CHAPTER

26

Fat Injections

Kotaro Yoshimura and Yuko Asano

Key Points

1. Moderate augmentation (100–200 ml) of the breast is successfully achieved by autologous lipojection without major complications if appropriately performed in selected patients.

2. Breasts augmented with fat injection are soft and show natural texture and appearance, and patients are free from daily stress and future concerns derived from foreign materials (such as complications and potential implant replacement or removal).

3. Surviving fat volume varies substantially among patients, and multiple factors are likely to affect the clinical results; patient factors include skin redundancy of the breast, quality (such as viability and progenitor richness) of fat grafts, and infiltration techniques are key determinants to clinical results of lipojection.

4. Aspirated fat tissue should be appropriately harvested and stored, and quickly processed and infiltrated with proper devices to avoid degradation of the graft tissues and to place aliquots of fat grafts as diffusely as possible.

5. Relative deficiency of adipose progenitor cells in aspirated fat tissue may lead to long-term atrophy of the grafts, and supplementation of vascular stromal fraction containing adipose progenitor cells may boost the efficacy and safety of lipojection to the breasts.

Introduction

Autologous fat transplantation is one of the promising cosmetic treatments for facial rejuvenation and soft-tissue augmentation due to the lack of an incisional scar and complications associated with foreign materials. However, certain problems remain, such as unpredictability and a low rate of graft survival due to partial necrosis. It has also been used in breast augmentation by a limited number of plastic surgeons, although the use of autologous fat for breast augmentation has been controversial due to the lack of consensus on whether it is safe and appropriate because of microcalcifications that may cause confusion in the evaluation of mammograms.

Implantation of prostheses has been a gold standard for breast augmentation, but complications with artificial materials such as capsular contracture remain to be resolved. The presence of the implant and capsules induced by implants could also affect breast tissue visualization in the mammogram. Furthermore, there is potential for rupture when pressure is exerted on the implant during mammography, and for this reason, hospitals in Japan reject women with breast implants to undergo mammography as a part of the annual social health examinations. Recently, autologous fat injection has been re-evaluated as a potential alternative to artificial implants for breast augmentation. This re-evaluation may reflect recent advances in autologous fat transfer and the radiological detection of breast cancer.

In this chapter, potentials of fat injection for breast augmentation or reconstruction are discussed as well as our novel approach of autologous fat grafting called cell-assisted lipotransfer (CAL); this is the concurrent transplantation of aspirated fat tissue and adipose progenitor cells.
Patient Selection

There are several patient factors that may affect the clinical result of conventional lipoinjection or CAL: skin redundancy of the breasts, age, body mass index (BMI), individual quality or character of the fat tissue, adhesive scars, breast implant and its capsule, systemic disease such as autoimmune disease, oral corticosteroids, etc.3,5 Good candidates are those who have sufficient fat at donor sites and redundant breast skin with healthy vascularity and without any scars.

Lipoinjection can be performed in any patient from their teens to 70s, but patients with low body mass index (BMI < 17) or athletes with little body fat are not good candidates due to the difficulty in harvesting a large volume of fat tissue. Patients who want a large-volume (250–400 ml) augmentation are not good candidates because augmentation volume achieved by a single session of lipoinjection is limited (100–200 ml).

Some patients are concerned about complications derived from foreign bodies and about possible surgical removal or replacement of implants in the future. Others do not want their history of breast surgery to come out. These patients want to avoid breast implants and do not want a great deal of breast augmentation; they are good candidates for this procedure.

Indications

Operative indications are described below according to three kinds of graft tissue preparations or operative purposes.

Graft tissue preparations

As for lipoinjection, we use conventional lipoinjection (micro-fat grafting) and a new technique; grafting of progenitor-enriched adipose tissue. We call the latter CAL; the concept and details of CAL are described later. There are two types of CAL: mini-CAL and full-CAL; only the fluid portion of liposuction aspirates are used for harvesting adipose progenitor cells in mini-CAL, while another similar volume of liposuction aspirates to graft tissues are used for the cell isolation in full-CAL (Fig. 26.1). Thus, full-CAL requires twice the volume of adipose tissue as the conventional lipoinjection or mini-CAL; approximately 700–800 ml of lipoaspirate are needed for conventional lipoinjection or mini-CAL for both breasts, while 1200–1500 ml lipoaspirate are required for the full-CAL procedure.

If BMI is less than 18 or body weight is less than 45 kg, conventional lipoinjection or mini-CAL is recommended. On the other hand, for patients with BMI > 25, full-CAL is easily performed without concerns for the donor site.

Operative purposes

Breast augmentation

Lipoinjection can be performed without combining any other procedures. We, however, also propose a secondary lipoinjection after 1 year implantation of breast prostheses if patients are very thin, have flat breasts and tight breast skin, because only a small volume lipoinjection can be performed in those cases due to high internal pressure and skin tension after injection.

Breast implant replacement

Lipoinjection can be performed at the same time as implant removal. The implants have to be removed through a periareolar incision, though they can be removed through an axillary incision if lipoinjection is performed separately from implant removal.

Breast reconstruction

Lipoinjection can be performed in any cases, but detailed assessment of tissue and skin conditions of the breasts is necessary before application. In most reconstruction cases with skin shortage, lipoinjection after tissue expansion is recommended; lipoinjection can be performed immediately after removal of the tissue expander. For patients who have sufficiently redundant skin and do not have severe scar tissue or adhesion of the skin to the underlying fascia or any other deep tissues, lipoinjection alone may work successfully. Lipoinjection can also be applied to patients with irradiated skin, though the injection volume is usually limited. Several sessions of lipoinjection can be performed to irradiated breasts with each interval being more than 12 months. Lipoinjection after tissue expansion is also applied to these irradiated patients, but expander implantation under the pectoralis major muscle and a careful expansion are recommended.
Inborn deformity

Simple hypoplastic breasts can be successfully treated with lipoinjection in most cases, but tubular deformity is hard to improve by lipoinjection alone. Funnel chest deformity can be corrected with lipoinjection, though the filling volume by a single surgery is limited and repetitive treatments are usually recommended.

Injection of progenitor-enriched fat tissue: principles and therapeutic concepts of CAL

Cell components of adipose tissue

Adipose tissue consists predominantly of adipocytes, adipose stromal cells (ASC), vascular endothelial cells, pericytes, fibroblasts and extracellular matrix. Adipocytes constitute more than 90% of tissue volume, but they are much larger in size than the other cells and the number of adipocytes is estimated to be less than 50% (Fig. 26.2). ASC are considered to be adipose tissue-specific progenitor cells (adipogenic and angiogenic progenitors), some of which have been shown to differentiate into multiple lineages and are now called adipose-derived stem cells. ASC contribute to adipose tissue turnover (adipose tissue is thought to turn over every 2–10 years) and provide cells for the next generation. ASC are currently being used in various clinical trials, including treatments for rectovaginal fistula (autologous cultured ASC) and graft-versus-host disease (non-autologous ASC). If ASC are harvested from a large volume (e.g., 500 ml) of liposuction aspirates, ASC can be used clinically without cell expansion because a sufficient number of cells can be obtained. The use of minimally manipulated fresh cells may lead to higher safety and efficacy in actual treatments.
Aspirated fat tissue versus intact fat tissue

We can use aspirated fat tissue as lipoinjection material but not excised fat tissue. Aspirated fat is fragile parts of the adipose tissue taken with negative pressure. Our research revealed that aspirated fat tissue contains only half the number of ASC compared to intact fat tissue (Fig. 26.3). The two main reasons for this relative deficiency of ASC contained in aspirated fat tissue are: (1) a major portion of ASC are located around large vessels (within the tunica adventitia) and left in the donor tissue, and (2) some ASC are released into the fluid portion of liposuction aspirates. Our histological study indicated that ASC are located not only between adipocytes but also around vessels. Large-sized vessels are located in the fibrous part of the tissue, which contains intact fat tissue but not aspirated fat tissue. Thus, aspirated fat tissue is regarded as relatively progenitor-poor fat tissue compared to intact fat tissue.

Stromal vascular fraction

Through collagenase digestion a heterogeneous cell mixture, which contains cell types other than adipocytes, can be extracted from adipose tissue as a cell pellet. This cell fraction is called the stromal vascular fraction (SVF) (Fig. 26.4), because they are basically stromal cells and contain vascular endothelial and mural cells. In the clinical setting SVF contains a substantial amount of blood-derived cells, such as leukocytes and erythrocytes, as well as adipose-derived cells such as ASC and vascular endothelial cells. Our study revealed that nucleate cells contained in the SVF are composed of 37% leukocytes, 35% ASC, 15% endothelial cells and other cells, though the percentage of blood-derived cells strongly depends on individual hemorrhage volume. In CAL, the freshly isolated autologous SVF is used as a supplementation for fat graft tissue without any manipulations such as cell sorting or cell culture.

Concept of CAL

Aspirated fat tissue has a significantly lower progenitor/mature-cell ratio as mentioned above, and this low ASC/adipocyte ratio may be the main reason for long-term atrophy of transplanted adipose tissue. There are at least three experimental studies, including ours, demonstrating that supplementation of adipose progenitor cells enhances the volume or weight of surviving adipose tissue. Enrichment of adipose progenitor cells by supplementation of SVF improves progenitor/adipocyte ratio; progenitor-poor aspirated fat tissue will be converted to progenitor-rich fat tissue. In CAL, freshly isolated SVF, which contains ASC, is supplemented to progenitor-poor aspirated fat tissue; the cells are attached to the aspirated fat with the fat acting as a living bioscaffold before transplantation (Fig. 26.5).

Transplanted adipose tissue undergoes ischemia and subsequent reperfusion as well as high internal pressure by edema and inflammatory changes in the host tissue. The microenvironments, injury-associated growth factors, and inflammation-associated cytokines and chemokines would influence ASC behaviors during the acute phase of tissue repair. Adipose grafts undergo adipocyte and capillary remodeling, and ASC are a main cell population functioning in the repairing process of the adipose tissue. The relative deficiency of ASC in aspirated fat tissue may affect the replacement process and lead to
postoperative atrophy of grafted fat, which is known to commonly occur during the first 6 months after lipoinjection.

**Operative Technique**

**Surgical procedures**

**Basic breast augmentation**

Donor sites are usually the thighs alone or the thighs and the abdomen or flanks, decided according to patient's preference and BMI. After the liposuction site is infiltrated with saline solution with epinephrine (0.0001%) under general anesthesia, adipose tissue is suctioned using a cannula with 2.5-mm inner diameter and a conventional liposuction machine. The lipoaspirates are centrifuged at 700 g for 3 min, and put into a metal jar (1000 ml) which is placed in water with crushed ice.

For the injection syringe, a 10 ml LeVeen inflator (Boston Scientific Corp., MA) or our original syringe (20 ml) is used because they are screw-type syringes (with a threaded plunger) and threaded connections that fit both the connecting tube and the needle, to allow for

---

**Fig. 26.3** Comparison of human intact fat tissue and aspirated fat tissue obtained from a single site of a single patient. Schematic views (top), electron microscopic and whole mount staining images (middle), and isolated progenitor cells (bottom). The basic structure of adipose tissue was preserved in the aspirated fat, while vascular vessels, especially those of large size, were significantly less detected in aspirated fat compared to the excised fat. It is well known that the honeycomb structures of vascular and neural perforator networks are left intact in aspirated sites after liposuction operation. ASC yield from aspirated fat tissue was significantly less (56 ± 12%) than that from excised fat tissue.
The stromal vascular fraction (SVF) can be obtained from adipose and fluid portions of liposuction aspirates through collagenase digestion. SVF contains 10–40% of adipose-derived stromal cells (ASC) (CD34+ CD31– CD45–), part of which have multipotency and can differentiate experimentally into several lineages in vitro. SVF contains also blood-derived cells (CD45+ cells) such as leukocytes. ASC are considered to physiologically differentiate into adipocyte and vessels.

Fig. 26.5 Scheme of cell-assisted lipotransfer (CAL). Relatively progenitor-poor aspirated fat tissue is converted to progenitor-rich fat tissue by supplementation with the stromal vascular fraction (SVF) isolated from one-half of the aspirated fat sample. (Strictly speaking, the source of SVF differs between mini-CAL and full-CAL; see also Figure 26.1 for the difference.) SVF cells are attached to the aspirated fat tissue, which acts as a scaffold in this strategy.
precise control during injection (Fig. 26.6). To reduce the
time of the procedure, two syringes are used; while one
syringe is being used for an injection, the other is filled
with the graft material in preparation for the next injec-
tion. A 16- or 18-gauge needle (150 mm long) is used
for lipoinjection and inserted subcutaneously from the
inframammary fold or areolar margin (Fig. 26.7). The
operator takes care to insert and place the needle hori-
zontally (parallel to the body), in order to avoid damag-
ing the pleura and causing a pneumothorax. The needle
is inserted in several layers and directions, and is con-
tinuously and gradually retracted as the plunger is
advanced (Fig. 26.7). This technique is used to obtain a
diffuse distribution of the graft material. The grafts are
placed into the fatty layers on, around, and under the
mammary glands (but not intentionally into the
mammary glands), and also into the pectoralis muscles.

Breast implant replacement

For patients with implants, lipoinjection can be per-
formed simultaneously with implant removal. Breast
implants are removed through a periareolar incision,
which is placed at the caudal third of the areola margin.
The lipoinjection is begun at the deepest layer under the
implant capsule and completed with the injection into
the most superficial subcutaneous layer. In the deepest
layer, the operator takes care to insert and place the
needle horizontally (parallel to the body), in order to
avoid damaging the pleura and causing a pneumothorax,
by inserting the operator’s finger into the implant capsule,
placing it on the bottom of the capsule, and recognizing

Fig. 26.6 Injection devices. A high-pressure injection can be
performed with a disposable syringe with a threaded plunger.
A 150 mm-long 16- or 18-gauge needle is connected to the
syringe with a connecting tube threaded at both ends. The
injection needle is rigidly manipulated by an operator, while an
assistant rotates the plunger according to the operator’s
instruction.

Fig. 26.7 Schematic diagram of the injection method. (Left) The
needle is inserted from either the areola margin or the
inframammary fold in variable directions and planes to achieve a
diffuse distribution. A small amount of fat tissue is injected as
small aliquots or a thin string with a long needle on a syringe with
a threaded plunger while the needle is continuously withdrawn
(right). Approximately 200–300 ml of fat tissue is usually injected
for cosmetic breast augmentation on each side. Fat is not injected
into the mammary glands, but into any other layers including the
pectoralis muscles.
Breast reconstruction

For breast reconstruction, lipoinjection is performed basically similar to basic breast augmentation. Centrifuged lipoaspirates are injected from the inframammary fold or scars. In patients who have substantial scar tissue or adhesion between skin and deep tissues, a tissue expander is inserted first and breast reconstruction with lipoinjection is performed as a secondary surgery immediately after removal of the tissue expander. The volume of injection is usually determined by skin tension of the reconstructed breast.

Preparation procedures of graft materials

Conventional lipoinjection

A volume of lipoaspirate is harvested by liposuction and centrifuged at 700 g for 3 min, and put into a metal jar (1000 ml) which is placed in water with crushed ice. As the centrifugation reduces the adipose volume by 25–30%, the volume reduction should be taken into account in tissue harvesting.

Full-CAL

In full-CAL, about twice the volume of lipoaspirate is harvested and half of the adipose portion and all of the fluid portion of the liposuction aspirate are used for isolation of SVF (Fig. 26.1). If a patient has BMI < 25, 1500 ml of aspirated fat tissue can be easily harvested from the abdomen and flanks or thighs. If BMI < 20, fat should be usually taken from both the abdomen and thighs.

About half of the collected liposuction aspirate (500–700 ml of aspirated fat tissue) is used for harvest of SVF. The SVF is isolated as described below and the cell processing procedure takes about 80 min. During the processing period, the other half of the lipoaspirate is harvested and prepared as a graft material with centrifugation at 700 g for 3 min. The freshly isolated SVF is added to the centrifuged fat tissue, followed by gentle mixing and a 10–15 min incubation to achieve appropriate cell adhesion to the centrifuged fat tissue.

Mini-CAL

In mini-CAL, the same volume of lipoaspirate is harvested as in the conventional lipoinjection; the adipose portion is centrifuged as the graft material, while the fluid portion of the liposuction aspirate is used for isolation of SVF. The cell processing process takes about 30 min. The freshly isolated SVF is added to the centrifuged fat tissue, followed by gentle mixing and a 10–15 min incubation to achieve appropriate cell adhesion to the centrifuged fat tissue.

Cell isolation procedure (cell processing for SVF isolation)

Processed lipoaspirate cells (PLA) cells and liposuction aspirate fluid (LAF) cells, both are so-called SVF, are sepa-

Fig. 26.8 Schematic illustration of the lipoinjection procedure for breast implant replacement. While injecting, operator’s fingers are inserted through a periareolar skin incision into the cavity of implant capsule to determine the location of the needle tip.
rated from the fatty and fluid portions of liposuction aspirates, respectively. Both cells are used for full-CAL, while only LAF cells are used for mini-CAL (Fig. 26.1).

For PLA cells, the suctioned fat is digested with 0.075% collagenase in phosphate buffered saline for 30 min on a shaker at 37°C after centrifugation. Mature adipocytes and connective tissues are separated from cell pellets by centrifugation (800 g, 10 min). Pellets were resuspended in erythrocyte lysis buffer (155 mM NH₄Cl, 10 mM, KHCO₃, 0.1 mM EDTA) and incubated for 5 min at room temperature. The pellets are resuspended and passed through a 100-mm mesh filter. To eliminate any remaining collagenase, the cell pellets are repeatedly washed by resuspending in Hanks buffer following centrifugation at least three times. For LAF cells, the suctioned fluid is centrifuged (400 g, 10 min), and the pellets are resuspended in erythrocyte lysis buffer. After 5 min incubation at room temperature, lysates were passed through a 100-mm mesh filter. The cell pellets are repeatedly washed by resuspending in Hanks buffer following centrifugation at least three times, and passed through a 100-mm mesh filter.

The whole procedure should be performed by well-trained physicians or technicians in an aseptic room (preferably at a level of ‘good manufacturing practice’) according to a designated standard operating procedure. Isolated cells should be strictly evaluated in quantity and quality. Cell counts for erythrocytes and nucleated cells are performed with a cell counter used for blood tests. The whole process of cell isolation takes about 80 min. It is also recommended that a fraction of the isolated SVF is seeded and cultured to make sure of cell viability and another fraction is frozen and stored in a deep freezer or liquid nitrogen for future cell tracing.

**Pitfalls and How to Correct**

**Pre- and postoperative evaluations**

For evaluation of clinical results, physical measurements (maximum and bottom breast circumferences, etc.), mammogram, magnetic resonance imaging (MRI) scan, echogram, photograph, and videograph are performed. We have also adopted a three-dimensional measurement system which enables a volumetric evaluation of the breast mound in a standing position (Fig. 26.9). An echogram is easy to perform at each visit and is sensitive enough to detect small cyst formation. Long-term follow up with an annual mammogram is recommended to detect abnormal signs such as calcification.

**Clinical results**

The total operation period is approximately 2–2.5 hours for conventional lipoinjection, 2.5–3 hours for mini-CAL, and 3.5–4 hours for full-CAL. The time of the injection process ranges from 35 to 60 min for both breasts. Subcutaneous bleeding and edema is usually seen on some parts of the breasts, and resolves in 1–2 weeks. Transplanted adipose tissue is gradually absorbed during the first 2 postoperative months (especially during the first month), and the breast volume shows a minimal change thereafter, although skin tension sometimes becomes looser between 2 and 6 months. The three-dimensional measurements showed that the surviving fat volume in full-CAL ranged from 100 to 250 ml at 12 months, meaning that the graft take ranged from approximately 40 to 80% (Fig. 26.10). Compared to breast augmentation with implants of the same size, augmentation with lipoinjection showed a lower height but more natural contour of breasts. Cyst formation depends on the volume and distribution of fat grafts. No cysts are palpable so long as the injection was correctly performed (see below), though cysts with a size of less than 5 mm may be detected by echogram. Patients are generally satisfied with the resulting texture, softness, contour and absence of foreign materials despite the limited size increase possible with autologous tissue. Computerized tomography scans and MRI show that transplanted fat tissue survives well and forms a significantly thick fatty layer, not only subcutaneously on and around the mammary glands but also between the mammary glands and the pectoralis muscles.

**Refinement of autologous fat graft techniques**

Surviving fat volume varies substantially among patients, and multiple factors are likely to affect the clinical results; patient factors include skin redundancy of the breast and technical factors include devices, graft fat preparation, and injection techniques. It is well accepted that adipose tissue should be placed as small aliquots, preferably within an area 3 mm in diameter. Since it takes a long time to perform the ideally diffuse placement of suc-
Fig. 26.9 Three-dimensional measurement system for breast volume. Breast volume can be measured by this system with the patient in a sitting position. A Perpendicular striped lights are shown on the breasts and photographed with a stereo-type digital camera. B The digital images are then analyzed with customized software; C, D the volume and projection of each breast above a standard plane designated by three fixed points (the shoulder, suprasternal notch, and xiphoid process) which do not usually shift after breast augmentation, are calculated, E, F Pre- and postoperative photos.
tioned fat in breasts,1 we use a disposable syringe with a threaded plunger and connections and a very long needle (150 mm); these devices are critical to performing large-volume lipoinjection safely and precisely in a short time3 (Fig. 26.6). We use a relatively large-sized suction cannula (2.5–3.5 mm inner diameter), centrifuge the aspirated fat, and keep it cooled until transplantation. It should be also noted that aspirated fat tissue should be injected as soon as possible, such as within 60 min after harvest. In our experience, clinical results (increase in breast size) appeared to be superior when centrifuged fat was used compared to non-centrifuged fat. This may be due to the improved adipocyte density and ASC/adipocyte ratio after centrifugation.16 ASC supplementation dramatically improves ASC/adipocyte ratio and is suggested to minimize adipose atrophy after transplantation, though further studies are needed to elucidate the effects of ASC.

Complications

If injection is performed incorrectly, problems deriving from fat necrosis will be seen: such as no augmentation effects, cyst formation, fibrogenesis, and calcification. Small cysts (<8 mm) detected by echogram usually disappear between 9 and 18 months, so no treatment is needed. Tiny calcifications may occur 1–2 years after surgery but they are very rare and easy to distinguish from malignant signs. To identify them on the mammogram is important and is useful to distinguish them from abnormal changes in the future. A larger volume of liposuction could induce postoperative donor site problems such as irregularity or seroma; lean patients are more susceptible to this, so preoperative selection of patients and careful procedures in liposuction are important.

Representative cases

Representative cases are demonstrated in Figs 26.11–26.15; one case from each category (breast augmentation...
Fig. 26.12 Case 2 (breast augmentation with full-CAL). A 30-year-old woman underwent breast augmentation with CAL (310 ml in each breast). A–C Preoperative views and D–F postoperative views at 24 months. Her breasts were dramatically augmented with an increase in breast circumference difference by 8.0 cm at 24 months. The breast mounds were soft with no subcutaneous indurations. An original inframammary fold on the left breast is slightly visible, but injection scars are not visible. G, H Mammograms 24 months after surgery show no abnormal signs.
Fig. 26.12, cont'd
Fig. 26.13 Case 3 (breast implant replacement with full-CAL). A 33-year-old woman who had 210 ml saline implants underwent implant removal and simultaneous CAL (260 ml in each breast). Preoperative (left panels) and postoperative (right panels) views at 12 months. Clinical views showed capsular contracture and upward displacement of the left implant A before surgery, B while the breasts had a natural and symmetrical appearance at 12 months. C, D T1-weighted MRI revealed that the transplanted adipose tissue survived and formed thick layers around and under the mammary gland at 12 months. E, F Mammograms showed that neither implant was ruptured before surgery and G, H that no calcification or other abnormal signs were visible in either breast at 12 months. Augmented breast mounds maintained sufficient breast volume even after implant removal, and were naturally soft without any subcutaneous indurations.
Fig. 26.14 Case 4 (inborn deformity treated with full-CAL). A 26-year-old woman, who had a hypoplastic breast along with thoracic deformity on the right side, underwent CAL augmentation for both sides (325 ml and 105 ml on the right and left, respectively) Preoperative (left panels) and postoperative (right panels) views at 12 months. Both breasts were significantly improved from A without any indurations detected. B A few small injection scars on the inframammary folds remain but are nearly invisible. C, D Pre- and postoperative CT scans show that adipose tissues were augmented not only subcutaneously but also under the mammary glands with no abnormal nodules or calcifications detected.

Fig. 26.15 Case 5 (breast reconstruction after mastectomy with full-CAL). A 43-year-old woman, who had undergone partial mastectomy followed by irradiation therapy, underwent breast reconstruction with full-CAL (240 ml on the left side). The right healthy side was also augmented by full-CAL of 160 ml. A Preoperative and B postoperative views at 12 months. Both breasts were augmented very well, and the reconstructed breast mound was soft and showed natural skin texture.
with conventional lipoinjection, breast augmentation with full-CAL, breast implant replacement with full-CAL, inborn breast deformity treated with full-CAL and breast reconstruction with full-CAL).

Postoperative Care

After surgery, the breasts should be kept in a good position with a well-fitting brassiere. Patients can take a shower on the next day. Massage of the breasts is prohibited during the first 3 months.

Conclusions

Increase in size obtained by lipoinjection is moderate, but patients can achieve soft and natural-looking breasts without any future concerns associated with foreign bodies. Major complications are not seen as long as the infiltration technique is correct. Our preliminary experiences of the CAL technique suggest the efficacy and safety of the ASC supplementation. Through further improvements of the technique and longer follow-up studies, autologous tissue transfer may become widely used for augmentation and reconstruction of the breasts in the future.

References