Using NIPAM gel dosimeter and concentric swing machine to simulate the dose distribution during breathing: A feasibility study

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Abstract.

\textbf{BACKGROUND:} Radiotherapy plays an important role in cancer treatment today. Successful radiotherapy includes precise positioning and accurate dosimetry.

\textbf{OBJECTIVE:} To use NIPAM gel dosimeter and concentric swing machine to simulate and evaluate the feasibility of lung or upper abdominal tumor dose distribution during breathing.

\textbf{METHODS:} We used a concentric swing machine to simulate actual radiotherapy for lung or upper abdomen tumors. A $4 \times 4 \text{ cm}^2$ irradiation field area was set and MRI was performed. Next, readout analysis was performed using MATLAB and the 3 mm, 3\% gamma passing rate > 95\% was used as a basis for evaluation.

\textbf{RESULTS:} The concentric dynamic dose curve for a simulated respiratory rate of 3 seconds/breath and $4 \times 4 \text{ cm}^2$ field was compared with $4 \times 4$, $3 \times 3$, and $2 \times 2 \text{ cm}^2$ treatment planning systems (TPS), and the 3 mm, 3\% gamma passing rate was 42.87\%, 54.96\%, and 49.92\%, respectively. Pre-simulation showed that the high-dose region dose curve was similar to the $2 \times 2 \text{ cm}^2$ TPS result. After appropriate selection and comparison, we found that the 3 mm, 3\% gamma passing rate was 97.92\% on comparing the $> 60\%$ dose curve with the $2 \times 2 \text{ cm}^2$ TPS.

\textbf{CONCLUSIONS:} NIPAM gel dosimeter and concentric swing machine use is feasible to simulate dose distribution during breathing and results conforming to clinical evaluation standards.

Keywords: NIPAM gel dosimeter, organ motion, MRI

1. Introduction

Radiotherapy plays an important role in cancer treatment today. Successful radiotherapy includes precise positioning and accurate dosimetry. Currently, common dosimetry tools include free-air ionization chambers, films, and thermoluminescent dosimetry.
Gel dosimeter is a chemical dosimeter that is also a relative dosimeter and has the advantage of providing three-dimensional relative dose distribution [1]. The action mechanism of gel dosimeter is the generation of free radicals by ionization of water molecules by radiation. Monomers and free radicals continuously bind to form polymer chain molecules and gelatin matrices to form a three-dimensional structure. After the gel dosimeter is irradiated, a readout instrument can be used to obtain image data to analyze the dose. Commonly used readout instruments include magnetic resonance imaging (MRI) [2], optical computed tomography (OCT) [3], and computed tomography (CT) [4]. MRI has the advantages such as having high resolution and high contrast; however, disadvantages such as high cost and temperature increase under long periods of scanning may result in dose differences.

Previous dosimetry studies mostly used static treatment target regions. However, the physiological motion and displacement of organs must be considered in some radiotherapy cases, of which breathing is a physiological motion that tends to cause organ motion. Bussels et al. used MRI measurement and found that significant motion of organs in the thorax and abdomen in the cranial-caudal direction will occur due to breathing [5]. Korin et al. used MRI to measure the displacement of upper abdominal organs in 15 volunteers and found that the diaphragm had the greatest cranial-caudal displacement and the motion range for normal breathing and deep breathing was 13 mm and 39 mm, respectively [6]. In addition, studies also mentioned that a higher dose may be needed in radiotherapy for various malignant tumors (such as lung cancer) due to organ motion caused by breathing so as to increase therapeutic efficacy and increase local control rate [7,8]. Therefore, accurate dose measurement in the target region is important when organ motion causes changes in the treatment target volume.

Dynamic phantoms have clinical value in simulating dosimetry during organ motion. In 2004, David et al. published a paper on quantitation of abdominal tumor motion caused by breathing using dynamic phantoms and its impact on dose distribution in intensity-modulated radiotherapy. The authors felt that interference caused by tumor motion must be decreased during treatment [9]. In 2009, Masset et al. employed dynamic phantoms and films to simulate the effects of normal breathing on planar doses. They found that there were differences in dose distribution in the longitudinal and axial directions under motion and found that simulation could be used to measure the consistency of real-time position management system [10]. In 2011, Boda-Heggemann et al. used dynamic phantoms and CBCT to simulate image-guided radiotherapy of lung tumors. The study results showed that simulation by using dynamic phantoms combined with breathing monitoring equipment and comparison with clinical images could increase the treatment accuracy for lung tumors [11].

The past research of NIPAM gel dosimeter has good reading results, and it is worth looking forward to the dose presentation in the three-dimensional space in the future [12]. However, most studies are static measurements, and a few dynamic measurements are studies on calibration tubes, which cannot present dynamic dose distribution. This is a feasibility study in which NIPAM monomer-based gel dosimeter was combined with simulated dynamic displacement of organs caused by a concentric swing machine used in the laboratory. A linear accelerator was used for irradiation, MRI was used as a measurement tool, and MATLAB was used for readout analysis to evaluate whether NIPAM gel dosimeter could be used to simulate dose distribution during breathing under dynamic irradiation.

2. Materials and methods

2.1. Preparation of NIPAM polymer gel

The NIPAM gel dosimeter used in this study was provided by the laboratory of the department of medical imaging and radiological sciences of central Taiwan university of science and technology. Its
Table 1
The specification of drugs used in preparing the gel

| Drug   | Specification/purity                | Chemical formula | Manufacturer   |
|--------|------------------------------------|------------------|----------------|
| Gelatin| Type A from porcine skin, 300 bloom | C₁₀₀H₁₄₁O₃₉N₃₁  | Sigma-Aldrich  |
| NIPAM  | 97% pure                           | C₆H₁₁NO          | Sigma-Aldrich  |
| THPC   | 80% solution in water              | C₄H₁₂O₄PCl       | TCI            |
| BIS    | For electrophoresis                | C₇H₁₀N₂O₂        | Mreck          |

formula included 87% deionized water, 5% gelatin, 5% NIPAM monomer, 3% bisacrylamide (BIS), and 5 mM Tetrakis Hydroxymethyl Phosphonium Chloride (THPC) was used as oxygen scavenger (Table 1). After preparation, the gel dosimeter was wrapped in an aluminum foil to prevent light from inducing polymerization and the gel dosimeter was stored in a temperature-controlled 4°C refrigerator for it to solidify. The gel dosimeter was used for irradiation 24 hours after the preparation.

2.2. Simulation of organ motion by using the concentric swing machine

In this study, the concentric swing machine that is commonly used in the laboratory was used to simulate organ motion in the body. The swing machine can cause the slider to swing back and forth in a concentric manner. The minimum swinging speed was 20 rpm, which was 3 second/revolution after conversion, and this conformed to lowest respiratory conversion frequency of 12–20 breaths/minute in adults mentioned by Korin et al. in 1992 [6].

2.3. Gel irradiation

Irradiation in this study was divided into two parts. The irradiation device was Varian Clinic 21 iX (Varian Medical Systems, USA). Before irradiation, the gel was used as a substrate without addition of other materials (such as cross-linking agent, monomer) to prepare calibration tubes and gel dosimeter. Following that, a CT simulator was used to acquire CT images and the Varian Eclipse™ treatment planning system (TPS) was used to generate a radiotherapy plan and estimated MU values. The first part was gel calibration tube irradiation. The calibration tube was a 10 cm long and 1.5 cm diameter glass test tube containing NIPAM gel. The irradiation conditions used were: gantry angle: 0°, photon energy: 6 MV, dose rate: 500 cGy/min, and field size: 20 × 20 cm². The gel test tube was placed in the center of a 30-cm long, 30-cm wide, and 4-cm thick acrylic phantom with a 1.5-cm hole, and a 10-cm thick solid-state water phantom was placed at its bottom for irradiation. During irradiation, the source to surface distance (SSD) was 95 cm, source to axis distance (SAD) was 100 cm, and irradiation doses were 0, 1, 2, 5, 8, and 10 Gy. This data was used to construct a dose calibration curve. The second part of the study was gel dosimeter irradiation. An empty cylindrical bottle was filled with NIPAM gel and irradiation was carried out after the concentric swing machine used for dynamic simulation was turned on. The gantry angle was 0°, photon energy was 6 MV, irradiation dose was 5 Gy, dose rate was 500 cGy/min, field size was 4 × 4 cm², and the gel dosimeter was fixed on the concentric swing machine, as shown in Fig. 1. The SSD was 95 cm and SAD was 100 cm. Based on the estimated concentric swing machine simulation results (Fig. 2), it was predicted that dose was concentrated in a 2 × 2 cm² region (yellow region in the Fig. 2) under the ideal conditions. Therefore, 2 × 2 cm² and 3 × 3 cm² treatment plans were prepared in advance for subsequent comparison.

2.4. MRI readout and image reconstruction

After irradiation, the gel dosimeter was first placed in the MRI room for 24 hours so that the dosimeter
and examination room ambient temperature reached an equilibrium. The temperature of the examination room was controlled at 22°C–24°C. The MRI used for measurement in this study was the GE Healthcare-Brochure 1.5T MR450w-with-GEM and the coil used was Geometry Embracing Method (GEM) head and neck unit. The imaging pulse sequence used is fast spin echo (FSE T2WI), and the scanning parameters such as repetition time (TR) and echo time (TE) are listed in Table 2. The image acquisition site was 1 image at 5 cm below the top of the container. After scanning was completed, the images acquired were sent to Advantage workstation 4.6 (GE Healthcare, Inc) and PACS system. On one hand, the T2 Map software in Functool was used for post-processing. On the other hand, the image was exported and MATLAB on the personal computer was used for readout analysis.
Table 2
| Parameter                        | Value                                      |
|---------------------------------|--------------------------------------------|
| Repetition time (TR)            | 3,000 ms                                   |
| Echo time (TE)                  | 22, 44, 66, 88, 110, 132, 154, 176, 198, 220, 242, 264, 286, 308, 330, 352 ms |
| FOV                             | 240 mm × 240 mm                            |
| Slice thickness                 | 5 mm                                       |
| Slice interval                  | 0.5 mm                                     |
| Resolution                      | 512 × 512                                  |

2.5. Gamma evaluation

After image reconstruction, the results measured by NIPAM gel dosimeters were compared with the TPS calculated data. The quantitative gamma evaluation proposed by Low et al. [13,14] was used for performance analysis. The gamma evaluation criteria were distance-to-agreement (DTA = 3 mm) and dose difference (DD = 3%). These criteria are commonly used for clinical dose verification.

3. Results

3.1. Gamma passing rates

The gamma passing rates with gamma evaluation criteria of DTA = 3 mm and DD = 3% at TPS and measurement of MATLAB. Actual comparison of 4 × 4 cm² TPS and readout results found that the passing rate was 42.87%. The objective of this study was to evaluate feasibility and the phantom swinging method used was different from commonly used phantoms. Therefore, the ideal dose distribution in a swinging state was first simulated for evaluation. From Fig. 2, it can be seen that under the ideal conditions, dose was most concentrated in the 2 × 2 cm² (yellow block) when the 4 × 4 cm² field was swinging in a concentric circle. The actual measurement results showed that the 100% dose curve was also close to 2 × 2 cm². Therefore, the 3 × 3 cm² and 2 × 2 cm² TPS were made and used for comparison with the readout results. The passing rates were 54.96% and 49.92%, respectively.

3.2. Dose distribution and isodose line

Figure 3 shows the dynamics of 2 × 2 cm², 3 × 3 cm², and 4 × 4 cm² and the isodose distribution of TPS. Figure 4 shows the gamma map. The clinical evaluation standard is whether the gamma passing rate is greater than 95%, and the aforementioned results cannot be met. This study mainly observes the dynamic dose distribution. From Figs 3 and 4, it can be found that although the pass rate is not up to the standard, 2 × 2 cm² has a high passing rate in the clinically important dose concentration area. Further evaluation of the isodose line distribution results found that due to the concentric swing, the dynamic isodose line distribution is circular and cross-shaped at the dose curve ≥ 60% and < 60%, respectively, as shown in Fig. 3a. Therefore, adjust the ROI so that the comparison area is close to 60% of the isodose line and compare the result with the 2 × 2 cm² TPS. Figure 5 shows the result, the gamma passing rate = 97.92%, which meets the clinical evaluation standard.

3.3. Gamma map of the adjustment

Figure 6 shows the gamma map of the 2 × 2 cm² TPS after the MATLAB ROI adjustment and
Fig. 3. Comparison of readout result and TPS. The top to bottom figures show the comparison for (a) $2 \times 2$ cm$^2$, (b) $3 \times 3$ cm$^2$, and (c) $4 \times 4$ cm$^2$, respectively.
Fig. 4. Gamma map of readout result and TPS. The top to bottom figures show the 3 mm, 3% gamma map for (a) $2 \times 2$ cm$^2$, (b) $3 \times 3$ cm$^2$, and (c) $4 \times 4$ cm$^2$, respectively.
Fig. 5. Comparison of readout results and dose curve at ≥ 60% for TPS 2 × 2 cm².

Fig. 6. 3 mm, 3% gamma map of readout results and dose curve at ≥ 60% for TPS 2 × 2 cm². From the figure, it can be seen that the rejected area mostly forms a cross-shaped dose region due to the concentric swing machine.

measurement. It can be seen from the figure that the rejected area is mainly the dose curve < 60%. As mentioned earlier, the concentric swing causes the dose curve to begin to appear cross-shaped, thereby reducing the passing rate.

4. Discussion

4.1. Selection of suitable readout tools

The main objective of this study was to examine the feasibility of clinical application of NIPAM gel dosimeter when irradiated under continuous motion. De Deene et al. proposed a complete insight on
how to use MRI to read out a gel dosimeter. Following their recommendation of multiple spin-echo in 2010 [15], they proposed a more complete scanner readout guideline in 2013, including selection of suitable gel dosimeters and MRI imaging sequence parameters, temperature control during gel dosimeter preparation, storage, and scanning, and comparison with treatment plan results [16]. Although there are scanning time and temperature control problems when MRI is used for gel dosimetry, MRI also has many advantages when compared with other readout techniques such as multiple options for parameter sequences, high resolution, and diverse changes in phantom container volume. Therefore, we believed that selection of suitable readout instruments should be one of the guidelines for gel dosimetry measurements.

4.2. Selection of optimal parameters

Detailed design is required when using MRI for gel dosimetry. In this study, multiple spin-echo was used for image scanning based on the recommendations of past papers, and T2 weighted image readouts were obtained under different TE conditions for the same TR. Following that, the MATLAB program was used to measure the pixel signal intensity in the area of interest and signal intensity was directly proportional to image brightness. Increase in TR increased the clarity of the phenomenon of protons jumping from a stable state to an excited stable, thereby increasing signal intensity and decreasing noise production. However, a disadvantage was that this increased scanning duration, thereby increasing gel temperature. In their article published in 2000, De Deene et al. mentioned that temperature elevation when MRI was used for gel dosimetry caused dose to be underestimated [17]. Therefore, TR was set as 3000 ms and only one slice of Dmax was acquired to decrease scanning time and temperature elevation. In addition, longer TE caused differences in gel that were irradiated with different doses and more TE sampling tended to lead to actual T2 attenuation. However, the shortcoming was that it increased total scanning duration. The data requirements of the post-processing workstation used in this study limited minimum TE to be 4 and the software recommended a TE of 8 to more accurately measure the T2 of the material. In order to obtain better measurement image results, we set up 16 sampling TE. Although scanning duration was longer, the calibrated 3 mm, 3% gamma passing rate was 97.92%, which met the clinical criteria of > 95%.

4.3. Dynamic dosimetry in previous papers

In 2015, Hsieh et al. used a calibration tube for dynamic and static dosimetry measurement comparisons in a study that used NIPAM gel dosimetry and results were consistent [18]. They employed an experimental method which placed the calibration field in the center of the field, such that the gel dosimeter was still in 100% of the irradiation area of dose distribution even under dynamic conditions. Therefore, the differences in measurements between the two conditions were not large. The distribution of the dose curve must be completely measured in order to further confirm its clinical applicability and compared with the treatment plan to completely evaluate whether the NIPAM gel dosimeter was suitable for dose analysis of dynamic phantoms. In this study, concentric swinging was used for gel dosimetry, which was different compared with common dynamic phantom swinging methods, so it was difficult to compare with treatment plan results. Therefore, pre-simulation was carried out followed by measurement comparison to estimate the differences between dynamic and static states when the gel dosimeter was used in actual clinical practice. The evaluation results found that the readout result and isodose curve in the treatment plan were similar in the high-dose region and gamma passing rate also met the clinical requirements.
4.4. Measurement results of other gel dosimeters

Thomas et al. published a paper in 2013 on employing 3D dosimetry to observe the effects of organ motion on radiotherapy and evaluated its interaction [19]. Their study used the PRESAGE gel dosimeter combined with CIRS dynamic phantom for dosimetry and observed two types of treatment plans, namely the intensity-modulated radiation therapy and the arc therapy (RapidArc). The study results found that the dose volume histogram (DVH) coverages for gross tumor volume (GTV) and clinical target volume (CTV) were similar and there were large differences in DVH coverage between internal target volume (ITV) and planning target volume (PTV). The authors believed that inaccuracy caused by organ motion could be minimized by considering leaf speed of the multileaf collimator, gantry rotation of the device, and breathing period. In this study, 3 mm, 3% gamma passing rate that is commonly used today was used for comparison. Predictions using concentric swing machine pre-simulation found that the GTV/CTV coverage was similar and ITV/PTV coverage difference was large. In addition, the gamma passing rate of dose curves $\geq 60\%$ and $< 60\%$ were different through selection of suitable comparison regions.

4.5. Study limitations and future developments

Although this study proved that it was feasible to use NIPAM gel dosimeter for dynamic dosimetry, the swinging method was different from commonly seen dynamic phantoms and the field used was not large, which differed from the field that is commonly used in clinical treatment. Therefore, we recommend that commonly used dynamic phantoms could be used in future studies and attempts should be made to quantify dose differences under dynamic states. Furthermore, although MATLAB is a commonly used program, it is not easy to write and apply the program in dose readout measurements. Regardless of whether MRI, OCT, or CT is used, subsequent readout analysis is required in addition to parameter setting for the device. Hence, developing a stable and complete analysis program should be one of the focus for consideration. In addition, although the NIPAM gel dosimeter has strengths in presenting three-dimensional dose distribution, it is still not widely used in clinical practice currently. Furthermore, gel dosimetry is a type of relative dosimetry. Besides comparing with the treatment plan results, we recommend comparing with other dosimetry tools that are commonly used in clinical practice, such as films and free-air ionization chambers to increase the reliability of the reference. Changes in dynamic phantom parameters, such as operating range and frequency, could also be added to obtain more data. We believe that this can be a direction for consideration for future researchers who are interested.

5. Conclusion

Gel dosimetry has the advantage of providing three-dimensional relative dose distribution and the use of dynamic phantom in simulation of dose measurement in organ motion has clinical reference value. Therefore, the combination of the two can simulate the distribution of clinical dynamic dose. Many previous studies on the NIPAM gel dosimeter used static treatment target zones. In this study, we carried out pre-simulation and comparison to confirm that it was feasible to employ NIPAM gel dosimeter and concentric swing machine in simulation of dynamic dose measurement. Appropriate selection and comparison was carried out and it was found that $\geq 60\%$ of dose curves were consistent with the simulation results and the 3 mm, 3% gamma passing rate was 97.92%, which met the clinical criteria of $> 95\%$. We recommend that common dynamic phantoms can be used and other parameter settings can be added in the future, and comparison with dosimetry tools commonly used clinically can be carried out to increase the reliability of the reference.
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Conflict of interest

None to report.

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