Anatomical and clinical outcomes of vaginally assisted laparoscopic lateral suspension in comparison with laparoscopic lateral suspension

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Background: In this retrospective study, we aimed to describe the surgical procedure for vaginally assisted laparoscopic lateral suspension (VALLS) and to compare its anatomical and clinical outcomes with laparoscopic lateral suspension (LLS). Methods: The surgical outcomes of 26 women with advanced-stage pelvic organ prolapse (POP) undergoing VALLS and 35 women with advanced-stage POP undergoing LLS were retrospectively analysed and compared. The surgical outcomes were documented according to the International Urogynecological Association guidelines. Complications were evaluated according to the Clavien-Dindo classification and classified using the joint project of the International Continence Society and the International Urogynecological Association Prosthesis/Graft Complication Classification System. Results: The results showed significant improvement in all POP Quantification measurements in both the VALLS and LLS groups, with overall objective cure rates of 88.4% and 80%, respectively (96.1% and 91.4%, respectively, for the apical compartment, 96.1% and 85.7%, respectively, for the anterior compartment). The median operation times for VALLS and LLS were 77 [66–90] minutes and 99 [82–125] minutes, respectively (p = 0.001). A significant improvement in POP symptoms was observed in both groups. Occult stress urinary incontinence (SUI) was detected in two (7.6%) VALLS patients, and de novo SUI developed in four (15.3%) VALLS patients post-operatively. Anterior compartment defects were detected in one VALLS and five LLS patients. Mesh erosion was found in one patient in each group. Discussion: VALLS appears to be an effective and reliable surgical method for patients with advanced-stage POP and can offer advantages in terms of operation time and POP recurrence rates.

Keywords
Laparoscopic lateral suspension; Minimally invasive surgery; Pelvic organ prolapse; POP surgery; Vaginally assisted laparoscopic lateral suspension

1. Introduction

Pelvic organ prolapse (POP) is the bulging of the uterus, anterior or posterior vaginal wall, or vaginal vault to a lower place or level [1]. The prevalence of POP in postmenopausal women is 3–6% according to symptoms and it may rise to 50% when based upon vaginal examination [2, 3]. A woman’s lifetime operation risk of surgery for POP is 12–19%. Moreover, 10–30% of women with POP require reoperation [4].

Various vaginal and abdominal surgical management methods using native tissue or mesh have been developed to treat POP. Following the notification of the U.S. Food and Drug Administration in 2011 and 2019 regarding POP repair with a vaginal mesh, trans-abdominal mesh operations have gained popularity [5, 6]. Sacrocolpopexy (SCP) is the mostly preferred technique for treating POP laparoscopically. However, SCP involves a long operative time and a steep learning curve. The procedure includes sacral area dissection, which can be extremely difficult, particularly in overweight women, and may result in major neurological, ureteral, or vascular injuries [7, 8].

A dissection at the level of the promontory/sacral area is not necessary in the laparoscopic lateral suspension (LLS) procedure established by Dubuisson et al. [9]. Moreover, the risk of major complications appears to be lower than that of SCP. LLS can be performed using a T-shaped synthetic mesh graft and may or may not be a uterus-sparing procedure [10]. Studies on LLS have documented more than 90% objective success rate in the anterior and apical compartments after one year [11, 12]. Thus, LLS may be an alternative surgical option to SCP for the management of apical compartment defects in women with POP [13, 14]. The most critical step of the LLS procedure is the mesh placement in the vesicovaginal space. The true vesicovaginal space should be reached by performing dissection of a full vaginal wall layer. The mesh should be placed flat without folding or overstretching, and fixation should be performed using the appropriate suture technique and material which requires advanced laparoscopic surgical skills [15].

The aim of this study was to describe the surgical steps of a vaginally assisted laparoscopic lateral suspension (VALLS) procedure which enables quick and effortless suture for the management of women with severe POP and to evaluate its effectiveness and reliability. To that end, we compared the preoperative characteristics and postoperative outcomes of patients undergoing VALLS and LSS.

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2. Methods

2.1 Ethical approval

This was a retrospective study conducted in the Department of Obstetrics and Gynecology of the Faculty of Medicine of Muğla Ştki Koçman University, Turkey. The study was approved by the Clinical Investigations’ Ethics Committee of Muğla University, Turkey (no. 4/I; 17 February 2021). The study was conducted in accordance with the provisions of the Declaration of Helsinki. Written informed consent was obtained from all patients before undergoing surgery.

2.2 Study design

The study included a study group of patients with stage 3–4 POP undergoing VALLS and a control group of patients with stage 3–4 POP undergoing LLS between February 2013 and March 2020. Patients with previous hysterectomy, SCP, or POP surgery with a vaginal mesh, patients older than 65 years, patients with diabetes, and patients with smoking habits were excluded. Transvaginal ultrasonography was performed in all patients before the operation. In addition, cervicovaginal smear scanning was performed in all patients. Urodynamic evaluation was done for the patients with pre- and post-operative urinary incontinence symptoms. According to the inclusion and exclusion criteria, 26 patients were included in the VALLS group, and 35 patients were included in the LLS group. The patients’ demographic and clinical characteristics, prolapse-related symptoms, and operational information were obtained from the hospital’s database and patient files. The physical examination results, reoperation and erosion rates, lower urinary tract symptoms (LUTS), and complications were retrieved.

All surgeries were performed by a single surgeon (EA). The patients were re-evaluated after a minimum of 12 months post-operatively. The pre- and post-operative examinations included Pelvic Organ Prolapse Quantification (POP-Q) measurements and staging. The degree of prolapse in points Ba, Bp, and C was determined according to the simplified POP-Q [1]. Urogynaecological examinations were performed with the patients in a semi-recumbent lithotomy position. The POP grade was evaluated by performing the Valsalva manoeuvre and/or coughing. The surgical outcomes were recorded according to the International Urogynaecological Association (IUGA) guidelines [16]. Satisfactory anatomic objective cure was defined as POP-Q sites Ba, C and Bp being less than -1 during follow-up. Complications were evaluated according to the Clavien-Dindo classification and classified using the joint project of the International Continence Society and the International Urogynaecological Association Prosthesis/Graft Complication Classification System [17, 18].

2.3 Operation technique

2.3.1 Operation technique for VALLS

A T-shaped mesh with a rectangular part of $4 \times 6$ cm in the middle and two long arms of $2 \times 18$ cm was cut from a $30 \times 30$ cm polypropylene macropore mesh (Parietene™, Covidien™, Trevoux, France) and prepared preoperatively. Surgery was performed under general anaesthesia in a lithotomy position. We used the four-port laparoscopic technique, a central 10-mm umbilical trocar for the 0-degree camera, two lateral ports, and a fourth ipsilateral port. A Foley catheter was inserted into the bladder, and a RUMI®II System (CooperSurgical, Trumbull, CT, USA) uterine retractor with a balloon tip was inserted into the uterine cavity for uterine manipulation and adequate exposure of the anterior and posterior vaginal fornices. The vesicovaginal space between the bladder and the anterior vaginal wall in the fascia plane was dissected until the lower third border of the vagina. The T-shaped mesh was placed in the abdominal cavity through a no.10 umbilical trocar. Combined with laparoscopic dissection, an approximately 4 cm circular incision was made vaginally 2 cm distal to the external cervical os to enter the vesicovaginal space and to reach the abdominal cavity. The middle part of the T-shaped mesh was placed in the vesicovaginal space and separately sutured vaginally with no. 2-0 Prolene® (monofilament polypropylene suture; Ethicon, Somerville, NJ, USA) suture to the anterior wall of the vagina, vesicovaginal fascia, pubocervical fascia, cervix, and isthmus of the uterus in such a way that there was no shrinkage in the mesh. The number of stitches was 10–14 sutures in all cases (Fig. 1).

![Fig. 1. The middle part of the T-shaped mesh with the dimensions of $4 \times 6$ cm is sutured to the vesicovaginal fascia with polypropylene suture material without wrinkling the mesh.](image-url)
Table 1. Preoperative demographic features and findings.

| Demographic features                           | VALLS n = 26 | LLS n = 35 | p-value  |
|------------------------------------------------|--------------|------------|----------|
| Age (year)                                      | Median [Min–Max] | Median [Min–Max] | 0.574*   |
| BMI (kg/m²)                                     | Mean ± SD    | Mean ± SD  | 0.311**  |
| Normal weight: 18.5–24.9                        | 14 (53.8)    | 19 (54.3)  |          |
| Overweight: 25–29.9                             | 8 (30.8)     | 6 (17.1)   |          |
| Obese: ≥30                                      | 4 (15.4)     | 10 (28.6)  |          |
| Parity, (n)                                     | 3 [1–9]      | 3 [2–7]    | 0.511*** |
| Number of vaginal deliveries                    | 3 [1–8]      | 3 [2–7]    | 0.510*** |
| Menopausal condition, n (%)                     |              |            | 0.563**  |
| Premenopausal                                   | 8 (30.8)     | 8 (22.9)   |          |
| Postmenopausal                                  | 18 (69.2)    | 27 (77.1)  |          |
| Previous POP surgery, n                         |              |            | 0.356**  |
| Colporraphy anterior                            | 4            | 3          |          |
| Colporraphy posterior                           | 2            | 3          |          |
| Manchester Fotergill                            | 1            | 0          |          |
| Previous stress urinary incontinence operations, n |            |            | 0.731****|
| Transobturator sub-urethral sling              | 1            | 0          |          |
| Tension free retropubic sling                   | 2            | 2          |          |
| Kelly-Kennedy                                   | 2            | 3          |          |
| POP-Q                                           |              |            | 0.570**  |
| Stage 3, n (%)                                  | 6 (23.1)     | 11 (31.4)  |          |
| Stage 4, n (%)                                  | 20 (76.9)    | 24 (68.6)  |          |

Values expressed as the mean ± standard deviation, Median [Min–Max] or number (%). SD, standard deviation; BMI, body mass index; POP-Q, pelvic organ prolapse quantification. *Independent Samples T Test. **ChiSquare Test. ***MannWhitney U Test. ****Fisher’s exact test.

The vaginal vault was closed vaginally with an absorbable no.0 Vicryl Rapid™ (polyglactin 910; Ethicon, Somerville, NJ, USA) suture. A 3-mm skin incision was made on both sides 2 cm above the iliac crest and 4 cm posterior to the anterior superior iliac spine. The forceps were initially advanced caudally in the retroperitoneal area, taking care to avoid the external iliac artery and vein, then advanced under the ligamentum rotundum. The distal tip of one long arm of the T shaped mesh was grasped and pulled out through the cutaneous incision. The lateral suspension procedure was performed in both sides. The lateral arms of the mesh were not fixed to the abdominal fascia, according to “tension free” repair principle. Peritonisation was also performed over the mesh inserted into the vesicovaginal space using a no. 2-0 absorbable Vicryl Rapid™ suture. The mesh was then cut at the skin level prior to the closure of the skin incision.

2.3.2 Operation technique for LLS

The only difference of LLS from VALLS was the mesh fixation to the vesicovaginal space. At this step no. 2-0 Prolene® was sutured intracorporeally with laparoscopy. As in the VALLS procedure, lateral arms of the mesh not sutured abdominal fascia in LLS procedure.

2.4 Statistical analysis

Statistical analysis was performed using IBM SPSS Statistics version 20.0 (IBM, Armonk, NY, USA) for Windows. The Shapiro-Wilk test was used to evaluate data normality. The data were expressed as the median and range for continuous variables, and binary variables were reported as numbers and percentages. For inter-group comparisons, the independent samples T-test was used for parameters with a normal distribution and the Mann Whitney U Test was used for parameters with non-normal distribution. A Pearson’s chi-square test or Fisher’s exact test was used for comparison of categorical data. A p value of ≤0.05 was considered statistically significant.

3. Results

The demographic and preoperative clinical characteristics of the patients in the VALLS and LLS groups are summarised in Table 1. The preoperative characteristics of the two groups were similar.

The post-operative anatomical outcomes of the patients in the two groups are summarised in Table 2.

The median post-operative follow-up duration was 22 (13–30) months in the VALLS group and 32 (14–69) months in the LLS group (p = 0.09). A significant improvement in POP-Q scores was observed in all compartments, with overall objective cure rates of 88.4% in the VALLS group (96.1% in the apical compartment and 96.1% in the anterior compartment) and 80% in the LLS group (91.4% in the apical
Table 2. Anatomic outcomes in patients undergoing uterus preserving VALLS and LLS.

| POP-Q              | VALLS n = 26 | LLS n = 35 | p-value |
|--------------------|--------------|------------|---------|
| Preop Point Ba     | Median [25%–75%] | Median [25%–75%] |
|                    | 5 [3–6]      | 4 [3–4]    | 0.110   |
| Preop Point C      | 6 [4–6]      | 6 [5–6]    | 0.982   |
| Preop Point Bp     | 3 [2–3]      | 2 [2–3]    | 0.159   |
| Postop Point Ba    | –3 [–3]–(–3) | –2 [–3]–(–2) | 0.001* |
| Postop Point C     | –6 [–7]–(–5) | –5 [–6]–(–3) | 0.087   |
| Postop Point Bp    | –2 [–4]–(–1) | –2 [–3]–(–1) | 0.465   |

*Significant at ≤0.05 level; Mann Whitney U Test.

Table 3. Comparison of preoperative and postoperative lower urinary tract symptoms of patients who underwent VALLS and LLS.

| Variables                      | Preop   | Postop  | p-value |
|--------------------------------|---------|---------|---------|
| Lower urinary tract symptoms   | VALLS n (%) | LLS n (%) | VALLS n (%) | LLS n (%) | p-value |
| Vaginal bulge                  | 26 (100) | 35 (100) | 0.955 | 1 (3.8) | 5 (14.3) | 0.179 |
| Urinary urgency                | 16 (61.5) | 24 (68.6) | 0.571 | 2 (7.7) | 10 (28.6) | 0.044* |
| Incomplete voiding             | 19 (73.1) | 27 (77.1) | 0.718 | 3 (11.5) | 7 (20) | 0.381 |
| Urinary frequency              | 22 (84.6) | 30 (85.7) | 0.906 | 18 (69.2) | 27 (77.1) | 0.491 |
| SUI                            | 2 (7.7) | 3 (8.6) | 0.902 | 3 (11.5) | 4 (11.4) | 0.989 |
| Constipation                   | 9 (34.6) | 11 (31.4) | 0.795 | 2 (7.7) | 2 (5.7) | 0.761 |
| Fecal Incontinence             | 4 (15.4) | 4 (11.4) | 0.653 | 2 (7.7) | 2 (5.7) | 0.760 |
| Sexual activity                | 7 (26.9) | 13 (37.1) | 0.404 | 16 (61.5) | 24 (68.6) | 0.571 |
| Dyspareunia                    | 6 (23.1) | 7 (20) | 0.733 | 3 (11.5) | 3 (8.6) | 0.703 |
| Pelvic pain                    | 14 (53.8) | 19 (54.2) | 0.973 | 10 (38.4) | 13 (42.8) | 0.917 |

*Significant at ≤0.05 level; Chi Square Test. SUI, stress urinary incontinence.

Table 4. Postoperative complications.

| Complications                      | VALLS n = 26 | LLS n = 35 | p-value |
|-----------------------------------|--------------|------------|---------|
| Recurrence, n (%)                 | 3 (11.5)     | 7 (20)     | 0.494* |
| Apical compartment                | 1 (3.8)      | 3 (8.5)    |
| Anterior compartment              | 1 (3.8)      | 5 (14.2)   |
| Posterior compartment             | 1 (3.8)      | 2 (5.6)    |
| Repeat surgery for recurrence, n (%) | 1 (3.8) | 7 (17.1) | 0.122* |
| Apical compartment                | 0            | 2 (5.6)    |
| Anterior compartment              | 1 (3.8)      | 4 (11.2)   |
| Posterior compartment             | 0            | 2 (5.6)    |
| Vaginal mesh erosion, n (%)       | 1 (3.8)      | 1 (2.8)    | 0.675* |
| Repeat surgery (Retropubic sling) for SUI, n (%) | 4 (15.3) | 5 (14.2) | 0.591* |

*Fisher’s exact test. SUI, stress urinary incontinence.

In our study, we found the median operative time as 77 [66–90] minutes for VALLS, and 99 [82–125] minutes for LLS (p < 0.001).

The pre- and post-operative LUTS of the patients in the two groups are displayed in Table 3. In both groups, the most common symptoms were palpable vaginal swelling and consequent walking difficulty. Considerable improvement in vaginal bulging, urgency, incomplete voiding, urinary frequency, and difficulty in defecation was observed post-operatively. In the VALLS group, occult stress urinary incontinence (SUI) was observed in two (7.6%) women, and de novo SUI developed in 4 (15.3%) women post-operatively. In the VALLS group, the rate of sexually active women increased from 7 (26.9%) to 16 (61.5%).

The post-operative complications in the two groups are shown in (Table 4). There were no major complications in either group (Clavien-Dindo grade 1). In one patient undergoing VALLS and two patients undergoing LLS, bladder perforation occurred during the dissection of the vesicovaginal compartment and 85.7% in the anterior compartment).
space, and suturing was performed intraoperatively. No pa-
tient required a blood transfusion or underwent laparotomy.

In terms of recurrence, an anterior compartment defect was
observed in one patient in the VALLS group, and ante-
rior colporrhaphy was performed. The patient had a BMI of
27 kg/m$^2$. In the LLS group, anterior compartment defects
were observed in five patients, one of whom was overweight
and four were obese. Concurrent apical and anterior com-
partment defects were observed in two patients in the LLS
group.

Grade 2 (>1 cm) anterior vaginal wall mesh exposure
occurred in one patient in each group, with similar mesh
erosion characteristics. The exposed part was detected be-
tween two and six months post-operatively and categorised
as 3BT3S1 according to the Prosthesis/Graft Complication
Classification System. In these patients, the exposed mesh
was partially excised, and the vaginal mucosa was primarily
repaired.

4. Discussion

POP is a very common condition, especially in obese
women. A restoration of the disrupted anatomical relation-
ships to normal anatomy contributes to a regression of LUTS
and may exert positive functional and psychological effects
[19]. Although several vaginal, abdominal, and endoscopic
surgical techniques have been developed for the treatment
of advanced POP, the optimal approach has yet to be deter-
dined. This study aimed to describe the surgical steps of a
VALLS procedure for the treatment of women with POP
stage ≥3 and evaluate its effectiveness by comparing it with
LLS. Our results showed that VALLS provided better rates of
anatomical support, symptomatic relief, patient satisfaction,
intraoperative and post-operative complications, and vaginal
erosion. The apical compartment cure rate in patients un-
dergoing VALLS was 96.1%, and the anterior compartment
cure rate was 96.1% during a follow-up period of, on aver-
age, 22 months. These results are consistent with Russo et al.
[20], who reported success rates of 94.1% in the apical com-
partment and 88.3% in the anterior compartment in robotic-
assisted LLS. Ganatra et al. [21] who reviewed 11 retrospec-
tive studies with a mean follow-up of 24.6 months, found
overall objective anatomical and subjective laparoscopic SCP
success rates of 92% and 94.4%, respectively.

Laparoscopic suturing is not an easy task, especially in a
deep and narrow area. Proper laparoscopic suturing of the
mesh may account for a large part of the operation time.
This procedure is particularly difficult and time-consuming
in obese patients. Moreover, in advanced POP cases, sutur-
ing a large and long mesh without curling in the vesicovaginal
area, including the pelvic fascia, may be possible only vagi-
nally [22].

In our study, the median operation time was significantly
shorter in the VALLS group than the LLS group. In a previ-
ous study with uterine sparing LLS, the mean operative time
was reported as 189.26 (± 44.62) minutes [23]. In another
study with LLS, mean operative time was reported as 120
(63–280) minutes [11]. In these studies, concomitant surgi-
ceries were performed in addition to LLS, so their operation
time was longer than ours. Moreover, VALLS may have
shortened the total operative time by reducing the suturation
time of the mesh to the anterior vaginal wall using laparo-
scopical access.

In this study, a significant amelioration of vaginal bulging,
urinary urgency, incomplete voiding, and urinary frequency
was observed in patients undergoing VALLS. An occult SUI
rate of 20% has been reported in patients with a POP diag-
nosis, which is even higher in patients with advanced-stage
POP [24]. Veit-Rubin et al. [14] reported a SUI rate of 6.6%
after LLS. In this current study, de novo SUI was observed
in four (15.3%) patients who underwent VALLS, and in five
(14.2%) patients who underwent LLS group.

In the past, hysterectomy was often performed in POP pa-
tients [25]. Later, surgical repair without hysterectomy was
preferred due to the belief that hysterectomy reduced libido
and impaired sexual function. Uterus-sparing LLS is asso-
ciated with higher satisfaction rates, better short-term sub-
jective outcomes, and lower post-operative constipation and
de novo SUI rates [23]. Avoiding hysterectomy may also
offer anatomical and functional benefits, such as maintain-
ing strong rectovaginal and vesicovaginal fascia and cervix
support [26]. It can also reduce morbidity and complication
rates. In relatively young and sexually active women, avoid-
ing hysterectomy and disruption of the vaginal axis may be
associated with better sexual function post-operatively [27].

Major causes of mesh-related complications, such as
erosion/extrusion, include aggressive dissection, intensive
catheterisation, poor suture techniques that disrupt the nutri-
tion of the mesh site, improper mesh placement, and shrink-
age of the mesh at the suture stage [28]. Traditional SCP and
LLS use non-absorbable or delayed absorbable and monofil-
ament sutures for mesh fixation to prevent detachment from
the vagina and reduce the risk of erosion [29]. The inclu-
sion of the pelvic fascia in the mesh suture, which is quite
easy in VALLS, can further reduce the risk of erosion. In pa-
tients in whom the mesh is stitched vaginally in laparoscopic
POP surgery, possible contamination of the mesh with the
vaginal flora may increase the risk of mesh erosion. In con-
trast, studies evaluating cases of vaginally assisted SCP have
found no increased risk of contamination caused by vaginal
suturing of the mesh [20, 22]. Follow-up period may influ-
ence the results of the mesh erosion rates [30]. Previous stud-
ies have reported variable time intervals to exposure, ranging
from 6 weeks to 8 years [14, 30, 31]. In a study related mesh
erosion after abdominal sacrocolpopexy median time to ero-
sion detection was 16.8 months [30], another study report
that mesh erosion median time to diagnosis after laparoscopic
lateral suspension was 20 months [32]. In the current study
the median post-operative follow-up duration was 22 (13–
30) months in the VALLS group and 32 (14–69) months in
the LLS group. Although there was no statistically significant
difference between the two groups, the follow-up period of the patients who underwent LLS was longer.

In our study, mesh erosion in the anterior compartment was observed in only one (3.8%) of the 26 patients undergoing VALLS. This is slightly lower than previously reported rates. A systematic review found a 4% rate after abdominal surgery using mesh [30]. Another study reported a 4.3% rate after 417 LLS procedures [14].

One reason for recurrence after LLS is the separation of the mesh from the tissue due to a failure to suture tightly [33]. We encountered recurrence of anterior compartment defect in five patients in the LLS group and one patient in the VALLS group. These patients were overweight or obese. Only one (3.8%) patient in the VALLS group required reoperation after more than 12 months, which is consistent with previously reported reoperation rates (3.4–11%) [34].

LLS is not preferred if significant apical and posterior defects are present together [35]. In LLS procedures using T-shaped synthetic mesh grafts, the lateral mesh arms do not close the Douglas space [14]. This may result in the progression of the posterior vaginal compartment defect [36]. However, the suspension axis of a lateral mesh does not lead to enterocele or space of Douglas hernia in patients treated for apical and anterior POP [11]. Reduced reoperation and recurrence risks have been reported in patients undergoing apical compartment defect repair with simultaneous posterior compartment repair [37]. We repaired posterior compartment defects using posterior colporrhaphy. We had one case of rectocele recurrence in VALLS group.

Limitations of our study are its retrospective, single-centre design, small sample size, and short follow-up period. On the other hand, a strength of this study is that all operations were standardized and performed by the same surgeon. Our study contributes to the literature by describing a new surgical method. Prospective studies including large numbers of patients and longer post-operative follow-up periods are needed to further evaluate this approach and compare it with other pelvic reconstructive surgery techniques.

5. Conclusions

VALLS appears to be a reliable and efficient modified method for women with POP scheduled for LLS. VALLS may reduce the POP recurrence rates, especially in obese patients. It also offers an advantage in terms of operation time and can be helpful to surgeons with modest laparoscopic suturing abilities. Moreover, it provides the opportunity for subcervical fascia plication and posterior colporrhaphy when necessary. Further studies are warranted to evaluate this technique.

Author contributions

EA conceptualized and designed the study, drafted the initial manuscript, performed surgical operations, and reviewed and revised the manuscript. BS and AAS designed the data collection instruments, collected data, carried out the initial analyses, and reviewed and revised the manuscript. All authors read and approved the final manuscript.

Ethics approval and consent to participate

This study was conducted in the Department of Obstetrics and Gynecology of Muğla Sıtkı Koçman University Faculty of Medicine in retrospective design. The study was approved by the Clinical Investigations’ Ethics Committee of Muğla University/Turkey (Approval date: February 17th, 2021 and No: 4/I). The written informed consent was obtained from all participants in accordance with Helsinki Declaration.

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Conflict of interest

The authors declare no conflict of interest.

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