Implementation and evaluation of a transit dosimetry system for treatment verification

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Abstract
Purpose: To evaluate a formalism for transit dosimetry using a phantom study and prospectively evaluate the protocol on a patient population undergoing 3D conformal radiotherapy.

Methods: Amorphous silicon EPIDs were calibrated for dose and used to acquire images of delivered fields. The measured EPID dose map was back-projected using the planning CT images to calculate dose at pre-specified points within the patient using commercially available software, EPIgray (DOSIsoft, France). This software compared computed back-projected dose with treatment planning system dose.

A series of tests were performed on solid water phantoms (linearity, field size effects, off-axis effects).

37 patients were enrolled in the prospective study.

Results: The EPID dose response was stable and linear with dose. For all tested field sizes the agreement was good between EPID-derived and treatment planning system dose in the central axis, with performance stability up to a measured depth of 18 cm (agreement within $\pm 0.5\%$ at 10 cm depth on the central axis and within $\pm 1.4\%$ at 2 cm off-axis). 126 transit images were analysed of 37 3D-conformal patients.

Patient results demonstrated the potential of EPIgray with 91% of all delivered fields achieving the initial set tolerance level of $D_D$ of 0 $\pm$ 5-cGy or $\%D_D$ of 0 $\pm$ 5%.

Conclusions: The in vivo dose verification method was simple to implement, with very few commissioning measurements needed. The system required no extra dose to the patient, and importantly was able to detect patient position errors that impacted on dose delivery in two of cases.

Introduction

It is now recommended by various national and international organisations that in vivo dosimetry monitoring should be undertaken by each radiotherapy centre [1–3]. In vivo dose measurements when compared to planned doses can spot errors in patient set-up, data transcription, machine fault or anatomical changes which may lead to over- or under-dosage of the tumour volume and unplanned toxicity. Point detectors such as diodes and thermoluminescent dosimeters are commonly used for these measurements. However, point detectors are sensitive to positioning errors particularly in highly modulated fields [4]. Such detectors are placed on external contours of patients, but it is known that external displacements can differ from internal tumour displacements up to approximately 2 cm [5,6].

Within the radiotherapy physics community EPID dosimetry is widely seen to have the potential to become an accurate and efficient means of large-scale patient specific dose verification [7–10]. A recent paper reported that between 2005 and 2009 the treatment plans of 4337 patients were verified using in vivo EPID dosimetry, among which 17 serious errors were detected [11]. Of these, 9 would have been missed if no treatment verification had been performed, thereby highlighting the importance of the method.

Current EPID technology for transit dosimetry is based upon passive, amorphous silicon (a-Si) flat panel imagers. Previous approaches have considered CCD camera based systems and liquid filled ionisation chamber matrixes. a-Si has proven popular due to its superior image quality and dose characteristics (linearity, uniformity, dose-rate dependence, field size dependence, relative dosimetry) [12–14].

Although of great use, EPID in vivo verification technology is still being developed and is not routinely available. A number of departments have implemented in vivo EPID dosimetry using
in–house solutions [15–17]. The availability of a new commercial software EPIgray (DOISsoft, France) has enabled us to use EPID devices as a portal dosimeter for in vivo dose verification. In this work the algorithm was implemented to calculate the point dose within any pre-specified location(s) within the patient, based on a recent publication which presents a method for back-projection of measured EPID fluence map to the plane of the patient [18]. A series of phantom measurements were undertaken to verify the algorithm’s performance under a range of beam conditions. A review board approved study was undertaken at the Royal Berkshire NHS Foundation Trust to prospectively evaluate the algorithm’s performance on a patient population, giving important test conditions that phantom studies cannot (namely anatomical changes including weight loss/gain, organ motion, tumour size changes etc.). This study was used to gain insight into clinical issues that come into play upon clinical implementation of EPID-based in vivo dose verification.

Materials and methods

Commissioning EPIgray

EPIgray version 2.0.3 software was used along with Elekta Synergy® and Precise linear accelerators running Integrity (Elekta, Crawley, UK) equipped with portal imaging system iViewGT a-Si panels (PerkinElmer, USA), and motorised 60° physical wedge. The sensitive layer consists of 1024 × 1024 pixels with a pitch of 400 μm, resulting in an active area of 409.6 × 409.6 mm² [19]. All measurements were performed with a nominal dose rate of 400 MU/min. Three linear accelerators were used for the patient measurements: the machines were matched to within Elekta specifications; for photons this is 1% for depth doses and output factors and 2% for profiles, although for most of the tested clinical fields upon commissioning, the match was found to be better than this. The treatment planning system (TPS) was Eclipse version 10.0 (Varian MS, Palo Alto, USA) with Analytic Anisotropic Algorithm (AAA). IMPAC/Mosaiq record and verify system version 2.41 (Elekta, Crawley, UK) was used to transfer plan parameters to the linear accelerator control system. Image datasets of patients and phantoms used were acquired with a GE LightSpeed 16 CT Scanner.

Prior to EPIgray commissioning the EPIDs were prepared for dosimetric measurement, according to manufacturer guidelines to ensure that all pixels have a similar response to a given irradiation and mechanical accuracy within 1 mm. The dosimetric properties of iViewGT EPIDs have been extensively investigated elsewhere [20].

The commissioning of EPIgray required configuration data for the CT, linear accelerator and EPID device. The HU to electron luminance calibration curve of the CT scanners was required. In order to model the linear accelerator the following were incorporated into the EPIgray beams library for each energy: beam profiles, per centage depth dose and quality index of open and wedged fields. A 10 × 10 cm² calibration image was taken for each energy with zero phantom thickness (through the treatment couch) for 100 MU to obtain a calibration factor (CF) that converted EPID signal to dose in water. A diagonal dose profile was input into the beams library to perform an equalisation correction for the EPID image, and linearity correction factors calculated to correct the non-linearity of the EPID with MU variation. EPID measurements were performed to acquire linearity factors at 10–500 MU, 10 × 10 cm² field, zero phantom thickness and calculated as:

\[
\text{Correction}_{\text{MU}} = \frac{\text{image value}_{\text{MU}}}{\text{image value}_{100 \text{ MU}}} \times \frac{100 \text{ MU}}{\text{MU}}
\]

(1)

Couch transmission correction was commissioned per manufacturer guidelines which involved (i) measurement of an open field at 0 gantry angle through 20 cm thick solid water, (ii) measurement of the same field through the same phantom thickness at 90 gantry angle. The latter measurement provided a couch transmission value through the entire width of the couch. An average couch transmission factor derived by the manufacturer from data from (i) and (ii) was applied to all 3D conformal fields as per the EPIgray algorithm.

Finally, finite Tissue Maximum Ratio (fTMR) measurements were acquired and input into the EPIgray library, as guided by the manufacturer. fTMR is the ratio between two doses measured in a phantom at \( d_{\text{max}} \). The numerator is the dose measured in the presence of an absorber of thickness ‘\( t \)’ and the denominator is the dose measured in the same conditions without an absorber [18]. fTMR is measured in the presence of an absorber of finite thickness as opposed to TMR where the phantom dimensions are infinite. Commissioning fTMR measurements consisted of ion chamber dose measurement at source to chamber distance of 160 cm (at EPID position) within a water tank at depth \( d_{\text{max}} \) for each energy. fTMR measurements were performed for three overlying absorber solid water set-ups: with SAD of 100 cm to (i) top, (ii) centre and (iii) bottom of the solid water. Commissioning measurements were over five square field sizes (2 × 2 cm² to 25 × 25 cm²), each for six solid water thicknesses (0–40 cm) across the three aforementioned solid water SDDs. In addition EPID images were acquired at each solid water/field size/energy configuration for set up (ii).

For all EPID images a dicon image and log file were needed; the latter provided the pixel scaling factor (PSF) for each image; this value held the dosimetric information of the pixel values, as iViewGT performed a greyscale normalisation to ensure that an image acquired with 5 MU has approximately the same grey level as a 500 MU image. Un-normalised pixel values \( s(x,y) \) were obtained from normalised values \( s’(x,y) \) as in Eq. (2):

\[
s(x,y) = \frac{(2^{16} - 1) - s’(x,y)}{\text{PSF}}
\]

(2)

EPIgray dose calculation process

The calibration factor CF measured during commissioning converted EPID dose to dose in water at \( d_{\text{max}} \). Dividing by the appropriate finite tissue maximum ratio (fTMR) removed the effect of the overlying patient. Within the software fTMR values were expressed in a look up table of coefficients values listed for patient thickness (5–40 cm), field size defined at the EPID (3.2 × 3.2 cm to 40 × 40 cm). An inverse square law correction was then used to recover dose at the position of the pre-specified dose calculation point P (at depth of maximum dose \( d_{\text{max}} \)) from dose at the EPID (SDD = 160 cm). A TMR correction shifted the reconstruction point from \( d_{\text{max}} \) to point P. The fluence matrices for TMR and fTMR were calculated by individually computing the primary and scatter component. The primary component was yielded by attenuation correction of the fluence matrix (modelled using PDD and fluence profile) using attenuation coefficients derived from the quality indices provided during commissioning. The software contained a look up table of computed scatter factors as a function of field size of subsector and depth of point P. Peak scatter factor (PSF) values set within the software data tables were used to calculate the overall scatter factors; PSF describes dose to scatter only (defined as: dose in tissue at \( d_{\text{max}} \)/dose at \( d_{\text{max}} \) due to primary radiation only). PSF values for our centre were provided by the manufacturer and obtained from the quality indices of each beam energy provided.
during commissioning. The algorithm divided each field into subsectors, and the scatter table searched using depth and field size for each subsector. Scatter values of each subsector were then summed.

CT images were acquired of the phantoms and patients and exported to the TPS in which a dose distribution was calculated over the set MU and field configurations. Reference dose calculation points were created within the TPS; it is to these predefined points that EPIgray reconstructs the dose and compares to TPS calculated dose. For phantom studies, a number of reference points were used to build up beam profiles and depth doses for EPIgray-TPS comparison.

The CT images, structures, RTPlan file and dose matrix were exported from the TPS to the EPIgray console. Validated treatment plans were exported to the record and verify system (Mosaïq) to enable treatment geometries to be reproduced and delivered by the accelerator. Treatment fields were delivered and MV portal images were acquired during treatment in single shot mode for 3D conformal fields. The appropriate iView image dataset was queried by the EPIgray software, and dose to the pre-specified reference point was calculated using the EPIgray software algorithm [21]. EPIgray automatically calculated the percentage difference between EPID-derived dose ($D_{\text{EPI}}$) and TPS-calculated dose ($D_{\text{TPS}}$), ($\%\Delta_D$, Eq. (2)) and used this value to either pass or fail the measurement according to thresholds preset by the user. The software had no function to automate and set thresholds on absolute dose difference ($\Delta_D = D_{\text{EPI}} - D_{\text{TPS}}$), and so $\%\Delta_D$ values were manually calculated in this study.

$\%\Delta_D = \frac{\left( D_{\text{EPI}} - D_{\text{TPS}} \right)}{D_{\text{TPS}}} \times 100\%$ (3)

Phantom measurements

The following measurements were undertaken using a 20 cm thick solid water phantom (25 × 25 cm) set at 90 cm SSD at 6 MV and 10 MV unless otherwise specified. A number of isocentric and off-axis reference points were created within the TPS, and $D_{\text{EPI}}$ was compared to $D_{\text{TPS}}$ for each reference point.

The reproducibility of the EPID devices were measured across ten repeat deliveries of a 10 × 10 cm$^2$ field delivering 100 MU across different days over a one month period at 6 MV and 10 MV. The linearity of dose response was tested: over 5–200 MU, field size 10 × 10 cm, at the isocentre. Field size dependence was tested for three open field sizes (5 × 5, 10 × 10, 15 × 15 cm) each exposed to 100 MU. Three dose reconstruction points were defined in the TPS: (i) 5 cm depth on central axis (cax), 10 cm depth on the central axis, 5 cm depth and 2 cm off axis. Off-axis dose calculation was tested: A 20 × 20 cm field was delivered (100 MU) and 21 dose reconstruction points were created (in 1 mm increments near field edge and 10–15 mm increments in field centre) across the beam profile at two different depths (5 cm and 10 cm deep). The effect of depth of reconstruction point $P$ was investigated: a 10 × 10 cm field (100 MU) was delivered at a range of SSDs (80, 90 and 100 cm SSD). Dose reconstruction points were created on the central axis at depths 1–18 cm in 1 cm increments. Dose calculation of wedged fields was measured: A 20 × 20 cm fully wedged field (100 MU) was delivered and dose reconstruction points were created in 10 mm increments across the beam profile at depth 5 cm to assess beam profile modelling, and at depths 1–18 cm on the central axis to assay depth effects.

Additional tests were performed to assess the sensitivity of EPIgray to detection of errors in (i) SSD, and (ii) patient thickness. (i) A 10 × 10 cm field (100 MU) was planned at 90 cm SSD, 20 cm thick solid water. The field was delivered with induced errors in SSD (80–100 cm SSD in 5 cm increments), (ii) The previous plan was delivered to (a) 10 cm, (b) 20 cm and (c) 30 cm thick solid water (matching the plan), and errors in patient thickness induced by varying the water thickness up to ±4 cm (keeping the SSD constant at 85 cm for 30 cm thickness, 90 cm for 20 cm thickness, and 95 cm for 10 cm thickness). The ability of EPIgray to detect both errors (i) and (ii) was evaluated through inspecting automatically calculated $\%\Delta_D$.

EPIgray performance with heterogeneity was investigated using a phantom designed by the IPSM Radiotherapy Topic Group, made of epoxy resin water equivalent material with an 8 cm diameter cylindrical lung equivalent insert of electron density 0.25 compared with water [22]. A 10 × 10 cm$^2$ field was delivered at 6 MV and 10 MV with the central axis centred on the lung insert. The central axis of the beam traversed 1.7 cm solid water, 8 cm lung insert and finally 14.3 cm solid water. Dose was reconstructed along the central axis at (i) the centre of the lung insert, and (ii) at an additional 5 cm depth within the solid water portion proximal to the lung insert.

Patient study

37 patients (2 abdominal, 10 brain, 5 head and neck, 4 lung, 3 pelvis, 10 rectal and 3 sarcoma patients) were included in this study. 119 first fraction iView images and additional 7 repeats were acquired and processed.

All patients were treated supine with the exception of rectal patients who were prone. Brain and head and neck patients were immobilised using a thermoplastic shell. Sarcoma patients were immobilised using vacuum bags. Lung patients were immobilised using wingboards. Each patient setup was verified and corrected using CBCT except for rectal patients who were MV imaged for setup. All pelvis patients were treated with a full bladder apart from the bladder patient who had an empty bladder.

The dose reconstruction point was placed at the isocentre for all plans: this was to mirror previously used diode positions and to profit from the better performance of EPIgray on the central axis. We used a preliminary action level of 0 ± 5% and 0 ± 5 cGy for % $\Delta_D$ and $\Delta_D$, respectively, based upon results from a different group [15].

Results

Phantom studies

EPIgray dose calculation was found to have a day to day reproducibility of 0.5% and 0.6% for 6 MV and 10 MV, respectively (quoted at the 1-sigma level), with mean ± 1 SD % $\Delta_D$ values of 10 identical open fields of −0.23 ± 0.48% and −0.90 ± 0.64%. EPIgray dose calculation was found to agree well with TPS dose calculation on the central axis over a range of MU (5–200 MU) for both 6 MV and 10 MV (Fig. 1) with a maximum % $\Delta_D$ of −1.9%.

Field size was found to have little effect on % $\Delta_D$ for dose reconstruction points on the central axis (P1 and P2 within Table 1) for the tested field sizes (5 × 5–15 × 15 cm$^2$). Smaller field sizes show greater difference between $D_{\text{EPI}}$ and $D_{\text{TPS}}$ for off-axis dose calculation points, as the dose point lies closer to the field edge; dose calculated to a point 2 cm off-axis showed a % $\Delta_D$ of +1.9% and +4.3% at 5 × 5 cm$^2$ field size, 2.4 and 3 times greater than the worse performing on-axis dose calculation for the larger fields at 6 and 10 MV, respectively.

Table 2 presents a series of measurements which demonstrate that EPIgray performed well near the central axis (% $\Delta_D$ values within −0.5% on the central axis and within −1.1% at 2 cm off-axis measured at 10 cm depth). Off-axis dose points showed greater values of % $\Delta_D$ up to −6.4% at 8 cm off-axis. Agreement between $D_{\text{EPI}}$ and $D_{\text{TPS}}$ worsened at shallower depths: % $\Delta_D$
increased by approximately 0.4% when reconstructing dose to a point 5 cm deep compared to 10 cm deep. Points reconstructed at $d_{\text{max}}$ demonstrated the greatest values of $\%\Delta_D$.

Fig. 2 shows good agreement between $D_{\text{EPID}}$ and $D_{\text{TPS}}$ up to a measured depth of 18 cm at each tested SSD for both 6 and 10 MV; all depths beyond the build-up region displayed $\%\Delta_D$ within $-1.4\%$, and within $-2.2\%$ for shallower regions less than 2 cm deep. EPIgray overestimated dose at depth of 100 cm SSD, 10 MV by on average 0.3%; this may be attributed to the reported over-sensitisation of EPIDs to lower energy scattered x-rays resulting from the shorter distance between the phantom and the EPID at closer agreement within the solid water portion of the phantom: dose discrepancies $\%\Delta_D$ of $-2.1\%$ ($-1.6\%$) and $-2.2\%$ ($-1.6\%$) were found within the solid water region 5 cm proximal to the lung insert for 6 MV and 10 MV, respectively. Fig. 4 demonstrates a marked difference between EPIgray and TPS reconstructed doses at the heterogeneous boundaries between the lung insert and solid water. All $\%\Delta_D$ values were within $+5.1\%$.

**Sensitivity to errors**

A range of errors in phantom thickness were induced from planned phantom thicknesses of 10 cm, 20 cm and 30 cm. EPIgray was sensitive to patient separation changes: a reduction in patient separation by 2 cm resulted in $\%\Delta_D$ values of 9.6% (7.6%), 8.5% (6.5%) and 6.1% (5.1%) for a planned patient thickness of 10 cm, 20 cm and 30 cm, respectively at 6 MV (10 MV). Likewise an increase in patient separation by 2 cm resulted in $\%\Delta_D$ values of $-9.3\%$ ($-6.7\%$), $-8.7\%$ ($-6.9\%$) and $-9.6\%$ ($-7.1\%$). Errors of 1 cm water equivalent path length resulted in a $\%\Delta_D$ value within tolerance ($\pm5\%$) for each of the three planned thicknesses and so would not have been detected. Results for each phantom thickness are presented in Fig. 5(a) and (b).

On delivering a 10 x 10 cm$^2$ field to a 20 cm thick solid water phantom over a range of incorrect SSDs different from the planned SSD of 90 cm, it was found that a ~10% error in SSD equated to a difference of ~2% in $\%\Delta_D$ (Fig. 5(c)). EPIgray, and transit dosimetry in general, is not sensitive to errors in SSD, and an additional SSD check should be in place.

**Patient study**

126 images were acquired and analysed across the recruited 37 patients. Table 3 lists the patient demographics. Seven fields could not be measured with EPIgray due to a large floor twist obstructing EPID panel extension (superior anterior oblique fields for seven brain patients). Diode measurements were made of these fields.

The EPID verification model agreed well with TPS dose for 3D conformal fields (Fig. 6); a mean of $-0.9\%$ lower than $D_{\text{TPS}}$ for all fields. Values of $\%\Delta_D$ and $\Delta_D$ for all fields measured for each ($\Delta_D$) of $+3.2\%$ (3.3 cGy) and $+4.2\%$ (4.2 cGy) were found within the centre of the lung insert for 6 MV and 10 MV, respectively. Closer agreement was found within the solid water portion of the phantom: dose discrepancies $\%\Delta_D$ of $-2.1\%$ ($-1.6\%$) and $-2.2\%$ ($-1.6\%$) were found within the solid water region 5 cm proximal to the lung insert for 6 MV and 10 MV, respectively.

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**Table 1**

$\%\Delta_D$ of a range of field sizes (5 x 5–15 x 15 cm$^2$) delivered to a 20 cm thick solid water phantom set at 90 cm SSD, 100 MU at 6 MV and 10 MV. Dose was calculated at three different reference points: P1 (cax, 5 cm depth), P2 (cax, 10 cm depth), P3 (2 cm off axis, 5 cm depth).

<table>
<thead>
<tr>
<th>Field Size</th>
<th>P1 (%)</th>
<th>P2 (%)</th>
<th>P3 (%)</th>
<th>P1 (%)</th>
<th>P2 (%)</th>
<th>P3 (%)</th>
<th>P1 (%)</th>
<th>P2 (%)</th>
<th>P3 (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>5 x 5 cm$^2$</td>
<td>6 MV</td>
<td>-0.1</td>
<td>0.0</td>
<td>1.9</td>
<td>0.0</td>
<td>-0.1</td>
<td>-0.5</td>
<td>0.0</td>
<td>-0.6</td>
</tr>
<tr>
<td></td>
<td>10 MV</td>
<td>-0.4</td>
<td>-0.5</td>
<td>4.3</td>
<td>-0.5</td>
<td>-0.6</td>
<td>-1.4</td>
<td>0.2</td>
<td>0.3</td>
</tr>
<tr>
<td>10 x 10 cm$^2$</td>
<td>6 MV</td>
<td>-0.1</td>
<td>0.0</td>
<td>1.9</td>
<td>0.0</td>
<td>-0.1</td>
<td>-0.5</td>
<td>0.0</td>
<td>-0.6</td>
</tr>
<tr>
<td></td>
<td>10 MV</td>
<td>-0.4</td>
<td>-0.5</td>
<td>4.3</td>
<td>-0.5</td>
<td>-0.6</td>
<td>-1.4</td>
<td>0.2</td>
<td>0.3</td>
</tr>
<tr>
<td>15 x 15 cm$^2$</td>
<td>6 MV</td>
<td>-0.1</td>
<td>0.0</td>
<td>1.9</td>
<td>0.0</td>
<td>-0.1</td>
<td>-0.5</td>
<td>0.0</td>
<td>-0.6</td>
</tr>
<tr>
<td></td>
<td>10 MV</td>
<td>-0.4</td>
<td>-0.5</td>
<td>4.3</td>
<td>-0.5</td>
<td>-0.6</td>
<td>-1.4</td>
<td>0.2</td>
<td>0.3</td>
</tr>
</tbody>
</table>

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**Table 2**

$\%\Delta_D$ of reference points calculated across a 20 x 20 cm$^2$ field delivered to 20 cm thick solid water (100 MU). Dose was reconstructed across the profiles of depth 10 cm, 5 cm and at $d_{\text{max}}$ for 6 MV and 10 MV.

<table>
<thead>
<tr>
<th>Depth</th>
<th>6 MV (%)</th>
<th>10 MV (%)</th>
<th>6 MV (%)</th>
<th>10 MV (%)</th>
<th>6 MV (%)</th>
<th>10 MV (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>P1</td>
<td>-0.5</td>
<td>-0.3</td>
<td>-0.9</td>
<td>-0.5</td>
<td>-1.4</td>
<td>-0.7</td>
</tr>
<tr>
<td>P2</td>
<td>-1.0</td>
<td>-1.1</td>
<td>-1.4</td>
<td>-1.2</td>
<td>-1.8</td>
<td>-1.2</td>
</tr>
<tr>
<td>P3</td>
<td>-2.4</td>
<td>-3.0</td>
<td>-3.0</td>
<td>-3.3</td>
<td>-3.5</td>
<td>-3.4</td>
</tr>
<tr>
<td>1 cm from field edge</td>
<td>-16.5</td>
<td>-8.9</td>
<td>+17.8</td>
<td>-11.0</td>
<td>-13.7</td>
<td>-8.2</td>
</tr>
</tbody>
</table>

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treatment site are presented in Table 4. Measurements with imager problems were excluded from analysis. The distribution of percentage (%\(D\)) and absolute (\(D\)) dose difference for all 126 fields is presented in Fig. 7: \(D\) displayed a tighter distribution than %\(D\). The mean difference between EPID and TPS doses was found to depend on treatment site: ranging from lung which displayed a mean 3.0% overestimation of dose compared to TPS dose, to sarcoma which was on average 3.9% less than TPS dose. Head and neck patients displayed the greatest standard deviation in %\(D\) followed by lung patients (SD of 6.6% and 5.4%, respectively) whereas brain and pelvic patients displayed a lower standard deviation of 2.6% and 2.5%, respectively. Results for each treatment site are displayed in Table 4.

Wedge fields delivered to patients performed well: all wedge fields were analysed together and displayed an average %\(D\) value of \(-0.5 \pm 5.4\%\) compared to \(-1.7 \pm 4.7\%\) for all open fields. Upon excluding patients with high levels of heterogeneity (lung and head and neck fields) the %\(D\) values of wedge and open fields came to a closer agreement (\(-2.1 \pm 3.9\%\) and \(-2.3 \pm 4.6\%,\) respectively).

7 image repeats were acquired and processed due to the following reasons for initial failure: (i) one left bicep which appeared to have a small amount of rotation of the arm upon inspecting iView versus digitally reconstructed radiograph verification images; (ii) one axilla sarcoma which failed due to the presence of a couch bar attenuating the exit beam; (iii) one rectum failed due to patient position; (iv) three due to imager failure; and (v) one larynx field failed also on repeat due to a large amount of air in the beam path and the failed field was re-measured with a diode and passed. In addition patient 18 was re-planned due to tumour shrinkage. For this patient two out of three fields failed on fraction 1: %\(D\) of 10.9% (9.6 cGy) and 10.1% (8.1 cGy) respectively; due to the irregular tumour shape the isocentre was out of tumour and not placed in soft tissue. Throughout the course of treatment the

![Figure 2](image-url) (a) EPIgray and TPS reconstructed doses over a range of depths along the cax: 10 × 10 cm² field (100 MU, 6 MV and 10 MV) delivered to a 20 cm thick solid water phantom at 90 cm SSD, (b) %\(D\) of reconstructed doses at a range of depths measured at 80 cm, 90 cm and 100 cm SSD.

![Figure 3](image-url) (a) EPIgray and TPS off axis dose reconstruction for 20 × 20 cm field (100 MU, 6 MV and 10 MV) delivered to a 20 cm thick solid water phantom at 90 cm SSD, (b) %\(D\) of off axis dose reconstruction for wedge fields, (c) EPIgray and TPS reconstructed doses over a range of depths along the cax: 20 × 20 cm² fully wedged field (100 MU, 6 MV and 10 MV) delivered to a 20 cm thick solid water phantom at 90 cm SSD, (d) %\(D\) of reconstructed doses at a range of depths for the wedge field.

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tumour shrank and became more regular which enabled a simpler plan with isocentre placed within the tumour soft tissue. All three fields passed on the first verification of the replan: $D_D$ of 5.6% (3.3 cGy), 5.0% (4.9 cGy) and 1.2% (1.5 cGy).

**Discussion**

A series of phantom measurements demonstrated that EPIgray performed well near the central axis ($\Delta_D$ values within $\pm 0.5\%$ at 10 cm deep), with performance stability up to a measured depth of 18 cm excluding the build-up region ($\%\Delta_D$ values within $\pm 1.4\%$). Doses calculated on the central axis were found to be unaffected by field size (across measured field sizes $5 \times 5$ and $10 \times 10$ cm).

### Table 3

Demographic information for the 37 patients enrolled on the study.

<table>
<thead>
<tr>
<th>Patient number</th>
<th>Anatomic site</th>
<th>Fields per patient (energy)</th>
<th>No. open (wedged) fields</th>
</tr>
</thead>
<tbody>
<tr>
<td>1</td>
<td>Abdomen</td>
<td>3 (6 MV)</td>
<td>0 (3)</td>
</tr>
<tr>
<td>2</td>
<td>Pancreas</td>
<td>3 (10 MV)</td>
<td>0 (3)</td>
</tr>
<tr>
<td>3–12</td>
<td>Brain</td>
<td>2–5 (6 MV)</td>
<td>0 (25)</td>
</tr>
<tr>
<td>13–16</td>
<td>Larynx</td>
<td>2–3 (6 MV)</td>
<td>1 (10)</td>
</tr>
<tr>
<td>17</td>
<td>Oesophagus</td>
<td>4 (10 MV)</td>
<td>1 (3)</td>
</tr>
<tr>
<td>18a</td>
<td>Right lung</td>
<td>3 (10 MV)</td>
<td>0 (3)</td>
</tr>
<tr>
<td>18b</td>
<td>Right lung re-plan</td>
<td>3 (10 MV)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>19–21</td>
<td>Left lung</td>
<td>3–4 (10 MV)</td>
<td>0 (10)</td>
</tr>
<tr>
<td>22</td>
<td>Bladder</td>
<td>3 (10 MV)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>23–24</td>
<td>Endometrium</td>
<td>4 (10 MV)</td>
<td>3 (5)</td>
</tr>
<tr>
<td>25–34</td>
<td>Rectum</td>
<td>3–4 (6 and 10 MV)</td>
<td>11 (26)</td>
</tr>
<tr>
<td>35</td>
<td>Sarcoma Lt bicep</td>
<td>3 (6 MV)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>36</td>
<td>Sarcoma Right flank</td>
<td>3 (6 and 10 MV)</td>
<td>1 (2)</td>
</tr>
<tr>
<td>37</td>
<td>Sarcoma Axilla</td>
<td>3 (6 and 10 MV)</td>
<td>3 (0)</td>
</tr>
</tbody>
</table>

![Figure 4](image-url)  
**Figure 4.** EPIgray and TPS dose reconstruction at depth within the IPSM phantom for $10 \times 10$ cm open beam, 90 cm SSD, measured at 6 MV and 10 MV along the central axis; (b) $\%\Delta_D$ of dose reconstruction at depth in the IPSM phantom. The central rectangular region corresponds to the low density lung insert region.

![Figure 5](image-url)  
**Figure 5.** The effect of error in phantom thickness on reconstructed dose presented as percentage difference between EPIgray and TPS calculated dose $\%\Delta_D$ for three phantom thickness (10 cm, 20 cm and 30 cm) at (a) 6 MV and (b) 10 MV. Phantom thickness error = (treated thickness – planned thickness)/planned thickness; (c) effect of error in SSD on reconstructed dose presented as percentage difference between EPIgray and TPS calculated dose $\%\Delta_D$. SSD error = (treated SSD – intended SSD)/intended SSD, where the intended SSD was 90 cm.

![Figure 6](image-url)  
**Figure 6.** $D_{EPI}$ and $D_{TPS}$ reconstructed doses of each measured patient field for 126 3D conformal beams. Dashed line: $D_{EPI} = D_{TPS}$.

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20 \times 20 \text{ cm}^2$) and good performance observed across a range of MU (stable within the tested 5–500 MU). EPIgray was found to be reasonably sensitive to patient thickness changes but unable to reliably detect thickness changes of less than 1 cm. EPIgray was not sensitive to SSD errors (Fig. 5); the latter is common to all transit dosimetry [16]. Wedged fields were found to be modelled less well within the phantom study (central axis $\Delta D$ values were within $\pm 4.5\%$ at 10 cm deep and within $\pm 6.9\%$ up to a measured depth of 18 cm excluding the build-up region). In addition, the dose to off-axis points ($>2$ cm off-axis) or points within 1 cm of the field edge had a greater $\Delta D$ value than central axis points in all cases. This corroborates well with previous TPS commissioning measurements: agreement between the TPS and machine delivery was measured using ion chambers and a 2D array across a range of open field parameters and was within $\pm 2\%$ for the majority of the points tested. However, for a number of off-axis, wedged and wedged off-axis points ion chamber measurement was found to be up to 3\% less than TPS dose. It is therefore recommended that dose points are created on the central axis or within 2 cm of the central axis where possible.

In order to test EPIgray performance with heterogeneity we measured 126 beams delivered to a cohort of 37 patients comprised of abdominal, brain, head and neck, lung, pelvis, rectal and sarcoma 3D conformal treatments. Patient results demonstrated the potential of EPIgray to verify delivered dose well within inhomogeneities; 91\% of all delivered fields achieved the initial set tolerance level of $\Delta D$ of 0 $\pm 5$ cGy or $\% \Delta D$ of 0 $\pm 5\%$. In addition a short study using an inhomogeneous phantom containing an 8 cm diameter cylindrical lung equivalent insert demonstrated that EPIgray overestimated dose within low density regions, by up to $\% \Delta D$ of $+5.1\%$. However dose reconstruction points were not placed within heterogeneities within the patient trial. Nevertheless dose reconstructed in solid water regions distal to the lung insert displayed $\% \Delta D$ values up to $-2.4\%$; when compared to central axis measurements measured up to 18 cm depth within homogeneous phantoms (which yielded $\Delta D$ within $-1.4\%$) it is evident that heterogeneity within the beam path affects EPIgray performance.

### Table 4

Summary of results of EPID patient transit dosimetry; $\% \Delta D$ and $\Delta D$ values for all measured fields per site, pass/fail status of fields and reasons for failure.

<table>
<thead>
<tr>
<th>Site</th>
<th>Patient number</th>
<th>No. fields measured</th>
<th>$% \Delta D \pm 1 \text{ SD}$</th>
<th>$\Delta D \pm 1 \text{ SD}$ (cGy)</th>
<th>Pass/fail status</th>
<th>Reason for fail</th>
</tr>
</thead>
<tbody>
<tr>
<td>Abdomen 1, 2</td>
<td>6</td>
<td>1.0 $\pm$ 2.1</td>
<td>1.1 $\pm$ 1.8</td>
<td>All pass</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Brain 3–12</td>
<td>25</td>
<td>-3.2 $\pm$ 2.6</td>
<td>-2.0 $\pm$ 1.8</td>
<td>All pass</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Head and neck</td>
<td>13–17</td>
<td>2.0 $\pm$ 6.6</td>
<td>1.4 $\pm$ 5.1</td>
<td>1 field failed; Repeat failed</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Lung</td>
<td>18–21</td>
<td>3.0 $\pm$ 5.4</td>
<td>2.1 $\pm$ 4.3</td>
<td>Patient 19 (2 fields) failed; Patient 19 re-planned and passed</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pelvis</td>
<td>22–24</td>
<td>-3.5 $\pm$ 2.5</td>
<td>-2.6 $\pm$ 1.7</td>
<td>2 fails, passed upon repeat</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Rectum</td>
<td>25–34</td>
<td>-0.9 $\pm$ 4.9</td>
<td>-0.3 $\pm$ 2.4</td>
<td>1 fail, passed upon repeat;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Sarcoma</td>
<td>35–37</td>
<td>-3.9 $\pm$ 4.1</td>
<td>-3.0 $\pm$ 3.7</td>
<td>1 fail, passed upon repeat;</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All wedged</td>
<td>103</td>
<td>-0.5 $\pm$ 5.4</td>
<td>-0.2 $\pm$ 3.9</td>
<td>Ray trace passed significant amount of air</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All open</td>
<td>23</td>
<td>-1.7 $\pm$ 4.7</td>
<td>-1.1 $\pm$ 3.1</td>
<td>Isocentre in heterogeneity. Replan: Isocentre in soft tissue</td>
<td></td>
<td></td>
</tr>
<tr>
<td>All open (excluding lung/H&amp;N)</td>
<td>20</td>
<td>-2.3 $\pm$ 4.6</td>
<td>-1.6 $\pm$ 2.9</td>
<td>2 x Imager malfunctions</td>
<td>Patient positioning error</td>
<td>Imager malfunction; Unplanned arm rotation</td>
</tr>
<tr>
<td>All 1–37</td>
<td>126</td>
<td>-0.9 $\pm$ 5.0</td>
<td>-0.4 $\pm$ 3.4</td>
<td>7 image repeats (including 3 imager problems); 1 re-plan</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

---

**Figure 7.** Histograms of $\% \Delta D$ and $\Delta D$ of 126 patient-delivered 3D conformal fields.
Importantly the system was able to detect errors in patient set up; patient 35 (left bicep) had a small amount of rotation of the arm in the first measurement as noted from verification images. On the repeat the expected and measured iView images coincided well and the field passed. This is a common issue for treatment of sarcomas in our centre as vacuum bags are used for immobilisation, and are less reproducible than thermoplastic shells. Patient 26 was found to have shifted on the order of 2 mm upon inspection of verification images; this translated to a \( \%A_D = 4.5\% \) value of the posterior field of \( -12.5\% \) \((-6.4\,\text{CGy}) \) for this fraction due to a highly modulated patient surface overlying the dose calculation point. On the repeat the verification images coincided well with planned images and the field passed \( \%A_D = 4.5\% \).

Other than patient errors, a number of factors also contributed to out of tolerance measurements for this measured cohort including: (i) EPID imager malfunction for 3 fields; (ii) Attenuation of the exit beam through devices, e.g. the couch bar present for patient 37. Such devices are accounted for in the TPS only for devices proximal to the beam entrance; (iii) Algorithm limitations: The TPS AAA algorithm and EPIgray algorithm deal with tissue heterogeneity differently.

The standard deviation of \( \%A_D \) varied with treatment site, and indicated the stability of conditions through which the central axis ray line traversed. Head and neck and lung fields showed the greatest standard deviations \((9.0\% \text{and } 5.4\%\), respectively), due to (i) tissue heterogeneity (and thus sensitivity to slight changes in patient position), and (ii) including regions of potential anatomical change. Brain is a more homogeneous site suffering less anatomical change and thus displayed a lower standard deviation of 2.6%.

Wedged fields delivered to patients performed well: all wedged fields were analysed together and displayed an average \( \%A_D \) value of \(-0.5\% \text{ and } 5.4\% \) compared to \( -1.7\% \text{ and } 4.7\% \) for all open fields. Upon excluding patients with high levels of heterogeneity (lung and head and neck fields) the \( \%A_D \) values of wedged and open fields came to a closer agreement \((-2.1\% \text{ and } 2.3\% \text{ respectively})\). It was expected that patient wedged fields performed better than the phantom results because the latter displayed results of a fully 60° wedged field, whereas patient fields generally had a lower wedge weighting. Wedged fields delivered to patients corroborated well with results for wedged fields within the phantom study, which found that central axis dose was in closer agreement to TPS for 10 MV fields than 6 MV fields \((\%A_D = -4.5\% \text{ and } -1.9\%)\) for 6 MV and 10 MV beams, respectively. A greater deviation from TPS dose was found for 6 MV patient wedged fields compared to 10 MV: \( \%A_D \) of \(-2.7\% \text{ and } 3.1\% \text{ and } -0.9\% \text{ and } 4.8\% \) respectively, analysed for all patients excluding head and neck and lung in order to reduce heterogeneity to closer model phantom conditions.

EPIgray tended to overestimate dose compared to TPS for treatment sites that involved beam transit through low density such as pelvis, brain and sarcoma \((\%A_D \text{ of } -3.5\%, -3.2\% \text{ and } -3.9\%\), respectively), whereas the solid water phantom study predicted better agreement \((\%A_D \text{ values within } -0.5\% \text{ at } 10 \text{ cm deep})\). This can be attributed to the difference in handling heterogeneity between the EPIgray algorithm and the TPS AAA algorithm. For this reason it may be important to consider setting asymmetric action levels, different for each treatment site, but no firm action levels could be set from this data as more patient numbers would be required.

Transit dosimetry cannot be used for all patient treatments: within this cohort 7 fields could not be measured due to inability to extend the EPID, and diode measurements were made of these fields. We suggest the following patient exclusion criteria: (1) patients requiring a floor twist, table lateral setting or immobilisation device that impedes panel extension, (2) patients with superior anterior oblique fields which impede panel extension, as evident in this cohort, (3) field lengths greater than 26 cm with a collimator twist. The latter is in place to reduce irradiation of EPID panel electronics, which can shorten the lifetime of the EPID. In addition there are cases where the isocentre is placed in the patient centre even for peripheral targets to ensure that CBCT or arc therapy gantry motion is possible without collision. For such cases off-axis dose calculation is required to enact treatment verification, and so further improvement to off-axis modelling would be beneficial to ensure EPIgray can be used in these cases.

We have found that the pass/fail status of a field was influenced by the location of the dose calculation point. Other groups have chosen points on the central axis at 5 cm deep \([15,16]\); however we decided to position our points at isocentric depth for dose calculation at tumour position, and Fig. 2 displayed that EPIgray performance was stable up 18 cm depth as measured in a water phantom. From the phantom measurements we recommend that the dose reference point should be placed at the isocentre (the protocol we have adopted in our centre), or in a position that meets the following criteria in order to reduce false positives: (i) lies >2 cm from a field edge, (ii) lies <2 cm from the central axis, (iii) must not lie in a high dose gradient, (iv) must not lie in a low dose area (<90% of prescribed dose), (v) must not lie beneath MLCs, this may not be avoidable for some IMRT segments but should be minimised as much as possible, or (vi) avoid positions where a slight change in lateral or sup-inf position of the point results in a large change of geometrical or radiological depth (i.e. regions of highly modulating surface contour or heterogeneity along the field projection), (vi) must not lie in air. Points (i), (ii), (iii) and (v) may have implications for small field segments in IMRT fields, by an amount related to the weighting of such small segments. However another group has demonstrated promising results for the use of EPIgray to measure IMRT fields of head and neck patients \((\text{mean } \pm 1 \text{SD } \%A_D = 1.53 \pm 2.46\%)\) and stated that field fails were mainly due to the dose calculation point lying within high dose gradient regions \([21]\). Testing EPIgray performance for these more complex fields is currently being undertaken at our centre.

There is a required difference between how the TPS and EPIgray must deal with couch attenuation. TPS requires a couch attenuation factor for fields that pass the couch proximal to the patient (which affects patient dose). Transit dosimetry requires a couch factor for fields that pass the couch either proximal or distal to the patient as couch absorption will cause an underestimation of patient dose if not accounted for. The EPIgray algorithm deals with this through application of an average couch factor to all fields. If a couch specific model is used within the TPS (as for IMRT patients at our centre) then care must be taken to ensure that EPIgray does not perform an additional couch attenuation correction. However for 3D conformal patients at our centre no TPS couch model is implemented (with manual correction for couch transmission applied to field MU) and so the EPIgray couch factor was applied. This average factor will cause an overestimate to lateral fields that do not traverse the couch, and an underestimate of oblique fields that traverse a greater thickness of couch, on the order of ±2%.

We have developed a decision tree for clinical workflow of EPIgray (Fig. 8). If fields fell outside of the set tolerance \((\%A_D \pm 5\% \text{ in this trial instance})\) an investigation was conducted by the radiotherapy physics department to find the reason for the difference. The final investigation in Fig. 8 is undertaken to investigate any potential patient anatomical changes such as bowel gas, weight loss/gain, and bladder filling that may have contributed to a discrepancy in reconstructed dose. Any fields that fail on the first repeat were re-measured using diodes.

Transit dosimetry is a sound final step in the treatment chain with potential to determine errors in dose delivery, without additional dose to the patient or time demands. It is beneficial that the
commercial software allowed flexibility in dose calculation point location, but a full 2D map of TPS – transit dose discrepancy would be of greater use in providing more insight into not only location but distribution of the error source. A new software feature enables calculation of the gamma value of the dose reconstruction point P across a local area of 30-mm centered around P for each field, comparing D_EPI at P to the TPS dose map (dose agreement and distance to agreement criterion can be manually set by the user). We are currently assessing this feature for our IMRT patients. Analysis was performed offline, but the ultimate goal would be to spot errors in real-time to reduce the impact of severe errors. The software algorithm used the planning CT to reconstruct the patient dose; for a true in vivo dosimetry measurement this should be performed using CBCT information acquired on-set such that dose is reconstructed on current patient geometry. However, it was not the primary goal of this technique to record in vivo doses, but to identify errors, and for this aim the planning CT was the ideal choice of patient information to magnify the impact of anatomical changes between the planning and treatment stages.

Conclusions

A commercially available system for patient dose verification was implemented and tested with a series of phantom and patient measurements. Commissioning was simple and implementation measurements were not excessive and could be performed in 5 h with an additional 2–3 h per linac. The commercial system was used to verify 126 3D conformal beams delivered to a cohort of 37 patients. The preliminary data suggested that difference action levels may be required for different treatment sites. Patient positioning errors could be spotted by this system. False positive errors were due to algorithm limitation, EPID imager malfunction, and attenuation of the exit beam through immobilisation devices. Through using an appropriate clinical workflow for investigation of out of tolerance results, true positives can be recognised from false positives, and thus real dose delivery errors can be minimised.

Acknowledgments

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References


