



Inlay graft of acellular dermal matrix to prevent incisional dehiscence after radiotherapy in prosthetic breast reconstruction

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Background As the indications for postmastectomy radiotherapy expand, innovative solutions are required to reduce operative complications and reconstructive failure after prosthetic breast reconstruction. In this study, we investigated the effectiveness of acellular dermal matrix (ADM) inlay grafts in preventing postoperative wound dehiscence of irradiated breasts in the context of prosthetic breast reconstruction.

Methods A retrospective analysis was conducted of 45 patients who received two-stage prosthetic reconstruction and radiotherapy following mastectomy. An ADM graft was placed beneath the incisional site during the second-stage operation in 19 patients using marionette sutures, whereas the control group did not receive the ADM reinforcement. Patient demographics and complications such as wound dehiscence, capsular contracture, peri-prosthetic infection, cellulitis, and seroma were compared between the two groups.

Results During an average follow-up period of 37.1 months, wound dehiscence occurred significantly less often in the ADM-reinforced closure group (0%) than in the non-ADM group (23.1%) (P = 0.032). There was no significant difference between the two groups in relation to other complications, such as capsular contracture, postoperative infection, or seroma.

Conclusions The ADM inlay graft is a simple and easily reproducible technique for preventing incisional dehiscence in the setting of radiotherapy after prosthetic breast reconstruction. The ADM graft serves as a buttress to offload tension during healing and provides a mechanical barrier against pathogens. Application of this technique may serve to reduce complications in prosthetic breast reconstruction after radiotherapy.

Keywords Mammaplasty / Breast implantation / Acellular dermis / Radiotherapy / Complications

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INTRODUCTION

Postmastectomy radiation therapy after total mastectomy has been shown to reduce the loco-regional recurrence rate and to increase overall survival in both early and advanced breast cancer patients [1,2]. While immediate breast reconstruction may improve quality of life for mastectomy patients, increased complication rates are seen when the reconstructed breast is subject-

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ed to radiotherapy [3,4]. In a prospective cohort of irradiated breast cancer patients, the implant-based reconstruction group had double the odds of any complication compared to the autologous breast reconstruction group at 2 years after surgery [3]. The observed complications include capsular contracture, infection, pain, skin necrosis, fibrosis, and impaired wound healing [5,6]. In particular, earlier studies reported that wound dehiscence occurred in 10%-23.5% of irradiated breasts, compared to 1%–2% in non-irradiated patients [6-8]. A higher rate of infection is intimately linked to wound dehiscence, as implants are susceptible to infection because seeding can occur even through a small opening. Despite this, implant-based reconstruction appears to be on the rise globally, which is attributable to several factors, such as patients' and physicians' preferences, changes in oncologic practice, and the increased frequency of bilateral mastectomies [9,10]. Therefore, developing strategies to mitigate these complications is of paramount importance.

Acellular dermal matrix (ADM) is widely used in implantbased breast reconstruction surgery to cover the inferolateral pole of an implant for superior aesthetic outcomes, as well as in efforts to prevent capsular contracture [11,12]. A systematic review found that ADM usage does not predispose irradiated patients to higher rates of complications when used to aid prosthetic breast reconstruction [13]. However, scant research has explored whether an ADM inlay graft can specifically support the incision site and prevent complications in irradiated breasts. In this study, the primary aim was to determine whether the use of an ADM inlay graft could prevent incisional dehiscence in irradiated breasts. The secondary aim was to determine whether the use of an ADM inlay graft was related to other complications, such as infection or seroma.

METHODS

Patient selection

A retrospective study was conducted of patients who underwent mastectomy followed by two-stage prosthetic breast reconstruction and postmastectomy radiotherapy between July 2012 and August 2016 at our institute. Two cohorts were examined: patients with (n = 19) and without (n = 26) an ADM inlay graft at the incisional site during the second-stage operation. After April 2015, ADM inlay grafts were applied when the amount of closing tension was high after implants were placed, with the intention of reinforcing the wound. Specifically, the incision was deemed to have high tension when it was located directly on or near the point of maximal projection after the permanent implant was inserted. Two plastic surgeons (DWL and SYS) performed all reconstructions. Patient information was collected from hospital electronic medical records, including demographic variables such as age, cancer stage, and body mass index, as well as information regarding the expanded volume of the tissue expander, permanent implant volume, chemotherapy, anti-estrogen therapy, target therapy, and complications. The institutional review board approved this study design and the requirement for informed consent was waived (IRB No. 4-2018-1183).

Reconstruction technique

All patients underwent total mastectomy or skin-/nipple-sparing mastectomy performed by a breast oncologic surgeon, followed by immediate breast reconstruction by a plastic surgeon. After mastectomy, a tissue expander was inserted into the subpectoral plane and the lower lateral portion of the implant was covered with an ADM sling. The expander was prefilled with approximately one-third of the mastectomy volume. Expansion began 2 weeks after the operation and proceeded every 2 or 3 weeks. Prior to radiation therapy, the expander was deflated to include only 150–250 mL of saline to make it less prominent in the supine position, thereby minimizing interference with postmastectomy radiotherapy [14]. Immediately following completion of radiation therapy, re-expansion was performed every 2 or 3 weeks until the desired breast volume was reached.

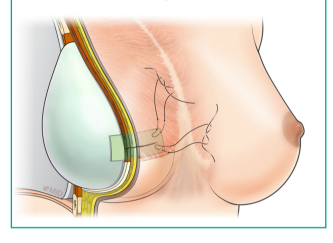
The second stage of reconstruction took place at least 3 months after the completion of radiation therapy and consisted of tissue expander removal with or without capsulectomy, followed by insertion of a permanent shaped cohesive-gel silicone implant. The incision was made as short as possible (6 cm on average), considering the impaired wound healing in irradiated tissues. ADM (MegaDerm; 4×7 cm and 0.8-1.4 mm in thickness; L&C Bio Corp., Seoul, Korea) was laid between the permanent implant and the expanded skin flap at the incisional site as an inlay graft to prevent incisional dehiscence in the ADM group. The ADM was fixed to the skin flap with Vicryl 3-0 pullout sutures. These marionette sutures were removed on postoperative day 7 (Fig. 1).

Radiation protocol

Radiation therapy commenced within 6 weeks after mastectomy or completion of the last cycle of chemotherapy. Prior to radiotherapy, all patients underwent simulation computed tomography scans for radiotherapy planning. The clinical target volume, including the ipsilateral chest wall, mastectomy scar, axillary nodes, internal mammary lymph nodes, and supraclavicular lymph nodes, was contoured. Depending on the protocol chosen by the radiation oncology team, patients received either conventional fractionation radiotherapy or hypofractionated radiotherapy. Patients with the former radiotherapy protocol were

Fig. 1. Acellular dermal matrix inlay graft

Acellular dermal matrix is laid between the permanent implant and the expanded skin flap on the incisional site in an inlay graft manner. The skin flap, pectoral muscle and previously grafted acellular dermal matrix are sectioned along the meridian line.



irradiated with 50.4 Gy in 28 fractions, while the prescription dose of the latter was 40.05 Gy in 15 daily fractions of 2.67 Gy.

Statistical analysis

When the data followed a normal distribution, continuous data were analyzed with the independent-samples t-test. Otherwise, the nonparametric Mann-Whitney test was used. The chi-square or Fisher exact test was performed to analyze categorical data. Significance was set at P < 0.05, using IBM SPSS Statistics for Windows, version 23.0 (IBM Corp., Armonk, NY, USA) for all computations.

RESULTS

Forty-five breast cancer patients who underwent two-stage prosthetic (expander and implant) reconstruction with postmastectomy radiotherapy were retrospectively examined. The demographic characteristics of patients in the non-ADM inlay graft and the ADM inlay graft group are illustrated in Table 1. The two groups were similar in age, body mass index, and preoperative breast volume (P > 0.05). There were also no significant differences between groups in the expanded expander volume, permanent implant volume, or the difference between those two values (P > 0.05). The two groups showed a similar distribution of additional therapy, such as neoadjuvant chemotherapy, adjuvant chemotherapy, anti-estrogen therapy, and target therapy (P > 0.05). However, there was a significant difference between the two groups in the radiation protocol (P = 0.005).

The rates of complications were compared between the two groups (Table 2). Wound dehiscence occurred in six cases in the

Table 1. Patients' demographic characteristics

Characteristics	Non-ADM group (n=26)	ADM group (n = 19)	P-value
Age (yr)	41.4 ± 10.6	41.1 ± 10.8	0.894
BMI (kg/m²)	21.6 ± 2.9	20.9 ± 2.5	0.179
Smoking	2 (7.7)	1 (5.3)	>0.999
Interval between second operation and radiation therapy (mon)	291.8 ± 312.2	216.8 ± 54.5	0.510 ^{a)}
Preoperative breast volume (mL)	365.0 ± 143.5	361.1 ± 105.7	0.921
Excised mastectomy skin (cm ²)	44.4 ± 33.2	52.9 ± 45.1	0.510
Expanded volume (mL)	443.2 ± 106.5	439.5 ± 101.7	0.910
Permanent implant volume (mL)	358.2 ± 89.0	313.2 ± 66.6	0.143 ^{a)}
Final stage			0.052
I	8 (30.8)	3 (15.8)	
II	11 (42.3)	9 (47.4)	
III	7 (26.9)	7 (36.8)	
IV	0	0	
Neoadjuvant chemotherapy	10 (38.5)	10 (52.6)	0.379
Adjuvant chemotherapy	14 (53.8)	10 (52.6)	>0.999
Anti-estrogen therapy	20 (76.9)	13 (68.4)	0.734
Target therapy	0	2 (10.5)	0.173
Radiation protocol			0.005
Conventional radiotherapy	11 (42.3)	16 (84.2)	
Hypofractionated radiotherapy	15 (57.7)	3 (15.8)	
Follow-up period (mon)	41.7 ± 16.7	30.7 ± 4.5	0.171

Values are presented as mean \pm SD or number (%).

ADM, acellular dermal matrix; BMI, body mass index.

^aThe statistical analysis was performed using the Mann-Whitney U test. The rest of the continuous variables were analyzed with the independent-samples t-test.

Table 2. Postoperative complications				
Complications	Non-ADM group (n=26)	ADM group (n = 19)	P-value	
Wound dehiscence	6 (23.1)	0	0.032	
Capsular contracture	9 (34.6)	4 (21.1)	0.507	
Cellulitis	1 (3.8)	3 (15.8)	0.295	
Peri-prosthetic infection	2 (7.7)	0	0.501	
Seroma/hematoma	0	0	-	
Total complications	18 (69.2)	7 (36.8)	0.031	
Needing major revisional surgery	5 (19.2)	0	0.063	
Values are presented as number (%).				

non-ADM inlay graft group, while no wound dehiscence was observed in the ADM inlay graft group (P = 0.032). Among the six operative revisions that were necessary in the non-ADM inlay graft group, two patients had implants downsized to reduce skin tension. Three other patients underwent reconstruction with a latissimus dorsi flap due to an unhealthy skin condition after repeated repairs. The last patient was treated with debridement, and an ADM inlay graft was used to repair the dehiscence site for therapeutic purposes. Nine cases of capsular contracture (34.6%) occurred in the non-ADM inlay graft group, and four

cases of capsular contracture (21.1%) occurred in the ADM inlay graft group [15]. In the non-ADM group, three patients (9.4%) suffered from postoperative infections, two of whom had a peri-prosthetic infection instigated by wound dehiscence, which required implant change and a latissimus dorsi flap. In the ADM graft group, three cases (15.8%) of cellulitis occurred, which were accompanied by redness and heating of the skin envelope; these resolved with intravenous antibiotic therapy within a week. No cases of seroma or hematoma were observed in either group. The total complication rate, excluding wound dehiscence, was 46.2% in the non-ADM graft group, while it was only 36.8% in the ADM graft group (P = 0.532). Furthermore, major revisional surgery to resolve the complications was needed in five patients (19.2%) in the non-ADM graft group and in none of the patients (0%) in the ADM graft group, but this difference did not achieve statistical significance (P = 0.063).

DISCUSSION

Previously, breast reconstruction with a prosthetic device combined with irradiation was a relative contraindication due to the high reconstruction failure rate of 37% and the corresponding negative impact on patient satisfaction [3,5,16]. Radiation not only increases complications, such as early wound healing problems and late capsular contracture, but also often results in a high-riding nipple and implant malposition [17]. Recently, complications and aesthetic results in implant-based reconstruction of irradiated breasts have improved to acceptable rates when compared to prior studies, with technical improvements including muscular coverage, use of ADM, newer prostheses, and modern radiation therapy protocols [18]. The use of ADM can provide aesthetic and psychological benefits, without affecting the complication rate in irradiated breasts [13]. Although Pestana et al. [19] suggested that ADM may increase complications requiring reoperation, they did not take into consideration whether radiation therapy took place before or after the reconstruction.

Although the volume of the permanent implants used in this study typically did not exceed that of the expander, tension upon wound closure can also be affected by the relative location of the incision in relation to the point of maximal projection. Wound dehiscence usually occurred at the maximal projection point of the implant, where the skin tension is the highest. Previously, Nahabedian [20] suggested making a new incision at the inframammary fold, instead of using the previous mastectomy incision to decrease the chance of wound dehiscence. Although the inferolateral inframammary fold incision is not remarkable, patients and reconstructive surgeons want to avoid additional scars if possible. In this study, application of ADM to the incisional site as an inlay graft decreased the rate of wound dehiscence. The grafted site of ADM was underneath the expanded skin flap, submuscular capsule, or previously taken ADM capsule, depending on the position of the incision. The ADM inlay graft is expected to take well regardless of the graft site, because ADM is grafted onto the capsule, which is a wellvascularized tissue.

Chronic injury of the soft tissue after radiation therapy is considered to occur due to the pathogenic fibroatrophic effect of radiation, which results in low tensile strength of the tissue [21]. Endothelial cell injury, vascular dysfunction, altered expression of bioactive cytokines including transforming growth factor-beta 1, and oxidative stress lead to excessive extracellular matrix deposition of collagen I and III, a persistent excess of myofibroblasts, and gradual fibroblast rarefaction combined with incomplete cell replacement [22]. As described by Cordeiro et al. [23], two-stage postmastectomy reconstruction with expanders and implants has a higher rate of failure at the second stage, but yields improved aesthetic outcome in the setting of postmastectomy radiation therapy compared with direct-to-implant reconstruction. The ADM inlay graft may be considered a strategy to mitigate these complications and reconstructive failure, as demonstrated by our results. We propose several speculative hypotheses regarding how an ADM inlay graft may prevent incisional dehiscence. First, the ADM inlay graft may act as a mechanical buttress to offload skin tension in a pants-over-vest fashion. The marionette stitches and ADM graft may relieve tension on the incision, improving blood supply to the wound. In a similar manner, ADM is often used for repair of ventral hernias to reinforce and offload tension on fascial closures [24]. Secondly, the ADM inlay graft may serve as a scaffold for wound regeneration, as ADM is already known to promote chronic wound healing [25]. Third, the ADM inlay graft can act as an additional barrier between the prosthetic and external environment. In non-grafted patients, wound complications may lead to severe peri-prosthetic infections that require operative interventions. The additional barrier provided by the ADM inlay graft reduced operative complications to 0%.

This is the first study, as far as the authors are aware, in which ADM was placed as an inlay graft manner to prevent wound dehiscence in irradiated breasts. The limitations of this study include its retrospective design, in which the patients were not randomly assigned to groups. Furthermore, as we analyzed a series of consecutive patients, the distribution of the radiation protocol was different between the ADM and non-ADM groups. Our radiation oncologists began using a hypofractionated radiation technique in the middle of the study period, resulting in an imbalance between the two groups in the radiation protocol. However, the relationship between hypofractionated radiotherapy and breast reconstruction has not been delineated [25]. Even if hypofractionated radiotherapy affected the complication rate of reconstructions, we believe that the ADM inlay graft is an ideal and easy to replicate technique for preventing incisional dehiscence in prosthetic reconstruction in the setting of postmastectomy radiation. Additional prospective trials with larger groups are required in the future for further verification of our findings.

NOTES

Conflict of interest

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

Ethical approval

The study was approved by the Institutional Review Board of Severance Hospital (IRB No. 4-2018-1183) and performed in accordance with the principles of the Declaration of Helsinki.

Author contribution

Conceptualization: Lee DW. Data curation: Ahn SJ. Formal analysis: Song SY. Funding acquisition: Lee DW. Methodology: Fan KL. Project administration: Lew DH. Visualization: Kim MJ. Writing - original draft: Ahn SJ. Writing - review & editing: Kim MJ. Approval of final manuscript: all authors.

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