Laparoscopic Sacral Uteropexy with Cravat Technique--Experience and Results

Murat Api¹, Semra Kayatas¹, Aysen Boza¹, Hakan Nazik², Hakan Aytan²

¹Zeynep Kamil Maternity and Children’s Training and Research Hospital, Department of Obstetrics and Gynecology, Istanbul and ²Adana Numune Training and Research Hospital, Department of Obstetrics and Gynecology, Adana, Turkey

ABSTRACT

Objective: The aim of the present study was to evaluate the safety and efficacy of a “Cravat” technique for the management of uterine prolapse in patients who want to preserve uterus, involving suspension of the uterus from the sacral promontory by using polypropylene mesh.

Materials and Methods: A prospective observational study between January 2011 and September 2013 was conducted. Prior to surgery, prolapse assessment was undertaken with Baden-Walker halfway system to grade the degree of prolapse at all sites. Patients with severe uterine prolapse (stage II-IV) who want to preserve uterus, were operated with Cravat technique. All patients were evaluated at 2 weeks and 6 weeks after surgery and followed for 6 months. Outcomes were evaluated objectively by vaginal examination using Baden-Walker halfway classification, and subjectively classifying patients as ‘very satisfied’, ‘satisfied’ and ‘not satisfied’ at the 6th month postoperatively.

Results: Sacral uteropexy was successfully performed by laparoscopy in 32/33 patients (one needed to be converted to laparotomy). Nine patients also had a concurrent procedure as colporaphy anterior, colporaphy posterior or transobturator tape. Postoperative recovery has been uneventful with subjective and objective cure rates were 96.9% and 93.9%, respectively at six month. One recurrence of total prolapse needed to be reoperated and two patients with sacrouteropexy still remained at stage 2 prolapse. There have been no cases of graft exposure, rejection or infection with a median follow-up of 23.9 months.

Conclusions: Laparoscopic sacral uteropexy with “Cravat technique” was found to be safe and simple procedure.

INTRODUCTION

Although pelvic organ prolapse (POP) is a common and disabling condition, the exact prevalence is difficult to ascertain due to different classification systems. Furthermore, many women do not seek medical attention despite of symptomatic POP. Population based studies report a 11 to 19 percent lifetime risk in women undergoing surgery for prolapse or incontinence (1,2).

A delay in childbearing to a later age leads many women to prefer operation with conserving uterus. Historically, the uterine preservation during prolapse surgery was first described as the Manchester procedure (3,4). The main aim of the pelvic floor reconstructive surgery should be to...
correct anatomical defects maintaining the uterus in normal anatomic position. Preservation of the uterus not only supports the pelvic floor, but also preserves fertility, improves sexual function and wellbeing. The increasing desire for uterine preservation provoked the development of new techniques with less morbidity and more patients’ satisfaction. Several conservative (pessary) or surgical treatment options were defined to preserve uterus in patients with POP (1,2). Surgical approach could be either vaginal or abdominal route.

The advancement of the minimal invasive surgery with respect to equipment and skills has provided the added option of endoscopic pelvic reconstructive surgery (1). Conventional laparoscopic and robot-assisted routes result in a shorter hospital stay, faster time to recovery, and lesser postoperative pain than laparotomy, with comparable short-term efficacy (5-8). The United Kingdom multicenter randomized equivalence trial found that after one year follow-up, there were no differences in anatomic or subjective pelvic floor outcomes compared to open and laparoscopic techniques; however, the blood loss, postoperative hemoglobin values, and hospital stay were better in the laparoscopic arm (9).

The objective of this study was to describe and evaluate the safety and efficacy of the surgical technique, laparoscopic sacrohysteropexy, a uterus preserving procedure for correction of uterine prolapse, involving suspension of the uterus with a mesh surrounding the isthmic portion of the uterus like cravat surrounding the neck.

**MATERIAL AND METHODS**

The investigation was designed as a prospective observational study from January 2011 through September 2013. The study protocol was conducted according to the revised Declaration of Helsinki and was approved by the Local Research and Ethics Committee of our hospital. All subjects provided written informed consent. Thirty-six women with symptomatic uterine prolapse (stage II-IV), who wanted to retain their uterus, underwent a laparoscopic sacro-uteropexy. Three of them were evaluated at 2 weeks and 6 weeks but could not be followed to 6 months so that 33 of them were included in the final analyses. Women with previous abnormal cervical cytological examination, abnormal uterine bleeding, significant uterine enlargement (e.g. uterine fibroids) and concomitant medical problems precluding general anaesthesia were excluded. The assessment of the prolapse was performed by the principal author (M.A) using the Baden–Walker halfway system (10). Urodynamic studies were performed in women with urinary incontinence complaint. Urodynamic results were evaluated in accordance with criteria established by the International Continence Society (11).

**Surgical Technique**

Surgery was performed under general anaesthesia with the woman in supine position and in low lithotomy position by the same senior author (M.A). After skin preparation, draping and catheterisation, Zinnanti uterine manipulator injector (ZUMI™, Cooper Surgical, Inc.) was used to obtain maximal anteversion of the uterus. A pneumoperitoneum was created and four laparoscopic ports were placed: 10mm umbilical port and three 5mm ports; two lateral and one left paraumbilical. The uterus was erected and the broad ligaments on each sides were opened from the posterior aspect above the levels of uterine vasculature, then the uterus bended posteriorly to open the peritoneal windows anteriorly on each side at the level of the cervico-uterine junction (Figure-1A). The vesico-uterine peritoneum was incised and the bladder was dissected distally by using blunt instrumentation (Figure-1B). The rectosigmoid colon was reflected to the left to expose the presacral area. The peritoneum over the sacral promontory was incised with monopolar scissors or harmonic scalpel and was bluntly dissected to the lateral sides. Ureters were securely observed to prevent damage (Figure-1C). A 25x1.5cm² prolene mesh (Ethicon, Inc. Johnson and Johnson, Somerville, NJ) was introduced through windows created in the broad ligaments (Figure-1D, Figure-1E) and was sutured anteriorly over the cervix with one polyglactin 910 suture (Ethicon, Inc., Johnson and Johnson, Somerville, NJ) (Figure-1F). Mesh was sutured to the sacral promontory under moderate tension to achieve adequate elevation of the uterus with at
least two, then 2/0 polypropylene suture and mesh arms were adjusted according to the need of uterine suspension by lifting the cervix at least 10cm above the level of introitus from the vaginal way (Figure-1G, Figure-1H). The rest of the mesh arms were trimmed over the sacral promontory after the fixation (Figure-1I). Some patients also underwent concomitant additional surgery as anterior and/or posterior colporrhaphy or transobturator tape (TOT).

All women were prospectively evaluated at 2 weeks and 6 weeks after surgery and followed up for at least 6 months. Operation time was defined as time from skin incision to final closure without including the concomitant surgery; duration of stay in hospital, complications, objective and subjective success rates were evaluated.

Postoperatively pelvic examination was performed to assess objective success rate defined as Baden-Walker grade 1 or 0 uterine prolapse. Subjective satisfaction of the patient was classified as ‘very satisfied’, ‘satisfied’and ‘not satisfied’.

Qualitative data are expressed in percentages (%) and quantitative data are expressed as the means ± standard deviation. Differences between the means in normally distributed variables were performed by using Student’s t-test.

Figure 1 - The surgical procedure of Cravat technique. A) Opening of left broad ligament. B) Incision of vesicouterine peritoneum. C) 25x1.5 cm² prolene mesh. D) Opening of peritoneum over the sacral promontorium. E) Mesh inserted to the windows in broad ligaments. F) Suturing of the mesh over the anterior cervix. G,H) Suturing of the mesh over the sacral promontory. I) Trimming of the rest of the mesh.

Bl = Broad ligament, Cx = Cervix, Ov = Over, P = Periton, M = Mesh, Sc = Sacrum, Sig.C = Sigmoid Colon, Ut = Uterus
chi-square test was done on categoric variables. Baden-Walker grades was analyzed by the Wilcoxon signed-rank test for pre and postoperative grades of related samples. Analyses was undertaken using Statistics Package for the Social Sciences 15 (SPSS Inc, Chicago, IL). A p value of < 0.05 was accepted as statistically significant.

**RESULTS**

A total of 33 patients underwent a sacro-uteropexy, a uterus sparing procedure for the management of uterovaginal prolapsus. The demographic characteristics of the patients are shown in Table-1 and operative data are given in Table-2. The mean age was 42.5 ± 7.4 years, mean parity was 3.2 ± 0.96 and mean body mass index was 25.3 ± 3.26 kg/m². Eight (24.2%) of them have a desire for children in the future. Preoperatively, all women had significant uterine descent of greater than or equal to grade 2, as measured using the Baden-Walker halfway system (Table-1). Thirteen women (39.3%) had grade 2 uterine prolapse, 18 women (54.5%) had grade 3 uterine prolapse, and the remaining 2 (6%) had grade 4 uterine prolapse. In addition, 3 (9%) of the women had grade 1-3 anterior vaginal wall prolapse, 2 (6%) had posterior vaginal wall prolapse and 2 (6%) of women had both of them.

Nine patients (27.2%) had additional surgery performed at the same time. Five patients (15.1%) with urodynamically confirmed urinary incontinence had TOT, 4 patients had colpography anterior, 6 of them had colpography posterior at the time of sacro-uteropexy.

Sacral uteropexy was performed successfully by laparoscopy in 32/33 patients; one of them was converted to laparotomy because of massive hemorrhage, but bleeding was successfully controlled and the same procedure was proceeded. Mean operation time was 46.39 ± 5.8 minute. The maximum and mean operation time decreased with increasing the surgeons’ experience on Cravat technique. The median hospital stay was 1.3 ± 0.4 day.

Major intraoperative vascular, genitourinary or gastrointestinal complications were not recorded. Only one patient developed middle sacral bleeding requiring laparotomy conversion.

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<th>Table 1 - The demographic characteristics of patients.</th>
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<td>Age (year ± SD)</td>
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<td>Parity (n ± SD)</td>
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<td>BMI (kg/m², ± SD)</td>
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<td>Sexual activity (n,%)</td>
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<td>SUI (n,%)</td>
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| Follow-up (month, mean ± SD) | 23.9 (± 4.7) |

SUI = stress urinary incontinence; BMI = body mass index; SD = standard deviation

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<th>Table 2 - The operation characteristics of Cravat technique.</th>
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<td>Operation time (minutes, mean, range)</td>
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<td>Duration of stay (day, mean ± SD)</td>
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<td>Concomitant procedure (n,%)</td>
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| CA | 0 |
| CP | 3 |
| CA+CP | 2 |
| TOT | 1 |
| CA+TOT | 2 |
| CP+TOT | 1 |

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<th>Complications (n)</th>
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<td>Transfusion</td>
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<td>Bowel injury</td>
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<td>Ureteric injury</td>
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CA = Colpography Anterior; CP = Colpography Posterior; TOT = Transobturator tape, SD = standard deviation
Postoperative recovery was uneventful. Among all participants, 96.9% (32/33) were either very satisfied 84.8% (28/33) or satisfied 12.1% (4/33). One (1.8%) patient reported not satisfied because of laparotomy conversion, blood transfusion and long hospital stay.

The mean postoperative Baden-Walker grades of the patients were statistically significantly lower compared to preoperative grades (p < 0.001). Subjective and objective cure rates were 96.3% and 93.9%, respectively at six months. There was no cases of graft exposure, rejection or infection with a median follow-up of 23.9 months and no case of recurrence of total prolapse.

**DISCUSSION**

The demand for uterine preservation during surgical management of uterovaginal prolapse is increasing and it is difficult to select the ideal uterus-sparing procedure for a given patient. Several alternative operations for prolapse repair with uterine preservation, using either a vaginal or an abdominal approach, have been proposed (1,2,12-20).

Vaginal operations are well defined in the literature; as in 2001, posterior intravaginal slingoplasty was first described with a reported mesh complications as infection and erosion (12). Other technique is the Manchester operation, that includes the vaginal shortening of the uterosacral and cardinal ligaments with cervical amputation which has a deleterious effect on fertility and also complicates with dyspareunia, dysmenorrhea, recurrent uterine prolapse and enterocele formation (3,4). Transvaginal sacrospinous fixation is the other vaginal operation that includes the fixation of cervix to the sacrospinous ligament by dissecting pararectal space down to the sacrospinous ligament. The proximity of the sacrospinous ligament to the sciatic nerve and pudendal vessels and nerves may cause significant buttock and leg pain and haemorrhage (13).

Abdominal operations performed for preserving the uterus have also been defined. The suspension of uterus from round ligaments (ventrosuspension) is associated with high recurrence rate. In a case series of nine women, recurrence of prolapse in eight women was reported within 3 months of surgery (2,14). This technique causes significant change of normal vaginal axis, hence it results in transmission of abdominal pressure to the cul-de-sac increasing the formation of enterocele (13,14).

Uterosacral plication is the other described technique in literature as placing three purse-string sutures from the uterosacral ligaments to the posterior cervix (15). Wu et al. reported a case series of seven women with no recurrence of prolapse at 9-17 months follow-up and also, Maher et al. reported an objective success rate of 79% in 43 women after a mean follow-up of 12 months (1,16). Lantzsch et al. described complications including massive haemorrhage, buttock pain and recurrent cystocele, and also Stepp and Paraíso described ureteral injury after uterosacral plication (17,18).

Suspension of prolapsed uterus from sacral promontorium by using polypropylene mesh was called sacrohysteropexy. This technique yields a satisfactory anatomic and functional results with normal vaginal axis (19). It was first described by Cutner et al. by passing Mersilene tape through uterosacral ligaments to resuspend the uterus to the sacral promontory bilaterally (20). Then this procedure was changed and non-absorbable mesh was started to be used by suturing the cervix posteriorly to the sacral promontorium with or without pelvic peritoneum closure via abdominal or laparoscopic way. The disadvantage of these techniques is the possible extrusion of mesh from cervix where it has been sutured before (21). Also, in sacrohysteropexy pulling the uterus from only one location seems to be less secure than wrapping the mesh around the uterus as described in our Cravat technique.

Price et al. used laparoscopic variation of the open procedure, involving suspension of the uterus from the sacral promontory by using bifurcated polypropylene mesh, which had been originally established by Leron and Stanton (2,19). In Price's technique, each broad ligament at the level of the cervico-uterine junction was opened, vesicouterine peritoneum was incised and bladder was dissected distally. Peritoneal relaxing incision was made down into the pelvis, laterally to the rectum,
medially to the right ureter, and also the peritoneum at the level of the insertion of uterosacral ligaments was mobilised to ease complete peritonisation of the mesh. Bifurcated polypropylene mesh was sutured anteriorly to the anterior cervical wall, posteriorly to the anterior longitudinal ligament over the sacral promontorium and then, complete peritonisation of mesh was performed (2). The difference from the bifurcated mesh described by Price et al. and our mesh was the 25x1.5 cm² tape. The advantage of our mesh was that it can be either available in every operation theatre or produced by slicing the 25x25 cm² square mesh into rectangular 25x1.5 cm² long pieces.

Price et al. connected the bifurcated tips of the mesh anterior to the uterine cervix and fixed these tips by 5-6 non-absorbable suture material (2). In our technique, the mesh was wrapped around the uterus anteriorly and the tips were connected on the sacral promontorium. For the anterior fixation we used one absorbable polyglactin 910 suture. We believe that mesh located between the vesica and cervix may potentially irritate the bladder, if mesh tips and 5-6 non-absorbable suture material are left in-situ in this area which may theoretically cause de-novo detrusor instability or bladder fistula afterwards. Absorbable suture material might not cause permanent irritation on the bladder but it is premature to say that absorbable material is superior to the non-absorbable one. Although we could not prove this fact by urodynamic examination, none of our subjects complained of overactive bladder symptoms in the postoperative follow-up period.

In Price’s technique, all of the operations were performed with a standardized mesh length placed under standardized tension. On the other hand, we adjusted the mesh length described as follows: we pulled the mesh between the prolapsed uterus and sacral promontory with a moderate tension in order to suspend the cervix at least above the interspinous level of the bony pelvis. Uterus was pushed with a ZUMI vaginally. The sacral promontory mesh was fixed to the anterior longitudinal ligament by non-absorbable suture and then the extra mesh was incised and taken out. We thought that every patient needs a different length of mesh because of the variation in length of the prolapsed portion, so putting the uterus in normal anatomic position by raising the cervix and adjusting the length of the mesh might be the key components of the our technique which increased the success rate.

In our technique, we did not make a peritoneal relaxing incision down into the pelvis, we only incised the peritoneum over the sacral promontory and avoided the bruise of mesh which might contribute to complications (22,23). We believe that suturing the peritoneum may cause direct/indirect ureteric injury (kinking) or inadvertent bowel injury (22). In addition, the insertion of sutures into the peritoneum may cause bleeding and the formation of haematoma (24). As a result of non-peritonisation of mesh in abdominal vault suspension operations done by Elneil et al. none of the patients developed problems with adhesions, only 3 out of 128 patients with a rate of 2.3% had vaginal mesh erosion (22).

Cravat technique had shorter mean operation time than other techniques (1,2). Surgical steps that likely contributed most to improve the operation time were eliminating the need to open the peritoneum from the sacral promontory to the cul-de-sac, non-peritonisation of mesh and since mesh was wrapped around the uterus, there was only one area that we should secure to mesh instead of two or more. The number and localisation of the sutures may also decrease the operation time; one suture rather than 6 and using anterior cervical wall rather than posterior one may fasten the operation.

Although this follow up period cannot provide conclusive evidence we are most encouraged that in long term follow-up more successful results might be expected from hysteropexy with Cravat technique with lower complication rate and shorter operation time.

**CONCLUSIONS**

Our results support the benefits of laparoscopic uterine preservation with “Cravat technique” in patients with POP. In a young patient with the desire of maintaining uterus, laparoscopic hysteropexy has the advantages of maintaining childbearing capacity, better anatomical restoration of the pelvic structures, improving sexual function, lesser intraoperative adhesions and shorter hospital stay. Although we had no control group, our laparoscopic sacral uteropexy technique was found to be relatively safe and simple. Nevertheless, the need
for larger randomized controlled trial evaluating its efficacy, safety and long term outcome measures still remains.

CONFLICT OF INTEREST

None declared.

REFERENCES


Correspondence address:
Aysen Boza, MD
Zeynep Kamil Maternity and Children’s Training and Research Hospital,
Department of Obstetrics and Gynecology,
Buranhatettin Ustunel Street, No: 2/3,
Uskudar, Istanbul, 34668, Turkey
Fax: + 90 216 391-0690
E-mail: arcke83@gmail.com