TECHNICAL NOTE

Development of an in vivo technique for dose verification at the prone breast board / skin interface

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Abstract

Due to the limited height of commercial prone breast boards, large or pendulous breasts may contact the base layer of the board during simulation and throughout the course of treatment. Our clinic has historically identified and marked this region of contact to ensure reproducible setup. However, this situation may result in unwanted hotspots where the breast rests atop the board due to electron scatter. In this study, we performed in-vivo dosimetric measurements to evaluate the surface dose in regions of contact with the immobilization device. The average dose and hotspot were identified and evaluated to determine whether plan modifications were necessary to avoid excess skin toxicity at the skin/breast board interface. The film method results were validated against a commissioned in vivo OSLD dosimetry system. Radiochromic film measurements agreed with OSLD readings (n = 18) overall within 1%, σ = 6.4%, with one deviation of >10%. Pertinent information for the physician includes the average, maximum, and minimum doses received at the film interface. Future readings will not require OSLD verification. Physicians now have access to additional spatial data to correlate skin toxicity with doses delivered at the skin/breast board interface. This new technique is now an established procedure at our clinic, and can inform future efforts to model enhanced methods to calculate the dosimetric effects from the prone breast board in the treatment planning system.

KEY WORDS

breast board, in vivo dosimetry, prone breast

1 | INTRODUCTION

Radiation therapy is an essential component of postlumpectomy treatment for breast cancer patients and can be delivered with patients either in the supine or prone position. Treatment in the prone position takes advantage of gravity and allows the breast, and hence the target, to fall away from the chest wall, ipsilateral lung, and heart. Prone breast tangent fields generally avoid primary beam interaction with the prone breast board; however, for patients where the breast contacts the base layer of the prone breast board, the primary beam must pass through the base layer to treat the entire breast (Fig. 1). This results in largely unaccounted for electron scatter that usually increases the surface dose and is difficult to accurately model in the treatment planning system (TPS). An in vivo dose measurement at the prone breast board/skin interface is warranted to accurately assess the dose to the surface of the breast.

The American Association of Physicists in Medicine (AAPM) Task Group (TG)-176 recommends fully modeling couchtop structures and...
immobilization devices in the TPS or to avoid irradiating through couchtop immobilization devices.\textsuperscript{1} Several subsequent studies have investigated dosimetric impacts of the prone breast board in general.\textsuperscript{2-4} While previous studies have investigated loss of skin sparing due to contact with the prone breast board, they are not sufficient to evaluate and address the clinical problem described here. A recent study rigorously measured couch rail and prone breast board transmission factors,\textsuperscript{3,4} finding that a clinical prone breast case generally receives less dose than the TPS predicts in the absence of proper modeling.\textsuperscript{3,4} Film studies were performed primarily through the sides of the prone breast board and directly through the couch structure. Surface dose measurements were underpredicted by the TPS on the order of 25–35%.\textsuperscript{3,4} These studies were performed in a phantom and used ion chamber measurements in a solid water block, but were not performed \textit{in vivo} and did not measure surface dose against the anterior contact surface of the breast, focusing instead on the medial edge of the breast at the chest wall (Fig. 2). Further work has been limited to radiochromic film measurements at the medial edge of the prone breast board in phantom\textsuperscript{5} and a case study using an anthropomorphic phantom to measure contact with the inferior and medial edge contact points near the inframammary fold.\textsuperscript{6}

The purpose of this technical report is to describe our method for measuring dose across the contact plane between the breast and the base layer of the prone breast board. The report also describes our verification of this method using an established \textit{in vivo} dosimetry system. The procedure detailed below is now clinically implemented for all prone breast patients whose treated breast contacts the base layer of the prone breast board. The information from this evaluation serves either to assure the physician that the treatment plan sufficiently minimizes risk of skin toxicity, or to guide the physician to request a new treatment plan designed explicitly to reduce the surface dose while maintaining all other planning goals.

\section{Materials and Methods}

During CT simulation for prone breast treatments in our clinic, we identify areas where the breast is in contact with the base layer of an Access ClearVue prone breast board (Qfix, Avondale, PA) by indexing a piece of white printer paper to the base layer under the breast (Fig. 3). Once the breast is positioned in a reproducible location, the contact plane is traced with a marker and the indexed position is recorded. The setup trace paper is in place for all fractions to guide radiation therapists toward a reproducible setup. Image guidance is performed for the first fraction, then every five subsequent fractions.

For this study, we replaced the setup trace paper with Gafchromic EBT3 (Ashland, Bridgewater, NJ) radiochromic film. Prior to the patient’s second fraction, the physicist copies the entire setup trace onto a piece of EBT3, which is then used by the radiation therapists in place of the setup paper. We chose fraction #2 for dosimetric evaluation to eliminate dose contamination from pretreatment imaging. Two strips are cut from the short edge of the film prior to irradiation for use with the one-scan protocol.\textsuperscript{7} One strip is irradiated to 120% of the expected dose to the film surface as predicted in the TPS, and the other is left unirradiated. Both film strips and the treatment film are scanned into an Epson 10000XL scanner set to 0.0 focus, 48-bit depth, and 72DPI resolution. Analysis is performed in FilmQA Pro (Ashland, Bridgewater, NJ) calibrated with tri-color optimization. Care is taken throughout the process to ensure that films and strips are scanned in a consistent alignment. All films are scanned the day after treatment to ensure that sufficient time was given for the film to polymerize.

![Figure 1](image1.png)

\textbf{Fig. 1.} “Prone Breast CT Slice.” Slice of prone breast treatment plan, demonstrating an intended reduction in absorbed dose near the breast board/skin interface.

![Figure 2](image2.png)

\textbf{Fig. 2.} “Prone Breast Board.” Prone breast board used in this study. This study measures the enhanced skin dose from the anterior contact surface of the breast with the base layer (solid green arrow) as opposed to the medial contact surface (dashed red arrow). Breast board image from Qfix.\textsuperscript{10}
A report is created for the physician with a custom standardized isodose map that is designed to be as similar as possible to the default isodose lines in the TPS (Fig. 4). The report highlights the average dose and maximum point dose, given the scanned resolution (~0.3 mm × 0.3 mm). These doses are displayed as a percentage of the prescribed dose to the breast.

This methodology was validated against our established in vivo dosimetry program, which uses nanoDot (Landauer Inc, Glenwood, IL) Optically Stimulated Luminescent Dosimeters (OSLDs). For this validation we taped one to four OSLDs, based on breast contact plane size, to the radiochromic film. Small opaque marks were made on the film at the three nondetecting corners of each OSLD (Fig. 5). Point doses measured in the detecting area were compared to the film doses. Our OSLD system was commissioned in accordance with the high-accuracy method of AAPM TG-191, except that our angular correction factors ($k_\theta$) were measured in phantom to be 1.03 and applied manually. OSLD standards were given the same dose as the irradiated film strip (above). To provide sufficient validation of the film dosimetry technique, the first seven instances of this protocol included OSLD verification measurements. It should be mentioned that any clinical decisions made as a result of this protocol were drawn from the established OSLD readings and not from the novel film measurements listed in Table 1.

3 | RESULTS AND DISCUSSION

We performed this evaluation eight times on seven patients. The first patient was a proof-of-concept without OSLDs that does not resemble the other measurements, so it is excluded from this study.
One patient was replanned and remeasured, leading to a total of seven readings from six patients used for the analysis presented here. These seven readings resulted in a total of 18 OSLD data points and seven sheets of film.

Table 1 summarizes the validation of the film method, noting the replanned measurement with \(^a\). A direct comparison of film readings to OSLD readings is provided — a value greater than 1 demonstrates a film measurement that is greater than what was measured on an OSLD at the same location. The mean film result is within 0.2% of the mean OSLD result, with a standard deviation of 6.4%. All film-to-OSLD comparisons were within the stated error of unscreened nanoDot OSLDs (10%) except for one measurement in patient 4, which skewed the average for that patient above 10%. Patient 4 is the only subject with film measurements that exceed 8% deviation from the corresponding OSLDs.

Hotspots found in the film are listed as a percentage of the prescription dose. Dose was prescribed to the entire breast volume and varied from 180 to 266 cGy/ fx. Film hotspots were found to be higher than the prescription dose in six of seven cases. In the patients marked with \(^b\), a field-in-field technique was used to shield the breast board/skin interface to maximum of 95% of the prescription dose. For example, reading 2 directly applies field-in-field shielding to the interface to the plan from reading 3, leading to a 6% of prescription improvement in the hotspot. Despite this effort, the average hotspot observed on the films was 112% of prescription.

This is a troubling result without an obvious cause — it is possible that our method of prone breast treatment planning is not well modeled in the TPS due to the heterogeneity interface between the breast board and the breast. Another working theory was that it was somehow correlated with CTV volume, but we found no correlation.

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**AUTHOR CONTRIBUTION STATEMENT**

Dr. Loughery: Lead author of manuscript, physics data collection and analysis, corresponding author. Dr. Chan: Coauthor of manuscript, clinical data collection and analysis. Dr. Burmeister: Study design, coauthor of manuscript, data analysis, and physics expertise. Dr. Dominello: Lead study design, data analysis, senior author of manuscript, and clinical expertise.

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