Mesh Surgery for Anterior Vaginal Wall Prolapse: A Meta-analysis

Cirurgia com tela para correção de prolapso de parede anterior: metanálise

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Abstract

Purpose Pelvic organ prolapse (POP) is a major health issue worldwide, affecting 6–8% of women. The most affected site is the anterior vaginal wall. Multiple procedures and surgical techniques have been used, with or without the use of vaginal meshes, due to common treatment failure, reoperations, and complication rates in some studies.

Methods Systematic review of the literature and meta-analysis regarding the use of vaginal mesh in anterior vaginal wall prolapse was performed. A total of 115 papers were retrieved after using the medical subject headings (MESH) terms: ‘anterior pelvic organ prolapse OR cystocele AND surgery AND (mesh or colporrhaphy)’ in the PubMed database. Exclusion criteria were: follow-up shorter than 1 year, use of biological or absorbable meshes, and inclusion of other vaginal wall prolapses. Studies were put in a data chart by two independent editors; results found in at least two studies were grouped for analysis.

Results After the review of the titles by two independent editors, 70 studies were discarded, and after abstract assessment, 18 trials were eligible for full text screening. For final screening and meta-analysis, after applying the Jadad score (> 2), 12 studies were included. Objective cure was greater in the mesh surgery group (odds ratio [OR] = 1.28 [1.07–1.53]), which also had greater blood loss (mean deviation [MD] = 45.98 [9.72–82.25]), longer surgery time (MD = 15.08 [0.48–29.67]), but less prolapse recurrence (OR = 0.22 [0.13–0.38]). Dyspareunia, symptom resolution and reoperation rates were not statistically different between groups. Quality of life (QOL) assessment through the pelvic organ prolapse/urinary incontinence sexual questionnaire (PISQ-12), the pelvic floor distress inventory (PFDI-20), the pelvic floor impact questionnaire (PFIQ-7), and the perceived quality of life scale (PQOL) was not significantly different.

Keywords
► anterior pelvic organ prolapse
► cystocele
► surgery
► mesh
► colporrhaphy
► meta-analysis

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Introduction

Pelvic organ prolapse (POP) is a major health issue worldwide, and even nowadays it represents a great challenge to modern gynecology. In epidemiologic surveys, “vaginal bulge” sensation reaches up to 6–8% of all female interviewees; and if pelvic examination alone is considered, the numbers rise to 30–60% of all women. POP diagnosis goes back to Antiquity, and multiple treatments and theories have been tried and abandoned over the years, with frequent recurrence and uncertain success rates.

The anterior vaginal wall is the most commonly affected, being responsible for up to 80% of all POP surgical procedures. The possible reasons for this dominance lie on its anatomical characteristics, in that the anterior vaginal wall lies on a more horizontal position in the female pelvis, suffering further gravitational pressure, without having any muscle support; moreover, the tissue separating the anterior vaginal wall from the bladder is comparatively thinner and more distensible than that enfolding the other vaginal walls.

Several surgical techniques have arisen and been discharged over the years. High recurrence rates and complications in POP surgery led to the development of meshes for the anterior vaginal wall since the 1950’s – from biological meshes, autologous (fascia) and heterologous (porcine), to synthetic ones, both absorbable and non-absorbable, with conflicting results and alarming complication rates in some series.

Recent studies revealed that synthetic meshes seem superior to biological ones, and both produce better anatomical outcomes when compared with simple colporrhaphy – however, with higher complication rates, such as larger intraoperative bleeding, slower surgical time and greater extrusion rates, and apparently without significant differences in the quality of life questionnaire assessment, although long follow-up studies on this issue are rare.

The main goal of this meta-analysis is to evaluate the effectiveness of POP surgery and its complications, emphasizing the anterior vaginal wall, with and without mesh use.
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Methods

A systematic review and meta-analysis was conducted on the effectiveness and complications regarding women treated for anterior vaginal wall prolapse surgically, comparing mesh treatment with traditional surgery.

For a complete original article search, the following steps were covered, according to the preferred reporting items for systematic reviews and meta-analyses (PRISMA) statement – search in the PubMed database with the medical subject headings (MESH) terms: ‘anterior pelvic organ prolapse OR cystocele AND surgery AND (mesh or colporrhaphy)’.

Inclusion criteria: Randomized controlled trials, in English, published in the past 15 years, including only women with anterior vaginal wall prolapse, comparing surgical treatment without mesh versus using synthetic absorbable meshes.

A total of 115 articles, after filtering results by species (human), gender (female) and language (English), were recovered. The last online search was performed on June 20th, 2015. After excluding duplicate series, identified by abstract reading, a total of 109 articles were included for final screening. The inclusion process was performed by two independent reviewers (EBC, and LCS).

Exclusion criteria were the following: articles regarding other vaginal walls or incontinence as a main result; follow-up shorter than one year; anatomic evaluation made without the use of the pelvic organ prolapse quantification (POP-Q) rating; literature or systematic reviews; studies using biological or absorbable meshes; and uncontrolled/non randomized trials.

Initially, the articles were selected by their titles/abstracts; afterwards, full text articles that potentially met all the inclusion criteria were assessed. Whenever there was lack of accordance between the two reviewers, a third researcher was consulted (CRT). After this initial screening, included articles were submitted to Jadad criteria, and only papers for which the Jadad score was greater than 2 were finally included.

In the papers where the inclusion criteria were met, assessed data were: objective cure; surgery time; intra-operative blood loss; recurrence rate; reoperation rate (for surgical failure); “vaginal bulge” complaint resolution rate; dyspareunia; and subjective cure through quality of life questionnaires – pelvic organ prolapse urinary incontinence sexual questionnaire (PISQ-12); pelvic floor distress inventory (PFDI-20); pelvic floor impact questionnaire (PFIQ-7); and perceived quality of life scale (P-QOL).

Outcomes verified in more than two articles were grouped for meta-analysis. A chart for data collection was created to extract the data of interest in each article, which were then retyped to a single database by two independent researchers, to avoid loss of data or mistyping of any kind. The bias risk was assessed by the use of a Cochrane Collaboration tool.

The statistical analysis for outcomes measured in proportion (dichotomous) used the Mantel-Haenszel test, assessing the risk ratio between the groups. The statistical analysis for quantitative outcomes (with averages and standard deviations) assessed the mean difference between the groups.

The heterogeneity between studies was assessed through Chi-square test of study homogeneity and through the I² index, which varies from 0–100%, where: 0–40%: there is no significant heterogeneity; 30–60%: there may be moderate heterogeneity; 50–90%: there may be substantial heterogeneity; 75–100%: there is plenty of heterogeneity between studies.

For studies with heterogeneity superior to 50% and p < 0.05, a random effect meta-analysis was performed. The computer system utilized was the Review Manager 5.3 (IKMD, Copenhagen, Denmark).

Results

Search strategy crossing the terms “Anterior pelvic organ prolapse OR cystocele AND surgery AND (mesh or colporrhaphy)” was conducted in the PubMed online database. A total of 115 articles were retrieved, which came down to 109 after the exclusion of repeated series through title/abstract analysis. Afterwards, 70 articles were discarded by the title due to failure to meet the inclusion criteria.

Two independent reviewers assessed the 39 remaining abstracts, and finally 18 papers were included for eligibility analysis. After full article evaluation, another 4 articles were excluded, with 14 remaining to be submitted to Jadad score assessment. Two more articles were excluded for Jadad scores under 3, with 12 articles included for final analysis (Table 1).

Objective Cure

For objective cure analysis, all 12 articles were included (1,540 women), with 86% heterogeneity (p < 0.00001). Patients in the mesh surgery group were considered cured in 551/781 cases, whereas 418/759 women were found cured in the surgery without mesh group, with 1.28 (confidence interval, CI = 1.07–1.53) greater objective cure risk in the group submitted to mesh surgery, after aleatory effect meta-analysis (Fig. 2).

Surgery Duration

For the analysis of surgery duration, 5 articles were included (779 women), with great heterogeneity. After aleatory effect meta-analysis, the group of women submitted to surgery without mesh was found to have less surgical time, with mid difference of 15.08 minutes (CI = 0.48–29.67).

Blood Loss

For blood loss assessment, 3 articles were included (631 women), with substantial heterogeneity. After aleatory effect analysis, the mesh surgery group was found to have greater blood loss, with a mean difference of 45.98 ml (CI = 9.72–82.25).

Prolapse Recurrence

Five articles were included for recurrence rate analysis (585 women), without significant heterogeneity among them. The
Fig. 1 Articles inclusion criteria or pelvic organ prolapse surgery and its complications.
Table 1  Articles that evaluated effectiveness of Pelvic Organ Prolapse surgery and its complications

<table>
<thead>
<tr>
<th>Articles</th>
<th>N</th>
<th>Mesh</th>
<th>Follow-up</th>
<th>Objective cure (POP-Q)</th>
<th>QOL</th>
<th>Subjective cure</th>
<th>Complications</th>
<th>Complication rates</th>
<th>Conclusion</th>
</tr>
</thead>
<tbody>
<tr>
<td>Altman et al¹⁴</td>
<td>410</td>
<td>Prolift</td>
<td>1 year</td>
<td>S = 2 / UD (question 16) 60.8 ± 34.5%</td>
<td>UDI</td>
<td>PIQ-12</td>
<td>“asymptomatic”</td>
<td>Duration 52.6 ± 13.5 min Blood loss 84.7 ± 35.4 mL Bladder perforation 7 ± 1 Pain 5 ± 0 Retention 16 ± 6 Gynecotomy 11 ± 1</td>
<td>Exclusion 3.2% Greater short time success with greater complications</td>
</tr>
<tr>
<td>de Tayrac et al¹⁵</td>
<td>163</td>
<td>Uptyflex</td>
<td>1 year</td>
<td>S = 2 / 93.9 %</td>
<td>UDI</td>
<td>PFDI-20</td>
<td>“satisfied”</td>
<td>Duration 99.1 ± 46 minutes Hospital stay 3.2 ± 3.3 days Abnormal bleeding 45 ± 30.8% Dyspareunia 5 ± 10%</td>
<td></td>
</tr>
<tr>
<td>Delnoy et al¹⁶</td>
<td>79</td>
<td>Naica TC</td>
<td>1 year</td>
<td>S = 2 / 82.5 ± 56.4%</td>
<td>UDI</td>
<td>P-QOL</td>
<td>General improvement (not significant)</td>
<td>Duration 75 ± 76 minutes Hospital stay 2.6 ± 2.6 days Blood loss 215.5 ± 219.5 mL Bladder perforation: 0</td>
<td>Exclusion 9.5% Greater anatomic success with slight increase in complication rates</td>
</tr>
<tr>
<td>El-Nazer et al¹⁷</td>
<td>40</td>
<td>Cynomesh</td>
<td>2 years</td>
<td>S = 2 / 95 ± 70% PFMS (perineal strength)</td>
<td>UDI</td>
<td>P-QOL</td>
<td>“surgical” resolution 95.3 ± 66.7% Voiding dysfunction 9% “Feels better”</td>
<td>Duration 73 ± 58 minutes Hospital stay 4.4 ± 4.6 days Abnormal bleeding 0 ± 3% Retention 0 ± 5%</td>
<td>Exclusion 5% Recurrence 4 ± 15%</td>
</tr>
<tr>
<td>Gutman et al¹⁸</td>
<td>65</td>
<td>Prolift</td>
<td>3 years</td>
<td>S = 2 / 45 ± 43%</td>
<td>UDI</td>
<td>PIQ-12</td>
<td>“surgical” resolution 92 ± 81% “Feels better” 88 ± 81%</td>
<td>Duration 69.7 ± 74.6 minutes Hospital stay 4.4 ± 4.6 days Abnormal bleeding 0 ± 3% Retention 0 ± 5%</td>
<td>Exclusion 15% Recurrence 13 ± 0%</td>
</tr>
<tr>
<td>Hiltunen et al¹⁹</td>
<td>202</td>
<td>Parietene</td>
<td>1 year</td>
<td>S = 2 / 93.3 ± 61.5%</td>
<td>UDI</td>
<td>PIQ-7</td>
<td>“surgical” resolution 93 ± 94% Voiding dysfunction 6% “Feels better” 23 ± 10%</td>
<td>Duration 73 ± 58 minutes Hospital stay 4.4 ± 4.6 days Abnormal bleeding 0 ± 3% Retention 0 ± 5%</td>
<td>Exclusion 17.3% Recurrence 4 ± 15%</td>
</tr>
<tr>
<td>Lamblin et al²⁰</td>
<td>68</td>
<td>Penigree</td>
<td>2 years</td>
<td>S = 2 / 100 ± 64.4%</td>
<td>UDI</td>
<td>PFDI-20</td>
<td>“surgical” resolution 95 ± 99% Voiding dysfunction 9% De novo incontinence 23 ± 10%</td>
<td>Duration 69.7 ± 74.6 minutes Hospital stay 4.4 ± 4.6 days Abnormal bleeding 0 ± 3% Retention 0 ± 5%</td>
<td>Exclusion 6% Recurrence 0 ± 11.7%</td>
</tr>
<tr>
<td>Nguyen et al²¹</td>
<td>76</td>
<td>Perigree</td>
<td>2 years</td>
<td>S = 2 / 89 ± 55%</td>
<td>UDI</td>
<td>PFDI-20</td>
<td>“surgical” resolution 95 ± 99% Voiding dysfunction 9% De novo incontinence 23 ± 10%</td>
<td>Duration 69.7 ± 74.6 minutes Hospital stay 4.4 ± 4.6 days Abnormal bleeding 0 ± 3% Retention 0 ± 5%</td>
<td>Exclusion 6% Recurrence 0 ± 3%</td>
</tr>
<tr>
<td>Nieminen et al²²</td>
<td>202</td>
<td>Parietene</td>
<td>3 years</td>
<td>S = 2 / 91 ± 65%</td>
<td>UDI</td>
<td>Non validated</td>
<td>Score “bulge” 1.16 ± 1.43 De novo incontinence 7 ± 5%</td>
<td>Exclusion 10% Recurrence 13 ± 41% Recurrence 11 ± 18%</td>
<td>Better anatomical results with equivalent symptom resolution, but high extrusion</td>
</tr>
<tr>
<td>Svalnoioglu et al²³</td>
<td>90</td>
<td>Parietene</td>
<td>1 year</td>
<td>S = 2 / 91 ± 72%</td>
<td>UDI</td>
<td>P-QOL</td>
<td>Voiding dysfunction 4.6 ± 0% De novo incontinence 0 ± 7%</td>
<td>Duration 69.7 ± 74.6 minutes Hospital stay 4.4 ± 4.6 days Abnormal bleeding 0 ± 3% Retention 0 ± 5%</td>
<td>Exclusion 6% Failure 9.3 ± 28.5%</td>
</tr>
<tr>
<td>Tamanini et al²⁴</td>
<td>100</td>
<td>Naica TC</td>
<td>1 year</td>
<td>S = 2 / 83.7 ± 55.5%</td>
<td>UDI</td>
<td>IOQ-7</td>
<td>Voiding dysfunction 2.3 ± 0% Bladder perforation (&lt; 500 ml) 2.3 ± 1.8%</td>
<td>Duration 73 ± 58 minutes Hospital stay 4.4 ± 4.6 days Abnormal bleeding 0 ± 3% Retention 0 ± 5%</td>
<td>Exclusion 9.5%</td>
</tr>
<tr>
<td>Vollebregt et al²⁵</td>
<td>125</td>
<td>Avista</td>
<td>1 year</td>
<td>S = 2 / 91 ± 41%</td>
<td>UDI</td>
<td>RIQ</td>
<td>“surgical” resolution 95 ± 99% Voiding dysfunction 9% De novo incontinence 23 ± 10%</td>
<td>Duration 69.7 ± 74.6 minutes Hospital stay 4.4 ± 4.6 days Abnormal bleeding 0 ± 3% Retention 0 ± 5%</td>
<td>Exclusion 4% Recurrence 0 ± 5%</td>
</tr>
</tbody>
</table>

Abbreviations: ICIQ-VS, international consultation of incoherence modular questionnaire vaginal symptoms; IIQ, incontinence impact questionnaire; MUH, urinary dysfunction measurement scale PGI-I, patient global impression of improvement; S, stage; UDI, urogenital distress inventory; PFDI, pelvic floor distress inventory; PFDI-20, 20 questions; PIQ, pelvic floor impact questionnaire; PIQ-7, seven questions; PFMS, pressure and fluid management symptoms; PIQ-12, pelvic organ prolapse/ urinary incontinence sexual questionnaire; POP-Q, pelvic organ prolapse quantification; QOL, quality of life; UDI, urogenital distress inventory; UDI-0, obstructive discomfort; UDI-5, irritative symptoms; UDI-5, stress symptoms;
mesh surgery group had 19 recurrences in 300 patients, while the group without mesh had 62 cases in 285 women – the mesh group had less recurrence: OR $= 0.22$ (CI $= 0.13–0.38$) ($\sim$Fig. 3).

**Prolapse Related Reoperation**

Three articles were considered for this analysis, totaling 357 women. Mesh surgery patients were reoperated 15/183 cases, whereas in the group without mesh, 21/174 patients had new surgical POP related procedures, with no significant difference between groups: OR $= 0.65$ (CI $= 0.83–4.51$) ($\sim$Fig. 4).

**Subjective cure**

For the analysis of subjective cure rates, considering “vaginal bulge” symptom resolution, 3 articles were selected (282 women). The mesh surgery group had vaginal bulge resolution in 135/144 patients, whereas 122/138 women in the group without mesh reported bulge resolution, without statistical difference: OR $= 0.65$ (CI $= 0.83–1.29$).

**Dyspareunia**

For dyspareunia rate evaluation, 6 articles were included (397 women), without significant heterogeneity. In the mesh group, dyspareunia was found in 15/193 patients, while it was found in 16/204 women in the group without mesh, showing no statistical difference: OR $= 0.94$ (CI $= 0.45–1.96$).

**Quality of life validated questionnaires ($\sim$Fig. 5)**

For sexual function analysis, through the PISQ-12 questionnaire, 3 articles were enrolled, with a total of 612 women. The mean difference between groups was $-0.38$ (from $-3.12$ to $2.37$), without statistical significance.

The subjective assessment through the P-QOL questionnaire included 2 articles, with 164 women. There was no significant heterogeneity among studies. The results showed no statistical difference in quality of life after surgery with or without mesh, with a mean difference of $-0.94$ (from $-3.31$ to $1.42$).

The PFIQ-7 quality of life questionnaire evaluation included 2 articles, totaling 137 women. There was no significant difference between groups, with a mean difference of $-0.77$ (from $-14.25$ to $12.74$) after aleatory effect analysis.

Finally, the PFDI-20 questionnaire was analyzed including 2 articles (139 women), also without statistical difference between groups, with a mean difference of $0.20$ (from $-19.95$ to $19.66$) after aleatory effect analysis.

**Discussion**

This meta-analysis showed that the use of meshes for the treatment of anterior vaginal wall prolapse is superior to native tissue surgical repair, when objective cure through POP-Q assessment is considered. All of the articles included in this meta-analysis considered as an objective cure criterion a post-surgical POP-Q stage inferior to 2 in the pelvic examination. This definition of cure, that is, when the prolapse’s maximum extent stays at least 1 cm above the hymenal ring, is based on the recommendations of the US National Institute of Health (NIH) workshop. However, this POP-Q based cure assessment has been criticized by many authors – many discrete or moderate prolapses are not classified using this criterion as cure or...
absence of prolapse, although they have no impact on women’s quality of life and, therefore, demand no intervention.  

For the prolapse patient, the main goal is symptom relief and quality of life improvement; accordingly, quality of life assessment in these cases seems more trustworthy than the POP-Q classification alone.

There are several different ways to assess quality of life, with the use of subjective criteria or validated questionnaires. This meta-analysis showed no significant results when evaluating the “vaginal bulge” symptom resolution, with no statistical difference between the groups with or without the use of mesh; but an aleatory effect analysis had to be made due to the great heterogeneity among studies.

Aleatory effect meta-analysis adds variance to the general effect proportionally to the variability of the results in the studies; that makes the confidence intervals of the estimated summary measure be bigger than those of the fixed effect meta-analysis.

When we evaluated quality of life using 4 different validated questionnaires (PISQ-12, P-QOL, PFIQ-7 and PFDI-20), we observed no significant difference in quality of life rating between women submitted to prolapse surgery with or without mesh use. A previous Cochrane Meta-analysis also concluded that although anterior vaginal mesh placement shows better anatomical results, it does not promote better functional outcomes.

Furthermore, our study showed that women submitted to mesh surgery have lower prolapse recurrence rates (80% less), without heterogeneity among analyzed articles, which increases this result’s credibility. The risk of a woman having to go through surgery for pelvic prolapse during her entire life is ~6–19%, with 30% chance of a new intervention for recurrence of the prolapse.

Moreover, reoperation rates after reconstructive prolapse surgery are very high (43–58%), which led to the use of synthetic meshes in the last decades, in an attempt to lower those flawed outcomes. This meta-analysis shows that the use of vaginal meshes is effective in diminishing recurrence rates; nevertheless, with increased surgery duration and blood loss, which may increase procedure-related morbidity.

The use of vaginal meshes is not free of risk. The main complications are: vaginal mesh erosion; infection; granulomas; dyspareunia; vesicovaginal fistula; and chronic pelvic pain. In 2008, the American Food and Drug Administration (FDA) released its first notification about the complications associated to the use of synthetic meshes, after receiving more than one thousand complication reports. In 2011, the FDA released two more statements about indications and security concerning mesh use.

This meta-analysis revealed a 7.4% medium rate of mesh erosion (3.2–19%). Mesh erosion or exposure rate varies with follow-up time, being observed from 6 weeks up to 7 years after the surgical procedure. Studies included in this meta-analysis had at least 1 year of follow-up; however, this may be considered a short time for this particular evaluation, explaining the low rate of extrusion observed.

Therefore, albeit the benefit of less prolapse recurrence, mesh surgery should be evaluated carefully, given the fact of morbidity elevation (longer surgeries and greater blood loss) and related extrusion rates. Furthermore, there was no difference in reoperation rates comparing mesh versus no mesh surgery – the most probable explanation for this fact is that maybe recurrence did not affect substantially those women’s quality of life to the point that a new surgical approach was needed.

There was no significant difference in de novo dyspareunia rates between mesh and no-mesh groups. Dyspareunia is, as stated above, one of the main complications associated with the use of vaginal meshes, and is a very important aspect to be considered in women with prolapse, especially those who wish to maintain a sexual healthy sexual life after surgery, being directly related to patient satisfaction and quality of life. This meta-analysis also showed that there was no statistical difference in quality of sexual life in women with and without mesh, evaluated through the PISQ-12 questionnaire. The fact that women submitted to mesh surgery did not show greater dyspareunia rates or significant mesh erosion in our report may have contributed to this sexual life equality between groups.
Conclusions

Female genital prolapses are a great cause of quality of life prejudice, with sexual, psychological and functional (defecation and micturition) impairment. Costs related to female POP surgery are high and growing worldwide. A 46% rise in corrective pelvic surgeries is expected in the next decades, due mainly to population aging.

The choice of an effective and safe treatment for genital prolapse is really important, and another fact to be taken into consideration, especially in the anterior vaginal wall defects, is that they are frequently associated with defects of the vaginal apex. Women with anterior vaginal stage 3 (POP-Q) prolapses have up to 98% chance of having a concomitant uterus/vault prolapse of at least the first stage – this association and its correction were not taken into consideration in many studies assessed in this meta-analysis.

Despite many strong points, such as inclusion of only randomized controlled trials of the past 15 years with minimum follow-up of 1 year, and with paper quality addressed by Jadad criteria, this meta-analysis has some limitations, mainly because of study heterogeneity, great difference in quality of life evaluation among studies, short follow-up for the appearance of certain late complications, such as mesh extrusion and recurrence, and the lack of a clear apex evaluation/treatment in the trials.

Future studies with longer follow-up and inclusion of an apex thorough evaluation and concomitant treatment when operating the anterior vaginal wall are necessary to better enlighten the definitive role of vaginal meshes in this scenario.

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