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Measurement of multi-slice computed tomography dose profile with the Dose Magnifying Glass and the MOSkin radiation dosimeter

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Keywords
moskin, glass, magnifying, profile, dose, tomography, dosimeter, computed, radiation, slice, multi, measurement

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Measurement of Multi-Slice Computed Tomography Dose Profile with the Dose Magnifying Glass and the MOSkin Radiation Dosimeter

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Abstract

This study describes the application of two in-house developed dosimeters, the Dose Magnifying Glass (DMG) and the MOSkin dosimeter at the Centre for Medical Radiation Physics, University of Wollongong, Australia, for the measurement of CT dose profiles for a clinical diagnostic 16-slice MSCT scanner. Two scanner modes were used; axial mode and helical mode, and the effect of varying beam collimation and pitch was studied. With an increase in beam collimation in axial mode and an increase of CT pitch in helical mode, cumulative point dose at scanner isocentre decreased while FWHM increased. There was generally good agreement to within 3% between the acquired dose profiles obtained by the DMG and the film except at dose profile tails, where film over-responded by up to 30% due to its intrinsic depth dose dependence at low doses.

Keywords: CT dosimetry; MOSFET; DMG.
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1. Introduction

CTDI_{100}, an existing dose quality assurance measure of CT scanner output, is defined as the average dose to a cylindrical homogeneous PMMA phantom, from a single axial scan with integration limits of ±50 mm, described by Equation 1.

\[
CTDI_{100} = \frac{1}{T} \int_{z_1}^{z_2} D(z) \, dz \quad \text{(mGy)} \quad \text{(Eq. 1)}
\]

where \( z_1 = -50 \) and \( z_2 = +50 \) are the limits of integration in mm, \( D(z) \) is the single slice dose profile and \( T \) is the nominal slice thickness in mm.

CTDI_{100} has been known to severely underestimate modern, wide cone beam doses (International Atomic Energy Agency, 2011) due to considerable contribution of doses from dose tails extending beyond the conventional 100 mm CT pencil ion chamber length frequently used in clinical CT quality assurance (DCT10, RTI Electronics). Recently, several authors have also reported on the enlarged longitudinal beam width (up to 160 mm) associated with the latest cone beam CT scanners (Geleijns et al., 2009; Gomà et al., 2011; Lin and Herrnsdorf, 2010; Mori et al., 2005; Nakonechny et al., 2005). It follows that besides the use of longer phantoms, a new kind of dosimetry tool and methodology is necessary for more accurate assessment of doses associated with these new wide cone beam CT scanners.

The present work is a proof of concept study towards improved wide beam MSCT dose assessment using the Dose Magnifying Glass (DMG) calibrated against the MOSkin dosimeter. The first aim of the study was to map the longitudinal z-axis CT dose profile in the CT scanner axial mode with different beam collimations (1.25 mm, 5 mm, 20 mm) and in the helical mode with different CT pitch values (0.625, 0.875, 1.35). The second aim of the study was to demonstrate the over-ranging phenomenon associated with CT helical scans compared to CT axial scans.
2. Methodology

2.1 CT scanner and dosimeters

The CT system used was a 16-slice General Electric (GE) Discovery 670 NM/CT SPECT/CT scanner. Table 1 shows the scan parameters used for this study which were designed based on commonly used clinical imaging parameters for diagnostic CT scans.

Table 1: Scan parameters of this study with a typical clinical CT tube potential of 120 kVp tube current of 300 mAs and a total scan range of 120 mm.

<table>
<thead>
<tr>
<th>CT Scan mode</th>
<th>CT Detector configuration</th>
<th>Nominal beam collimation (mm)</th>
<th>Scan length (mm)</th>
<th>Pitch</th>
</tr>
</thead>
<tbody>
<tr>
<td>Axial</td>
<td>2 x 0.625mm</td>
<td>1.25</td>
<td>29.375</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>4 x 1.25 mm</td>
<td>5</td>
<td>30</td>
<td>-</td>
</tr>
<tr>
<td></td>
<td>8 x 2.5 mm</td>
<td>20</td>
<td>30</td>
<td>-</td>
</tr>
<tr>
<td>Helical</td>
<td>8 x 2.5 mm</td>
<td>20</td>
<td>30</td>
<td>0.625</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td></td>
<td>0.875</td>
</tr>
</tbody>
</table>

The MOSkin dosimeter, designed and developed at the Centre for Medical Radiation Physics (CMRP), University of Wollongong, Australia, has previously been investigated for applications in radiation therapy (Hardcastle et al., 2008; Kwan et al., 2009) and recently in CT (Lian et al., 2012). It was found to be tissue equivalent at depth in clinical kilovoltage beams (Lian et al., 2011) and was therefore selected as our gold standard absolute dosimeter in this work.

The DMG is a high-spatial resolution relative dosimeter developed at the CMRP. The DMG comprises an array of 128 n⁺ strips fabricated on a p-type silicon substrate of 375 µm thickness. Each n⁺ strip is a silicon diode with width 20 µm and length 2 mm, wire-bonded onto kapton. An electronic system allows simultaneous real time readout of its 128 channels. Figures 1 and 2 illustrate the topological schematic diagram and photographs of the DMG respectively. Further details on its construction, design and applications in clinical therapy can be found in (Hardcastle et al., 2012 and Wong et al. (2010, 2011))

In this study, the beam profile data acquired by the DMG was compared with the dose profile obtained with a new type of Gafchromic film, XR-QA2 film (International Specialty Products, 2011), specially developed for dosimetry in radiology.
Figure 1: Topology of DMG, not-to-scale (From Wong et al 2010).

Figure 2: Photograph of (a) the 25.6 mm silicon detector array of the Dose Magnifying Glass (DMG) (b) DMG mounted on 1 cm thick solid water with its connector at one end which can be inserted into a phantom.

2.2 Experimental setup

Figure 3 illustrates our experimental setup. The DMG was placed between seven slabs of 1 cm thick solid water blocks (RMI 457 Gammex, Middleton, WI, USA), each of 30 cm x 30 cm dimensions. In addition, the DMG was surrounded by 1 cm thick blocks of solid water. The longitudinal axis of the silicon diode array was in coincidence with the z-axis of the CT.
scanner. The DMG was connected by two long ribbon cables to a computerized DAQ system outside of the CT room for real-time data acquisition and processing.

Figure 3: Experimental setup of the CMRP DMG system. (a) Photograph showing the DMG aligned to the isocentre of the CT scanner using the laser positioning reference markers. (b) Side view of DMG placed between solid water slabs, 8 cm above and below the sensitive detector array of the DMG.

2.3 Dosimeter Calibration

With the DMG set up as shown in Figure 3, the relative sensitivity response of each channel of the DMG was obtained with the 120 kVp CT beam (HVL=0.8 mm Cu) in scout mode (stationary x-ray tube), at a nominal beam width of 20 mm over a longitudinal total z-axis scan distance of 120 mm. The DMG was manually shifted and re-positioned 6 times, in a so-called “shift-and-measure” technique from -60 mm to +60 mm along the longitudinal z-axis of the CT scan couch, first described in Wong et al. (2011). The shift-and-measure technique was applied in the calibration to create a flat calibration x-ray CT field for all 128 channels of the detector as much as possible in this study.

After equalization of the 128 channels of the 25.6 mm length DMG diode array in the CT scout scan mode, the DMG sensitivity response was found to have a 3.6% coefficient of variation (COV) in a single scan. With 3 repeated x-ray CT scans, an average COV of 7% was obtained in the overall plotted CT dose profile. We attribute this overall variation to the reproducibility of the CT beam and the uncertainties associated with DMG positioning alignment.

The MOSkin dosimeter was separately calibrated in a superficial 150 kVp x-ray beam (HVL=0.6 mm Cu) on a Gulmay D3300 superficial/orthovoltage therapy unit on the surface of a 30 cm x 30 cm x 20 cm solid water phantom. At this beam quality, the MOSkin calibration factor obtained was 0.67 mV/mGy.
Similarly, for the XR-QA2 film, small film pieces of 3 cm x 3 cm each were cut to dimensions, and calibrated on the Gulmay D3300 therapy unit at 150 kVp tube potential (HVL=0.6 mm Cu) on the surface of a 30 cm x 30 cm x 20 cm solid water phantom. Doses ranging from 0 to 200 mGy were applied and the exposed films were digitized with a scanner (Microtek ScanMaker i800, 9600 x 4800 DPI CCD, 48-bit color) at least 24 hours after x-ray irradiation. Each film piece was scanned before and after the exposure at the same position on the scanner bed to ensure readout measurement consistency. In addition, the scanner was allowed to warm up at least 10 minutes prior to its first scan.

The images obtained from film digitization were analysed with ImageJ software (National Institutes of Health, MD) and Photoshop software. For each film piece, a central region of interest (ROI) of 1 cm x 1 cm dimensions was specified and analysed. The following film scan protocol was applied: Reflective scan mode, RGB mode 48-bit color, resolution 300 dpi, 100% image scaling, with no color correction nor filters applied. Each film piece was scanned three times to minimize scanner response variations. The red channel was applied for image analysis. Finally, a calibration curve was plotted to determine the relationship between absorbed doses to optical density (i.e. degree of film darkening after irradiation).

2.4 Depth dose characterization

Figure 4 shows the comparison of the dose measured with MOSkin, Gafchromic XR-QA2 film and Marcus parallel plate ionization chamber (IC) at different depths in a 30 x 30 cm square solid water phantom measured in a 150 kVp (0.6 mm Cu HVL) superficial therapy beam on a Gulmay D3300 superficial therapy machine. The MOSkin dosimeter was reproducible to within 3% of ion chamber measurements for doses at depths larger than 80 mm in solid water. At depths larger than 80 mm, the XR-QA2 film was found to over-respond by up to 30% compared to the IC and the MOSkin dosimeter. For depths less than 80 mm, the average agreement between MOSkin and film was 4%. For depths close to and at the surface with steep dose gradients (dose buildup), the larger disparity between MOSkin and film readouts was attributed to the different Water Equivalent Depth (WED) of the dosimeters.
3. Results and Discussion

3.1 Axial scan mode

For the axial scan mode (Figures 5 to 7), the effect of increasing beam collimation (1.25 mm, 5 mm, 20 mm) was studied. It was seen from the reconstructed dose profiles that with increasing beam collimation, FWHM increases while cumulative point dose measured at beam isocentre decreases. This may be explained by the spreading of the beam resulting in different scattering percentage contribution to cumulative central point dose.

MOSkin point dose and the film dose profiles generally agreed to within 3%, and the dose profiles obtained with the film and DMG generally agreed except at the dose tails where the film over-responded by up to 30%. From the depth dose characterization results presented in Figure 4, the observed 30% film over-response at the dose tails is to be expected.
Figure 5: Axial scan mode with 1.25 mm nominal beam collimation. FWHM (DMG) = 31.2 mm, FWHM (film) = 37.3 mm, % difference in FWHM (film-DMG) = 16%. MOSkin point dose measurement at beam isocentre= 55.1 mGy

Figure 6: Axial scan mode with 5 mm beam collimation. FWHM (DMG) = 44.0 mm, FWHM (film) = 55.9 mm, % difference in FWHM (film-DMG) = 21%. MOSkin point dose measurement at beam isocentre= 36.2 mGy
Figure 7: Axial scan mode with 20 mm beam collimation. FWHM (DMG)= 43.0 mm, FWHM (film)= 47.8 mm, % difference in FWHM (film-DMG)=10.0%. MOSkin point dose measurement at beam isocentre= 30.4 mGy

3.2 Helical scan mode
Figures 8 to 10 show the effect on dose profile with increasing CT pitch 0.625, 0.875 and 1.35. With an increase in CT pitch, the cumulative point dose measured by the MOSkin at beam isocentre decreased. This is to be expected since couch movement per rotation increases with collimation kept constant. Isocentre MOSkin point dose measurement agreed with film to within 3% for all helical scans.

The measured FWHM for the helical scan (Figure 8) was 1.5 times wider than that obtained for the axial scan (Figure 7) for the same 20 mm beam collimation. This result verifies the application of the DMG for the measurement of the known over-ranging effect in helical beams (Van der Molen and Geleijns, 2007).
Figure 8: Helical scan mode at 20 mm nominal beam collimation, pitch 0.625. FWHM (DMG)= 61.4 mm, FWHM (film)= 70.7 mm, % difference in FWHM (film-DMG)=13.0%. MOSkin point dose measurement at beam isocentre= 55.1 mGy.

Figure 9: Helical scan mode at 20 mm nominal beam collimation, pitch 0.875. FWHM (DMG)= 70.0 mm, FWHM (film)= 74.6 mm, % difference in FWHM (film-DMG)=6.0%. MOSkin point dose measurement at beam isocentre= 34.8 mGy.
A major limitation in this study was the use of square solid water blocks surrounding the DMG instead of the conventional cylindrical 16 cm (head) or 32 cm (body)-diameter PMMA phantoms during CT dose scanning since square phantoms do not allow the correct assessment of the CTDI$_{100}$. Moreover the current packaging of the DMG results in an asymmetrical response in comparison with that of a cylindrical phantom with the conventional CT pencil ion chamber placed in the central axis. However, the major objective of this study was a demonstration of a proof of concept, that the DMG calibrated by the MOSkin was feasible for MSCT dose profile acquisition. Verification of scanner CTDI$_{100}$ was beyond the scope of the present study.

It follows that future verification studies with the DMG should initially be focused on fabricating a custom cylindrical PMMA phantom that incorporates the DMG. This will then allow the direct comparison of CTDI$_{100}$ measurements acquired with the DMG to that acquired by the conventional CT pencil-ion chamber for the further verification of narrow CT beam widths. Use of customised phantoms incorporating the DMG will also facilitate the future measurement of CTDI$_{\infty}$ for wide cone beam widths.
4. Conclusion

This study has shown the practical application of the DMG calibrated using the MOS\textit{skin} dosimeter, for the measurement of diagnostic x-ray CT dose profiles in a typical 16-slice diagnostic x-ray MSCT scanner. The beam profile in the CT axial and helical scan modes were successfully acquired with the DMG. There was generally good agreement between the acquired dose profiles obtained by the DMG and the film to within 3\% except at the dose profile tails, where film over-responded by up to 30\% due to the inherent depth dose response of film at low doses. The over-ranging phenomenon typically associated with helical scans compared to axial scans was also verified by the DMG and MOS\textit{skin} measurements.

Based on the results obtained in this study, we recommend future application of both the MOS\textit{skin} dosimeter and DMG for the dose profile assessment of longer beam widths associated with next generation wide cone beam CT scanners in order to overcome present limitations associated with the existing CT dose metric, CTDI\textsubscript{100}. Future work will be focused on fabricating a custom cylindrical phantom that incorporates the DMG.

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