

Radiobiological and dosimetric impact of RayStation pencil beam and Monte Carlo algorithms on intensity-modulated proton therapy breast cancer plans

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Abstract

Purpose: RayStation treatment planning system employs pencil beam (PB) and Monte Carlo (MC) algorithms for proton dose calculations. The purpose of this study is to evaluate the radiobiological and dosimetric impact of RayStation PB and MC algorithms on the intensity-modulated proton therapy (IMPT) breast plans.

Methods: The current study included ten breast cancer patients, and each patient was treated with 1–2 proton beams to the whole breast/chestwall (CW) and regional lymph nodes in 28 fractions for a total dose of 50.4 Gy relative biological effectiveness (RBE). A total clinical target volume (CTV_{Total}) was generated by combining individual CTVs: AxI, AxII, AxIII, CW, IMN, and SCVN. All beams in the study were treated with a range shifter (7.5 cm water equivalent thickness). For each patient, three sets of plans were generated: (a) PB optimization followed by PB dose calculation (PB-PB), (b) PB optimization followed by MC dose calculation (PB-MC), and (c) MC optimization followed by MC dose calculation (MC-MC). For a given patient, each plan was robustly optimized on the CTVs with same parameters and objectives. Treatment plans were evaluated using dosimetric and radiobiological indices (equivalent uniform dose (EUD), tumor control probability (TCP), and normal tissue complication probability (NTCP)).

Results: The results are averaged over ten breast cancer patients. In comparison to PB-PB plans, PB-MC plans showed a reduction in CTV target dose by 5.3% for D_{99%} and 4.1% for D_{95%}, as well as a reduction in TCP by 1.5–2.1%. Similarly, PB overestimated the EUD of target volumes by 1.8–3.2 Gy(RBE). In contrast, MC-MC plans achieved similar dosimetric and radiobiological (EUD and TCP) results as the ones in PB-PB plans. A selection of one dose calculation algorithm over another did not produce any noticeable differences in the NTCP of the heart, lung, and skin.

Conclusion: If MC is more accurate than PB as reported in the literature, dosimetric and radiobiological results from the current study suggest that PB overestimates the target dose, EUD, and TCP for IMPT breast cancer treatment. The overestimation of dosimetric and radiobiological results of the target volume by PB needs to be further interpreted in terms of clinical outcome.

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KEY WORDS

breast cancer, dose calculation algorithms, EUD, IMPT, Monte Carlo, NTCP, TCP

1 | INTRODUCTION

Intensity-modulated proton therapy (IMPT) is used for the treatment of breast cancer at many proton centers across the world. Literature^{1,2} has shown that proton therapy for breast cancer could potentially reduce normal tissue complication probability (NTCP) by reducing side effects such as cardiac and pulmonary toxicities. It is paramount that the reduction of NTCP must be accompanied by an increase in tumor control probability (TCP) to prevent tumor recurrence. Both the TCP and NTCP are calculated based on the absorbed dose in disease sites and normal tissues, respectively. Hence, the accuracy of proton dose calculation algorithm is critical in estimating the absorbed dose in tumors and organs at risk (OARs).

RayStation (version 6.1.1.2; RaySearch Laboratories, Stockholm, Sweden) employs analytical pencil beam (PB) and Monte Carlo (MC) algorithms for proton dose calculations. Several studies^{3–8} have highlighted the limitation of PB algorithm within RayStation for dose calculation, especially in the presence of range shifter and inhomogeneities. For instance, Saini et al.³ found that MC is superior to PB when a range shifter is employed with oblique beams, large air gaps, and/or heterogeneous media. Taylor et al.⁴ demonstrated that MC calculations are more accurate than PB calculations when compared to physical measurements. Shirey et al.⁶ showed better accuracy of MC compared to PB when treatment involves the range shifter and superficial lesions.

Although superior dose prediction accuracy of MC over PB has been well established,^{3–8} literature investigating the impact of RayStation PB and MC on IMPT breast cancer treatment is scarce. The investigation of proton dose calculation algorithms for breast treatment is particularly important due to the presence of a tumor at a shallower depth and range shifter in the proton beam path. The addition of range shifter at the end of the nozzle exit reduces the beam energy. This is necessary to achieve a full dose modulation of the tumor volume, but the range shifter creates an air gap between its downstream and patient surface, thus increasing in-air spot size.⁹

Tommasino et al.⁷ included five breast cancer patients and the treatment plans were optimized with PB and recalculated with PB and MC. Additionally, Tommasino et al.⁷ performed the phantom measurements to demonstrate better accuracy using MC than using PB. Liang et al.⁸ did a more comprehensive dosimetric study by including 20 breast cancer patients. In their study,⁸ the authors used both PB and MC for plan optimization as well as dose calculations, whereas MC for plan optimization was not addressed by Tommasino et al.⁷ It is worth noting that both studies evaluated PB and MC on IMPT breast plans using dosimetric indices. However, at the time of

writing this paper, the radiobiological impact of RayStation PB and MC on IMPT breast plans is yet to be investigated.

The goal of this study is twofold. First, we investigated the radiobiological impact of RayStation PB and MC algorithms on IMPT breast plans. Specifically, treatment plans are evaluated in terms of equivalent uniform dose (EUD), TCP, and NTCP. Our study included plans optimized using PB and MC as well as dose calculated using PB and MC. Second, since there is only one dosimetric study⁸ from a single institution investigating the use of RayStation MC for plan optimization of IMPT breast cancer planning, independent research from another institution on this topic is essential. Our study aims to supplement the work of Liang et al.⁸ by comparing the dosimetric results of PB and MC for IMPT breast cancer treatment. Additionally, for each case, we have presented a more comprehensive analysis of plan robustness and computational time — results of these two parameters were not provided in detail in a previous publication.⁸

2 | MATERIALS AND METHODS

2.A | Patients, CT simulation, and contouring

The current study consisted of ten female left breast cancer patients who have been treated using IMPT at our center between 11/2017 and 01/2019. All ten patients received treatment to the chest wall (CW) or whole breast. For all ten patients, the treatment also included regional lymph nodes. Patients were simulated on Siemens computed tomography (CT) Scanner (Siemens Healthcare, Forchheim, Germany) in head-first supine treatment position with arms above their heads based on our institutional protocol. This includes a vacuum and wing board for immobilization devices and a free breathing CT scan with a slice thickness of 2 mm.

For contouring of target volumes and OARs, CT images were transferred either to RayStation or Velocity (Varian Medical Systems, Palo Alto, CA, USA). A total clinical target volume (CTV_{Total}) was generated by combining individual CTV structures: breast or CW, axillary level I–III nodes (Axl-III), internal mammary nodes (IMN), and supraclavicular nodes (SCVN). The OARs included the heart, left lung, right lung, esophagus, left anterior descending artery (LAD), and skin (either 3 mm (CW) or 5 mm (whole breast) inward from the body surface).

2.B | Dose prescription and treatment planning

All ten patients were treated for a total dose of 50.4 Gy relative biological effectiveness (RBE) in 28 fractions on a ProteusPLUS PBS

proton therapy system¹⁰ (Ion Beam Applications, Louvain-la-Neuve, Belgium). Treatment plans were generated in RayStation (v6.1.1.2) using 1–2 beams, and each beam included the range shifter of 7.5 cm water equivalent thickness made up of lucite. A 5 mm setup uncertainty on CTV was used for the robust optimization for a total of seven scenarios. All treatment plans were robustly optimized with the goal of 95% of CTV receiving at least 95% of the prescription dose while minimizing dose to the OARs. All plans were computed with a dose calculation grid size of 3 mm. For each case, three plans were generated using identical beam angles, air gap, optimization structures, optimization constraints, and optimization settings. A sampling history of 10,000 ions/spot was used for MC optimization, and a statistical uncertainty of 0.5% was used for MC dose calculation.

1. PB-PB Plan: The plan was optimized using PB followed by dose calculation using PB.
2. PB-MC Plan: The plan was optimized using PB followed by dose calculation using MC.
3. MC-MC Plan: The plan was optimized using MC followed by dose calculation using MC.

2.C | Dosimetric analysis

The CTV_Total was evaluated for the mean dose (D_{mean}), the dose received by 99% of the volume ($D_{99\%}$), 95% of the volume ($D_{95\%}$), and 2% of the volume ($D_{2\%}$). The D_{mean} was calculated for the left anterior descending artery (LAD), heart, and esophagus, whereas the dose received by 0.03 cc (D_{max}) was calculated for the skin. The ipsilateral lung (i.e., left lung) was evaluated for the relative volume that received 20 and 5 Gy(RBE) (V_{20} and V_5 , respectively), whereas the contralateral lung (i.e., right lung) was evaluated for the V_5 .

2.D | EUD Analysis

Equivalent uniform dose evaluation was performed using the cumulative dose volume histograms (DVHs) of the proton treatment plans (PB-PB, PB-MC, and MC-MC). EUD is based on the Niemierko's phenomenological model.¹¹

The EUD^{11,12} is defined as.

$$\text{EUD} = \left(\sum_{i=1}^M (v_i \text{EQD}_i^a) \right)^{\frac{1}{a}} \quad (1)$$

$$\text{EQD} = D \times \frac{\left(\frac{\alpha}{\beta} + \frac{D}{n_f} \right)}{\left(\frac{\alpha}{\beta} + 2 \right)} \quad (2)$$

In eq. (1), a is a unit less model parameter that is specific to the normal structure or tumor of interest, and v_i is unit less and represents the i^{th} partial volume receiving dose D_i in Gy.^{11,12} Since the relative volume of the whole structure of interest corresponds to 1, the sum of all partial volumes v_i will equal 1.^{11,12} The EQD is the biologically equivalent physical dose of 2 Gy. In eq. (2), n_f and $d_f = D/n_f$ are the number of fractions and dose per fraction size of the treatment course,

respectively. The α/β is the tissue-specific linear-quadratic (LQ) parameter of the organ being exposed. The EUD calculations in this study are based on the parameters listed in Table 1.^{14–16}

2.E | TCP Analysis

The Poisson linear quadratic (PoissonLQ) radiobiological model¹³ employed within RayStation was used to estimate the TCP of CTV_Total, CTV_breast, CTV_AxI, CTV_AxII, CTV_AxIII, CTV_IMN, and CTV_SCVN. The TCP-PoissonLQ model is defined as¹³:

$$\text{TCP}(D) = \prod_{i=1}^M \left[\exp \left(-N_0 \exp \left(\sum_{k=1}^n \left\{ -\alpha d_{k,i} - \beta d_{k,i}^2 \right\} \right) \right) \right]^{\frac{v_i}{V_{\text{ref}}}} \quad (3)$$

$$\text{TCP}(D) = \prod_{i=1}^M \left[\exp \left(- \exp \left[e_{\gamma} - \frac{\text{EQD}_{2,i}}{D_{50}} (e_{\gamma} - \ln(\ln(2))) \right] \right) \right]^{\frac{v_i}{V_{\text{ref}}}}$$

where, M , total number of voxels; D , total dose; $D_{k,i}$, dose to the k^{th} fraction to voxel i ; n , total number of fractions; N_0 , initial number of cells; α , parameter of LQ model; β , parameter of LQ model; v_i/V_{ref} , relative volume of voxel i compared to the reference volume; D_{50} , dose giving a 50% response probability; γ , maximum normalized gradient of the dose response curve; $\text{EQD}_{2,i}$, equivalent dose in voxel i given in 2 Gy-fractions.

The values of radiobiological parameters^{14–16} used for TCP calculations are provided in Table 1.

2.F | NTCP Analysis

The Lyman-Kutcher-Burman (LKB) model employed within RayStation¹³ was used to calculate the NTCP of the heart, lung (ipsilateral), and skin. The LKB model is defined as¹³:

$$\text{NTCP}(D) = \frac{1}{\sqrt{2\pi}} \int_{-\infty}^t e^{-\frac{x^2}{2}} dx \quad (4)$$

$$t = \frac{D_{\text{eff}} - D_{50}}{m \cdot D_{50}}$$

$$D_{\text{eff}} = \sum_{i=1}^M \left(\frac{v_i}{V_{\text{ref}}} \text{EQD}_i^{1/n} \right)^n \quad (5)$$

where, D , total dose; D_{50} , dose giving a 50% response probability; m , slope of the response curve; M , total number of voxels; n , parameter reflecting the biological properties of the organ specifying volume dependence; v_i/V_{ref} , relative volume of voxel i compared to the reference volume; EQD_i , equivalent dose in voxel i given in 2 Gy-fractions.

TABLE 1 Radiobiological parameters of EUD & TCP calculations for the breast cancer plans.

Parameter	Values	Reference
D_{50} (Gy(RBE))	30.89	14,15
γ	1.3	14,15
α/β	4	16
a	-7.2	14,15

EUD, equivalent uniform dose; TCP, tumor control probability; RBE, relative biological effectiveness.

TABLE 2 Radiobiological parameters of NTCP calculations for the breast cancer plans.

Structure	D ₅₀ (Gy(RBE))	m	n	Reference
Heart	48	0.1	0.35	17,18
Lung (ipsilateral)	37.6	0.35	0.87	18,19
Skin	39	0.14	0.38	20

NTCP, normal tissue complication probability; RBE, relative biological effectiveness.

The values of radiobiological parameters^{17–20} used for NTCP calculations are provided in Table 2.

2.G | Statistical analysis

In order to test the statistical significance of dosimetric and radiobiological results in the current study, the Mann-Whitney U-test was performed. A p value of less than 0.05 was considered to be statistically significant.

3 | RESULTS

3.A | Dosimetric analysis

Table 3 shows the dosimetric results of nominal PB-PB, PB-MC, and MC-MC plans. The results are averaged over ten breast cancer patients. The recalculation of PB plans with MC showed the reduction in dose to the CTV_Total by the average differences of 5.3% for D_{99%} ($P = 0.001$), 4.1% for D_{95%} ($P < 0.001$), 2.7% for D_{mean} ($P < 0.001$), and 1% for D_{2%} ($P = 0.112$). The doses to the CTV_Total in MC-MC and PB-PB plans were comparable with no statistical significance ($P > 0.05$). Specifically, on average, the difference in

CTV_Total dose between MC-MC and PB-PB plans was less than 0.5% for both D_{95%} and D_{mean} and 1.4% for D_{99%}. The D_{2%} was similar in both MC-MC and PB-PB plans. The current study used the treatment planning goal of CTV_Total D_{95%} = 95% of the prescription dose. On average, the CTV_Total D_{95%} was 97.5%, 93.5%, and 97.1% in PB-PB, PB-MC, and MC-MC plans, respectively. For PB-PB plans, nine patients had CTV_Total D_{95%} > 95% and one patient had CTV_Total D_{95%} = 94.8%. MC-MC plans also exhibited similar results such that eight patients had CTV_Total D_{95%} > 95% and two patients had CTV_Total D_{95%} = 94.6% and 94.8%. However, PB-MC plans produced inferior results, and there was only one patient with CTV_Total D_{95%} = 95.1%, and the other nine patients had CTV_Total D_{95%} results ranging from 91.4% to 94.5%. Figure 1 shows a sample DVH of CTV_Total in PB-PB, PB-MC, and MC-MC plans of an example patient. Figure 2 shows the dose distributions in PB-PB, PB-MC, and MC-MC plans of an example patient.

The average difference in D_{mean} to the heart, LAD, and esophagus among different plans (PB-PB, PB-MC, and MC-MC) was less than 0.5 Gy(RBE) ($P > 0.05$). The difference in D_{max} to the skin between PB-MC and PB-PB plans ranged from –2.3% to 3.7% with no statistical significance ($P = 0.290$). The positive difference means PB-MC plan has higher D_{max} than PB-PB plan. MC-MC plans produced consistently higher D_{max} to the skin except in one case. The difference in D_{max} to the skin between MC-MC and PB-PB plans ranged from –0.3% to 1.8% with no statistical significance ($P = 0.406$). For the ipsilateral lung, the average V₂₀ among PB-PB, PB-MC, and MC-MC plans was similar (12.4% vs. 13.9% vs. 22.3%, respectively). However, in comparison to PB-PB plans, the difference in V₅ of the ipsilateral lung was slightly higher in PB-MC plans (2.7% to 8%; $P = 0.082$) and MC-MC plans (0.7% to 13.8%; $P = 0.049$). The average V₅ of the contralateral lung was similar among PB-PB (1.7%), PB-MC (0.9%), and MC-MC (1.1%) plans ($P > 0.05$).

TABLE 3 Dosimetric results in nominal PB-PB, PB-MC, and MC-MC plans of breast cancer. The results are averaged over ten breast cancer patients.

		PB-PB Avg. (range)	PB-MC Avg. (range)	P-value	MC-MC Avg. (range)	P-value
CTV_Total	D _{99%} (%)	95.5 (91.6–98.8)	90.4 (87.9–92.6)	<0.001	94.1 (92.1–98.1)	0.174
	D _{95%} (%)	97.5 (94.8–99.6)	93.5 (91.4–95.1)	<0.001	97.1 (94.6–99.3)	0.385
	D _{mean} (%)	100.3 (98.4–101.5)	97.6 (95.3–98.6)	<0.001	100 (98.2–101.2)	0.290
	D _{2%} (%)	102.6 (100.2–104.3)	101.6 (98.0–104.1)	0.112	102.7 (100.1–104.6)	0.821
Heart	D _{mean} (Gy(RBE))	0.45 (0.12–1.07)	0.47 (0.1–0.95)	0.762	0.47 (0.11–1.05)	0.970
LAD	D _{mean} (Gy(RBE))	3.37 (0.43–9.21)	3.10 (0.43–9.06)	0.764	3.64 (0.33–11.17)	0.910
Esophagus	D _{mean} (Gy(RBE))	6.02 (3.33–17.24)	5.94 (3.63–17.53)	0.705	6.28 (3.71–19.26)	0.850
Skin	D _{max} (%)	100.2 (94.6–103.1)	101.2 (95.2–105.8)	0.290	100.9 (94.8–104)	0.406
Left lung	V ₂₀ (%)	12.4 (2.2–22.7)	13.9 (2.6–25.3)	0.406	13.1 (4.9–24.5)	0.597
Left lung	V ₅ (%)	32.8 (21.6–47.5)	37.6 (26.7–53.4)	0.082	36.8 (31.1–51.9)	0.049
Right lung	V ₅ (%)	1.7 (0–9.4)	0.9 (0–4.1)	0.597	1.1 (0–4.2)	0.940

PB-PB, PB optimization followed by PB dose calculation; PB-MC, PB optimization followed by MC dose calculation; MC-MC, MC optimization followed by MC dose calculation; CTV, clinical target volume; LAD, left anterior descending artery; RBE, relative biological effectiveness.

^aP-value for PB-MC vs. PB-PB.

^bP-value for MC-MC vs. PB-PB.

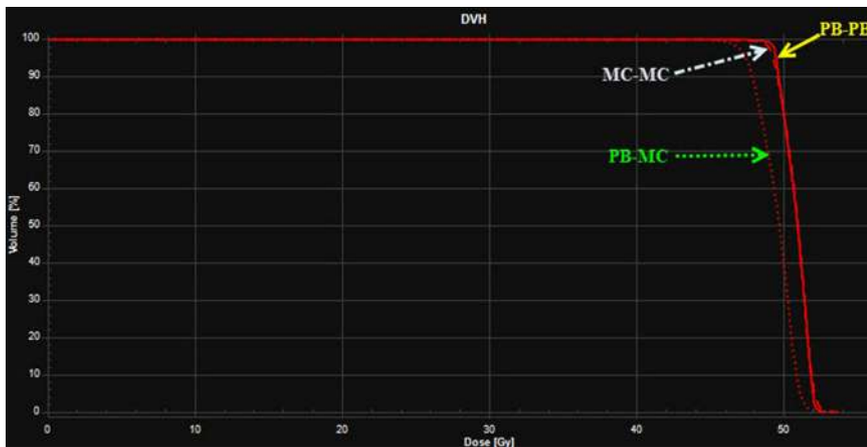


Fig. 1. A sample dose volume histograms of clinical target volume (CTV_Total) in PB optimization followed by PB dose calculation, PB optimization followed by MC dose calculation, and MC optimization followed by MC dose calculation plans of an example patient. Treatment planning goal: CTV_Total D95% = 95% of prescription dose (50.4 Gy relative biological effectiveness (RBE)).

3.B | Robust analysis

The robust analysis was carried out in all three sets of plans (PB-PB, PB-MC, and MC-MC), and each plan was evaluated for a total of

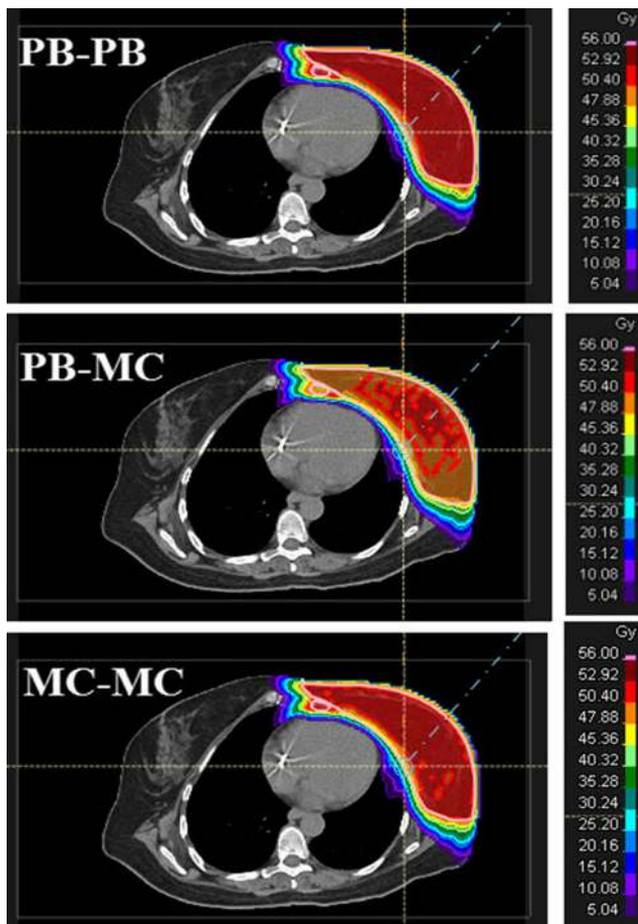


Fig. 2. Dose distributions in PB optimization followed by PB dose calculation, PB optimization followed by MC dose calculation, and MC optimization followed by MC dose calculation plans of an example patient. Treatment planning goal: clinical target volume (CTV_Total) D95% = 95% of prescription dose (50.4 Gy relative biological effectiveness (RBE)).

eight scenarios. Range uncertainty was evaluated for $\pm 3.5\%$ and isocenter shift was evaluated for $X = \pm 5$ mm, $Y = \pm 5$ mm, and $Z = \pm 5$ mm. The acceptable robustness criteria for IMPT breast treatment was 95% of the CTV_Total is covered by at least 90% of the prescribed dose (i.e., $D_{95\%} \geq 45.36$ Gy(RBE)). Figure 3 illustrates the robust evaluation of the CTV_Total for all ten patients. The results showed that all three sets of plans (PB-PB, PB-MC, and MC-MC) achieved the robustness criteria for the IMPT breast treatment ($D_{95\%} \geq 90\%$ of prescription dose) in all ten patients in the current study.

3.C | EUD analysis

Tables 4 and 5 show the EUD results in PB-PB, PB-MC, and MC-MC plans. For all seven target volumes of each patient in the current study, a reduction in EUD ($p \leq 0.001$) was noticed when PB optimized plans are calculated using MC. On average, EUD in PB-MC plans was reduced by 2.3 Gy(RBE) for CTV_AxI, 1.8 Gy(RBE) for CTV_AxII, 1.8 Gy(RBE) for CTV_AxIII, 1.9 Gy(RBE) for CTV_CW/breast, 3.2 Gy(RBE) for CTV_IMN, and 3.1 Gy(RBE) for CTV_SCVN. The comparison between MC-MC and PB-PB plans showed that EUD was comparable ($P > 0.05$): 50.7 Gy(RBE) vs. 51.0 Gy(RBE) for AxI, 50.6 Gy(RBE) vs. 51.0 Gy(RBE) for AxII, 50.4 Gy(RBE) vs. 50.7 Gy(RBE) for AxIII, 50.1 Gy(RBE) vs. 50.4 Gy(RBE) for CTV_CW/breast, 50.1 Gy(RBE) vs. 50.5 Gy(RBE) for CTV_IMN, and 50.2 Gy(RBE) vs. 50.5 Gy(RBE) for CTV_SCVN. EUD results of CTV_Total are displayed in Fig. 4. In comparison to PB-PB plans, EUD of CTV_Total was comparable in MC-MC plans (50.4 Gy(RBE) vs. 50.2 Gy(RBE); $P = 0.430$), whereas PB-MC plans showed reduction ($P < 0.001$) in EUD of CTV_Total by an average difference of 1.9 Gy (RBE).

3.D | TCP analysis

Table 6 and Figs. 4 and 5 show the TCP results in PB-PB, PB-MC, and MC-MC plans. The results are averaged over ten breast cancer patients. In comparison to PB-PB plans, PB-MC plans consistently showed the reduction in TCP by an average difference of 1.9%

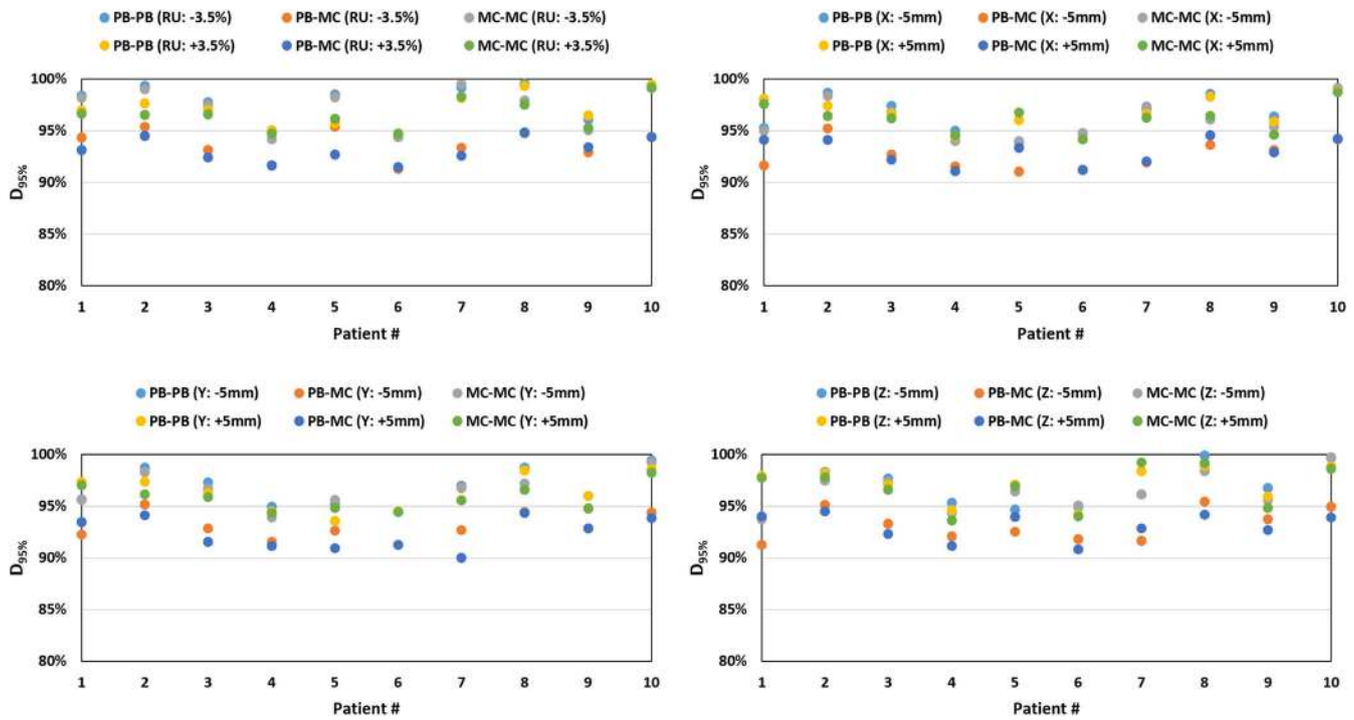


FIG. 3. Robust evaluation of the D_{95%} of the total clinical target volume (CTV_{Total}) in PB-PB (PB optimization followed by PB dose calculation), PB-MC (PB optimization followed by MC dose calculation), and MC-MC (MC optimization followed by MC dose calculation) plans for ten breast cancer patients. Each plan was evaluated for range uncertainty of ±3.5% and isocenter shifts of X = ±5 mm, Y = ±5 mm, and Z = ±5 mm. The acceptable robustness criteria for the intensity-modulated proton therapy breast was 95% of the CTV_{Total} is covered by at least 90% of the prescribed dose (i.e., D_{95%} ≥ 45.36 Gy(RBE)).

TABLE 4 EUD of CTV_{AxI}, CTV_{AxII}, and CTV_{AxIII} in nominal PB-PB, PB-MC, and MC-MC plans of breast cancer.

Patient #	CTV _{AxI}			CTV _{AxII}			CTV _{AxIII}		
	EUD (Gy(RBE))			EUD (Gy(RBE))			EUD (Gy(RBE))		
	PB-PB	PB-MC	MC-MC	PB-PB	PB-MC	MC-MC	PB-PB	PB-MC	MC-MC
1	51.7	49.6	51.7	51.6	50.4	51.6	50.9	50.0	50.9
2	51.8	49.9	51.4	51.7	49.8	51.3	51.4	50.0	51.1
3	50.6	48.2	50.5	50.6	48.8	50.5	50.6	49.1	50.5
4	50.5	48.9	49.5	50.5	49.3	49.5	49.9	47.3	49.7
5	51.6	49.2	51.4	51.8	50.3	51.7	51.3	49.5	50.8
6	49.9	47.7	49.8	49.8	47.4	49.8	49.9	47.3	49.7
7	50.8	47.1	50.9	50.8	48.6	50.8	50.8	48.6	50.8
8	51.2	49.0	50.3	51.2	49.4	50.3	51.2	49.6	50.2
9	51.0	49.2	50.2	50.9	49.2	50.0	50.8	49.2	49.8
10	50.8	48.6	50.8	50.7	48.8	50.7	50.7	48.6	50.7
Average	51.0	48.7	50.7	51.0	49.2	50.6	50.7	48.9	50.4
SD	0.6	0.9	0.7	0.6	0.9	0.7	0.5	1.0	0.5
P-value		<0.001	0.271		<0.001	0.272		0.001	0.162

PB-PB, PB optimization followed by PB dose calculation; PB-MC, PB optimization followed by MC dose calculation; MC-MC, MC optimization followed by MC dose calculation; CTV, clinical target volume, EUD, equivalent uniform dose; RBE, relative biological effectiveness.

^aP-value for PB-MC vs. PB-PB.

^bP-value for MC-MC vs. PB-PB.

($P < 0.001$) for CTV_{AxI}, 1.5% ($P = 0.002$) for CTV_{AxII}, 1.7% ($P = 0.001$) for CTV_{AxIII}, 1.7% ($P = 0.002$) for CTV_{CW/breast}, 2.8% ($P = 0.003$) for CTV_{IMN}, 2.9% ($P < 0.001$) for CTV_{SCVN}, and 1.8%

($P < 0.001$) for CTV_{Total}. In contrast, MC-MC plans achieved TCP results similar to the ones in PB-PB plans. On average, TCP results were similar in PB-PB and MC-MC plans for all CTV structures:

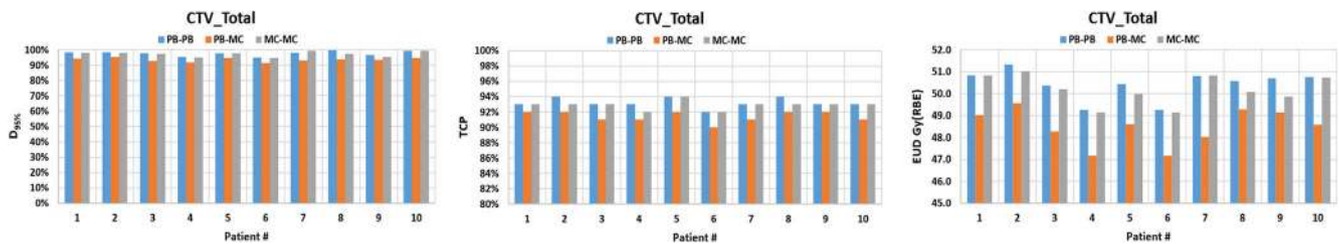
TABLE 5 EUD of CTV_CW/Breast, CTV_IMN, and CTV_SCVN in nominal PB-PB, PB-MC, and MC-MC plans of breast cancer.

Patient #	CTV_CW/Breast			CTV_IMN			CTV_SCVN		
	EUD (Gy(RBE))			EUD (Gy(RBE))			EUD (Gy(RBE))		
	PB-PB	PB-MC	MC-MC	PB-PB	PB-MC	MC-MC	PB-PB	PB-MC	MC-MC
1	50.7	48.9	50.7	51.1	49.1	51.1	51.3	48.1	51.3
2	51.3	49.6	51.0	51.0	48.3	51.0	51.2	48.8	51.7
3	50.4	48.3	50.2	50.2	44.7	49.7	50.2	47.3	50.0
4	49.1	47.1	49.0	49.3	46.1	49.1	49.0	45.8	48.8
5	50.2	48.5	49.2	51.4	47.8	50.7	51.4	48.3	50.8
6	49.1	47.1	49.0	49.3	46.1	49.1	49.0	45.8	48.8
7	50.9	48.4	51.0	52.1	48.9	52.0	51.0	47.1	50.6
8	51.0	49.4	50.1	51.1	48.1	50.3	51.1	48.2	49.9
9	50.7	49.2	49.9	48.9	46.0	47.8	50.7	47.6	49.8
10	50.8	48.7	50.7	50.5	47.5	50.4	50.6	47.2	50.6
Average	50.4	48.5	50.1	50.5	47.3	50.1	50.5	47.4	50.2
SD	0.7	0.9	0.8	1.0	1.5	1.2	0.9	1.0	1.0
P-value		<0.001	0.289		<0.001	0.447		<0.001	0.327

PB-PB, PB optimization followed by PB dose calculation; PB-MC, PB optimization followed by MC dose calculation; MC-MC, MC optimization followed by MC dose calculation; CTV, clinical target volume; CW, chestwall; EUD, equivalent uniform dose; IMN, internal mammary nodes; SCVN, supraclavicular nodes; RBE, relative biological effectiveness.

^aP-value for PB-MC vs. PB-PB.

^bP-value for MC-MC vs. PB-PB.

**FIG. 4.** $D_{95\%}$, tumor control probability, and equivalent uniform dose of the total clinical target volume (CTV_Total) for breast cancer patients ($n = 10$) in PB-PB (PB optimization followed by PB dose calculation), PB-MC (PB optimization followed by MC dose calculation), and MC-MC (MC optimization followed by MC dose calculation) plans generated by intensity-modulated proton therapy (IMPT) technique.**TABLE 6** Tumor control probability results in nominal PB-PB, PB-MC, and MC-MC plans of breast cancer. The results are averaged over ten breast cancer patients.

		PB-PBAvg. (\pm SD)	PB-MCAvg. (\pm SD)	P-value	MC-MCAvg. (\pm SD)	P-value
CTV_Total	TCP (%)	93.2 (\pm 0.6)	91.4 (\pm 0.7)	<0.001	92.9 (\pm 0.6)	0.345
CTV_CW/Breast	TCP (%)	92.9 (\pm 0.9)	91.2 (\pm 1.2)	0.002	92.7 (\pm 0.7)	0.535
CTV_AxI	TCP (%)	93.4 (\pm 0.5)	91.5 (\pm 1.0)	<0.001	93.2 (\pm 0.6)	0.545
CTV_AxII	TCP (%)	93.3 (\pm 0.7)	91.8 (\pm 0.9)	0.002	93.1 (\pm 0.7)	0.571
CTV_AxIII	TCP (%)	93.4 (\pm 0.5)	91.7 (\pm 0.9)	0.001	93.1 (\pm 0.6)	0.326
CTV_IMN	TCP (%)	93.0 (\pm 1.3)	90.2 (\pm 2.1)	0.003	92.9 (\pm 1.4)	0.940
CTV_SCVN	TCP (%)	93.4 (\pm 0.7)	90.5 (\pm 1.2)	<0.001	93.1 (\pm 0.7)	0.385

PB-PB, PB optimization followed by PB dose calculation; PB-MC, PB optimization followed by MC dose calculation; MC-MC, MC optimization followed by MC dose calculation; CW, chestwall; EUD, equivalent uniform dose; IMN, internal mammary nodes; SCVN, supraclavicular nodes; TCP, tumor control probability.

^aP-value for PB-MC vs. PB-PB.

^bP-value for MC-MC vs. PB-PB.

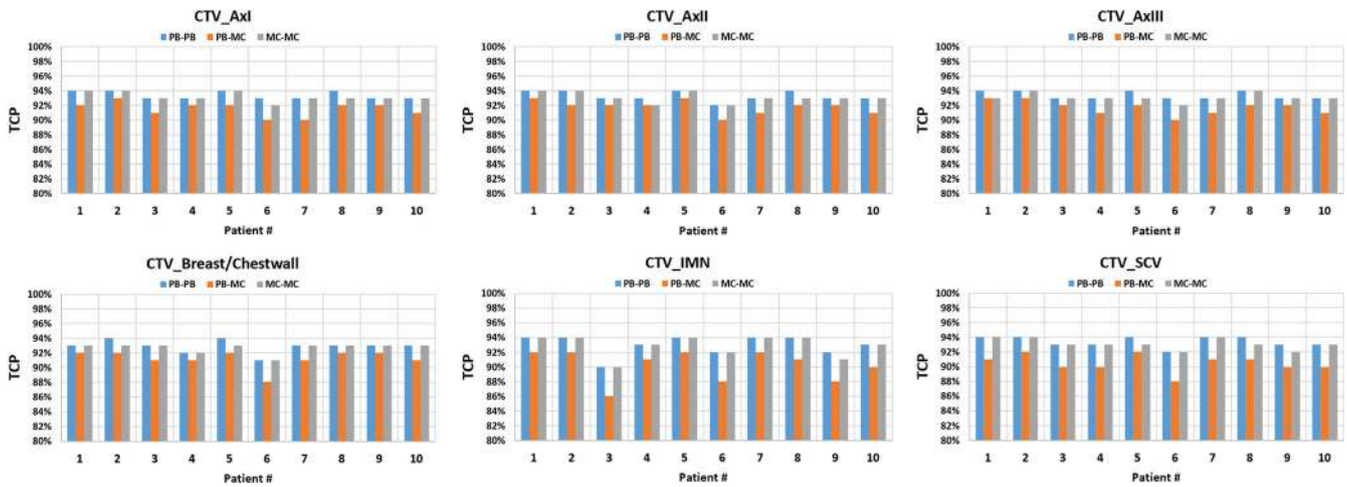
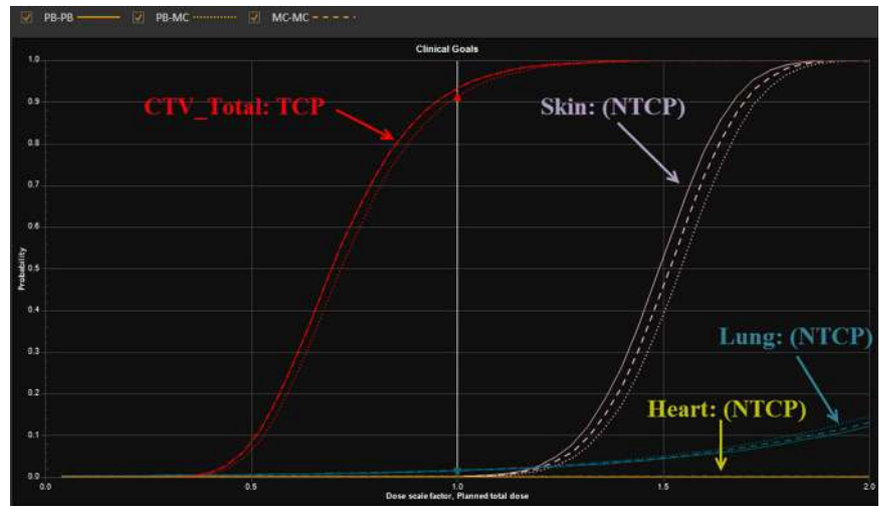


FIG. 5. Tumor control probability of the total clinical target volumes for breast cancer patients (n = 10) in PB optimization followed by PB dose calculation, PB optimization followed by MC dose calculation, and MC optimization followed by MC dose calculation plans generated by intensity-modulated proton therapy technique.

FIG. 6. Tumor control probability clinical target volume (CTV_Total) and normal tissue complication probability (heart, skin, and left lung) in PB optimization followed by PB dose calculation, PB optimization followed by MC dose calculation, and MC optimization followed by MC dose calculation plans of an example patient.



CTV_AxI (93.4% vs. 93.2%; $P = 0.545$), CTV_AxII (93.3% vs. 93.1%; $P = 0.571$), CTV_AxIII (93.4% vs. 93.1%; $P = 0.326$), CTV_CW/breast (92.9% vs. 92.7%; $P = 0.535$), CTV_IMN (93.0% vs. 92.9%; $P = 0.940$), CTV_SCVN (93.4% vs. 93.1%; $P = 0.385$), and CTV_Total (93.2% vs. 92.9%; $P = 0.345$). Figure 6 shows the TCP of CTV_Total and in PB-PB, PB-MC, and MC-MC plans of an example patient.

3.E | NTCP analysis

Table 7 shows the NTCP results for the heart, ipsilateral lung, and skin. There was no clear distinction among PB-PB, PB-MC, and MC-MC plans in terms of NTCP results. Based on the LKB model and published radiobiological parameters used in this study, the NTCPs were 0% for the heart, $\leq 0.2\%$ for the skin, and 0.4% to 1.9% for the ipsilateral (left) lung. Figure 6 shows the NTCP of the heart, left lung, and skin in PB-PB, PB-MC, and MC-MC plans of an example patient.

3.F | Patient-specific quality assurance (QA) analysis

Patient-specific QA measurement was done for PB-PB plans of all ten patients in a water tank using DigiPhant-PT (IBA Dosimetry, Schwarzenbruck, Germany) and MatriXX-PT (IBA Dosimetry, Schwarzenbruck, Germany). A 2D gamma analysis was performed between the calculated and measured 2D dose distributions using patient-specific QA module implemented within myQA software platform (IBA Dosimetry, Schwarzenbruck, Germany). For 2D gamma evaluation, we utilized 3% and 3 mm criteria and low-dose threshold of 10%. A gamma passing rate of $\geq 90\%$ was considered to be an acceptable level. Table 8 shows the gamma evaluation results of all ten patients. The average 2D gamma was $94.0\% \pm 2.9\%$ with a minimum of 90.1% and a maximum of 98.9%.

TABLE 7 NTCP of heart, lung, and skin in nominal PB-PB, PB-MC, and MC-MC plans of breast cancer.

Patient #	Heart			Skin			Ipsilateral Lung		
	NTCP (%)			NTCP (%)			NTCP (%)		
	PB-PB	PB-MC	MC-MC	PB-PB	PB-MC	MC-MC	PB-PB	PB-MC	MC-MC
1	0.0	0.0	0.0	0.0	0.0	0.0	0.9	1.0	1.0
2	0.0	0.0	0.0	0.0	0.0	0.0	0.7	0.8	0.8
3	0.0	0.0	0.0	0.1	0.0	0.1	0.8	0.9	0.9
4	0.0	0.0	0.0	0.0	0.0	0.0	0.6	0.7	0.6
5	0.0	0.0	0.0	0.0	0.0	0.0	0.4	0.4	0.5
6	0.0	0.0	0.0	0.0	0.0	0.0	1.7	1.9	1.8
7	0.0	0.0	0.0	0.0	0.0	0.0	0.9	1.0	0.9
8	0.0	0.0	0.0	0.0	0.0	0.0	0.7	0.8	0.8
9	0.0	0.0	0.0	0.2	0.1	0.1	0.9	1.0	0.8
10	0.0	0.0	0.0	0.1	0.1	0.1	1.5	1.6	1.6
Average	0.0	0.0	0.0	0.0	0.0	0.0	0.9	1.0	1.0
SD	0.0	0.0	0.0	0.1	0.0	0.0	0.4	0.4	0.4
P-value		0.968 ^a	0.968 ^b		0.674 ^a	0.936 ^b		0.384 ^a	0.674 ^b

PB-PB, PB optimization followed by PB dose calculation; PB-MC, PB optimization followed by MC dose calculation; MC-MC, MC optimization followed by MC dose calculation; NTCP, normal tissue complication probability.

^aP-value for PB-MC vs. PB-PB.

^bP-value for MC-MC vs. PB-PB.

TABLE 8 2D gamma evaluation results from patient-specific QA. Calculated (PB-PB plans) and measured 2D dose distributions were compared using patient-specific QA module implemented within the myQA software platform.

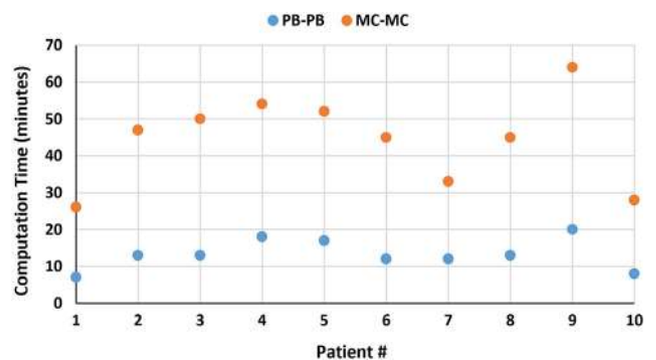
Patient #	Gamma passing rate (%)	
	Field 1	Field 2
1	95.4	92.9
2	90.6	98.9
3	93.5	98.3
4	96.0	
5	94.2	98.4
6	91.1	90.1
7	92.5	90.3
8	94.9	
9	94.2	
10	92.8	

PB-PB, PB optimization followed by PB dose calculation; QA, quality assurance.

Gamma evaluation criteria = 3% and 3 mm; Low-dose threshold = 10%; Accepted gamma passing rate $\geq 90\%$.

4 | DISCUSSION

Previous studies on RayStation proton dose calculation algorithms were mostly focused on the dosimetric impact of algorithms involving either phantom³⁻⁷ or disease sites.^{5,7,8} To our best knowledge, at the time of writing this paper, there is currently no literature that has addressed the radiobiological impact of RayStation PB and MC in IMPT breast treatment plans. It is essential to investigate how

**FIG. 7.** Computation time in minutes for intensity-modulated proton therapy breast plans (PB optimization followed by PB dose calculation and MC optimization followed by MC dose calculation) of ten breast cancer patients.

dosimetric accuracy of dose calculation algorithms can be translated to the radiobiological differences in clinical patient cases. Hence, the current study was undertaken to demonstrate the radiobiological impact of RayStation PB and MC in terms of EUD, TCP, and NTCP in IMPT breast cancer treatment plans.

For breast cancer treatment, a tumor volume is often situated at a shallower depth. This necessitates the use of a proton beam with a smaller range. However, our proton system has a minimum of 4.0 cm range in water. Hence, for the treatment of shallower target such as in the case of CW/breast, a range shifter is typically used to reduce the energy of the proton beam. Although the use of range shifter allows to achieve full dose modulation of the tumor volume that may be extended close to the skin, an accurate modeling of algorithms accounting for angular distribution of a pencil proton

beam after traversing the range shifter and translation of angular distribution into a geometric spread of proton beam's cross-section at the detector/patient surfaces is critical.^{9,21}

It has been reported that recalculation of PB plans using MC will result in decrease in target dose and coverage. In a breast study by Tommasino et al.,⁷ the MC recomputed average dose to the planning target volume (PTV) was 7.1% lower than the prescription dose of 50 Gy. Liang et al.⁸ reported the reduction of CTV D_{mean} by 2.1% of the prescription dose in MC recomputed plans compared with PB plans. In the current study, we observed the reduction of the CTV_Total D_{mean} in PB-MC plans by an average difference of 2.7% (range, 1.6–3.5%) when compared to PB-PB plans. Liang et al.⁸ also reported the reduction of CTV $D_{99\%}$ and CTV $D_{95\%}$ by 3.7% and 3.4%, respectively, when PB plans are recalculated with MC. The findings from the current study agree with that of Liang et al.⁸ such that we noticed the reduction of the CTV_Total $D_{99\%}$ and $D_{95\%}$ in PB-MC plans by an average difference of 5.3% and 4.1%, respectively, when compared to PB-PB plans.

RayStation has made MC available for plan optimization. MC-optimized plans offered optimal CTV coverage and dose distribution similar to the ones in the PB plan. The current study agrees with the results from the Liang et al.⁸ such that CTV_Total dose between MC-MC and PB-PB plans was found to be minimal (<0.5% for both $D_{95\%}$ and D_{mean} ; 1.4% for $D_{99\%}$). The differences between Liang et al.⁸ and our study may be attributed to the planning techniques. For instance, Liang et al.⁸ normalized PB optimized plans and MC optimized plans such that 95% of the PTV was covered by 95% of the prescription dose, whereas plan normalization technique was not utilized in our study.

If MC is more accurate than PB for IMPT treatment of shallower targets requiring range shifter as reported in the literature,^{3–7} dosimetric results comparing PB-PB vs. PB-MC plans in the current study suggest that PB overestimates the target dose and coverage. Based on the radiobiological results in the current study, PB slightly overestimated the TCP for all CTV structures by 1–3% for Axl, 1–2% for AxII, 1–3% for AxIII, 1–3% for CW/breast, 2–4% for IMN, 2–4% for SCVN, and 1–2% for CTV_Total. Similarly, PB overestimated the EUD of target volumes by 1.8–3.2 Gy(RBE). However, the choice of optimization and dose calculation algorithms did not produce any noticeable differences in the NTCP of the heart, lung, and skin. It has been reported that breast cancer patients treated with proton therapy could have the risk of acute skin toxicities.^{22,23} The NTCP of skin for the clinical endpoint of severe acute toxicity in our study was $\leq 0.2\%$ for all ten patients. This was calculated using the LKB model and radiobiological parameters predicted by Pastore et al.²⁰ It must be noted that radiobiological evaluation in our study was carried out based on radiobiological parameters that are derived from the conventional mega-voltage X-ray (photon) therapy. This is a limitation of our study. As more breast cancer patients are being treated using proton therapy and enrolled in clinical trials, there is a need for proton derived NTCP models correlating to the tissue toxicities of breast cancer patients. Due to lack of proton derived radiobiological parameters, researchers continue to use photon-derived NTCP

models for proton therapy.^{12,14,25} Recently, Blanchard et al.²⁴ validated photon-derived NTCP models that can be used to select head and neck patients for proton treatment.

The current study assumed constant RBE value of 1.1. Several publications^{26,27} have demonstrated the existence of variable RBE for proton therapy and depend on the cell type, endpoint, LET, radiation dose, etc. The variability in RBE could lead to different α/β values, thus impacting EUD, TCP, and NTCP.²⁸ In this study, we did not explore the impact of variable RBE on IMPT breast plans. Our future work will investigate how the combination of variable RBE and proton dose calculation algorithm can affect the radiobiological results.

One of the challenges associated with MC plan optimization is the treatment planning efficiency. Figure 7 illustrates the computation time in minutes for PB-PB and MC-MC plans of all ten patients. For PB-PB plans, the average computation time was 13.3 ± 4.1 min (range, 7–20 min), whereas the average computation time for MC-MC plans was 44.4 ± 12.1 min (range, 26–64 min). Overall, PB-PB plans had higher computation efficiency, with an average factor of 3.4 when compared to MC-MC plans. It must be noted that IMPT plan optimization time is dependent on several factors such as computing hardware and software resources, robustness scenarios, number of optimization structures and their constraints, and optimization settings (number of iterations and sampling history – number of ions/spot).

5 | CONCLUSION

If RayStation MC is more accurate than PB as reported in the literature, dosimetric and radiobiological results from the current study suggest that PB overestimates the target dose, EUD, and TCP for IMPT breast cancer treatment. The overestimation of dosimetric and radiobiological results of the target volume by PB needs to be further interpreted in terms of clinical outcome. The use of RayStation MC for both plan optimization and dose calculation of IMPT breast cancer plans can provide optimal target coverage and radiobiological results (EUD and TCP for target volumes) with better accuracy.

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CONFLICT OF INTEREST

No conflicts of interest.

REFERENCES

1. Mast ME, Vredevelde EJ, Credoe HM, et al. Whole breast proton irradiation for maximal reduction of heart dose in breast cancer patients. *Breast Cancer Res Treat.* 2014;148:33–9.

2. Taylor CW, Wang Z, Macaulay E, et al. Exposure of the heart in breast cancer radiation therapy: a systematic review of heart doses published during 2003 to 2013. *Int J Radiat Oncol Biol Phys.* 2015;93:845–53.
3. Saini J, Maes D, Egan A, et al. Dosimetric evaluation of a commercial proton spot scanning Monte-Carlo dose algorithm: comparisons against measurements and simulations. *Phys Med Biol.* 2017;62:7659–7681.
4. Taylor PA, Kry SF, Followill DS. Pencil beam algorithms are unsuitable for proton dose calculations in lung. *Int J Radiat Oncol Biol Phys.* 2017;99:750–756.
5. Maes D, Saini J, Zeng J, Rengan R, Wong T, Bowen SR. Advanced proton beam dosimetry part II: Monte Carlo vs. pencil beam-based planning for lung cancer. *Transl Lung Cancer Res.* 2018;7:114–121.
6. Shirey RJ, Wu HT. Quantifying the effect of air gap, depth, and range shifter thickness on TPS dosimetric accuracy in superficial PBS proton therapy. *J Appl Clin Med Phys.* 2018;19:164–173.
7. Tommasino F, Fellin F, Lorentini S, Farace P. Impact of dose engine algorithm in pencil beam scanning proton therapy for breast cancer. *Phys Med.* 2018;50:7–12.
8. Liang X, Li Z, Zheng D, Bradley JA, Rutenberg M, Mendenhall N. A comprehensive dosimetric study of Monte Carlo and pencil-beam algorithms on intensity-modulated proton therapy for breast cancer. *J Appl Clin Med Phys.* 2019;20:128–136.
9. Titt U, Mirkovic D, Sawakuchi GO, et al. Adjustment of the lateral and longitudinal size of scanned proton beam spots using a pre-absorber to optimize penumbræ and delivery efficiency. *Phys Med Biol.* 2010;55:7097–106.
10. Rana S, Bennouna J, Samuel EJJ, Gutierrez AN. Development and long-term stability of a comprehensive daily QA program for a modern pencil beam scanning (PBS) proton therapy delivery system. *J Appl Clin Med Phys.* 2019;20:29–44.
11. Niemierko A. Reporting and analyzing dose distributions: a concept of equivalent uniform dose. *Med Phys.* 1997;24:103–10.
12. Rana S, Cheng C, Zheng Y, et al. Dosimetric study of uniform scanning proton therapy planning for prostate cancer patients with a metal hip prosthesis, and comparison with volumetric-modulated arc therapy. *J Appl Clin Med Phys.* 2014;15:4611.
13. RaySearch Laboratories. *RayStation 6 Reference Manual.* Stockholm, Sweden: RaySearch Laboratories; 2017.
14. Okunieff P, Morgan D, Niemierko A, et al. Radiation dose-response of human tumors. *Int J Radiat Oncol Biol Phys.* 1995;32:1227–37.
15. Horton JK, Halle JS, Chang SX, et al. Comparison of three concomitant boost techniques for early-stage breast cancer. *Int J Radiat Oncol Biol Phys.* 2006;64:168–75.
16. Owen JR, Ashton A, Bliss JM, et al. Effect of radiotherapy fraction size on tumour control in patients with early-stage breast cancer after local tumour excision: long-term results of a randomised trial. *Lancet Oncol.* 2006;7:467–71.
17. Luxton G, Keall PJ, King CR. A new formula for normal tissue complication probability (NTCP) as a function of equivalent uniform dose (EUD). *Phys Med Biol.* 2008;53:23–36.
18. Oinam AS, Singh L, Shukla A, Ghoshal S, Kapoor R, Sharma SC. Dose volume histogram analysis and comparison of different radiobiological models using in-house developed software. *J Med Phys.* 2011;36:220–9.
19. Semenenko VA, Li XA. Lyman-Kutcher-Burman NTCP model parameters for radiation pneumonitis and xerostomia based on combined analysis of published clinical data. *Phys Med Biol.* 2008;53:737–755.
20. Pastore F, Conson M, D'Avino V, et al. Dose-surface analysis for prediction of severe acute radio-induced skin toxicity in breast cancer patients. *Acta Oncol.* 2016;55:466–73.
21. Gottschalk B, Koehler AM, Schneider RJ, Sisterson JM, Wagner MS. Multiple Coulomb scattering of 160 MeV protons. *Nucl Instrum Methods Phys Res B.* 1993;74:467–490.
22. Liang X, Bradley JA, Zheng D, et al. Prognostic factors of radiation dermatitis following passive-scattering proton therapy for breast cancer. *Radiat Oncol (London, England).* 2018;13:72.
23. Verma V, Shah C, Mehta MP. Clinical outcomes and toxicity of proton radiotherapy for breast cancer. *Clin Breast Cancer.* 2016;16:145–154.
24. Blanchard P, Wong AJ, Gunn GB, et al. Toward a model-based patient selection strategy for proton therapy: external validation of photon-derived normal tissue complication probability models in a head and neck proton therapy cohort. *Radiother Oncol.* 2016;121:381–386.
25. Hall DC, Trofimov AV, Winey BA, Liebsch NJ, Paganetti H. Predicting patient-specific dosimetric benefits of proton therapy for skull-base tumors using a geometric knowledge-based method. *Int J Radiat Oncol Biol Phys.* 2017;97:1087–1094.
26. Paganetti H. Relative biological effectiveness (RBE) values for proton beam therapy. Variations as a function of biological endpoint, dose, and linear energy transfer. *Phys Med Biol.* 2014;59:R419–472.
27. Lühr A, von Neubeck C, Krause M, Troost EGC. Relative biological effectiveness in proton beam therapy – Current knowledge and future challenges. *Clin Transl Radiat Oncol.* 2018;1:35–41.
28. Paganetti H. Relating the proton relative biological effectiveness to tumor control and normal tissue complication probabilities assuming interpatient variability in alpha/beta. *Acta Oncol.* 2017;56:1379–1386.