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## **Implementation of an optimization method for parotid gland sparing during inverse planning for head and neck cancer radiotherapy**

### **Implémentation d'une méthode optimisée pour l'épargne des glandes parotides lors de la planification d'arcthérapie volumétrique modulée (VMAT) des cancers ORL**

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Conflit d'intérêt : aucun

Titre courant : Optimization method for parotid gland sparing

## Résumé

Objectif de l'étude : Afin d'épargner les glandes parotides lors d'une planification dosimétrique de la sphère ORL en arcthérapie volumétrique modulée (VMAT), un modèle basé sur la superposition de volumes cible prévisionnel (PTV) et des organe à risque (Moore et al.) a été développé et évalué.

Matériels et méthodes : les dossiers de 160 patients traités pour un cancer ORL localement évolué ont été considérés. Une optimisation du modèle a été réalisée avant son évaluation. Trente cas ont été planifiés avec et sans utilisation du modèle. La variabilité entre les opérateurs a été évaluée par 12 opérateurs sur un cas. Les doses moyennes dans les glandes parotides ( $D_{\text{moy}}$ ), l'indice d'homogénéité et le nombre d'unités moniteur (UM) ont été relevés.

Résultats : au total, 89 % des  $D_{\text{moy}}$  dans les glandes parotides prédites par le modèle ont été atteintes. L'utilisation du modèle a significativement réduit la  $D_{\text{moy}}$  dans les glandes parotides :  $-6,1 \pm 4,3$  Gy. Avec le modèle, une moins bonne homogénéité dans le PTV ainsi qu'une augmentation des UM (+10,5 % en moyenne) ont été obtenues. Pour la variabilité entre les opérateurs, les histogrammes dose-volume (HDV) des glandes parotides étaient significativement différents avec et sans utilisation du modèle ; l'écart type de la  $D_{\text{moy}}$  est passé de 2,2 Gy à 1,2 Gy et de 2,9 Gy à 0,8 Gy respectivement pour les glandes parotides homolatérale et controlatérale.

Conclusion : Lors d'une planification inverse en ORL, le modèle optimisé guide l'opérateur en fournissant une valeur de  $D_{\text{moy}}$  atteignable la plus faible possible pour les glandes parotides, permettant ainsi une diminution significative de la  $D_{\text{moy}}$  de -6,1 Gy. Cette méthode permet de réduire significativement la variabilité entre les patients et entre les opérateurs lors de la planification dosimétrique.

Mots clés : Planification dosimétrique ; prédiction de la dose ; VMAT ; contrôle qualité

## **Abstract**

**Purpose:** To guide parotid gland (PG) sparing at the dose planning step, a specific model based on overlap between PTV and organ at risk (Moore et al.) was developed and evaluated for VMAT in head-and-neck (H&N) cancer radiotherapy.

**Materials and methods:** One hundred and sixty patients treated for locally advanced H&N cancer were included. A model optimization was first performed (20 patients) before a model evaluation (110 patients). Thirty cases were planned with and without the model to quantify the PG dose sparing. The inter-operator variability was evaluated on one case, planned by 12 operators with and without the model. The endpoints were PG mean dose ( $D_{\text{mean}}$ ), PTV homogeneity and number of monitor units (MU).

**Results:** The PG  $D_{\text{mean}}$  predicted by the model was reached in 89% of cases. Using the model significantly reduced the PG  $D_{\text{mean}}$ :  $-6.1 \pm 4.3$  Gy. Plans with the model showed lower PTV dose homogeneity and more MUs (+10.5% on average). For the inter-operator variability, PG dose volume histograms without the optimized model were significantly different compared to those with the model; the  $D_{\text{mean}}$  standard deviation for the ipsilateral PG decreased from 2.2 Gy to 1.2 Gy. For the contralateral PG, this value decreased from 2.9 Gy to 0.8 Gy.

**Conclusion:** During the H&N inverse planning, the optimized model guides to the lowest PG achievable mean dose, allowing a significant PG mean dose reduction of -6.1 Gy. Integrating this method at the treatment planning step significantly reduced the inter-patient and inter-operator variabilities.

Keywords: Treatment planning; prediction dose; VMAT; quality control

## Introduction

Intensity modulated radiotherapy (IMRT) is the standard treatment for head-and-neck (H&N) cancer, sparing the parotid glands (PGs) with sharp dose gradients, and thereby limiting xerostomia [1–3]. However, there is variability during the treatment planning optimization process depending on the clinical case complexity, treatment planning system (TPS), modulation degree, or operator experience. A dose planning only based on the standard dose recommendations does not provide the optimal treatment planning for every patient [4]. Since 2006, several studies have proposed dose and/or geometric indexes/tools based on their own experience to limit treatment planning variability [5–11]. In 2011, Moore et al. proposed a method based on a mathematical model to guide operators during the dose planning process about the compromise between the delivered dose to the target volume and the organ at risk (OAR) dose constraints [5]. The mathematical model (based on a knowledge of prior patients) allows prediction of the lowest achievable mean dose ( $D_{\text{mean}}$ ) to an OAR knowing the prescribed dose to the PTV and the overlap volume between the PTV and considered OAR. Moreover, they shown that this method better saves healthy tissues while decreasing inter-operator variability. However, the proposed model was not specific for tumor localization, and was conceived as part of the specific practices from the authors' institution (Washington University School of Medicine, St Louis, MO). To our knowledge, this type of method has not been developed in another institution, with an optimized model for PG sparing during H&N

cancer radiotherapy. In addition, the inter-operator variability with the implementation of such a method has not been evaluated.

In the context of volumetric modulated arc therapy (VMAT) for H&N cancer radiotherapy, the objectives of this study were:

- to optimize the published model of Moore et al. [5] for PG dose sparing, with our institution practices
- to evaluate the optimized model's performance in a large cohort of patients
- to quantify the PG dose sparing with the use of the optimized model (compared to the standard approach)
- to evaluate inter-operator variability

## **Methods**

### ***Patient characteristics, image acquisition and delineation***

A cohort of 160 consecutive patients with histologically confirmed locally advanced H&N squamous cell carcinoma treated from December 2015 to September 2017 was used for this study. All patients had bilateral cervical lymph node irradiation. Thirty-eight patients included in the first phase of the study (optimization and evaluation of the mode) had post-operative radiotherapy. Computed tomography (CT) contrast-enhanced images, indexed every 2-3 mm, were acquired, from the vertex to the carina (Brilliance, Big Bore, Philips, Netherlands). Target volume and OARs were delineated slice-by-slice on CT images. The gross tumor volume (GTV) was defined by clinical examination, CT, and FDG-PET scan. The GTV encompassed all visible and palpable primary and nodal disease. In post-operative cases, areas of involved margins or lymph node stations containing lymph nodes with extracapsular extension were considered to be at very high risk. High-risk clinical target volume (CTV) was defined as areas potentially containing

microscopic disease limited by anatomic barriers or by the GTV plus a 10 mm margin, when an anatomic barrier was not clearly identified (e.g., base of the tongue), or when the lymph node levels containing involved lymph nodes and neighbouring node levels were considered at risk of subclinical involvement greater than 15-20%. Low-risk CTV encompassed the remaining lymph node areas at risk of potential microscopic spreading greater than 5%. The PTV, designed to account for setup uncertainties, was defined using an additional margin of 5 mm. OARs included spinal cord, brainstem, mandible, larynx, bilateral inner ears, oesophagus, pharyngeal constrictors, oral cavity, contralateral PG (CPG), ipsilateral PG (IPG), and submandibular glands (SMGs). These OARs were delineated by radiotherapists regarding international recommendations [12]. The term ipsilateral or contralateral makes reference to the relationship between the OAR and the highest dose level PTV.

### ***Dose planning***

All patients received definitive or post-operative external beam radiotherapy using VMAT technique. Patients were immobilized in the supine position with custom aquaplast masks holding both the neck and shoulders (Posicast-Lite PR5, CIVCO, Iowa, US). Two or three dose levels in simultaneously integrated boost (SIB) were used, depending on the clinical situation (Table 1). Treatment planning was carried out on the Pinnacle v.9.10 (Philips) TPS using 6 MV photon beams from a Synergy or VERSA HD linear accelerator (Elekta, Stockholm, Sweden) equipped with an Agility (160 leaves) multileaf collimator. The collapsed cone convolution algorithm and a dose grid size of  $3 \times 3 \times 3 \text{ mm}^3$  were used for dose calculation. Two  $358^\circ$  clockwise and counter-clockwise arcs (control point every  $2^\circ$ ) were used. The highest priority was assigned to PTVs with the following constraints: more than 95% of any PTV should receive more than 95% of the prescribed dose, or more than 98% of any PTV should receive more than 90% of the prescribed dose; no more than 2%

of any PTV could receive more than 110%. The following OAR dose constraints were used [12–15]: brainstem and spinal cord maximum doses ( $D_{2\%}$ ) were 50 Gy and 45 Gy, respectively; for the PG, the mean dose ( $D_{\text{mean}}$ ) was  $<26$  Gy or the maximum dose received by 50% of the volume was  $<30$  Gy; for the SMG,  $D_{\text{mean}}$  was  $\leq 39$  Gy; for the oral cavity,  $D_{\text{mean}}$  was as low as possible; for the larynx,  $D_{\text{mean}}$  was  $\leq 45$  Gy; for the pharyngeal constrictors,  $D_{\text{mean}}$  was  $\leq 40$  Gy; for the oesophagus,  $D_{\text{mean}}$  was  $\leq 35$  Gy; for the internal and inner ears,  $D_{2\%}$  was  $\leq 40$  Gy; and for the mandible,  $D_{2\%}$  was  $\leq 65$  Gy.

All dose plans were performed by physicists or dosimetrists, and validated by a radiation oncologist.

### ***Moore et al. optimization model***

Moore et al. defined a mathematical model allowing for prediction of a radiation plan to achieve the lowest possible mean dose to an OAR [5]. This model depends on the prescribed dose to the PTV and the overlap volume between the PTV and considered OAR [5]. The equation of the model was the following:

$$D_{\text{achievable\_mean}} = D_{\text{prescribed}} \left( A + B \left( C - e^{-D \times V_{\text{overlap}}/V_{\text{PG}}} \right) \right) \quad (1)$$

$V_{\text{overlap}}$  is the intersection between PTV and PG.

This model was created from 17 H&N patients (overlap between PGs and PTV) and 8 prostate patients (overlap between rectum and prostate) static IMRT plans, with  $A = 0.2$ ;  $B = 0.8$ ;  $C = 1$ ;  $D = 3$ .

An adapted Moore's model was fitted based on 20 H&N standard treatment plannings from our institution. A script was developed in the TPS to extract volumes of PTV, PGs and overlaps between PTV and each PG. Delivered mean doses to the PGs were collected. Using these data, the ratio of the PG  $D_{\text{mean}}$  to the prescription dose to PTV



Optimization method for parotid gland sparing – N. Delaby

( $D_{\text{prescribed}}$ ) was plotted against the ratio of the overlap volume between PTV and PG ( $V_{\text{overlap}}$ ) to the PG volume ( $V_{\text{PG}}$ ), for each patient PG. Moore's equation coefficients (A, B, C and D) were modified by a dichotomy process, so that the curve of the model fits along the lower bound of our local data, representing the optimal average PG dose achievable in our selected cohort of patients ( $N = 20$ ).

When the optimized model was used, the mean dose objective of the entire PG was adjusted in the TPS according to the  $D_{\text{achievable\_mean}}$  (obtained from  $D_{\text{prescribed}}$ ,  $V_{\text{overlap}}$  and  $V_{\text{PG}}$ ).

***Evaluation of the optimized model (difference between the achievable PG mean dose calculated by the model and the planned PG mean dose)***

The robustness of the adapted Moore model was assessed for a cohort of 110 H&N patients. Treatment plans were optimized knowing lowest achievable PG mean dose from the optimized model, from the equation (1). When the PG  $D_{\text{predicted}}$  was not reached, an analysis was performed to understand the model limits.

***Comparison between the standard planning and the use of the optimized model for achievable PG mean dose prediction***

A comparison between the standard planning and the planning with the use of achievable PG mean doses was performed in a cohort of 30 H&N patients treated with definitive radiotherapy. Patients with post-operative irradiation were not included in this phase of the study to avoid biases in the data analysis. Dose planning was first performed with standard planning dose constraints [12–14]. Then, three planners retrospectively replanned the 30 H&N cases using the lower achievable PG mean dose provided by the optimized model during the dose planning process.

***Inter-operator variability with standard planning and the use of the optimized model for achievable PG mean dose prediction***

The impact of the model application on inter-operator variability was evaluated in one H&N representative case of our clinical practices, with three dose levels (IPG overlap with PTV: 3.3 cm<sup>3</sup>; CPG overlap with PTV: 1.7 cm<sup>3</sup>). Twelve operators (medical physicists and dosimetrists) with different dosimetric experience (from 3 months to 10 years, median = 2.75 years) performed two plans of the same case: one with the standard planning and one using PG achievable mean dose provided by the optimized model.

***Endpoints and statistical analysis***

a. Dose endpoints

Dose distributions were assessed with dose volume histograms (DVHs) and particular dosimetric points, including the volume of the PTV receiving at least 95% of the prescribed dose ( $V_{95\%}$ ), the mean dose to PG, and the maximum dose ( $D_{2\%}$ ) to the spinal cord.

In addition, the delta index ( $\delta$ ) was computed to evaluate the ability of the model to be respected [5]. It was defined as:

$$\delta = \frac{D_{mean} - D_{predicted}}{D_{predicted}} \quad (2)$$

Ideal  $\delta$  value is 0 (no difference between mean doses predicted and planned).  $\delta$  index allows to define the deviation between the predicted dose value by the model ( $D_{predicted}$ ) and the final calculated dose ( $D_{achieved}$ ).

NTCP values were calculated for PGs with the parameters defined by Dijkema et al. [16] and Houweling et al. [17]: TD50 = 39.9 Gy; n = 1; m = 0.40.

b. Treatment plan quality

Plans were assessed by conformity [18], and homogeneity [19] indexes, and complexity metrics. These indexes are defined as follows:

$$\textit{Conformity index (CI)} = \frac{V_{PTV\ 95\%}}{V_{PTV}} \quad (3)$$

where  $V_{PTV95\%}$  is the volume of PTV receiving 95% of the prescribed dose.  $V_{PTV}$  is the volume of the PTV. Ideal CI value is 1.

$$\textit{Homogeneity index} = \frac{D_{2\%} - D_{98\%}}{D_{50\%}} \quad (4)$$

where  $D_{2\%}$ ,  $D_{98\%}$  and  $D_{50\%}$  represent the doses to 2%, 98% and 50% of the PTV, respectively. Ideal HI value is 0.

Complexity of treatment plans was calculated with different metrics described by Du et al. [20]:

- number of monitor units (MU) planned (1 UM = 1 cGy in reference conditions: 10 cm x 10 cm square field, Skin Surface Distance = 95 cm, depth = 5 cm)
- aperture area (computed as the total area of all MLC segments openings, in cm<sup>2</sup>)
- aperture irregularity (calculated the non-circularity of the aperture, equal to 1 in case of circular aperture)
- beam modulation (taking into account aperture area and MU number associated to each segment, equal to 0 with a treatment plan without modulation)

Quality assurance of treatment plans was performed with EBT3 Gafchromic films (Ashland) in an Octavius 40043-CT homogeneous phantom (PTW). Films were scanned using the Epson 11000XL desktop flat-bed scanner in association with FilmQAPro

software (Ashland). Comparison between delivered and calculated dose distributions was performed by local gamma analysis [21] (Verisoft software v7.0.1, PTW) with 3%/3 mm and 30% low dose threshold criteria.

c. Statistical analysis

Wilcoxon (non-parametric) tests were performed to compare standard planning with planning using the optimized model. A p-value  $\leq 0.05$  was considered as significant. Correlation test (Spearman) was performed between dose difference (optimized model minus standard method) and  $V_{\text{overlap}}/V_{\text{PG}}$ .

## Results

### *Moore et al. model optimization: determination of the minimum PG achievable mean dose*

Figure 1 displays the PG mean dose normalized to the prescribed doses for PTV in function of the relative overlap between PG and PTV. The mathematical model defined by equation (4) showed better fitting PG sparing considering overlap with the PTV and the prescribed dose to the PTV (solid curve in Fig. 1 representing the inferior shape).

$$D_{\text{mean}} = D_{\text{prescribed}} \left( 0.17 + 0.8 \left( 1.02 - e^{-V_{\text{overlap}}/V_{\text{PG}}} \right) \right) \quad (5)$$

where  $D_{\text{mean}}$  is the achievable PG mean dose, prescribed dose is the dose prescribed to the PTV,  $V_{\text{overlap}}$  is the overlap volume between PG and PTV, and  $V_{\text{PG}}$  is the PG volume.

Then, the  $D_{\text{mean}}$  provided by this optimized model for each PG was used during the planning process. These values were implemented as optimization goals with a script.

***Evaluation of the optimized model (difference between the achievable PG mean dose calculated by the model and the planned PG mean dose)***

Figure 2 displays the normalized mean doses ( $D_{\text{mean}}/D_{\text{prescribed}}$ ) to PGs plotted against the ratio between volume overlap between the PTV and the PG, within a cohort of 110 patients.

The mean ( $\pm$  SD)  $\delta$  index value was 0.13 ( $\pm$  0.11) for the IPG and 0.9 ( $\pm$  0.12) for the CPG. The  $\delta$  index value of 26 PGs was outside the range [mean  $\delta \pm 1$  SD], including 15 IPGs and 12 CPGs (circles in Figure 2).

***Evaluation of PG sparing with the optimized model compared to the standard method***

Figure 3 and Figure 4 presents the mean PTV and PG DVHs for treatment plans performed with and without the optimized model, within a cohort of 30 patients. Table 2 presents the delivered doses to PTVs and OARs,  $\delta$  index and NTCP. Plans performed with the use of optimized model showed an increase of mean PTV coverage and an increase of PTV maximum dose (+1.1 Gy on average). On average, the IPG mean dose decreased of -6.4 Gy, and the CPG mean dose of -5.8 Gy. Mean doses were similar with standard planning for oral cavity and SMG ( $p > 0.05$ ) and significantly different for larynx and spinal cord ( $D_{2\%}$ ), but accepted by physician. Using the optimized model on the PGs,  $\delta$  index values were significantly lower and NTCP values significantly lower ( $p$ -value  $< 0.01$ ).

Figure 5 displays box plots of the PG mean dose with the standard planning, the achievable PG mean dose predicted by the optimized model and the final PG mean dose reached using the optimized model. Dispersion (minimum value – maximum value) was larger with standard method than optimized one: 43.8 Gy versus 36 Gy for IPG and 35.8 Gy versus 28.6 Gy for CPG. Figure 6 shows “ $D_{\text{mean}}/D_{\text{prescription}}$ ” plotted against

“ $V_{\text{overlap}}/V_{\text{PG}}$ ” for the 20 patients for the IPG (Figure 6a.) and for the CPG (Figure 6b.) obtained with the optimized model strategy. “ $D_{\text{mean}}/D_{\text{prescription}}$ ” plotted against “ $V_{\text{overlap}}/V_{\text{PG}}$ ” for the standard planning has also been reported on Figure 6. No correlation ( $r_s$  (IPG) = 0.02 and  $r_s$  (CPG) = -0.46) was found between dose difference (optimized model minus standard method) and  $V_{\text{overlap}}/V_{\text{PG}}$ . Table 3 gathers also the conformity, homogeneity and complexity indexes and the results of agreement between TPS dose calculation and measurement. Plans performed with the optimized model were more heterogeneous for high- and middle-dose PTV than with standard planning. Plans performed with the optimized model were more complex than those performed with standard planning. In fact, significant smaller field sizes (mean of 50.1 cm<sup>2</sup> versus 56.8 cm<sup>2</sup> with standard planning) and significant MU increase (745 MU versus 673 MU with standard planning) were obtained with the use of the optimized model. Agreement between the TPS dose calculation and measurement was not significantly different between the planning methods.

***Evaluation of inter-operator variability with standard planning and the use of the optimized model for achievable PG mean dose prediction***

Figure 7 displays the minimum and maximum envelopes of the PG DVH for the H&N case planned by 12 different operators, with the standard planning (red colour) and with the use of optimized model (green colour). With the standard planning, the standard deviations of the PG mean doses were 2.2 Gy for the IPG and 2.9 Gy for the CPG. With the use of the optimized model the standard deviations significantly decreased to 1.2 Gy for the IPG and 0.8 Gy for the CPG. Table 4 shows the PTV CI and HI indexes. The PTV CI indexes were not significantly different between the standard planning and the use of the optimized

model. However, treatment plans performed with the optimized model provided HI indexes significantly higher for high and middle dose PTVs and not significantly different for low dose PTV.

## **Discussion**

For H&N treatment planning, we showed the dosimetric benefit of using a simple, personalized and realistic model to predict achievable dose to the PGs. With this model, the PG mean doses decreased by 6.1 Gy on average compared to a standard planning, without compromising PTV coverage and without OAR overdose (Table 2). Moreover, the inter-operator variability was divided by 2 and by 3 for IPG and CPG mean doses, with the optimized model. In average, for standard method, IPG mean doses were above GORTEC recommendation (32.4 Gy for 26 Gy recommended) [12]. Using optimized method, mean IPG dose was equal to GORTEC recommendation. For CPG, optimized method reduced in average PG mean dose below GORTEC recommendation (26.9 Gy to 21 Gy).

PG mean doses obtained using the optimized model were consistent with Moore et al. [5] results. In their study, the model was applied to rectal and H&N IMRT cases. For H&N, the  $\delta$  index varied from 0.28 +/- 0.24 without the model, to 0.13 +/- 0.10 with the model. In our study, for H&N VMAT cases,  $\delta$  index varied from 0.42 +/- 0.28 to 0.09 +/- 0.14 (Table 2). This method seems useful for all localizations with overlap between the PTV and OARs requiring a dose compromise. Powis et al. [22] recently defined their own model based on Moore's method for rectum sparing in case of prostate VMAT.

Guidelines recommend to deliver a mean dose lower than 26 Gy to the CPG and as low as possible to the IPG [4]. The way to achieve these constraints is strongly dependant on operator experience, especially for the CPG. In this study, regardless of operator

experience, the inter-operator variability significantly decreased using the optimized model (Figure 7).

In this study, only 10% of 110 treatment plans realized with the optimized model were outside the arbitrary range [mean  $\delta$  index  $\pm$  1 SD] (represented with points on Figure 2). Some cases could be explained both by the small overlap volume between PGs and PTV, and a PG location on the whole length close to the PTV. For such geometry, more sophisticated models might provide better dose prediction, as the overlap volume histogram (OVH) proposed by Wu et al. [7]. The OVH describes the fractional OAR volume that is within a specified distance to the PTV. This concept could be more suitable for non-overlapping cases but is more complex to implement in clinical routine. Another cause of deviation (between predicted and planned doses) can appear in case of PTV proximity to OAR requiring stringent dose constraints, such as brainstem or spinal cord.

We stress that implementation of such model must be institution-specific because TPS, treatment techniques, or OAR dose constraints can affect the performance of the model. Institution-specific trade-offs have also a major influence on the model. For now, we have only optimized the model for H&N VMAT treatment planning. Even in our institution, the model will need to be re-evaluated to be applied on another clinical localization or irradiation technique.

The optimized model used in this study increased the treatment plan complexity (Table 3). Improving PG sparing generated thinner and smaller segments and increased the number of MUs. Planners should be attentive to avoid too complex treatment plans, by using complexity index thresholds for example. Higher modulation complexity may result in lower dose accuracy. In that way, we performed quality controls to check the dose accuracy. Plans with the optimized model did not show significant difference (Table 3).



Our standard method did not provide any index to inform planner if the lowest OAR dose value is achieving, while covering PTV. Since several years, new automated planning tools were commercialized (AutoPlanning module for Pinnacle (Philips) TPS, RapidPlan for Varian TPS, etc.). Those tools can improve OAR sparing and/or PTV coverage, especially for new planners [23]. Implementation of such tools, needs time to be configured (especially for knowledge-based methods) and evaluated to be confident with. Several studies evaluated the impact of those commercial solutions on PG sparing [23–26]. Compared to manual planning, the use of RapidPlan in a H&N study including 20 patients, allowed a mean PG decrease of 2.2 Gy [24]. With AutoPlanning software, a H&N study on 10 patients reported on average a PG mean dose decrease of 3 Gy [25], and another study on 10 patients found a PG mean dose reduction of 1.5 Gy [23]. Such tools appear attractive but are always a TPS add-on with additional cost. Our implementation of optimized model required only methodology and time. The two approaches, commercial tool and optimized model, are not rivals and can be implemented together [26,27].

Mean dose PG sparing is a crucial objective for H&N IMRT/VMAT techniques. Indeed, due to large anatomical variations during treatment course, the delivered dose may differ from the initial planned dose, especially a dose increase for the PGs [28–30]. Adaptive radiotherapy (ART), using one or several replannings [31,32], has been proposed to spare the PGs [33–37]. Weekly replannings can reduce the PG mean dose by 5 Gy, translating a reduction of 11% in the NTCP risk for xerostomia [30]. Thus, ART workflow can provide PG dose sparing and clinical benefit for some patients (30 to 65% of patients [29]). However, it was known as very time consuming and requiring resources strategy. In this context, the integration of an adapted Moore's method to local dosimetric protocols could reduce PG mean dose easier than an ART strategy.

This study has some limitations. In the model optimization and robustness phase of this work, 38 patients with post-operative bilateral neck irradiation were included to reflect our institutional daily practice. However, in the comparison phase (optimized model versus standard method) only patients with radical radiotherapy were included to avoid biases in the data analysis. Because the achievable lowest dose delivered to the PGs are driven by the prescribed dose to the PTV and the  $V_{\text{overlap}}$  between this PTV and the PG, we may assume that the results obtained for patients with radical radiotherapy would be similar to those with the post-operative radiotherapy. Indeed, our radical radiotherapy protocol include dose levels quite similar to our post-operative protocol for the high risk and elective target volumes. The optimized model was based on a cohort of 20 patients and would likely be more robust with a larger cohort, including more complex geometries to improve the dose prediction of non-standard cases. The inter-operator variability was assessed only on one case, and could be extended on more clinical cases. As using the optimized model increases treatment plan complexity, a constraint in term of minimum segment area or maximum MU could have been introduced. Moreover, this study could be supplemented by a multi-centric study to extent methodology to other practices.

## **Conclusions**

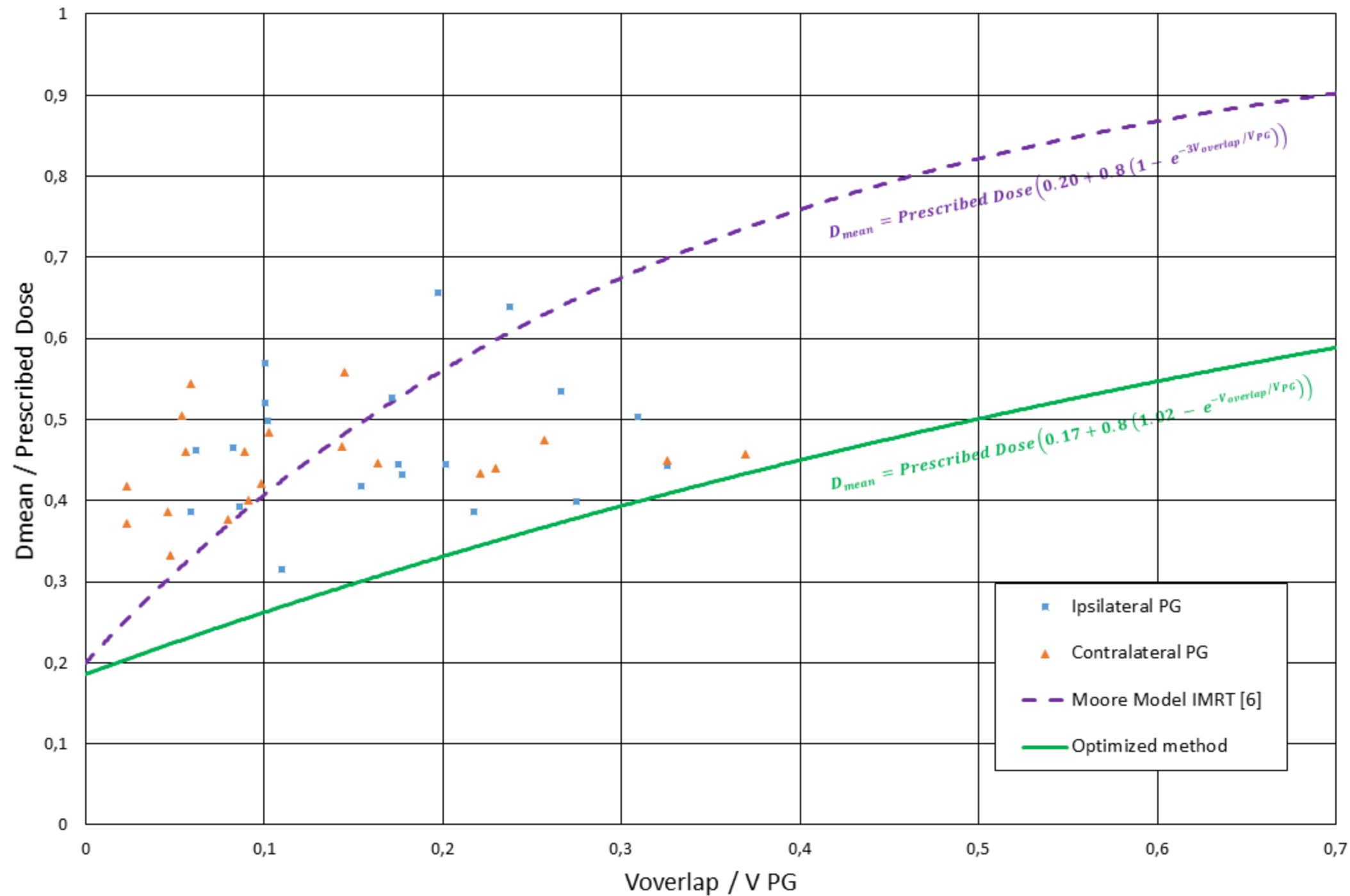
A simple and optimized mathematical model was implemented in our department to spare the PGs during H&N treatment dose planning. Using the achievable PG mean dose provided by the optimized model, the PG mean dose decreased of 6.1 Gy on average. Moreover, the optimized model allows to significantly reduce the inter-operator variability, whatever the planner's experience. The optimized model has been used in our daily clinical practices for the last two years. The benefits of PG dose sparing need to be evaluated in terms of clinical benefits (e.g., xerostomia, local control).

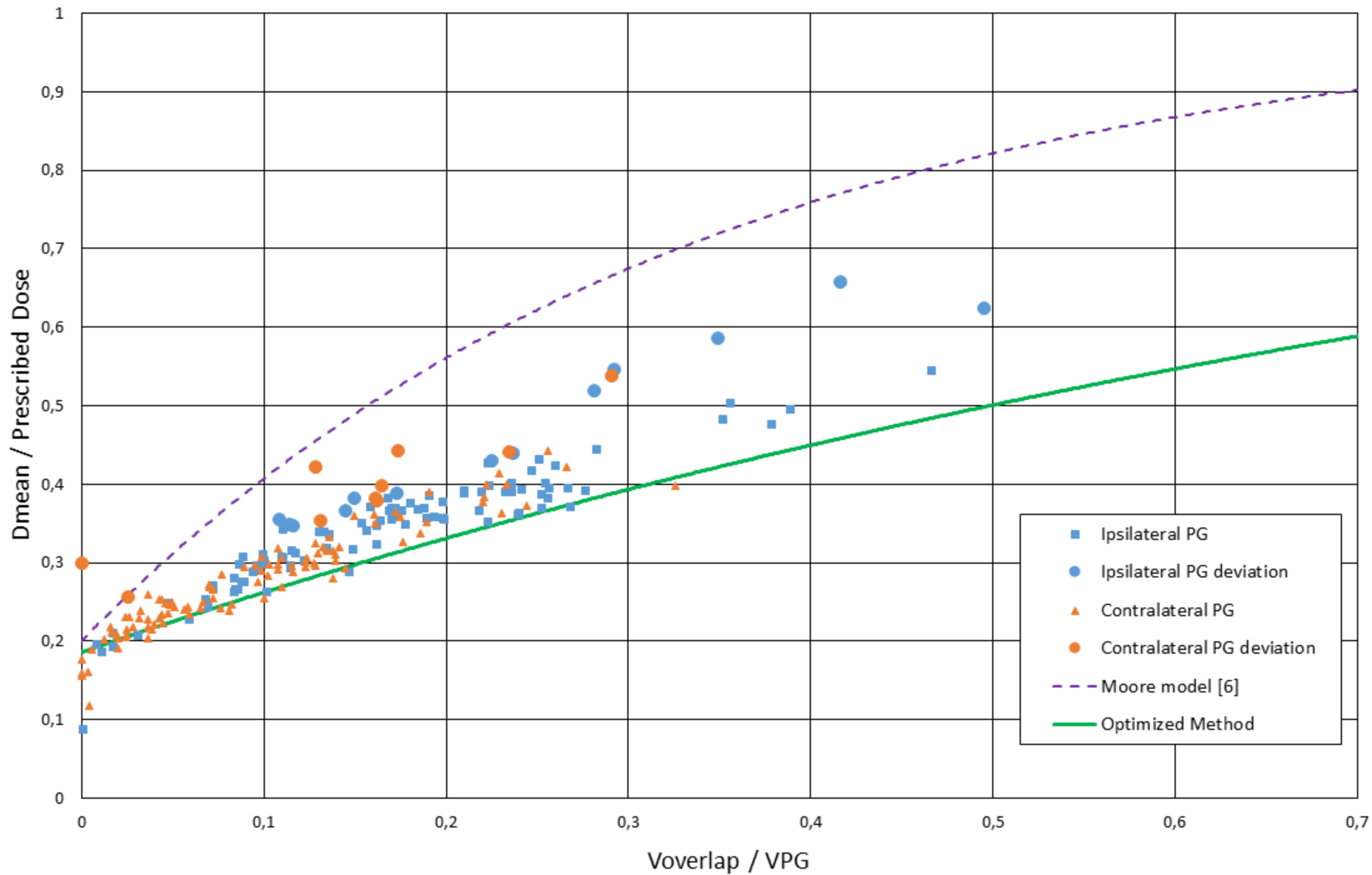
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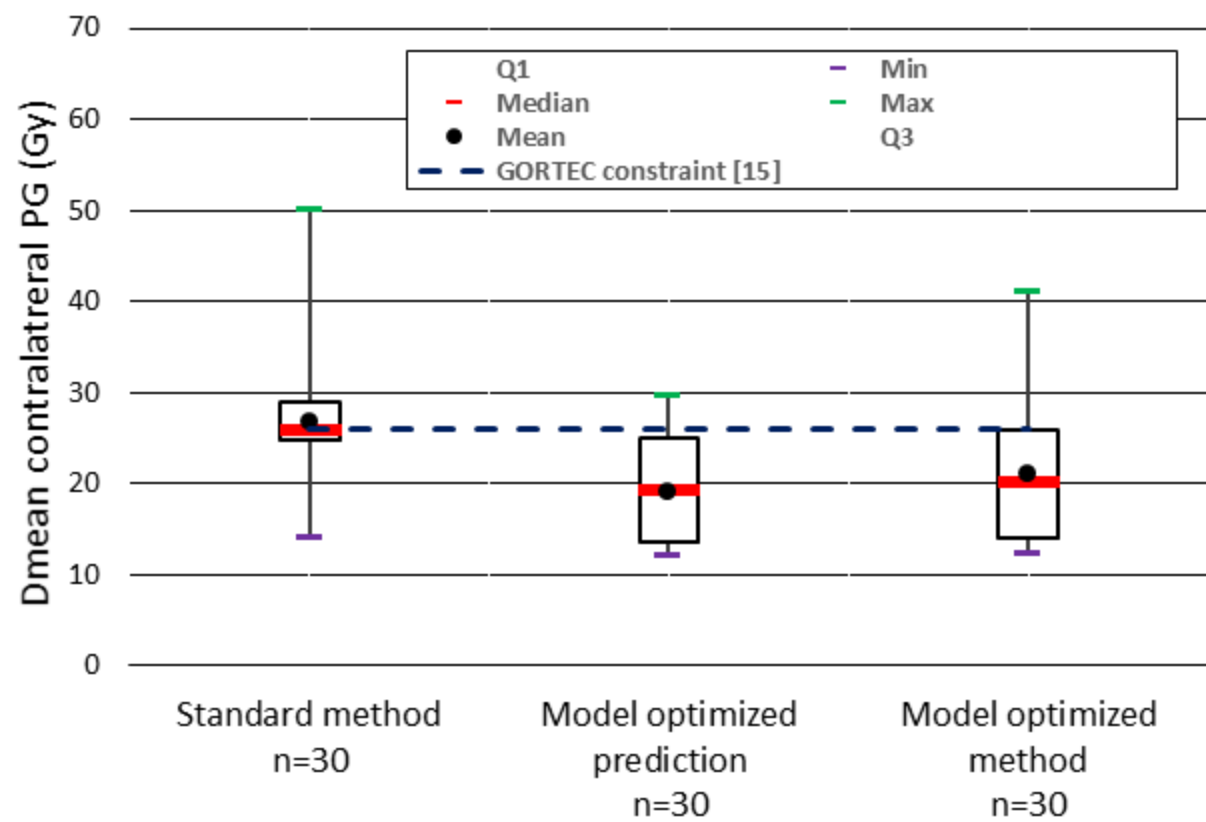
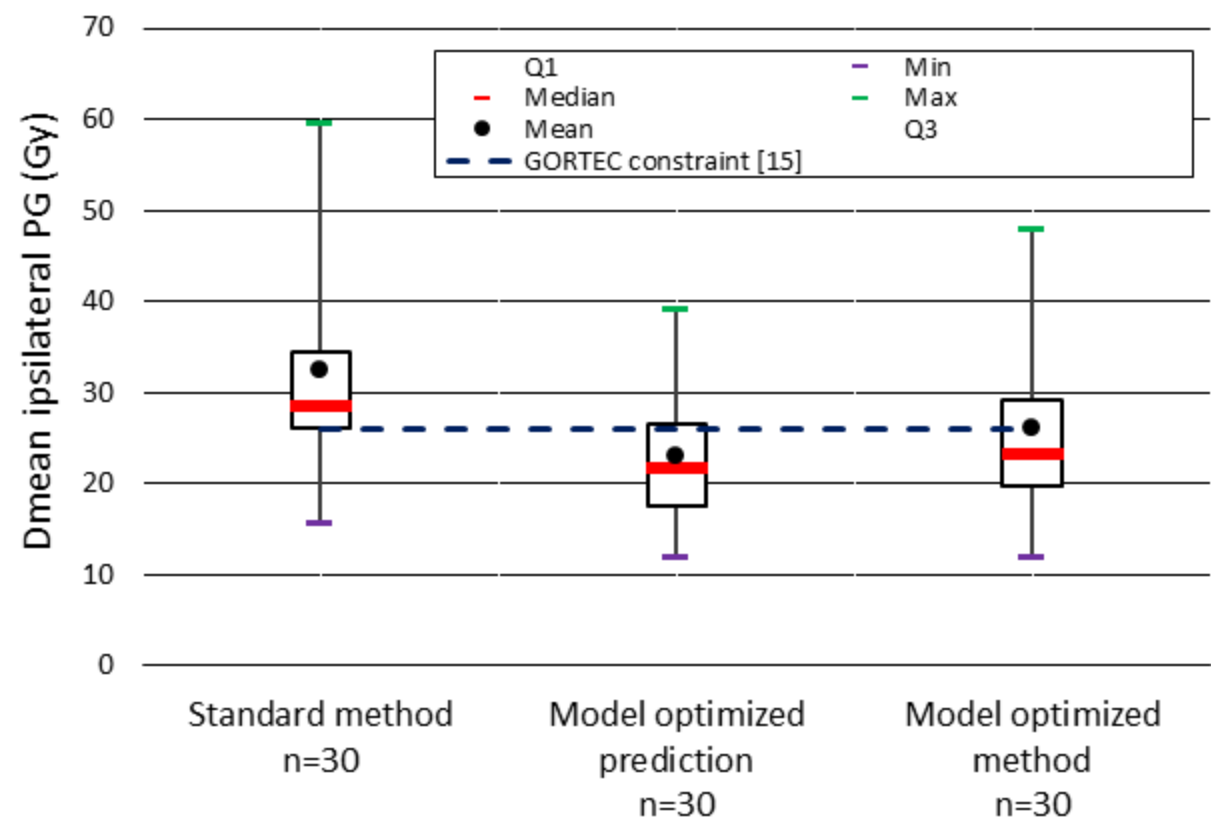




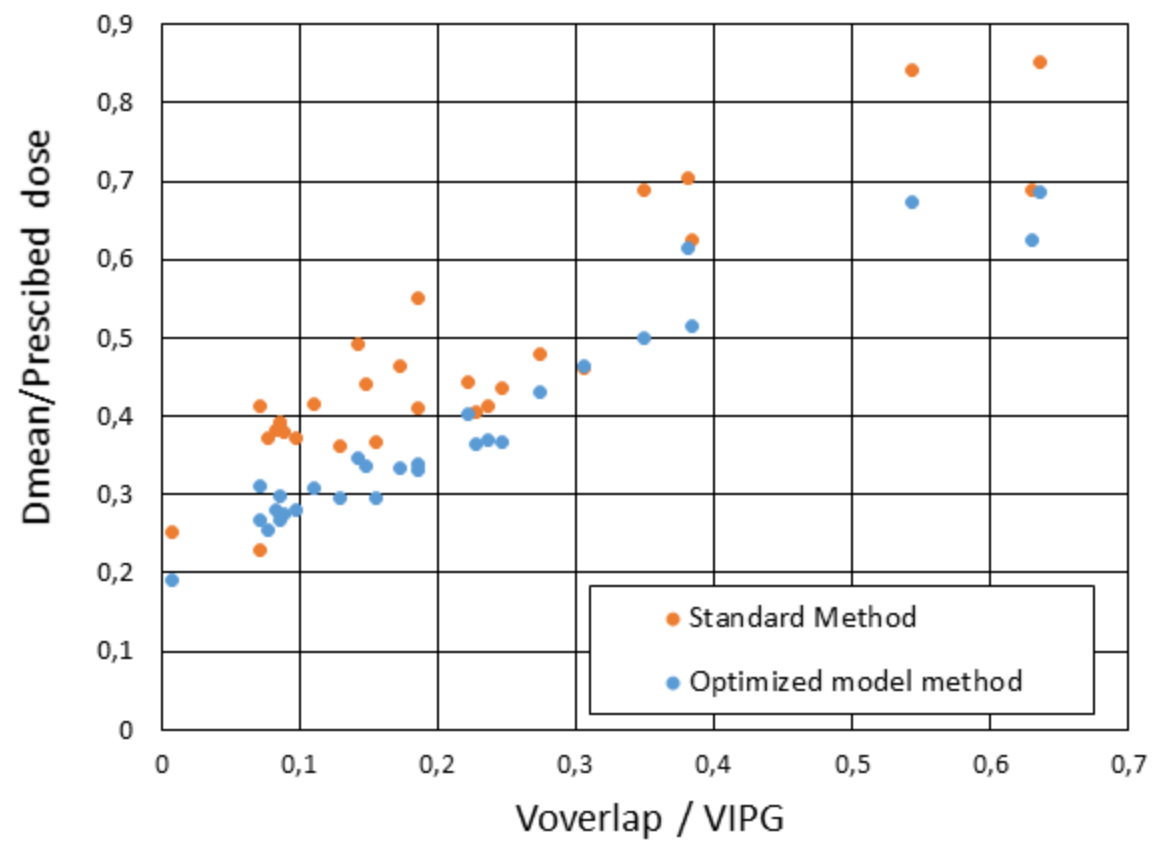




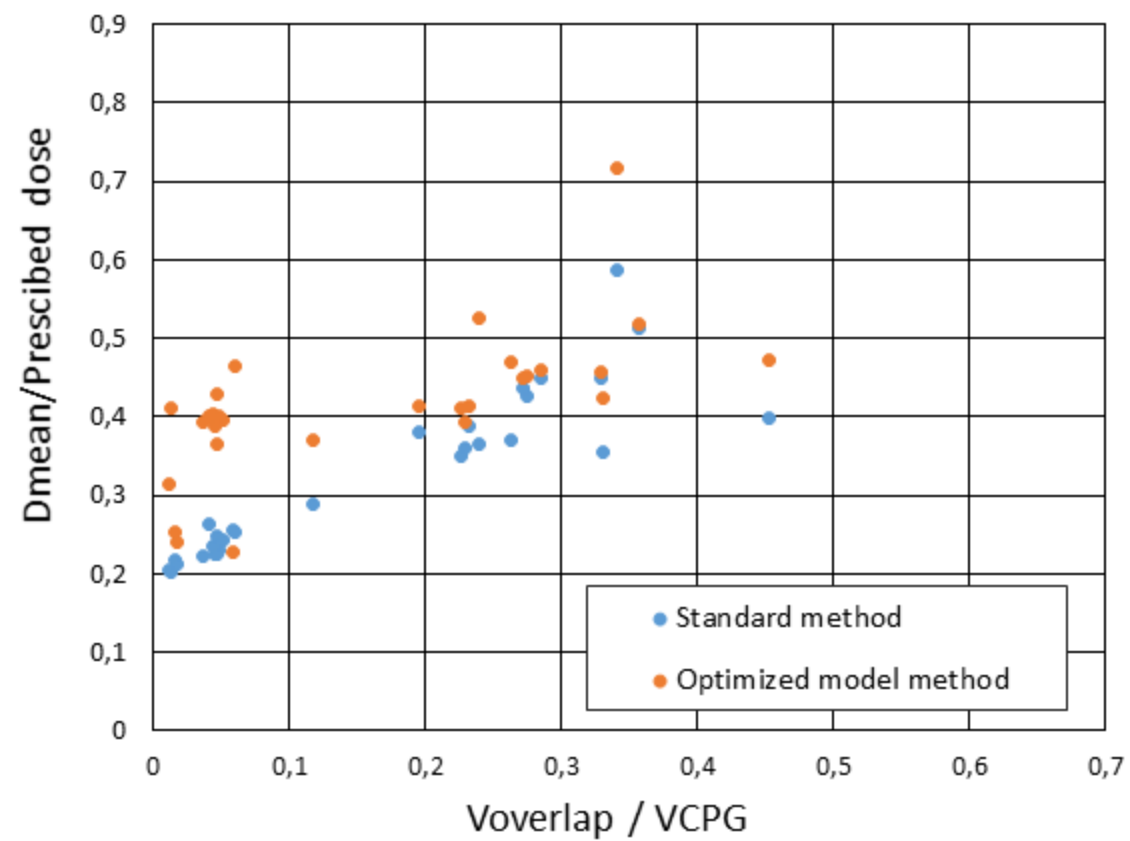


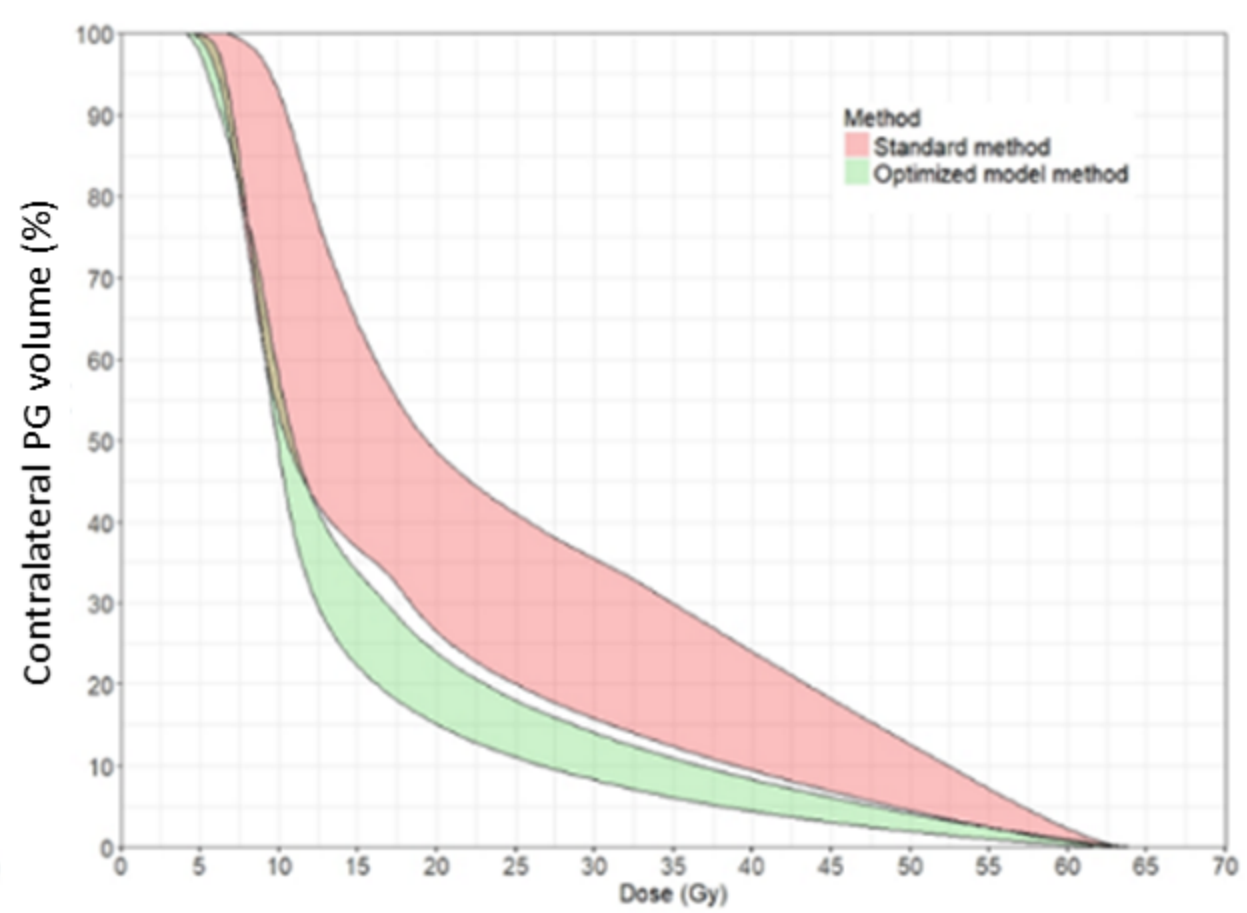
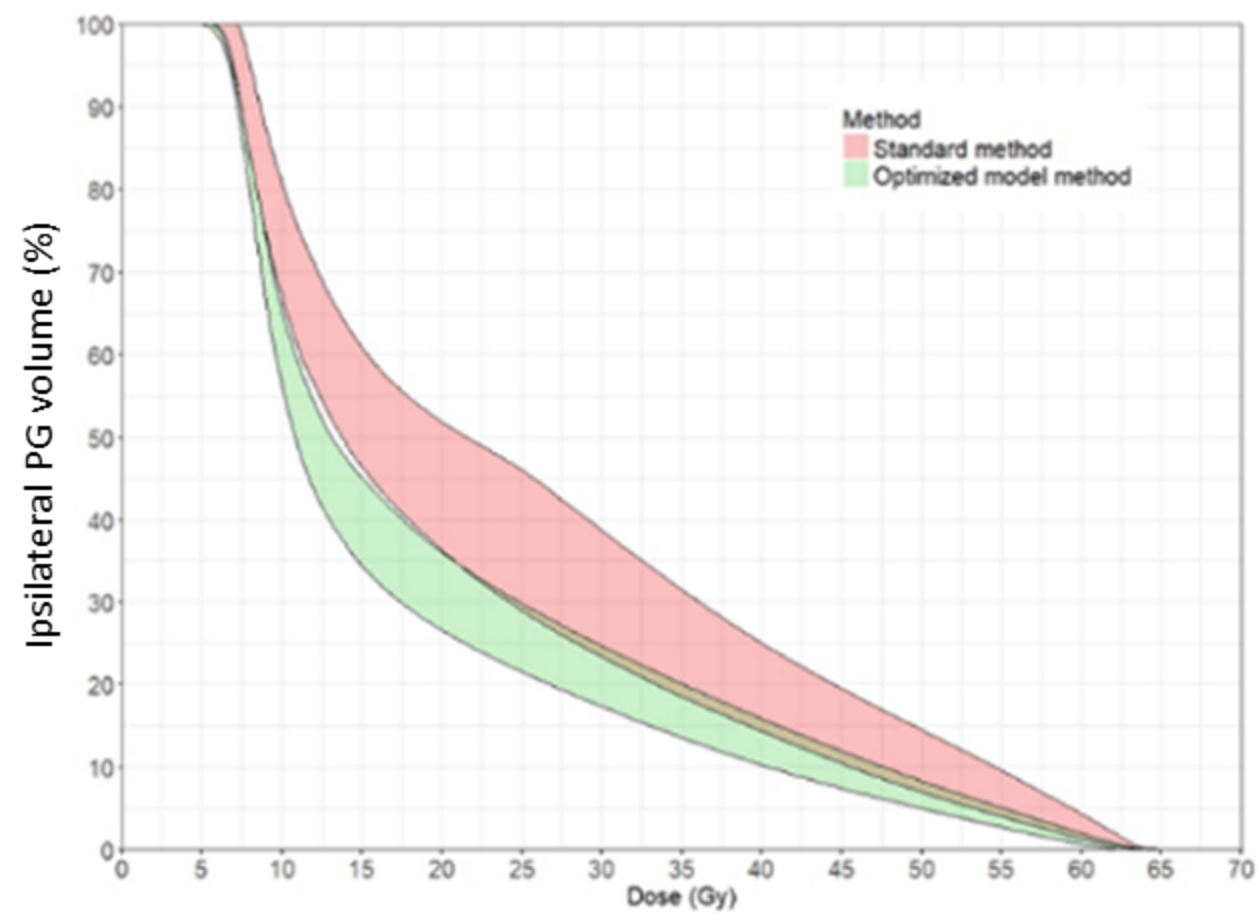


a/



b/





	Cohort size (number of patients)	Dose prescription (number of patients)			PG volume (mean $\pm$ SD, in cm <sup>3</sup> )		PG overlap (mean $\pm$ SD, in cm <sup>3</sup> )	
		70–63-56 Gy	66-59.4–54.1 Gy	60–54 Gy	IPG	CPG	IPG	CPG
<b>Model optimization</b>	20	10	4	6	28.9 $\pm$ 10.2	29.1 $\pm$ 9.3	4.8 $\pm$ 2.6	3.4 $\pm$ 2.1
<b>Robustness</b>	110	82	20	8	29.6 $\pm$ 8.0	30.3 $\pm$ 9.1	5.0 $\pm$ 3.0	3.0 $\pm$ 2.1
<b>Standard vs optimized model</b>	30	30	0	0	27.5 $\pm$ 10.7	26.1 $\pm$ 11.0	4.9 $\pm$ 3.0	3.0 $\pm$ 1.9
<b>Total</b>	160	122	24	14	29.1 $\pm$ 8.1	29.4 $\pm$ 9.6	4.9 $\pm$ 2.9	3.1 $\pm$ 2.1

**Table 1. Patient’s study characteristics: patient cohort size, dose prescription, PG volume and, overlap between PG and PTV**

IPG: ipsilateral parotid gland

CPG: contralateral parotid gland

		<b>Standard planning</b> Mean [min; max]	<b>With optimized model</b> Mean [min; max]	<b>p-value</b>
<b>Planned dose to PTVs and OARs (in Gy)</b>	High-dose PTV ( $D_{\text{mean}}$ )	<b>69.8</b> [67.9; 71.8]	<b>70.1</b> [67.6;71.8]	<b>0.11</b>
	Middle-dose PTV ( $D_{\text{mean}}$ )	<b>62.7</b> [62.1; 65.1]	<b>63.1</b> [61.2;65.9]	<b>0.03</b>
	Low-dose PTV ( $D_{\text{mean}}$ )	<b>56.3</b> [54.6; 57.5]	<b>56.7</b> [54.8; 57.9]	<b>0.02</b>
	Spinal Cord ( $D_{2\%}$ )	<b>34.6</b> [30.6; 37.5]	<b>38.1</b> [34.7; 41.7]	<b>&lt;0.001</b>
	Oral cavity ( $D_{\text{mean}}$ )	<b>54.9</b> [26.8 ; 67.7]	<b>54.2</b> [28.7; 66.7]	<b>0.09</b>
	Larynx ( $D_{\text{mean}}$ )	<b>56.7</b> [21.1; 83.2]	<b>57.2</b> [23.6; 71.8]	<b>0.03</b>
	Ipsilateral SMG ( $D_{\text{mean}}$ )	<b>63.0</b> [33.3; 70.8]	<b>61.9</b> [37.9; 72.2]	<b>0.41</b>
	Contralateral SMG ( $D_{\text{mean}}$ )	<b>57.9</b> [31.3; 71.1]	<b>58.6</b> [32.8; 71.7]	<b>0.18</b>
	Ipsilateral PG ( $D_{\text{mean}}$ )	<b>32.4</b> [15.9; 59.7]	<b>26.0</b> [12.0; 48.0]	<b>&lt;0.001</b>
	Contralateral PG ( $D_{\text{mean}}$ )	<b>26.9</b> [14.3; 50.2]	<b>21.1</b> [12.5; 41.2]	<b>&lt;0.001</b>
<b><math>\delta</math> index (ideal value = 0)</b>	Ipsilateral PG	<b>0.41</b> [-0.05; 0.72]	<b>0.12</b> [-0.03; 0.40]	<b>&lt;0.001</b>
	Contralateral PG	<b>0.46</b> [-0.02; 1.10]	<b>0.09</b> [-0.02; 0.41]	<b>&lt;0.001</b>
<b>NTCP</b>	Ipsilateral PG	<b>0.349</b> [0.060; 0.832]	<b>0.251</b> [0.038; 0.668]	<b>&lt;0.001</b>
	Contralateral PG	<b>0.290</b> [0.099; 0.575]	<b>0.197</b> [0.042; 0.504]	<b>&lt;0.001</b>

**Table 2. Comparison of planned doses,  $\delta$  index and NTCP (for parotid glands) between standard planning and use of optimized model during planning**

A cohort of 30 patients was considered.

			<b>Standard planning</b> Mean [min; max]	<b>Optimized model method</b> Mean [min; max]	<b>p-value</b>
<b>Dosimetric indexes</b>	<b>CI</b> (ideal value: 1)	High-dose PTV	<b>0.98*</b> [0.86; 0.99]	<b>0.98*</b> [0.86; 0.99]	<b>0.78</b>
		Middle-dose PTV	<b>0.97</b> [0.88; 1.00]	<b>0.97</b> [0.92; 0.99]	<b>0.04</b>
		Low-dose PTV	<b>0.99</b> [0.96; 1]	<b>0.99</b> [0.97; 1]	<b>0.50</b>
	<b>HI</b> (ideal value: 0)	High-dose PTV	<b>0.07</b> [0.03; 0.15]	<b>0.09</b> [0.05; 0.12]	<b>0.02</b>
		Middle-dose PTV	<b>0.17</b> [0.06; 0.22]	<b>0.21</b> [0.14; 0.25]	<b>&lt;0.001</b>
		Low-dose PTV	<b>0.20</b> [0.10; 0.31]	<b>0.20</b> [0.11; 0.29]	<b>0.52</b>
<b>Complexity indexes</b>	<b>MU</b>		<b>673.4</b> [558.0; 845.1]	<b>744.6</b> [559.9; 865.0]	<b>&lt;0.001</b>
	<b>Area (cm<sup>2</sup>)</b> (ideal value: large as possible)		<b>56.8</b> [44.0; 78.5]	<b>50.1</b> [33.1; 70.1]	<b>&lt;0.001</b>
	<b>Irregularity</b> (ideal value: 1)		<b>13.1</b> [8.5; 17.4]	<b>14.0</b> [9.6; 16.2]	<b>0.002</b>
	<b>Modulation</b> ([0;1], ideal value: 0)		<b>0.8</b> [0.73; 0.84]	<b>0.8</b> [0.76; 0.85]	<b>&lt;0.001</b>
<b>Agreement between dose calculation and measurement</b>	<b>Gamma index passrate (%)</b> (ideal value in our institution: > 85%)		<b>94.1</b> [88.8; 97.6]	<b>94.6</b> [84.5; 97.4]	<b>0.32</b>

\* = normalization point

**Table 3. Dosimetric, complexity indexes and gamma index evaluation for standard planning and use of optimized model**

Dosimetric and complexity indexes are mean [min; max], for 30 patients. Local gamma analysis (criteria: local, 3%/3 mm, low dose threshold = 30%) was performed between calculation and measurement for standard planning and use of the optimized model (mean [min; max] for 15 patients).

		<b>Standard planning</b>	<b>With optimized model</b>	<b>p-value</b>
CI (ideal value: 1)	High-dose PTV	<b>0.99 ± 0.01</b>	<b>0.98 ± 0.01</b>	<b>0.16</b>
	Middle-dose PTV	<b>0.96 ± 0.01</b>	<b>0.95 ± 0.02</b>	<b>0.02</b>
	Low-dose PTV	<b>0.97 ± 0.01</b>	<b>0.98 ± 0.01</b>	<b>0.20</b>
HI (ideal value: 0)	High-dose PTV	<b>0.07 ± 0.01</b>	<b>0.09 ± 0.01</b>	<b>0.02</b>
	Middle-dose PTV	<b>0.17 ± 0.02</b>	<b>0.20 ± 0.03</b>	<b>0.03</b>
	Low-dose PTV	<b>0.14 ± 0.01</b>	<b>0.15 ± 0.01</b>	<b>0.08</b>

**Table 4. Dosimetric indexes with standard planning and use of optimized model for the inter-operator variability assessment**

Dosimetric indexes (conformity and homogeneity) with the standard planning and the use of optimized model are mean ± standard deviation, for 12 different operators.