# Tissue Reinforcement in Implant-based Breast Reconstruction

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<td>doi:10.1097/GOX.0000000000000140</td>
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Tissue Reinforcement in Implant-based Breast Reconstruction

Michael Scheflan, MD*  Amy S. Colwell, MD, FACS†

Background: Tissue reinforcement with allogeneic or xenogeneic acellular dermal matrices (ADMs) is increasingly used in single-stage (direct-to-implant) and 2-stage implant-based breast reconstruction following mastectomy. ADMs allow surgeons to control implant position and obviate the need for submuscular implant placement. Here, we review the benefits and risks of using ADMs in implant-based breast reconstruction based on available data.

Methods: A comprehensive analysis of the literature with focus on recent publications was performed. Additional information regarding the proper use of ADMs was based on our institutional experience.

Results: ADM use may improve definition of the lateral confines of the breast and lower pole projection. It may facilitate direct-to-implant procedures and improve aesthetic outcomes. The effect of ADMs on complication rates remains controversial. Known patient risk factors such as obesity, smoking, and radiotherapy should be considered during patient selection. For patients with healthy, well-vascularized skin envelopes, ADM-assisted direct-to-implant reconstruction is a safe and cost-effective alternative to 2-stage implant reconstruction, with low complication rates. ADMs may be used to treat capsular contracture, and limited available data further suggest the possibility that ADMs may reduce the risk of capsular contracture. Novel synthetic or bio-synthetic tissue reinforcement devices with different physical and ease-of-use properties than ADMs are emerging options for reconstructive surgeons and patients who seek to avoid tissue products from human or mammalian cadavers.

Conclusions: ADM-assisted implant-based breast reconstruction may improve aesthetic outcomes. However, appropriate patient selection, surgical technique, and postoperative management are critical for its success, including minimizing the risk of complications. (Plast Reconstr Surg Glob Open 2014;2:e192; doi: 10.1097/GOX.0000000000000140; Published online 4 August 2014.)

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reconstructions, implant-based reconstructions are simpler, take less time to perform, are less invasive, support faster patient recovery, and avoid the need for donor site surgery, which may cause significant postoperative deterioration of physical well-being. In addition, they are believed to have more favorable third-party reimbursement limits.

The increasing availability of acellular dermal matrices (ADMs) prepared from human or animal cadavers for tissue reinforcement provides plastic surgeons with a unique tool to improve aesthetic outcomes of implant-based reconstruction by expanding and reshaping the implant pocket, while further reducing the invasiveness of surgical intervention. The substantial pain caused by the serratus anterior elevation necessary for total or partial muscle coverage can be avoided, as lateral coverage is provided by the ADM. However, results of a prospective randomized study in 70 patients who underwent immediate ADM-assisted or conventional submuscular tissue expander/implant reconstruction after mastectomy revealed no significant differences between these patient groups in immediate postoperative pain or pain during the expansion phase. In addition to providing mechanical stability, ADMs facilitate cellular and vascular infiltration during wound healing and tissue regeneration through incorporation of the matrix. Since the first publication of ADM-assisted primary breast reconstruction in 2005, an increasing number of products have become available in the United States (Table 1).

In Israel, AlloDerm (LifeCell Corp., Branchburg, N.J.), the prototypical ADM of human dermal origin, has been available since 2005 and SurgiMend (TEI Biosciences Inc., Boston, Mass.), a fenestrated ADM derived from fetal bovine dermis, has been available since 2008. Allogeneic ADMs are not extensively used in many European countries because of regulatory restrictions on human tissue products and cost considerations favoring alternative products.

In this review, we describe the benefits of tissue reinforcement in implant-based breast reconstruction, such as improved aesthetic outcomes, provide cost considerations, and discuss the risk of postoperative complications. Moreover, we provide recommendations based on our own experience for how the risk of complications can be minimized by appropriate patient selection, surgical technique, and postoperative management.

**IMMEDIATE IMPLANT-BASED RECONSTRUCTION WITH AND WITHOUT TISSUE REINFORCEMENT**

**Conventional Implant-based Reconstruction**

Conventional implant-based reconstruction after mastectomy requires the creation of an implant pocket beneath the pectoralis major muscle for total or partial muscle implant coverage (Table 2). For total muscle coverage, elevation of the serratus anterior muscle is required for lateral coverage and support. Total muscle coverage limits the possible anterior and inferior projection and thus may lead to less than optimal aesthetic outcomes. Furthermore, because the size of the implant pocket that can initially be created with this approach is constrained by available skin and muscle tissue after the mastectomy, immediate reconstruction generally requires a 2-stage procedure, in which a tissue expander is implanted first, then gradually expanded over several months, and eventually replaced by a permanent implant.

**Implant-based Reconstruction with ADMs**

The use of ADMs enables the formation of larger implant pockets and optimal implant positioning without the need for serratus anterior muscle elevation. Provided the skin is sufficiently healthy, ADM use in 2-stage reconstruction allows for more predictable tissue expander position, larger intraoperative expander fill volumes, and fewer expansions compared with submuscular placement.

An important aspect of ADM use is that it facilitates immediate direct-to-implant reconstruction.

### Table 1. Allogeneic and Xenogeneic Soft Tissue Reinforcement Devices Available in the United States

<table>
<thead>
<tr>
<th>Product</th>
<th>Year Introduced</th>
<th>Manufacturer</th>
<th>Material Origin</th>
<th>Sterile</th>
<th>Hydration Time</th>
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<tbody>
<tr>
<td>AlloDerm</td>
<td>1994</td>
<td>LifeCell Corp.</td>
<td>Human dermis</td>
<td>No (aseptically processed)</td>
<td>10–40 min</td>
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<tr>
<td>AlloDerm Ready To Use</td>
<td>2012</td>
<td>LifeCell Corp.</td>
<td>Human dermis</td>
<td>Yes</td>
<td>≥2 min</td>
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<tr>
<td>DermaMatrix</td>
<td>2005</td>
<td>Synthes, Inc. (Westchester, Pa.)</td>
<td>Human dermis</td>
<td>No (aseptically processed)</td>
<td>3 min</td>
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<tr>
<td>FlexHD</td>
<td>2007</td>
<td>Ethicon, Inc.</td>
<td>Human dermis</td>
<td>No (aseptically processed)</td>
<td>None</td>
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<tr>
<td>ALOMAX</td>
<td>≥2009</td>
<td>Davol Inc. (Warwick, R.I.)</td>
<td>Human dermis</td>
<td>Yes</td>
<td>“Rapidly”</td>
</tr>
<tr>
<td>Repriza</td>
<td>2010</td>
<td>Specialty Surgical Products, Inc. (Victor, Mont.)</td>
<td>Human dermis</td>
<td>Yes</td>
<td>None</td>
</tr>
<tr>
<td>Strattice</td>
<td>2008</td>
<td>LifeCell Corp.</td>
<td>Porcine dermis</td>
<td>Yes</td>
<td>≥2 min</td>
</tr>
<tr>
<td>Veritas Collagen Matrix</td>
<td>2001</td>
<td>Synovis (St Paul, Minn.)</td>
<td>Bovine pericardium</td>
<td>Yes</td>
<td>None</td>
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<tr>
<td>SurgiMend</td>
<td>2006</td>
<td>TEI Biosciences Inc.</td>
<td>Fetal bovine dermis</td>
<td>Yes</td>
<td>60 s</td>
</tr>
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</table>
Table 2. Surgical Techniques of Total and Partial Muscle Coverage and of ADM-assisted Direct-to-implant Reconstruction

| Total muscle coverage | The pectoralis major muscle is elevated lateral to medial by electrocautery after the lateral edge has been identified. The inferior and medial origins are maintained. The serratus anterior muscle is elevated over the fourth, fifth, and sixth ribs by electrocautery. The superior portion of the rectus abdominis fascia or muscle may be included in the dissection of the muscular pocket to facilitate positioning of the tissue expander and help prevent superior malposition. One or 2 closed-suction drains are placed to facilitate drainage and help prevent seroma. |
| Partial muscle coverage | The pectoralis muscle is elevated and partially released inferiorly to allow greater lower pole expansion. The medial most sternal origin should be released inferiorly to allow the implant to fit medially and obtain cleavage. This typically corresponds to the 4 and 8 o’clock positions on the chest wall but may be advanced to the 3 or 9 o’clock positions if necessary. Sutures are typically used to prevent retraction of the pectoralis muscle, and the serratus muscle is often elevated to help cover the expander laterally and prevent lateral malposition. |
| ADM-assisted direct-to-implant reconstruction | In one technique, the pectoralis major muscle is elevated from the chest wall inferiorly up to the 3 or 9 o’clock position medially, and a tailored ADM is placed in the pocket and sewn to the pectoralis muscle. In another technique, the pectoralis major muscle is elevated up to the 4 or 8 o’clock position medially, and a rectangular ADM is sewn to the inframammary fold, if present and well preserved, or to the thoracic fascia to create a new fold. After placement of the implant into the subpectoralis/sub-ADM pocket, a drain each is placed into the pocket along the inframammary fold and into the lateral pocket margin. |

(Table 2, Fig. 1) in appropriately selected patients by allowing precise positioning of a full-sized permanent implant, with favorable aesthetic outcomes and minimal risk of implant displacement, visibility, rippling, or extrusion. Moreover, ADM materials that are not completely resorbed help prevent pectoralis muscle retraction and offer additional soft tissue coverage in the lower pole of the breast. Experience from 331 consecutive immediate direct-to-implant reconstructions performed with AlloDerm at the Massachusetts General Hospital suggests that this approach is associated with favorable aesthetic outcomes and low complication rates in patients with thick, well-vascularized skin flaps after skin-sparing or nipple-sparing mastectomy. Similarly, the use of a SurgiMend in 341 consecutive...
PATIENT RISK FACTORS FOR COMPLICATIONS AND RECONSTRUCTIVE FAILURE

A number of database analyses consistently identified high body mass index and smoking as independent risk factors for complications and/or implant loss. For 1170 two-stage breast reconstructions performed over a 2-year period, smoking, obesity, and hypertension each increased the odds of reconstructive failure by factors of 5, 7, and 4, respectively. For more than 14,000 reconstructions with or without ADM captured in the Tracking Outcomes and Operations in Plastic Surgery (TOPS) database between 2008 and 2011, high body mass index, smoking, and diabetes were independent risk factors for expander/implant loss. ADM use seems to have no substantial influence on patient-related risk factors. A recent analysis of data from the American College of Surgeons National Surgical Quality Improvement Program, which identified smoking and body mass index as independent risk factors for short-term complications, found no statistically significant risk differences between immediate ADM-assisted (n = 1717) and submuscular (n = 7442) tissue expander reconstruction.

ADM USE AND POSTOPERATIVE COMPLICATIONS

A persistent concern among plastic surgeons is whether ADMs increase the risk of short-term complications, given the conflicting findings from retrospective studies, systematic reviews, and meta-analyses. However, although meta-analyses (level III evidence) found increased risks of infection, seroma, and/or implant loss associated with ADM use, some recent large studies (level II or III evidence) not included in these analyses found similar or lower complication rates for ADM-assisted versus traditional 2-stage reconstruction. For example, a comparative study of 479 implant-based reconstructions found no difference in total complication rates or rates of infection and seroma between ADM-assisted direct-to-implant reconstruction compared with 2-stage tissue expander/implant reconstruction without ADM (Table 3). Similarly, in a recent prospective cohort study, ADM use was associated with significant reductions in expander/implant loss (Table 2) and unexpected returns to the operation room. These findings suggest that ADM use itself is not an independent risk factor for complications and that the large discrepancies in findings among different institutions may be attributable to other factors. This view is supported by the results of recent, large-scale National Surgical Quality Improvement Program and TOPS analyses. Although the TOPS analysis showed that ADM use (versus no ADM use) was associated with a statistically significant increase in the risk of expander/implant loss (odds ratio, 1.42; 95% confidence interval, 1.04–1.94; P = 0.026), the absolute risk increase was only 0.7%.

In our experience, the vast majority of complications attributed to ADM-assisted reconstruction are avoidable by appropriate patient selection and surgical technique. An emerging consensus on the importance of these factors is reflected in the recently published joint guidelines from the Association of Breast Surgery and the British Association of Plastic, Reconstructive and Aesthetic Surgeons and recommendations by other experts in the field. Effective coordination of mastectomy and reconstructive surgery to ensure optimal viability of the skin envelope may further improve outcomes. Familiarity of the reconstructive surgeon with optimal device-specific techniques also is crucial for avoiding complications. In 331 consecutive ADM-assisted 1-stage implantations conducted between 2006 and 2009 at the Massachusetts General Hospital, increasing experience of the surgeons and better communication with the breast surgeon substantially reduced the incidence of skin necrosis, resulting in a significant 2-fold reduction in total complication rates from the surgeons’ first to subsequent years of performing the procedure (21.4%–10.9%; P < 0.02). However, if skin viability is questionable at the time of the mastectomy, total muscle coverage (rather than
Table 3. Complication Rates with ADM-assisted and Conventional Implant-based Breast Reconstruction in Controlled Studies

<table>
<thead>
<tr>
<th>Source</th>
<th>No. Reconstructions (Patients)*</th>
<th>ADM</th>
<th>Non-ADM</th>
<th>ADM</th>
<th>Non-ADM</th>
<th>P</th>
<th>ADM</th>
<th>Non-ADM</th>
<th>P</th>
<th>ADM</th>
<th>Non-ADM</th>
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<td>ADM</td>
<td>Non-ADM</td>
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<tr>
<td></td>
<td></td>
<td>Total Complications, %</td>
<td>Infections, %</td>
<td>Seroma, %</td>
<td>Skin Necrosis, %</td>
<td>Reconstructive Failure or Implant Loss, %</td>
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<td>ADM-assisted direct-to-implant versus 2-stage tissue expander/implant with partial or total submuscular placement</td>
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<td>148</td>
<td>14.8</td>
<td>19.6</td>
<td>0.180</td>
<td>3.0</td>
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<td>0.152</td>
<td>1.5</td>
<td>1.9</td>
<td>0.810</td>
<td>9.1</td>
<td>10.1</td>
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<td>0.19</td>
<td>7.0</td>
<td>4.3</td>
<td>0.17</td>
<td>4.0</td>
<td>2.0</td>
<td>0.18</td>
<td>8.5†</td>
<td>6.6†</td>
<td>0.41</td>
</tr>
</tbody>
</table>

*Complication rates were based on the number of reconstructions, or on the number of patients where patient numbers are indicated.
†Major flap or skin necrosis.
‡Patient selected for ADM due to thin skin envelope.
§Major complications only.
¶Requiring intravenous antibiotics.
‖Seroma or hematoma.
**Reconstructions may have included direct-to-implant procedures.
‖‖Explantation due to infection, seroma, or extrusion.
ND indicates not determined.
partial muscle release with or without ADM) should be considered even if this may negatively affect the final aesthetic outcome (Fig. 2).

**THE RISK OF CAPSULAR CONTRACTURE**

Capsular contracture is a common risk of implant-based reconstruction, particularly in the setting of radiotherapy. A variety of factors may contribute to development of capsular contracture, including but not limited to insufficient sterility during surgery, hematoma, mechanical strain on the inferior skin envelope, type and surface properties of the implant, and radiation therapy. It has been suggested that the use of ADMs may minimize capsular contracture by reducing pressure on the inferior breast skin envelope, which in turn may reduce fibroblast stimulation and inflammation. In a retrospective comparison of 2-stage reconstruction with and without ADM use during expander implantation in 203 patients, the capsular contracture rate was significantly lower for ADM-assisted reconstructions (3.8% versus 19.4%; P < 0.001) at a mean follow-up after implant exchange of 29 months. Moreover, a remarkably low capsular contracture rate of only 0.4% was observed in an 8-year study of 466 ADM-assisted direct-to-implant breast reconstructions with a mean follow-up of 29 months, and ADM use, including complete implant coverage, has been used successfully in the treatment of capsular contracture. However, given that the risk of implant-related capsular contracture increases over time, it remains to be demonstrated whether ADMs reduce the long-term risk of capsular contracture. Furthermore, it is presently unclear whether non-ADM-based tissue reinforcements can influence the incidence of capsular contracture.

**EFFECT OF RADIOTHERAPY**

A major concern with implant-based breast reconstruction is the effect of radiation therapy on complication rates. Radiation therapy given before or after mastectomy has been associated with significantly increased rates of major complications, including implant removal or replacement and capsular contracture. However, a recent systematic review found that ADM did not increase the complication rate in the setting of radiotherapy. Similarly, another recent literature review concluded that ADM use essentially had a neutral effect on postoperative complications among patients who received adjuvant radiation therapy after implant-based reconstruction, and results of a recent retrospective study even suggested that ADM use may significantly reduce the odds of complications (including explantation) in the setting of postmastectomy radiation therapy.

Among patients who received a total of 479 ADM-assisted direct-to-implant or conventional 2-stage reconstructions at the Massachusetts General Hospital, radiotherapy was associated with an increased rate of early complications. Among patients who received radiation, the highest complication rate was seen in the setting of preoperative irradiation and conventional 2-stage reconstruction (41.1%), whereas the lowest rate was seen in direct-to-implant reconstructions with postoperative radiation (16.7%). The importance of timing of postmastectomy radiotherapy was also demonstrated in a prospective, controlled study of 257 patients undergoing subpectoral 2-stage breast reconstruction. Patients who received radiotherapy on the tissue expander had a significantly higher failure rate (40%) than those who received radiotherapy on the permanent implant (6.4%) or received no radiotherapy (2.3%; P < 0.0001).

Radiation therapy also seems to affect capsular contracture rates after ADM-assisted implant-based

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breast reconstruction. The overall rate of clinically significant capsular contracture (grade III/IV) among 341 reconstructions at the Assuta Medical Center was only 2.0%. Remarkably, capsular contracture occurred exclusively in patients who previously received preoperative or postoperative radiation therapy at a rate of 12.3%. In a study of ADM-assisted 2-stage reconstruction in 289 women, radiation therapy before mastectomy and at expander stage resulted in dramatically increased rates of infection (53% and 73%, respectively, versus 1.4% without radiation) and grade III/IV capsular contractures (41% and 61%, respectively, versus 1.4% without radiation), although eventual explantation was avoided in most cases. Thus, although the use of ADM in implant-based reconstruction may reduce the risk of capsular contracture, it remains to be demonstrated whether this benefit extends to patients who receive pre- or postoperative radiation therapy.

**COST CONSIDERATIONS**

Recent cost analyses including the costs of probable complications in addition to physician and hospital fees estimate that ADM-assisted direct-to-implant reconstruction may result in moderate to substantial cost savings compared with traditional 2-stage implant reconstruction. At the Massachusetts General Hospital, overall hospital charges of the 2 procedures did not differ significantly (P = 0.8) because the substantially lower professional fees charged by anesthesiologists and surgeons for ADM-assisted direct-to-implant reconstruction were largely offset by higher hospital charges. An important cost factor when using tissue reinforcement can be the costs of the tissue support itself, particularly if AlloDerm is used. However, these costs may decrease with the increasing availability of lower cost xenogenic ADMs and alternative tissue reinforcement devices. Furthermore, overall cost savings may potentially be larger than currently estimated if future research demonstrated that the use of ADM-assisted procedures substantially reduces the incidence of implant loss, capsular contracture, time off from work, and corrective surgery. The usage of ADM in 2-stage reconstruction increases the material costs for the procedure. For cost savings to be realized in this setting, a decreased need for revisions, shorter operative time, and/or lower complication rates would need to be demonstrated in a cost–benefit analysis.

**NON-ADM OPTIONS FOR SOFT TISSUE REINFORCEMENT**

ADMs may vary in their chemical composition and physical properties, with the potential to affect the quality and timing of tissue regeneration, and the risk of complications. Differences in material-associated risk of inflammation, thickness, requirements for hydration, and sterility may affect the handling facility of ADMs and their ability to simultaneously provide adequate structural support and sufficient pliability. Patients who do not accept cadaver material being part of their reconstructed breasts would benefit from alternative products with appropriate physical and ease-of-use properties that are either synthetic or made from biomaterials other than ADMs.

TiLOOP Bra (pfm medical titanium, Nuremberg, Germany), a nonabsorbable titanium-coated polypropylene mesh approved for breast reconstruction in Europe, was retrospectively evaluated in 231 breast reconstructions. Explantation (7.8%) was the most common major complication, with skin necrosis and capsule fibrosis identified as significant risk factors in multivariate analysis.

**CONCLUSIONS**

Tissue reinforcement in implant-based breast reconstruction may reduce the invasiveness of implant-based procedures and improve aesthetic outcomes. Appropriate patient selection, proper surgical technique, and adjusted postoperative management are critical for the success of ADM-assisted reconstruction, including minimizing the risk of complications. For qualifying patients with healthy, well-vascularized skin envelopes, ADM-assisted direct-to-implant
reconstruction is a safe and cost-effective alternative to 2-stage implant reconstruction that can provide excellent aesthetic results. Known patient risk factors for implant-based reconstruction, such as obesity and smoking, also apply to ADM-assisted reconstruction and should be considered during patient selection.

The availability of novel synthetic or biosynthetic tissue reinforcement devices that have different physical and ease-of-use properties than ADMs may enhance the ability to refine surgical techniques to further optimize aesthetic outcomes and minimize complications.

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