Development of a treatment technique for the AeroForm™ tissue expander breast implant system—A single department experience

Abstract

Purpose: Air tissue expander systems offer breast cancer patients a number of advantages but present an unusual challenge when treating patients post-mastectomy with external beam radiation therapy. The primary aim of this study was to describe the planning protocol developed for external beam locoregional radiation therapy (RT) in post-mastectomy patients with in situ air expanders, and to quantify the effect of the AeroForm™ (AirXpanders®, Palo Alto, California) implant on target volumes and treatment factors. Secondary aims included an evaluation of clinical outcomes including acute side-effects.

Methods and materials: This is a single institution review of eight female patients treated between March 2016 and August 2018. All patients received a prescription of 5000 centigray (cGy) in 25 treatments delivered five times per week. Specific density overrides were applied to critical structures in order to achieve the most accurate dose distribution. Acute side-effects were assessed at weeks two, four and six during radiation therapy, and followed-up at week six after the final treatment.

Results: A three-dimensional (3D) conformal radiation therapy (CRT) tangential wedged-pair technique, with additional contouring and density overrides applied to critical structures, is suitable for patients with breast tissue expanders. 5mm–10mm of bolus over the entire chest wall throughout treatment enhances treatment delivery accuracy to the superficial skin layers without increasing early acute skin reactions.

Conclusion: It is feasible to plan radiation therapy in patients with air tissue expanders requiring chest wall (with or without nodes) irradiation using an appropriate planning technique and density overrides applied to critical structures. The addition of bolus to the entire chest wall increases dose distribution accuracy. Early clinical results show that this treatment method is reproducible, has acceptable early toxicity, and results in similar acute toxicity to those seen in patients without air expanders. Our air expander protocol will continue to be adapted as additional prospective data is collected.

Keywords: radiation therapy, breast cancer, mastectomy, air tissue expanders, planning

Introduction

Breast cancer is the most common cancer in women worldwide. With nearly 1.7 million new cases diagnosed in 2012, it is the second most common cancer overall.1 Post-mastectomy radiation therapy (PMRT) to the chest wall and locoregional nodes has been shown to reduce the incidence of locoregional and distant recurrence, and improve overall survival in patients with node positive breast cancer. These same benefits from PMRT were also seen in women with one to three positive nodes treated with axillary dissection and systemic therapy.2,3

Reconstruction can have lasting effects on women’s quality of life,4–7 particularly as the majority of these women will be long-term survivors of breast cancer. In a recent meta-analysis, patients undergoing breast reconstruction were found to benefit psychologically, having less anxiety and depression compared to those who undergo mastectomy alone.4 Little is known, however, about the psychological benefits of immediate versus delayed breast reconstruction, and data from available studies is conflicting. It is generally felt that there might be an advantage in choosing immediate reconstruction if it can be done safely from an oncological perspective.8

Tissue expander systems enable suitable patients to begin the reconstruction process immediately after mastectomy. Traditional expanders use a saline-filled implant that is expanded over a number of months via percutaneous injections of saline solution into the implant.10 A newer air tissue expander system, the AeroForm™ implant (AirXpanders®, Palo Alto, California), is a needle free, patient-controlled, wireless device which eliminates expansion visits to the clinic and provides faster expansion compared to saline-filled expanders.10 The device consists of an implantable silicone tissue expander containing a metal reservoir of compressed carbon dioxide (CO₂). It is programmed to allow expander filling via an external hand-held remote control of up to 10 cubic centimetres (cc) of CO₂ three times per day in three-hourly intervals to a maximum dose of 30cc per day, and stops expanding once capacity is reached.10

Concerns about the morbidity of implant reconstructions in patients who have undergone radiation therapy or require PMRT have been raised as implant reconstruction gains popularity.11,12 Radiation therapy appears to increase breast complications and impair patient-reported satisfaction with breasts among patients receiving implant reconstruction but not those receiving autologous reconstruction.11 Immediate reconstruction with tissue expanders also poses new...
challenges for the radiation therapy (RT) team from a treatment planning perspective.

In this single institution report, we set out to describe the planning protocol developed for post mastectomy radiation therapy (PMRT) in patients with an in situ AeroForm™ expander, and to quantify the effect of this expander on target volumes and treatment factors. As a secondary aim, we included an assessment of acute side-effects with a focus on skin toxicity. The skin toxicity was graded according to Radiation Therapy Oncology Group Common Toxicity Criteria (RTOG CTC).

**Methods and materials**

Figure 1 shows an example of an air expander in situ on a CT transverse slice. Between March 2016 and August 2018, data was collected prospectively from eight female patients fitted with the AeroForm™ implant who were referred to our institution for post-operative radiotherapy. The objective was to develop a treatment planning protocol using the Pinnacle® Treatment Planning System (TPS) v9.10 (Philips, USA). All patients underwent a computed tomography (CT) scan as per our standard breast treatment simulation protocol on the Optima CT 580 RT (GE Healthcare) scanner. All treatment plans were generated for the Elekta Synergy linear accelerator with Agility™ head (Elekta, Sweden), and collapsed cone algorithm beam modelling was carried out in the TPS. The slice thickness of all CT scans was 2.5 millimetres (mm), and these were calculated on a 3mm dose grid size in the TPS. The radiation oncologist defined the treatment area by marking the superior level, inferior level, medial border and posterior border on the patient. These borders defined the beam shape of the multi-leaf collimator (MLC) using the beams eye view in the TPS.

All patients received a standard planning target volume (PTV) prescription of 5000 (cGy) in 25 treatments delivered five times per week. The PTV, as defined by the radiation oncologist on the planning CT scan, included the skin, subcutaneous tissue and pectoral fascia (Figure 2). The planning dosimetry goal was to achieve 95% of the prescribed dose to 95% of the PTV. The following parameters were evaluated on the dose volume histogram (DVH) for assessment of plan quality: PTV receiving 95% and 90% of the prescribed total dose (TD), along with volume (in cubic centimeters) receiving more than 10% of the prescribed TD. The organs at risk (OAR) assessed included: lungs (combined) volume receiving >2000cGy, >1000cGy, >500cGy and mean dose (cGy); left and right lung volumes individually receiving >3000cGy, >2000cGy, >1000cGy, >500cGy and mean dose (cGy); liver volume receiving >3500cGy, and heart volume receiving >2500cGy, >1000cGy and mean dose (cGy). Prescribed OAR for standard breast patients included the heart, liver, left and right lungs. The following additional contours were required due to artifact present on the CT scan: artefact_tissue, artefact_lung, artefact_heart, capsule of the implant and canister (Figure 2).

Density overrides were established by the planning team, to achieve the most accurate plan dosimetry as possible (Table 1). The values in Table 1 were dependent on CT density and the planning beam model. A three-dimensional conformal radiotherapy (3D CRT) wedged pair tangential planning technique was subsequently used to minimize the uncertainties (Figure 3).

**Table 1** Structure contours and density overrides

| Structure     | Structure definition                                                                 | Override density Values (g/cm^2) |
|---------------|--------------------------------------------------------------------------------------|----------------------------------|
| Artefact_tissue | Any artefact within patient normal tissue that has a density of >1.2g/cm^2             | 1.0                              |
| Artefact_heart | Any artefact within patient heart tissue that has a density of >0.95 g/cm^2           | 0.95                             |
| Artefact_lung  | Any artefact within patient lung that has a density of >0.1 g/cm^2                   | 0.15                             |
| Expander      | Ensure to exclude the canister volume once outlined.                                  | 0                                |
| Canister      | The metal CO₂ reservoir in the central area of the implant.                            | 5.8                              |

**Figure 1** AeroForm™ implant on a transverse slice CT scan. Main components visible are the centrally placed CO₂ canister encompassed by a permeable silicone casing.

**Figure 2** Additional contours required to be created and overridden as per Table 1 guidelines to account for artefact present on the CT scan.

**Figure 3** Transverse slice of 3D CRT wedged tangential beams applied for planning in Pinnacle3.
Daily imaging was performed throughout RT as per the department standard breast protocol. Two-dimensional electronic portal images (2D EPIs) were taken from the medial and lateral breast tangent beam angles to assess the lung depth treated at the central axis of the beam. The position of the metal canister and other expander changes were assessed daily. Kilovoltage (kV) planar images were taken on treatment days one through three to evaluate the set up position of the supraclavicular field and to check for any positional changes. These were repeated once a week to check positional accuracy. If positional or other changes were visible on the EPI, they were reported to the planning department. A repeat CT scan could then be performed on the patient and fused with the original CT to assess the overall dosimetric impact. Acute skin reactions were assessed at weeks two, four and six. A follow-up appointment was scheduled for six weeks after the last day of RT.

**Results**

Table 2 shows the prospectively collected data from eight female patients as they were planned and treated. The development of the treatment protocol was carried out in conjunction with the data collection. All patients, with the exception of Patient 4 (due to unrelated complications), received a standard PTV prescription of 5000cGy in 25 treatments delivered five times per week. The final dose distribution using the air tissue expander protocol was found to vary significantly from that of a standard breast plan due to the varying densities affecting the dose (Figure 4).

The potential structural, positional and volumetric changes throughout treatment, as assessed by daily EPIs, resulted in no significant visible changes in any patient, irrespective of the planning technique. Repeat CT scans were therefore not required to reassess the treated volume.

The dose to OAR according to the planning data for Patients 1-8 is outlined in Table 3. Different OAR were assessed depending on the laterality of the patient. The heart was of concern for patients being treated for left-sided breast cancer, while the liver was a priority for those receiving radiation to the right breast.

![Table 2 Air breast expander data collection](image)

| Pt number | Pt age | Volume treated | Laterality | Max point dose 0.1 cc (cGy) | Planning technique | Photon energy (MV) | Highest acute toxicity score | HI=(D2-98)/Dp |
|-----------|--------|----------------|------------|-----------------------------|--------------------|-------------------|----------------------------|----------------|
| 1         | 48     | CW+nodes       | Left       | 5895                        | Open tangential fields (no wedge) | 6                 | Erythema                  | 24.52%          |
| 2         | 77     | CW             | Right      | 5476                        | Wedged tangential pair        | 6                 | Erythema                  | 26.01%          |
| 3         | 49     | CW+nodes       | Left       | 5714                        | Wedged tangential pair        | 6                 | Moderate desquamation     | 27.62%          |
| 4         | 46     | CW+nodes       | Left       | 5714                        | Open tangential fields (no wedge) | 6                 | N/A                       | 26.28%          |
| 5         | 51     | CW+nodes       | Right      | 5786                        | Wedged tangential pair        | 6                 | Erythema                  | 27.64%          |
| 6         | 54     | CW+nodes       | Left       | 5814                        | Wedged tangential pair        | 6                 | Moderate desquamation     | 22.21%          |
| 7         | 68     | CW             | Right      | 5655                        | Wedged tangential pair        | 6                 | Moderate erythema         | 21.89%          |
| 8         | 54     | CW+nodes       | Left       | 5674                        | Wedged tangential pair        | 6                 | Moderate erythema         | 20.82%          |

![Table 3 Dose to organs at risk (%)](image)

| Patient   | 1 | 2 | 3 | 4 | 5 | 6 | 7 | 8 |
|-----------|---|---|---|---|---|---|---|---|
| Laterality| Left | Right | Left | Right | Left | Right | Left | Right |
| Left lung | V3000cGy | 28.1 | 0 | 11.5 | 23.1 | 0 | 16.5 | 0 | 27.2 |
|           | V2000cGy | 32.7 | 0 | 16.6 | 28.8 | 0 | 20.1 | 0 | 19.5 |
|           | V1000cGy | 39.6 | 0 | 24.5 | 38.0 | 0 | 25.5 | 0 | 14.7 |
|           | Mean dose (cGy) | 1530 | 30 | 850 | 1380 | 50 | 990 | 20 | 990 |
| Right lung| V3000cGy | 0 | 10.5 | 0 | 0 | 9.7 | 0 | 10.1 | 0 |
|           | V2000cGy | 0 | 12.6 | 0 | 0 | 13.8 | 0 | 12.7 | 0 |
|           | V1000cGy | 0 | 16.5 | 0 | 0 | 21.4 | 0 | 16.7 | 0 |
|           | V500cGy | 22.9 | 0 | 0 | 0 | 8 | 30 | 0 | 0 |
| Mean Dose (cGy) | 30 | 680 | 30 | 40 | 20 | 40 | 650 | 30 |
| Lungs combined | V2000 cGy | 13.6 | 6.5 | 8.0 | 13.9 | 7.4 | 9.0 | 7.2 | 9.4 |

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Table continued...

| Patient | 1  | 2  | 3  | 4  | 5  | 6  | 7  | 8  |
|---------|----|----|----|----|----|----|----|----|
| Mean dose (cGy) | 650 | 360 | 420 | 690 | 460 | 470 | 380 | 490 |
| Heart   |    |    |    |    |    |    |    |    |
| V2500cGy | 1.6 | 0  | 0.1 | 0.5 | 0  | 0  | 0.6 |    |
| V1000cGy | 2.5 | 0.1 | 0.8 | 1.4 | 0  | 0  | 0.2 |    |
| Mean Dose (cGy) | 20 | 110 | 140 | 160 | 40 | 130 |    |    |
| Liver   |    |    |    |    |    |    |    |    |
| V3500cGy | 0  | 2.9 | 0  | 0  | 0  | 3.8 | 0  |    |

Figure 4 Isodose distribution of an air tissue breast expander patient plan.

Discussion

Previous studies have shown that AeroForm™ air tissue breast expanders can function as intended with radiation doses of up to 70Gy and yet their presence in situ creates a challenge for the RT planning team when treating post-mastectomy patients. We identified planning challenges associated with the device, and then prospectively collected data on a series of patients using the implant expander with the aim of developing a specific treatment protocol for this patient subset.

- The four anticipated key planning challenges included:
  - The presence of a metallic cylinder in the center of the expander which leads to CT scan artefacts (with associated contouring difficulties), radiation scatter and dose inhomogeneities, and uncertainties with dose calculation (Figure 1).
  - How to achieve adequate dose distribution to the rind of chest wall stretched over the gas filled silicone shell of the expander. The thickness of this chest wall is often very thin-usually 5mm or less.
  - Appropriate beam arrangement when there is a large breast on the contralateral side, as seen in bilateral mastectomies and reconstructions.
  - Detection of volume changes during the course of RT and the need to compensate for small volume changes.

The thin thickness of the chest wall over the gas filled expander is a complex scenario for the TPS as it has to calculate dose to the buildup region while also missing backscatter. To raise the thickness of the layer, five to ten millimetres of bolus was planned conformally over the entire treatment field throughout the RT course, enabling a more accurate dose calculation. This likewise increased the dose to the subcutaneous tissue-a potential area of clinical failure in the chest wall after mastectomy. The density override of 5.8g/cm³ to the canister structure was established by transmission measurements in a solid water phantom at several depths with an air expander up-stream in a tangential-like setup.

No patients required additional CT simulation scans due to positional or volumetric changes of their implants throughout the treatment course. However, the 2D EPI images were not high quality and were not able to detect small changes in the position of the canister nor changes in volume. In addition to the poor image quality, research indicates that the EPI image registration can underestimate the actual bony anatomy setup error in breast cancer patients by 20% to 50%. A 3D cone beam CT (3D CBCT) imaging technique would be superior at detecting volumetric changes as well as setup uncertainties significantly. At the time of writing, we had not performed 3D CBCT daily imaging as all patients thus far had supraclavicular fields. The isocentre placement also created a significant collision risk between the gantry head of the linear accelerator and the patient and couch. It is our intention, as further data is collected, to investigate repositioning the isocentre to ensure there is no collision risk. This would enable 3D CBCT images to be performed daily to assess the volume more accurately.

The isodose distribution was found to vary significantly from a standard breast plan due to the composition of the AeroForm™ implant as shown in Figure 4. The 3D wedged tangential pair technique eliminated dose uncertainties that were of concern using our standard breast intensity modulated radiation therapy (IMRT) planning technique that additionally would have required a 3D imaging technique. The results in Table 2 from the analysis of the homogeneity index (HI) show a range from 20.8% to 27.6% due to the wedged pair technique not being able to compensate for the shape and density variations (the latter mainly the shielding effect of the canister).

As a secondary aim, we evaluated clinical outcomes specific to skin toxicity. Research indicates the use of chest wall bolus is associated with greater acute and late effects including moist desquamation and skin telangiectasia. With this knowledge, higher skin toxicity reactions than that of breast treatment without bolus were expected. Five out of seven patients experienced grade I or grade II erythema as an acute skin toxicity according to Radiation Therapy Oncology Group Common Toxicity Criteria (RTOG CTC), and two patients had moist desquamation (RTOG CTC grade III). However, in these patients, the area of concern was localized within the nodal region and was therefore unrelated to the implant. Patient 4 did not complete treatment due to unrelated complications and no clinical outcomes data is available for this patient.

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Conclusion

In patients with AeroForm™ air tissue expanders, it is feasible to plan with a 3D conventional planning technique. The addition of 5mm - 10mm of bolus placed over the entire chest wall increases the accuracy of dose distribution, as well as the superficial skin dose. Early clinical results show that the treatment method is reproducible and generates acceptable early toxicity. The clinical outcomes between breast patients undergoing PMRT with or without AeroForm™ expanders do not seem to differ with the use of the protocol, however we acknowledge that our findings are based on a small patient series. Data will continue to be collected prospectively to shape the protocol’s ongoing development. Further research into treatment imaging techniques would improve the accuracy of the canister position and implant volume evaluations.

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Conflict of interest

None.

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