Evaluation of Microscopic Tumour Extension in Localized Stage Non-Small-Cell Lung Cancer for Stereotactic Radiotherapy Planning

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Abstract: Background: Stereotactic radiotherapy for localised stage non-small-cell lung carcinoma (NSCLC) is an alternative indication for patients who are inoperable or refuse surgery. A study showed that the microscopic tumour extension (ME) of NSCLC varied according to the histological type, which allowed us to deduce adapted margins for the clinical target volume (CTV). However, to date, no study has been able to define the most relevant margins for patients with stage 1 tumours. Methods: We performed a retrospective analysis including patients with adenocarcinoma (ADC) or squamous cell carcinoma (SCC) of localised stage T1N0 or T2aN0 who underwent surgery. The ME was measured from this boundary. The profile of the type of tumour spread was also evaluated. Results: The margin required to cover the ME of a localised NSCLC with a 95% probability is 4.4 mm and 2.9 mm for SCC and ADC, respectively. There was a significant difference in the maximum ME distance based on whether the patient had chronic obstructive pulmonary disease (COPD) (p = 0.011) for ADC. Multivariate analysis showed a statistically significant relationship between the maximum microextension distance and size with the shrinkage coefficient. Conclusion: This study definitively demonstrated that the ME depends on the pathology subtype of NSCLC. According to International Commission on Radiation Units and Measurements (ICRU) reports, 50, 62 and 83 CTV margins, proposed by these results, should be added to the GTV (Gross tumour volume). When stereotactic body radiation therapy is used, this approach should be considered in conjunction with the dataset and other margins to be applied.

Keywords: non-small-cell lung carcinoma; adenocarcinoma; squamous cell carcinoma; stereotactic body radiation therapy
1. Introduction

Non-small-cell lung carcinomas (NSCLCs) represent 85% of lung cancer diagnoses [1]. Only 15% to 25% of tumours are diagnosed early [2,3], and their management is considerably improved with the development of stereotactic body radiation therapy (SBRT), mini-invasive surgery and interventional radiology.

The standard treatment for stage I NSCLC is surgery, reaching 3-year and 5-year overall survival rates of 77.9–79% and 66.1–84%, respectively [4–6]. However, approximately one in four patients is not eligible for surgery due to medical contraindications or because the patient refuses the operation.

Currently, the approved therapeutic alternative is SBRT, allowing a 3-year local control rate of 89–96%, a specific survival rate of 66–82% and an overall survival rate of 32–91.8% [7–11].

To delineate targets in radiotherapy, several volumes have been defined by the International Commission on Radiation Units and Measurements (ICRU) reports 50, 62 and 83, including the gross tumour volume (GTV), the clinical target volume (CTV) including microscopic extension (ME) and the planning target volume (PTV). CTV remains a disputable topic in SBRT studies.

A study of CTV margins for lung cancer treated at the time of tridimensional radiotherapy was performed based on pathological examination data [12]. However, to date, no studies have defined specific margins for tumours meeting the criteria for SBRT. These margins remain to be specified to optimise the definition of irradiated volumes, to improve local control and to limit side effects [13,14]. The objective of our study was therefore to use anatomopathological slices to evaluate bronchopulmonary tumour cell extension beyond the macroscopically visible tumour to determine the best CTV margins for SBRT.

2. Materials and Methods

2.1. Ethical Approval

This study follows the mandatory French laws required by the CNIL (Commission Nationale de l’informatique et des libertés), was declared to this French institution by the MR004 form and was recorded in the HDH (Health Data Hub) and was approved by our Institutional Review Board.

2.2. Patients

One hundred and twelve patients with localised pT1N0 or pT2aN0 adenocarcinoma (ADC) and 42 patients with localised pT1N0 or pT2aN0 squamous cell carcinoma (SCC), who underwent surgery from January 2014 to December 2018 and from January 2013 to December 2018, respectively, were retrospectively included. Patients with neoadjuvant chemotherapy or without a preoperative computed tomography (CT) scan with iodinated contrast injection were excluded. A keyword search in the computerised database of anatomo-clinical reports (DIAMIC) was performed to identify cases that matched the following inclusion criteria over the chosen period: “(p) T1 (a b c) T2a” AND “adenocarcinoma lung” OR “squamous cell carcinoma” OR “non small cell lung cancer”. For each group of pathologies, the patients’ characteristics are reported in Table 1.

Table 1. Patient characteristics.

|                     | Adenocarcinoma | Squamous Cell Carcinoma | Total |
|---------------------|----------------|-------------------------|-------|
|                     | n   | %   | n   | %   | n   | %   |
| Patients            | 112 | 72  | 42  | 28  | 154 |
| Slides              | 341 | 73  | 127 | 27  | 468 |
| Age (mean, in years)| 65.5| 65.8|     |     |     |
| Gender              |     |     |     |     |     |
| Male                | 52  | 46  | 33  | 79  | 85  | 55  |
| Female              | 60  | 54  | 9   | 21  | 69  | 45  |
| pTNM 2017           |     |     |     |     |     |     |
Table 1. Cont.

|                          | Adenocarcinoma | Squamous Cell Carcinoma | Total |
|--------------------------|----------------|-------------------------|-------|
|                          | n   | %  | n  | %  | n  | %  |
| pT1mi N0                 | 7   | 6  | 0  | 0  | 7  | 4  |
| pT1a N0                  | 7   | 6  | 8  | 19 | 15 | 10 |
| pT1b N0                  | 43  | 39 | 13 | 31 | 56 | 36 |
| pT1c N0                  | 28  | 25 | 7  | 17 | 35 | 22 |
| pT2a N0                  | 27  | 24 | 14 | 33 | 44 | 28 |
| Architecture             |     |    |    |    |    |    |
| Acinar                   | 60  | 54 |    |    |    |    |
| Lepidic                  | 18  | 16 |    |    |    |    |
| Mucinous                 | 12  | 11 |    |    |    |    |
| Solid                    | 10  | 9  |    |    |    |    |
| Papillary                | 7   | 6  |    |    |    |    |
| Micropapillary           | 3   | 3  |    |    |    |    |
| No Other Specified       | 1   | 1  |    |    |    |    |
| Atelectasis              |     |    |    |    |    |    |
| Yes                      | 13  | 12 | 9  | 21 | 22 | 15 |
| No                       | 98  | 88 | 33 | 79 | 131| 85 |
| Site                     |     |    |    |    |    |    |
| Proximal                 | 16  | 14 | 8  | 19 | 24 | 16 |
| Peripheral               | 95  | 85 | 34 | 81 | 129| 84 |
| Margins                  |     |    |    |    |    |    |
| Nodular                  | 32  | 29 | 16 | 38 | 48 | 31 |
| Spiculated               | 79  | 71 | 26 | 62 | 105| 62 |
| Insufflation quality     |     |    |    |    |    |    |
| Good                     | 55  | 49 | 24 | 57 |    |    |
| Medium                   | 47  | 42 | 17 | 40 |    |    |
| Poor                     | 10  | 9  | 1  | 3  |    |    |
| Angioinvasion            |     |    |    |    |    |    |
| Yes                      | 10  | 8.9| 7  | 17 |    |    |
| No                       | 101 | 90 | 34 | 81 |    |    |
| Lymphatic invasion       |     |    |    |    |    |    |
| Yes                      | 2   | 2  | 1  | 2  |    |    |
| No                       | 109 | 97 | 41 | 98 |    |    |
| Fibrosis                 |     |    |    |    |    |    |
| Yes                      | 34  | 30 | 13 | 31 |    |    |
| No                       | 78  | 70 | 29 | 69 |    |    |
| Haemorrhage              |     |    |    |    |    |    |
| Yes                      | 44  | 39 | 17 | 40 |    |    |
| No                       | 68  | 61 | 25 | 60 |    |    |
| Inflammation             |     |    |    |    |    |    |
| Yes                      | 16  | 14 | 11 | 26 |    |    |
| No                       | 96  | 86 | 31 | 74 |    |    |
| Mode of extension        |     |    |    |    |    |    |
| AIL                      | 2   | 2  | 1  | 2  | 3  | 2  |
| AIV                      | 1   | 1  | 1  | 2  | 2  | 1  |
| STAS                     | 24  | 21 | 10 | 24 | 34 | 22 |
| Interstitial             | 31  | 28 | 4  | 10 | 35 | 22 |
| Tabacco                  |     |    |    |    |    |    |
| Yes                      | 87  | 78 | 39 | 93 | 126| 82 |
| No                       | 24  | 21 | 3  | 7  | 27 | 18 |
| COPD                     |     |    |    |    |    |    |
| Yes                      | 34  | 30 | 19 | 45 | 53 | 46 |
| No                       | 78  | 70 | 23 | 55 | 101| 64 |
| TILs                     |     |    |    |    |    |    |
| 0–10%                    | 79  | 71 | 10 | 24 | 89 | 59 |
| 20–40%                   | 27  | 24 | 25 | 60 | 52 | 33 |
| 50–90%                   | 6   | 5  | 7  | 17 | 13 | 8  |

AIL: adjacent lymphatic invasion; AIV: adjacent vascular invasion; COPD: chronic obstructive pulmonary disease; STAS: spread through air space; TILs: tumour-infiltrating lymphocytes.
Surgery consisted of a segmentectomy, lobectomy or pneumonectomy by thoracotomy or videothoracoscopy with extemporaneous examination and lymph node dissection. Surgical specimens were sent fresh and oriented to the pathology department. The superoinferior, anteroposterior, medial–lateral, and maximal dimensions of each tumour were measured on the preoperative CT scan by a radiologist and a radiation oncologist using optimal window/level settings and then compared with the tumour size measured on pathology.

2.3. Histology

The resected specimens were fixed in 10% buffered formalin for 24 h, dehydrated in successive alcohol baths and then embedded in paraffin. The paraffin blocks were cut with a microtome, and the bands were spread on slides and then stained with haematoxylin-eosin. The thickness of the slices was 4 microns. All slices involving the tumour were analysed, representing one to 6 slices per patient with an average of 3 slices. The tumour was delineated with the naked eye using a marker (Figure 1). Then, the microscopic tumour extension (ME) was measured on the slices from the boundary between the tumour and healthy tissue to the most distant tumour cell using a micrometre. For each patient, only the “longest” distance from the available and studied slices was retained for the final analysis. The measurement performed was weighted by a factor corresponding to the shrinkage coefficient of the tissue related to the different fixation and inclusion steps. In a prospective monocentric study, Park et al. described that formalin fixation caused 4.06% shrinkage in 46.8% of tumours. The overall mean tumour size change after formalin fixation was 0.77 mm (SD: 1.02 mm), and the percentage difference in tumour size was 4.06% (range: 0–26.1%; SD: 5.15%) [15]. Measured size, type of architecture according to the World Health Organization (WHO) 2015 classification [16], presence or absence of angioinvasion, lymphatic invasion, presence of fibrosis, inflammation or haemorrhage, and tumour-infiltrating lymphocytes (TILs) were also analysed. TIL rates were used to define three groups: 0–10%, 20–40%, and 50–90% [17].

![Figure 1. Example of microinvasion measurement. (A). Tumour stained with hematoxylin and eosin and delineated with marker pen. (B). Microscopic tumour extension measured from the boundary between the tumour and healthy tissue to the most distant tumour cell using a micrometre.](image-url)
2.4. Statistical Analysis

Statistical analysis was performed using R v3.6.0 software. The Student’s t test was used to analyse the relationship between the ME and the different clinical or histological parameters. The Chi² or Fisher exact test was used to compare qualitative parameters. The analysis of the margins required to recover the ME was performed by comparing the percentage of patients with an ME less than or equal to the value of the analysed margin. In the case of a recovery of 95% of the sample, the margin is considered acceptable. Concerning the multivariate analyses, the distribution of the residuals did not follow a normal distribution, and we calculated confidence intervals and p values using the bootstrap method (1000 iterations).

3. Results

3.1. Patient and Tumour Characteristics

The patient and tumour characteristics are presented in Table 1. The mean age at diagnosis of the ADC and SCC groups was 65.5 years (SD 8.29) and 65.8 years (SD 7.84), respectively. In the ADC and SCC groups, the F/M ratios were 1.15 and 0.27, respectively. In the ADC and SCC groups, 34 patients (30%) and 19 patients (45%) developed chronic obstructive pulmonary disease (COPD), respectively.

The mean sizes of ADC and SCC tumours on the CT scan were 2.15 cm (SD 0.9; min = 0.8; max = 5.3) and 2.33 cm (SD 1.1; min = 0.3; max = 5.4), respectively (Table 2).

| Table 2. Tumour size and microscopic extension according to histology. |
|-----------------------------------------------|
| Mean (Standard Deviation) | Median (Q25–75) | Min | Max | n |
|---------------------------|-----------------|-----|-----|---|
| ADC size (cm)             | 1.97 (0.848)    | 1.87 (1.35; 2.52) | 0.21 | 3.95 | 112 |
| ADC radiologic size (cm)  | 2.15 (0.909)    | 2.00 (1.50; 2.60) | 0.80 | 5.30 | 112 |
| SCC size (cm)             | 2.46 (1.00)     | 2.19 (1.50; 3.41) | 1.04 | 4.16 | 42  |
| SCC radiologic size (cm)  | 2.33 (1.10)     | 2.20 (1.57; 2.80) | 0.30 | 5.40 | 42  |
| ADC ME (mm)               | 0.734 (1.12)    | 0 (0; 1.30)       | 0    | 6.12 | 112 |
| SCC ME (mm)               | 0.737 (1.49)    | 0 (0; 0.615)      | 0    | 5.94 | 42  |

ADC: adenocarcinoma; ME: microscopic extension; SCC: squamous cell carcinoma.

Three hundred forty-one slices of ADC were derived from 112 tumours, and 127 slices of SCC were derived from 42 tumours. The mean sizes of ADC and SCC tumours were 1.97 cm (SD 0.85; min = 0.21; max = 3.95) and 2.46 cm (SD 1.00; min = 1.04; max = 4.16), respectively (Table 2).

3.2. Radio-Histologic Correlations

Comparative analysis showed a significant correlation between radiologic size and histologic size with shrinkage coefficients for ADC and SCC, respectively ($p < 0.001$ and $p < 0.01$). A Bland–Altman plot was made respectively for each histological type between radiologic size and histologic size with shrinkage coefficients (Figure 2).

3.3. Adenocarcinoma

A mean distance of 0.73 mm (standard deviation: 1.12), a median of 0 mm (0; 1.30), and minimum and maximum values of 0 mm and 6.12 mm, respectively, were observed for the ME (Table 2). Considering a margin of 3 mm, the ME coverage rate was 96.4%, which was not significantly different from 95% ($p = 0.488$). To cover exactly 95% of the sample, a margin of 2.86 mm would be required. The affected lobe, the presence of TILs, and the presence of vascular or bronchial contact did not significantly influence the ME. A statistically significant linear correlation was noted between the maximum ME distance and the number of slides (correlation coefficient: 0.233, 95% CI (0.0493; 0.401); $p = 0.014$) (Figure 3). A significant difference in distance from the ME depending on whether the
The mean sizes of ADC and SCC tumours on the CT scan were 2.15 cm (SD 0.9; min = 0.8; max = 5.3) and 2.33 cm (SD 1.1; min = 0.3; max = 5.4), respectively (Table 2).

Table 2. Tumour size and microscopic extension according to histology.

|                      | Mean (Standard Deviation) | Median (Q25–75) | Min | Max | n  |
|----------------------|---------------------------|-----------------|-----|-----|----|
| ADC size (cm)        | 1.97 (0.848)              | 1.87 (1.35; 2.52) | 0.21 | 3.95 | 112|
| ADC radiologic size (cm) | 2.15 (0.909)             | 2.00 (1.50; 2.60) | 0.80 | 5.30 | 112|
| SCC size (cm)        | 2.46 (1.00)               | 2.19 (1.50; 3.41) | 1.04 | 4.16 | 42 |
| SCC radiologic size (cm) | 2.33 (1.10)               | 2.20 (1.57; 2.80) | 0.30 | 5.40 | 42 |
| ADC ME (mm)          | 0.734 (1.12)              | 0 (0; 1.30)      | 0   | 6.12 | 112|
| SCC ME (mm)          | 0.737 (1.49)              | 0 (0; 0.615)     | 0   | 5.94 | 42 |

ADC: adenocarcinoma; ME: microscopic extension; SCC: squamous cell carcinoma.

Three hundred forty-one slices of ADC were derived from 112 tumours, and 127 slices of SCC were derived from 42 tumours. The mean sizes of ADC and SCC tumours were 1.97 cm (SD 0.85; min = 0.21; max = 3.95) and 2.46 cm (SD 1.00; min = 1.04; max = 4.16), respectively (Table 2).

### 3.2. Radio-Histologic Correlations

Comparative analysis showed a significant correlation between radiologic size and histologic size with shrinkage coefficients for ADC and SCC, respectively ($p < 0.001$ and $p < 0.01$). A Bland–Altmann plot was made respectively for each histological type between radiologic size and histologic size with shrinkage coefficients (Figure 2).

### 3.3. Adenocarcinoma

A mean distance of 0.73 mm (standard deviation: 1.12), a median of 0 mm (0; 1.30), and minimum and maximum values of 0 mm and 6.12 mm, respectively, were observed for the ME (Table 2). Considering a margin of 3 mm, the ME coverage rate was 96.4%, which was not significantly different from 95% ($p = 0.488$). To cover exactly 95% of the sample, a margin of 2.86 mm would be required. The affected lobe, the presence of TILs, and the presence of vascular or bronchial contact did not significantly influence the ME. A statistically significant linear correlation was noted between the maximum ME distance and the number of slides (correlation coefficient: 0.233, 95% CI (0.0493; 0.401); $p = 0.014$) (Figure 3). A significant difference in distance from the ME depending on whether the patient had COPD ($p = 0.011$). The mean rank of ME was not significantly different based on the GOLD stage ($p = 0.062$). To cover 95% of the ADC sample with COPD, a margin of 3.74 mm would be required. To cover 95% of the ADC sample without COPD, a margin of 2.27 mm would be required. At a risk of 5%, when adjusting for COPD, the number of slides analysed, TILs and GTV, we were unable to show a statistically significant relationship between the maximal distance ME and size with the shrinkage coefficient (Table 3).

**Figure 2.** Bland-Altman plots. Radiologic size and histologic size with shrinkage coefficients for adenocarcinoma (A) and squamous cell carcinoma (B).
Figure 3. Correlation between maximal distance microextension and tumour size for adenocarcinoma (A) and squamous cell carcinoma (B).
Table 3. Results of multivariate analysis to determine the statistical relationship between maximum distance ME and tumour size with the shrinkage coefficient for adenocarcinoma.

| Coefficients | p       | p Global |
|--------------|---------|----------|
| Tumour size (mm) | 0.0209 (−0.00802; 0.0595) | 0.16 | 0.16 |
| COPD 1 vs. 0 | 0.434 (−0.0260; 1.00) | 0.065 | 0.065 |
| Number of slides | 0.166 (−0.0575; 0.504) | 0.15 | 0.15 |
| TILs (%) 20–40 vs. 0–10 | 0.0757 (−0.0418; 0.517) | 0.74 | 0.69 |
| 50–90 vs. 0–10 | −0.340 (−0.965; 0.377) | 0.45 | - |
| GTV volume (cm³) | −0.0304 (−0.0813; 0.0101) | 0.21 | 0.21 |

COPD: chronic obstructive pulmonary disease; GTV: gross target volume; TILs: tumour-infiltrating lymphocytes.

3.4. Squamous Cell Carcinoma

A mean distance of 0.74 mm (standard deviation: 1.49), a median of 0 mm (0; 0.62), and minimum and maximum values of 0 mm and 5.94 mm, respectively, were observed for the ME (Table 2). Considering a margin of 3 mm, the ME recovery rate was 92.9%, which was not significantly different from 95% ($p = 0.524$). To cover exactly 95% of the sample, a margin of 4.43 mm is required. COPD and the affected lobe, and the presence of vascular or bronchial contact did not significantly influence the presence and distance of the ME. The mode of dissemination, including vascular, lymphatic or endobronchial dissemination, did not influence the distance of the ME. The presence of TILs was significantly inversely correlated with the distance from the ME. Post hoc analysis of the comparison of the three TIL groups showed a significant difference in maximum ME distance between the 0–10% and 50–90% ($p < 0.05$) TIL groups. To cover 95% of the samples in the TIL 0–10%, 20–40% and 50–90% groups, margins of 4.97 mm, 2.13 mm and 0 mm, respectively, would be required. With a 5% risk, by adjusting for TILs, COPD, the number of slides analysed and GTV volume, a statistically significant relationship was noted between the maximal distance ME and size with the shrinkage coefficient. Distance ME was also significantly linked to TILs and GTV volume for SCC (Table 4). To cover exactly 95% of SCCs greater than 1.5 cm long on the CT, a margin of 4.92 mm is required. To cover exactly 95% of SCC less than or equal to 1.5 cm long axis on CT, a margin of 0.74 mm is required.

Table 4. Results of multivariate analysis to determine the statistical relationship between maximum distance ME and tumour size with the shrinkage coefficient for squamous cell carcinoma.

| Coefficients | p       | p Global |
|--------------|---------|----------|
| Tumour size (mm) | −0.0461 (−0.0681; −0.0170) | <0.01 | <0.01 |
| TILs (%) 0–10 vs. 20–40 | 0.979 (0.159; 2.41) | 0.015 | 0.015 |
| 50–90 vs. 20–40 | −0.524 (−1.04; −0.172) | 0.25 | - |
| COPD 1 vs. 0 | 0.125 (−0.584; 1.08) | 0.79 | - |
| Number of slides | −0.264 (−0.976; 0.287) | 0.3 | - |
| GTV volume (cm³) | 0.109 (0.0647; 0.169) | <0.001 | - |

COPD: chronic obstructive pulmonary disease; GTV: gross target volume; TILs: tumour-infiltrating lymphocytes.

4. Discussion

There are limited data to directly determine whether the radiologic tumour size, as defined by computed tomographic (CT) imaging, correctly represents the gross pathological tumour size in NSCLC. Giraud et al. reported that without ME, the radiologic size of a lung tumour was very close to its gross pathological size [12]. Similarly, Li et al. demonstrated that the three-dimensional measurement of GTV on CT approximated its pathologic size, not including ME [18]. In contrast, Chan et al. suggested that the radiographic tumour size overestimates the pathologic size [19]. The correlation between radiologic size and pathologic size validates the marked pen limit that we delineated and beyond which we measured ME.
To our knowledge, we present the largest study in terms of patient number and in terms of analysed relevant slices to evaluate tumour cell diffusion beyond the carcinoma boundaries. Few data have been published on the ME or the most appropriate CTV margins during lung irradiation. Giraud et al. showed that margins of 8 mm and 6 mm around the GTV for ADC and SCC, respectively, covered 95% of the ME. This study included 70 patients with ADC (32 patients) or SCC (38 patients), stage I to IV. A mean of 5 slices were analysed per patient (354 slices in total). Notably, in the study of Giraud et al., only 46% of patients had a stage I tumour. Consequently, the majority of tumours were not indicated for SBRT. Furthermore, if the ME is related to tumour size, the study by Giraud et al., with larger tumours, may overestimate the ME of stage I tumours [12]. Li et al. showed that for a 95% coverage of the ME, the margins to be applied for CTV were 7 mm and 5 mm for ADC and SCC, respectively. The content of the article was only accessible to Chinese-reading people, limiting data analysis; this study included a total of only 43 patients [18]. The study by Yuan et al. was conducted to quantify the extent of ME beyond nodal GTV. The extent of nodal extracapsular extension (ECE) on pathologically dissected lymph nodes of 243 patients with NSCLC was measured, and the correlation between ECE and lymph node sizes, histological type and tumour cell differentiation was studied. A 3 mm- and 8 mm-GTV margin was proposed for lymph node sizes $\leq 20$ mm and $\geq 20$ mm, respectively [20]. Van Loon et al. demonstrated a correlation between ME and the volume of GTV delineated on the scanner in a study including 34 NSCLC patients. Histological types were variable: 18 ADCs, six SCCs, four large cell carcinomas, three mixed adeno-squamous carcinomas, one bronchioloalveolar tumour and two NSCLC not otherwise specified. Furthermore, this study included NSCLCs with a large range of diameters from 11.1 to 84.8 mm [21]. Grills et al. showed that tumour grade was correlated with ME in 35 patients with ADC. However, this study did not consider tissue shrinkage due to fixation and included T1N0 tumours. However, looking at the range of tumour sizes, we noticed that the minimum and maximum sizes were 8 mm and 48 mm, respectively, which corresponds to T1 to T2 tumours [22]. As our study included only stage I NSCLC, these correlations between tumour size and ME could explain the difference observed between our results and those reported by Giraud et al., and Li et al. [12,18]. The hypothesis that can explain the lower ME measured in the current study is the relationship between tumour size and the size of the ME. Because the range of size of the tumour included in this current study was not so large and due to limited sample size, we failed to demonstrate a significant correlation between tumour size and ME for adenocarcinoma. We found a significant association between histologic tumour size and ME for SCC. Having also shown a significant correlation between radiographic and histologic size, we evaluated the margin required to cover 95% of MEs when the longest tumour axis measured on CT is greater than or less than 1.5 cm.

Significant heterogeneity in the definition of volumes and dose prescription for stereotactic pulmonary radiotherapy is noted. The ICRU 91 has not specified a value for the margin defining CTV; however, it recommends defining a CTV in each case. The European Society of Radiation Oncology (ESTRO) 2017 recommendations showed that of the 11 institutions surveyed, all but one did not apply a margin for CTV and applied a median margin of 5 mm (3-7 mm) for PTV [23]. Senthi et al. also highlighted a significant heterogeneity of target volumes in their literature review of 20 studies. PTV was defined as a 3- to 11-mm margin from internal target volume (ITV). They were sometimes split into 3- to 8-mm margins from GTV to CTV and 3- to 5-mm margins from CTV to PTV [24]. Recently, in a retrospective study, Trémolières et al. showed that personalised PTV margins of 4 mm can be applied for upper lobe lesions located above the carina if only a single 4DCT is performed [25].

The systematic review by Chi et al. highlighted significant heterogeneity in the prescription of the total dose, fractionation and isodose. A biological equivalent dose (BED) at the isocentre of at least 100 Gy$_{10}$ was associated with better survival and local control. However, the authors also showed that the peripheral BED ranged from 37.5 to
211.2 Gy\textsubscript{10} and was less than 100 Gy\textsubscript{10} in many studies [26]. In contrast, Wulf et al. found that the peripheral BED of the target volume was an independent factor influencing local control [27]. Klement et al. showed that the average between near-minimum and near-maximum doses (BED\textsubscript{ave}) was better correlated with tumour control probability than either BED\textsubscript{max} or BED\textsubscript{min}, and was highly correlated with the mean GTV dose. The authors concluded that BED\textsubscript{ave} may be used as a prescription target and proposed and suggested that more attention should be given to high mean doses within the GTV than to the coverage of the PTV [28]. These differences in margins and dose prescription may lead to variations in ME coverage. Of note, the use of a margin for ITV or PTV could cover these microextensions; however, it is important to remember that the goal of the PTV is to consider the different errors and uncertainties and not the ME, which is the role of the CTV or the differences in motion, which is the role of ITV. Finally, the application of a heterogeneous dose prescription with its dose fall-off at the CTV/PTV border is also a relevant point to discuss. Grills et al. showed that the variability of ME coverage can be substantially dependent on the planned dose distribution [22]. Finally, even if the CTV is generally undefined, assuming coverage of the ME by the penumbra, ICRU 91 recommends that the CTV be formally defined. Moreover, ICRU 91 reminds us that the penumbra can be asymmetrical and that two treatment plans delivering the same dose to the GTV can deliver two different doses to the CTV [29].

The current results demonstrate an inverse correlation between TILs and ME in the SCC group. The presence of TILs is a prognostic factor for prolonged survival in several cancers, including NSCLC [30–33]. The literature review by Bremnes et al. revealed a recent interest in TILs. Of 17 studies on the impact of TILs in NSCLCs, 13 were published after 2011. Donnem et al. showed that CD8+ T cell density was a positive and independent prognostic factor for relapse-free, specific and overall survival [34]. The meta-analysis by Geng et al. analysing 29 articles for a total of 8600 patients showed that CD8+ T cell infiltration of the tumour stroma and microenvironment was associated with improved overall survival. CD4+ T cell infiltration of the tumour stroma was also associated with improved overall survival [35]. Our results suggest that ME is blocked by TILs; consequently, the CTV should be adjusted according to the TIL rate. TIL biopsy research could help to adjust CTV margins. Few studies have compared TIL rates on biopsy and resection specimens. In the breast, two studies have shown that the TIL rate evaluated on biopsy may be representative of the whole tumour [36,37]. However, discordant results were found in colorectal cancer [38]. To our knowledge, no such comparison has been made in NSCLC. Furthermore, biopsy raises a new issue. Aspiration specimens are contaminated by blood, and the degree of contamination varies. The number of lymphocytes in the smear does not always reflect the grade of infiltration of TILs. Nakahara et al. showed that the ratio of the number of lymphocytes to the number of lymphocytes and neutrophils was correlated with the TIL rate [39], so this could be a solution for patients with only a biopsy. However, the biopsy is not systematic before NSCLC SBRT, and a radiomics approach should be considered. Many studies have demonstrated the feasibility of radiomic prediction of the immune microenvironment using CT imaging in NSCLC [40–42]. These methods could make it possible to adapt CTV by describing the tumour microenvironment without invasive methods. However, before the validation of radiomics in microenvironmental analysis by large trials, we propose to use the greater margins when the TIL rate is not known.

The impact of COPD status on ME in NSCLC patients has already been demonstrated in other studies. Maeda et al. showed less differentiated, more invasive histological profiles for stage 1 NSCLC with more frequent vascular and pleural invasions and a solid component more frequently found in ADCs. COPD promotes tumour proliferation and angiogenesis through the production of chemokines and cytokines, such as tumour necrosis factor-\(\alpha\), interleukin-1 and interleukin-6 [43]. Several studies have shown that the presence of a solid component is indicative of a more invasive profile [44,45]. One hypothesis is that when a tumour forms near an emphysematous or inflammatory lesion, the surrounding inflammation, including numerous cytokines (interleukin-6, tumour necrosis factor-\(\alpha\) and
interleukin-1β) [46] and chemokines (CXCL8 and CXCL1), alters the autocrine and paracrine interactions between malignant cells and invading leukocytes. Biton et al. showed that CD8 TIL exhaustion is correlated with COPD severity, whereas CD4 TIL levels remain stable [47]. This notion could explain the significant link between COPD status and ME. The CTV should be adjusted according to the COPD status.

We believe that it might be prudent to apply a margin corresponding to a CTV according to the histological type, the COPD status for ADC, the rate of TILs and the tumour size for SCC. As an example, we propose a decision tree that could be a basis for deciding which margins to define. (Figure 4).

Our study is not exempt from bias, and the retraction of tissue during paraffin inclusion is a bias to consider. During the inclusion phase of surgical specimens, tissue shrinkage occurs. This retraction could lead to a decrease in the measured ME. Several other studies have shown tumour shrinkage at inclusion in NSCLC, head and neck cancers and breast cancers [15,48–50]. In a study of 100 head and neck cancer specimens, the average decreases in length, width and depth after formalin fixation were 4.40%, 6.18%, and 4.10%, respectively [48]. To overcome this uncertainty, we conducted a prospective study in 2017 that included 126 NSCLC surgical specimens that showed an average shrinkage of 4.06% after fixation [15]. We then applied this shrinkage coefficient to our measurements. The limited number of pathological slices per patient is also a limitation of our study; however, this finding is consistent with the literature [12,22]. Finally, the small number of patients in the subgroup analysis based on the TIL rate is also a limitation. Of note, we included only ADC and SCC, and the other histological types were not included based on the limited numbers. We were unable to study the relationship between ME and mutations, given the small number.

The mean value of ME was 2.18 mm for adenocarcinoma (ADC) and 1.33 mm for squamous cell carcinoma (SCC) (p = 0.001). However, considering 95% of the ME, margins of 7 mm and 5 mm must be allowed for ADC and SCC, respectively. The contribution of these CTV margins should be balanced against the good local controls in the literature [8,9,51,52].

5. Conclusions

The mean microscopic extension for all the samples examined was much lower than the CTV adopted by some previous studies and did not differ between the two histologic types studied.

In the case of stereotactic radiotherapy, in the light of our results, we believe that it might be prudent to apply a margin corresponding to a CTV according to the histological type, the COPD status for adenocarcinomas, the rate of TILs and the tumour size for

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**Figure 4.** Decision tree according to histological type, COPD status and TIL rate to cover 95% of the distance of diffusion probability. ADC: adenocarcinoma; COPD: chronic obstructive pulmonary disease; CT: computed tomography; NSCLC: non-small-cell lung carcinoma; SCC: squamous cell carcinoma.
squamous cell carcinomas. Larger prospective trials that take into consideration the tumour movement related to respiration by performing quadridimensional scanning and maximal intensity projection sequences should be performed.

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**Abbreviations**

ADC adenocarcinoma  
BED Biological Equivalent Dose  
COPD Chronic Obstructive Pulmonary Disease  
CT Computed Tomographic  
CTV Clinical Target Volume  
DIAMIC computerized database of anatomoclinical reports  
ECE Extra Capsular Extension  
ESTRO European Society of Radiation Oncology  
GTV Gross Tumour Volume  
ICRU International Commission on Radiation Units and Measurements  
ITV internal target volume  
ME microscopic tumour extension  
NSCLC Non-Small-Cell Lung Cancer  
PTV Planning Target Volume  
SBRT Stereotactic Body Radiation Therapy  
SCC Squamous Cell Carcinoma  
TILs Tumour-infiltrating Lymphocytes  
WHO World Health Organization

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