Prevention and management of postoperative urinary retention after urogynecologic surgery

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Abstract: Postoperative urinary retention (POUR) is a frequent consequence of gynecologic surgery, especially with surgical correction of urinary incontinence and pelvic organ prolapse. Estimates of retention rates after pelvic surgery range from 2.5%–43%. While there is no standard definition for POUR, it is characterized by impaired bladder emptying, with an elevation in the volume of retained urine. The key to management of POUR is early identification. All patients undergoing pelvic surgery, especially for the correction of incontinence or prolapse, should have an assessment of voiding function prior to discharge. There are several ways to assess voiding function – the gold standard is by measuring a postvoid residual. Management of POUR is fairly straightforward. The goal is to decompress the bladder to avoid long-term damage to bladder integrity and function. The decision regarding when to discontinue catheter-assisted bladder drainage in the postoperative period can be assessed in an ongoing fashion by measurement of postvoid residual. The rate of prolonged POUR beyond 4 weeks is low, and therefore most retention can be expected to resolve spontaneously within 4–6 weeks. When POUR does not resolve spontaneously, more active management may be required. Techniques include urethral dilation, sling stretching, sling incision, partial sling resection, and urethropily. While some risk of POUR is inevitable, there are risk factors that are modifiable. Patients that are at higher risk – either due to the procedures being performed or their clinical risk factors – should be counseled regarding the risks and management options for POUR prior to their surgery. Although POUR is a serious condition that can have serious consequences if left untreated, it is easily diagnosed and typically self-resolves. Clinician awareness of the condition and vigilance in its diagnosis are the key factors to successful care for patients undergoing surgical repair.

Keywords: incontinence, prolapse, postoperative, urinary retention

Introduction
Postoperative urinary retention (POUR) is a frequent consequence of gynecologic surgery, especially with surgical correction of urinary incontinence and pelvic organ prolapse. 1–2 Estimates of retention rates after incontinence and prolapse surgery range from 2.5%–24% and are as high as 43% after tension-free transvaginal mesh sling placement. 2,3 When managed properly, POUR does not carry a high risk of morbidity. However, failure to identify this condition can lead to serious clinical sequelae, as prolonged bladder distention can lead to urinary tract infection (UTI), detrusor dysfunction, and even damage to the surgical repair. 4 A significant challenge in the diagnosis of POUR comes from the fact that there is no universal definition of urinary retention, either postoperative or otherwise. When using the broader definition of “voiding dysfunction” to characterize postoperative bladder function, estimates are even higher,
with a range of 39%-84%. These higher rates include any transient voiding dysfunction that is documented in the postoperative period, as early as in the recovery room. Prolonged retention – that lasting 4 weeks or longer after surgery – is much rarer, with rates of 2%-4% for a transvaginal mesh sling.

Traditional pubovaginal fascial slings and the Burch urethropexy both carry higher rates of prolonged retention (4%-10% and 4%-22%, respectively) but are less commonly performed today. Transurethral injection of a bulking agent such as collagen, carbon beads, or hydroxylapatite has a 15% rate of transient retention and a very low risk of prolonged retention. For surgical repair of pelvic organ prolapse, the rate of any postoperative retention has been quoted at 29%. In many cases, women undergo a combination of procedures to treat both incontinence and prolapse, which likely increases their probability of experiencing postoperative urinary retention.

**Definition**

Normal bladder function involves both successful storage and emptying. Either of these phases can be affected by pelvic surgery and can lead to postoperative voiding dysfunction. Urodynamic studies show that normal bladder sensation begins at around 150 mL, with first urge occurring at 250 mL, strong urge at around 400 mL, and capacity ranging from 400–700 mL. Micturition occurs when the bladder wall is distended, which activates stretch receptors, which in turn send afferent signals up the spinal cord to the cerebral cortex. If the cortex determines it is an appropriate time to allow micturition, it sends an efferent signal to the pontine micturition center, which in turn activates parasympathetic motor neurons that cause detrusor contraction, and it inhibits somatic effenter motor neurons, which leads to urethral sphincter relaxation. Postsurgical changes that lead to edema, inflammation, damage to peripheral nerve endings, and pain can all affect bladder sensation and the micturition pathway, leading to POUR.

**Risk factors**

In addition to the physiologic factors described above, known risk factors for POUR include the type of surgery performed, the type of anesthesia used, use of postoperative opioid pain medication, patient age and sex, and baseline bladder dysfunction. (Table 1) Any surgery has the risk of POUR, but we find that this risk is increased with gynecologic surgery and anorectal surgery, likely due to heightened sensitivity in these regions that can lead to increased pain, as well as to the disruption of small peripheral pelvic nerve branches that play a role in bladder sensation. POUR after anorectal surgery has been reported to occur in up to 21.9% of hemorrhoidectomy cases, with an overall rate of 16.7% for all anorectal cases. This corroborates with the idea that both pain and peripheral neuropathy play a role in POUR. Pain medication can also exacerbate bladder dysfunction and increases the rate of POUR. Increased use of opioid pain medication has demonstrated an almost 1.5-times-greater risk of developing postoperative retention (odds ratio [OR] =1.3). This may be a combination of greater pain requiring greater opioid use, as well as the effects of the opioid itself. Furthermore, the same study demonstrated that immediate postoperative prophylactic analgesia with indomethacin suppository decreased the rate of POUR significantly (7.9% versus 25.0%, P=0.0005). Another study demonstrated that vaginal bupivacaine use at the time of sling placement increased the rate of POUR, with a higher PVR in the bupivacaine group (225 mL versus 140 mL, P=0.043). Interestingly, in this study bupivacaine did not reduce pain scores or pain medication use. Another study comparing pudendal block with spinal block found a significantly higher rate of POUR

**Table 1 Risk Factors for POUR**

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<thead>
<tr>
<th>Demographic risk factors</th>
<th>Surgical risk factors</th>
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<tr>
<td>Age &gt;50</td>
<td>Pelvic surgery (gynecologic and colorectal)</td>
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<tr>
<td>Female sex</td>
<td>Spinal anesthesia</td>
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<tr>
<td>Lower body mass index (BMI)</td>
<td>Intraoperative fluid administration &gt;750 mL</td>
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<tr>
<td>Advanced pelvic organ prolapse</td>
<td>Estimated blood loss &gt;100 mL</td>
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<tr>
<td>Baseline bladder dysfunction</td>
<td>Postoperative opioid use</td>
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<tr>
<td>Previous incontinence surgery</td>
<td>Postoperative UTI</td>
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**Abbreviations:** UTI, urinary tract infection; POUR, postoperative urinary retention.
in the spinal group (7.5% versus 69.6%, $P<0.001$). It was notable that, compared to the spinal group, the pudendal group had lower pain scores in the postoperative period (2.7 versus 5.2, $P<0.001$) and decreased use of other analgesics (20% versus 55%, $P<0.001$). In a logistic regression model, pain was an independent risk factor for postoperative retention (OR = 1.26, $P=0.02$).

Incontinence and prolapse surgery have the highest rates of POUR compared with other benign pelvic surgery. Incontinence surgery is designed to create some degree of urethral obstruction, but the key is to avoid overcorrection of the urethral angle. Likewise, prolapse repair can also lead to changes in the urethra–vesical junction (UVJ) that can affect voiding function. Furthermore, surgery in the vaginal and retropubic spaces is more likely to cause disruption of perforating nerve branches, leading to transient neuropathy, which can affect bladder sensation and micturition. In one study evaluating POUR after resection of endometriosis, the majority of patients (87.5%) with resection of at least one branch of the inferior hypogastric nerve demonstrated retention and need for self-catheterization.

Clinical factors affecting POUR include age, female sex, lower body mass index, previous incontinence surgery, advanced-stage prolapse (stage 3–4), and postoperative UTI. Whereas age is an independent risk factor for POUR, it is exacerbated by the fact that women undergoing incontinence and prolapse repair are generally older than the average patient undergoing benign gynecologic surgery. Keita et al$^{21}$ found that women over the age of 50 years had a 2.4 times higher likelihood of experiencing POUR. Toyonaga et al$^{22}$ found that women undergoing anorectal surgery were 2–8 times as likely to experience POUR compared with men (OR = 2.1 for hemorrhoidectomy and OR = 8.3 for fistulectomy).

Intraoperative risk factors for POUR have also been demonstrated. Keita et al$^{21}$ found that intraoperative fluid administration $\geq 750$ mL (OR = 2.3) and bladder volume $\geq 270$ mL on entry to the postanesthesia care unit (OR = 4.8) significantly increase the risk of POUR. Joelsson-Alm et al$^{23}$ found that preoperative ultrasound monitoring of bladder volume significantly decreased the rate of POUR (17.0% versus 27.1%, $P=0.045$). Toyonaga et al$^{22}$ confirmed the effect of fluid and bladder volume, showing that perioperative fluid restriction decreased the rate of POUR (7.9% versus 16.7%, $P<0.0001$, OR = 4.05). Hakvoort et al$^{10}$ found that the presence of a high-grade cystocele (OR = 2.5, 95% confidence interval [CI] = 1.3–4.7), performing a levator plication (OR = 4.3, 95% CI = 2.0–9.3), performing a Kelly plication (OR = 5.1, 95% CI = 1.7–15.5), and the amount of intraoperative blood loss (OR = 1.4 per 100 mL, 95% CI = 1.1–1.8) were all independent risk factors for POUR after vaginal prolapse surgery.

Multiple studies have confirmed the association between neuraxial anesthesia and POUR. Choi et al$^{24}$ performed a systematic review of the effect of anesthesia on bladder function and found that the duration of bladder dysfunction is correlated with the dose of intrathecal local anesthesia as well as the potency of the anesthetic used. By denervating the regional pelvic nerves for pain control, neuraxial anesthesia also leads to denervation of the bladder for a transient period of time. Local anesthetics in the spinal bolus block both the afferent and efferent pathways of the voiding mechanism. The addition of opioids to this bolus has been shown to enhance the degree of bladder dysfunction, with increased bladder capacity and decreased detrusor contractility due to both local action at the spinal nerves and also in the pontine micturition center.$^{25}$ The degree of denervation from neuraxial anesthesia can range from mild to severe.

The strength of the effect of neuraxial anesthesia on bladder function depends on the type of intrathecal local anesthetic used.$^{21}$ Incidence rates were greater than 20% in studies using the more potent drugs tetracaine or bupivacaine; those using procaine or lidocaine had retention rates less than 20%. Time to return of normal voiding also varied, ranging from 103 minutes for 2-chloroprocaine to 462 minutes for bupivacaine, with a rate of 260 minutes for lidocaine. Likewise, the effect of intrathecal opioids also varied depending on potency. Using morphine in the spine increases retention rates to 25%–36%, while rates of retention after use of fentanyl or sufentanil were 0%–25%.$^{24}$ However, even these less potent opioids can demonstrate an effect on bladder dysfunction. One study showed that the addition of sufentanil to intrathecal lidocaine prolonged the length of bladder dysfunction by over an hour and a half (233 minutes versus 332 minutes, $P<0.05$).$^{25}$ Similar effects have been seen with epidural anesthesia, although less data exist. Rates of bladder dysfunction after the use of long-acting opioids in epidural anesthesia ranged from 9.2%–79.5%, compared to 0%–40% for short-acting opioids.$^{23}$

Predicting POUR

It has been long held that preoperative urodynamic parameters have been considered useful in predicting POUR, but more recent studies have called these findings into question. Some studies have reported an associated between specific preoperative urodynamic findings and POUR, specifically peak flow rate, detrusor pressure during voiding, and the
presence of straining to void. Bhatia et al identified three types of voiding mechanisms on preoperative urodynamics for women undergoing Burch colposuspension: 1) detrusor contraction with urethral relaxation (DC + UR), which is considered normal function; 2) urethral relaxation with absence of detrusor contraction) (UR w/o DC); and 3) urethral relaxation with absence of detrusor contraction and presence of valsalva voiding (UR + VV). They found significantly increased length of retention in the latter two groups that had abnormal voiding: 4.8 days versus 6.5 days versus 10 days, respectively (P < 0.05), with a 12-times-greater risk of retention in the VV group. Further, 20% of patients (6/30) required prolonged catheterization (>7 days): one patient in the UR w/o DC group and five patients in the UR + VV group. No patients with normal baseline voiding (DC + UR) demonstrated prolonged retention. Interestingly, only one of these six patients admitted to preoperative voiding problems.

PVR in the prolonged catheterization group ranged from 150–335 mL, with a mean of 210 mL.

A more recent study has shown contradictory results when assessing the utility of preoperative urodynamics to predict postoperative voiding dysfunction. Lemack et al reported on women randomized to traditional pubovaginal fascial sling versus Burch colposuspension, with voiding dysfunction more strictly defined as either the need for a catheter after 6 weeks postsurgery or surgical revision. Women with baseline PVR >150 mL or those with obstructed voiding without significant prolapse were excluded. Rates of voiding dysfunction were significantly higher in the fascial sling group: 17.2% (49/285) versus 2.7% (8/294) (P < 0.05). However, there were no differences in preoperative urodynamics parameters between those with and without voiding dysfunction. Mean maximum flow during noninvasive flowmetry values were similar among women with and without voiding dysfunction (23.4 mL/sec versus 25.7 mL/sec, P = 0.16), as well as when assessed separately in the Burch colposuspension group (25.8 mL/sec versus 25.7 mL/sec, P = 0.98) and in the pubovaginal sling group (23.1 mL/sec versus 25.7 mL/sec, P = 0.17). Voiding pressures and degree of abdominal straining were also similar and not associated with POUR. However, this study was very stringent in its exclusion criteria, with exclusion of women with preoperative PVR >150 mL and those with obstructed voiding at baseline. It is possible that a more generalizable population would demonstrate preoperative voiding dysfunction on urodynamics that could predict POUR.

Management

Identification

The key to management of POUR is early identification. All patients undergoing surgery for the correction of incontinence and/or prolapse should have an assessment of voiding function prior to discharge. There are several ways to assess voiding function—the gold standard is by measuring a PVR. The assessment of voiding function in the postoperative period is often called a voiding trial. Various methods exist, ranging from minimally involved techniques to strict protocols. On the minimal side is documentation of ability to void, with no measurement of voided volume or PVR. A more stringent voiding trial will include measurement of voided volume, followed by measurement of PVR. There are two common techniques for performing this type of voiding trial—spontaneous fill or retrograde fill. The key is to have an adequate bladder volume prior to starting the voiding trial. If the bladder volume is too low, it does not provide an accurate assessment of voiding function and may not fully assess for POUR. This could lead to a scenario wherein a patient successfully voids a small amount of urine in the postanesthesia care unit and is discharged without a catheter. After she arrives home she develops larger bladder volumes due to hyposensory and then experiences POUR because she is unable to successfully void at higher volumes. To avoid this type of false-negative test, the bladder should contain at least 300 mL prior to attempting to void. With the spontaneous fill method, the bladder is allowed to fill passively and the patient voids when she feels fullness. However, the downside to this method is that it may lead to an inadequate void if she her starting bladder volume is low. There is no way to ensure an

Table 2 Management of POUR

<table>
<thead>
<tr>
<th>Early POUR (&lt;6 weeks post-surgery)</th>
<th>Prolonged POUR (&gt;6 weeks post-surgery)</th>
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<tbody>
<tr>
<td>1. Identification of incomplete emptying</td>
<td>Surgical options:</td>
</tr>
<tr>
<td>2. Rule out overcorrection of UVJ angle</td>
<td>- sling stretching, sling release, partial sling resection, or urethrolysis</td>
</tr>
<tr>
<td>3. Passive bladder drainage</td>
<td>Nonsurgical options:</td>
</tr>
<tr>
<td>4. Reassess voiding intermittently*</td>
<td>- acupuncture, moxibustion</td>
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</tbody>
</table>

Note: *If using indwelling foley, can repeat voiding trial weekly; if patient is self-catheterizing, can keep a bladder diary and assess PVR for improvement/resolution.

Abbreviations: UVJ, urethra-vesical junction; POUR, postoperative urinary retention.
adequate voided volume with the spontaneous fill method. With a retrograde fill voiding trial, the bladder is filled retrograde via a transurethral catheter to a set volume (typically 300 mL) and then the catheter is removed and the patient is asked to void. With either technique, the void is followed by measurement of PVR—either via in-and-out catheterization or bladder scan. Is it important to note with either filling technique that the PVR should be measured within 10–15 minutes of the void, in order to ensure accuracy.

A passing score on the voiding trial is determined by the PVR. The definition of passing varies considerably from clinician to clinician. Generally, a PVR less than 100–200 mL is considered passing; but some practitioners will use a percentage of total volume, with a PVR less than one-half to one-third of the total bladder volume. Thus, if a patient is retrograde filled to 300 mL and voids 200 mL, she would be considered to have passed her voiding trial, because she has a PVR of 100 mL, which is also one-third of the total bladder volume. The stricter definition of passing will use a smaller PVR cut-off (ie, 100 mL or one-third of total volume), while a more lenient definition of passing will use a larger PVR (ie, 150–200 mL or one-half of total volume). Use of percentages rather than specific cutoff values for PVR is less reliable, as it is dependent on the starting bladder volume. For example, if starting bladder volume is 600 mL, a PVR of one-half is still 300 mL which most would agree is too high to be considered normal. Obviously, with a stricter definition for passing the voiding trial, overall failure rates will be higher, but the false negative rate will be lower. The goal is to balance the rate of false negatives (patient is thought to void adequately but may develop POUR at home) with the rate of unnecessary catheterizations (false positives).

One study assessed the accuracy of the spontaneous and retrograde fill techniques using strict criteria for failure (PVR ≥100 mL). Patients undergoing midurethral mesh sling and/or prolapse repair underwent both voiding trial techniques but were randomized as to which one was performed first. The rate of POUR (defined as failure of voiding trial) on day of hospital discharge was predictably higher on the basis of strict definition of PVR: 84% for the spontaneous fill method and 62% for the retrograde method. The retrograde method had 94.4% sensitivity and 58.1% specificity to detect prolonged POUR lasting at least 7 days, compared with the spontaneous method with 100% sensitivity and 25.8% specificity. Positive and negative predictive values for the retrograde method were 56.7% and 94.7%, respectively, compared with the spontaneous method results of 43.9% and 100%. The retrograde was both more accurate in predicting prolonged retention and was preferred by patients. For this study, PVR was measured with in-and-out catheterization.

An alternative to catheterization for measurement of PVR is bladder scanning, where a transabdominal ultrasound probe is used to calculate bladder volume by imaging pockets of fluid. Older studies found this method to be less accurate than catheterization, with ultrasound PVR measurements tending to underestimate actual residual volume. One study found the ultrasound PVR measurements were most accurate (60.6%) when the readings were below 50 mL and least accurate (10%) when readings were higher than 150 mL; this is concerning if one is trying to rule out POUR on the basis of an elevated PVR. Readings between 50–150 mL were found to be 27% accurate. Newer studies have shown improved accuracy of bladder scanning to assess PVR, likely due to improvements in both technology and technique. Al-Shaikh et al assessed three-dimensional ultrasound in comparison to catheterization and found good correlation between the two techniques (r=0.79, P<0.001). The accuracy in this trial may have been improved not only by the use of a three-dimensional scanner, but also by the fact that four separate scans were performed and averaged. Typically only one or two scans are performed. It is important to recognize that technique does play a role and proper training is important in order to obtain accurate measurements.

**Early versus late POUR**

In addition to assessing for immediate postoperative POUR, a separate but equally important issue is delayed retention occurring after discharge from the hospital. Depending on the PVR cutoff value used to determine voiding trial pass rates, a number of patients who initially pass their voiding assessment will experience delayed POUR at home. This can occur when patients have longer intervals between voids, leading to larger bladder volumes, often due to residual hyposensation from surgery and the use of narcotics for pain control, and also in the face of a postoperative UTI. Whereas complete urinary retention is usually obvious, partial retention may be subtler to diagnose. Clinical symptoms that may provide indication of delayed partial urinary retention include urinary hesitancy, slow stream, the need to lean forward to void, the need to perform a Crede maneuver, straining to void, double-voiding, a feeling of incomplete emptying, or overflow incontinence. A history of each of these can be elicited by any caregiver.

If delayed POUR is suspected, the patient should be brought in for assessment. Physical examination should include visual inspection of the urethra, with placement of a sterile cotton swab into the urethra up to the level of the UVJ
Figure 1. An overcorrected UVJ angle may be demonstrated as a hyperacute urethral resting or straining angle (a negative cotton swab angle or less than 0° from the horizontal), or by a rigid or fixed urethra. Difficult placement of the cotton swab may also reveal a kinked urethra or a urethral stricture, which can occur with either incontinence or prolapse repair. A vaginal examination should also include an assessment for other causes of POUR, including hematoma or abscess, pelvic floor spasm, and any specific areas of tenderness or pain. In addition to gathering a history and performing a physical examination, obtaining a PVR is essential. Ideally, a voiding trial would be performed, but at the very least a PVR would allow the clinician to assess the degree of retention.

Bladder drainage
Management of POUR is fairly straightforward. The goal is to decompress the bladder to avoid long-term damage to bladder integrity and function. Immediate catheterization is always the first step. This may be performed either with in-and-out catheterization or with placement of an indwelling foley catheter. Although placement of an indwelling foley catheter is easier, there are several drawbacks to prolonged
use of this method. Indwelling catheters lead to increased rates of UTI compared to intermittent catheterization. In a trial of 87 women undergoing vaginal prolapse repair, patients were randomized to clean intermittent catheterization (CIC) versus indwelling catheter for the management of POUR.32 Women with CIC had significantly less risk of both bacteruria (14% versus 38%, P=0.02) and UTI (12% versus 33%, P=0.03) compared to women with an indwelling catheter. In addition, CIC led to faster resolution of POUR compared with indwelling catheter (18 hours versus 72 hours, P<0.001), likely due to the fact that every CIC provides a PVR to measure return of bladder function. It is likely that increased length of indwelling catheter use increases the likelihood of UTI. One trial comparing three periods of catheter use after pelvic surgery found no statistical difference in UTI rates when comparing 1 day (12.2%) versus 3 days (7.9%) versus 5 days (23.1%) of catheter use.33 However, the number of subjects in each group was low and the study was likely underpowered to find a difference. A large retrospective trial of more than 35,904 patients undergoing cardiac, vascular, abdominal, or orthopedic surgery also assessed length of catheter use.34 While 86% of patients had some form of perioperative catheter use, 50% maintained a catheter longer than 2 days postoperatively, with mean catheter use of 3.5 days across all groups. Those who had a catheter longer than 2 days were twice as likely to develop a UTI (P=0.02) and also had an increased 30-day mortality risk (P<0.001). The results of this trial led to the National Surgical Infection Prevention guideline for removal of all transurethral catheters by postoperative day 2 unless clinically indicated for surgical site healing.35

Another option for prolonged bladder drainage in the postoperative period is a suprapubic catheter (SPC). This type of catheter is typically placed intraoperatively at the time of surgical procedures that are more likely to cause retention, by inserting the catheter directly through the abdomen in the midline of the suprapubic area (just superior to the pubic bone) and directly into the bladder. While the risk of bowel injury is very low, techniques to decrease the risk of complications include bladder distention to increase target size (300–500 mL), placement of the catheter within 1–2 cm above the pubic symphysis, cystoscopic guidance, Trendelenburg position, and aspiration of catheter contents to confirm patent drainage of urine.36 SPC has several advantages over a transurethral catheter, including lower infection rates, less maintenance, earlier ambulation, more comfort, and ability to assess bladder function with the catheter in place.36–38 A systematic review found that patients were more than three times as likely to develop a UTI with a transurethral catheter than with an SPC (OR =3.22, 95% CI =1.95–5.4).39 Although there was an increased rate of complications in the SPC group (29% versus 11%, P=0.01), these were related to tube malfunction, with no visceral injuries among 1,300 patients. When comparing SPC to CIC, there is no difference in the rate of UTI.39,40 Some studies have reported no difference in patient preference between these two techniques, while others have shown a preference for SPC.39,40 An SPC is surprisingly well-tolerated by patients. The major benefit of an SPC over both CIC and indwelling transurethral catheter is that a voiding trial can be conducted with the SPC in place. To conduct a voiding trial, the SPC tube is clamped in order to allow the bladder to fill spontaneously; alternatively, the bladder can be filled retrograde through the SPC. The patient can then void with a full bladder and the PVR can be measured by simply unclamping the SPC and measuring the residual volume. No additional catheterization is needed.

Discontinuation of bladder drainage
The decision for when to discontinue catheter-assisted bladder drainage in the postoperative period can be assessed in an ongoing fashion by measurement of PVR. As discussed earlier, the rate of prolonged POUR beyond 4 weeks is low, ranging from 2%–4% for a tension-free transvaginal mesh sling. Therefore, transient retention can be expected to resolve spontaneously within 4–6 weeks with the use of some form of bladder drainage to prevent damage to the bladder. Studies have shown the use of an indwelling catheter can actually prolong catheter use by the very fact that bladder function cannot be assessed with an indwelling catheter in place.32,39 Both CIC and SPC methods allow for measurement of PVR while still maintaining bladder drainage. With any method, the decision to discontinue bladder drainage is based on PVR measurements. Generally, the PVR will trend downward as bladder function returns. In our practice, a PVR ≤100 mL two times in a row and at least a 200 mL voided volume are criteria for passing. Patients are nonetheless cautioned to avoid bladder distention in the weeks following a passed voiding trial, as they may still have some hyposensation.

Surgical management
When POUR does not resolve spontaneously, more active management may be required. (Table 2) Because prolonged retention is more common with incontinence repair than with prolapse repair, the relevant options usually include sling stretching, sling release, partial sling resection, or urethralysis. Although some clinicians favor intervention
early in the postoperative period, especially in the setting of complete retention, it is reasonable to wait at least 4 weeks postsurgery as most cases of retention will resolve spontaneously during that time. Early intervention runs the risk that the intervention may be completely unnecessary and that any associated complications would be regrettable. Molden et al\textsuperscript{41} retrospectively assessed outcomes after sling revision for both retropubic and transobturator slings. Techniques included urethral dilation, stretching of the sling, incision to release the sling, resection of the central portion of the sling, and urethrolysis. Of 175 patients with prolonged bladder dysfunction, obstructive voiding symptoms were seen in 23\% (27/119) with a retropubic sling compared to 12\% (6/51) with a transobturator sling (\(P=0.04\)). While voiding dysfunction resolved in 80\% of patients, 21\% developed de novo SUI. There was no difference in outcomes on the basis of the type of revision performed. Interestingly, while the median time to sling revision was 62 days, early sling revision (\(\leq 14\) days) was associated with less de novo SUI compared to delayed sling revision (15–90 days or >90 days). However, this may be difficult to translate into practice as most revisions would be accomplished by 4–6 weeks, which cannot be teased out of the 15–90 day group.

South et al\textsuperscript{42} assessed sling revision for both retropubic fascial slings and tension-free mesh slings and found that early release had a higher success rate for improvement in voiding function compared to late release (91\% versus 71\%, \(P=0.01\)), however early and late were classified as 6 months and 33 months, respectively. The rate of de novo SUI in this study was 8.9\%, with no difference between early and late sling release (9.5\% versus 7.9\%, \(P=0.99\)).

Segal et al\textsuperscript{43} explored outcomes after sling release versus vaginal urethrolysis versus retropubic urethrolysis for various incontinence procedures. Of 44 patients who underwent either a sling release or urethrolysis, 93.2\% (41/44) had it performed for obstructive symptoms, with resolution in 73.2\% (30/41) of cases; 34\% (15/44) of subjects developed de novo SUI after sling release or urethrolysis. There was no difference in outcomes between the three techniques. Rardin et al\textsuperscript{44} reported on 1,175 patients who underwent placement of a tension-free retropubic mesh sling and found that 1.9\% (23/1,175) underwent sling release for either POUP (20/1175) and/or urge urinary incontinence (10/1,175). Mean time to sling release was 17 weeks (range, 2–69 weeks), and all patients with POUP had resolution, with 13\% developing de novo SUI. Although various techniques exist, sling release offers the least invasive approach with equivalent outcomes to sling resection or urethrolysis. Timing of sling release should not be significantly delayed past the time when spontaneous resolution of POUP would be expected, ie, 4–6 weeks postsurgery.

**Nonsurgical management**

Alternative nonsurgical treatments have been attempted for POUP, including medications such as alpha adrenergic antagonists and anxiolytics, and less conventional techniques such as moxibustion and acupuncture. (Table 2) Alpha blockers have shown to have little benefit in women for the relief of obstructive retention when compared to men, and there is no evidence that this class of drug helps in the setting of POUP. There is an older report of success with oral anxiolytics for the treatment of POUP, but the data are several decades old and the study did not assess current incontinence procedures.\textsuperscript{45} More recent evidence from a randomized placebo-controlled trial of preoperative lorazepam showed no effect on the rate of POUP.\textsuperscript{46} Moxibustion offers an interesting option that has shown some benefit in treating POUP. A randomized trial comparing moxibustion versus hot compress versus infrared radiation for patients with POUP after hemorrhoid and fistula surgery showed significantly higher rates of improvement with moxibustion (90\% versus 40\% versus 55\%, respectively; \(P<0.05\)).\textsuperscript{47} Another study assessing two techniques of acupuncture found recovery of bladder function in both groups (90\% and 73\%, respectively, \(P<0.05\)) compared to baseline, with average PVR after two treatments of 93.5±15.5 mL and 102.5±15.7 mL, respectively.\textsuperscript{48}

**Prevention**

Whereas some risk of POUP is inevitable, certain risk factors are modifiable. Women planning to undergo incontinence and/or prolapse repair should be counseled about the high probability of transient POUP, but they can be reassured by the low rate of prolonged retention. They should be prepared for the possibility of needing to catheterize in some fashion and can even be taught CIC prior to surgery. Patients with specific risk factors for POUP, namely female sex, older age (>50 years), lower body mass index, previous incontinence surgery, and advanced-stage prolapse, should be counseled regarding these risks. Managing intraoperative fluid administration and bladder volume, along with minimizing blood loss, will decrease the risk of POUP. Although we know that spinal anesthesia can increase the risk of POUP, the surgeon can request that the anesthesiologist use shorter-acting intrathecal local anesthetics and minimize the use of intrathecal opioids. In
the recovery room, the risk of retention can be reduced by minimizing bladder distention as well as minimizing the use of intravenous and oral opioids, while still managing postoperative pain (perhaps with intravenous acetaminophen and nonsteroidal inflammatory drugs).

Urodynamic testing of bladder function does not predict POUR in the average patient; however, preoperative urodynamic assessment should be considered for patients with baseline voiding dysfunction, mixed incontinence, or advanced prolapse. For patients with stress-predominant urinary incontinence and no prior incontinence procedures, performing a standing stress test and measuring a PVR prior to surgical repair of incontinence and pelvic organ prolapse is often adequate. 25

**Conclusion**

Postoperative urinary retention is a common perioperative issue, not only in gynecologic procedures, but also in a wide range of abdominal, pelvic, and orthopedic procedures. Patients who are at higher risk for POUR – either due to the procedures being performed or their clinical risk factors – should be counseled regarding the risks and management options for POUR prior to their surgery. Being informed and prepared can go a long way in helping patients deal with a condition that, while not life-threatening, can be very stressful and can affect activities of daily living for both themselves and their family. The key to management of POUR is early identification, in order to avoid serious and preventable sequelae. With many options available to assess bladder function, there is no excuse for not performing some type of postoperative assessment of voiding function prior to patient discharge.

Having a POUR protocol both within one’s practice and in the surgical center can significantly increase the success of properly assessing all patients and setting up proper management for treatment. 29 This protocol can also help ease patient anxiety with management of their retention, by giving them a treatment plan with time points for reassessment of the condition and for moving from passive to active management, i.e., surgical intervention. Having patient buy-in can decrease nursing time spent on reassurance as well as decrease needless early intervention, given that most cases of POUR will resolve spontaneously. Although POUR is a serious condition that can have serious consequences if left untreated, it is easily diagnosed and typically self-resolves. Clinician awareness of the condition and vigilance in its diagnosis are the key factors in successful surgical care for this phenomenon.

**Disclosure**

This work has not been previously published in any form. The author reports no other conflicts of interest in this work.

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