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Optimising the tension of an autologous fascia pubovaginal sling to minimize retentive complications

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Abstract Aim: To determine the optimal degree of pubovaginal slings (PVS) tension,

measured by lax sling dimensions to minimize the risk of urinary retention. Methods: This prospective study analyzed female patients undergoing PVS for stress urinary incontinence (SUI) by two surgeons over 24 months from January 2016.

Intra-operative measurements of lax sling dimensions tented over rectus fascia were recorded. Logistic regression was used to analyse the likelihood of urinary retention (more than 3 months of intermittent self-catheterisation (ISC) or surgical revision) for given PVS dimensions. The secondary analysis assessed for an association between PVS measurements and persistent SUI.

Results: Fifty-one patients were recruited with a median age of 53 (34-78) and follow-up of 11 (3-20) months. All but one patient reported improvement of SUI. Ten (19.6%) patients developed postoperative urinary retention. Five (9.8%) resolved after a temporary period of ISC. The other five (9.8%) required ongoing ISC or sling division. A strong association existed between short sling height and prolonged urinary retention (P = 0.00). Receiver operating characteristic (ROC) curve analysis showed a sling height of 40 mm had a sensitivity of 100% and specificity of 51% for retentive complications (area under curve [AUC] = 0.90). Lax sling height up to 60 mm was not associated with persistent SUI.

Conclusions: Stretching the sling suspension sutures at least 40 mm above the rectus fascia was associated with a lower risk of urinary retention than less than 40 mm. This simple technique would appear to be worth evaluating in a larger sample. A looser sling did not compromise the cure of SUI at a mean follow-up of 11 months.

KEYWORDS

autologous fascia pubovaginal sling, stress urinary incontinence, urinary retention, voiding dysfunction

INTRODUCTION 1

Stress urinary incontinence (SUI) in women impacts greatly on physical, sexual, and psychosocial health. It is an undeniably common problem with prevalence estimates in Australia ranging between 13% and 46%,¹

increasing with age. Only 53%² achieve subjective cure with conservative treatments, with many subsequently choosing to opt for surgical correction.

Debate exists over the gold-standard surgical treatment: autologous fascia pubovaginal slings (PVS) vs synthetic mid-urethral slings (MUS). A recently updated

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meta-analysis³ failed to prove the superiority of one suburethral sling over the other; comparing both effectiveness and complication rates.

MUS has a successful dry/cure rate of 81% to 84%.⁴ This; however, is offset by 6% risk of bladder perforation, as well as up to an 8% incidence of mesh-related complications (extrusion, infection, scarring, dyspareunia, chronic pain, and erosion).⁴ Due to the comparative simplicity and speed of MUS insertion, it has become the procedure of choice for treating uncomplicated SUI by most incontinence surgeons.

By comparison, PVS utilizes native tissues and is thus spared from mesh specific complications. A metaanalysis found the SUI cure rate with PVS was 82% to 90% in patients with more than 4-year-follow-up.⁴ Nonetheless, it is a longer and more complex procedure with a slower recovery, requires both vaginal and abdominal incisions with inherent wound complications (8%), and has higher rates of obstructive voiding patterns. However, since the FDA issued a warning on transvaginal mesh used for pelvic organ prolapse (2008-2011), MUS slings have also come under the spotlight. As a result, more PVS surgery is now being done, with a corresponding decrease in MUS surgery.⁵

Historically, PVS was used for intrinsic sphincter deficiency (ISD, more severe urinary incontinence), associated with greater leakage and tendency to urethral fixation. Given the increasing awareness of complex longterm mesh complications, indications for PVS have broadened to include "index" SUI. PVS remains the preferred treatment for complicated SUI (following failed prior incontinence surgery, to support a urethral diverticulum repair or fistula repair).

There is little randomized controlled data comparing MUS with PVS. Nonrandomised comparisons without controlling for risk factors show that the rates of sustained voiding dysfunction (lasting more than 1 month or requiring intervention) occur in 8% (4%-15%) of women undergoing PVS, compared with 3% (2%-4%) for MUS.4

The goal of PVS is to restore the proximal urethral support to address urethral hypermobility and ISD elements. Historical slings involved tunnelling of attached and mobilized rectus fascia with high rates of voiding dysfunction. Once it became evident that a free graft of fascia suspended on monofilament sutures could provide long term cure rates, that has become the standard. The method by which the suspension sutures are tied together so that slings do not result in urinary retention or voiding dysfunction is in evolution. Midurethral placement of a sling of any fabric is less likely to cause voiding dysfunction. Proximal urethral placement may, however, be preferable in cases where the proximal

urethra is wide open at rest-a condition that may be diagnosed on fluoroscopy. Such preoperative characterization is rarely available. Where urethral resupporting is required rather than correction of ISD, a loose sling only may suffice.

Over the years multiple attempts have been made to develop a reliable way to appropriately tension a PVS. These include an intra-operative rectal ultrasound to measure the urethrovesical angle, using a perurethral cotton bud to maintain a specific urethrovesical angle and using a spring scale to measure the sling tension.⁶⁻⁸ However, these methods have not become standard practice. As such, it is now commonplace for incontinence surgeons to tension a PVS according to a simple heuristic rule: tie the suspension suture knot 2 or 3 fingers breadth above the rectus fascia⁹ (not too tight, not too loose, and lacking reproducibility).

The purpose of this study is to analyse sling suture dimensions with a view to improving the technique of tensioning PVS. We aimed to develop a reliable approach to the objective measurement of sling height (rather than subjective tensioning) which would minimize retentive complications and still achieve excellent continence rates.

2 PATIENTS AND METHODS

A prospective study was performed, analyzing clinical outcomes of all female patients undergoing autologous fascia sling for SUI by two Melbourne-based primary operators (HOC, JG) over a 24-month period commencing January 2016. Both surgeons were experienced with PVS, and used an agreed standardized surgical technique. All eligible women aged over 18 years were recruited from both the surgeons' private and public practices. Institutional ethics approval was obtained for this study (HREC/16/WH/201).

SUI was defined as the involuntary leakage of urine with physical activity, coughing, or sneezing. Only women who voided with detrusor contractions were included; pre-existing urinary retention or those who voided with abdominal straining were excluded. Women with mixed incontinence (ie both SUI and urge incontinence) and women with a previous failed SUI surgery or previous bulking procedure performed for SUI were eligible for study inclusion. Patients with concomitant prolapse were excluded, as were women who underwent an autologous fascia sling procedure for a primary reason other than SUI. Our study excluded patients with an acontractile bladder and those who strained to void (unknown PdetQmax). All patients had urodynamic studies (UDS) before surgery. Women who demonstrated

bladder contractility (any rise in Pdet) during the voiding phase were classified as those with detrusor contractions and were included in the study.

Both primary surgeons used a consistent operative technique. The initial technique was developed with the collaboration of both surgeons (the first 12 cases) to confirm uniform technique before separate independent implementation.

The procedure was performed through a combined abdominal and vaginal approach. With the patient in a dorsal lithotomy position, a 6 to 8 cm transverse abdominal incision is made about 3 cm above the pubic bone. Autologous tissue was harvested from the anterior rectus sheath $(8 \times 1.5 \text{ cm})$ and placed transvaginally as a suburethral sling at the bladder neck without causing upward distortion or hyper-suspension. The sling length and width of 8×1.5 cm was chosen because of the surgeons' familiarity and previous experience. The current literature describes different fascial sizes and variation in certain steps of the surgical technique.¹⁰ There is also a paucity of data in the literature if the sling length or width can alter the clinical outcome. The sling was placed at the level of the bladder neck in all cases: the proximal border of the sling was placed at the level where the catheter balloon started (indicating the level of the bladder neck).

Monofilament, polypropylene Prolene sutures were secured to each end of the harvested fascia when the fascia was still not fully removed from the patient, as the authors found that this partial attachment stabilized the fascia for easy placement of the Prolene sutures, as compared with placing them after the fascia was fully removed and on the nurse's instrument table. The Prolene sling arms were then passed through retropubic tunnels and delivered just lateral to the rectus muscles, through the inferior leaf of the rectus fascia.

The operation then proceeded to tie the ends of the fascia sling via the traditional method of "2-3 fingerbreadths" above the fascia. The fascia sling was lifted upwards by its knot so that its greatest height was achieved without resistance from, or causing distortion to, the abdominal wall tissue. Based on the traditional tensioning method and forming an isosceles triangle, the base and height of this tented sling were recorded in millimeters (Figure 1). These measurements were used to reflect the laxity of the PVS. During the measurement of the sling height, a 14 F Foley catheter was in place. The urethra and bladder neck were observed at this time as the vaginal incision was still not closed. Measurement of the sling height did not cause distortion of the urethra in all cases.

It is important that the sling was not hyper-suspending the bladder neck before tying the knot as this may



FIGURE 1 Measured sling height and base dimensions

cause the knot to retract back later. The vaginal incision was not closed until after the knot was tied. This allowed the surgeons to check that the sling arms had not been pulled up too tight inadvertently, just before the final tensioning. If the bladder neck was observed to be hypersuspended, the sling arms could be readjusted via the vaginal incision, before the final tensioning was done (Figures 2 and 3).

Cystoscopy was then done after tensioning to ensure no bladder injury but was not routinely used as an aid for sling tensioning.

The measurement technique was found to have good inter-rater reliability during the initial collaborative cases and can be easily passed on to other surgeons. After independent implementation, the primary surgeon who did the tensioning measured the sling height. Potential bias and variability can occur here but this was reduced with ensuring that the sling arms were not pulled upwards when tying the knot.

Routine postoperative management was trial-of-void on day 2. All women were routinely reviewed at 6 weeks and 3 months with a uroflow study, urogenital distress inventory 6 (UDI-6) and Patient Global Impression of



FIGURE 2 Study endpoints



FIGURE 3 Distribution of measured sling dimensions

Improvement (PGI-I) questionnaire. These surveys provided a component-based "symptom severity score" and an overall appraisal of the clinical outcome, both validated for use in incontinence research.^{11,12}

Long-term voiding dysfunction was the primary clinical endpoint, defined as prolonged intermittent self-catheterisation (ISC) beyond 3 months or division/ revision of autologous fascia sling.

The collected dataset also included: age, body mass index (BMI), pre and postoperative self-reported pad usage, valsalva leak point pressure (VLPP) from preoperative UDS, postoperative postvoid residual volumes, and encountered complications.

Categorical variables are presented as percentage number of patients, and continuous variables as median with range. Logistical regression was used to analyse the likelihood of a sling being "too tight" for different suprarectus sling height and base lengths measured intra-op. Subjective outcomes were assessed via UDI-6 and PGI-I.

3 | RESULTS

Five patients were lost to complete follow-up (9%). Three patients declined to be part of the study. A total of 10 patients were excluded from the study: two patients were excluded as they were already doing clean intermittent catheterization before the sling surgery for neurogenic reasons. Eight other patients were also excluded as their sling was placed not for the primary reason for treating SUI (two had urethral diverticulectomy, six had reinforcement of cystocele repair with a fascial sling).

Fifty-one women that completed the full study had a median follow-up of 11 (3-20) months. Median age was 53 (34-78), median VLPP 60 (25-145) cmH₂O, and median BMI

of 28 (18.4-43). The surgeons had a split caseload of 20:31.11 (21.6%) women had pre-existing detrusor overactivity (DO) with mixed lower urinary tract symptoms (LUTS), 5 (9.8%) underwent PVS as a salvage procedure following failed SUI surgery (four tension-free vaginal tapes [TVT], one Burch colposuspension) and 4 (7.8%) had PVS surgery for recurrent SUI postdivision of a tight TVT.

A total of 10 (19.6%) patients developed postoperative urinary retention. A total of 5 (9.8%) self-resolved after a temporary period of ISC less than 3 months. The remaining 5 (9.8%) women had persistent long-term retentive complications requiring indefinite ISC or revision surgery (division or loosening of the sling). Out of the five patients who had revision surgery, only one (the first one) was noted to have a change in the height of the knot (from 3.5 to 3.0 cm), discovered during the revision surgery. The tensioning during the initial surgery may have been done with the sling arms inadvertently pulled up before the tying of the knot which then retracted back down when the wound was closed, to a lower height. As this case was seen very early on during the study, the surgeons had been particularly careful not to accidentally pull up on the sling arms or hypersuspend the bladder neck, before the knot was tied in subsequent cases. The vaginal incision was not closed before sling tensioning, for this reason, to allow for assessment and readjustment of the sling if needed. The following four patients who needed revision surgery did not have a change in height of the knot.

Acknowledging the UDI-6 questionnaire's limitation for assessing overactive bladder (OAB), in our study, eight patients had worse/new urge incontinence post-op, so potentially 8/51 (15.7%) patients experienced de novo or worsening OAB.

The range of PdetQmax for the population was 9 to $41 \text{ cmH}_2\text{O}$ (median $22 \text{ cmH}_2\text{O}$). Pre-op PdetQmax was

TABLE 1 Demographics

Retentive complications	No (n = 41)	Yes (n = 10)	P value
Age (y)	53 (34-74)	54 (41-78)	0.86
BMI (kg/m ²)	29.4 (18.4-43)	25 (20.6-32)	0.08
VLPP (cmH ₂ O)	70 (25-145)	43 (30-65)	0.08
Pre-existing DO/ Mixed LUTS (n)	10	1	0.43
Recurrent SUI after previously divided sling (n)	2	2	0.17
Previous failed MUS/SUI surgery (n)	5	0	0.57
Primary operator ratio	17:24	3:7	0.72
Duration of follow-up (mo)	9 (3-20)	14 (3-19)	0.19

Abbreviations: BMI, body mass index; SUI, stress urinary incontinence; VLPP, valsalva leak point pressure.

not associated with retentive complications in our study (P = 0.85; Table 1).

On logistic regression (Table 2), short lax sling height was associated with postoperative urinary retention (OR: 0.74 [95% CI: 0.60-0.91]; P = 0.004) and unresolved voiding dysfunction and eventual sling revision (OR: 0.74 [95% CI: 0.61-0.90]; P = 0.003). Sling base length was not associated with postoperative retention (P = 0.22) or long-term retention (P = 0.15).

Age (P = 0.72), duration of follow-up (P = 0.20), and BMI (P = 0.10) were not confounders for urinary retention on univariate logistic regression. Nor was an operator (P = 0.51), VLPP (greater vs 60 cmH₂O or less, P = 0.34), pre-existing DO (P = 0.43), previous divided sling/TVT (P = 0.17) or previous failed SUI surgery (P = 0.57) using Fisher's exact test.

A sling height of 40 mm on receiver operating characteristic (ROC) curve analysis (Figure 4) had a sensitivity of 100% and specificity of 51% for retentive complications (area under curve [AUC] = 0.90). Thirty-three percent of those with a sling height of 40 mm or less encountered retentive complications, compared with 0% of those with a sling height greater than 40 mm (Fishers exact P = 0.003).

Other encountered post-op complications were: 3 (6%) with urinary tract infection, 4 (8%) with wound issues (haematoma, seroma, and superficial wound dehiscence), 1 (2%) with unsatisfactory cosmetic result, and 1 (2%) with sexual dysfunction.

Overall the PVS success rate was high, with a subjective resolution of exertion-related leakage in 82% (42/51), improvement in a further 16% (8/51) of women, and a failure rate of 2% (1/51). Of the eight women who

had a mild degree of residual SUI, only two needed to use a safety liner throughout a 24-hour period. One patient underwent transurethral bulking injection (Bulkamid) with good effect. A secondary analysis was performed on this subgroup to see if measured lax sling lengths also demonstrated the predictive ability for procedure failure and persistent SUI. Logistic regression compared sling height to the binary outcome of any SUI reported via UDI-6 question 3 ("Do you experience urine leakage related to physical activity, coughing or sneezing?"). Sling height was also compared with the self-reported need for ongoing pad usage postoperatively. Neither definition was statistically significant (P = 0.51 and 0.62, respectively). However, univariate analysis (Table 3) demonstrated an association between sling failure/persistent SUI and BMI (P = 0.045). No other variable (Table 2) was statistically significant including lax sling height (P = 0.56), VLPP (60 cmH₂O or less, 0.51), pre-existing DO (P = 0.34), salvage PVS after previous failed SUI surgery (P = 0.08), and re-do procedure after previous divided MUS (P = 0.48).

Regarding overall satisfaction with the procedure (assessed by PGI-I), 87% of women described their condition as being either "very much better" or "much better". A total of 11% of women said that it was "a little better," whereas only 2% stated their overall condition had worsened. In assessing the main determinants of global satisfaction, a positive correlation was only found between PGI-I score and responses from the retention domain question from UDI-6 (Spearman's $\rho = 0.34$, P = 0.02); indicating that patients' dissatisfaction was related to voiding dysfunction but not persistent SUI.

TABLE 2 Logistic regression of lax sling height and base measurements (Odds Ratios)

	Postoperative urinary retention	Unresolved voiding dysfunction
Sling height (mm)	$0.74 \ (P = 0.004)$	$0.74 \ (P = 0.003)$
Sling base (mm)	1.05 $(P = 0.22)$	1.07 $(P = 0.15)$

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FIGURE 4 ROC Curve of lax sling height and urinary retention

4 | DISCUSSION

Recognition that synthetic MUS are associated with low but very significant foreign body morbidity and chronic pain has redeveloped interest in alternatives. Free grafts of autologous rectus fascia (PVS) became a standardized procedure in the 1980s and popularised in the 1990s at a similar time to the rising popularity of synthetic MUS. Whilst the majority of surgeons had been offering the relatively ineffective anterior colporrhaphy and colposuspension, a major conversion to synthetic MUS occurred across the globe. The rise was related to relatively low operative morbidity and greatly improved success relative to colposuspension and colporrhaphy, boosted by industry support for education and operative training opportunities. Relatively low growth occurred in the number of PVS which were generally reserved for complex urethral disorders rather than "index" SUI.

The position of any sling in the proximal urethra is associated with greater voiding dysfunction that when placed in the mid-urethra. Curing or at worst, significantly improving severe sphincter dysfunction is the goal of PVS and any tensioning technique would not want to thwart that primary goal. Notwithstanding, if it is possible to reduce voiding dysfunction rates without compromising SUI cure, that would be desirable.

This pilot study of 51 patients demonstrated that a standardized approach to measuring intra-operative sling height during autologous rectus fascia PVS insertion may reduce voiding dysfunction. A wide range of sling heights (20 to 60 mm) was observed using the traditional method of tying the knot a couple of fingerbreadths above rectus; thus demonstrating the inherent difficulty in reproducing consistent tension from this method. The ROC curve analysis found a sling height greater than 40 mm was associated with a reduced incidence of voiding dysfunction. The significance of sling height is explained by the relative obstruction of the sling pulling the proximal urethra and bladder neck anteriorly, increasing the vesical pressure required to effect voiding.

Unlike sling height; however, sling base measurements (ranging from 40 to 80 mm), did not influence the incidence of postoperative urinary voiding dysfunction. Formed by the upward piercing sling through the lateral borders of the rectus muscle, sling base differences were attributed to the normal anatomical variation of rectus width. Quadratic modeling (assuming fixed points where PVS traverses the lower leaf of the rectus fascia), demonstrated that the base dimension negligibly contributes to laxity; doubling the base of a sling, affects its mobility by only 1 to 2 millimeters.

Higher rates of voiding dysfunction have been demonstrated in autologous fascia PVS patients who were dependent on Valsalva manoeuvre to void preoperatively by Iglesia et al¹³. Therefore, our study only included women who voided from detrusor contractions confirmed on UDS. Subgroup analyses were performed on enrolled patients with a history of previous incontinence surgery and those with pre-existing mixed incontinence. Neither of these factors appeared to affect clinical outcomes in this modest-sized cohort (P > 0.05).

PVS is now used for correction of both urethral hypermobility and ISD. In our series, the incidence of ISD on the basis of VLPP measurement was relatively high (51.4%). Although other studies have demonstrated that ISD has slightly poorer cure rates than hypermobility,¹⁴ VLPP (with <60 cmH₂O indicative of probable ISD)

TABLE 3 Univariate analysis of persistent postoperative SUI subgroup

Continuous variable	Log regression	Dichotomous variable	Fishers test
Sling height (mm)	0.56	VLPP <60 cmH ₂ O	0.51
Sling base (mm)	0.85	Pre-existing DO/mixed LUTS	0.34
Age (y)	0.32	Recurrent SUI after previously divided sling	0.48
BMI (kg/m ²)	0.05	Previous failed MUS/SUI surgery	0.08
Follow-up (mo)	0.41	Operator	0.98

Abbreviations: BMI, body mass index; SUI, stress urinary incontinence.

was not associated with higher rates of voiding dysfunction nor procedure failure/persistent SUI in our study. Continuing the study with larger numbers and a cohort with less severe SUI (more cases of "index" SUI) may help determine whether this technique could have greater application.

Modifications in traditional autologous fascia PVS technique have been described in an effort to minimize the inherent risk of urinary retention with PVS surgery. Linder et al¹⁵ published a series of 33 women who underwent a transobturator approach to autologous fascia sling placement. NIne percent of patients required short term ISC and nil required the release of sling for ongoing voiding dysfunction. Notably; however, 15% of patients required a salvage operation within the 14.9 months of average follow-up for ongoing SUI. Khan et al¹⁶ published results of mid-urethrally placed autologous fascia slings, with improvement rates of 90% and 75.4%, and dry rates of 48% and 50.8% at 1 and 10 years, respectively. ISC was performed by 6.5% of patients and division of sling occurred in 3.3% of cases compared with synthetic MUS slings.

Assessed through secondary analysis, PVS laxity was not found to predict sling failure (persistent SUI) with heights up to 60 mm. Subjective rather than objective cure rates were chosen for this analysis as self-reporting of improvement/cure must be considered of primary importance to patients. Despite subjective cure rates being typically lower than objective cure rates, an inability to detect a small correlation could readily be explained by the study being underpowered. Nonetheless, this study at least suggests that any correlation between a "loose" sling (40-60 mm height) and persistent SUI is insignificant compared with the correlation between a "tight" sling (<40 mm height) and voiding dysfunction; both statistically and in terms of clinical consequence. Patients reported worse PGI-I score because of voiding dysfunction complications and not because of mild residual SUI. For this reason, this study supports a prudent approach to tensioning fascia slings.

Consistent with other contemporary research, this study demonstrated that age does not appear to be a factor in cure or complication rates with sub-urethral slings,¹⁷ an important factor worthy of further study in an aging population. Obesity was the only variable found to have an association with persistent SUI. This highlights the importance of emphasizing relevant conservative measures in holistic SUI treatment: weight loss, minimizing alcoholic and caffeinated beverages, pelvic floor exercises, and treating constipation.¹⁸

Given the recent controversy in both academic and lay literature about mesh use in reconstructive pelvic surgery, patients are increasingly looking to alternatives.¹⁹ With a median follow-up of 11 months, this study is limited by the duration of follow-up and being a consecutive series of only 51 patients. The value of the study is the effort to standardize sling tension to decrease voiding dysfunction which may prove to be of significance as patients move to robust native alternatives.

Although we used the UDI-6 survey, another questionnaire such as OAB-q SF could also be used to assess for OAB, as UDI-6 did not ask about urgency. However, we chose to use UDI-6 because of its ease and incorporation of questions pertaining to urge incontinence, stress incontinence, and urinary retention, all in one questionnaire. In addition, the cough stress test was only done during the UDS preoperatively but not postoperatively. Ideally, this would be included for a more objective measure of SUI rather than just relying on the UDI-6 questionnaire.

Other limits of this study include its heterogenous nature (mixed incontinence and previous failed SUI surgery), although subgroup analyses failed to demonstrate any statistical impact. Furthermore, any conclusions drawn from this study must be tempered by the modest cohort size and low event rate (10/51). Potential confounders include the health of periurethral tissues, nature of vaginal closure, the position of the sling along the urethra, and degree of fibrosis that occurred during healing. Admittedly these factors may impact on final sling tension, but they are hard to control for and measure objectively. Unfortunately, menopausal and smoking status were not obtained as part of the study protocol. These are potentially useful data for any future study, as they may affect the fascia quality and the healing ability of the patient.

We also note that sling tension has to be adjusted according to each individual patient's degree of leakage. In this study, the exact sling height was not preplanned before the surgery, but rather we looked at the variability one can get from the traditional "2-3 fingerbreadths" method. A further study in the future should certainly look at randomizing the lax sling height to patients with different VLPP, and then correlated with outcomes such as success rate and urinary retention.

We demonstrated the presence of a strong association between lax sling height and reduced voiding dysfunction. Height less than 40 mm on ROC curve analysis had a sensitivity of 100% and specificity of 51% for retentive complications. In addition, a height of 40 to 60 mm was not found to be associated with poorer SUI cure rates. Given there was no observable detriment, we are encouraged to continue this standardized approach to sling tension. This serves as a proof-of-concept study to guide a larger, more detailed experimental study on the topic.

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5 | CONCLUSIONS

A higher lax sling height of greater than 40 mm above rectus fascia may reduce the risk of voiding dysfunction and the need for sling revision.

ETHICAL APPROVAL

Ethics approval was provided by the Western Health Low-Risk Human Research Ethics Panel (HREC/16/WH/201).

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