Reduction Mammaplasty

Effective: January 1, 2021

Next Review: July 2021
Last Review: December 2020

IMPORTANT REMINDER

Medical Policies are developed to provide guidance for members and providers regarding coverage in accordance with contract terms. Benefit determinations are based in all cases on the applicable contract language. To the extent there may be any conflict between the Medical Policy and contract language, the contract language takes precedence.

PLEASE NOTE: Contracts exclude from coverage, among other things, services or procedures that are considered investigational or cosmetic. Providers may bill members for services or procedures that are considered investigational or cosmetic. Providers are encouraged to inform members before rendering such services that the members are likely to be financially responsible for the cost of these services.

DESCRIPTION

Reduction mammaplasty is the surgical excision of a substantial portion of the breast, including the skin and underlying glandular tissue, until a clinically normal size is obtained.

MEDICAL POLICY CRITERIA

Notes:

- This policy is not applicable when there has been a prior mastectomy for which the Women's Health & Cancer Rights Act applies. The Reconstructive Breast Surgery/Mastopexy, and Management of Breast Implants policy (Surgery, Policy No. 40 – see Cross References) may be applicable. Please refer to the Surgery, Policy No. 40 for reconstruction after partial or complete mastectomy.
- This policy is not intended to address treatment of gender dysphoria which is addressed in the Transgender Services medical policy (Medicine, Policy No. 153 – see cross references), which may be applicable.

I. Reduction mammaplasty may be considered medically necessary when one or more of the following are met:
A. As a preparatory first stage procedure preceding a nipple-sparing mastectomy, when the amount of breast tissue removed from each breast is at least the minimum in grams per breast for the patient’s body surface area (in meters squared using the Mosteller formula) according to the Schnur Sliding Scale (see Policy Guidelines for body surface area/breast weight table); or

B. When all of the following criteria (1. - 3.) are met:

1. The patient is aged 18 years or older; and

2. The amount of breast tissue removed from each breast, not including fat removed by liposuction, must be at least the minimum in grams per breast for the patient’s body surface area* according to the Schnur Sliding Scale (see Policy Guidelines), or, in cases of asymmetry where one breast meets criterion but the other breast does not, the combined weight of the tissue removed from both breasts must total at least twice the Schnur Sliding Scale minimum for the patient’s body surface area (the health plan may review medical records to confirm the amount of breast tissue removed during the procedure); and

3. Two or more of the following clinical indications have been present for at least 12 months and have failed to respond to appropriate conservative therapy:

   a. Pain in the upper back, neck, shoulders, and/or arms, with all of the following documented in the medical records by the referring physician or provider:

      i. The pain is of long-standing duration and increasing intensity; and

      ii. The pain has been evaluated to determine that it is not associated with another diagnosis such as arthritis, if applicable; and

      iii. The pain is not relieved by at least three months of conservative therapy such as an appropriate support bra with wide straps, exercises, heat/cold treatments and appropriate non-steroidal anti-inflammatory agents/muscle relaxants.

   b. Dermatitis of the shoulder or shoulder grooving not responding to at least three months of conservative treatment including a support bra or appropriate dermatologic treatments, (e.g. taking steps to eliminate friction, heat, and maceration by keeping skin cool and dry and where appropriate, topical agents).

   c. Intertrigo between the pendulous breasts and the chest wall persisting despite at least three months of conservative dermatologic treatments (e.g. taking steps to eliminate friction, heat, and maceration by keeping skin cool and dry and where appropriate, antimycotic agents).

   d. Kyphosis documented by x-ray.

   e. Ulnar paresthesia not relieved by at least three months of conservative therapy such as an appropriate support bra with wide straps, range of motion exercises, physical therapy, and appropriate non-steroidal anti-inflammatory agents/muscle relaxants.
II. Reduction mammaplasty is considered not medically necessary when Criteria I. is not met.

III. Reduction mammaplasty for gynecomastia is considered not medically necessary.

IV. The use of liposuction as an additional procedure with breast reduction surgery is considered not medically necessary.

V. The use of liposuction as the sole procedure for breast reduction is considered investigational.

NOTE: A summary of the supporting rationale for the policy criteria is at the end of the policy.

POLICY GUIDELINES

Mosteller formula: body surface area (m²) = ( [height (cm) x weight (kg)] / 3600 )½ [1]

Click here for link to Body Surface Area Calculator

Schnur Sliding Scale

<table>
<thead>
<tr>
<th>Body Surface Area (m²)</th>
<th>Grams per Breast of Minimum Breast Tissue to be Removed</th>
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<tbody>
<tr>
<td>1.350-1.374</td>
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<td>1.625-1.649</td>
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NOTE: When BSA is < 1.350 minimum is 199 grams
LIST OF INFORMATION NEEDED FOR REVIEW

It is critical that the list of information below is submitted for review to determine if the policy criteria are met. If any of these items are not submitted, it could impact our review and decision outcome:

1. Total amount of breast tissue to be removed, include if L/R or bilateral
2. Height and weight
3. Any two of the following detailed in chart notes, history and physical, physical therapy notes, radiologic exams, dermatology treatments notes, and/or any other clinical notes:
   A. Medical records by the referring physician, which include pain in the upper back, neck, shoulders and/or arms with documentation of long standing pain, and detailed notes regarding treatment with at least three months of conservative therapy, and that the pain is not associated with another diagnosis such as arthritis;
   B. Documentation of shoulder grooving or dermatitis of the shoulder with description of at least three months of conservative treatment with dermatology notes and outcome;
   C. Intertrigo despite three months detailed documentation of conservative therapy;
   D. X-ray showing kyphosis;
   E. Ulnar paresthesia despite three months documentation of conservative therapy and outcome with chart notes detailing specific treatment.

CROSS REFERENCES

1. Gender Affirming Interventions for Gender Dysphoria, Medicine, Policy No. 153
Female breast hypertrophy, or macromastia, is the development of abnormally large breasts in the female. This condition can cause significant clinical manifestations when the excessive breast weight adversely affects the supporting structures of the shoulders, neck and trunk. Macromastia is distinguished from large, normal breasts by the presence of persistent symptoms such as shoulder, neck, or back pain, shoulder grooving, or intertrigo. This condition can be improved and the associated signs and symptoms can be alleviated by reduction mammoplasty surgery.

The following literature appraisal is focused on the investigational technique of reduction mammoplasty by liposuction alone. In order to understand the impact on health outcomes of reduction mammoplasty by liposuction alone, prospective clinical trials are needed, comparing liposuction with standard reduction mammoplasty. These comparisons are necessary in order to understand the safety and efficacy of liposuction and to determine whether liposuction offers advantages over conventional surgical procedures with respect to patient satisfaction, complications, durability, and cosmesis.

While there are some published articles concerning the use of liposuction as the sole procedure for breast reduction, none compare the outcomes of liposuction alone to standard excisional reduction mammoplasty.[2-9] Examples of these articles are detailed below:

Moskovitz (2007) conducted a study of liposuction alone for treatment of macromastia in twenty-four African-American women due to their high risk for complex scar formation following standard excision mammoplasty.[8] The mean aspirate was 1075 cc of fat per breast; however, the before and after liposuction pictures indicate that the participants continued to support large breasts. Outcome measures included the SF-36, EuroQol, Multidimensional Body-Self Relations Questionnaire, McGill Pain Questionnaire and Breast-Related Symptoms Questionnaire. Statistical analysis demonstrated a significant improvement in breast-related symptoms and pain. This was a relatively small, non-randomized trial and patients were not blinded to the intervention. Conclusions concerning the effect of liposuction alone on breast-related symptoms in patients with macromastia cannot be made.

Jakubietz (2011) reported the indications and limitations of this procedure compared to conventional surgical excision.[9] Advantages included selective removal of fat, ease of procedure, and the advantages of less invasive procedures such as faster recovery time and reduced scarring. One disadvantage of liposuction alone included the inability to correct shape and ptosis, making aesthetic results optimal only for young patients. In addition, there are concerns about the extent to which subsequent breast imaging may be impaired, and the possible spread of cancer cells. The authors recommended caution when considering use of this technique.

In summary, high quality evidence on the use of liposuction for reduction mammoplasty has not been identified; comparative trials of sufficient size and duration are needed before any conclusions can be made about the use of this technique for breast reduction.
In 2011, the American Society of Plastic Surgeons (ASPS) released an evidence-based clinical practice guideline on the use of reduction mammaplasty.[10] Several clinical questions were addressed, including whether women who did not meet standard health insurance criteria for volume of breast resection experience postoperative relief. On the basis of a single study which compared satisfaction outcomes of women who met standard insurance criteria with women who did not meet such criteria, the society concluded that, “resection volume is not correlated to the degree of postoperative symptom relief.” The society recommended extending the option of reduction mammaplasty to this category of patient. However, among women not meeting standard criteria for resection volume, no comparisons were made between surgical and standard conservative treatment, limiting interpretation of the above findings. Additionally, these recommendations did not specifically address the safety and effectiveness of reduction mammaplasty by liposuction.

Female breast hypertrophy, or macromastia, is the development of abnormally large breasts in the female, which can cause medical problems. There is enough research to show that reduction mammaplasty can improve health outcomes for certain patients with this condition. Therefore, reduction mammaplasty may be considered medically necessary when policy criteria are met. Reduction mammaplasty as treatment for macromastia is considered not medically necessary when policy criteria are not met.

There is not enough research to show that liposuction mammaplasty can improve health outcomes more than traditional mammaplasty techniques. Therefore, reduction mammaplasty by liposuction alone is considered investigational.

Gynecomastia refers to the benign enlargement of the male breast, mainly due to excessive growth of glandular tissue. Reduction mammaplasty (partial removal) for the treatment of gynecomastia is considered not medically necessary as the current standard of care is for the removal of most or all glandular tissue.

REFERENCES


**CODES**

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*Date of Origin: January 1996*