Three-dimensional printer-generated patient-specific phantom for artificial in vivo dosimetry in radiotherapy quality assurance

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ABSTRACT

Pretreatment intensity-modulated radiotherapy quality assurance is performed using simple rectangular or cylindrical phantoms; thus, the dosimetric errors caused by complex patient-specific anatomy are absent in the evaluation objects. In this study, we construct a system for generating patient-specific three-dimensional (3D)-printed phantoms for radiotherapy dosimetry. An anthropomorphic head phantom containing the bone and hollow of the paranasal sinus is scanned by computed tomography (CT). Based on surface rendering data, a patient-specific phantom is formed using a fused-deposition-modeling-based 3D printer, with a polyactic acid filament as the printing material. Radiophotoluminescence glass dosimeters can be inserted in the 3D-printed phantom. The phantom shape, CT value, and absorbed doses are compared between the actual and 3D-printed phantoms. The shape difference between the actual and printed phantoms is less than 1 mm except in the bottom surface region. The average CT value of the infill region in the 3D-printed phantom is $-6 \pm 18$ Hounsfield units (HU) and that of the vertical shell region is $126 \pm 18$ HU. When the same plans were irradiated, the dose differences were generally less than 2%. These results demonstrate the feasibility of the 3D-printed phantom for artificial in vivo dosimetry in radiotherapy quality assurance.

1. Introduction

Intensity-modulated radiation therapy (IMRT) and volumetric-modulated arc therapy (VMAT) have recently been adopted for various cancer sites, contributing to improvements in survival outcomes with fewer treatment-related side effects [1–4]. Previous reports recommend that patient-specific quality assurance (QA) be performed for patients having IMRT and VMAT to reduce the likelihood of dose errors and assure patient safety during radiotherapy [5,6]. Detecting delivered dose errors in IMRT and VMAT is especially important, as these involve complex procedures that contain many potential errors. In general, patient-specific QA is performed on the direct dose measurements or independent dose calculation. Independent dose calculation systems have recently progressed rapidly; one of their limitations, however, is its lack of validation of the deliverability of a treatment plan on the actual treatment device [7]. Therefore, these validations must be performed in combination with thorough quality control of accelerator and multileaf collimator performance. Moreover, it is strongly recommended to perform experimental verification for a relatively large number of patients to prevent systematic dose errors during initial clinical implementation of a new IMRT technique [8]. For this reason, based on institutional policies or protocols, patient-specific QA is often conducted using experimental dose measurements. Experimental dose measurements are usually performed using simple rectangular or cylindrical phantoms with various detectors (e.g., an ionization chamber or radiochromic film) [9–11]. In some cases, pretreatment QA results may reveal disagreement between the measured and treatment planning system (TPS)-calculated phantom doses. In these cases, we need to judge whether these disagreements are at acceptance or action levels by considering other metric values, such as the distance to the agreement.
and gamma index. However, recent work has discussed the insufficient sensitivity of these conventional pretreatment QA results in predicting the clinical impact on the patient from changes in the dose-volume histograms for the clinical target volume and organs at risk [12,13]. One reason for this poor sensitivity is that the shape of the phantoms used for pretreatment QA and that of patients are entirely different. In fact, it is difficult to estimate the magnitude and location of a dose error in a patient from the phantom dose error. Ideally, the magnitude and location of the errors in the patient may provide more important information than the evaluation metrics (e.g., pass rate) based on conventional pretreatment QA, and would be more critical in clinical judgment. From these viewpoints, to estimate the in vivo dose to the patient is a challenge in pretreatment QA.

Recent studies have demonstrated that 3D printing offers flexibility in design and manufacturing for auxiliary equipment in radiotherapy [14]. Ju et al. suggested a new system for manufacturing a range compensator in proton therapy, using an industrial 3D printer with ultraviolet curable acrylic plastic [15]. Park et al. fabricated a patient-specific customized bolus, using a low-cost personal 3D printer with a polylactic acid (PLA) filament, and evaluated its feasibility for post-total-mastectomy electron conformal therapy [16]. Further, Ehler et al. demonstrated the construction of a patient-specific phantom, using a 3D printer with an acrylonitrile butadiene styrene (ABS) filament for dosimetry [17]. ABS warps on cooling, preventing this phantom from being printed as a 100% solid object; thus, their approach was to print the exterior shape of the phantom and fill it with a uniform mixture of a modified M3 mix. PLA filaments are generally more robust to warping during cooling than ABS filaments [16,18,19]. The aim of this study is to construct a system for the fabrication of patient-specific phantoms, using a 3D printer with a PLA filament and to evaluate the feasibility of this 3D-printed phantom for artificial in vivo dosimetry in radiotherapy QA. To simplify this procedure for clinical use, the phantom is fabricated using only 3D printing—without filling the inner cavity with a liquid material such as the modified M3 mix. Considering the fabrication cost of the phantom, we have adopted a low-cost personal 3D printer.
2. Materials and methods

2.1. Image acquisition, data processing, and printing procedure

The overall fabrication process is summarized in Fig. 1. An anthropomorphic head phantom containing the bone and hollow of the maxillary sinus (QualitA Co, Ltd., Nagano, Japan) was selected as the mock patient. The fabrication target object was one section of the phantom (Fig. 1, process 1), and the maximum lengths were about 25 mm, 150 mm, and 195 mm in the superior–inferior, left–right, and anterior–posterior directions, respectively. The phantom was scanned using a computed tomography (CT) scanner (Asteion 4; Toshiba Medical Systems, Tokyo, Japan). The CT data were reconstructed in a 300-mm field of view on a 512 × 512 grid with a slice thickness of 1.0 mm and exported in DICOM format.

Data processing was performed per the following three steps: (i) creation of the 3D surface rendering of the exterior shape of the patient, (ii) formation of the insertion slots for dosimeters, and (iii) creation of the control code containing the build information for 3D printing.

OsiriX Lite ver. 8.0.2 (Pixmeo Inc., Bernex, Switzerland) was used for 3D surface rendering of the patient’s exterior shape. The threshold value for surface rendering was set to –500 Hounsfield units (HU), which matches the OsiriX default surface rendering value of the skin surface and that of previous research [17]. Surface rendering data were saved in a stereo-lithography (STL) format file.

The STL file was imported into 3D modeling software (Shade3D professional ver. 15; Shade3D Co., Ltd., Tokyo, Japan) to prepare slots for dosimeter insertion. A GD-302 M radiophotoluminescence (RPL) glass dosimeter (Asahi Techno Glass Corporation, Shizuoka, Japan) and an FGD-1000 readout system (Asahi Techno Glass Corporation, Shizuoka, Japan) were selected for the dosimeter for several reasons: the uniformity and reproducibility of the signal from 50 RPL glass dosimeters are each ± 1.1% (one standard deviation); good dose linearity is maintained for doses ranging up to 30 Gy; energy dependence is within 2% for photon energies of a 60Co beam, 4 MV, and 10 MV X-ray beams; and angular dependence, ranging from 0° (along the long axis of the RPL glass dosimeter) to 120°, is within 1.5% [20]. RPL glass dosimeters also exhibit less fading (within 1.7% after five months) and can be read repeatedly [21]. The diameter of the RPL glass dosimeter, stored in a dedicated plastic holder, was measured with a micrometer (NSK Ltd, Tokyo, Japan) as 2.78 mm. Prior to final 3D printing, trial printing to determine the slot diameter was performed because contraction was expected to occur during cooling. Trial 3D models were also created with Shade3D software, an example of which is depicted in Fig. 2. The exterior shape was a 20 mm × 20 mm × 20 mm cube; slot diameters of 2.8 mm, 2.9 mm, 3.0 mm, and 3.1 mm were validated.

The STL files were then compiled into G-code, which is used to run the commands that modulate the position, velocity, temperature, and extrusion timing of the 3D printer, using Slic3r Ver. 1.2.9 [22] slicing software; the printing parameters are summarized in Table 1.

The patient-specific phantom was generated based on the G-code data using a FDM-200W personal 3D printer (Ninjabot, LCC, Shizuoka, Japan) with a PLA filament, which is a fused-deposition-modeling (FDM)-based 3D printer.

2.2. Irradiation plans and evaluation methods

In this study, all measurement are based on the use of 6-MV photon...
beams produced using a Clinac iX linear accelerator (Varian Medical Systems, Palo Alto, CA). To evaluate basic properties, the phantoms were irradiated using a single beam with a 260° gantry angle and a field size of 8 cm × 8 cm. The monitor unit was set to 100 MU. The positional relationships between the irradiation beam and the RPL glass dosimeters are shown in Fig. 3a. The dose distributions were calculated with Eclipse TPS (Ver. 13.6; Varian Medical System, Palo Alto, CA). All dose calculations were performed using the anisotropic analytical algorithm (AAA; Varian Medical Systems, Palo Alto, CA) and a dose calculation grid size of 1.0 mm × 1.0 mm. In this study, to minimize the angular dependence of the RPL glass dosimeter, all dosimeters were placed perpendicular to the beam axis, and their readout centers coincided with the isocenter plane of the linear accelerator.

A clinical case was evaluated using a two-arc VMAT plan. A maxillary sinus cancer patient (coinciding with the anatomical site of the 3D-printed phantom) who had previously undergone VMAT at our institution was selected. The study was approved by the Institutional Review Board of the Nagoya University Hospital and all patients provided written informed consent. Two-arc VMAT was performed using 181°–179° clockwise and 179°–181° counterclockwise rotations; the positional relationships between the irradiation beam and the RPL glass dosimeters are depicted in Fig. 3b.

Dose readings of the RPL glass dosimeters were performed automatically. We determined the calibration rate between the average raw output value of ten RPL glass dosimeters and the dose measured by the calibrated ionization chamber (Farmer-type ionization chamber type 30013; PTW, Freiburg, Germany). This calibration was performed using 181°–179° clockwise and 179°–181° counterclockwise rotations; the positional relationships between the irradiation beam and the RPL glass dosimeters are depicted in Fig. 3b.

Dose readings of the RPL glass dosimeters were performed automatically. We determined the calibration rate between the average raw output value of ten RPL glass dosimeters and the dose measured by the calibrated ionization chamber (Farmer-type ionization chamber type 30013; PTW, Freiburg, Germany). This calibration was performed using 6-MV photon beam with a 10 cm × 10 cm field size in a solid water phantom (Tough Water, Kyoto Kagaku Co., Ltd., Kyoto, Japan), at a depth of 10 cm [20,23]. The final outputs were obtained by multiplying the raw output values of the RPL glass dosimeters by the calibration rate. To eliminate the fading effect, the readout was performed within 24 h after irradiation. Measurements of the single static beam and the two-arc VMAT plan were repeated thrice.

To evaluate the feasibility of using the 3D-printed phantom for dosimetry, the phantom shape and CT value were compared with the anthropomorphic head phantom. Furthermore, the similarity of their external shapes was determined by calculating their Dice similarity coefficient [24]. These evaluations were performed using Eclipse TPS. The average CT value was calculated from 20 randomly selected points of interest. The dose differences, obtained from the RPL glass dosimeters, between the anthropomorphic head and 3D-printed phantoms were then estimated. Finally, the differences between the measured and TPS-calculated doses were determined. Moreover, in one case, the CT values of the 3D-printed phantom were converted to 374 HU, which corresponds to the PLA density of 1.25 g/cm³, and dose comparisons were continued.

3. Results

In a trial print to verify the slot diameter for inserting the RPL glass dosimeters, slot diameters of 2.8 mm, 2.9 mm, 3.0 mm, and 3.1 mm were tested; the best size in our environment was 3.0 mm. Diameters of 2.8 mm and 2.9 mm could not be inserted into and 3.1 mm was loose. Fig. 4 shows images of the anthropomorphic head and 3D-printed phantoms. The total printing time for this part was approximately 20 h. The shape difference between the anthropomorphic head phantom and the 3D-printed phantom was less than 1 mm, except for the bottom surface region (Fig. 5). The white arrows in Fig. 5b represent geometric errors exceeding 1 mm. A maximum difference of 2.0 mm was exhibited on the bottom surface of the posterior region in the patient. The Dice similarity coefficient for the external shape was 0.974. A comparison of the CT value profiles is depicted in Fig. 6. Except for the bone region, the profile shape of the 3D-printed phantom coincided with that of the anthropomorphic head phantom. The maximum difference in CT values for the internal region was within ± 100 HU, and that of the border region between tissue and air was within ± 250 HU. The average CT value of the soft tissue region in the anthropomorphic head phantom

![Fig. 5. CT images of the anthropomorphic head phantom (left), 3D-printed phantom (center), and the overlaid images of the anthropomorphic head phantom (green) and the 3D-printed phantom (red) (right): (a) axial, (b) coronal, and (c) sagittal views. The white arrow represents region in which the geometric error exceeds 1 mm.](image-url)
was 12.7 ± 17.5 HU. The average CT value of the infill region in the printed phantom was −5.8 ± 18.8 HU and that of the vertical shell region was 126.1 ± 17.5 HU.

The measured dose differences between the anthropomorphic head phantom and the 3D-printed phantom are presented in Table 2. For single static field irradiation, the measured dose difference between the 3D-printed and anthropomorphic phantoms was 0.6 ± 1.9%; for the two-arc VMAT plan, the measured dose difference was 1.5 ± 1.2%. The measured doses on the 3D-printed phantom were slightly increased compared to those of the anthropomorphic phantom.

The percentage dose differences between the measured and TPS-calculated doses (measurement - calculation) relative to the TPS-calculated dose are indicated in Table 3. For single static field irradiation, the average percentage dose differences of the anthropomorphic head phantom, 3D-printed phantom, and 3D-printed phantom with converted CT values were 0.9 ± 2.5%, 2.3 ± 1.1%, and 6.3 ± 4.5%, respectively. These average percentage dose differences did not include the results of the dosimeter location number seven, which was outside the radiation field. For the two-arc VMAT plan, the average percentage dose difference of the anthropomorphic head phantom, 3D-printed phantom, and 3D-printed phantom with converted CT values were −1.2 ± 1.6%, −2.1 ± 1.9%, and 1.3 ± 2.7%, respectively. All percentage dose differences between the anthropomorphic head phantom and 3D-printed phantom were within ±5%. However, the results of the 3D-printed phantom with converted CT values were poorer than those without converted CT values.

4. Discussion

Objects printed using FDM tend to warp during cooling, a characteristic that may depend on the extruder temperature, bed temperature, object size, area, and printing material. Several previous works had employed ABS filament because its physical properties closely resemble those of water [17,25,26]. However, ABS filament can undergo considerable deformation during cooling [16,18,19]; thus, we employed PLA filament. The results of this study demonstrate slight geometric errors between the CT scanned image of the anthropomorphic head phantom and the 3D-printed phantom, mostly in the bottom region. This geometric error also contains the uncertainty of the partial volume effect in the CT images. Except for the bottom region, the geometric error was less than 1 mm. In addition, the Dice similarity coefficient for the external shape was equivalent to that in Ehler’s work [17]. Thus, with respect to the geometrical reproducibility of the 3D-printed phantom, it would be possible to use a patient-specific 3D-printed phantom created using a PLA filament for radiotherapy dosimetry. Moreover, fabrication quality can be improved by optimizing the printing parameters and employing high-performance devices and software.

The average HU of the 3D-printed phantom infill region in this study is −6 ± 18 HU, lower than those in the previous works. Burleson et al. found the average HU of a 3D-printed PLA bolus to be approximately 160 HU [26], whereas, Zue et al. declared it to be approximately 106 HU [27]. In contrast, the HU values of the vertical shell region in this work are approximately 125 HU, which is comparable to those in previous work [26,27]. This tendency resembles the control results of the fill density parameter reported by Madamesila et al. [28]. The most obvious cause of this change in HU seems to be the variation in the actual fill density, since the material composition of the infill and shell region are identical. Therefore, different 3D printers and software packages may produce different actual material densities for the same fill density parameter, and it is important to evaluate the actual fill density before clinical use.

Table 2

<table>
<thead>
<tr>
<th>Location</th>
<th>Single static field</th>
<th>Two-arc VMAT plan</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Anthropomorphic</td>
<td>3D-printed</td>
</tr>
<tr>
<td>1</td>
<td>1.01 ± 0.013</td>
<td>1.03 ± 0.019</td>
</tr>
<tr>
<td>2</td>
<td>0.81 ± 0.006</td>
<td>0.83 ± 0.007</td>
</tr>
<tr>
<td>3</td>
<td>0.67 ± 0.001</td>
<td>0.66 ± 0.010</td>
</tr>
<tr>
<td>4</td>
<td>1.03 ± 0.004</td>
<td>1.01 ± 0.006</td>
</tr>
<tr>
<td>5</td>
<td>0.79 ± 0.011</td>
<td>0.80 ± 0.003</td>
</tr>
<tr>
<td>6</td>
<td>0.60 ± 0.005</td>
<td>0.62 ± 0.005</td>
</tr>
<tr>
<td>7</td>
<td>0.03 ± 0.000</td>
<td>0.03 ± 0.001</td>
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</tbody>
</table>

Fig. 6. CT value profiles and their differences along the white dotted arrows in Fig. 5a in the (a) sagittal and (b) coronal directions.
density of each 3D printer and software package before using a 3D-printed phantom for dosimetry.

RPL glass dosimeters were employed in this study. To minimize dose uncertainty caused by angular dependency, energy dependency, and fading, we carefully handled the dosimeters throughout, with particular adherence to previous studies [20,23]. In Japan, postal dose audits have been performed on radiation therapy units using RPL glass dosimeters since 2007 [29] and previous studies have employed RPL glass dosimeters for radiotherapy dose verification. Hus et al. compared measurements from an RPL glass dosimeter with those from a thermoluminescent dosimeter and radiographic film, and found differences less than 5%, which is clinically acceptable accuracy [30]. Nakaguchi et al. also reported that the accuracy of the RPL glass dosimeter measurement was almost 5% and that angular dependency and geometric uncertainties should be considered [23]. They also raised the issue of geometric uncertainties in which the insertion slots in the phantom were larger than the RPL glass dosimeter. This is an advantage of 3D-printed phantoms, in that they may reduce the geometric uncertainty caused by a diameter mismatch between the slots and the dosimeter.

In our environment, the maximum dose differences between the anthropomorphic head phantom and the 3D-printed phantom were 3.9%, and most of the measurement points agreed within 2%. As per our expectations, notable dose differences should have been obtained because this study employed PLA filament. However, the dose difference between the anthropomorphic head phantom and the 3D-printed phantom was almost within the measurement uncertainty for the RPL glass dosimeter [23,30]. A factor contributing to this result may be the low CT value in the infill region of the 3D-printed phantom, as discussed in the previous section. Differences in the equipment and software used may affect this result; thus, it is necessary to verify it in each environment. Moreover, the dosimetric impact of bone structures, described in previous studies [31,32], was not noticeable from the dose differences between the anthropomorphic head phantom and the 3D-printed phantom. Ehler et al. also described differences in doses at a bone–tissue interface [15]. As for the necessity of considering bone structure, we agree with previous studies, and future investigation is needed. One potential reason for the low sensitivity of the dosimetric impact of the bone structure in this study is the energy dependency of the RPL glass dosimeter. The RPL glass dosimeter has a higher response to low-energy photons [33,34]. It is possible that because of extra low-energy scattered photons and electrons inside the bone material [32], the doses measured by the RPL glass dosimeter were inaccurately accounted for by energy dependency.

Dosimetric comparison between the calculated TPS and the measured dose was carried out in this study, and the percentage dose differences between the anthropomorphic head phantom and 3D-printed phantom for single static field and VMAT plans were within ±5%, which is within the measurement uncertainty for the RPL glass dosimeter [21,28]. Note that measured doses were slightly lower in the VMAT cases than in the static field cases. We consider that this tendency may be caused by field-size dependence [33,34]. The average field size decreases by VMAT, low-energy scattered radiations in the phantom decreases, and the outputs of the RPL glass dosimeters decrease. Therefore, the field-size correction factor described by Mizuno et al. [33] may be useful for obtaining more accurate IMRT and VMAT treatment plan verification results. In many cases, when comparing the TPS-calculated and measured doses, the phantom density should be overridden in the TPS with a suitable value [26]. In this study, the CT values of the 3D-printed phantom was converted to the corresponding PLA density value of 1.25 g/cm³. However, the results of the 3D-printed phantom with the converted CT values deteriorated compared to those without the converted CT values. In our experimental environment, converting the CT values with the PLA density can be regarded as overcorrection. This tendency does not contradict the result of low CT value in the infill region of the 3D-printed phantom, as mentioned in the above section.

The initial cost of the 3D printer used in this work was $2500 and that of the filament was $40 per 750 g. This system, for producing patient-specific 3D-printed phantoms, was confirmed to have permissible geometric accuracy and is cost effective. However, the total printing time to fabricate the part was approximately 20 h, which needs improvement. If the extruder head speed is identical, fabrication time depends on the distance that the extruder head moves. One of the methods for reducing fabrication time involves the usage of larger nozzle diameters (e.g. 0.6 mm). Accordingly, it would be possible to use greater layer-height and extrusion-width settings to reduce the distance that the extruder head moves. However, this may affect the geometric fidelity and dosimetric properties of the phantom and requires additional evaluation.

5. Conclusion

We have constructed a system for producing patient-specific phantoms, using a low-cost personal 3D printer with a PLA filament. The modeling accuracy of the 3D-printed phantom was acceptable. The dose differences between the anthropomorphic head phantom and the 3D-printed phantom were approximately within 2%, in our experimental environment. These results demonstrate the feasibility of the 3D-printed phantom for artificial in vivo dosimetry in radiotherapy QA.

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