Comparison between different methods of breast implant volume choice and degree of postoperative satisfaction

Introdução

Aumentação mammaplastia é atualmente a mais frequentemente realizada cirurgia cosmética mundialmente¹. No Brasil, é a segunda cirurgia mais frequentemente realizada de plástica estética, correspondendo a 13,64% de procedimentos¹. Assim, as taxas de complicações, que são relativamente baixas, tornam-se altas em números absolutos. Haveria muitos artigos na literatura sobre o tema, no entanto, a maioria deles são estudos retrospectivos ou multicentro e não se concentram no tamanho do implante. Adicionalmente, quando elas descrevem a escolha do implante, eles o fazem por meio de uma metodologia única, sem nenhuma comparação prospectiva entre os outros métodos existentes²-⁷. A população estudada em estes estudos geralmente consiste em mulheres do hemisfério Norte, onde a cultura e as costumes estão fortemente correlacionadas com as aspirações e perspectivas sobre a cirurgia²,³,⁷,¹⁰. Embora o Brasil tenha sido o segundo país do mundo em cirurgia plástica estética em 2016, poucas estudos realizados que avaliaram este aspecto cultural no país¹.

O estudo de aumento mammaplastia com implante de mama foi altamente satisfeito entre os pacientes. No entanto, não houve diferença no grau de satisfação no período pós-operatório entre os três métodos de medicação do volume mamário.
Comparison between different methods of breast implant volume choice and degree of postoperative satisfaction

METHODS

This was a prospective study, carried out at the University Hospital Pedro Ernesto of the State University of Rio de Janeiro (UERJ), with 94 women from Rio de Janeiro, aged between 18 and 49 years, submitted to augmentation mammoplasty due to hypomastia. The non-inclusion factors were: patients under 18 years of age, those with indication for mastopexy, smokers, patients with psychiatric disorders, those with a prior history of breast surgery and those with systemic diseases. The exclusion factors were: breastfeeding or pregnancy during the study, failure to perform the preoperative study and loss of postoperative follow-up.

The implants were introduced in the retroglandular space through an inframammary access. Patients were systematically divided into three groups, for convenience: Control Group with 44 patients, MamaSize® Experimental Group with 25 patients and Silicone Experimental Group with 25 patients. All implants were textured, with a round base and high projection.

In the control group, breast implants were chosen through anthropometric measurements. By measuring the basis and thickness of the patient’s breast parenchyma, the silicone implant basis is calculated. Subsequently, the product that corresponds to that base is chosen.

In the MamaSize® Experiment group, the implants were chosen according to the MamaSize® meter, where the mold is placed behind a bra without a cup, in front of a full-length mirror (Figure 1). The intersection between the mold of the patient’s breast size (vertical axis) and that chosen by the patient (horizontal axis) shows the volume to be placed (Figure 1). Aiming to be similar to silicone molds, the following correlation was made between the volume chosen through MamaSize® and the available volume to be placed: 170=175mL, 220=215mL, 240=235mL, 260=255mL, 290=285mL, 300=305mL, 330=325 mL, 360=355 mL.

In the Experimental Silicone Group, the following volume molds were used: 175mL, 195mL, 215mL, 235mL, 255mL, 285mL, 305 mL, 325mL and 355mL. The patient chose the volume using the breast implant measurer that reproduced them in their shapes and dimensions, using a bra without a cup, in front of a full-length mirror. After the choice, new tests with volumes were performed, one above and one below the chosen one, for the ratification of the decision.

Satisfaction questionnaires were applied in the pre- and postoperative periods by the same evaluator, using a visual analogue scale (Figure 2), where 0 meant
Comparison between different methods of breast implant volume choice and degree of postoperative satisfaction

very unsatisfied and 100 meant very satisfied for the four variables: shape, size, symmetry, and consistency. For the scar, only the postoperative period was evaluated\textsuperscript{15,16}.

![Visual analog scale](image)

\textit{In the preoperative period}

\textit{According to the scale above, grade questions 1 through 4 according to your degree of satisfaction:}

1. How do you feel about your breast shape today?
2. How do you feel about your breast size today?
3. How do you feel about your breast symmetry today?
4. How do you feel about your breast consistency today?

\textit{In the postoperative period of the 1\textsuperscript{st} month / 6\textsuperscript{th} month / 12\textsuperscript{th} month}

\textit{According to the scale above, grade questions 1 through 4 according to your degree of satisfaction:}

1. How do you feel about your new breast shape today?
2. How do you feel about your new breast size today?
3. How do you feel about your new breast symmetry today?
4. How do you feel about your new breast consistency today?
5. How do you feel about your breast scar today?

\textbf{Figure 2. Visual Analog Scale and Questionnaire}

The study, classified as a longitudinal and analytical case-control one, has a convenience sample, in which the definition of cases and controls is systematized. We analyzed 25 cases as Silicone, 25 cases as MamaSize\textsuperscript{®} and 44 cases as Control. The first analysis used descriptive statistics. Frequency, relative frequency and 95\% confidence intervals were used to depict the variables in descriptive tables, aiming to understand the groups' profile in relation to the performed research. All variables were tested in relation to their normality, i.e., to verify whether they come from a population with normal distribution, using the Shapiro-Wilks test. To verify whether the anthropometric variables and the research variables were from the same population, regardless of the group (Silicone, MamaSize\textsuperscript{®} and Control), the ANOVA statistical test was used, or Kruskal-Wallis test, when the data were not from the Normal population. To verify the existence of significant changes during the follow-up period, the \textit{t} test was used, or the Wilcoxon test, when the data are not from the Normal population. For the assessment of the statistical tests, a significance level of 0.05 (5\%) was used, in which \textit{p}-value <0.05 was considered significant for the analyses.

The computer programs Microsoft Excel 2010 and Software R, version 3.3.1 (R Core Team 2015, Vienna, Austria) were used for data organization, creation of tables/charts and statistical analysis.

The study was submitted to \textit{Plataforma Brasil}, under CAAE number 13986513.2.0000.5259 version 1, and was approved on 05/21/2013, under Opinion Number 285716. All patients signed the free and informed consent form for the surgical procedure and for the study participation.

\textbf{RESULTS}

The mean values of the patients' age, BMI and mammary basis did not show statistical difference between the three groups. The mean age of the groups was 28 years, the mean BMI was 21.91 (kg/m\textsuperscript{2}) and the mean value of the mammary basis was 11.62cm. The results of the mean implant volume, when statistically evaluated,
showed no difference between them: MamaSize® Group: 284.04mL; Implant Group: 280.83mL; Control Group: 287.85mL (p-value: 0.6761).

Table 1 shows the comparison of the variables shape, size, symmetry and consistency in the preoperative and postoperative periods (12th month), showing a statistical difference. However, when the groups were compared between them regarding the four variables in the 12th postoperative month, there was no statistical difference. When comparing the patients’ scores for the scar variable between the 1st and 12th month in all three groups, no statistical significance was observed.

<table>
<thead>
<tr>
<th>VARIABLES</th>
<th>GROUP</th>
<th>PERIODS</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td></td>
<td>Preoperative</td>
<td>12th postoperative</td>
</tr>
<tr>
<td></td>
<td></td>
<td></td>
<td>Month</td>
</tr>
<tr>
<td>Shape</td>
<td>Control</td>
<td>38.07 (30.25-46.86)</td>
<td>98.29 (96.78-99.80)</td>
</tr>
<tr>
<td></td>
<td>Silicone</td>
<td>37.60 (25.67-49.53)</td>
<td>96.40 (93.77-99.03)</td>
</tr>
<tr>
<td></td>
<td>MamaSize®</td>
<td>42.00 (31.54-52.46)</td>
<td>97.60 (95.13-100.07)</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>21.82 (16.61-28.13)</td>
<td>95.00 (91.56-98.44)</td>
</tr>
<tr>
<td>Size</td>
<td>Silicone</td>
<td>22.40 (13.30-31.50)</td>
<td>93.40 (89.79-97.01)</td>
</tr>
<tr>
<td></td>
<td>MamaSize®</td>
<td>19.20 (10.87-27.53)</td>
<td>87.60 (80.77-94.47)</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>68.07 (58.38-79.25)</td>
<td>90.92 (86.25-95.59)</td>
</tr>
<tr>
<td>Symmetry</td>
<td>Silicone</td>
<td>62.80 (48.28-77.32)</td>
<td>92.80 (87.40-98.62)</td>
</tr>
<tr>
<td></td>
<td>MamaSize®</td>
<td>62.16 (48.81-75.51)</td>
<td>95.20 (90.72-99.68)</td>
</tr>
<tr>
<td></td>
<td>Control</td>
<td>52.73 (43.11-63.21)</td>
<td>97.50 (94.81-100.19)</td>
</tr>
<tr>
<td>Consistency</td>
<td>Silicone</td>
<td>63.20 (49.52-76.88)</td>
<td>94.00 (89.38-98.62)</td>
</tr>
<tr>
<td></td>
<td>MamaSize®</td>
<td>58.80 (46.00-71.60)</td>
<td>98.80 (96.99-100.61)</td>
</tr>
</tbody>
</table>

Figure 3 shows the preoperative and the 12th month postoperative aspects of a patient, in the control group, with a 285mL implant. Figure 4 shows a patient in the preoperative and in the 12th month postoperative periods, with a 285mL implant, and the methodology of choice with the silicone mold in the Silicone group. Figure 5 shows a patient in the preoperative and in the 12th month postoperative periods, with a 285mL implant, and the methodology of choice with the MamaSize® mold.

**Figure 3.** Control group: preoperative (A); postoperative (B).

**Figure 4.** Silicone Group: Preoperative (A), 12th month (B), Mold (C).
DISCUSSION

Because it is the second most often performed esthetic plastic surgery in Brazil and the first in the USA, augmentation mammoplasty reoperation rates due to volume exchange, which would firstly be relatively low (1.9% to 5.4%)⁹,¹², are significantly higher in absolute numbers. Therefore, predicting this volume and, thus, avoiding reoperations, in addition to adding less morbidity to the patient, would avoid an expense that, in the US for instance, is around US$.5770.00 per reoperation⁹,¹². The literature shows that up to 20% of patients in the postoperative period of breast augmentation surgery complain of breast volume, although not all of them want to reoperate¹⁷.

There are articles that study preoperative types of breast implant volume measurement. However, despite showing good results, no studies were found comparing them prospectively⁴,⁵,¹³,¹４,¹⁷-¹⁹. Thus, what would be the measurement methodology of preoperative breast implant volume with the lowest cost and the best benefit? Is there any difference in the postoperative evaluation by the patients, if they use these different preoperative methods discussed in the literature? As satisfaction is closely related to the patient’s expectations, and these vary according to the local culture, we believe that the Brazilian patients should be studied. Therefore, this study was then created, which utilized three easy, inexpensive methodologies widely used in the literature to compare the degree of patient satisfaction in the pre- and postoperative periods.

The sample results confirm the stereotype of these patients: they are young, with a mean age of 28 years and normal BMI, whose main dissatisfaction is breast size. The mean mammary basis was 11.62cm, with an anticipated implant volume of approximately 285mL. Therefore, anthropometrically similar patients were studied and compared.

When patient satisfaction was evaluated before and one year after the surgery, there was a significant increase in the degree of satisfaction in all groups, with statistical significance. This fact supports studies in the literature that show the excellent results of this intervention²,³,⁵. However, when comparing the degree of satisfaction in the postoperative evaluation between the three studied groups, there was no statistical difference, which shows that the method used to choose the implants does not interfere with the degree of satisfaction.

The scar is an important variable to be explained to the patient in the preoperative consultation, considering the change in location according to one’s culture: in the USA and Brazil, inframammary scars are more common; in China the axillary scar is more frequent⁶,⁸,⁹. In our study, in the 1st, 6th and 12th months, the evaluation mean was higher than 85. Additionally, there was no statistical difference when comparing the periods. This confirms the patients’ acceptance of the scar, even in the first postoperative month.

The choice made through anthropometric measures has been the routine in our service for several years. On the other hand, methods that directly include the patient in this choice have proved equally effective and can facilitate decision-making, as well as the sharing of the choice responsibility. This is likely to reduce reoperation rates for breast volume dissatisfaction after a few years. On the other hand, it does not reduce
mandatory reoperation rates in rare complications such as a late seroma. Therefore, guiding patients about all risks inherent to the procedure is essential.

New third-dimensional preoperative measurement techniques have been introduced in the market. These are devices with high added value, requiring company-specific software and hardware. On the other hand, these devices are still under evaluation, and a dissatisfaction degree with the final result of around 25% has already been verified.

Our study allowed us to conclude that augmentation mammoplasty with breast implant has a high index of satisfaction among the patients, but there was no difference in satisfaction in the postoperative period between the three breast volume measurement methods.

**RESUMO**

**Objetivos:** avaliar o grau de satisfação de pacientes submetidas à mamoplastia de aumento e comparar três métodos diferentes, fáceis, baratos e universais, de escolha pré-operatória de volume de implante mamário. **Métodos:** estudo prospectivo, realizado no Hospital Universitário Pedro Ernesto da Universidade do Estado do Rio de Janeiro, em 94 mulheres naturais do Rio de Janeiro, com idades entre 18 e 49 anos, e submetidas à cirurgia de mamoplastia de aumento com implante, por hipomastia. Todos os implantes eram texturizados, com base redonda e projeção alta e foram introduzidos na loja retroglândular, por via inframamária. As pacientes foram divididas em três grupos: Controle, Silicone e MamaSize®, com 44, 25 e 25 pacientes, respectivamente. Foram realizados questionários de satisfação nos períodos pré e pós-operatórios pelo mesmo avaliador, através de escala analógico-visual, em que 0 significava muito insatisfeita e 100 significava muito satisfeita para as quatro variáveis: forma, tamanho, simetria e consistência. No período pós-operatório avaliou-se também o grau de satisfação com a cicatriz cirúrgica. **Resultado:** quando comparados os graus de satisfação do pré-operatório com os do pós-operatório, houve diferença em todas as variáveis dos três grupos, com significância estatística. Entretanto, quando comparados os dados dos pós-operatórios entre si, não houve diferença significativa. O grau de satisfação com a cicatriz cirúrgica foi elevado. **Conclusão:** a mamoplastia de aumento com implante teve um grande índice de satisfação entre as pacientes. No entanto, não houve diferença no grau de satisfação no período pós-operatório entre as três metodologias de mensuração de volume mamário.


**REFERENCES**


Received in: 26/07/2017
Accepted for publication: 23/11/2017
Conflict of interest: none.
Source of funding: none.

Mailing address:
Rafael Daibert de Souza Motta
E-mail: rdsmotta@hotmail.com / rafaelsmotta@gmail.com

© by