Overall and GTV subvolumes tumour control probability (TCP) for head and neck cancer treated by 3D-IMRT with inhomogeneous dose distribution

Leszek Hawrylewicz¹, Bogusław Maciejewski², Klaus Rudiger Trott⁵, Andrzej Tukiendorf⁴, Leszek Miszczyk², Magdalena Markowska³

¹Department RT Planning, M. Skłodowska-Curie National Research Institute of Oncology, Gliwice Branch, Poland
²Department Radiotherapy, M. Skłodowska-Curie National Research Institute of Oncology, Gliwice Branch, Poland
³Division Research Programmes, M. Skłodowska-Curie National Research Institute of Oncology, Gliwice Branch, Poland
⁴Faculty of Health Sciences, Department of Public Health Medical University, Wroclaw, Poland
⁵Klinik und Poliklinik für Strahlentherapie der Technischen Universität München, Germany

Introduction. In this study, an original model has been developed to estimate the real TCP that is a product of the TCPs calculated for GTV subvolumes of head and neck cancer based on 3D-IMRT dose planning.

Material and methods. Retrospective pilot group consist of 16 cases of oropharyngeal cancer in stage T1–2N0 previously treated with 3D-IMRT with at least 3-year follow-up. The total dose (TD) was 60–70 Gy in 2.0 Gy fractions delivered over 42–49 days. Within GTV two subvolumes were marked out: SVA with the planned 100% TD, and underdosed (90–95%) SVB. The TCP for both was calculated using the original formula developed by Withers and Maciejewski.

Results. During 3-year follow-up, 8 local recurrences (LR) occurred. In about 70% of SVB “dose cold spots” encompassed more than 50% GTV volume. This resulted in the TCP_{SVB} decrease to 60%. Thus, the real overall TCP was much lower than a priori predicted, and in these cases local recurrences occurred.

Discussion. Both cold spot SVB volumes and their dose deficit strongly correlated with a high risk of LR.

Conclusions. In conclusion the magnitude of dose deficit and the size of cold subvolume within GTV have an independent negative impact on real TCP and demand dose re-planning.

Key words: 3D-IMRT planning, cold spots within GTV, estimates of partial TCPs within GTV subvolumes

How to cite:
• the slope of the exponential decrease of the fraction of surviving tumour stem cells within the irradiated volume (which depends on the dose per fraction, the intrinsic tumour stem cell radiosensitivity, and which may also be influenced by micro-environmental factors).

The dependence of the curative dose (TCP-50) on the tumour volume has been investigated in experimental tumours (in particular by Suit [2] and by Guttenberger [3]) and in clinical studies, (e.g. Maciejewski et al. [4], Dubben et al. [5] and Magee et al. [6]). The analysis of these data suggests that a ten-fold difference in the absolute number of tumour stem cells between tumours of the same type and T-stage (which may be due to differences in gross tumour volume, tumour stem cell fraction at the start of radiotherapy or accelerated repopulation during radiotherapy) may represent a difference in TCD-50 of around 7 Gy. However, the relationship between TCP and TCD-50 is much more complicated when the dose in the GTV is heterogeneously distributed. Theoretical calculations of the impact of dose inhomogeneity within the PTV/GTV have been published, yet little clinical evidence to support these calculations has been presented so far.

When the 3D-IMRT was introduced into daily practice it became obvious that a dose gradient within the target leads to non-uniform dose distribution in the tumour volume. Tomé and Fowler [7, 8] calculated an increase in the TCP for tumour subvolumes boosted to higher dose, and TCP loss within under-dosed sub-volumes (“cold spots”). It was concluded that the clinical impact of a dose deficit would depend not only on the magnitude of the deficit but also on the size of subvolume [9–11].

More than 15 years ago, Withers and Maciejewski developed a radiobiological model for changes in the TCP estimates for subvolumes and their dependence on initial tumour stem cell number represented by the size of the respective subvolumes and the total doses delivered (unpublished). However, at that time, 2D radiotherapy with homogenous dose distribution within the target was the standard and dose differences in the GTV subvolumes were not a problem. Nowadays, 3D-IMRT with heterogeneous dose distribution within the GTV is widely used, which may impact on the TCP [12–15].

Material and methods

Dose planning data

For the present study, a pilot set of 16 consecutive 3D-IMRT treatment plans for T1–T2N0M0, sq.c.c of the oral cavity, oropharynx and supraglottic larynx, all with at least 3-year follow-up and with apparent inhomogeneous dose distribution within the GTV were selected from the treatment planning data bank in our institution. Inhomogeneous dose distribution was defined as sub-volumes larger than 5% of the GTV in which the total dose was reduced by >5%. Treatment plans with homogenous dose distribution D100 or D95 covering the whole GTV were not taken into account.

Radiotherapy

For all 16 patients, the 3D treatment plans and DVHS were developed using the Eclipse Planning System (version 8.6 or 13, Varian). Using the Clinac 2300 accelerator with 120 MLC and 3D-IMRT technique, conventional 2.0 Gy daily fractions were delivered 5 days a week to a total dose ranging from 60 Gy in 42 days to 70 Gy in 48 days. There were no extensions of overall treatment time, and therefore the time factor is not considered in the analysis.

Tumour volume measurements

For the purpose of this study, tumour volumes were estimated from the data bank of the CT/MRI sequential scans spaced by 2–3 mm as proposed by Johnson et al. [11]. The primary tumour was outlined in each scan at the TPS workstation and the tumour volume was calculated by a computer-based analysis system. The primary tumour volume was defined as GTV, which ranged from 2.5 cm$^3$ to 29.2 cm$^3$.

For the purpose of the present analysis, the tumour volume was subdivided into two subvolumes:

• SVA – the volume of the GTV covered by 100% isodoses of the planned total dose (D100);
• SVB – the volume of the GTV covered by on average 90–95% of the planned total dose (D90–95). The total dose for this subvolume was converted into biologically normalised total dose (BNTD) if given in 2.0 Gy fractions using the L-Q model with = 10 Gy.

Initial stem cell number ($K_i$)

Following the assumptions made by McBride and Withers [9], a tumour of 1 cm in diameter ($v = 0.52$ cm$^3$) was assumed as a standard volumetric unit, containing $10^6$ cells [9, 16, 17], among which 1% possesses stem cell potential ($10^2$ tumour stem cells). Then, the initial stem cell number in each specific primary tumour volume ($V_i$) would be:

$$K_i = 107 \times (V_i/0.52) \quad \text{(1)}$$

Stem cell numbers in subvolumes SVA and SVB were calculated using the same equation (1), using SVA and SVB volumetric parameters.

Tumour cure probability (TCP)

The relationship between the number of surviving tumour stem cells ($K_i$), tumour volume ($V_i$) and total dose (TD) approximates simple Poisson statistics [18, 19]:

$$TCP = \exp (-K_i) \quad \text{(2)}$$

where $K_i$ is equal:

$$K_i = K_s \times SF_i \quad \text{(3)}$$
in which \(K_i\) is initial stem cell number and \(SF_i\) is the surviving fraction after the total dose (TD).

This equation can be rearranged as follows:

\[ TCP_i = \exp (-K_i \times SF_i) \]  

Surviving fractions may be estimated using various methods, such as SF2.0 (surviving fraction after a dose of 2.0 Gy), or effective \(D_{10}\), which is the dose that reduces survival to \(e^{-1}\) for a particular fraction regimen or the LQ model. These three methods are mainly used in experimental radiobiology, but are not very practical for daily clinical radiotherapy.

Mc Bride and Withers [9] suggested that the surviving fraction can more easily be determined in terms of \(eD_{10}\) i.e. the dose which reduces stem cell survival by one decade to 10%. In our study this parameter was used. An approximate value for \(eD_{10}\) for 2.0 Gy fractions was suggested as about 7 Gy [1, 9, 16]. Therefore using \(eD_{10} = 7\) Gy, for a tumour treated with the total dose TD, the absolute number of surviving functional stem cells (not surviving fractions) would be reduced to \(10^{-7TD/eD_{10}}\).

Combining equations the subvolume TCP\(_i\) can be calculated from equation:

\[ TCP_i = \exp [-(10^7 \times (V_i/0.52) \times (10^{-TD/eD_{10}}))] \] [5]

where \(10^7\) is approximately the number of stem cells in a tumour 1 cm in diameter (0.52 cm\(^3\)), \(V_i\) – is tumour subvolume, and TD\(_i\) is the delivered total dose.

TCP\(_i\) values were calculated using the previously given parameters for the GTV and subvolumes A and B. Finally, the real TCP\(_{pl}\) was calculated as a product of the TCP\(_A\) and the TCP\(_B\):

\[ TCP_{pl} = TCP_A \times TCP_B \] [6]

For all 3D-IMRT plans, TCP\(_{pl}\) and TCP\(_{pl}\) were compared and finally related with 3-year follow-up clinical results (local recurrence or disease-free survival).

**Clinical data**

After completing the results of TCP\(_{pl}\) and TCP\(_{pl}\) calculations, they were compared with retrospective 3-year treatment outcomes of the selected 16 patients previously treated with 3D-IMRT. The outcome end-points, i.e. local tumour control (LTC) and local recurrence (LR) were considered. There was no incidence of distant metastases.

**Results**

Table I shows initially planned TCP\(_{pl}\) estimated from equation [5] for the data taken from treatment planning charts of the group of 16 cases. Dose planning and delivery had been prescribed by individual radiation oncologists generally based on the T stage criterion, even though tumour volumes differed by about 10 times (2.5 cm\(^3\)–29.2 cm\(^3\)). Although there were no extensions in overall treatment time and the standard fraction of 2.0 Gy was given regularly, 5 days a week, in hindsight, the choice of the total doses for some cases seems illogical, e.g. TD of 70 Gy was given to 4.55 cm\(^2\) (pt. no. 2) whereas the tumour volume 2.5 larger (case no. 11) received only 60 Gy and the largest one in this series (case no. 16) received 63 Gy.

Nevertheless, except for two cases (no. 7 and no. 16), estimates of the planned TCP\(_{pl}\) are within an acceptable range and predicted a high probability of local tumour control.

The analysis of the impact of the subvolumes A and B within GTV on estimated values of the TCP (tab. II) shows that TCP\(_A\) estimated for SVA were generally very high. However they do not correlate with the incidence of local recurrence.

In contrast with SVA, the size of subvolumes SVB, and derived NTD and partial TCP\(_B\) values had a strong impact on the estimated real TCP values, which were decreased by 3–74% compared to the initial TCP\(_{pl}\) calculated from the treatment plans. Three-dimensional least square (20, 21) planes for dose-volume-TCP relationship are presented in figure 1.

The spatial distribution of these three parameters estimated prior to therapy appear to be of little use in predicting the risk of local recurrence (fig. 1 a). The correlation was even weaker when the SVA was analysed (fig. 1 b). Local recurrence was observed in patients who received the prescribed TD. In contrary, figure 1 c shows a significant impact of "cold" dose in SVB on TCP\(_{pl}\) which was particularly strong when, within

**Table I.** Planned TCP values for all 16 patients and gross tumour volume (GTV) and the calculated number of tumour stem cells and prescribed total dose (NTDp). Black dots indicate that a local tumour recurrence occurred during 3-year follow-up.

<table>
<thead>
<tr>
<th>Pts No</th>
<th>T Stage</th>
<th>VOL. (GTV) cm(^3)</th>
<th>Log(_{10}) K (p)</th>
<th>NTDp izobio Gy(_{2})</th>
<th>Planned TCP (p)</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>T1</td>
<td>2.5</td>
<td>7.7</td>
<td>60</td>
<td>~ 88%</td>
</tr>
<tr>
<td>2</td>
<td>T1</td>
<td>4.55</td>
<td>7.95</td>
<td>70</td>
<td>~ 99%</td>
</tr>
<tr>
<td>3</td>
<td>T1</td>
<td>5.4</td>
<td>8</td>
<td>60</td>
<td>~ 77%</td>
</tr>
<tr>
<td>4</td>
<td>T1</td>
<td>5.6</td>
<td>8</td>
<td>60</td>
<td>~ 77%</td>
</tr>
<tr>
<td>5</td>
<td>T2</td>
<td>6.2</td>
<td>8.1</td>
<td>60</td>
<td>~ 71%</td>
</tr>
<tr>
<td>6</td>
<td>T2</td>
<td>6.2</td>
<td>8.1</td>
<td>60</td>
<td>~ 71%</td>
</tr>
<tr>
<td>7</td>
<td>T2</td>
<td>8.1</td>
<td>8.2</td>
<td>60</td>
<td>~ 65%</td>
</tr>
<tr>
<td>8</td>
<td>T2</td>
<td>9.5</td>
<td>8.5</td>
<td>66</td>
<td>~ 93%</td>
</tr>
<tr>
<td>9</td>
<td>T2</td>
<td>11.0</td>
<td>8.33</td>
<td>66</td>
<td>~ 92%</td>
</tr>
<tr>
<td>10</td>
<td>T2</td>
<td>11.5</td>
<td>8.34</td>
<td>66</td>
<td>~ 92%</td>
</tr>
<tr>
<td>11</td>
<td>T2</td>
<td>12.5</td>
<td>8.4</td>
<td>60</td>
<td>~ 91%</td>
</tr>
<tr>
<td>12</td>
<td>T2</td>
<td>14.0</td>
<td>8.43</td>
<td>66</td>
<td>~ 90%</td>
</tr>
<tr>
<td>13</td>
<td>T2</td>
<td>15.0</td>
<td>8.46</td>
<td>70</td>
<td>~ 97%</td>
</tr>
<tr>
<td>14</td>
<td>T2</td>
<td>19.0</td>
<td>8.56</td>
<td>66</td>
<td>~ 87%</td>
</tr>
<tr>
<td>15</td>
<td>T2</td>
<td>22.0</td>
<td>8.63</td>
<td>70</td>
<td>~ 96%</td>
</tr>
<tr>
<td>16</td>
<td>T2</td>
<td>29.2</td>
<td>8.74</td>
<td>63</td>
<td>58%</td>
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</table>
During 3D radiotherapy planning, heterogeneous dose distribution within the target volume needs detailed searching for possible 'cold spots' and 'cold doses', not so much in the CTV and PTV but above all in the GTV. The treatment plan and the dose distribution should be revised by a mathematically simple calculation of the realistic TCP<sub>RL</sub> and compared with the conventionally determined TCP<sub>pl</sub>. For this task we recommend using equation [6]. It is a simple and non-time-consuming procedure. If an unacceptable decrease in real TCP<sub>RL</sub> compared with the planned TCP<sub>pl</sub> is found, the dose distribution within specified volumes needs to be corrected, which should lead to as uniform a dose distribution as possible at least in the GTV.

Table III shows an option of corrections of SVA and SVB and realistic TCP as a product of both estimates (SVA is covered by TD<sub>100</sub> and SVB by TD<sub>90–95</sub>). Black dots indicate that a local tumour recurrence occurred during 3-year follow-up.

### Table II. Estimates of the TCP for subvolumes SVA and SVB within GTV, and realistic TCP as a product of both estimates (SVA is covered by TD<sub>100</sub> and SVB by TD<sub>90–95</sub>). Black dots indicate that a local tumour recurrence occurred during 3-year follow-up

<table>
<thead>
<tr>
<th>Pts No</th>
<th>%VOL&lt;sub&gt;GTV&lt;/sub&gt; (%)</th>
<th>SUBVOLUME A</th>
<th>SUBVOLUME B (V&lt;sub&gt;90–95&lt;/sub&gt;)</th>
<th>TCP&lt;sub&gt;ESTM&lt;/sub&gt; (TCP&lt;sub&gt;A&lt;/sub&gt; X TCP&lt;sub&gt;B&lt;/sub&gt;)</th>
<th>TCP&lt;sub&gt;PL&lt;/sub&gt;</th>
<th>TCP&lt;sub&gt;RL&lt;/sub&gt;</th>
<th>3-year follow-up</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>1.5</td>
<td>V&lt;sub&gt;41&lt;/sub&gt; 60 Gy 94%</td>
<td>V&lt;sub&gt;12&lt;/sub&gt; 56.8 Gy 78%</td>
<td>73%</td>
<td>-15%</td>
<td>DFS</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>1.0</td>
<td>V&lt;sub&gt;43&lt;/sub&gt; 70 Gy 99.5%</td>
<td>V&lt;sub&gt;27&lt;/sub&gt; 61.1 Gy 95%</td>
<td>94%</td>
<td>-5%</td>
<td>DFS</td>
<td></td>
</tr>
<tr>
<td>3</td>
<td>1.0</td>
<td>V&lt;sub&gt;42&lt;/sub&gt; 60 Gy 88%</td>
<td>V&lt;sub&gt;17&lt;/sub&gt; 55.8 Gy 53%</td>
<td>47%</td>
<td>-41%</td>
<td>LR</td>
<td></td>
</tr>
<tr>
<td>4</td>
<td>1.0</td>
<td>V&lt;sub&gt;44&lt;/sub&gt; 60 Gy 94%</td>
<td>V&lt;sub&gt;29&lt;/sub&gt; 56.8 Gy 52%</td>
<td>49%</td>
<td>-28%</td>
<td>LR</td>
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<td>5</td>
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<td>V&lt;sub&gt;46&lt;/sub&gt; 60 Gy 98%</td>
<td>V&lt;sub&gt;94&lt;/sub&gt; 56.7 Gy 41%</td>
<td>40%</td>
<td>-21%</td>
<td>LR</td>
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<tr>
<td>6</td>
<td>1.0</td>
<td>V&lt;sub&gt;49&lt;/sub&gt; 60 Gy 78%</td>
<td>V&lt;sub&gt;21&lt;/sub&gt; 57.4 Gy 80%</td>
<td>67%</td>
<td>-4%</td>
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<tr>
<td>7</td>
<td>1.0</td>
<td>V&lt;sub&gt;51&lt;/sub&gt; 60 Gy 76%</td>
<td>V&lt;sub&gt;29&lt;/sub&gt; 55.9 Gy 62%</td>
<td>47%</td>
<td>-18%</td>
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</tr>
<tr>
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<td>1.0</td>
<td>V&lt;sub&gt;62&lt;/sub&gt; 66 Gy 95%</td>
<td>V&lt;sub&gt;18&lt;/sub&gt; 63.2 Gy 97%</td>
<td>92%</td>
<td>-3%</td>
<td>DFS</td>
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<td>1.0</td>
<td>V&lt;sub&gt;66&lt;/sub&gt; 66 Gy 99%</td>
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<td>-15%</td>
<td>DFS</td>
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<td>-37%</td>
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<td>V&lt;sub&gt;39&lt;/sub&gt; 56.3 Gy 65%</td>
<td>38%</td>
<td>-13%</td>
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<td>1.0</td>
<td>V&lt;sub&gt;40&lt;/sub&gt; 70 Gy 99%</td>
<td>V&lt;sub&gt;60&lt;/sub&gt; 56.5 Gy 23%</td>
<td>23%</td>
<td>-74%</td>
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<td>71%</td>
<td>-16%</td>
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<td>83%</td>
<td>-13%</td>
<td>DFS</td>
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<td>33%</td>
<td>-25%</td>
<td>LR</td>
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</table>

**Discussion**

Many authors have emphasised that both tumour volume (TV) and tumour dose define tumour control probability (TCP) [5, 11–14, 17, 22]. Tumour stage (T), however, fails to provide reliable information of tumour volume and TCP. Therefore tumour staging cannot replace measurement of tumour volumes. Even within one tumour stage, TV can vary considerably as shown in table I: In the group of T1–T2N0M0 treatment plans for oral cavity and oropharynxgeal cancer investigated, there was a 10-fold difference in the TV.

Analysing a survey of cervix, breast, head and neck and melanoma clinical data Dubben et al. [5] produced a series of steep TCP-TV-curves. Because in our model study, 16 treatment plans were randomly chosen from our clinical data bank, we cannot explain why some small TV (case no. 2) were treated with 70 Gy whereas much larger TV received 60 Gy. It was the individual choice of different radiation oncologists, who prescribed total doses according to the T-stages of tumours.

The absolute number of tumour stem cells has been shown to be proportional to the tumour volume in most rodent and human cancers (unless there are large necrotic volumes found [1, 2, 11, 12]). For the purpose of our study, we assumed that 1% of tumour cells are tumour stem cells [9].

A local control rate of 90% results if on average 0.1 tumour stem cells survive, or in other words, if one in ten irradiated tumours contains one or more tumour stem cell. In a tumour with about 10<sup>10</sup> tumour stem cells such as #16, the stem cell surviving fraction has to be about 10<sup>-10</sup> to achieve a local control rate of 90%. Using eD<sub>0</sub> of 7.0 Gy assumed in our model, it would require a total dose of approximately 70 Gy instead of the 63 Gy given to increase TCP from approximately 40% to 90%. The observed local recurrence thus had to be expected.
Table III. Examples of re-planning of dose distribution within the GTV subvolumes SVA and SVB in all patients to get similarly high TCPs in both, and also high realistic overall TCP.<br><br>Figure 1. 3D-least square planes for dose-volume-TCP relationships: A – for planned parameters; B – for subvolume SVA; C – for subvolume SVB; black dots indicate local tumour recurrence occurred during 3-year follow-up.
Besides the absolute number of tumour stem cells, other factors such as hypoxia, clonal radio-resistance, intercellular communication, and repopulation rate may increase inter-tumour or intra-tumour heterogeneity of stem cell density and of the resulting tumour radiosensitivity. Brenner [22] and Johnson et al. [11] suggested that although some deviations in cellular characteristics of the tumour might modify the volume response to radiation it would unlikely be of crucial importance. Daily fractionation with 2.0 Gy in all tumours was given which, if at all, might lead to a similar impact on repopulation, which is known to be a major factor causing local recurrences in head and neck cancer. Currently, there is no way to determine heterogeneity of repopulation rates and starting times between tumours. Thus, the contribution of this factor to the findings of our study cannot be properly evaluated. The intra-tumour heterogeneity of tumour stem cell density cannot, at present, be seriously discussed because of the lack of reliable data, however, histopathological studies on stem cell marker distribution may enable us in the future to determine stem cell density in tumours.

Particularly in 3D-IMRT there is a high risk of minor dose inhomogeneity because of the relatively steep gradient of dose within a narrow distance from the centre of the tumour. Tome and Fowler [7, 8], Withers [9, 10] and other authors discussed in detail the physical and clinical aspects of "cold spots" and "cold doses". Whereas GTV can be precisely contoured using radiological images, CTV and PTV can only be individually surmised based on the experience of the radiation oncologist because there is no chance to image small conglomerates of tumour (stem) cells outside the GTV. Therefore we focused on underdosed cold spots within the GTV. At the edge of the SVB the dose may even be a bit lower, but we used an average value to simplify our model. With constant number of fractions, the dose per fraction is also reduced. To compare biological effectiveness of the total doses in both SVA and SVB, mean total doses (NTD) in the SVB were normalised to the dose given in 2.0 Gy fractions using the L-Q model with α = 10 Gy and listed as NTD(izoGy). The relationship between planned and delivered NTDₜ for SVA and SVB is presented in Table II. The results show that the size of the SVA which received 100% of the planned total dose ranged from 5% to 82% but the mean TD in SVA was high enough to correspond with high TCPₐ except case no.11 for which the planned TD was too low to eradicate the SVA. For the SVB, the situation was worse. In 11 cases, the SVB was larger than the SVA. The real TCP values were estimated by multiplying TCPₐ and TCPₜ calculated for SVA and SVB. The real TCPₚ values significantly differ from the planned TCPₚ values. All local recurrences occurred in those cases in which a significantly reduced real TCPₚ was calculated.

Our results in the present study support the suggestions of other authors that the biological impact of heterogeneous dose distribution and dose deficit in tumour subvolumes depends not only on the dose deficit but also on the extent of the cold spot(s). Tome, Fowler, Withers [7–10] and other authors postulated that a cold spot of 20–40% of the target volume underdosed by 10% of the prescribed TD would cause the loss in TCP by about 15% or more. Our observations are in agreement with those theoretical predictions. Yet, we also agree with Tome and Fowler [7, 8] and Goitstein and Niemierko [19] that a significant decrease in the TCP depends steeply on dose even for small cold volumes, and that such a deficit cannot be rectified by boosting the dose to the relatively large volume of the PTV.

It is obvious that using IMRT and other 3D-conformal techniques, some dose inhomogeneity in the GTV is unavoidable. The efficacy of these radiotherapy techniques cannot only be dealt with on the basis of physical parameters alone, disregarding radiobiological principles [10]. TCP should be
considered as a function not only of dose but also of the initial number of tumour stem cells, indirectly expressed by tumour subvolumes but not by tumour stages. The treatment outcome is strongly influenced by unaccounted differences in a spatial dose distribution. The hazard of cold spots has been clearly documented and intuitively, even a cubic millimetre of receiving a low dose may lead to recurrence. Such a risk significantly increases when the size of a cold spot enlarges from millimetres to cubic centimetres. It must be estimated as a priori as an essential part of treatment planning. Our model involves the simple assumption of constant stem cell density, and uniform dose distribution in each of the two subvolumes (more than two SV can also be analysed). This model should be taken only as example of what might occur in practice.

Whereas complex TCP equations defined by Tome [7, 8] and Goitein and Niemierko [19] may be useful for mathematically sophisticated analyses, they are useless for daily planning by radiation oncologists. Our proposition of TCP estimation (i.e. our equation no. 6) is simple and can easily be used even by a mathematically inexperienced radiation oncologist, and it takes only about one minute using a simple calculator with Ln and Log functions.

The unacceptable discrepancies between the planned (tab. I) and real TCP_{RL} (tab. II) which occurred in our study, need re-planning procedures with the aim of enlarging the D100 subvolume (SVA) and minimising the size of the underdosed cold subvolume (SVB) as much as possible. Examples of such correction of the IMRT planning are shown in table III.

Conclusions

In 3D-IMRT and other conformal radiotherapy techniques, inhomogeneous dose distributions are unavoidable. Therefore the hazard of underdosed cold spot(s) within the target volume (at least GTV) should be accounted for. The efficacy of these radiotherapy techniques expressed by local tumour probability cannot be considered based on physical parameters alone, disregarding radiobiological principles. Tumour volume (but not tumour stage) is an appropriate though approximate measure of initial number of tumour stem cells which is the most relevant predictor of the TCP. The biological impact of any dose deficit in the cold spot(s) on the TCP depends not only on the magnitude of the deficit but on the size of the cold spot subvolume. Instead of the 95% isodose criterion, mapping V_{100} within the target receiving 100% of the planned dose is recommended, which should be as large as possible, minimising the biological impact of the underdosed cold subvolume(s). The real TCP_{RL} is the product of the TCP_{A} for the V_{100} and TCP_{RL} for the cold subvolume. Any serious discrepancy between the real TCP_{RL} and the planned TCP_{RL} requires precise re-planning and correction of dose distribution within GTV subvolumes.

This paper is dedicated to the memory of Rod Withers who initiated the concept of the present work.

Conflict of interest: none declared

Boguslaw Maciejewski
M. Skłodowska-Curie National Research Institute of Oncology, Gliwice Branch
Department Radiotherapy
Wybrzeze Armii Krajowej 15
44-102 Gliwice, Poland
e-mail: boguslaw.maciejewski@io.gliwice.pl

Accepted: 2 Jun 2020

Received: 22 May 2020

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