



Assessment of Temporary Radioactivation for Tissue Expanders in Breast Radiation Therapy: Preliminary Study

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ABSTRACT

Background: As breast tissue expanders consist of metallic materials in the needle guard and ferromagnetic injection port, irradiation can produce radioactivation.

Materials and Methods: A CPX4 (Mentor Worldwide LLD) breast tissue expander was exposed using the Versa HD (Elekta) linear accelerator. Two photon energies of 6 and 10 MV-flattening filter free (FFF) beams with 5,000 monitor units (MU) were irradiated to identify the types of radiation. Furthermore, 300 MU with 10 MV-FFF beam was exposed to the CPX4 breast tissue expander by varying the machine dose rates (MDRs) 600, 1,200, and 2,200 MU/min. To assess the instantaneous dose rates (IDRs) solely from the CPX4, a tissue expander was placed outside the treatment room after beam irradiation, and a portable radioisotope identification device was used to identify the types of radiation and measure IDR.

Results and Discussion: After 5,000 MU delivery to the CPX4 breast tissue expander, the energy spectrum whose peak energy of 511 keV was found with 10 MV-FFF, while there was no resultant one with 6 MV-FFF. The time of each measurement was 1 minute, and the mean IDRs from the 10 MV-FFF were 0.407, 0.231, and 0.180 $\mu\text{Sv/hr}$ for the three successive measurements. Following 10 MV-FFF beam irradiation with 300 MU indicated around the background level from the first measurement regardless of MDRs.

Conclusion: As each institute room entry time protocol varies according to the working hours and occupational doses, we suggest an addition of 1 minute from the institutes' own room entry time protocol in patients with CPX4 tissue expander and the case of radiotherapy vaults equipped with a maximum energy of 10 MV photon beams.

Keywords: Breast Radiation Therapy, Breast Tissue Expander, Ferromagnetic Material, Radioactivation

Original Research

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Introduction

For most women who undergo total mastectomy, breast reconstruction is a preferable option with the advantages of relative simplicity, low morbidity, and cosmetic outcomes [1, 2]. The most widely used reconstruction process involves two stages: temporary tissue expansion and permanent implantation [3], accounting for approximately 61% of all breast reconstruction procedures in the United States [4]. Tissue expansion is intended to reserve sufficient space by stretching the skin and muscles when excessive

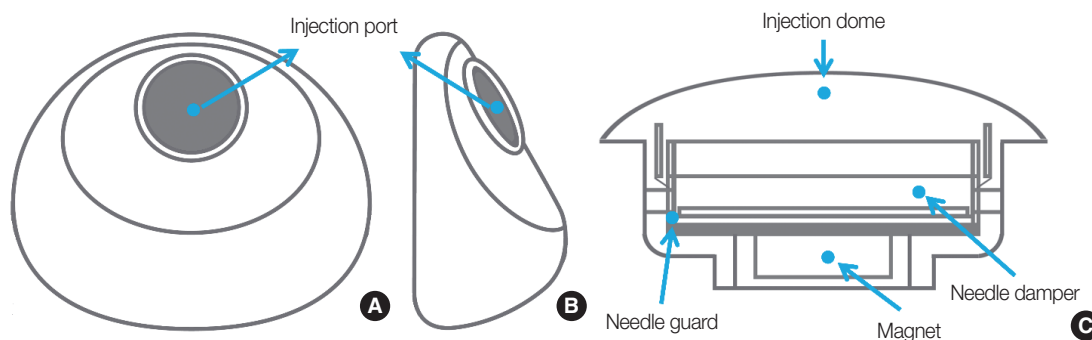


Fig. 1. Projection view of CPX4 (Mentor Worldwide LLD) breast tissue expander for (A) side view, (B) frontal view, and (C) subcomponents of the injection port.

resections are made without sufficient breast tissue [5]. The breast tissue expander is a balloon-like inflatable device with a silicone outer shell and a superficial port [6]. Once a tissue expander is inserted during the surgical stage, saline is injected on a regular basis [6]. Once they attain the desired shape and size, the tissue expander is removed and replaced with permanent saline or silicone breast prostheses [6].

Nowadays, breast radiation therapy is usually administered by an intensity modulated radiation therapy technique using low-energy photon beams (usually around 6 MV) because this enables a substantial reduction in toxicity while providing sufficient target coverage [7, 8]. However, it has been reported that patients who received postmastectomy radiotherapy (RT) in the presence of tissue expanders tend to be less effective and have an increased risk of complications than those who received permanent implants [9] or autologous tissue reconstruction [10]. One reason for this is the perturbation of the dose distribution, and this is mainly because the extremely high-density of the metallic port of the tissue expander causes a discrepancy between the calculated and measured dose distribution [11–15]. There are metallic materials in the needle guard and ferromagnetic injection port of the CPX4 breast tissue expander (Mentor Worldwide LLD), consisting of 316 L stainless steel and neodymium, respectively. Although it varies with the Hounsfield unit modeling coverage of the high-density materials in the treatment planning system, it has been reported that the amount of dose mismatch can be mitigated using higher photon (> 10 MV) energies along with a build-up compensating bolus [15–17]. Furthermore, metal artifacts in computed tomography images caused by metallic components often hinder the correct segmentation of the breast and adjacent tissues [14, 18]. As distinct from the above-mentioned issues, there is a possibility of temporary radioactivation (TRA) induced from

Table 1. Physical Dimensions of Components Consisted of CPX4 Breast Tissue Expander

Variable	Materials	Diameter (cm)	Thickness (cm)
Injection dome	Cured silicone	5.385	0.838
Needle damper	Thermoplastic	3.493	0.051
Needle guard	Stainless steel (316 L)	3.650	0.025
Magnet	Neodymium	1.270	0.475

the tissue expander with radiation exposures. Radioactive nuclides are generated when high-energy photon beams irradiate high-density materials owing to photonuclear reactions [19]. Although many studies have focused on the radioactivation of high-atomic number materials in the RT vault [20–23], to the best of our knowledge, no previous studies have demonstrated the TRA of the tissue expander.

In this study, we utilized a portable radioisotope identification device (RIID) to identify the types of radiation and measured the instantaneous dose rates (IDRs) when the CPX4 breast tissue expander was irradiated with respect to the machine dose rate (MDR) and incident energy. We also identified the duration at which the measured dose was below a safe level.

Materials and Methods

1. Metallic Injection Port on Tissue Expander

As shown in Fig. 1, a CPX4 breast tissue expander consists of an injection dome, needle damper, needle guard, and magnet area. The physical properties of the CPX4 breast tissue expander components are summarized in Table 1.

2. Experiment Design

Recently, breast RT is widely conducted with hypofractionation of which a daily dose of 2.7 Gy, although it could

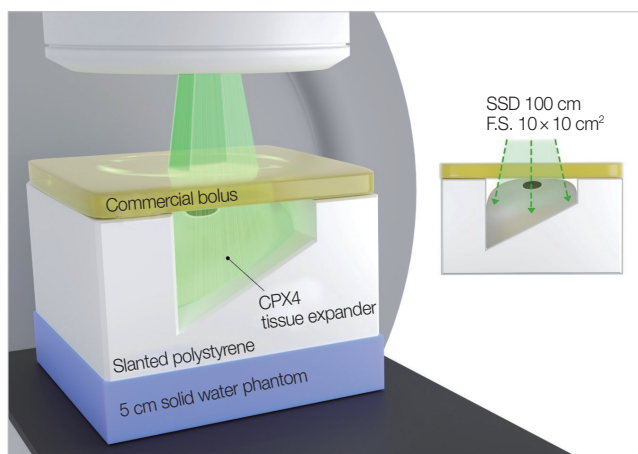


Fig. 2. Experimental setup to evaluate the types of radiation and to measure instantaneous dose rates after irradiations to CPX4 (Mentor Worldwide LLD) breast tissue expander. SSD, source to surface distance; F.S., field size.

vary according to the pathological stage of the tumor [24, 25]. To cover a common daily dose, we designed to deliver 3 Gy to the surface of the breast tissue expander. Two photon energies of 6 and 10 MV-flattening filter free (FFF) beams from the Versa HD linear accelerator (LINAC; Elekta) were used in this study. We used an FFF beam rather than a beam with flattening filter to assess the IDR according to the MDR.

The experimental setup is depicted in Fig. 2. A 5-cm solid water phantom was placed on the table-top for backscatter and slanted polystyrene to adjust the tilted geometry of the CPX4 breast tissue expander. Slanted polystyrene was used to make the beam incidence perpendicular to the tissue expander, thus delivering a uniform dose. Above a tissue expander, the energy-dependent thickness of the commercial bolus (1.5 cm for 6 MV-FFF and 2.5 cm for 10 MV-FFF) was also placed for build-up. The source-to-surface distance was set to 100 cm, and the field size was set to $10 \times 10 \text{ cm}^2$.

To demonstrate the types of radiation according to the energy, 5,000 monitor units (MU) were delivered with 6 and 10 MV-FFF. To demonstrate the TRA according to the MDR, 300 MU with 10 MV-FFF beams were irradiated by varying the MDRs of the 600, 1,200, and 2,200 MU/min.

3. Measurement of Radioactivation

A RIIDEye X-GN detector (Model 42508/85; Thermo Fisher Scientific Messtechnik GmbH) was used to detect the presence of any types of radiation. The detector was calibrated using the test sources ^{137}Cs , ^{226}Ra , and ^{252}Cf , and each peak energy was all within the tolerance limits ($\pm 2\%$ for ^{137}Cs and

^{226}Ra , and $\pm 20\%$ for ^{252}Cf). The RIIDEye X-GN consisted of a $5.08 \text{ cm} \times 5.08 \text{ cm}$ NaI gamma detector with a typical resolution of 7.5% at 662 keV and detectable gamma energy ranging from 25 keV to 3 MeV. Furthermore, an $18 \text{ mm} \times 34 \text{ mm}$ crystal was internally separated from the gamma detector, which enabled the detection of neutron count rates up to 2,000 counts per second. Data from all measurements were stored in Ansi N42 format. When all experiments were completed, RIIDView software version 1.0 (Thermo Fisher Scientific Messtechnik GmbH) was used to analyze IDR, measured energy spectrum, and isotopes estimated using a 256 channel-based quadratic compression conversion algorithm. Additionally, another software, InterSpec version 1.0.11 (National Technology & Engineering Solutions of Sandia LLC), was used because the RIIDView software does not support exporting raw spectrum data. This software supported the export of each spectrum, thus facilitating direct comparisons of the energy spectrums across measurements. Based on the spectrum, the software suggests the most well-matched nuclide by searching for it in a predefined nuclide database.

Prior to beam irradiation, the RIIDEye X-GN underwent background correction. After beam irradiation, a tissue expander was placed outside the treatment room to assess the IDRs solely from the CPX4 breast tissue expander not from the LINAC head, and the TRA was measured for 60 seconds. If any nuclides were detected or the IDR was measured to be higher than the background level, the measurement continued until there were no nuclides or the IDR decreased to the background level. All measurements were repeated twice to demonstrate the statistical uncertainties.

Results and Discussion

In general places far away from the treatment room, RIIDEye X-GN showed a mean IDR of $0.181 \mu\text{Sv/hr}$, which was set as the background level. As shown in Fig. 3, the IDRs after delivering 5,000 MU with 10 MV-FFF were 0.407 ± 0.011 , 0.231 ± 0.001 , and $0.180 \pm 0.004 \mu\text{Sv/hr}$ for three sequential measurements. The IDR reached the background level except for the first measurement. Fig. 4 presents the first measured energy spectrum after 5,000 MU delivered for 10 MV-FFF showing negligible counts within 3 minutes. There was notable energy spectrum whose peak energy of 511 keV was found with 10 MV-FFF, while there was no obvious one with 6 MV-FFF. No neutron fragments were detected in either case.

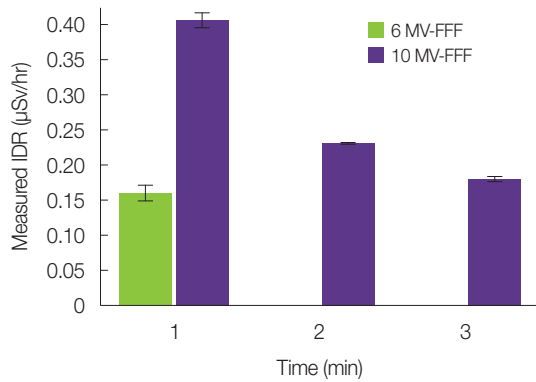


Fig. 3. Measured instantaneous dose rates (IDRs) according to the energy after 5,000 monitor units irradiation to CPX4 (Mentor Worldwide LLD) breast tissue expander. FFF, flattening filter free.

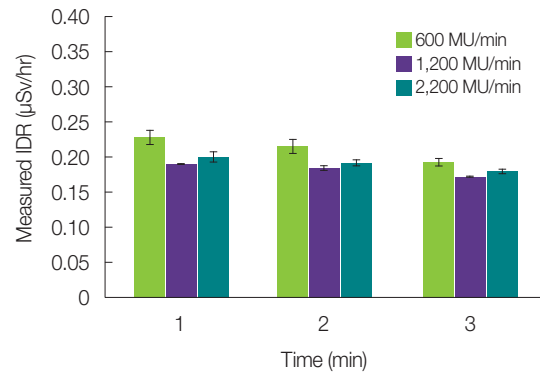


Fig. 5. Measured instantaneous dose rates (IDRs) according to the machine dose rate after irradiation to CPX4 (Mentor Worldwide LLD) breast tissue expander. MU, monitor unit.

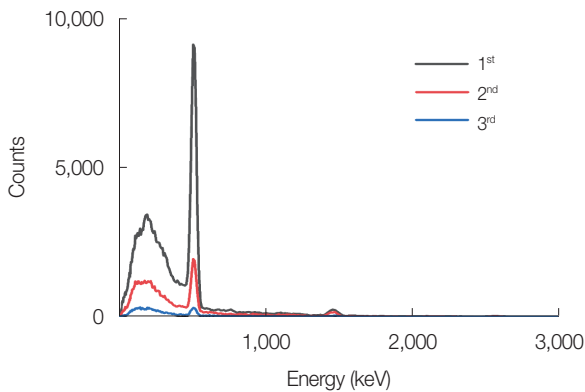


Fig. 4. Three successive energy spectrum measurements after 5,000 monitor units delivery with 10 MV-flattening filter free photon beam to CPX4 (Mentor Worldwide LLD) breast tissue expander.

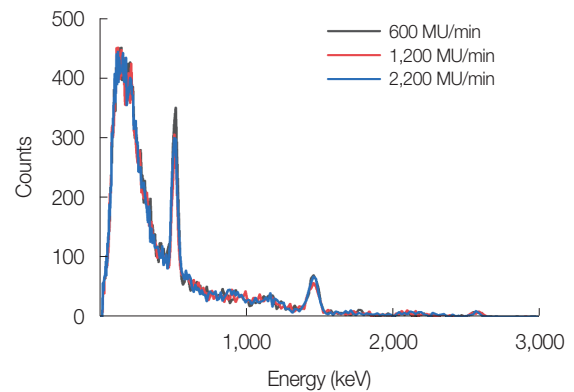


Fig. 6. Measured energy spectra after 300 monitor units (MU) beam irradiation to CPX4 (Mentor Worldwide LLD) breast tissue expander with 10 MV-flattening filter free photon beams according to the dose rates.

The IDR measurements with 300 MU according to MDR are presented in Fig. 5, and are shown around the background level from the first measurement. Furthermore, it was not significantly affected by the MDR (600, 1,200, and 2,200 MU/min). No neutron fragments were detected in either experiment. Although a peak energy of 511 keV was detected when 300 MU was delivered to the CPX4 tissue expander, the corresponding counts were negligible, as shown in Fig. 6.

It is a matter of course that the measured IDRs from a tissue expander was much lower than that of LINAC head. Previous research has demonstrated that the IDR was 4.14 $\mu\text{Sv/hr}$ immediately after 1,000 MU delivery with 15 MV, and decreased to 0.65 $\mu\text{Sv/hr}$ after 5 minutes [26]. Another study showed that the measured dose rate was 0.702 $\mu\text{Sv/hr}$ right after 500 MU delivery with 10 MV, and decreased to 0.406 $\mu\text{Sv/hr}$ after 1 minute [27]. Although it could vary depending on the LINAC products and incident energy, it is definitely greater than those

from tissue expanders. The LINAC head-generated neutrons are mainly due to (γ, n) reactions, and the produced neutron might cause nuclear reactions itself and the tissue expanders [28, 29]. Although the half-life of a single neutron in air is microseconds, the radioactivity remains and should be considered its accumulation [30]. Even though the specific activation process is unknown because of its multicomponents of tissue expanders, authors could assume that the majority of activation was induced from high-density materials, i.e., stainless steel and neodymium magnet, whose physical densities are 8.1 and 7.01 g/cm^3 , respectively.

Furthermore, the mean occupational radiation doses for medical fields in South Korea and those from the United Nations Scientific Committee on the Effects of Atomic Radiation annual report in 2020 were 0.38 and 0.57 mSv, respectively, which could be regarded as keeping well as low as reasonably achievable practice when applying occupational dose

limit [31–33]. Referring to both IDRs from LINAC head and national occupational radiation dose statistics, tissue expander-induced IDRs may not be a great concern because even slightly higher IDRs of 0.392 $\mu\text{Sv/hr}$ was caused by extremely high MU delivery. Moreover, the prescription greater than 10 Gy could not be a real clinical situation. Even if hypofractionated RT to breast and supraclavicular lymph nodes often employs monitor units greater than 1,000 MU [34], the MU should not be evaluated alone because the delivered dose was determined not only by the amount of MU but also by the other beam parameters such as field size and source-to-axis distance [35]. Despite the 5,000 MU delivery with a 10 MV-FFF photon beam with a field aperture of 10 cm \times 10 cm, the IDR was reduced to the background level after 1 minute.

The optimal room entry time was discussed from the literatures and it varied a lot [30, 36]. As they varied upon the number of patients and the proportion of treatment sites related to the used energy, each institute established the room entry time protocols considering both working hours and occupational equivalent doses.

Figs. 4, 6 indicate that while 10 MV irradiation to the CPX4 tissue expander could generate an energy spectrum whose peak energy was around 511 keV, this corresponded to the background level within 1 minute. In this regard and considering that the experiment was performed outside the treatment room, an additional delay time of 1 minute from institutes' own protocols after treatment of patients with CPX4 tissue expander could guarantee that there is little harm to RT staff, and it was not influenced by the amount of irradiation dose. There was no notable energy peak with 6 MV-FFF regardless of the dose, indicating that no additional delay time due to the use of tissue expanders is necessary for the RT staff.

This study had several limitations. First, other kinds of tissue expanders exist. Tissue expander products from the same company (Mentor Worldwide LLD) vary according to size, shape, and product type. We expect that in-depth experiments, including other kinds of tissue expanders with different sizes, shapes, and constituent materials, could provide insights into clinical protocol establishment or radiation safety perspectives. Second, the measurements could not be performed with 15 MV in this study because of the absence of 15 MV with this LINAC. While the use of a 15 MV photon beam is known to reduce dose heterogeneity and improve dose coverage in patients with tissue expanders [11, 16], it is also known that photoneutron generation and radioactivation

are more prevalent at higher photon energies. Any institute with LINACs equipped with 15 MV or higher should verify the types of isotope, amount of radioactivation, and their influence on energy and dose rate in advance treatments for patients with breast tissue expanders. Nonetheless, our study seems significant from a clinical protocol establishment or radiation safety perspective because the study is the first attempt to reveal the potential of TRA when performing breast RT in patients with tissue expanders.

Conclusion

This study demonstrated TRA of the CPX4 tissue expander when the expander was irradiated. While irradiation of 6 MV-FFF beams to the CPX4 tissue expander generated no resultant nuclides, those with 10 MV-FFF beams produced a spectrum with peak energies of 511 keV. However, experiments with 3 Gy delivery, exhibited IDRs of around background level within 1 minute, regardless of the MDR. In patients with CPX4 tissue expander and the case of RT vaults equipped with a maximum energy of 10 MV photon beams, the staffs are recommended to enter the treatment room by adding at least 1 minute from their own protocols because a room entry time protocol varies according to each institute.

Conflict of Interest

No potential conflict of interest relevant to this article was reported.

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Ethical Statement

This article does not contain any studies with human participants or animals performed by any of the authors.

Author Contribution

Conceptualization: Lee H. Methodology: Lee H, Chun M. Data curation: Lee H, Yoo L. Formal analysis: Lee H, Oh DH,

Chun M. Supervision: Lee H, Oh DH, Chun M. Funding acquisition: Chun M. Project administration: Chun M. Investigation: Lee H, Oh DH, Yoo L, Chun M. Visualization: Yoo L, Chun M. Validation: Lee H, Chun M. Writing - original draft: Lee H, Chun M. Writing - review and editing: Oh DH, Chun M. Approval of final manuscript: all authors.

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