Dosimetric and radiobiological comparison in different dose calculation grid sizes between Acuros XB and anisotropic analytical algorithm for prostate VMAT

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Abstract

To investigate feasible treatment planning parameters, we aimed to evaluate the dosimetric and radiobiologic impact of the dose calculation algorithm and grid size in the volumetric modulated arc therapy (VMAT) plan for prostate cancer. Twenty patients were selected, and the treatment plans were initially generated with anisotropic analytical algorithm (AAA) and recalculated with Acuros XB (AXB) algorithm. Various dose grids were used for AXB (1, 2, and 3 mm) and AAA (1, 3, and 5 mm) plan. Dosimetric parameters such as homogeneity index (HI) and conformity index (CI), and radiobiological parameters such as tumor control probability (TCP) and normal tissue complication probability (NTCP) were calculated. Significant differences were observed in the planning target volume (PTV) coverage between both algorithms, and the V₉₅% of AAA were significantly affected by grid (p < 0.01). On 1 mm grid, the mean rectal dose difference between both algorithms was 2.87% of the prescription dose (p < 0.01), which was the highest among the critical organs. The TCP and NTCP of the AAA were higher than those of AXB (p < 0.01). Compared to AXB with 1 mm grid, the 2 mm grid showed comparable dose calculation accuracy with short calculation time. This study found that the PTV and rectum show significant differences according to dose calculation algorithm and grid. Considering the dose calculation performance for heterogeneous area, we recommend AXB with 2 mm grid for improving treatment efficiency of prostate VMAT.

Introduction

Clinical use of volumetric-modulated arc therapy (VMAT) [1] has grown extremely since its debut in 2008. The application of VMAT for prostate cancer has been well-demonstrated for both plan quality and efficiency [2, 3]. A well-known advantage of VMAT is its ability to
deliver faster treatment, which makes treatments patient-friendly and improves treatment accuracy because intrafractional motion is reduced. A previous study reported that VMAT produces comparable and higher conformal dose distribution and accurate dose delivery in comparison to intensity modulated radiotherapy (IMRT) [4–6].

In order to obtain effective treatment outcomes, minimal toxicity and complications should be achieved through reducing the radiation dose to organs at risks (OARs) near the prostate, such as the bladder, the femur, and the rectum. The endorectal balloon (ERB) for prostate radiotherapy can be implemented for sparing the rectal wall and reducing the intrafraction motion [7–10]. However, its use might introduce the heterogeneous region due to the generation of an air cavity in rectum, which will compromise the treatment efficacy, and therefore, accurate dose calculation is extremely important.

The Anisotropic Analytical Algorithm (AAA; Varian Medical Systems, Palo Alto, CA, USA) [11] is a convolution/superposition model for photon dose calculation. The AAA has improved accuracy of calculation compared to previous algorithms and is widely used at various treatment sites. The AAA calculates a dose accounting for lateral electron transport and, performs simplified density scaling of the kernels calculated in water. The AAA uses a kernel of water instead of a kernel with medium-specific characteristics, therefore it has limitation on accuracy of dose calculation. The Acuros XB (AXB; Varian Medical Systems, Palo Alto, CA, USA) [12], advanced dose calculation algorithm has been implemented for clinical use. The AXB uses the multiple-source model derived for the AAA, and performs dose calculation by explicitly solving the linear Boltzmann transport equation (LBTE). A mass density as well as material type was reflected to each given voxel in the AXB dose calculation. In other words, the AXB can reflect the radiation interaction according to each material property, and have been reported more accurate dose prediction in the heterogeneous region than the AAA in previous studies. Koo et al. [13] evaluated the dosimetric effect of AXB and AAA algorithms for prostate cancer and reported that dose calculation in the air cavity with AXB was more accurate than that of AAA. Kroon et al. [14] also investigated the performances of the dose calculation algorithm in the air cavity of non-small cell lung cancer (NSCLC). They have shown that the AXB with 2.5 mm calculated dose accurately, compared to the AAA. Kan et al. [15] analyzed dosimetric impacts of AXB and AAA on intensity modulated stereotactic radiotherapy (IMSRT) plans for nasopharyngeal carcinoma (NPC), and recommended the AXB as a standard reference for IMSRT boost for NPC. In most of studies, AXB showed generally low target coverage and OAR dose. In addition, previous studies have reported that dose distribution with AXB is more accurate and consistent with the Monte Carlo simulation in the heterogeneous region than that of AAA [16–18].

Since AAA uses the discretized kernel in lateral directions [19] and AXB also discretizes solution variables (space, angle, and energy) with material composition of voxels from the CT Hounsfield unit, both dose calculation algorithms are affected by the dose grid size. Although the error due to discretization may be smaller than the error obtained by the algorithm, the influence may be different depending on the algorithm and also be significant in some clinical cases. In general, the calculated dose distribution at fine grid is considered more accurately, however it is time consuming and may not feasible in the clinic. It is important to optimize the dose calculation algorithm and grid size at a reasonable calculation time without significantly compromising accuracy.

In other words, the calculated dose distributions of treatment planning system (TPS) are affected by the dose grid size, and the presence of a dosimetric influence according to the calculated grid size has been reported in IMRT and VMAT [20, 21]. Huang et al. [22] assessed the potential impact of grid size on dose calculation using AXB and AAA algorithms for stereotactic body radiotherapy for NSCLC, and showed that the dose difference between the two algorithms
using a 2.5 mm grid was greater than that of a 1 mm grid on low density planning target volume (PTV). Previous studies have reported that the dose difference can be significant for radiobiological factors such as normal tissue complication probability (NTCP) and tumor control probability (TCP) [20, 23], which may affect radiotherapy quality and treatment success. Shiv P. Srivastava et al. [20] analyzed the radiobiological effects and dosimetric impacts of plans, according to calculation grid sizes, in head and neck cancers. They showed that a smaller calculation grid provides superior dosimetric outcomes with improved TCP and reduced NTCP.

In our previous study, we've confirmed the AXB and AAA performance in a rectangular acryl phantom with an air cavity by EBT3 film dosimetry [13]. As a result, we confirmed that AXB had more similar results than AAA to film measurements in the air cavity and air-material interface in phantom. Unlike the phantom, patients have various anatomies, and the treatment plan for patient is more complicate than the phantom study. The influence of dose calculation grids and algorithms were not clearly identified in some clinical cases, and it is important for efficient patient care. No previous study has evaluated the impact on prostate VMAT plans by different dosimetric and radiological effects (TCP, NTCP) obtained by different algorithms under different grid sizes. In this study, we investigate the dosimetric and radiological impact on prostate VMAT plans with ERB from the dose calculation algorithms with different grid sizes.

Materials and methods

Patient selection and treatment planning

All experimental methods of this study were performed in accordance with the relevant guidelines and regulations. The Institutional Review Board of the Seoul National University Bundang Hospital approved the data collection and analysis (B-1802/451-106), and informed consent of the participants was waived by the IRB. All patient data used in the study were anonymized. Twenty patients who received prostate VMAT in our institution from April 2016 to April 2017 were randomly selected retrospectively. The average age of the patients was 64 (58–84) and the average weight was 73.1 kg (49.4–82.5 kg).

Prior to the planning computed tomography (CT) simulation, all patients were asked to drink 300 ml of water before the start of the one-hour of simulation, to ensure that their bladders were completely filled. An EBR was inserted into the rectum and inflated with approximately 70 cm³ with air. After 1 minute, the EBR was pulled toward the patient’s anal sphincter to the pre-marked position on the EBR catheter [24]. CT was performed using a Philips Big Bore CT scanner (Philips Medical Systems, Amsterdam, Netherlands).

The clinical target volume (CTV) on the planning CT was delineated from the prostate volume using magnetic resonance imaging. CTV included gross tumor and subclinical microscopic disease. The PTV was created from the CTV by expanding 5 mm posteriorly and 7 mm elsewhere. The rectum, bladder, left femoral head, and right femoral head were delineated as the OARs. Varian couch is modelled in our TPS and was inserted into each treatment plan (used for dose calculation).

The prostate VMAT plans were created using Eclipse TPS (version 11.0.34, Varian Medical System, Palo Alto, CA, USA) under the same dose prescription (78 Gy/39 fractions) and dose–volume criteria (Table 1). The aim of the planning optimization was to cover at least 95% of the PTV with 95% of the prescription dose. Two full arc techniques using a 10 MV beam from a Varian TrueBeam STX linear accelerator were used for optimal target coverage.

All prostate VMAT plans were optimized initially with the AAA algorithm and recalculated with the AXB algorithm. In our TPS, the grid size can be up to 3 mm for AXB, and up to 5 mm for AAA. In order to investigate the trend of dosimetric and radiobiological parameters as the grid increases, a grid size of 1, 3, and 5 mm for AAA and 1, 2, and 3 mm for AXB were adopted.
Dosimetric and radiobiological parameters

In order to evaluate the dosimetric and radiobiological parameters, cumulative dose-volume histograms (DVHs) were calculated for each plan. Dosimetric parameters such as median, mean, maximum, minimum dose and \( V_{95\%} \) (percent volume irradiated by 95% of the prescription dose) for PTV were analyzed. \( V_{95\%} \) of PTV was used as a measure of the target coverage in this study. To evaluate the target dose of each VMAT plan, homogeneity index (HI), conformity index (CI), and conformation number (CN) were calculated for PTV. HI was calculated by Eq (1):

\[
HI = \frac{D_{2} - D_{98}}{D_{50}}
\]  

Where \( D_{2}, D_{98}, \) and \( D_{50} \) represent the dose to 2%, 98%, and 50% volume for the PTV, respectively. A lower HI means that the plan has a more homogeneous target dose. CI was calculated by Eq (2):

\[
CI = \frac{V_{RI}}{TV}
\]

Where \( V_{RI} \) is the volume of reference isodose on body, and TV is the physical volume of PTV. The CI refers to the degree of isodose conformity, and it is ideal for the CI to remain close to 1.

To assess conformity to target dose and the healthy tissue irradiation, CN was evaluated by Eq (3):

\[
CN = \frac{TV_{RI}}{TV} \times \frac{TV_{RI}}{V_{RI}}
\]

Where \( TV_{RI} \) represents the PTV volume covered with the reference isodose. The first term of

### Table 1. Dose volume constraints for prostate volumetric modulated arc therapy plans.

| Structure    | Constraints                  |
|--------------|------------------------------|
| Rectum       | \( V_{30\%} < 7000 \text{ cGy} \) |
|              | \( V_{50\%} < 5430 \text{ cGy} \) |
| Bladder      | \( V_{30\%} < 7000 \text{ cGy} \) |
|              | \( V_{50\%} < 5430 \text{ cGy} \) |
| Femoral heads| \( V_{5\%} < 5430 \text{ cGy} \) |
| GTV          | \( V_{99\%} > 7800 \text{ cGy} \) |
| PTV          | \( V_{9\%} < 8190 \text{ cGy} \) |
|              | \( V_{2\%} < 8100 \text{ cGy} \) |
|              | \( V_{97\%} > 7650 \text{ cGy} \) |
|              | \( V_{99\%} > 7410 \text{ cGy} \) |

### Table 2. Parameters used to calculate tumor control probability (TCP) and normal tissue complication probability (NTCP).

| Type          | Organ       | \( a \) | \( \gamma_{50} \) | TCP50  | TD50  | \( a/\beta \) |
|---------------|-------------|--------|----------------|--------|-------|--------------|
| Tumor         | Prostate    | -13    | 2.2            | 67.5   | 1.5   |
| Critical Organ| Rectum      | 8.33   | 2.66           | 80     | 5.4   |
|               | Bladder     | 2      | 3.63           | 80     | 7.5   |
|               | Lt femur head| 13     | 2.7            | 65     | 3     |
|               | Rt femur head| 13     | 2.7            | 65     | 3     |

https://doi.org/10.1371/journal.pone.0207232.t001

https://doi.org/10.1371/journal.pone.0207232.t002
CN refers to the target coverage, and the second terms indicate the degree of delivered dose on normal tissue.

For OARs, dosimetric parameter included the median, maximum, and minimum dose and a set of $V_{x\%}$, which is the volume of the organ receiving $x\%$ or more of the prescription dose.

In order to investigate the radiobiological impact on the PTV and various OARs, the values of TCP and NTCP were calculated from the DVH of planning data using a different dose calculation grid size and algorithm [25]. Equivalent uniform dose (EUD) is defined as the dose that when distributed uniformly over a structure would produce the same effect as the dose specified by the DVH. EUDs were calculated using Niemierko’s phenomenological model [26] by Eq (4):

$$EUD = \left(\sum_{i=1}^{n} (v_i D_i^a)\right)^{\frac{1}{a}}$$  \hspace{1cm} (4)

The EUD model can be used in both PTV and normal tissue by applying different input parameters. The $a$ is a unitless parameter derived specifically from normal tissue or tumor properties. The $v_i$ represents the relative sub-volume of the $i$-th that received a dose of $D_i$ in Gy units. Therefore, the sum of all $v_i$ is equal to 1 in the above EUD formula. Differential DVHs were obtained from a given VMAT plan to obtain the $D_i$ and $v_i$ at each structure. NTCP and TCP are expressed by Eqs (5) and (6):

$$NTCP = \frac{1}{1 + \left(\frac{TD_{50}}{EUD}\right)^{\frac{1}{a}}}$$ \hspace{1cm} (5)

$$TCP = \frac{1}{1 + \left(\frac{TCD_{50}}{EUD}\right)^{\frac{1}{a}}}$$ \hspace{1cm} (6)

The $TD_{50}$ is the tolerance dose for 50% complication probability within a specific time interval. The $TCD_{50}$ is the tumor dose to control 50% of the tumor when irradiated homogeneously, and $\gamma_{50}$ is a unitless parameter derived from the slope of the dose-response curve that is specific to the organ or tumor. Table 2 lists the input parameters for calculating TCP and NTCP, and these parameters were referenced to other studies [23, 25].

In order to evaluate the feasibility of grid sizes and algorithms in clinical practice, we calculated the deviations of the mean, median, maximum, and minimum dose to the PTV and OARs. The deviations were compared between the VMAT plans and evaluated with the dose calculation times. The deviations are expressed by Eq (7):

$$Deviation = \left|\frac{Ref - Eval}{Ref}\right| \times 100$$ \hspace{1cm} (7)

Where, $Ref$ represents the reference dosimetric parameter and $Eval$ refers to the dosimetric parameters of evaluated plans. The parameters with the AXB and a 1 mm grid size (AXB1) were selected as $Ref$, which is the combination for the most accurate algorithm and dose grid in this study. AXB have been reported as superior dose calculation algorithm in heterogeneous region and the general consensus is that finer dose calculation grid would produce more accurate dose. $Eval$ were rest of plans such as AAA1, AAA3, AXB2, et cetera.

To clearly show the dose difference according to the algorithms, the AXB1 and AAA1 plans of one patient were imported into the Computational Environment for Radiotherapy Research (CERR version 4.6) [27], and we subtracted the dose distribution of AAA1 from that of AXB1. CERR is a programming package of MATLAB (MathWorks, Inc., Natick, USA), and has many functions for radiotherapy research such as the CT slice viewer, contouring tool, DVH.
calculation, dose distribution subtraction tool, et cetera. In addition, we subtracted the distribution of the 3 mm grid from that of the 1 mm grid for both algorithms to evaluate the grid effect on the VMAT plan.

**Statistical analysis and comparison of dose calculation time**

Statistical analysis were performed using SPSS Statistics 19 (IBM SPSS, Chicago, IL) to assess statistical significance between algorithm type and grid size. The Wilcoxon signed rank test [28] was used in this study, and differences were considered statistically significant at p-values < 0.05. The calculation time of all AAA and AXB plans were also recorded to investigate the feasible grid sizes and algorithms in prostate VMAT.

**Results**

**Dosimetric comparison**

Table 3 shows the mean and standard deviation of dosimetric parameters to PTV on plans using different dose calculation grid sizes and algorithms. For PTV, the $V_{95\%}$ with the AXB1 was 96.65%, whereas the $V_{95\%}$ was 99.03% with AAA1. The difference in average $V_{95\%}$ between AAA1 and AAA3 mm was 1.22%, and that between AXB1 and AXB3 was 0.29%. The $V_{95\%}$ for AXB2 was 96.60%, and the $V_{95\%}$ difference between AXB1 and AXB2 was small (only 0.05%). The median dose slightly decreased over the change from AAA1 to AAA5. Contrary to the AAA trends, the median dose increased by 0.66 Gy (0.84% of prescription) over the variation from AXB1 to AXB3. A similar trend was observed for the mean and maximum dose of PTV over the transition from AAA1 to AAA5, and the mean dose decreased by 1.25 Gy accordingly. The CI value of AXB3 was 1.03, the closest to one in all cases. The HI of AAA5 was 0.14, which was the highest among all cases. Fig 1 shows the average DVH of PTV and different OARs on prostate VMAT plans using various calculation grid sizes and algorithms.

For different OARs, the average dosimetric parameters are listed in Table 4, and the $V_{X\%}$ are shown in Fig 2. As shown in Table 4, the median and mean dose of rectum was higher in AAA1 than in AXB1, and the differences were 2.07% and 2.87% of prescription dose by simply subtracting those values in Table 4. Using a 1 mm dose grid, the difference of $V_{X\%}$ of rectum dose between the two algorithms was high in the region ranging from $V_{40\%}$ to $V_{90\%}$ (2.63% – 5.78%).

| V95 (%) | AAA – Mean(%) | AXB – Mean(%) |
|---------|---------------|---------------|
| 1 mm    | 99.03 (0.43)  | 96.65 (0.73)  |
| 3 mm    | 97.81 (0.42)  | 96.60 (0.72)  |
| 5 mm    | 91.28 (1.15)  | 96.94 (0.71)  |

| Dmedian (Gy) | AAA – Mean(%) | AXB – Mean(%) |
|--------------|---------------|---------------|
| 80.25 (0.20) | 79.54 (0.19)  | 79.53 (0.18)  |
| 80.16 (0.20) | 79.51 (0.19)  | 79.53 (0.18)  |
| 79.66 (0.27) | 79.54 (0.19)  | 79.54 (0.18)  |

| Dmean (Gy) | AAA – Mean(%) | AXB – Mean(%) |
|------------|---------------|---------------|
| 79.85 (0.18) | 79.10 (0.20) | 79.01 (0.18) |
| 79.51 (0.18) | 78.60 (0.23) | 79.10 (0.20) |
| 78.60 (0.23) | 79.54 (0.19) | 79.53 (0.22) |

| Dmax (Gy) | AAA – Mean(%) | AXB – Mean(%) |
|-----------|---------------|---------------|
| 84.42 (0.70) | 84.98 (0.76) | 84.24 (0.73) |
| 83.30 (0.39) | 82.70 (0.53) | 84.45 (0.77) |
| 82.70 (0.53) | 84.98 (0.76) | 84.24 (0.73) |

| Dmin (Gy) | AAA – Mean(%) | AXB – Mean(%) |
|-----------|---------------|---------------|
| 64.99 (5.13) | 58.34 (4.95) | 59.48 (4.79) |
| 64.65 (4.77) | 58.34 (4.95) | 59.48 (4.79) |
| 61.17 (3.87) | 58.34 (4.95) | 59.48 (4.79) |

| CI | AAA – Mean(%) | AXB – Mean(%) |
|----|---------------|---------------|
| 1.09 (0.02) | 1.04 (0.02) | 1.03 (0.02) |
| 1.04 (0.02) | 1.04 (0.02) | 1.03 (0.02) |
| 1.04 (0.02) | 1.04 (0.02) | 1.03 (0.02) |

| CN | AAA – Mean(%) | AXB – Mean(%) |
|----|---------------|---------------|
| 0.90 (0.01) | 0.90 (0.01) | 0.91 (0.01) |
| 0.90 (0.01) | 0.90 (0.01) | 0.91 (0.01) |
| 0.90 (0.01) | 0.90 (0.01) | 0.91 (0.01) |

| HI | AAA – Mean(%) | AXB – Mean(%) |
|----|---------------|---------------|
| 0.09 (0.01) | 0.12 (0.01) | 0.11 (0.01) |
| 0.10 (0.00) | 0.14 (0.01) | 0.11 (0.01) |
| 0.14 (0.01) | 0.12 (0.01) | 0.11 (0.01) |

SD: standard deviation

https://doi.org/10.1371/journal.pone.0207232.t003
V_{20\%}, V_{30\%}, and maximum dose was lower in AAA compared to AXB, and all other parameters were higher in AAA than in AXB at the same grid size. The difference in bladder dosimetric parameters for the two algorithms was less than 1% of the reference values, which were the bladder volume for V_{x\%} and the prescription dose for the other parameters. In left and right femoral heads, only V_{20\%} to V_{40\%} showed a difference greater than 1% of reference values between AAA1 and AXB1, and only V_{30\%} showed a difference greater than 1% between AAA3 and AXB3. AAA predicted higher dosimetric parameters than AXB for both femoral heads.

**Table 4. Dosimetric data of organs at risk.**

|                | Bladder                  |                | Rectum                  |
|----------------|--------------------------|----------------|-------------------------|
|                | Maximum dose (%)         | Minimum dose (%) | Mean dose (%)          | Median dose (%) | Maximum dose (%)         | Minimum dose (%) | Mean dose (%) | Median dose (%) |
| **AAA**        |                          |                |                         |                |                          |                |                |                |
| 1 mm           | 107.33                   | 1.40           | 27.97                   | 14.27          | 104.54                   | 2.33           | 51.69         | 49.14         |
| 3 mm           | 106.19                   | 1.44           | 28.18                   | 14.83          | 103.71                   | 2.38           | 51.50         | 48.94         |
| 5 mm           | 105.18                   | 1.47           | 28.27                   | 15.53          | 103.01                   | 2.42           | 50.91         | 48.37         |
| **AXB**        |                          |                |                         |                |                          |                |                |                |
| 1 mm           | 107.19                   | 1.34           | 27.49                   | 13.73          | 105.16                   | 2.21           | 48.82         | 47.07         |
| 2 mm           | 106.55                   | 1.34           | 27.48                   | 13.80          | 104.41                   | 2.21           | 49.02         | 47.14         |
| 3 mm           | 106.70                   | 1.36           | 27.79                   | 14.18          | 104.81                   | 2.26           | 49.59         | 47.52         |
| **Right femoral head** |                  |                |                         |                |                          |                |                |                |
| AAA            |                          |                |                         |                |                          |                |                |                |
| 1 mm           | 47.71                    | 5.63           | 27.83                   | 28.58          | 47.17                    | 4.90           | 26.30         | 27.25         |
| 3 mm           | 47.54                    | 5.93           | 27.87                   | 28.56          | 46.88                    | 5.09           | 26.36         | 27.26         |
| 5 mm           | 47.21                    | 6.33           | 27.76                   | 28.39          | 46.59                    | 5.33           | 26.28         | 27.06         |
| AXB            |                          |                |                         |                |                          |                |                |                |
| 1 mm           | 47.11                    | 5.27           | 27.25                   | 28.00          | 46.48                    | 4.64           | 25.74         | 26.71         |
| 2 mm           | 47.11                    | 5.36           | 27.25                   | 27.99          | 46.44                    | 4.67           | 25.74         | 26.71         |
| 3 mm           | 47.23                    | 5.58           | 27.46                   | 28.19          | 46.42                    | 4.81           | 25.95         | 26.89         |

https://doi.org/10.1371/journal.pone.0207232.t004
The deviation of average dosimetric parameters depending on the algorithms and grid size is shown in Table 5. Almost AXB2 parameters were within 1% deviation, except the minimum dose to the PTV (1.95%) and right femoral head (1.71%). In the comparison with AXB3, the minimum dose of all structures except the bladder and the median dose to the bladder were different with AXB1 over 2%. However, most of the parameters of OARs were in excess of 2% different in comparison with AAA.

The subtracted, AXB1 and AAA1 dose distribution of one patient are shown in Fig 3. Likewise, Figs 4 and 5 show the subtracted distributions between 1 and 3 mm dose grid plan of AAA and AXB, respectively. The red, blue, yellow, green, and orange lines refer to the PTV, femoral heads, rectum, bladder, and body contour, respectively.

**Radiobiological comparison**

The average TCP and NTCP values with respect to algorithm and grid size are shown in Table 6. The difference of TCP values between AAA1 and AXB1 was 4.05%, and the difference between AAA3 and AXB3 was 0.9%. The largest NTCP of the rectum was 6.11% for AAA1, and the lowest NTCP was 3.23% for AXB1. In the bladder and the two femoral heads, no apparent NTCP values were observed, and all average NTCPs of the bladder and femoral heads were less than 1%, in all cases.

**Statistical analysis and dose calculation time comparison**

The p values for various comparative groups are shown in Table 7, and p values < .05 were observed in several dosimetric and radiobiological parameters. In the comparison between AXB1 and AAA1, most of the parameters had p-values of less than 0.05 and showed
statistically significant differences. In the comparison between AAA3 and AXB3, the p-values of the mean and median PTV doses exceeded 0.05 compared to the 1 mm group. In the AAA plan comparison of grid size, a statistically significant difference between the 1 mm grid and the others were found for all PTV parameters. In OARs, the p-value of $V_{30\%}$ of rectum was higher than 0.05 according to grid size in the AAA case. The median dose and $V_{95\%}$ of the PTV for the AXB plans were not statistically significant between 1 and 2 mm grids, and a statistically significant difference was found for all rectum parameters between AXB1 and AXB2.

The average calculation times according to type of algorithm and grid size are shown in Table 8. With AXB, it took 4061 seconds to calculate the dose at the 1 mm grid size, which was 1850 seconds longer than the 2211 seconds of AAA. In the 3 mm grid, the calculation time difference between the two algorithms is 17 seconds.

**Discussion**

In several studies, AXB has been reported to predict more accurate doses in heterogeneous medium than AAA, considering tissue specific interactions of photons [29–31]. In our results, the $V_{95\%}$ of the PTV was higher in the AAA than the AXB, and the differences for both algorithms were statistically significant (p values < 0.01). By using the ERB, an air cavity is usually generated in the rectum structure, which may cause a region of overlap between the PTV and air cavity. As a result, the AAA predicted a higher dose to the air cavity within the PTV and
could consequently estimate better target coverage than AXB. PTV may contain a portion of the ballooned rectum, where AAA overestimates the dose. However, this portion did not actually contain much dose, because this was air. Therefore, target coverage was estimated better than AXB when using the AAA.

Although the differences of median and mean doses to the PTV were statistically significant with respect to the type of algorithm at 1 mm grid size, these were less than 1% of the prescribed dose. At 3 mm grid size, the differences were not statistically significant and less than 1% of the prescription dose. The average $V_{95\%}$ of the PTV for the AAA plan decreased with increasing grid size, whereas $V_{95\%}$ for the plan with AXB increased slightly. For dose calculation, AXB discretizes the resolution in space, angle, and energy, and requires the material composition of voxels in the CT image by converting the Hounsfield unit to material derived mass density value. The automatic material composition is implemented with five biological materials divided according to the Hounsfield unit. When this material composition is performed on each dose voxel, the conversion of CT values might be affected by the size of the grid systematically. In our results, the p-value of the $V_{95\%}$ between AXB1 and AXB3 was less than 0.01, and the increased $V_{95\%}$ was 0.29% of the PTV volume. This means that the difference of dose grid size can produce systematic errors on target coverage of the PTV. However, it was not apparently changed. We think that the dose grid size may affect on each of the sub-procedure of the AXB dose calculation. Further study is needed to clarify these issues. The plans using AAA are more strongly influenced by the dose grid size compared to those using AXB. Due to the dramatic dose falloff near the PTV, the dose grid factors such as discrete sampling and volumetric averaging effects can work differently between both algorithms. As shown in Fig 4, a high dose difference was observed around the PTV, which is expressed in yellow. In general, AAA1 achieves a more precise dose prediction than AAA3 by using a fine dose grid, and this caused a sampling and volumetric averaging effect. Therefore, AAA1 may have a higher dose in the
dose falloff region than AAA3. In comparison to Figs 4 and 5, the grid effect appears to have a relatively larger influence on AAA than on AXB around the PTV, which affects target coverage relative to the grid size.

To compare plan qualities in greater detail, the CI and HI showed statistically significant differences according to algorithms using the same grid size. The average CI of AAA was higher than that of AXB and far from the ideal value of one. This means that the volume above the prescribed dose is greater than the target volume when using AAA, and treatment plan could be evaluated to have worse conformity with AAA than AXB. However, the HI of AAA was less than that of AXB, and the AAA plan evaluations were more homogeneous than those of AXB. This, specifically, was the reason for the low dose in the air cavity of the PTV when using AXB. High doses in PTV air caused the increase in D98, leading to the decrease in HI. However, the dose overestimated by AAA in PTV air was not appropriate, and HI reduction of AAA did not guarantee the quality of a better treatment plan. Finally, AAA and AXB did not significantly differ with respect to the CN at the same grid size, as the average value was not apparently different between both algorithms.

As shown in Fig 3, the rectal dose was higher in AAA1 than in AXB1, and the dose of overlapped region between the PTV and the rectum was also higher in AAA1 compared with AXB1. In the subtracted dose distributions, dose difference between AAA1 and AXB1 were
apparently large in only the rectum region, and AXB1 showed higher doses than AAA1 in the upper region of the rectum. Other studies have reported that the accuracy of AXB was superior to that of AAA in low density media [14, 15, 17, 18] and that AXB predicted higher doses of penumbra than the AAA in the air cavity [32, 33]. Based on the LBTE, AXB can consider the

Fig 5. Dose distribution subtracted the plan with 1 mm grid from that of 3 mm in AXB. Each image are axial plane of one patient from the top of planning target volume to the bottom of that at 12 mm intervals.

Table 6. Average and standard deviations of tumor control probability (TCP) and normal tissue complication probability (NTCP) values depending on the algorithms and grid size.

|               | AAA 1 mm Average (SD) | AAA 3 mm Average (SD) | AAA 5 mm Average (SD) | AXB 1 mm Average (SD) | AXB 2 mm Average (SD) | AXB 3 mm Average (SD) |
|---------------|------------------------|------------------------|------------------------|------------------------|------------------------|------------------------|
| TCP (%)       | Prostate (PTV)         | 87.49 (2.02)           | 86.59 (2.14)           | 82.37 (3.07)           | 83.44 (3.06)           | 84.32 (2.44)           | 85.69 (2.23)           |
| NTCP (%)      | Bladder                | 0.05 (0.18)            | 0.05 (0.19)            | 0.04 (0.15)            | 0.05 (0.18)            | 0.05 (0.18)            | 0.05 (0.19)            |
|               | Rectum                 | 6.11 (3.18)            | 5.55 (2.79)            | 4.77 (2.52)            | 3.23 (1.58)            | 3.42 (1.66)            | 3.99 (1.93)            |
|               | Rt femoral head        | 0.04 (0.11)            | 0.04 (0.11)            | 0.04 (0.10)            | 0.03 (0.08)            | 0.03 (0.09)            | 0.03 (0.09)            |
|               | Lt femoral head        | 0.05 (0.14)            | 0.05 (0.13)            | 0.04 (0.12)            | 0.03 (0.10)            | 0.03 (0.10)            | 0.04 (0.11)            |

SD: standard deviation

https://doi.org/10.1371/journal.pone.0207232.t006
Table 7. Statistical analysis results depending on the dose calculation algorithms and grid size. The comparison groups are (AAA1 vs AXB1), (AAA3 vs AXB3), (AAA1 vs AAA3), (AXB1 vs AXB2), (AXB1 vs AXB3) from the left.

| Algorithm | AAA vs AXB | AAA | AXB |
|-----------|------------|-----|-----|
|           | Grid size  | 1 mm | 3 mm | 1 vs 3 mm | 1 vs 5 mm | 1 vs 2 mm | 1 vs 3 mm |
| PTV       | Mean dose  | < .01 | 0.58 | < .01     | < .01     | < .01     | < .01     |
|           | Median dose| < .01 | 0.52 | < .01     | < .01     | 0.26      | < .01     |
|           | TCP        | < .01 | < .01| < .01     | < .01     | < .01     | < .01     |
|           | V95%       | < .01 | < .01| < .01     | < .01     | 0.16      | < .01     |
|           | HI         | < .01 | < .01| < .01     | < .01     | < .01     | < .01     |
|           | CI         | < .01 | < .01| < .01     | < .01     | < .01     | < .01     |
|           | CN         | 0.391 | 0.852| < .01     | < .01     | < .01     | < .01     |
| Bladder   | Mean dose  | < .01 | < .01| < .01     | < .01     | 0.56      | < .01     |
|           | Median dose| < .01 | < .01| < .01     | < .01     | 0.01      | < .01     |
|           | Maximum dose| 0.42 | 0.01| < .01     | < .01     | < .01     | < .01     |
|           | V30%       | < .01 | < .01| < .01     | < .01     | < .01     | < .01     |
|           | V50%       | < .01 | < .01| 0.25      | 0.01      | < .01     | < .01     |
|           | V70%       | < .01 | < .01| < .01     | < .01     | 0.09      | 0.37      |
|           | NTCP       | 0.59  | < .01| < .01     | 0.26      | 0.03      | 0.52      |
| Rectum    | Mean dose  | < .01 | < .01| < .01     | < .01     | < .01     | < .01     |
|           | Median dose| < .01 | < .01| < .01     | < .01     | 0.02      | < .01     |
|           | Maximum dose| 0.01 | < .01| < .01     | < .01     | < .01     | < .01     |
|           | V30%       | 0.71  | 0.911| 0.50      | 0.20      | 0.02      | 0.63      |
|           | V50%       | < .01 | < .01| < .01     | < .01     | < .01     | < .01     |
|           | V70%       | < .01 | < .01| < .01     | < .01     | < .01     | < .01     |
|           | NTCP       | < .01 | < .01| < .01     | < .01     | < .01     | < .01     |
| Right femoral head | Mean dose | < .01 | < .01| < .01     | 0.02      | 1.00      | < .01     |
|           | Median dose| < .01 | < .01| < .01     | 0.16      | < .01     | 0.32      | < .01     |
|           | Maximum dose| < .01 | < .01| 0.02      | 0.01      | 0.90      | 0.51      |
|           | V30%       | < .01 | < .01| 0.17      | < .01     | < .01     | < .01     |
|           | V50%       | 0.04  | 0.04 | 0.35      | 0.43      | 0.35      | 0.08      |
|           | V70%       | 0.18  | 0.18 | 0.66      | 0.66      | 0.18      | 0.18      |
|           | NTCP       | 0.01  | < .01| 0.86      | 0.09      | 0.04      | < .01     |
| Left femoral head | Mean dose | < .01 | < .01| < .01     | 0.40      | 1.00      | < .01     |
|           | Median dose| < .01 | < .01| < .01     | 0.59      | < .01     | 1.00      | < .01     |
|           | Maximum dose| < .01 | < .01| < .01     | 0.16      | < .01     | 0.38      | 0.72      |
|           | V30%       | < .01 | < .01| 0.77      | < .01     | < .01     | < .01     |
|           | V50%       | 0.04  | 0.04 | 0.89      | 0.04      | 0.08      | 0.04      |
|           | V70%       | 0.18  | 0.18 | 0.18      | 0.18      | 0.18      | 0.18      |
|           | NTCP       | 0.01  | < .01| 0.17      | 0.06      | 0.02      | < .01     |

Table 8. Average and standard deviation of dose calculation times recorded at different algorithms and grid size.

| Algorithm | AAA | AXB |
|-----------|-----|-----|
|           | 1 mm Average (SD) | 3 mm Average (SD) | 5 mm Average (SD) | 1 mm Average (SD) | 2 mm Average (SD) | 3 mm Average (SD) |
| Time (s)  | 2211 (155) | 245 (27) | 130 (10) | 4061 (922) | 671 (91) | 262 (26) |

SD: standard deviation
transport of photons and electrons through matter. Unlike AAA that uses a kernel of water with simplified density scaling for heterogeneous region, AXB use material library including five biologic materials (lung, adipose tissue, muscle, cartilage, and bone) and 16 non-biologic materials with a maximum density of steel. By using this, AXB can reflect characteristics of a various materials on dose calculation. Therefore, AXB is possible to correctly estimate the dose to a heterogeneous region such as the rectum. However, the dose calculation of AAA was associated with lateral and depth-directed components [34], and this algorithm cannot properly approximate the dose in the air due to the use of lateral scattering terms derived from Monte Carlo simulations in water [17]. The AAA compensates the heterogeneity by the inhomogeneity correction factor based on media density. As a result, AXB1 showed the predicted dose more correctly and estimated the higher rectum dose of the outside field, in comparison with AAA1.

The dosimetric difference of rectum was the largest among all OARs, and this largely depended on the algorithm. The average median and mean doses to the rectum were more than 2% higher in AAA1 compared with AXB1, and this dosimetric discrepancy has been reported to significantly influence radiobiological evaluation [35]. Likewise, we observed that the average NTCP in the rectum was heavily influenced by the respective algorithm. The rectum \( V_{40\%} \) to \( V_{90\%} \) showed large differences that depended on the algorithm rather than the dose grid. This was caused by the overlap between the PTV and rectum, which was induced in the optimization process to receive a high dose. In the bladder, most parameters showed statistically significant changes with both algorithms, however the changes were less than 1% of reference. Both femoral heads showed significant differences according to the algorithm and grid size, however the differences were not apparent.

In our results, the TCP differences according to algorithm and grid size were statistically different, and the TCP decreased as grid size increased with AAA. In contrast, the TCPs of AXB1 and AXB2 showed similar values, and the TCPs increased about 1%, in comparison with AXB1, as dose grid size increased to 3 mm. Under the same grid size, the average TCP of AAA was higher than that of AXB. By using the AAA, the target coverage (\( V_{95\%} \)) was superior to that of AXB. Therefore, the calculated TCP of AAA was relatively higher than that of AXB, and the TCP differences among the algorithms showed a quite large variations (4.05%), which were statistically significant (p-value <0.01). We used EUD-based TCP models. As described above, AAA predicted doses with overestimated errors in the air inside PTV than for AXB. This leads to the increase in EUD and TCP of AAA compared with AXB. The NTCP difference of the rectum was the largest among all the OARs, and this depended on the algorithm. The average NTCP of AAA was higher than that of AXB, and the average NTCP of the rectum decreased with increasing dose grid in the case of AAA, and inversely increased with AXB. In addition, all p-values of the rectum were less than 0.01, depending on the algorithm and grid size. This means that the rectum was influenced significantly by both algorithm and dose grid. Although we performed the statistical analysis depending on the algorithms and grid sizes, the impact of these factors on the bladder and both femoral heads was not clearly predictable due to very low NTCP values.

Rectal complications in radiation therapy are mainly caused by rectal tissue damage, and the dose deposited in air may not contribute to rectal toxicity from a clinical viewpoint. Our results for NTCP and dosimetric parameters in rectum contoured with air include may not have a direct impact on the clinical significance. Streller et al. [36] evaluated the impact of ERB on rectal wall dose by using AAA algorithm. However, dosimetric properties of two algorithms were different at the interface between air and matter, and the AXB showed more similar dose distribution to film measurement than AAA at the interface [33]. In addition, the dose rebuild-up has been reported with AXB [33]. The dose rebuild-up causes a large dose increase
from air to matter, which can have a significant impact on dose calculations. Therefore, as a future study, additional analysis of the rectal well according to the algorithm and grid is necessary to perform monitoring of the control/complications of patients.

The International Commission on Radiation Units recommends 5% of overall absorbed dose delivery accuracy, and the American Association of Physicists in Medicine recommends a 2% dose calculation accuracy [35, 37]. Considering the calculation time, the use of a 1 mm grid is not feasible in the clinical practice. In the previous reported studies, the use of a 2.5 mm grid should be recommended to reduce the dose distribution error, and 2 mm should be recommended at least in high dose gradients [38, 39]. Therefore, from our results, we recommended a 2 mm grid with AXB, considering the relatively short dose calculation time and calculation accuracy for prostate VMAT.

Conclusion
The AXB and AAA showed statically significant difference in dosimetric and radiobiological parameters. Our study demonstrated that the calculated grid size worked sufficiently enough to influence the plan quality evaluation. As previously reported, the AXB and fine dose grid provided significantly accurate dose calculations in air cavity regions, and our results indicated that AXB2 had sufficient accuracy and was reasonable, in terms of dose calculation time, compared the AXB1. So, we suggest employing AXB algorithm and 2 mm grid for improving treatment efficiency of VMAT plans for prostate cancer.

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References
1. Otto K. Volumetric modulated arc therapy: IMRT in a single gantry arc. Medical physics. 2008; 35 (1):310–7. https://doi.org/10.1118/1.2818738 PMID: 18293586
2. Jiang X, Li T, Liu Y, Zhou L, Xu Y, Zhou X, et al. Planning analysis for locally advanced lung cancer: dosimetric and efficiency comparisons between intensity-modulated radiotherapy (IMRT), single-arc/partial-arc volumetric modulated arc therapy (SA/PA-VMAT). Radiation Oncology. 2011; 6(1):140.
3. Rong Y, Evans J, Xu-Welliver M, Pickett C, Jia G, Chen Q, et al. Dosimetric evaluation of intensity-modulated radiotherapy, volumetric modulated arc therapy, and helical tomotherapy for hippocampal-avoidance whole brain radiotherapy. PloS one. 2015; 10(4):e0126222. https://doi.org/10.1371/journal.pone.0126222 PMID: 25894615

4. Zhang W-Z, Zhai T-T, Lu J-Y, Chen J-Z, Chen Z-J, Li D-R, et al. Volumetric modulated arc therapy vs. c-IMRT for the treatment of upper thoracic esophageal cancer. PloS one. 2015; 10(3):e0121385. https://doi.org/10.1371/journal.pone.0121385 PMID: 25815477

5. Ostheimer C, Hübisch P, Janich M, Gerlach R, Vordermark D. Dosimetric comparison of intensity-modulated radiotherapy (IMRT) and volumetric modulated arc therapy (VMAT) in total scalp irradiation: a single institutional experience. Radiation oncology journal. 2016; 34(4):313. https://doi.org/10.3857/roj.2016.01935 PMID: 27951625

6. Quan EM, Li X, Li Y, Wang X, Kudchadker RJ, Johnson JL, et al. A comprehensive comparison of IMRT and VMAT plan quality for prostate cancer treatment. International Journal of Radiation Oncology• Biology• Physics. 2012; 83(4):1169–78.

7. van Lin ENT, van der Vught LP, Witjes JA, Huisman HJ, Leer JW, Visser AG. The effect of an endorectal balloon and off-line correction on the interfraction systematic and random prostate position variations: a comparative study. International Journal of Radiation Oncology• Biology• Physics. 2005; 61(1):278–88.

8. Vanneste BG, van Wijk Y, Lutgens L, Van Limbergen E, van de Beek K, et al. Dynamics of rectal balloon implant shrinkage in prostate VMAT Dynamiken eines schrumpfenden rekten Ballonimplantats während Prostata-VMAT. Strahlentherapie und Onkologie. 2018; 194(1):31–40. https://doi.org/10.1007/s00066-017-1222-x PMID: 29038832

9. Park JM, Park S-Y, Kim J-i, Carlson J, Kim JH. The influence of the dose calculation resolution of VMAT plans on the calculated dose for eye lens and optic pathway. Australasian physical & engineering sciences in medicine. 2017; 40(1):209–17.
22. Huang B, Wu L, Lin P, Chen C. Dose calculation of Acuros XB and Anisotropic Analytical Algorithm in lung stereotactic body radiotherapy treatment with flattening filter free beams and the potential role of calculation grid size. Radiation Oncology. 2015; 10(1):53.

23. Kang SW, Suh TS, Chung JB, Eom KY, Song C, Kim IA, et al. Comparison of dosimetric and radiobiological parameters on plans for prostate stereotactic body radiotherapy using an endorectal balloon for different dose-calculation algorithms and delivery-beam modes. Journal of the Korean Physical Society. 2017; 70(4):424–30.

24. Kim JS, Chung JB, Kim IA, Eom KY. Dosimetric effects of endorectal balloons on intensity-modulated radiation therapy plans for prostate cancer. Journal of the Korean Physical Society. 2013; 63(8):1637–43.

25. Gay HA, Niemierko A. A free program for calculating EUD-based NTCP and TCP in external beam radiotherapy. Physica Medica: European Journal of Medical Physics. 2007; 23(3):115–25.

26. Niemierko A, Michael G. Modeling of normal tissue response to radiation: the critical volume model. International Journal of Radiation Oncology• Biology• Physics. 1993; 25(1):135–145.

27. Deasy JO, Blanco AI, Clark VH. CERR: a computational environment for radiotherapy research. Medical physics. 2003; 30(5):979–85. https://doi.org/10.1118/1.1568978 PMID: 12773007

28. Wilcoxon F. Individual comparisons by ranking methods. Biometrics bulletin. 1945; 1(6):80–83.

29. Kan MW, Leung LH, So RW, Yu PK. Experimental verification of the Acuros XB and AAA dose calculation adjacent to heterogeneous media for IMRT and RapidArc of nasopharyngeal carcinoma. Medical physics. 2013;40(3).

30. Kan MW, Leung LH, Peter K. Dosimetric impact of using the Acuros XB algorithm for intensity modulated radiation therapy and RapidArc planning in nasopharyngeal carcinomas. International Journal of Radiation Oncology• Biology• Physics. 2013; 85(1):e73–e80.

31. Rana S, Rogers K. Dosimetric evaluation of Acuros XB dose calculation algorithm with measurements in predicting doses beyond different air gap thickness for smaller and larger field sizes. Journal of medical physics/Association of Medical Physicists of India. 2013; 38(1):9.

32. Kang S-W, Suh T-S, Chung J-B, Eom K-Y, Kim I-A, Kim J-S, et al. Dosimetric accuracy of AAA and acuros XB dose calculations within an air cavity for small fields of a 6-MV flattening filter-free beam. Journal of the Korean Physical Society. 2015; 67(12):2138–45.

33. Kang SW, Chung JB, Lee JW, Kim MJ, Kim YL, Kim JS, et al. Dosimetric accuracy of the Acuros XB and Anisotropic analytical algorithm near interface of the different density media for the small fields of a 6-MV flattening-filter-free beam. International Journal of Radiation Research. 2017; 15(2):157–65.

34. Tillikainen L, Helminen H, Torsti T, Siiljamäki S, Alakuijala J, Pyyry J, et al. A 3D pencil-beam-based superposition algorithm for photon dose calculation in heterogeneous media. Physics in Medicine & Biology. 2008; 53(14):3821.

35. Papakonstantinou N, Battista JJ, Boyer AL, Kappas C, Klein E, Mackie TR, et al. Tissue inhomogeneity corrections for megavoltage photon beams. AAPM Task Group. 2004; 65:1–142.

36. Strreller T, Rusch U, Herraiz Lablanca MD, Minneken I, Najafi Y, Shrestha B, et al. The effect of endorectal balloon on anorectal dose during postoperative volumetric arc radiotherapy of prostate cancer. Radiotherapy and Oncology. 2017; 123(3):454–8. https://doi.org/10.1016/j.radonc.2017.04.014 PMID: 28464996

37. Report 24. Journal of the International Commission on Radiation Units and Measurements. 1976;os13 (1):NP-NP.

38. Chung H, Jin H, Palta J, Suh TS, Kim S. Dose variations with varying calculation grid size in head and neck IMRT. Physics in Medicine & Biology. 2006; 51(19):4841.

39. Dempsey JF, Romeijn HE, Li JG, Low DA, Palta JR. A Fourier analysis of the dose grid resolution required for accurate IMRT fluence map optimization. Medical physics. 2005; 32(2):380–8. https://doi.org/10.1118/1.1843354 PMID: 15789583
