

## I-Stop TOMS Transobturator Male Sling, a Minimally Invasive Treatment for Post-prostatectomy Incontinence: Continence Improvement and Tolerability

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<b>OBJECTIVE</b>	To prospectively evaluate the efficacy and tolerability of the I-STOP TOMS transobturator male sling in patients with post-prostatectomy stress urinary incontinence. Minimally invasive techniques, such as slings, are becoming the standard of care for mild to moderate post-prostatectomy incontinence.
<b>METHODS</b>	From March 2007 to June 2009, 122 patients with post-prostatectomy stress urinary incontinence were treated with the I-STOP TOMS sling and followed up for 1 year in the Phase IV HOMme INContinence trial. The preoperative and postoperative evaluation included daily pad use, pad test, questionnaires evaluating urinary function and bother (University of California, Los Angeles, Prostate Cancer Index – urinary function short form, and International Consultation on Incontinence Modular Questionnaire – urinary incontinence short form) and uroflowmetry, including the post-void residual urine volume. Patient satisfaction and perineal pain were also assessed.
<b>RESULTS</b>	A total of 103 patients were followed up for 12 months. The surgical procedure was considered easy to perform. The mean daily pad use decreased significantly from 2.4 to 0.6 at 12 months of follow-up; 87.0% of the patients reported improved continence (59.4% completely dry, 20.3% 1 pad/d, 7.3% >1 pad/d), and 13.0% reported no improvement. All quality-of-life scores (University of California, Los Angeles, Prostate Cancer Index – urinary function short form, and International Consultation on Incontinence Modular Questionnaire – urinary incontinence short form) improved significantly after sling implantation. Treatment satisfaction was >90%. The post-void residual urine volume did not increase substantially, and acute urinary retention did not occur. The perineal pain scores were very low at follow-up. Wound infection was seen in 2 patients at the 1-month follow-up visit.
<b>CONCLUSION</b>	The I-STOP TOMS is a good treatment option for patients with post-prostatectomy stress urinary incontinence. With follow-up ≤12 months, most patients were continent or had improved continence. The intervention was well tolerated, with few infections. UROLOGY 79: 458–464, 2012. © 2012 Elsevier Inc.

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\*A complete list of the HOMme INContinence Study Group can be found in the Appendix.

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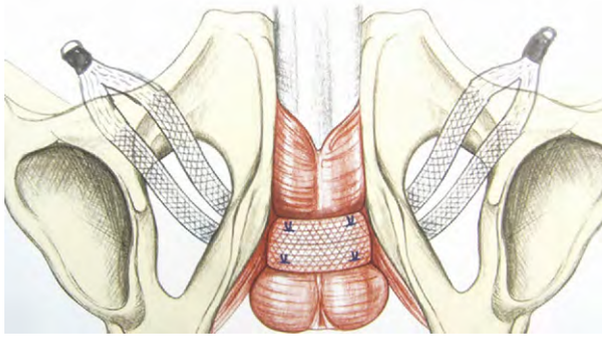
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**S**tress urinary incontinence (SUI) is common after prostatectomy, although the reported prevalence of this condition is highly variable (0.8%-87.0%) owing to the numerous definitions of post-prostatectomy continence.<sup>1,2</sup> Male SUI is mainly caused by sphincter deficiency; however, urethral support deficiency and increased mobility of the bulbar and membranous urethra can also be involved. Even mild post-prostatectomy SUI can strongly affect patients' quality of life. The initial treatment consists of pelvic floor muscle training and behavioral therapy, although the evidence on the efficacy of these treatments is rather weak. The artificial urinary sphincter remains the reference standard for severe SUI, which is often related to major sphincter deficiency.<sup>1</sup>



**Figure 1.** Suburethral I-STOP TOMS is a monofilament polypropylene (macropores  $>75 \mu\text{m}$ ) nonextensible 4-arm sling (2 arms on each side). Dimensions are  $45 \text{ cm} \times 1.4 \text{ cm}$ , with a 2.8-cm central part placed over urethra.

This procedure involves high costs, carries the risk of erosion and infection, and patients can be hesitant to have a mechanical implant or be unable to use it.

Therefore, other minimally invasive treatment options could be an alternative for patients with mild to moderate post-prostatectomy SUI. These include sling procedures, implantation of compressive adjustable balloons, or injection of bulking agents. The current guidelines from the International Consultation on Incontinence (ICI)<sup>3</sup> do not recommend the latter 2 options, for which multiple sessions are often required. Slings are becoming the standard of care for mild and moderate male SUI.<sup>3,4</sup> Although all available slings are placed under tension to occlude the urethra at rest and during stress maneuvers, they differ in the materials used, the methods of fixation and the position of the support.<sup>5</sup>

The 4-arm I-STOP TOMS transobturator male sling (CL Medical) is an adapted version of the 2-arm TOMS bulbar sling (CL Medical).<sup>6</sup> It is a monofilament polypropylene (macropores  $>75 \mu\text{m}$ ) nonextensible 4-arm large sling (Fig. 1). The dimensions are  $45 \text{ cm} \times 1.4 \text{ cm}$ , with a 2.8-cm central part placed over the urethra. The aim of the present trial was to evaluate the improvement in continence and quality of life and the tolerability of patients with post-prostatectomy SUI treated with the I-STOP TOMS transobturator male sling.

## MATERIAL AND METHODS

### Patients and Study Design

The eligible patients had SUI related to prostatectomy (radical or transurethral resection of the prostate) performed  $>6$  months before study entry. In addition, they were unresponsive to, or refused, urinary physiotherapy, and had a urinary incontinence score of 4-16 using the ICI Modular Questionnaire-urinary incontinence short form (score range 0-21).<sup>7</sup>

Patients with bladder outlet obstruction, bladder overactivity (assessed urodynamically), low compliance, or urethral or anastomotic stenosis (assessed by urethroscopy or urethrography) were excluded. Furthermore, patients were excluded if they had progressive prostate cancer as assessed by the prostate-specific antigen level, a history of prostate radiotherapy, neurologic

disorder, or chronic urinary retention with overflow incontinence. Current urinary tract infection resulted in temporary exclusion until the infection had resolved.

Patients were asked to stop any urinary incontinence medication, in particular anticholinergic agents, during the study.

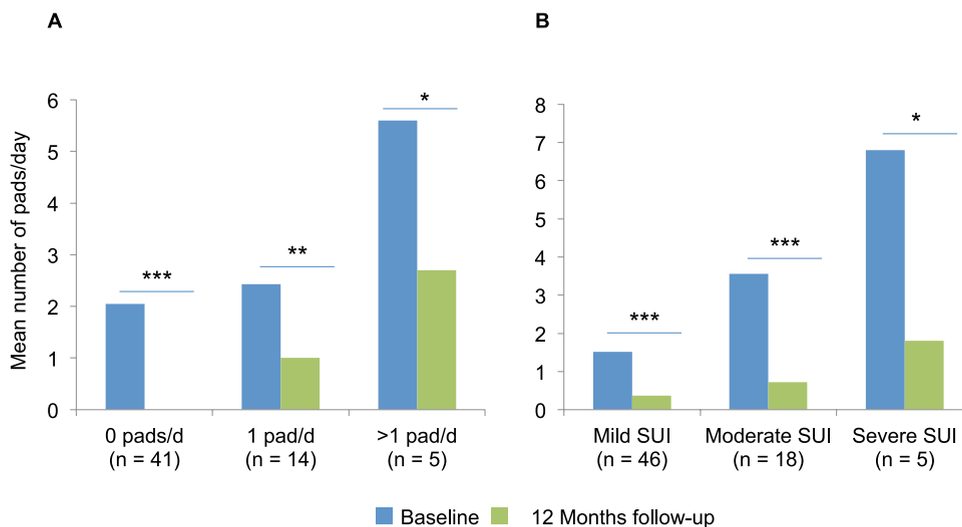
In the present prospective, multicenter, interventional, open-label, nonrandomized, Phase IV study, data were obtained at inclusion, just after sling implantation, and 1, 3, 6, and 12 months after implantation. Longer patient follow-up could be decided by each center. The present study, the HOMme INContinence study, was approved by the local research ethics committee (Comité Consultatif de Protection des Personnes se prêtant à des Recherches Biomédicales, Haute-Normandie, May 4, 2006, extended to increase the number of patients and centers on November 8, 2007) and conducted in compliance with good clinical practice guidelines and the Declaration of Helsinki. All patients provided written informed consent. The study was registered at ClinicalTrials.gov (trial registration number NCT00442078).

### Implantation Procedure

The sling was attached at each end to a clip to connect it to a Hemet or helical needle, according to surgeon preference. All surgeons were experienced in transobturator sling procedures. Implantation was performed with the patient under spinal or general anesthesia, and a 16F Foley urethral catheter was inserted. The patient was placed in the lithotomy position, and a 6-cm median vertical perineal incision below the inferior border of the pubic symphysis was performed to expose the bulbospongiosus muscle. Next, the perineal aponeurosis at the top of the triangular space was delimited laterally by each ischio-cavernosus muscle and medial to the bulbospongiosus. A short 2-mm incision through the pelvic fascia afforded access to the obturator muscle just under the ischiopubic ramus bone. A stab incision was made at the top of the thigh, 4 cm from the median line and 4 cm below the major adductor longus muscle. The transobturator puncture was preferentially outside-inside using a Hemet needle. The endpoint of the puncture was the opening of the pelvic fascia. After sling attachment to the needle, pulling back the needle implanted the 2 arms of the sling in the same passage. The same procedure was repeated on the other side. The sling was sutured to the bulbospongiosus muscle with nonabsorbable sutures and then pulled firmly from each side to obtain a 2-mm visible mark on the bulbospongiosus muscle. The perineal body was not dissected. However, in the case of rolling of the inferior edge of the sling on the bulb, the bulb was dissected just enough to place it under the sling and then sutured to the sling. No retrograde urethral pressure adjustment was performed. The incision was closed without drainage, and the urethral catheter was left indwelling for 2 days.

### Patient Assessments

All patients completed their continence status and 2 validated questionnaires to evaluate urinary function and bother at baseline and 1, 3, 6, and 12 months after implantation. The urinary function short form (Prostate Cancer Index, University of California, Los Angeles) consists of 4 questions on urinary function (range 0-100) and 1 on urinary bother (range 0-100, with a low score indicating a worse outcome).<sup>8</sup> The second questionnaire, the urinary incontinence short form (ICI Modular Questionnaire), has a score range of 0-21, with a low score indicating mild incontinence.<sup>7</sup> A short-term pad test was assessed at



**Figure 2.** Mean number of pads used daily at baseline and at 12 months of follow-up according to **(A)** continence status at 12 months in improved patients and **(B)** severity of SUI at baseline (n = 69). Mild SUI, 1-2 pads/d; moderate SUI, 3-5 pads/d; and severe SUI, >5 pads/d. \* $P \leq .05$ , \*\* $P \leq .01$ , \*\*\* $P \leq .001$ .

baseline and 3 and 12 months after implantation. The test measured the weight of the pads after bladder filling with 200 mL saline and after a short set of exercises of leakage provocation, as recommended by the ICI guidelines.<sup>9,10</sup> In addition, postoperatively, at 1, 3, 6, and 12 months, the patients indicated their satisfaction with the intervention and their new health status (1 question for each item, with 4 answer options, from 'not satisfied at all' to 'very satisfied') and evaluated the perineal pain (visual analog scale; range 0-10). The maximal flow rate and post-void residual (PVR) urine volume, low stream, urinalysis, and adverse events were also assessed at these points.

The primary endpoint evaluated was the number of pads used at 12 months of follow-up. The secondary efficacy endpoints included changes in the number of pads used, a change in the pad test outcome, improvements in continence and urinary bother scores, and changes in the satisfaction with the intervention and the new health status.

### Statistical Analysis

The statistical analysis included the efficacy and tolerability results for those patients with 12 months of follow-up. The surgical and tolerability results were descriptively reported per event. A comparison of between-group baseline and efficacy outcomes was performed using a Kruskal-Wallis test. Within-group comparisons were analyzed using the Wilcoxon matched-pairs signed-ranks test. Secondary analyses of the number of pads used within 24 hours were also performed, taking into account the SUI level at baseline (i.e., mild SUI [1-2 pads/d], moderate SUI [3-5 pads/d], and severe SUI (>5 pads/d)).<sup>11</sup>  $P < .05$  was considered statistically significant for all comparisons. The statistical analyses were performed using SAS software (SAS Institute, Cary, NC).

## RESULTS

A total of 122 patients were included from 30 centers in France from March 2007 to June 2009. The duration of follow-up was 12 months for 103 patients (84.4%), and

19 patients were lost to follow-up, 8 (6.6%) at 6 months, 6 (4.9%) at 3 months and 5 (4.1%) at 1 month. The additional results reported included only those patients with 12 months of follow-up. Of the 122 patients, 72.9% had undergone open radical prostatectomy, 22.2% laparoscopic radical prostatectomy, and 5.1% transurethral prostate resection. The mean interval from previous prostate surgery was 41.5 months, and 94.8% had undergone prostate surgery  $\geq 12$  months before sling implantation. The mean patient age was  $69.4 \pm 6.1$  years. The rate of urinary infection in the previous 3 months before study inclusion was 4.9%. Implantation was performed with the patient under general anesthesia in 83.7% of the patients and locoregional anesthesia was used in 13.3%. The mean intervention time was 36.3 minutes. In 75.7% of the procedures, an outside-inside transobturator puncture was used; in 24.3%, it was an inside-outside puncture. For 93.9% of the procedures, the surgeon indicated that the perineal dissection was easy to perform. Performing the left puncture, the right puncture, and passing the sling were indicated as easy in 89.9%, 94.9%, and 97.0% of the procedures, respectively.

### Efficacy

The number of pads used daily at baseline and at 12 months was available for 69 patients. At 12 months, 60 (87.0%) of the 69 patients had improvement in the number of pads used daily: 41 (59.4%), 14 (20.3%), and 5 (7.3%) patients reported 0, 1, and >1 pad/d, respectively. However, 9 patients (13.0%) had no improvement, with 7 (10.1%) and 2 (2.9%) reporting 1 and >1 pad/d, respectively. Pad use at 12 months had decreased significantly compared with that at baseline (mean 0.6 vs 2.4,  $P \leq .001$ ; n = 69). In patients with improved continence status, the mean decrease in daily pad use at 12 months was 2.1, 2.1, and 3.0 in patients with 0, 1, and

**Table 1.** Improvement in self-reported incontinence rate/bother (questionnaires) at 12 months of follow-up

Variable	Baseline	Endpoint	Difference	P Value
UCLA-PCI-specific HRQOL (n = 101)				≤.0001
Leakage	1.0 ± 1.1	37.2 ± 7.9	35.3 ± 8.1	
Urinary control	41.5 ± 3.9	71.0 ± 4.4	29.3 ± 5.6	
Pads used daily	36.1 ± 4.6	80.4 ± 6.0	43.1 ± 6.9	
Problem of urine dripping	14.9 ± 3.8	72.3 ± 5.9	56.9 ± 6.6	
Problem of urine function	19.6 ± 5.0	75.0 ± 5.9	55.0 ± 6.8	
ICIQ-UI (n = 102)				≤.0001
Frequency of leakage	3.9 ± 0.1	2.0 ± 0.3	-1.9 ± 0.3	
Quantity of leakage	3.3 ± 0.2	1.5 ± 0.2	-1.8 ± 0.3	
Bother	6.6 ± 0.4	2.1 ± 0.4	-4.5 ± 0.5	

UCLA-PCI, University of California, Los Angeles, Prostate Cancer Index; HRQOL, health-related quality of life; ICIQ-UI, International Consultation on Incontinence modular Questionnaire—Urinary Incontinence.

Data presented as mean improvement ± standard error.

UCLA-PCI: question 1, leakage (answer, 'everyday', 'once a week', 'less than once a week', 'not at all' [score 0, 33, 66, 100, respectively]); question 2, urinary control (answer, 'no control', 'frequent dribbling', 'occasional dribbling', 'total control' [score 0, 33, 66, 100, respectively]); question 3, pads used daily (answer, '3 pads/d', '1-2 pads/d', 'no pads/d' [score 0, 50, 100, respectively]); question 4, how big a problem is dripping urine or wetting pants (answer, 'no problem', 'very small problem', 'small problem', 'moderate problem', 'big problem' [score 100, 75, 50, 25, 0, respectively]); question 5, how big a problem is your urine function (answer, 'no problem', 'very small problem', 'small problem', 'moderate problem', 'big problem' [score 100, 75, 50, 25, 0, respectively]).

ICIQ-UI: question 1, frequency of leakage (answer, 'never', 'once a week maximum', '2-3 times a week', 'about once daily', 'several times daily', 'always' [score 0-5]); question 2, quantity of urinary leakage (answer, 'none', 'small', 'median', 'important quantity' [score 0-6]); question 3, bother about leakage (answer, 'not at all' to 'important bother' [score 0-10]).

>1 pad/24 d, respectively (Fig. 2A). Comparing the patients with different SUI levels at baseline, the decrease in the number of pads used daily was statistically significant compared with at baseline for both patients with mild and moderate SUI and those with severe SUI (Fig. 2B). The absolute difference in pad use between baseline and follow-up tended to be larger in patients with severe SUI (Fig. 2B). The mean daily pad use at 12 months of follow-up was 0.4, 0.7, and 1.8 for mild, moderate, and severe SUI, respectively. The outcomes in pad weight of the short-term pad test were only available for 36 patients, because all the clinicians did not report the pad test results. The results were significantly improved at 12 months compared with at baseline (mean 11.3 g vs 105.1 g;  $P \leq .01$ ).

At 12 months of follow-up, all symptoms and bother scores, as assessed by the urinary function short form (Prostate Cancer Index, University of California, Los Angeles, CA) and the urinary incontinence short form (ICI Modular Questionnaire), were significantly improved statistically compared with at baseline (Table 1). The functional scores improved from 'everyday' or 'frequent problem' to 'once a week' or 'occasional'. The bother scores improved from 'big' or 'moderate problem' to 'very small problem'. In addition, 91.2% of the patients were 'satisfied' or 'very satisfied' with the treatment and 87.8% with their new health status from the post-operative period to the end of follow-up at 12 months. The satisfaction was stable over time.

### Perioperative Complications and Tolerability

No complications, such as bladder perforation, intraoperative bleeding (>200 mL), or nerve, bowel, or vascular injury occurred during the intervention, except for wounding of the corpus cavernosum (4.0% of the patients).

Micturition at removal of the catheter 48 hours after surgery occurred in 98.9% of the patients. Hematoma and wound infection were very rare, and the mean perineal pain visual analog scale score were low (Table 2). Of the patients, 97.3%-100% were free of urinary tract infection at the different follow-up visits, and 96.5%-100.0% of the patients had not experienced urinary tract infection in the month before the visits. The maximal urinary flow rates were similar before and after surgery. The PVR urine volume was increased after surgery and was normal at 30 days; a low stream was reported by some patients (Table 2). Acute urinary retention (AUR) did not occur.

### COMMENT

This is the first study presenting prospective data on the efficacy and tolerability of the 4-arm large I-STOP TOMS male sling. Prospective studies on sling implantation for post-prostatectomy SUI in series including 100 patients are exceptional.

In the present study, 87.0% of the patients reported improved continence, with 59.4% completely dry at 12 months of follow-up; 13% reported no improvement but the absence of worsening. The quality of life had improved significantly at early follow-up, and the improvements were maintained through the follow-up period. The patients were highly satisfied with the intervention and with their new health status. Tolerability was high. Some patients experienced a low stream but in the absence of an increased PVR urine volume and without additional risks.

One limitation of the present study was the length of follow-up, which was 12 months. Long-term data on the efficacy and tolerability are needed. In addition, a short pad test was used instead of a 24-hour pad test<sup>9</sup>; however, the test was performed according to the recommenda-

**Table 2.** Tolerability of the sling procedure

Variable	Baseline	Immediate Postoperatively	Follow-Up Visit (mo)			
			1	3	6	12
Hematoma (n)	—	8/92	2/96	0/96	0/79	0/76
Wound infection (n)	—	0/97	2/96	0/89	0/79	0/76
Current UTI (n)	0/103	—	2/85	2/89	2/73	0/69
UTI in previous 30 days (n)	—	—	3/85	3/89	2/73	1/69
Difficulty voiding/low stream (n)	5/101	—	22/97	16/87	8/74	9/69
Perineal pain (VAS)						
Patients (n)	99	94	102	96	82	102
Mean score $\pm$ SD	0.4 $\pm$ 1.2	2.7 $\pm$ 1.9	1.2 $\pm$ 1.8	0.4 $\pm$ 1.0	0.2 $\pm$ 0.8	0.1 $\pm$ 0.4
Qmax (mL/s)						
Patients (n)	90	60	74	74	60	56
Mean $\pm$ SD	23.4 $\pm$ 10.7	20.1 $\pm$ 8.7	19.4 $\pm$ 9.9	21.4 $\pm$ 12.4	20.8 $\pm$ 9.7	21.5 $\pm$ 8.9
PVR urine volume (mL)						
Patients (n)	84	63	71	73	55	53
Mean $\pm$ SD	6.0 $\pm$ 14.7	14.0 $\pm$ 28.3	11.7 $\pm$ 21.1	10.2 $\pm$ 24.3	11.0 $\pm$ 19.5	10.5 $\pm$ 21.5

UTI, urinary tract infection; VAS, visual analog scale; Qmax, maximal flow rate; PVR, post-void residual.

tions of the ICI.<sup>10</sup> The combination of regular follow-up visits with patient-reported outcomes using validated patient-completed questionnaires is a strong point of the present study.<sup>7,8</sup> We observed that safety reporting was lower compared with efficacy reporting with longer follow-up. We believe this is a common situation in the case of good tolerability.

Male slings have been used for a few decades; however, the slings developed in recent years are not comparable to those from earlier years. The study inclusion criteria for patients and the outcomes definitions, such as the success rate have varied highly, making it difficult to compare trials quantitatively.<sup>4</sup> The published success rates of studies, with a mean follow-up of 6-24 months using male slings of all types, have ranged from 38% to 76%, depending on the outcome measures used, with the best results achieved in patients with low to moderate incontinence who had not undergone radiotherapy.<sup>1,12</sup>

The most common complications are infection, erosion, and urinary retention. If the sling is placed with insufficient pressure, incontinence will remain. However, if the pressure on the sling is too high, it could result in obstruction, leading to AUR, although other factors, such as the sling location, could also be involved. This could explain the significant proportion of patients developing AUR in some studies. In our study, none of the patients had experienced AUR at  $\leq$ 1 year after implantation. Indicative of obstruction is a decrease in the maximal urinary flow rate and/or an increased PVR urine volume. Both outcomes remained similar to the baseline measurements in the present trial.

The occurrence of adverse events can also correspond with the fixation technique used during implantation. The retropubic fixation procedure has raised concerns regarding the risk of bladder perforation or erosion.<sup>13</sup> Using the perineal route, a sling can be fixed by bone screws to the pubic bone or using the transobturator technique. The latter approach gained wide popularity for the treatment of SUI in women<sup>14</sup> and was first described for male sling implantation

in 2005.<sup>15</sup> Generally, the transobturator technique is perceived as being easier to perform and reproducible, with a low rate of complications.<sup>6,16</sup> Data on 4 transobturator male slings are available.<sup>6,11,13,17</sup>

The male perineal sling InVance<sup>TM</sup> is implanted in the same position as the I-STOP TOMS; however, the former is anchored to the pubic bone with bone screws. InVance<sup>TM</sup> study with 50 patients showed that about half of the treated patients reached complete continence; however, an AUR rate of 12% and persistent perineal pain rate of 12% were reported at a median follow-up of 6 months.<sup>18</sup> In a representative study of 62 patients with the InVance<sup>TM</sup> sling, the infection rate was 4.8% and the urinary retention rate was 3.2%.<sup>19</sup>

Adjustable slings typically require reintervention (38.6% for Argus, >80% for the Male Reemex System), and complications have been relatively common. Sling removal because of urethral erosion or infection was described in 5.9%-15.8% of patients.<sup>20,21</sup>

The bulbomembranous AdVance<sup>TM</sup> sling is positioned deeper than the I-STOP TOMS and more posterior to reposition the membranous urethra. However, the urethral wall is thin in this location, and the location of the sling is close to the sphincter. Comparing our results with those from a study of 124 patients with mild to moderate SUI after radical prostatectomy treated with the transobturator AdVance<sup>TM</sup> sling, the efficacy rates were comparable, with 59.4% completely dry at 12 months of follow-up in our series compared with 55.7% at 12 months in their study.<sup>22</sup> However, the postoperative complication rate, in particular AUR, was greater than in our series (12.9% with the AdVance<sup>TM</sup> sling).<sup>22</sup> A recently published AdVance<sup>TM</sup> series reported 2.8%-5.7% of patients with posturinary retention (mean follow-up of 35 weeks vs follow-up of 12 months).<sup>23-25</sup> In addition, the mean interval needed to implant the AdVance<sup>TM</sup> sling was 97 minutes, longer than that required in our series (36.3 minutes).<sup>16</sup>

Overall, the results of the present study have shown that the I-STOP TOMS achieves adequate suburethral

positioning to obtain good continence in most patients without causing obstruction or other adverse events.

## CONCLUSIONS

The I-STOP TOMS is an appropriate choice for nonradiated patients with mild to moderate post-prostatectomy SUI. At  $\leq 12$  months, most patients were continent or had improved continence. The operation time was short, and the intervention was well tolerated, with minimal postimplantation pain and few cases of infection.

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## APPENDIX

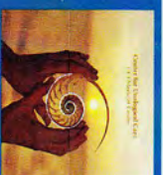
The HOMme INContinence study group consisted of the following participants: I. Bah-Clozel, Clinique Pasteur, Guilherand Granges; G. Bochereau, Clinique Saint-Augustin, Nantes; B. Chavrier, Clinique de la Sauvegarde, Lyon; P. Coeurdacier, Polyclinique Sévigné, Cesson Sévigné; F. Collet, Clinique Trénel, Sainte-Colombe; E. David, Polyclinique du Grand Sud, Nîmes; A. Delannoy, Centre Hospitalier Avranches-Granville, Avranches; O. Delbos, Clinique du Millénaire, Montpellier; L. Drelon, Clinique des 2 Caps, Coquelles; D. Dupuy, Clinique Ambroise Paré, Toulouse; R. Faye, Clinique de l'Anjou, Angers; J. Grall, Clinique de Fontaine, Dijon; E. Gremmo, Polyclinique Synergia, Carpentras; P. Grise, Hôpital Charles Nicolle, Rouen; O. Lan, Polyclinique Synergia, Carpentras; B. Le Portz, Clinique Océane, Vannes; F. Levigne, Centre de l'Hospitalisation Privée de la Loire, Saint-Étienne; J. Lienhart, Clinique Trénel, Sainte-Colombe; P. Lille, Clinique Saint Odilon, Moulins; A. Manunta, CHU Pontchaillou, Rennes; B. Marc, Polyclinique Saint-Privat, Boujan-sur-Libron; O. Marecaux, Clinique Sainte-Catherine, Sainte-Catherine-les-Arras; D. Mianne, Clinique de Provence, Orange; B. Njinou, Clinique des Ormeaux, Le Havre; C. Olivier, Polyclinique du Sidobre, Castres; P. Paulhac, Clinique des Emailleurs, Limoges; Y. Perraud, Centre de l'Hospitalisation Privée de la Loire, Saint Étienne; O. Rousseau, Clinique du Cèdre, Bois-Guillaume; J. Sarkissian, Hôpital Jean Mermoz, Lyon; C. Saussine, Centre Hospitalier Universitaire-Hôpital Civil, Strasbourg; J. Vannier, Clinique Saint-Augustin, Tours; R. Vautherin, Clinique Trénel, Sainte-Colombe; and G. Ybert, Clinique Chirurgicale Marie Immaculée, Bourges.



# Transobturator Male Slings: Comparison of Two Meshes

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## PURPOSE

This study aims to determine if two different transobturator male slings (TOMS) have similar outcomes. Additionally, this study attempts to detect any areas of superiority of one sling type compared to the other.

## MATERIALS and METHODS

Forty patients who received TOMS were studied from November 2007-February 2011. Data analyzed included demographics, pre- and post-operative pad usage, urodynamic parameters, etiology of SLI, presence of prior urethral insult, and adverse events. Patients were also asked to complete the patient global impression of improvement (PGI-I) at each follow up visit. Patients were evaluated for success (dry or  $\geq 50\%$  improvement in pad usage) and failure ( $< 50\%$  improvement or increased pad usage). Separately, PGI-I was evaluated for success (score of 1-2) and failure (score  $\geq 4$ ).

## Male Slings



Advance (AMS)



I-Stop (C.L. Medical)

Total patients (n=40)	Advance (n=20)	I-Stop (n=20)	P-Value
SLI (Mean & SD)	20 (5.13)	21 (5.13)	ns
Prevalence (Mean & SD)	3.9 (2.65)	3.2 (2.12)	ns
Age (Mean & SD)	90.9 (28.1)	111.3 (42.8)	ns
ADL (Mean & SD)	97.8 (20.4)	104.0 (28.8)	ns
IL (Mean & SD)	3.6 (1.9)	3.6 (1.1)	ns
UOP (Mean & SD)	2 (10.9%)	11 (54.9%)	0.002
Urethral disease	6 (30.0%)	8 (40.0%)	ns

BMI= Body Mass Index, ALP= Abdominal Leak  
 Post-Pressure, MUP= Maximal Urethral Pressure  
 FL= Functional Profile Length, DO= Detrusor  
 Overactivity, SC= Standard Deviation, IQR= Interquartile Range

## RESULTS

Twenty patients received AdvVance<sup>®</sup> male sling and 20 received I-Stop<sup>®</sup> male sling with mean follow up time of 14.9 ( $\pm 8.25$ ) months and 8.3 ( $\pm 6.92$ ) months respectively. Mean age was 70.7 years ( $\pm 8.09$ ). Detrusor overactivity in the I-Stop group was the only pre-operative parameter to demonstrate significance. Both groups demonstrated significantly decreased pad usage post-operatively. Successful outcomes defined by pad usage were 70% and 75% with complete dryness rates of 40% and 45% for AdvVance<sup>®</sup> and I-Stop<sup>®</sup> respectively. Successful outcomes defined by PGI-I were 55% and 70% for AdvVance<sup>®</sup> and I-Stop<sup>®</sup> respectively. All adverse events were managed conservatively with no long-term sequelae.

Table 2. Prostatic Procedures

Type of Procedure	Group A	Group I	P-Value
Open	10 (50%)	7 (35%)	ns
Laparoscopic	8 (40%)	6 (30%)	ns
Endoscopic	1 (5%)	6 (30%)	0.04
Radio/Brachy therapy	5 (25%)	3 (15%)	ns

Table 3. Adverse Events

Adverse Event	Group A	Group I
Urinary retention	15% (3/20)	25% (5/20)
Urinary tract infection	10% (2/20)	0% (0/20)
Pain (Scrotum/Perineum/Groin/Thigh)	20% (4/20)	20% (4/20)
Numbness (Scrotum/Perineum/Groin/Thigh)	15% (3/20)	5% (1/20)

## CONCLUSIONS

I-Stop<sup>®</sup> male sling showed equally successful outcome of complete dryness, significant decrease in pad usage, and patient satisfaction. It also consistently showed a trend toward superiority when used in difficult cases with urethral damage.

# The I-STOP® TOMS® transobturator male sling, a minimally invasive treatment of post-prostatectomy incontinence: continence improvement and tolerability

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## Abstract

The I-STOP TOMS transobturator male sling, a minimally invasive treatment of post-prostatectomy incontinence, continence improvement and tolerability. Philippe Grise<sup>a</sup>, Renaud Vautherin<sup>b</sup>, Bertin Njinou<sup>c</sup>, Ghislain Bochereau<sup>d</sup>, Jean Lienard<sup>e</sup>, Christian Saussine<sup>f</sup>. Corresponding author: Department of Urology, Charles Nicolle Rouen University Hospital, rue de Gemme 76031 Rouen, France. Tel: 2 32 866173. Fax: 2 32 866205. E-mail address: philippe.grise@univrouen.fr

**INTRODUCTION AND OBJECTIVES:** Post-prostatectomy incontinence (PPI) is a bothersome problem, affecting the quality of life of patients. Minimally invasive techniques are becoming the standard of care for mild to moderate PPI. The I-STOP TOMS transobturator male sling is a four arms polypropylene macroporous non-extensible sling. For this type of sling, improvement of continence, quality of life and tolerability have never been evaluated in patients with mild to moderate PPI.

**METHODS:** Between 2007 and 2009, we set up a prospective multicenter trial to evaluate the improvement of continence, quality of life and tolerability of the I-STOP TOMS in patients with mild to moderate PPI. Patients with bladder outlet obstruction, bladder overactivity or low compliance, or with a history of radiation were excluded. The transobturator puncture was done outside-inside or inside-outside according to the surgeon's preference. The sling was applied to the bulbar urethra with a pressure on the corpus spongiosum. The primary endpoint was complete continence (zero pad/day) or improved continence (0 to 1 pad/day) at day 90 and day 360. Secondary endpoints were continence scores, quality of life and patient satisfaction up to 12 months. To evaluate tolerability Visual Analog pain scale (VAS), urinary infection, urine flow rate and post void residual (PVR) were assessed.

**RESULTS:** In total, 122 patients from 30 centers with a mean age of 69 (54-88) years were included. After surgery 43% (52/122) patients were completely continent at 90 and 360 days, respectively. 58% (71/122) and 86% (105/122) had improved continence at 90 and 360 days, respectively. Quality of life had improved significantly at all assessed time points. Patients were highly satisfied with the intervention and with their new health status. Only two patients experienced urinary tract infection at day 30, without recurrence. Mean VAS pain scores were low (0.3, 1 and 0 before, post-surgery, at day 30 and at day 90, respectively). Mean urine flow rates were similar before, post and 30 days after surgery. PVR was increased post-surgery and was normal at 30 days.

**CONCLUSIONS:** The I-STOP TOMS is an appropriate choice for patients with mild to moderate PPI. Up to 12 months, most patients were completely continent or had improved continence. The intervention was well tolerated with low morbidity.

Source of Funding: CL Medical for assistance to data management

## INTRODUCTION AND OBJECTIVE

Post-prostatectomy incontinence (PPI) is a bothersome problem, affecting the quality of life of patients. Minimally invasive techniques are becoming the standard of care for mild to moderate PPI.

The I-STOP® TOMS® transobturator large bulbar male sling was evaluated prospectively in patients with PPI.



TOMS® sling is a large four-arms, four corners, polypropylene macroporous non-extensible sling.

TOMS® sling is applied with tension on the bulbar urethra covered with the bulbospongiosum muscle, below the pubis, without section of the perineal body.

On each side, the arms are connected to the same needle. They are conducted through the trans-obturator foramen with a hemet or helical needle.

## METHODS

Between 2007 and 2009, a prospective multicenter trial evaluated the result on incontinence, quality of life and tolerability of the I-STOP TOMS in patients with mild to moderate PPI. Patients with bladder outlet obstruction, bladder overactivity or low compliance, or with a history of radiation were excluded. The transobturator puncture was done outside-inside or inside-outside according to the surgeon's preference. The sling was applied to the bulbar urethra with a pressure on the corpus spongiosum until to have a visible mark on the corpus spongiosum. No retrograde pressure adjustment was realized.

The primary endpoint was complete continence (no pad/day) or improved continence (0 to 1 pad/day) at day 90 and day 360. Secondary endpoints were continence scores, quality of life (assessed by UCLA-PCI urinary domain and ICIQ-UI) and patient satisfaction up to 12 months. To evaluate tolerability, Visual Analog pain scale (VAS), urinary infection, urine flow rate and post void residual (PVR) were assessed.

## RESULTS

122 patients from 30 centers with a mean age of 69 (54-88) years were included. The surgery was easy to perform in all the cases, operative time : 23 minutes. No per-operative complications were reported.

Complete continent patients at 90 and 360 days were: 43% (36/84) and 58% (71/76).

Improved continence patients at 90 and 360 days were: 86% (72/84) and 93% (71/76).

Quality of life was improved significantly at all assessed time points: 90% of the patients were satisfied or very satisfied with the intervention, and 85% with their new health status.

Only two patients experienced urinary tract infection at day 30, no recurrence..

Mean VAS pain scores were low the day after surgery: 2.7, 30 days: 1.2, 90 days: 0.4

Mean urine flow rates were similar before: 23.2mL/s, 90 days after surgery: 20.5, 360 days: 21.5

Mean PVR was non significant after catheter removal: 16.6 mL, 90 days: 12.7, 360 days: 10.5

No urethral erosion were reported

## DISCUSSION

Different male slings were described but some complications were observed i.e. perineal pain, infection or erosion. Good tolerance of the polypropylene mesh is now well demonstrated.

Trans-obturator route limits the risk of bladder puncture, and avoid the pain and osteitis with bone screw procedure.

The TOMS sling is not applied on membranous urethra in order to limit the risk of erosion, but applied on the bulbar portion where the urethra wall is larger and covered with spongiosum tissue.

The four arms design allows a tension equally applied on the four corners of the mesh, avoiding a string effect.

Although the mechanism is not sufficiently documented, a support effect associated with a compressive is proposed. No retention was observed but a minor obstructive effect may be possible although no pressure-flow study was conducted.

## CONCLUSION

The I-STOP TOMS® is an appropriate choice for patients with mild to moderate PPI for patients demanding of improvement on their day life comfort.

The large 4 corners TOMS bulbar male sling was easy to insert and well tolerated.

Up to 12 month, most of the patients were completely continent or had improved continence.

## TRANS-OBTURATOR MALE SLING TOMS FOR THE TREATMENT OF URINARY STRESS INCONTINENCE IN MEN.

### Aims of study

Despite improvement in the surgical technique of radical prostatectomy, stress urinary incontinence (SUI) remains a problem that affects the quality of life of many patients. The prevalence of post-prostatectomy depends on the definition of incontinence. In order to minimize surgical morbidity and cost of the artificial sphincter, sling procedures were described with or without bone anchor. With the experience of the female trans-obturator polypropylene sling, a new trans-obturator male sling (TOMS) was developed, the first results with minimal one year experience are reported.

### Material and methods

A prospective multicenter clinical study was conducted on male patients suffering from post prostatectomy incontinence and failure of physiotherapy. Patients with minor or moderate SUI were included with minimal 12 month follow-up. Exclusion criteria were pre or post-operative radiation, less than one year interval from surgery, bladder outlet obstruction, bladder overactivity or hypocompliance. Pre operative assessments included clinical study questionnaire, urodynamics (uretrocystometry, uroflowmetry, bladder residual), a pad test short form, ICIQ and SF36 questionnaire, Visual analogic pain scale (VAS). Post-operative evaluations were at 1, 6, 12, 18 and 24 months using the same evaluation except uretrocystomanometry. Pre and post-operative hazards were recorded on a case report form. The sling was made of polypropylene macroporous non extensible, 1 cm large, connected at each end to a needle attachment device. Hemet or helicoidal needle was used according to surgeon preference. The surgical technique was done under spinal or general anaesthesia, a 16 F Foley urethral catheter was inserted then a median perineal incision exposed the angle between each corpus cavernosum and corpus spongiosum. The trans-obturator puncture was outside-inside similar to the female procedure. The sling was applied to the urethra then pulled firmly from each side until to have a visible mark on the corpus spongiosum. No retrograde pressure adjustment was realized. The urethral catheter was left for 1day. Before hospital discharge, a uroflowmetry, a residual, and a pelvic pain evaluation on VAS were obtained.

### Results

A total of 14 patients, age 70.8 (57-87) years old underwent surgery. Mean follow-up was 18 (12-23) months. The surgery was easy to perform in all the cases. No per-operative complications were reported, no significant intra-operative bleeding (>200ml) occurred or nerve, bowel or vascular injury. One patient experienced temporary urinary retention, one a low stream, and residual was always less than 100 ml. Mean pad use modified from 2.4 to 0.9, and 50% of them used no pad. Pad test mean weigh decreased from 62g to 8 g. The SF36 score improved from a median of 126 (57-325) to 317 (92-500), the Mann-Whitney test was significant with  $p < 0.001$ . The ICI-Q score improved from 13 (6-16) to 7 (0-15). The pain mean value was 2.5 post-operatively, 0.9 at one month and 0.1 at one year.

### Interpretation of results

Patients with minimal and moderate incontinence are demanding of improvement and even only one pad a day affects their quality of life. Many new minimal invasive techniques (injectable biomaterials, balloons, sling with bone screw attachment or with retropubic puncture) have been proposed for managing SUI in males but they had adverse side effects or poor results. The artificial sphincter remains the gold standard technique, but the cost and the erosion or infection rate limit the indication for severe incontinence. The transobturator sling has no screw fixation and minimal well tolerated polypropylene biomaterial, this may explain the good tolerance and minimal pain. This route is an alternative to retropubic puncture that limit the risk of bladder perforation. This short series demonstrate the feasibility, the good tolerance, and marked improvement in continence.

### Concluding message

The trans-obturator male sling (TOMS) is a new attractive surgical procedure for moderate or minor post-prostatectomy SUI. The implanted biomaterial is non mechanical, easy to insert and well tolerated. Most of the patients are improved or continent with one year follow-up. In properly informed patients, this sling may afford an improvement in their quality of life.

<b>Specify source of funding or grant</b>	<b>no funding or grant</b>
<b>Is this a clinical trial?</b>	<b>Yes</b>
<b>Is this study registered in a public clinical trials registry?</b>	<b>Yes</b>
<b>Specify Name of Public Registry, Registration Number</b>	<b>Haute autorité de Santé</b>
<b>What were the subjects in the study?</b>	<b>HUMAN</b>
<b>Was this study approved by an ethics committee?</b>	<b>Yes</b>
<b>Specify Name of Ethics Committee</b>	<b>CCPPRB Rouen France</b>
<b>Was the Declaration of Helsinki followed?</b>	<b>Yes</b>
<b>Was informed consent obtained from the patients?</b>	<b>Yes</b>