Impact of head and neck cancer adaptive radiotherapy to spare the parotid glands and decrease the risk of xerostomia

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Abstract

Background: Large anatomical variations occur during the course of intensity-modulated radiation therapy (IMRT) for locally advanced head and neck cancer (LAHNC). The risks are therefore a parotid glands (PG) overdose and a xerostomia increase.

The purposes of the study were to estimate:
- the PG overdose and the xerostomia risk increase during a "standard" IMRT (IMRTstd);
- the benefits of an adaptive IMRT (ART) with weekly replanning to spare the PGs and limit the risk of xerostomia.

Material and methods: Fifteen patients received radical IMRT (70 Gy) for LAHNC. Weekly CTs were used to estimate the dose distributions delivered during the treatment, corresponding either to the initial planning (IMRTstd) or to weekly replanning (ART). PGs dose were recalculated at the fraction, from the weekly CTs. PG cumulated doses were then estimated using deformable image registration. The following PG doses were compared: pre-treatment planned dose, per-treatment IMRTstd and ART. The corresponding estimated risks of xerostomia were also compared. Correlations between anatomical markers and dose differences were searched.

Results: Compared to the initial planning, a PG overdose was observed during IMRTstd for 59% of the PGs, with an average increase of 3.7 Gy (10.0 Gy maximum) for the mean dose, and of 8.2% (23.9% maximum) for the risk of xerostomia. Compared to the initial planning, weekly replanning reduced the PG mean dose for all the patients (p < 0.05). In the overirradiated PG group, weekly replanning reduced the mean dose by 5.1 Gy (12.2 Gy maximum) and the absolute risk of xerostomia by 11% (p < 0.01) (30% maximum). The PG overdose and the dosimetric benefit of replanning increased with the tumor shrinkage and the neck thickness reduction (p < 0.001).

Conclusion: During the course of LAHNC IMRT, around 60% of the PGs are overdosed of 4 Gy. Weekly replanning decreased the PG mean dose by 5 Gy, and therefore by 11% the xerostomia risk.

Keywords: Head and neck cancer, Anatomical variation, Adaptive RT, Xerostomia
**Introduction**

The treatment of unresectable Head & Neck Cancer (HNC) consists of a chemoradiotherapy [1,2]. One of the most common toxicity of this treatment is xerostomia, inducing difficulties in swallowing and speaking, loss of taste, and dental caries, with therefore a direct impact on patient quality of life. Xerostomia is mainly caused by radiation induced damage mainly to the parotid glands (PG), and to a lesser extend to the submandibular glands [3]. Intensity modulated radiotherapy (IMRT) permits to deliver highly conformal dose in complex anatomical structures, while sparing critical structures. Indeed, three randomized studies have demonstrated improving (PG) sparing by using IMRT compared to non-IMRT techniques, resulting in better salivary flow and decreased xerostomia risk [4-6]. However, large variations can be observed during the course of IMRT treatment, such as body weight loss [7,8], primary tumor shrinking [7], and PG volume reduction [9]. Due to these anatomical variations and to the tight IMRT dose gradient, the actual administered dose may therefore not correspond to the planned dose, with a risk of radiation overdose to the PGs (Figure 1) [10,11]. This dose difference clearly reduces the expected clinical benefits of IMRT, increasing the risk of xerostomia. Although bone-based image-guided radiation therapy (IGRT) allows for setup error correction, the actual delivered dose to the PGs remains higher than the planned dose [12], due to the fact that IGRT does not take shape/volume variations into account. By performing one or more new planning during the radiotherapy treatment, adaptive radiotherapy (ART) aims to correct such uncertainties. ART has been already shown to decrease the mean PG dose during locally advanced head and neck cancer IMRT [13], but no surrogate of the PG dose difference and of the dosimetric benefit of ART has yet been identified. In the context of IMRT for locally advanced HNC, this study sought to:

- estimate the difference between the planned dose and the actual delivered dose (without replanning) to the PGs, i.e., the PG overdose;
- estimate the PG dose difference with replanning and without replanning to spare the PGs while keeping the same planning target volume (PTV), i.e., the benefit of ART;
- identify anatomical markers correlated with these dose differences (PG overdose and ART benefit).

**Materials and methods**

**Patients and tumors**

The study enrolled a total of 15 patients with a mean age of 65 years (ranging from 50 to 87 years). Patient, tumor, and treatment characteristics are provided on Table 1. All tumors were locally advanced (Stage III or IV, AJCC 7th ed). The mean PG volume was 25.3 cc (ranging from 16.6 cc to 52.1 cc, standard deviation (SD): 8.1 cc).

**Treatment and planning**

All patients underwent IMRT using a total dose of 70 Gy (2 Gy/fraction/day, 35 fractions), with a simultaneous integrated boost technique [14] and concomitant chemotherapy. Planning CTs (CT0) with intravenous contrast agents were acquired with 2 mm slice thickness from the vertex to the carina. A thermoplastic head and shoulder mask with five fixation points was used. PET-CT and MRI coregistration was used for tumor delineation. Three target volumes were generated. Gross tumor volume (GTV) corresponded to the primary tumor along with involved lymph nodes. Clinical target volume 70 Gy (CTV$_{70}$) was

![Figure 1 Illustration of the anatomical variations on the dose distribution.](image-url)

IMRT dose distributions at different times for a given patient, showing the PG overdose without replanning (B) and the benefit of replanning (C). A: Planned dose on the pre-treatment CT (CT0). B: Actual delivered dose without replanning during the treatment (Week 3). C: Adaptive planned dose with replanning to spare the parotid glands (PG) at the same fraction (Week 3). PGs are shown by the red line. The full red represents the Clinical Target Volume (CTV70). The arrow show the head thickness. Figure 1B and 1C compared to 1A shows that the PGs and the CTV70 volumes and the neck thickness have decreased. These anatomical variations have led to dose hotspots in the neck, close to the internal part of the two PG (Figure 1B). Replanning (Figure 1C) allowed to spare the PG even better than on the planning (Figure 1A).
equal to GTV plus a 5 mm 3D margin, which was adjusted to exclude air cavities and bone mass without evidence of tumor invasion. CTV\textsubscript{63} corresponded to the area at high-risk of microscopic spread, while CTV\textsubscript{56} corresponded to the prophylactic irradiation area. GTV, CTV\textsubscript{63}, CTV\textsubscript{56}, and all organs at risk were manually delineated on each CT slice. Adding a 5 mm 3D margin around the CTVs generated the PTVs. PTV expansion was limited to 3 mm from the skin surface in order to avoid the build-up region and to limit skin toxicity [15]. All IMRT plans were generated using Pinnacle V9.2. Seven Coplanar 6-MV photon beams were employed with a step and shoot IMRT technique. The prescribed dose was 70 Gy to PTV\textsubscript{70}, 63 Gy to PTV\textsubscript{63}, and 56 Gy to PTV\textsubscript{56}. The collapsed cone convolution/superposition algorithm was used for dose calculation. The maximum dose within the PTV was 110% (D\textsubscript{2%}). The minimum PTV volume covered by the 95% isodose line was 95%. Dose constraints were set according to the GORTEC recommendations [16]; a mean dose (D\textsubscript{mean}) <30 Gy and a median dose <26 Gy for contralateral PGs.

Patients were treated as planned on CT0 and no changes were applied to dose distribution during treatment. During the treatment course, weekly in-room stereoscopic imaging corrected set-up errors >5 mm. All patients signed an informed consent form. The study was approved by the institutional review board (ARTIX study NCT01874587).

Table 1 Patient, tumor, and treatment characteristics at the initial planning (CT0)

| ID | Gender | Age | Tumor localization | TNM | Volume (cc) | D\textsubscript{mean} (Gy) | Xerostomia NTCP (%) [21] |
|----|--------|-----|--------------------|-----|-------------|------------------|-------------------------|
|    |        |     |                    |     | CTV\textsubscript{70} | HLP  | CLP | HLP | CLP | HLP | CLP |
| 1  | M      | 86  | Tonsil             | T3N1 | 45.2        | 52.1 | 48.6 | 30.2 | 31.1 | 26.5 | 28.3 |
| 2  | F      | 63  | Tonsil             | T2Nx | 26.3        | 31.1 | 27.5 | 31.4 | 26   | 29.0 | 18.7 |
| 3  | M      | 74  | Oropharynx         | T3N2c| 181.5       | 24.9 | 20.7 | 37.9 | 31.1 | 44.3 | 28.4 |
| 4  | F      | 66  | Oropharynx         | T2N2c| 107.2       | 27.8 | 23.4 | 32.9 | 27.9 | 32.3 | 22.0 |
| 5  | M      | 57  | Velum              | T3N0 | 62.4        | 20.7 | 18.0 | 28.1 | 27.8 | 22.4 | 21.7 |
| 6  | M      | 67  | Oropharynx         | T3N2c| 156.2       | 24.5 | 22.7 | 30.8 | 29.4 | 24.7 | 21.4 |
| 7  | M      | 52  | Oropharynx         | T4N2 | 165.1       | N/A  | 21.6 | N/A  | 28.7 | N/A  | 23.4 |
| 8  | M      | 67  | Trigone            | T4N1 | 139.3       | 22.0 | 19.3 | 30.7 | 29.2 | 27.4 | 24.4 |
| 9  | F      | 65  | Oropharynx         | T3N3 | 237.5       | 23.9 | 20.2 | 42.4 | 31.1 | 55.2 | 28.2 |
| 10 | F      | 65  | Oropharynx         | T4N3 | 257.9       | N/A  | 24.5 | N/A  | 35.2 | N/A  | 37.7 |
| 11 | M      | 50  | Oropharynx         | T4N2c| 434.5       | N/A  | 17.7 | N/A  | 36.3 | N/A  | 40.3 |
| 12 | M      | 53  | Oropharynx         | T3N0 | 14.4        | 16.6 | 23.3 | 41.3 | 24.2 | 52.9 | 15.9 |
| 13 | M      | 73  | Oropharynx         | T3N2c| 147.0       | 29.4 | 29.2 | 54.6 | 32.2 | 81.7 | 30.7 |
| 14 | M      | 56  | Larynx             | T3N0 | 14.0        | 22.8 | 29.2 | 19.7 | 9.2  | 10.1 | 2.7 |
| 15 | M      | 52  | Hypopharynx        | T2N2 | 76.3        | 20.3 | 22.4 | 29.4 | 29.1 | 25.0 | 24.4 |

M: male; F: female; CT0: initial planning; CTV\textsubscript{70}: clinical target volume receiving 70 Gy; PGs: parotid glands; HLP: homolateral PGs; CLP: contralateral PGs; D\textsubscript{mean}: mean dose at initial planning; N/A: not applicable (PGs included in the CTV); NTCP: normal tissue complication risk of xerostomia defined as a salivary flow ratio <25% of the pretreatment one [21].

Weekly dose estimations, in cases of replanning and without replanning

During the treatment, each patient underwent six weekly CTs (CT1 to CT6) according to the same modalities as CT0, except for the intravenous contrast agents (not systematically used, particularly in case of cisplatin based chemotherapy). For each patient, the anatomical structures were manually segmented on each weekly CT by the same radiation oncologist. In case of complete response, initial macroscopically-involved areas were still included in the CTV, which was adjusted to exclude any air cavities and bone mass without evidence of initial tumor invasion.

Actual weekly doses (Figure 2, Step 1A) were estimated by calculating the dose distribution on the weekly CT, using treatment parameters and isocenter from CT0. Weekly re-planned doses (Figure 2, Step 1C) were calculated by generating a new IMRT plan on each weekly CT in accordance with the dose constraints described for the initial planning. PTV coverage did not differ between initial planning and weekly re-planned CT. The dose constraints for the organs at risk have respected the GORTEC recommendations at the initial planning and in all replanning.

Total cumulated dose estimations by deformable registration

Cumulated doses were estimated for the two scenarios, with or without replanning (Figure 2 Steps 1B and 1D), according to the following deformable image registration.
procedure (contour-guided Demons registration algorithm) [17]. For PGs on each CT, a signed distance map was generated to represent the squared Euclidean distance between each voxel and the PG surface. Distance maps of each PG were then registered using the Demons registration algorithm [18]. The resulting deformation fields were employed to map the weekly dose distributions to the planning CT using tri-linear interpolation. Next, the mapped dose distributions were summed to estimate the cumulated dose for each PG. The average Dice score for PG registration, from the weekly CT to each planning CT was computed as followed:

\[
\text{Dice score} = \frac{2 \times |A \cap B|}{|A| + |B|},
\]

where: A is the delineated PG contour on the weekly CT, B is the planning contour propagated by the registration and \(|.|\) denotes the number of voxels encompassed by the contour. The Dice score ranges from 0 (worst case: no match between both contours) to 1 (perfect match) [19]. A 3D dose difference in the PG was calculated between the cumulated dose distribution and planned dose distribution.

Anatomical variation description
Anatomical variations (between CT0 and weekly CTs) were characterized by variations in CTV\(_{70}\) and PG volumes, in the distances between PGs and CTV\(_{70}\) and in the thickness of the neck (at the level of the geometrical centers of the PGs). The distance between PGs and CTV\(_{70}\) corresponded to the minimal distance between the surfaces of the two contours (PG-CTV\(_{70}\)), computed using an Euclidean distance map of the first contour, iteratively considering all the points of the second contour and keeping the resulting minimal distance.

Statistical analysis
The impact of the anatomical variations on PG dose was analyzed considering D\(_{\text{mean}}\) and the full DVH. Their impact on the risk of xerostomia was estimated by using the LKB NTCP model (\(n = 1, m = 0.4, \text{ and TD50} = 39.9\)) [20,21], the complication being defined as a salivary flow ratio <25% of the pretreatment one [22].

The mean PG dose differences between the weekly doses (with and without replanning) and the planned dose were calculated (Figure 2). The PG overdose was assessed as the difference between the dose without replanning (at the fraction or cumulated) and the dose at the planning. The benefit of weekly replanning was assessed as the difference between the doses with replanning and without replanning (at the fraction or cumulated). Linear mixed-effects models were used to test if the following parameters were correlated with the PG overdose or the benefit of the weekly replanning: initial volumes of the CTV\(_{70}\) and of the PGs, decreasing (between the weekly CT and the planning CT) of the volume of the CTV\(_{70}\) (in cc and %) and the PGs (in cc and %), shortening of the distance between PGs and CTV\(_{70}\), reduction of the head thickness and the time between the CT0 and the beginning of treatment. All dose and volume comparisons were performed using nonparametric tests (Wilcoxon test). Statistical analysis was carried out using the Statistical Package for the Social Sciences V 20.0.

Results
Since 3 ipsilateral PGs were completely included within the PTV (Patient number 7, 10 and 11), they were excluded from the analysis, resulting in a total of 27 PGs analyzed. The average Dice score [19] for PG registration,
from the weekly CT to each planning CT was 0.92 (from 0.83 to 0.95).

Quantification of anatomical variations during the 7 weeks of treatment
From CT0 to CT6, the PG volumes decreased by a mean value of 28.3% (ranging from 0.0 to 63.4%, SD 18%), corresponding to an average decrease of 1.1 cc/week (ranging from 0.0 to 2.2 cc/week). The CTV70 decreased by a mean value of 31% (ranging from −73% to −13%, SD 28%).

The distance between the PGs and the CTV (PG-CTVds) decreased in 74% of the PGs by 4.3 mm on average (ranging from 0.1 to 12 mm, SD 3.7 mm), whereas it increased in the other 26% of the PGs by 3.2 mm on average (ranging from 1.1 to 6.3 mm, SD 2.1 mm).

The thickness of the neck decreased for 78% of the patients by a mean value of 7.9 mm (ranging from 0.1 to 26.6 mm, SD 6.2 mm).

Dose comparison between planned dose, and doses with or without replanning in PGs
The per-treatment PG doses (with or without replanning) were analyzed, first considering the weekly fractions and then, using the cumulated doses from all weekly fractions, for all the 15 patients. The results are shown in Table 2.

PG dose comparisons at the per-treatment weekly fraction
In order to assess the PG overdose, comparison was first made between the dose at the fraction without replanning (Figure 2 Step 1A) and the planned dose. For 67% of the plans, the Dmean increased on average by 4.8 Gy (up to 24.9 Gy, SD 4.6 Gy). In the other 33% of plans, the Dmean decreased by 3.9 Gy (up to 10.7 Gy, SD 2.9 Gy). The variation of the mean PG dose during the treatment was showed Figure 3 for two representative patients.

Then, to assess the benefit of replanning, comparison was made between the dose with (Figure 2 step 1C) and without replanning (Figure 2 Step 1A). In 85% of the plans, replanning decreased the Dmean on average by 4.6 Gy (up to 23.8 Gy, SD 4.0 Gy).

| Dmean (Gy),mean (Min-max;SD) | p-value |
|-----------------------------|---------|
| Planned dose (1)            | 30.9 (9.2-54.6; 7.9) | - |
| Without replanning (2)      | 33.0 (7.7-61.2; 9.9) | - |
| With replanning (3)         | 29.4 (4.1-51.7; 8.3) | - |
| PG overdose (4) = (2)-(1)    | 1.8 (−10.6-24.9; 5.8) | <0,001 |
| Replanning benefit (5) = (3)-(2) | 3.8 (0-23.8; 4.0) | <0,001 |
| Cumulated doses             |          |        |
| Without replanning (2)      | 32.0 (8.7-57.6; 9.3) | - |
| With replanning (3)         | 28.6 (4.6-51.2; 8.4) | - |
| PG Overdose (4) = (2)-(1)   | 1.1 (−7.9-10.0; 4.1) | 0.1 |
| Replanning benefit (5) = (3)-(2) | 3.6 (0-12.2; 3.3) | <0,001 |

Table 2 Parotid gland overdose and replanning benefit assessments, based on the fraction or the cumulated doses, for all the 15 patients

PGs: parotid glands; Dmean: First, the mean PG dose was calculated for each patient and each week (DmeanWeekly). Then, the mean of the DMeanWeekly was calculated for each patient (DmeanPt). Finally, the mean of the DmeanPt was calculated for the whole population (Dmean).

p values are calculated using the Wilcoxon test, to test if the Dmean in (1) and (2), and if the Dmean in (2) and (3) are statistically different.
Anatomical parameters correlated with PG overdose or replanning benefit
PG overdose and replanning benefit (at the fraction or cumulated) increased with the CTV70 shrinkage and the reduction of neck thickness (p < 0.01). At the fraction, a reduction of 10 cc of the CTV70 or of 1 mm of the neck thickness leads to an increase of the mean PG dose of 0.3 Gy. The PG volume variation has no impact on the mean PG dose.

Discussion and conclusion
The main goal of definitive chemoradiotherapy in locally advanced HNC is to improve locoregional control, while
keeping a high quality of life. Reducing the dose in the PGs during the whole course of IMRT and therefore xerostomia is a major challenge. Indeed, we found the majority of the PGs (59%) being overirradiated of a mean dose of 4 Gy (up to 10 Gy), resulting to an absolute increase risk of xerostomia of 8% (up to 24%). The ART strategy appears to benefit not only to the over-irradiated PG patients, reducing the mean dose of 5 Gy (up to 12 Gy) and the xerostomia risk of 11% (up to 30%), but also to the non-over-irradiated PGs. These results suggest thus a large use of ART for the majority of locally advanced HNC patients. In our study, four patients (N° 4, 7, 10 and 12) have not clear benefit from replanning. These patients were presented a spontaneous decrease of the mean PG dose during the treatment. No more gain was possible with the replanning due to the other constraints (homogeneity,
spinal cord, brainstem ...). The GORTEC dose volume constraints has been respected for all the replanning.

The dosimetric benefit of ART has been shown in a limited number of studies, and not exclusively for the PGs. In a series of 22 patients, Schwartz et al. evaluated the impact of one and two replanning using daily CT on rails [23]. The mean PG dose was decreased of 3.8% for contralateral PGs and of 9% for ipsilateral PGs, with possible sparing of the oral cavity and larynx. In another series of 20 patients, a single replanning performed at the 3rd or 4th week of treatment decreased the mean PG dose of 10 Gy [9]. On the other hand, Castadot et al. didn’t show any dosimetric benefits for PGs when using four replanning in a series of 10 patients, however reducing the spinal cord dose and improving the CTV50 dose conformation [24].

The optimal number and time of replanning are unclear. Wu et al. concluded that one replanning decreased the mean PG dose by 3%, two replanning by 5%, and six replanning by 6% [13]. A “maximalist” weekly replanning strategy was considered feasible in our study, as in an ongoing randomized study (ARTIX) comparing one IMRT based planning to a weekly based IMRT replanning. The benefit of such strategy has to be demonstrated compared to other replanning strategies. Ongoing study (like ARTFORCE trial) test the benefit of only one replanning [24]. The benefit of each supplementary weekly replanning has to be evaluated. A true adaptive RT strategy should be personalized to each patient, ranging potentially from no re-planning to a maximalist weekly replanning. Ideally, replanning decisions may likely be based on either geometrical criteria or cumulated dose monitoring corresponding to the dose-guided RT approach. Replanning is also particularly time-consuming, complete delineation taking up to 2.5 hours in our experience and that of others [25-28].

Deformable image co-registration software can be used to propagate the OAR contours from the initial planning CT to the per-therapeutic planning CT, reducing the delineation time by approximately a factor 3 [26,28]. The CTV delineation should be however carefully checked, to prevent recurrence due to inadequately reduced CTV. Indeed, the goal of ART in our study was to spare the PGs during treatment as they were spared at the planning, while keeping the same appropriate CTV coverage (and not to reduce the CTV coverage).

The analysis of the anatomical variations occurring within the course of IMRT is crucial to understand the overdose of the PGs and to identify early the sub-group of the overdosed PGs (59%). We found that mean volumes decreased by 28% for the PGs and 31% for the CTV, in agreement with the literature reporting values of 15% to 28% for the PGs, 69% for the GTV, and 8% to 51% for the CTV [7,9,13,23,26,29]. We found that the PGs overdose (without replanning) and the dosimetric benefit of replanning increased with the tumor shrinkage and the reduction of head thickness. The last one is likely explained by loss of weight, tumor shrinking and decrease of the PG volume. The reduction of the head thickness leads consequently to the occurrence of dose hotspot in the neck, close or within the PGs (Figure 1). Other studies also found that reduction of the neck diameter increases the risk of over-irradiation [30,31]. The variation of the mean PG dose was more important between the CT0 and the CT1 than between each weekly CT. This difference may be explained by the delay between CT0 and the first weekly CT. In our study, the PG dose differences between the fraction and the initial planning are likely related to both the set-up error (we did not quantified) and the anatomical structures volume/shape variations. Systematic set-up errors may increase the mean PG dose by around 3% by mm of displacement [32]. This point suggests, for a daily practice, to combine both a daily bone registration to correct the set-up errors and replanning to correct the anatomical variations.

Fraction comparison only provides information for a specific moment and there is a need for full treatment dose evaluation and comparison. Deformable registration enables dose fraction accumulation [33]. Since PG shape and volume variations were limited, our study’s Dice scores were relatively high (0.92). However, the Dice score does not provide any information regarding the registration’s anatomical “point to point” correspondence accuracy. Moreover, the possibility of PG defects observed over the course of radiotherapy [34] should prompt careful consideration of this cumulated dose approach, thereby justifying an independent "fraction to fraction" dose analysis. Our results, based on both weekly fraction and cumulated dose, were consistent. The 3D dose visualization and differential DVH of the dose difference between the cumulated dose and planning dose (Figure 1) revealed moreover the heterogeneity of hotspot distribution in PGs, which may also impact on the xerostomia risk. The cranial part of the PGs seems to be more critical [35,36], maybe due to the presence of an important concentration of salivary gland stem cells at this level [37]. The possible heterogeneity of the radiosensitivity within the PG could be therefore more carefully investigated in order to consider to spare not only the full gland (represented by a mean dose end-point) but also subparts of the gland. Indeed, relatively small dose (10 Gy) within the PG may cause severe loss of function [38], and dose greater than 20 Gy may cause up to 90% loss of the acinar cells [39]. It seems also that radiation-induced gland dysfunction are due to membrane damage, causing secondarily necrosis of acinar cells and atrophy of the lobules [40].
tumor localization. Even if CTs from a single patient were always delineated by the same radiation oncologist, intra-observer variabilities in organ delineation are also potentially responsible for uncertainties. Moreover, the clinical benefit of the weekly replanning has been estimated and was not reported in the study.

In conclusion, an ART strategy combining a daily bone registration and a weekly replanning may be proposed for locally advanced HNC, with an expected benefit to decrease xerostomia. This PG-sparing strategy appears however particularly complex and should be therefore assessed within prospective trials, with a special attention for CTV delineation. The optimal number and time of replanning are unclear. The benefit of a weekly replanning strategy versus other replanning strategies have to been demonstrated.

Competing interests
The authors declare that they have no competing interests.

Authors’ contributions
JC, GL, EC, FJ, KB and OH were responsible for patients treatments and care. AS, MN, PH and GC performed the deformable registration and the dose accumulation. JDO, JC, and RDC performed all statistical analysis. JC, AS and RDC wrote the manuscript. All authors helped, read and approved the final manuscript.

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