT lymphoma patients achieving, and those not achieving, 2-year PFS.](image1)

Fig. 2. Receiver operating characteristic (ROC) curves for the four [18F]FDG-PET-derived quantitative parameters SUVmean, SUVmax, TMTV, and TLG with regard to separation of MALT lymphoma patients achieving, and those not achieving, 2-year PFS.

![Kaplan-Meier estimates for the 2-year PFS suggest that risk stratification by total lesion glycolysis (TLG) is feasible in MALT lymphoma patients treated with CD20-antibody-based immunotherapy, but not in a more heterogeneous population of MALT lymphoma patients treated with different therapeutic regimens.](image2)

Fig. 3. Kaplan-Meier estimates for the 2-year PFS suggest that risk stratification by total lesion glycolysis (TLG) is feasible in MALT lymphoma patients treated with CD20-antibody-based immunotherapy, but not in a more heterogeneous population of MALT lymphoma patients treated with different therapeutic regimens.
progression). Furthermore, PFS is recognized by the FDA as a valid surrogate endpoint for non-Hodgkin lymphomas (https://www.fda.gov/downloads/Drugs/DevelopmentApprovalProcess/DevelopmentResources/UCM614369.xlsx). For DLBCL, a recent analysis of multiple randomized trials with a total sample size of 7507 patients also clearly supported the use of PFS, and showed that 2-year PFS significantly correlates with OS [20].

While our study is limited by its modest cohort size, this was a hypothesis-generating study, as [18F]FDG/PET is not considered a routine procedure for MALT lymphoma according to the ICML guidelines [9]. The fact that—contrary to our own institutional standard of care—CT is still the standard test for MALT lymphomas in the vast majority of institutions, despite its known limitations for treatment response assessment in lymphomas [21], prevented us from using a multi-centric approach that would have increased the number of study participants. For the survival analysis, we did not further subdivide our patient population (e.g., into gastric and non-gastric lymphomas), because only 12/61 patients overall, and 6/35 patients in the CD20-antibody group, showed progression. Finally, [18F]FDG/PET data were obtained from a PET/CT scanner on the one hand, and a PET/MRI scanner on the other hand, which probably affected SUV as well as SUV-based measurements. To correct for the latter, we applied a post-reconstruction harmonization technique that was specifically designed for this task [13]; still, remaining minor differences between the datasets of the two scanners cannot be completely ruled out.

**Conclusion**

In conclusion, our study results suggest that TLG may have prognostic meaning in MALT lymphoma patients treated with CD20-antibody-based immunotherapy. More specifically, TLG appears to have a high negative predictive value for 2-year PFS, and may, therefore, improve early risk stratification and, possibly, risk-adapted management of MALT lymphoma patients treated with CD20-antibody-based immunotherapy. For the general population of MALT lymphomas that receive different types of treatment, the prognostic value of [18F]FDG-PET-based quantitative parameters remains questionable. Larger patient cohorts are required to validate our results, and longer observation periods that will enable assessment of OS in addition to PFS.

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**Compliance with Ethical Standards.** All procedures performed in studies involving human participants were in accordance with the ethical standards of the institutional and/or national research committee and with the 1964 Helsinki declaration and its later amendments or comparable ethical standards.

**Conflict of Interest**

Marius E. Mayerhoefer received honoraria for lectures as well as research support (not related to the present study) from Siemens Healthineers. Barbara Kiesewetter and Markus Raderer received honoraria for lectures from Celgene, Ipsen, and Novartis.

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