A comparative analysis of Acuros XB and the analytical anisotropic algorithm for volumetric modulation arc therapy

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ABSTRACT

Background: This study aimed to verify the dosimetric impact of Acuros XB (AXB) (AXB, Varian Medical Systems Palo Alto CA, USA), a two model-based algorithm, in comparison with Anisotropic Analytical Algorithm (AAA) calculations for prostate, head and neck and lung cancer treatment by volumetric modulated arc therapy (VMAT), without primary modification to AAA. At present, the well-known and validated AAA algorithm is clinically used in our department for VMAT treatments of different pathologies. AXB could replace it without extra measurements. The treatment result and accuracy of the dose delivered depend on the dose calculation algorithm.

Materials and method: Ninety-five complex VMAT plans for different pathologies were generated using the Eclipse version 15.0.4 treatment planning system (TPS). The dose distributions were calculated using AAA and AXB (dose-to-water, AXBw and dose-to-medium, AXBm), with the same plan parameters for all VMAT plans. The dosimetric parameters were calculated for each planning target volume (PTV) and involved organs at risk (OAR). The patient specific quality assurance of all VMAT plans has been verified by Octavius®-4D phantom for different algorithms.

Results: The relative differences among AAA, AXBw and AXBm, with respect to prostate, head and neck were less than 1% for PTV D95%. However, PTV D95% calculated by AAA tended to be overestimated, with a relative dose difference of 3.23% in the case of lung treatment. The absolute mean values of the relative differences were 1.1 ± 1.2% and 2.0 ± 1.2%, when comparing between AXBw and AAA, AXBm and AAA, respectively. The gamma pass rate was observed to exceed 97.4% and 99.4% for the measured and calculated doses in most cases of the volumetric 3D analysis for AAA and AXBm, respectively.

Conclusion: This study suggests that the dose calculated to medium using AXBm algorithm is better than AAA and it could be used clinically. Switching the dose calculation algorithm from AAA to AXB does not require extra measurements.

Key words: anisotropic analytical algorithm; planning target volume; volumetric modulated arc therapy; multi-leaf collimator; gamma index

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Introduction

The Anisotropic Analytical Algorithm has been broadly utilized for dose calculation in the Eclipse treatment planning system. Van Esch et al. [1] reported that AAA improves the accuracy of dose calculations, compared to the single pencil beam (SPB) algorithm, and can achieve 5% agreement with measurements in thoracic phantom. In spite of this, AAA has been noticed to significantly overesti-
mate the dose near air-tissue interfaces [2]. A novel
dose calculation algorithm called Acuros XB, which
explicitly solves the Linear Boltzmann Transport
Equation (LBTE), has been applied for clinical prac-
tice [3]. LBTE is the leading equation that describes
the distribution of radiation particles resulting from
their interactions with matter. AXB discretizes the
space, angle, and energy variables of the LBTE into
grids and computes the energy fluence variation of
electrons and scattered photons in a substance.

Han et al. [4] and Bush et al. [5] have shown that
AXB could reach accuracy comparable to that of
Monte Carlo methods, which are broadly consid-
ered the gold standard for precise dose calculation
used in radiation therapy in phantom experiments,
assuming the existence of homogeneous water
and heterogeneous media. Several clinical studies
have been performed for dosimetric comparison of
VMAT and intensity modulated radiation therapy
(IMRT) plans between AXB and AAA, indicating
that AXB underestimated the doses to PTV or nor-
tmal tissues in the cases of prostate, lung, head and
neck and pelvis treatment, compared to AAA [6–9].
Compared to these results, another study has re-
vealed that AAA underestimated the dose to the
spine [7]. The AXB and AAA difference depends on
the treatment site and beam energy. AXB provides
two dose reporting methods: dose-to-water (AXB_w)
and dose-to-medium (AXB_m). For the AXB_w en-
ergy dependent fluence-to-dose response functions
are based on water, whereas for the AXB_m they are
based on each material. It was a subject of debate
whether to select AXB_w or AXB_m for clinical prac-
tice [8]. Walters et al. [8] have determined that the
dose-to-water method offers a better evaluation of
the dose to sensitive tissue in the bone, compared
to the dose-to-medium one.

The purpose of this study is to ascertain that
implementing the new dose computation algorithm
will not majorly change the clinical treatment plans
in our department. The study clearly helps new cen-
ters that are willing to switch the TPS from AAA to
AXB. The pre-treatment patient-specific quality as-
surance with the Octavius system will additionally
boost up the clinical results.

Materials and methods

Ninety-five patients, who had undergone treat-
ment in our institute during April–December 2019,
were included into this study as shown in Table 1.

Dose calculation and planning

For all the three types of pathologies, VMAT
plans were generated by the Eclipse TPS for a Clin-
ac iX accelerator, equipped with a Millennium 120
multileaf collimator (MLC) (Varian Medical Sys-
tems. Palo Alto. CA. USA), using 6 and 18 MV pho-
ton beams. The VMAT plan was created in AXB_m.
For the prostate cases, the total dose prescribed to
the PTV was 76 Gy, with a daily dose of 2.0 Gy in 38
fractions. The prostate VMAT plans were generated
using two full arcs with 6 & 18 MV photon beams,
as needed. The head and neck VMAT plans were
generated by using two or three full arcs with a 6
MV photon beam and the total prescription dose to
PTV was 70 Gy (2.0 Gy/fraction). The prostate and
head and neck cases were treated in simultaneous
integrated boost (SIB) fractionation schemes. The
greatest dose was mentioned for the study. For lung
SBRT cases, the prescription dose to PTV was 48 Gy
in 4 fractions of 80% of isodose. The VMAT plans
for lung SBRT were made using two or three partial
arcs with 6 MV photon beam. The dose calculation
grid used in this study was 2.5 mm, except for 1.0
mm for lung SBRT cases. Then, each plan was recal-
culated for AAA and AXB_m, while maintaining the
AXB_m calculated monitor units, leaf motion, and
beam arrangement. The beam models in AAA and
AXB were based on the same physical data.

Evaluation of dosimetric parameters

For the sake of comparison, the relative dose and
volume differences in the corresponding dose-vol-
umetric parameters, obtained by AAA and AXB for the same case, were calculated as follows [10]:

\[
\text{Relative difference (\%)} = \frac{\text{Value of AXB}_x - \text{Value of AAA}}{\text{Value of AAA}} \times 100
\]

where: AXBx is selected between AXB_w and AXB_m depending on what dose reporting mode should be compared.

**Evaluation of the plans and statistical analysis**

For the PTV, the evaluation parameters included D_{95\%}, D_{98\%}, minimum dose and the mean dose. For the OARs, the analysis included the mean dose and a set of appropriate V_x values, V_x being the volume of the organ getting a dose of x or more. In the case of OARs, the analysis included an appropriately selected dose or volume parameter. For the rectum and bladder, V_{30}, defined as the volume that receives more than 30 Gy, was analyzed. The absolute doses in this study were presented in Gy and all the numerical data were rounded to the nearest tenth.

**Octavius phantom**

The 2D-Array, together with Octavius®-4D (PTW-Freiburg, Germany), have been widely described in the literature [11–13]. Figure 1 shows the normal setup of Octavius®-4D with the 2D array detector. An inclinometer mounted on the gantry ensures that the rotation unit always rotates along with the gantry, always keeping the 2D array perpendicular to the beam axis. The beam always hits the detector array in a perpendicular way, because the same face of the detector follows the gantry, so no correction factors are required.

To evaluate the dosimetric agreement between the measured and the calculated dose, the gamma evaluation method, implemented in the Verisoft 7.2.0 version, was used, where the measured dose matrix was used as reference. This calculation of the gamma index is based on the concept Low [14]. A two-sample paired t-test was used to compare the results for the average dose D_{95\%} and D_{98\%} and D_{\text{max}} of PTV prostate, head and neck and lung pathologies. A p-value < 0.05 was considered statistically significant.

**Results**

The relative differences among AAA, AXB_w and AXB_m, with respect to prostate, head and neck and lung cases, are shown in Figure 2. The PTV D_{95\%} and D_{98\%} were within 1.0 ± 0.23% difference between AXB_w and AXB_m to the AAA algorithm, for the prostate and head and neck cases, however, the value increases up to 3.23 ± 0.26% for the lung cases treatment by SBRT. Bladder V_{30} shows 1.3 ± 0.31% and 2.6 ± 0.29% difference between AAA and AXB_w, AXB_m respectively. The maximum difference was found to be 4% for esophagus mean dose and 2.12% in D_{\text{max}} spinal cord in the treatment of head and neck pathologies. The maximum dose to the thoracic wall was underestimated with maximum value of the relative dose difference of 3.95% from AAA during the treatment of SBRT.

Several factors may explain the fact that no significant dose differences were found at the level of the spinal cord. Unlike the lungs, the spinal cord receives most of its dose by scatter. Additionally, the much lower density of the lungs, compared to water, can easily result in higher difference in different dose calculation algorithms because of increased electron interaction paths. Whether dose computation algorithms compute significantly different
All the plans were analyzed [14] and the γ-pass rate with the 3% dose tolerance and 3 mm distance was calculated to harmonize the relation to the treatment planning system. Table 3 summarizes the results of the average γ-passing rate for the different pathologies examined in the present study, by 2D and 3D analysis for AAA, AXBw and AXBm. The results demonstrated that the gamma pass rates for all the plans were higher than ≤ 97.0 % by volumetric 3D analysis, where a 10% low-dose threshold was fixed [15]. Our clini-

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Table 2. The planning target volume (PTV) dose (D_{95\%}, D_{98\%} and D_{max}) for prostate, head and neck and lung cases. Statistical error indicates standard deviation. p_{a-m}: p-value between AAA and AXB_{m}. p_{a-w}: p-value between AAA and AXB_{w}

| PTV    | AAA       | AXB_{w}   | AXB_{m}   | p_{a-m} | p_{a-w} |
|--------|-----------|-----------|-----------|---------|---------|
| Prostate |           |           |           |         |         |
| D_{95\%} | 75.53 ± 0.6 | 74.69 ± 0.4 | 74.96 ± 0.3 | 0.01    | 0.006   |
| D_{98\%} | 74.28 ± 0.5 | 73.89 ± 0.3 | 74.68 ± 0.3 | 0.02    | 0.004   |
| D_{max}  | 79.43 ± 0.4 | 78.25 ± 0.4 | 80.59 ± 0.6 | 0.01    | 0.001   |
| Head and neck |           |           |           |         |         |
| D_{95\%} | 68.97 ± 0.8 | 68.58 ± 0.6 | 68.88 ± 0.6 | 0.037   | 0.026   |
| D_{98\%} | 68.52 ± 0.6 | 68.59 ± 0.3 | 68.64 ± 0.8 | 0.022   | 0.005   |
| D_{max}  | 74.84 ± 1.2 | 73.86 ± 0.9 | 74.59 ± 1.0 | 0.031   | 0.007   |
| Lung     |           |           |           |         |         |
| D_{95\%} | 50.01 ± 0.9 | 48.68 ± 0.5 | 48.51 ± 0.4 | 0.04    | 0.06    |
| D_{98\%} | 48.37 ± 1.1 | 46.93 ± 0.8 | 46.81 ± 0.6 | 0.02    | 0.05    |
| D_{max}  | 60.21 ± 0.4 | 60.12 ± 0.5 | 60.06 ± 0.5 | 0.01    | 0.02    |

Table 3. Average γ-passing rates for AAA, AXB_{w} and AXB_{m} for different pathologies in planar 2D and volumetric 3D analysis. Statistical error indicates standard deviation

| Pathologies | Planar analysis | 2D | Volumetric 3D analysis |
|-------------|----------------|----|------------------------|
|              | Coronal | Sagittal | Transversal | Coronal | Sagittal | Transversal |
| Gamma passing rates (%) for AAA |           |         |         |           |         |         |
| Prostate     | 95.8 ± 1.7 | 96.5 ± 0.2 | 96.2 ± 0.4 | 97.9 ± 1.2 |
| Head and neck| 97.2 ± 0.3 | 97.3 ± 1.1 | 98.1 ± 0.8 | 98.1 ± 1.1 |
| Lung         | 91.8 ± 1.9 | 91.2 ± 0.5 | 94.7 ± 1.1 | 97.4 ± 1.4 |
| Gamma passing rates (%) for AXB_{w} |           |         |         |           |         |         |
| Prostate     | 96.6 ± 0.3 | 95.8 ± 1.1 | 96.2 ± 0.6 | 97.5 ± 0.4 |
| Head and neck| 96.2 ± 0.3 | 96.9 ± 0.5 | 98.2 ± 0.5 | 98.7 ± 0.5 |
| Lung         | 99.1 ± 0.1 | 99.2 ± 0.3 | 99.6 ± 0.1 | 99.7 ± 0.3 |
| Gamma passing rates (%) for AXB_{m} |           |         |         |           |         |         |
| Prostate     | 96.8 ± 1.5 | 96.5 ± 1.3 | 97.4 ± 0.8 | 99.4 ± 0.6 |
| Head and neck| 97.4 ± 1.4 | 97.8 ± 1.5 | 97.6 ± 1.0 | 99.6 ± 0.3 |
| Lung         | 95.8 ± 1.1 | 95.1 ± 1.1 | 99.6 ± 0.3 | 99.7 ± 0.1 |

A practical standard of 95% or greater for the gamma index percentage was standardized and was achieved for all the plans.

Discussion

For many years, the methods used to inter-compare the calculated doses to be delivered to different media have been subject to scientific debate [8]. This debate has generally addressed what is the best quantity to score (dose to medium or dose to water) with respect to the biological effect of radiation. Table 4 shows an overview of literature comparisons to the study.

Rana et al. [6] used Rapid Arc plans to perform a planning study of prostate cancer patients in which the clinical dosimetric effects of AAA and AXB were compared. Our results confirm their findings (range –0.21–0.67%) concerning the PTV D_{95\%}.

Kan et al. [16] reported that the mean and minimum doses to the PTV calculated by AXB were reduced by AAA for nasopharyngeal carcinoma. They found –2.0% and 4.0% discrepancies for...
AXB_m and AXB_w in bone content. The results of the current study showed that AAA computed, on average, an up to 1.01% higher maximum PTV dose than AXB_m, which was used for generating the treatment plans. Hirata et al. [17] determined that AXB showed agreement with the measurements within 2.6% at the high-density area, while AAA and PBC calculations overestimated the dose by more than 4.5% and 4.0%, respectively. These findings are in agreement to those of Fogliata et al. [18]. They showed the difference in the doses calculated using AXB and AAA was within 3% in lung planning target volumes. Bassi et al. [19] validated AXB in the presence of inhomogeneities for VMAT. They reported up to 1.8% dose calculated uncertainties with AXB. The absolute dose measured with GafChromic EBT in the heterogeneous phantom in the abdominal region proved that the dose differences after the air calculated by AXB are less than 3% while with AAA differences up to 11% can be obtained. Soh et al. [20] displayed ± 1.5% difference between AAA and AXB algorithm predicted depth dose, excluding the surface and buildup doses.

Han et al. [4] reported that AAA and AXB dose calculations agreed well with RPC lung phantom to TLD and film measurements for IMRT and VMAT plans. The cause for this small difference between AXB and AAA is attributed to the modeling of the heterogeneity of lung tissue in the AXB, compared to AAA, as reported by other studies [21]. Robinson D [22] has demonstrated that AAA overestimates the doses to the heterogeneity interface. Supporting this finding, Liu et al. [23] have also reported lower conformity (~2.1%), compared to AAA. Tajaldeen et al. [24] showed 2.3%, 1.3% and 0.7% discrepancies between AAA, AXB, and AXB_m algorithms and measured dose, respectively. Kumar et al. [25] found less than 1% discrepancy between AAA and AXB for mean PTV dose in deep-inspiration breath-hold respiratory techniques used for the treatment of left breast cancer.

In 2002, Liu [8] asserted that the dose-to-medium method allows a closer relationship between tissue response and dose, while Keall Paul argued against this statement and specified that all clinical knowledge and dosimetry protocols are based on the dose-to-water mode.

Eclipse AXB is based on the macroscopic cross sections of the media assigned from the CT scan for transport of photons and electrons. A biological material, such as the lung, adipose tissue, muscle, cartilage, or bone, can be assigned for voxels with a density < 3 g/cm³. The voxels are assigned a material design corresponding to the weighted proportion of the materials, when the density ranges overlap (e.g., adipose and muscle). AXB_m used the energy deposition cross section or restricted electronic stopping power from the medium at that point. AXB_m is then intrinsically computed so that the TPS reference dose should be specified in water, because the difference between water and tissue is inherently captured [26].

The results emphasize that several factors affect plan evaluation when using the Octavius®-4D phantom. They are especially improved if the more limiting local y-index computation approach is used. Indeed, the global y-index produces more homogeneous results with higher passing rates, because its tolerance level is computed with respect to the value of the maximum dose. The 2D approach considers each slice as independent of the surrounding volume, with the drawback that the

### Table 4. An overview of literature comparisons to current study

| Literature data | Pathologies | Treatment technique | AAA vs. AXB_m | AAA vs. AXB_w | Current study |
|-----------------|-------------|---------------------|---------------|---------------|---------------|
| Fogliata A et al. 2012 | Lung | VMAT | < 0.5% | 2.6% | 3.0% |
| Rana S et al. 2013 | Prostate | VMAT | 0.5% | 0.67% | -0.21% |
| Kan MW et al. 2013 | Head and neck | VMAT | -2.0% | 0.14% | 1.01% |
| Han T et al. 2013 | Lung phantom | IMRT and VMAT | -2.2% | 2.6% | 3.0% |
| Lui et al. 2014 | Lung | SBRT | -2.1% | 2.6% | 3.0% |
| Hirata K et al. 2015 | Head and neck | VMAT | 2.6% | 0.14% | 1.01% |
| Kumar L et al. 2020 | Breast | Breath hold | 1.0% | NA | NA |
| Bassi S et al. 2020 | Phantom study | Static | 1.8% | NA | NA |

VMAT — volumetric modulated arc therapy; IMRT — intensity modulated radiation therapy; SBRT — stereotactic body radiation therapy; NA — not applicable.
results are strongly dependent on the chosen plane, without a certain significant correlation between the magnitude of errors of different plans [27]. The 3D analysis allows a slice-by-slice evaluation, also considering the neighboring slices. Our results confirmed that the single slice evaluation (2D) always had an inferior agreement, compared to the 3D analysis and volumetric γ-index. Pulliam et al. [28] compared the two gamma results, using Monte Carlo computation as reference dose distribution, and quantified the increase of the passing pixels percent up to 3.2% in the 3D analysis, confirming our findings. Moreover, some problems that would be identified with 3D analysis might be missed in 2D individual planes. The 3D analysis could also highlight local regions where problems exist.

Conclusion

In the present study, the results showed that doses calculated to medium by AXB\textsubscript{m} could be used clinically for VMAT application. Moving from AAA to AXB does not require extra measurements because the dose difference between AAA and AXB were small in prostate and head and neck pathologies. However, the appropriateness of switching the dose calculation algorithm from AAA to AXB should be confirmed carefully from a clinical viewpoint in lung pathologies. AXB\textsubscript{m} results in advantage not only for prostate and head and neck but also for lung pathologies over AAA.

Conflict of interest
None declared.

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