Measuring the dose in bone for spine stereotactic body radiotherapy

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A B S T R A C T

Purpose: Current quality assurance of radiotherapy involving bony regions generally utilises homogeneous phantoms and dose calculations, ignoring the challenges of heterogeneities with dosimetry problems likely occurring around bone. Anthropomorphic phantoms with synthetic bony materials enable realistic end-to-end testing in clinical scenarios. This work reports on measurements and calculated corrections required to directly report dose in bony materials in the context of comprehensive end-to-end dosimetry audit measurements (63 plans, 6 planning systems).

Materials and methods: Radiochromic film and microDiamond measurements were performed in an anthropomorphic spine phantom containing bone equivalent materials. Medium dependent correction factors, \( k_{\text{med}} \), were established using 6 MV and 10 MV Linear Accelerator Monte Carlo simulations to account for the detectors being calibrated in water, but measuring in regions of bony material. Both cortical and trabecular bony material were investigated for verification of dose calculations in dose-to-medium (\( D_{\text{m,m}} \)) and dose-to-water (\( D_{\text{w,w}} \)) scenarios.

Results: For \( D_{\text{m,m}} \) calculations, modelled correction factors for cortical and trabecular bone in film measurements, and for trabecular bone in microDiamond measurements were 0.875(±0.1%), 0.953(±0.3%) and 0.962(±0.4%), respectively. For \( D_{\text{w,w}} \) calculations, the corrections were 0.920(±0.1%), 0.982(±0.3%) and 0.993(±0.4%), respectively. In the audit, application of the correction factors improves the mean agreement between treatment plans and measured microDiamond dose from −2.4%(±3.9%) to 0.4%(±3.7%).

Conclusion: Monte Carlo simulations provide a method for correcting the dose measured in bony materials allowing more accurate comparison with treatment planning system doses. In verification measurements, algorithm specific correction factors should be applied to account for variations in bony material for calculations based on \( D_{\text{m,m}} \) and \( D_{\text{w,w}} \).

1. Introduction

The complexity of treatment delivery has rapidly increased over recent years with a particular shift towards stereotactic treatments including stereotactic body radiation therapy (SBRT) [1–4]. The potential advantages to patients of effective local control with fewer hospital visits can only be realised if these treatments can be delivered safely. In contrast to conventional radiotherapy, stereotactic techniques employ smaller treatment margins and higher doses per fraction, greatly increasing the requirements for accurate dosimetry delivered with precision [1,5]. For accurate SBRT treatments the correct vendor supplied tools need to also be underpinned with a strong physics programme.

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Technological advances in image-guided radiation therapy, treatment planning and modulated delivery modalities have facilitated safe ablative treatment of extra cranial target volumes with increased conformity and increased geometric precision [1,5]. Nonetheless, there remains the weakness that quality assurance of SBRT treatments is commonly performed on homogeneous water or PMMA phantoms [6–8], ignoring the complexities of heterogeneities where dosimetry issues are likely to occur. Improvements in modern phantom materials including synthetic bone mean that suitable heterogeneous phantoms are now readily available [9,10]. However, measurements in non-water material are challenging as detector calibrations are only provided for dose to water. This work seeks to fill this gap by providing the modelled corrections for measuring in bone, facilitating improved end-to-end testing on more realistic heterogeneous phantoms for local hospital commissioning and dosimetry audit programmes.

The spine is the most common site of skeletal metastases, affecting almost 40% of all cancer patients who develop metastatic disease [11–13]. SBRT offers effective local control in the case of spine metastases, making it the treatment of choice for many patients with this disease [5,13,14]. SBRT spine treatment plans use complex delivery systems such as intensity-modulated radiotherapy (IMRT), volumetric modulated arc therapy (VMAT) or robotic radiosurgery (Cyberknife).

End-to-end testing and dosimetry audits become increasingly important to ensure systemic errors are not present in the patient delivery pathway. The SBRT spine dosimetry audit by Imaging and Radiation Oncology Core (IROC) [9] found 60% of audit failures were due to systematic dose errors, highlighting the need for accurate dosimetry in end-to-end testing. Previously described SBRT dosimetry audits have addressed the need for non-water material corrections in the measurement of dose in bone. In the end-to-end testing for Spine SBRT described by Hardcastle et.al [15], the authors chose to avoid the bony materials all together and perform measurements in homogenous phantoms, as they lacked correction factors that were required for film measurements in bony regions. In the study by Lee et al. [16], measurements of SBRT spine were 3.9–5.3% higher than the planned dose when measuring in bony materials, with the discrepancies attributed to non-water like materials and calibration of detectors. Traditionally, radiotherapy detectors are calibrated using a normalised dose to water approach, in a water phantom. Currently no primary standards dosimetry laboratories offer calibrations in non-water media such as bone. Even if a treatment planning system (TPS) calculates dose to water, a measurement in a bone phantom and a detector calibrated in a water phantom will not give the correct answer, as the secondary electrons from the surrounding bone increase the dose compared to calibration conditions in water. In this paper, we measure and calculate the corrections required to directly measure in bony materials and allow comparison to TPS algorithms that calculate either dose to water, in water \(D_{\text{w},\text{w}}\) or dose to medium, in medium \(D_{\text{m},\text{m}}\).

To calculate the corrections required, the definitions of dose to water and dose to medium firstly need to be clearly defined. Modern TPS used in radiotherapy compute dose to a patient using a variety of dose computation algorithms and mathematical approximations, leading to inconsistencies in the clinical data. TPS have historically calculated radiotherapy dose as dose to water, in water with variable electron density \(D_{\text{w},\text{w}}\). This method accounts for radiation transport through differing densities of patient tissues but not the tissue types themselves [17]. In the \(D_{\text{w},\text{w}}\) case where bony materials are present, the TPS treats the bone as density water, whereas in reality the TPS treats it as water with a density of that of the bone. Since the introduction of Monte Carlo and Linear Boltzmann Solver based calculations, dose can be reported as dose to medium-in-medium \(D_{\text{m},\text{m}}\) or as dose to water-in-medium \(D_{\text{m},\text{m}}\). For \(D_{\text{m},\text{m}}\) calculations, the TPS accounts for radiation transport through a number of patient like materials and the density of such materials. For \(D_{\text{m},\text{m}}\) calculations, the TPS first performs a \(D_{\text{m},\text{m}}\) calculation, and then converts the dose to \(D_{\text{w},\text{w}}\) via stopping power ratios for Monte Carlo algorithms [18,19]. For Linear Boltzmann Solver algorithms, such as AcurosXB [20], calculations of electron fluence are performed in patient like materials, regardless of whether \(D_{\text{m},\text{m}}\) or \(D_{\text{w},\text{w}}\) is selected. For the final calculation step, the TPS uses medium or water cross sections and densities depending on the reporting quantity. For patient like materials with atomic numbers close to water these processes result in very similar calculations of dose [18]. Large discrepancies of approximately 10% can be observed between the different reporting methods in regions of high density, such as cortical bone [17–19,21]. Andreo [18] recommended avoiding the conversion from \(D_{\text{m},\text{m}}\) to \(D_{\text{w},\text{w}}\) via stopping power ratios due to the uncertainties involved. Gladstone et.al [22] also recommend reporting \(D_{\text{m},\text{m}}\) for consistency in NRG clinical trials. For the purposes of this study, the \(D_{\text{w},\text{w}}\) reporting mode has not been investigated, as all AXB and MC plans included in the analysis were reported as \(D_{\text{m},\text{m}}\). Absorbed dose is traditionally defined as \(D_{\text{w},\text{w}}\) [23] and radiotherapy detectors are traditionally calibrated in terms of absorbed dose to water \(D_{\text{w},\text{w}}\) which creates challenges in dose verification for calculations in bony material. With the aim of a consistent definition of dose, and in particular, only one definition of absorbed dose to water, we propose a methodology for measuring dose in bone as either as \(D_{\text{m},\text{m}}\) or \(D_{\text{w},\text{w}}\) based on the primary reporting mode of the calculation algorithm [24]. The methodology is applied in the context of an end-to-end clinical dosimetry audit of SBRT spine treatments. Independent dosimetry audits are recommended by international SBRT guidelines for validation of treatment delivery of such complex techniques [1,2,25]. The Australian Clinical Dosimetry Service (ACDS) [26] conducts independent dosimetry audits across Australian and New Zealand Radiation Oncology facilities. The ACDS introduced the SBRT modality into the Level III end-to-end audit program in 2018. The Level III audit is a dosimetric intercomparison where an anthropomorphic phantom undergoes all steps within the patient treatment pathway [27]. The ACDS SBRT audit consists of multiple cases replicating the most common tumour sites treated with SBRT; lung, spine and soft tissue. Measurements are conducted on-site at each facility by ACDS representatives, using an anthropomorphic thorax phantom (CIRS, Norfolk, VA, USA), which includes trabecular and cortical bone, inhale lung and soft tissue equivalent materials. The primary detectors used in the audit are Gafrichrom EBT3 radiochromic film (Ashland, Bridgewater, NJ USA) and a PTW 60019 microDiamond (PTW Freiburg, Germany). This paper discusses the measurement of dose in the SBRT spine treatment case of the audit with the application of Monte Carlo modelled correction factors for the bony materials in the phantom, including 63 measured SBRT treatment plans from six treatment planning systems.

2. Materials and methods

2.1. Theoretical considerations

Discrepancies between the measured and planned doses in the bony regions of the spine were expected due to the detectors being calibrated in terms of absorbed dose to water. This was independent of the algorithm calculating either dose to medium \(D_{\text{m},\text{m}}\) or dose to water \(D_{\text{w},\text{w}}\). In the \(D_{\text{m},\text{m}}\) case, the plan is calculating the dose to bone, but the detector is calibrated in terms of dose to water and so requires a correction. In the \(D_{\text{w},\text{w}}\) case where bone is present, the planning system is calculating dose to high density water. Whilst the plan is calculating dose to water, the detector in the phantom is surrounded by high Z bone material which does not match the calibration conditions of the detector measuring in a water phantom. To account for these discrepancies, we propose the use of correction factors \(k_{\text{med}}\) for both the film and microDiamond detectors when measuring in the different bony material as per Eqs. (1) and (2).

\[
D_{\text{b},\text{b}} = M_b \times N_{\text{D},\text{w}} \times k_{\text{med},\text{b},\text{b}}
\]  

\[
D_{\text{H},\text{D},\text{w}} = M_b \times N_{\text{D},\text{w}} \times k_{\text{med},\text{D},\text{D},\text{w}}
\]
In the \( D_{m,m} \) algorithm scenario, the quantity of interest is the dose in the bony material (\( D_{b,b} \)) which is calculated through measurement in bone (\( M_b \)), the detector calibration factor (\( N_{det,b} \)), and a medium dependent correction factor (\( k_{med} \)). A schematic of modelled correction factors for the \( D_{m,m} \) scenario, as measured from a detector in bone (\( D_{det,b} \)) to the dose to bone in bone (\( D_{b,b} \)), is shown in Fig. 1. Four situations are modelled for the \( k_{med,bb} \) correction factors. Fig. 1a shows the definition of what the detector is calibrated to; i.e., calibration to absorbed dose to water, in a homogeneous water phantom. Fig. 1b then shows the detector calibration conditions; where the detector is in a homogeneous water phantom. The audit measurement conditions are shown in Fig. 1c, with the detector measuring in bone material. Finally, Fig. 1d shows the TPS calculation of the dose to a voxel of bone, surrounded by bony material. The dose at the centre voxel in each scenario is used to calculate the \( k_{med} \) as per Eq. (3).

\[
k_{med,bb} = \frac{D_{b,b}}{D_{det,b}} \frac{D_{det,w}}{D_{det,b}}\tag{3}
\]

For the \( D_{w,w} \) algorithm scenario, the quantity of interest is the dose in the bony region, which is treated as ‘high density water’ by conventional TPS. The correction for conventional TPS calculations using water materials with the density of bone (\( D_{b,b} \)) is achieved by replacing the bone in the final schematic Fig. 1d with high density water. The proposed \( k_{med} \) for the \( D_{w,w} \) scenario is given in Eq. (4).

\[
k_{med,wb} = \frac{D_{b,w}}{D_{w,w}} \frac{D_{det,w}}{D_{det,b}}\tag{4}
\]

### 2.2. Monte Carlo modelling

Monte Carlo simulations were performed for Gafchromic Film and stylised microDiamond geometries obtained from the manufacturers’ specifications [28,29]. All Monte Carlo modelling was performed in the EGSnrc [30] user code DOSXYZnrc [31] with the 2020 version. The source used in these simulations was a phase space file generated in BEAMnrc [32] for \( 4 \times 4 \) cm\(^2\) fields from a previously validated 6 MV and 10 MV Elekta Synergy Linear Accelerator model [33]. A \( 4 \times 4 \) cm\(^2\) field represents typical field sizes for the SBRT spine plans included in the audit. In all simulations, the incident beam was from above the modelled geometries at 80 cm SSD. Density correction files generated with the ESTAR [34] program were used for CIRS bony materials and diamond material. All default transport options were used, unless specified otherwise in Table 1. In all cases, the unit being scored is dose, and the uncertainties represent one standard deviation uncertainties calculated with history-by-history statistical analysis. All simulations were performed on the National Computational Infrastructure (NCI) high powered computer cluster with Xeon Platinum 8274 CPUs, and the total computation time to generate the various \( k_{med} \) factors took between 310 and 1680 h.

EBT3 Gafchromic film consists of 28 \( \mu m \) active layer, between two polystyrene layers of 125 \( \mu m \) [28]. This structure was modelled parallel to the incident beam, in the axial plane in the centre of a 25 mm bone cube. The bone cube consisted of a 15 mm cube of CIRS Trabecular Bone (density 1.197 g/cm\(^3\)) surrounded by 5 mm outer shell of CIRS Cortical Bone (density 1.91 g/cm\(^3\)). This geometry is a simplistic representation of the body of the phantom vertebrae. The film and bone cube were placed at the centre of a \( 30 \times 30 \times 30 \) cm\(^3\) cube of water. The resolution of the central bone cube was modelled in 1.0 mm\(^3\) voxels to evaluate the interface effects, with the surrounding water voxels at 13.75 cm\(^3\). The density and material of the bone cubes were varied to obtain each of the scenarios detailed in Eqs. (3) and (4), for both 6MV and 10MV.

The microDiamond was modelled based on manufacturer provided specifications [29], with the stem of the detector as a 6.9 \( \times \) 6.9 \( \times \) 40 mm cubic of RW3 (polystyrene/epoxy, density 1.045 g/cm\(^3\)), and the active diamond layer as a 3.1 \( \times \) 3.1 \( \times \) 0.001 mm cubic of Carbon (density 3.53 g/cm\(^3\)), located 1 mm from the end of the stem. Surrounding the microDiamond was a 20 \( \times \) 20 mm\(^3\) of CIRS trabecular bone, replicating the measurement point of the detector in the SBRT phantom. The microDiamond and bone cube were placed at the centre of a \( 30 \times 30 \times 30 \) cm\(^3\) cube of water. The density and material of the bone cubes were varied to obtain each of the scenarios detailed in Eqs. (3) and (4) for both 6MV and 10MV. This is a somewhat simplified representation of the microDiamond, which may contribute additional uncertainty to the modelling.

Based on manufacturer supplied material composition data [35], density correction files for the CIRS cortical and trabecular bone were created using the ESTAR program [34]. These density correction files were used to generate the PEGS4 electron stopping power data files for the simulations. For \( D_{m,m} \) simulations, the mass density of the CIRS cortical and trabecular bones was 1.91 g/cm\(^3\) and 1.197 g/cm\(^3\) respectively. For \( D_{w,w} \) calculations, (where the voxels are simulated as water with a density of bone), the mass density of water was defined as 1.769 g/cm\(^3\) and 1.156 g/cm\(^3\) respectively for CIRS cortical and trabecular bone respectively, in order to obtain equivalent electron densities [35].

### 2.3. SBRT spine dosimetry audit

The ACDS SBRT audit is performed on a customised CIRS Thorax phantom. The phantom consists of a plastic water body, inhale lung material and a spine composed of CIRS Trabecular Bone and CIRS

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**Table 1**

<table>
<thead>
<tr>
<th>EGSnrc Monte Carlo simulation transport settings.</th>
<th>Option</th>
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<tbody>
<tr>
<td>Global electron cut off energy (ECUT)</td>
<td>521 keV</td>
</tr>
<tr>
<td>Global photon cut off energy (PCUT)</td>
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<tr>
<td>Boundary crossing algorithm</td>
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<tr>
<td>Global SMAX</td>
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<tr>
<td>Skin depth for boundary (MFP)</td>
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<tr>
<td>Bremsstrahlung angular sampling</td>
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<tr>
<td>Photon cross sections</td>
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<tr>
<td>Bremsstrahlung cross sections</td>
<td>BH</td>
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**Fig. 1.** Dose to bone correction factor schematic: (a) shows the calibration definition of dose to water in a water phantom, (b) shows the calibration conditions of the detector in water, (c) shows the measurement conditions of the detector in bone, and (d) shows the TPS calculated dose to medium (bone).
Cortical Bone [36] (Fig. 2). Weeks prior to the audit, the phantom is mailed to a radiation oncology facility where it undergoes a CT scan, treatment planning and quality assurance according to the local SBRT protocol. ACDS staff then attend the site to perform measurements of the plan. Gafchromic EBT3 Film is used to measure dose at the centre of the target volume in the transverse plane, and two PTW 60019 micro-diamond detectors are used for point dose measurements in both the target (trabecular bone) and the spinal cord. Both detectors are calibrated for absolute dosimetry using a normalised dose to water approach (D_{w,m}) at the Australian Primary Standards Laboratory.

Included in the measured film plane are both cortical bone and trabecular bone materials. Using the modelled film correction factors for both cortical and trabecular bone, a correction factor ‘spine mask’ for both D_{w,m} and D_{w,w} was generated based on the geometry of the anthropomorphic phantom. The resolution of actual CT images of patients as well as the audit phantom leads to blurring in the voxels at the interface of the materials. This is handled in the audit process through the application of a film mask which averages each voxel by the correction factors in the surrounding 3 mm voxels. The resolution of the film mask was 72dpi, corresponding to pixel size of 0.3 × 0.3 cm². The D_{m,m} or D_{m,w} correction factor spine mask was applied to the films measured in the audit, as per the primary reporting mode of the TPS algorithm used in the plan. For analysis of ACDS SBRT audit plans, the measured film is localised to a physical position in the phantom via the facility CT scan. Spatial accuracy of the delivered plan is then assessed using metrics such as distance-to-agreement between planned and measured isodoses and gamma criteria. For this study, the primary objective is to show the difference in planned vs. measured dose due to the bony material. As such, for the purposes of this study only, the measured film and the plan were aligned for best fit to eliminate the spatial inaccuracies in the delivery. The primary metric reported for the purposes of this study is absolute local dose difference between the plan and the measurement. Dose differences were analysed in the PTW VeriSoft software v6.1. For the microDiamond point dose measurements, the modelled k_{meq} correction factors were applied to the measured data and compared to the point dose within the plan.

To date, 63 SBRT spine treatment plans have been measured in the audit. Plans were submitted from six TPS: Eclipse (Varian Medical Systems, Palo Alto, CA, USA), Monaco (Elekta AB, Stockholm, Sweden), Pinnacle (Philips Radiation Oncology Systems, Milpitas, CA, USA), RayStation (RaySearch Laboratories, Stockholm, Sweden), iPlan (Brainlab AG, Munich, Germany) and Precision (Accuray, Sunnyvale CA, USA). Table 2 lists the number of plans per algorithm that were included in the SBRT audit, and the calculation method employed. Of the audited plans, 90% were completed with 6MV or 6FFF.

Fig. 2. Transverse CT slice of the ACDS thorax phantom, with trabecular and cortical bone spine, and microdiamond measurement points in spine and spinal cord. A typical planned dose distribution in the film plane is shown.

### Table 2

<table>
<thead>
<tr>
<th>SBRT spine audit plans included in the study.</th>
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<tbody>
<tr>
<td>Calculation type</td>
</tr>
<tr>
<td>D_{w,m} Dose to medium, in medium</td>
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<tr>
<td>D_{w,w} Dose to water, in water</td>
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*CCC is dependent on specific implementation of algorithm, and in this work is investigated as both a D_{w,m} and D_{w,w}.

The Collapsed Cone Convolution (CCC) algorithm implemented in Pinnacle and RayStation TPS, and the Adaptive Convolve (AC) algorithm implemented in Pinnacle TPS are challenging to categorize as they report a mix of D_{w,m} and D_{w,w}. These algorithms partially account for material of the voxels by applying material specific mass attenuation coefficients for photon attenuation, which adjusts the water based kernel. Due to the lack of clarity in the literature [21,22,24,37,38], these algorithms have been analysed using both D_{w,m} and D_{w,w}.

As the literature recommends avoiding the stopping power/energy deposition ratio based conversion of dose to water in a medium (D_{w,m}) [17,18,22,24], we have not modelled correction factors for this methodology. Additionally, no SBRT spine plans were submitted to this study using this calculation method.

### 3. Results

#### 3.1. Material validation

Validation of the PEGS4 data for the CIRS cortical and trabecular bone materials was performed by comparing transmission measurements and TPS calculations with Monte Carlo modelled results. A C13 ionisation chamber (IBA Dosimetry, Schwarzenbruck, Germany) and a PTW microDiamond were used to measure the dose from the 6 MV Elekta Synergy 4 × 4 cm² field, at 15 cm depth in a 30 × 30 × 30 cm³ CIRS solid water phantom. Measurements were repeated with 15 × 15 × 2 cm³ slabs of CIRS cortical and trabecular bone to a maximum thickness of 8 cm placed on top of the water phantom. All measurement scenarios were replicated in the Monaco TPS, using the Monte Carlo algorithm, D_{m,m} reporting mode and a combined statistical uncertainty of 0.5% per plan. The measurement scenarios were also replicated in simple Monte Carlo models using the EGSSrc user code DOSXYZnrc. Table 3 shows the difference in transmission ratios for the water phantom vs. water phantom with 2 cm slab of cortical/trabecular bone. The transmission ratio in the EGSSrc monte carlo simulations has been compared to calculations in Monaco TPS (Monte Carlo algorithm) and measurements with CC13 and microDiamond detectors. For cortical bone, the ratio of the dose in the water and water/bone models for

### Table 3

<table>
<thead>
<tr>
<th>Measurement</th>
<th>Monte Carlo vs. Measurement</th>
<th>Monte Carlo vs. Monaco MC (D_{w,m})</th>
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<tbody>
<tr>
<td>CC13</td>
<td>Trabecular bone 0.5%</td>
<td>0.6%</td>
</tr>
<tr>
<td></td>
<td>Coritical bone −0.5%</td>
<td>−0.3%</td>
</tr>
</tbody>
</table>

The difference in the transmission dose ratios in water phantom, vs water phantom with 2 cm cortical and trabecular bone slabs.
thickness of cortical bone ranging from 2 cm to 8 cm was found to be within 1.0% of CC13 measurements and 0.7% of microDiamond measurements. For the trabecular bone, measurements were performed using a single 2 cm bone slab due to availability of equipment. The combined uncertainty in the Monte Carlo models was 0.4%.

3.2. Algorithm specific corrections

The 2D dose maps in each of the simulated Gafchromic film scenarios are shown in Fig. 3. $k_{med}$ correction factors were calculated for each voxel in the $D_{m,m}$ (Fig. 3a-d) and $D_{HDw,w}$ (Fig. 3f-i) scenario DOSXYZnrc input files as per Eqs. (2) and (4). Calculations were performed for both 6MV and 10MV beams. The resulting correction factor maps for 6MV are shown in Fig. 3e and j. For the Gafchromic Film, the final CIRS cortical and trabecular bone $k_{med}$ correction factors were averaged across the 3 mm voxels at the centre of each bone region, avoiding the voxels at the direct interface. The final correction factors for Gafchromic film are summarised in Table 4. The uncertainty in the MC correction factors only accounts for the statistical uncertainty in the MC calculations. For $D_{m,m}$ scenario, the difference between 6 MV and 10 MV was <0.2% for both cortical and trabecular bone. For the $D_{w,w}$ scenario, the difference between 6 MV and 10 MV was 1.1% and 0.9% for cortical and trabecular bone respectively.

The results of the simulated Gafchromic film and PTW 60019 microDiamond correction factors are summarised in Table 4. For the microDiamond, the correction factors were determined from the voxels in the active layer and the 1 mm tip of the detector. The difference for all modelled correction factors for the microDiamond between 6 MV and 10 MV was <0.2%.

Fig. 4a and b show the sagittal central axis profiles across the Gafchromic film $k_{med}$ correction factor maps for the $D_{m,m}$ and $D_{w,w}$ scenarios. Sharp changes in dose, and resulting correction factors, are seen at the interface regions between the cortical and trabecular bone, particularly for the $D_{m,m}$ scenario. Smoothed profiles were created to reflect the blurring of interfaces which occurs in clinical and audit plans based on CT resolution. The profiles were smoothed across 3 mm, to show the handling of the blurring in the spine mask.

3.4. Audit results – PTW 60019 microDiamond point dose

Fig. 7 shows the per algorithm results of the PTW 60019 microDiamond point dose in trabecular bone in the spine target volume. The 6MV correction factors were applied to all measurements, as there was no significant energy dependence shown in the microDiamond correction factor simulations. Algorithms with fewer than 5 audit plans have been excluded from the analysis. The mean and standard deviation for the local dose discrepancy of the uncorrected measurements was −2.4% and 3.9% respectively. Application of the algorithm specific correction factors brings the mean local dose discrepancy to 0.4%, with a standard deviation 3.7%.

4. Discussion

With the introduction of modern TPS algorithms, there has been much debate in the literature around the clinical impacts of calculations based on patient like materials. For patient media with atomic numbers close to water, calculations of $D_{m,m}$ and $D_{w,w}$ show only very minor differences. As a large proportion of clinical radiotherapy focuses on the $k_{med}$ factors listed in Table 4, according to the primary reporting mode of the algorithm used in each plan. The 6MV correction factors were applied to plans using 6MV and 6FFF, while the 10MV correction factors were applied to plans using 10MV and 10FFF. The average uncorrected and corrected local dose difference maps in the SBRT spine audit are shown per algorithm in Fig. 5 and Fig. 6. Table 5 shows the average local dose difference across the scored area of the film, encompassing both cortical and trabecular bone. Algorithms with fewer than 5 audit plans have been excluded from the analysis. The large dose difference seen in the cortical bone region in the top row is notably improved after the film mask is applied.

Fig. 6 shows the average local dose difference maps for the CCC algorithm; uncorrected, corrected for $D_{m,m}$ and corrected for $D_{w,w}$. The classification of Pinnacle CCC is more complex than other algorithms, as this algorithm reports a mix of $D_{m,m}$ and $D_{w,w}$. However following AAPM Task Group 329 [24] it can be considered $D_{m,m}$ in this context. We have included results from correcting by both dose to medium and dose to water for completeness. The dose to medium correction appears to over-correct in the cortical bone region, which is also evident in the average local dose differences in Table 5.
treatment of tumours in soft tissue regions, the impact of $D_{m,m}$ calculations will be rather insignificant. However, in patient materials with high density such as cortical bone, the differences between $D_{m,m}$ and $D_{w,w}$ calculations can be up to 10% [17–19,21]. One of the concerns with the $D_{m,m}$ reporting mode is the challenges of verifying dose in non water-like media such as bone. Traditional patient specific QA often employs the use of a homogenous phantom for verification of dose delivery. The radiation detectors used for patient specific QA are calibrated dose-to-water, which leads to large discrepancies in measured dose for plans with regions of bony material. The Monte Carlo simulations in this paper provide a method for correction of measured dose in bone like materials in an SBRT spine phantom. Whilst the bone like materials included in

### Table 4

<table>
<thead>
<tr>
<th>Calculation type</th>
<th>Algorithm</th>
<th>$k_{med}$ EBT3 Film</th>
<th>$k_{med}$ PTW 600019 microDiamond</th>
</tr>
</thead>
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<tr>
<td>$D_{m,m}$ Dose to medium, in medium</td>
<td>AcurosXB (AXB)</td>
<td>0.875 (±0.1%)</td>
<td>0.962 (±0.4%)</td>
</tr>
<tr>
<td></td>
<td>Monte Carlo (MC)</td>
<td>0.876 (±0.1%)</td>
<td>0.963 (±0.2%)</td>
</tr>
<tr>
<td></td>
<td>Collapsed Cone Convolution (CCC)</td>
<td>0.953 (±0.3%)</td>
<td>0.955 (±0.2%)</td>
</tr>
<tr>
<td></td>
<td>Adaptive Convolve (AC)</td>
<td>0.955 (±0.3%)</td>
<td>0.993 (±0.2%)</td>
</tr>
<tr>
<td>$D_{w,w}$ Dose to water, in water</td>
<td>Anisotropic Analytical Algorithm (AAA)</td>
<td>0.920 (±0.1%)</td>
<td>0.994 (±0.2%)</td>
</tr>
<tr>
<td></td>
<td>Ray Tracing (RT)</td>
<td>0.910 (±0.1%)</td>
<td>0.994 (±0.2%)</td>
</tr>
</tbody>
</table>

Fig. 4. (a) Central axis profiles in $D_{m,m} k_{med}$ correction factor map; raw results showing the sharp changes at the CIRS cortical and trabecular bone interfaces at 5 mm and 20 mm depths and smoothed profile showing the averaging applied in the spine mask. (b) Central axis profiles in $D_{w,w} k_{med}$ correction factor map; the raw and smoothed profiles show better agreement in the high density water scenario.

### Table 5

<table>
<thead>
<tr>
<th>Algorithm</th>
<th>Average local dose difference (%) Raw</th>
<th>Corrected $D_{m,m}$</th>
<th>Corrected $D_{w,w}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Eclipse AXB</td>
<td>-1.4%</td>
<td>1.2%</td>
<td></td>
</tr>
<tr>
<td>Monaco MC</td>
<td>-1.8%</td>
<td>0.7%</td>
<td></td>
</tr>
<tr>
<td>Eclipse AAA</td>
<td>-1.1%</td>
<td>0.1%</td>
<td></td>
</tr>
<tr>
<td>Collapsed Cone</td>
<td>0.6%</td>
<td>3.4%</td>
<td>1.2%</td>
</tr>
</tbody>
</table>

Fig. 5. (a)CIRS trabecular and cortical bone materials in the phantom and (b) the ‘blurred’ correction factor mask. The uncorrected and corrected average local dose difference maps (film vs. planned dose) of the SBRT spine audit for (c,d) Eclipse AXB, (e,f) Monaco MC, (g,h) Eclipse AAA.

Uncorrected and corrected average local dose difference (%) (film vs. planned dose) across scored area of film for SBRT spine audit (encompassing both cortical and trabecular bone).

treatment of tumours in soft tissue regions, the impact of $D_{m,m}$ calculations will be rather insignificant. However, in patient materials with high density such as cortical bone, the differences between $D_{m,m}$ and $D_{w,w}$ calculations can be up to 10% [17–19,21]. One of the concerns with the $D_{m,m}$ reporting mode is the challenges of verifying dose in non water like media such as bone. Traditional patient specific QA often employs the use of a homogenous phantom for verification of dose delivery. The radiation detectors used for patient specific QA are calibrated dose-to-water, which leads to large discrepancies in measured dose for plans with regions of bony material. The Monte Carlo simulations in this paper provide a method for correction of measured dose in bone like materials in an SBRT spine phantom. Whilst the bone like materials included in
this study simulate patient tissue, traceability of non-water materials is not established. According to manufacturer data, the linear attenuations of simulated bone materials in the phantom are within 1% of actual attenuation in bone [36]. This was also verified by in-house attenuation measurements compared to MC modelled results, however due to the obvious ethical and technical challenges involved, no measurements were done with actual human bone for comparison as would typically be done when commissioning synthetic/plastic water.

The MC corrections for film measurements of treatment plans using $D_{m,m}$ calculations were found to be 4.7% for trabecular bone and 12.5% for cortical bone. The corrections for algorithms using $D_{w,w}$ calculations were 1.8% for trabecular and 8.0% for cortical bone. When these corrections were applied to the film results from the audit, the large dose differences seen in the cortical bone region were significantly improved. Similar corrections were calculated for the microDiamond measuring in trabecular bone of 0.7% and 3.8% for $D_{w,w}$ and $D_{m,m}$ respectively. When the correction was applied to the audit results the overall average dose difference between planned and measured doses improved from 2.4% to 0.4%. In this study we have used a simplified microDiamond geometry in the MC simulations. For the SBRT audit, the microDiamond detector for point dose verification is a secondary measurement, with film being the primary detector. The microDiamond local point dose discrepancies show substantial uncertainty, with a standard deviation of 3.7%. This is largely due to the steep gradients seen in the SBRT spine plans. The point dose measurement is designed to offer a coarse check of the plan in real time, with the film serving as the primary detector due to the information it provides on both dosimetric and spatial accuracy of the delivery. The simple model does appear to provide a good approximation for the purposes of a dose in medium correction and shows significant improvement in the average local point dose differences measured in the audit scenario.

A $4 \times 4$ cm$^2$ field size was chosen in this study to ensure full coverage of the 2.5 cm bone cube at the centre of the modelled geometries. A $4 \times 4$ cm$^2$ field also represents typical field sizes for the SBRT spine plans included in the audit. A limitation of this study is the use of a single field in the Monte Carlo simulations, while the vast majority of the audit plans were delivered with a VMAT technique. The modelled correction factors for bony materials may also be field size dependent; however, both field size and delivery technique differences are outside the scope of the current work, and could be the subject of future investigations. Another limitation of the study was the use of a 6MV and 10MV beam for the simulations, whilst many of the audit plans were delivered with flattening filter free beams (FFF). There may be differences in the correction factors for flat beams vs. flattening filter free beams that have not been investigated. In this study we have applied the 6MV correction factors to 6FFF treatment plans, and 10MV to 10FFF plans. For the $D_{m,m}$ calculation modes, the difference between all correction factors for 6MV and 10MV was < 0.2%, within the uncertainty of the Monte Carlo simulations. For the $D_{w,w}$ calculation mode, the difference between the correction factors Gafchromic film for 6MV and 10MV was ~1.0%.
measurements, algorithm specific correction factors should be applied to account for variations in bony material for calculations based on $D_{m,M}$ and $D_{w,W}$.

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References


