Complications of laparoscopic sacrocolpopexy: as harmless as they seem?
Complicações da sacrocolpexia laparoscópica: serão assim tão inofensivas?

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INTRODUCTION

Abdominal sacrocolpopexy (ASC) has been shown to be more effective than a vaginal approach surgery on the treatment of apical vaginal prolapse\(^1,2\), with reported long-term success rates of 68–100%\(^3\). Furthermore, it also allows simultaneous treatment of other compartment defects\(^4\). Laparoscopic sacrocolpopexy avoids the need for a large abdominal incision and minimizes bowel manipulation, potentially leading to less postoperative pain and shorter recovery time\(^5\).

The use of synthetic mesh, while essential for the success rate, is associated to a complication unique to this kind of repair: the possibility of erosion through adjacent tissue. Mesh erosion rates after sacrocolpopexy range from 2% to 10% according to literature\(^6\)–\(^9\), with a wide range of consequences. Despite the widespread concern about mesh related complications, their treatment is not extensively discussed.

Risk factors associated with mesh erosions include smoking\(^7\), prior surgical scarring\(^10\), oestrogen deficiency\(^11\), concomitant hysterectomy\(^7\), mesh type\(^12\) and a transvaginal placement of the mesh\(^13\).

Mesh erosion is one of the most worrisome complications related to sacrocolpopexy, especially because of the impact on quality of life. However, a series of other complications may be associated to this procedure.

Complications of sacrocolpopexy may be broadly divided in two main groups\(^8\):

- general surgical complications (procedure or surgeon-based) – related to surgical technique, as dissection...
or haemostasis; although not related to the mesh itself, they can influence surgery outcome.

• mesh-related complications (product-based) – directly attributed to the presence of mesh inside the patient’s body.

The purpose of this study is to alert to possible complications after laparoscopic sacropexy, as well as treatment options, by analysing all cases of patients who needed surgical treatment of complications after laparoscopic sacropexy in our department. At the same time, other surgery-related complications are described, with the aim of alerting surgeons to their occurrence, and therefore contributing to their prevention.

METHODS

After receiving institutional review board approval, a retrospective analysis of electronic medical records was performed for all patients who underwent surgical treatment for Grade III-b complications (according to Clavien-Dindo Grading System for the Classification of Surgical Complications) after sacropexy (sacro-cervicopexy, sacrocolpopexy or sacrohysteropexy) with mesh to treat pelvic organ prolapse by laparoscopic approach, from January 1998 to December 2013. Sacropexies were performed by 5 different experienced laparoscopic surgeons. Only complications treated in our institution were considered.

Medical records of these patients were retrieved to collect clinical data (BMI, menopausal status, hormonal replacement therapy (HRT) usage, topical oestrogens usage, prior hysterectomy, smoking), and operative details were abstracted (concomitant hysterectomy – total or subtotal –, type of mesh used, fixation technique, peritonization, perioperative antibiotic prophylaxis), along with complications and its management. Type of complications, clinical presentation, delay before surgical management, initial and eventual subsequent approaches and postoperative data were assessed.

RESULTS

A total of 1238 sacropexies were performed between January 1998 and December 2013, in our Department. Mean follow up time was 41.3 months, with 16 patients lost to follow up.

A total of 33 patients (2.7%) had a Grade III-b complication (N = 33).

NOTE: Women for whom data on any field were not recorded were excluded from the descriptive analysis of the respective parameter. Ratios (n/N) and percentages (%) are presented according to available data.

Patients’ characteristics

Mean age was 52.0 ± 8.3 [42-75] years.

There were 2/28 (7.1%) obese patients, and 9/28 (32%) were overweight. Mean body index mass was 24.1 ± 3.5 [18-32] Kg.m⁻². Most of the patients (24/33) were postmenopausal (72.7%), and 13/24 (54.2%) were using systemic hormonal replacement treatment. Six out of 31 (19.4%) had current smoking habits.

Fourteen out of 33 (42.4%) had undergone a previous hysterectomy (total abdominal hysterectomy in 7 patients, total laparoscopic hysterectomy in 6 and subtotal laparoscopic hysterectomy in 1).

Intraoperative relevant findings/incidents

In surgical reports, 4/33 (12.1%) cases of through-and-through vaginal stitches were found. This fact was noticed intraoperatively and immediately corrected, either by replacing the suture laparoscopically or by vaginally cutting the visible suture. Two cases of difficult vesicovaginal dissection were reported, as well as 2 cases of adhesions between the sigmoid and the vaginal vault before the beginning of the procedure, 2 cases of uterine perforation at the time of cannulation and 1 pelvic endometriosis.

Complications

Complications are presented in Table I.

There were 6 general surgical complications (2 haematomas, 2 peritonitis, 1 mechanical bowel obstruction and 1 vaginal wall necrosis); and 34 mesh related complications (27 vaginal mesh exposures, 4 periprosthetic abscesses, 1 vesicovaginal fistula, 1 rectovaginal fistula and 1 symptomatic mesh retraction). When accessing the interval of time to diagnosis, 8 early postoperative complications (during the first month) were reported (2 peritonitis, 2 haematomas, 1 mechanical bowel obstruction, 1 vesicovaginal fistula, 1 vaginal wall necrosis, 1 vaginal mesh exposure) along with 32 late postoperative complications (diagnosed more than one month after surgery) (26 vaginal mesh exposures, 4 periprosthetic abscesses, 1 rectovaginal fistula and 1 symptomatic mesh retraction).

Clinical presentation

Clinical presentations are presented in Table II.
## TABLE I. COMPLICATIONS

<table>
<thead>
<tr>
<th>Patient</th>
<th>Type of prosthesis</th>
<th>Fixation</th>
<th>Hysterectomy</th>
<th>Intraoperative relevant findings</th>
<th>Postoperatively topical estrogen usage</th>
<th>Type of complication</th>
<th>Type of approach on complication solving/time to surgery, according to complication (months)</th>
<th>Total no. of surgeries needed</th>
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<tbody>
<tr>
<td>1</td>
<td>Polyester</td>
<td>Suture</td>
<td>Concomitant STLH</td>
<td>Uterine perforation</td>
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<td>Exposure</td>
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<td>Vaginal exposure</td>
<td>Periprosthetic abscess *</td>
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<td>Vaginal / 8</td>
<td>Vaginal / 8</td>
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*Continues on the next page*
## TABLE I. COMPLICATIONS

<table>
<thead>
<tr>
<th>Patient</th>
<th>Type of prosthesis</th>
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<th>Hysterectomy</th>
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<th>Postoperatively topical estrogen usage</th>
<th>Type of complication</th>
<th>Type of approach on complication solving/time to surgery, according to complication (months)</th>
<th>Total no. of surgeries needed</th>
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<td>Laparoscopy / 1</td>
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<td>Vaginal / 11</td>
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<td>Concomitant STLH</td>
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<tr>
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<td>Suture</td>
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<td>Concomitant STLH</td>
<td>-</td>
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<td>Vaginal exposure + infection</td>
<td>Vaginoscopy / 23</td>
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<td>27</td>
<td>-</td>
<td>-</td>
<td>(No previous hysterectomy)</td>
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<td>Vaginal / 48</td>
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<td>-</td>
<td>Concomitant STLH</td>
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<td>-</td>
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<td>Vaginoscopy / 39</td>
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<td>Previous TAH</td>
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<td>Peritonitis Recurrence</td>
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<td>Polypropylene</td>
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<td>Laparoscopy / 0.17, Laparoscopy / 2</td>
<td>2</td>
<td></td>
</tr>
</tbody>
</table>

Notes: ‘complication subsequent to complication-solving surgery.’

Literature described risk factors for vaginal mesh exposure are signaled in bold.
Management of complications
Mean time between prolapse and first surgery for complication treatment was 18.2 ± 25.3 months [3 days–131 months]. A mean of 1.8 ± 1.5 [1–6] surgeries were needed for complication management, with eleven patients having more than one procedure. Considering only these 11 patients, the mean time between the prolapse surgery and the last surgery for complication resolution was 64.7 ± 49.1 [2–162] months, and the mean number of surgeries 3.4 ± 1.6 [2–6].

Twenty-two out of 27 patients (81.5%) had a pure non-infected vaginal mesh exposure. In 14/22 (63.6%) patients, an initial medical treatment was attempted (with topical oestrogens and antiseptics), though unsuccessful, with persistence of mesh exposure. For 13/14 (92.9%) patients, a vaginal approach was the next step, with exposed mesh removal and vaginal wall resuture. Among these patients, 5/12 (41.7%) needed a subsequent laparoscopic approach, due to persistent exposure; 4/5 with superimposed infection. At the end, of those 22 patients initially diagnosed with non-infected vaginal mesh exposure, 11 patients (50%) had the vaginal mesh exposure resolved after vaginal surgery, while 10 patients (45.5%) needed at least one laparoscopic surgery and 1 patient (4.5%) needed a laparotomy, due to heavy bleeding during mesh removal, and difficulties in the identification of anatomical structures. A total of 6 local infections were posteriorly diagnosed among the patients initially diagnosed with vaginal mesh exposure without associated infection.

The remaining 5/27 patients (18.5%) presented a vaginal mesh exposure with concomitant signs of local infection at the time of diagnosis (vaginal malodorous discharge, vaginal wall erythema). Two out of these 5 patients were initially treated with topical treatment, 2 with a vaginal approach and only 1 with an initial laparoscopic approach. At the end, just a single patient had its complication resolved after vaginal surgery; all the remaining needed, at least, one laparoscopic surgery (Figure 1).

The initial approach for fistulas (vesicovaginal and rectovaginal), cases of peritonitis or periprosthetic abscesses, bowel incarceration, haematomas, and mesh retraction, was laparoscopic; the same applied to cases of vaginal wall necrosis without mesh exposure.

Whenever a laparoscopic approach was performed, partial or total mesh removal was carried out (21/33 patients; 63.6%). Total mesh removal was attempted in all cases of concomitant infection. Five patients needed subsequent surgeries to remove remaining mesh pieces causing persistent infection.

Two patients also needed subsequent laparoscopies to resolve complications related to the mesh removal (one partial necrosis of the right ureter, needing reimplantation, and one haematoma of the right pararectal fossa).

All patients with suspected or confirmed infection were studied with abdominal CT or MRI, searching for eventual abscess collection or spondylodiscitis. No cases of spondylodiscitis were found.

DISCUSSION
Pelvic organ prolapse (POP) is common and seen on examination in 40% to 60% of parous women, potentially diminishing their quality of life. In the United States, the updated lifetime risk of undergoing prolapse surgery is 12.6%.

According to a recent Cochrane review, sacrocolpopexy was associated with a lower rate of recurrent vault prolapse on examination and painful intercourse than sacrospinous suspension, and a higher success rate and lower reoperation rate than high vaginal uterosacral suspension and transvaginal polypropylene mesh. A recent publication actually states ASC as the most effective treatment for apical vaginal prolapse, with reported long-term success rates of 68–100%. Plus, an abdominal approach allows a simultaneous correction of the three pelvic floor compartments defects: anterior, apical and posterior, preserving vaginal...
integrity. The laparoscopic approach represents an alternative to open surgery, with comparable outcomes, while benefitting patients with the well-recognized advantages of minimally invasive surgery. The characteristics of this completely minimally invasive surgery, as well as its potential benefits for sexual function (preservation of vaginal length and axis and lower rates of dyspareunia), make this procedure a better option for younger, sexually active women.

Being the main responsible for the effectiveness and durability of the procedure, the use of synthetic mesh carries, nevertheless, a set of possible related complications, unique to this procedure. Those complications frequently require surgical treatment. A reoperation is not necessarily related to mesh placement; it can be secondary to dissection, which is necessarily wider for abdominal mesh placement, comparing to vaginal approach. The rate of type III-b complications according to the Dindo classification seems to be higher after vaginal than abdominal mesh insertion (7% vs 4.8%). Understanding the subjacent mechanisms of mesh complications should help us to reduce their occurrence.

Before discussing mesh-related complications, general surgical complications will be considered. The distinction between general surgical complications vs mesh related complications seems reasonable and facilitative of complication analysis. The first group will be mandatorily specific to the abdominal approach, needing transperitoneal dissection, and thus required to be considered separately from mesh related complications.

Two cases of peritonitis occurred, both reoperated on day 3. In both patients previous abdominal hysterectomy and need for adhesiolysis was reported, which may have contributed to unnoticed bowel injury during dissection. In a study focusing on gastrointestinal complications of laparoscopic sacrocolpopexy, Warner et al. report a rate of 1.3% for intraoperative bowel injury. Surprisingly, prior abdominal surgery was positively associated with functional gastrointestinal complications, but not with bowel injury. Nonetheless, it is prudent to consider that those patients that present with previous surgeries, particularly total abdominal hysterectomy, and with multiple adhesions, mainly at vaginal cuff level, have an increased risk for bowel injury at the time laparoscopic sacrocolpopexy.

Simultaneously, the risk of vaginal wall necrosis may also be increased. The increased fragility of vaginal wall after dissection (always difficult in these cases), aligned with excessive bipolar coagulation, may explain this. Preoperative local oestrogen treatment should be considered, especially in these patients, as it is proved that
increases vaginal wall thickness\textsuperscript{11}.

Two patients presented early haematomas. One patient was diagnosed with vaginal suture exposure (later complicated by infection) more than 8 years after the index surgery, and the other patient presented a subsequent periprosthetic abscess, both demanding complete mesh removal, achieved laparoscopically. The overall median rate of bleeding complications was described by Nygaard as 4.4\%\textsuperscript{3}. The presence of a haematoma had already been described as a potential risk factor for mesh erosion\textsuperscript{19}.

A patient needed a laparoscopy on day 3 due to a small bowel incarcerated hernia in the Retzius space, dissected for a paravaginal defect repair. The incomplete closure of the space at the end of the paravaginal repair was the probable mechanism behind bowel incarceration. Later, the patient also presented a vaginal mesh exposure.

Mesh erosion is a well-known complication of using synthetic mesh. Erosions may be asymptomatic and inconsequential or they may present with severe infection or result in fistula. The analysis of the prevalence and identification of risk factors among the patients with vaginal mesh erosion is beyond the scope of this study, which intends to describe complications and its management. Nevertheless, an overview of Table I allows us to quickly notice that at least one of the literature reported risk factors for mesh erosion was present in every single patient who hereafter presented this complication (previous or concomitant total laparoscopic hysterectomy, through-and-through vaginal stitches, but it became quickly evident that a “new” type of complication was emerging. Three of the patient presented mesh exposure with no associated infection, probably an immune reaction to a foreign body, as described by one of the hypothesis of de Tayrac\textsuperscript{20}. According to this author, mesh erosion may result from a combination of bacterial infection and devascularization of the vaginal wall. De Tayrac describes 3 hypothesis to explain mesh-related complications, such as erosion, shrinkage or pain: an immune reaction to a foreign body; a prolonged inflammatory response (oxidative attack); a chronic infection. According to this paper, biomaterial implantation is followed immediately by a “race for the surface”, a contest between tissue cell integration and bacterial adhesion to that same surface that the bacteria win. The surface is occupied and is thus less available for tissue integration. In a study performed by Boulanger et al.\textsuperscript{24} bacterial contamination was found in all meshes removed for complications after surgical management of urinary incontinence or pelvic organ prolapse, and even if quantification was often low, its exact role is not yet clear. Vollebregt et al. also described as much as 83.6\% of colonized vaginal implanted meshes\textsuperscript{25}. This colonization justifies the systematic antibiotic prophylaxis, respected in most of the patients of this series. Nonetheless, this measure was not sufficient to prevent the cases of mesh exposure and/or infection.

Concerning infected meshes, two scenarios may occur: a localized infection, limited to the vaginal wall, or retroperitoneal extension of the infection up to the abdominal cavity, frequently causing a periprosthetic abscess. In both scenarios, the most reasonable one is that the mesh is infected at the time of its placement, as described above. In the first case, and if polypropy-
Porous, monofilament, soft polypropylene, type I mesh was used, a partial transvaginal removal of the mesh may be attempted. In the second scenario, complete removal of the infected mesh is required, especially if multifilament polyester mesh was used. Polyester mesh was used in 17/32 (53.1%) patients, especially in the beginning of our study. It is interesting to remark that in patients who presented longer intervals between surgery and diagnosis of periprosthetic abscesses, a polyester mesh had been used. Two occurred after haematoma and 1 after uterine perforation.

More frequent was the diagnosis of vaginal mesh exposure with no associated infection, which presented in most cases as vaginal bleeding or vaginal discharge (Table II). A conservative treatment for this condition was initially attempted in 14 patients (with topical oestrogens and antiseptic agents) with disappointing results, all of them needing subsequent surgical treatment. Transvaginal surgical excision and transvaginal endoscopic technique may be the next step\(^{26}\), with partial removal of the exposed mesh. When insufficient, an anterior insufficiency, with the creation of a bacterial biofilm, which allowed bacteria to remain quiescent during the initial period\(^{16}\). An intercurrent event, such as persistent exposition to the vaginal environment, may have contributed to bacterial multiplication and clinical infection. The fact that postoperative topical oestrogens were initially prescribed to only half of the patients may have contributed to the development of vaginal mesh exposure\(^{11}\). Five occurred in patients with polyester meshes, which supports the use of a macro-porous, monofilament, soft polypropylene, type I mesh\(^{22}\), and that conservative vaginal treatment should only be attempted in cases of polypropylene meshes, particularly in cases of small abscesses or asymptomatic patients. Nonetheless, our experience has shown that conservative treatment is rarely sufficient. An infected mesh should be completely removed, especially in case of deep infection and/or a polyester mesh. The particularity of the attachment to the promontory makes it mandatory to proceed with extreme care, because of the risk of spondylodiscitis.

Figure 1 presents the treatment for vaginal mesh exposures. It demonstrates the frequent need for several re-interventions, certainly affecting these patients quality of life. An initial less invasive treatment should be always balanced with the risk of re-intervention.

This is especially important when, as recently reinforced by a Chamsy and Lee publication\(^{28}\), “laparoscopic excision procedures of sacrocolpopexy mesh are typically challenging, even in the hands of experienced surgeons”.

Late exposure occurrence should not be neglected. Our series reports cases of a vaginal mesh exposure diagnosed 8 years after prolapse surgery, and Nygaard et al\(^{35}\) recently reinforced the importance of a long follow-up.

The only case of mesh retraction was on patient 8, who presented severe constipation, needing a surgical approach 9 months after the initial surgery. At the time, no evident mesh retraction was noticed but a marked fibrosis of the uterosacral ligaments was perceived. That under tension area was surgically released. Two years later, the patient was reoperated for a posterior mesh retraction evident at vaginal examination, persistent constipation and vaginal exposure.

Other relevant question addresses the treatment of recurrences after complete mesh removal. In our series, 2 patients needed subsequent surgeries for recurrent prolapse. On patient 21, a second laparoscopic sacrocolpopexy was performed, since a previous complete mesh removal had been necessary. This patient had a previous polyester mesh, and a polypropylene mesh was placed 10 months after the removal of the first one. Patient 33, on the other hand, had needed multiple pelvic adhesiolysis during laparoscopic sacrocolpopexy and presented an acute peritonitis on day 3, which would contribute to a more difficult insertion of a new mesh. A transvaginal approach was preferred for recurrent vaginal vault prolapse treatment in this case.

Limitations of this study are primarily related to its retrospective nature and small sample size. The fact that not all clinical data were accessible also limited our analysis. Even the technique evolved along the years. Follow-up intervals were not equivalent between all subjects, so it is possible that with time, new complications will be revealed. The retrospective nature of this study made it difficult to assess subjective symptoms of complications, so the clinical presentation is mostly based on signs.

In 2007, our center published a series on complete laparoscopic treatment of genital prolapse\(^{30}\), with no major complications, 5% of vaginal mesh erosion and 1% of mesh removal due to infectious complications. The authors intention was never to present a further statistical analysis of mesh-related complications, but rather a comprehensive review of their experience, by
considering all cases of patients who needed subsequent surgery for treating complications after sacroectomy.

In conclusion, when performing laparoscopic sacroectomy, surgeons should be aware of the risk factors for complications such as vaginal opening, post-hysterectomy status (complicating dissection and contributing to tissue devascularization), uterine perforation, adhesiolysis, through-and-through vaginal stitches and hypoestrogenism. Those situations should alert to the fact that need of re-intervention is more likely. In order to prevent this complication, some measures may be taken:

- Avoidance of aggressive coagulation of the vaginal wall during dissection on vault prolapse correction.
- Avoiding vaginal opening.
- Vaginal preparation with topic oestrogens.
- Systematic check for transfixing vaginal suture and stitches and hypoestrogenism. Those situations should be never neglected. The longer the follow-up, the later the diagnosis of complications can happen. It seems reasonable to assume that a patient with a mesh implanted is always at risk for mesh exposure.

After a mesh placement, patients are at risk for requiring multiple surgeries to resolve complications, surgeries that can be ineffective and expose the patient to new complications. Major complications can occur, as great vessels or urinary tract injury, and dissection can be challenging - during prolapse surgery or during complication-resolving surgery. Laparoscopic treatment of complications is feasible, but not always easy. The ability to resolve surgery and mesh-related complications should be considered a pre-requisite to perform prolapse surgery.

AUTHORS’ CONTRIBUTION

S Campos: project development, data collection, data analysis, manuscript writing
V Billone: data collection, data analysis
M Durão: manuscript writing
M Beguinot: data collection, data analysis
N Bourdel: responsible surgeon
B Rabischong: responsible surgeon
M Canis: responsible surgeon, manuscript writing
R Botchorishvili: project development, data collection, data analysis, responsible surgeon, manuscript writing

All patients gave their informed consent prior to their inclusion in the study.

The authors have full control of all primary data and they agree to allow the journal to review their data if requested.

CONFLICT OF INTEREST

The authors declare that they have no conflict of interest and received no funding.

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