Overactive bladder syndrome (OAB) is a common condition affecting up to 43% of women in the United States [1]. For many patients, symptoms of OAB include a loss of self-esteem, restriction of activities and psychological distress [2]. The American Urological Association (AUA) and the Society of Urodynamicists, Female Pelvic Medicine and Urogynecology Reconstruction (SUFU) recommends patients try medication and behavioral modification before proceeding to third line OAB treatments which includes percutaneous tibial nerve stimulation (PTNS), OnabotulinumtoxinA (Botox®), and sacral neuromodulation (SNM) [3].

Behavioral therapy includes fluid management (limiting caffeine and alcohol), bladder retraining (trying to defer the urge), and pelvic floor physical therapy. With these changes, we can see up to 50% symptom improvement. The next step is OAB medications. There are currently 8 OAB medications available in the U.S. – six anticholinergic, and two β3 agonist. β3 agonist medications are newer overall and cause relaxation of the detrusor muscle, in contrast to anticholinergic medications that block detrusor contraction. While common side effects of anticholinergic medications include dry mouth, dry eyes, and constipation in an average of 20% of patients, β3 agonist medications do not cause these side effects [4]. β3 agonist medications can increase blood pressures in some patients, so they should be used with caution in patients with uncontrolled hypertension. Overall, medications work 50% of the time to improve symptoms. We often have to try behavioral changes and at least two medications prior to insurance approval of third-line therapies.

OAB medications are primarily anticholinergic with side effects including dry mouth, blurred vision, and constipation and have been associated with cognitive changes in elderly patients [5]. Up to 40% of patients stop OAB medication therapy due to side effects, with 66.7% stopping medication therapy within two years [6]. Common OAB medications, such as oxybutynin, have reported efficacy rates of 32-60%, making OAB medication therapy ineffective for many patients [7]. Once eligible for third-line treatments, many patients persist on OAB medication therapy despite poor efficacy. Poor efficacy, decreased satisfaction, decreased patient use of OAB medications over time, and side effects make medication therapy not suitable for everyone.

Posterior tibial nerve stimulation (PTNS) is an electrical stimulation of the posterior tibial nerve. The nerve can be accessed at a point near the ankle. A thin needle is placed through the skin to the level of the nerve. The needle is then connected to a battery that electrically stimulates the tibial nerve. The initial protocol developed by Peters [8] that we use includes tibial nerve stimulation for 30 minutes one time a week for 12 weeks. Peters found in 2010, when compared to a sham procedure, PTNS improved both subjective and objective symptoms in 55% of patients [8]. When compared to tolterodine (OAB medication), both improved symptoms in 50% of patients. The greatest symptom improvement was in patients undergoing PTNS while continuing medication treatment [9]. If the 12-week treatment is successful, patients can opt to continue monthly maintenance visits. While there are frequent clinic visits, there is little to no risk, making this option palatable to many patients.

Injection of OnabotulinumtoxinA (Botox®) into the detrusor muscle with cystoscopy in the office setting works by paralyzing part of the detrusor muscle. Additionally, it also decreases sensory afferent signals from the bladder to help with urgency and pain symptoms. It was first approved in the U.S. by the FDA in 2011 for neurogenic patients and expanded in 2013 to include non-neurologic OAB. We perform this in the office after injecting local anesthetic into the bladder, starting at 100 units. On average improved symptoms last for 9 months, or 14 months using a higher dose. When compared to OAB medications, both OAB medications and OnabotulinumtoxinA reduced daily urge incontinence episodes by 3.4 episodes per day, but OnabotulinumtoxinA was more likely to result in no incontinence at all [10]. The main concern for patients is that when it works too well, 1-2% of patients have temporary urinary retention and need to perform self-catheterization until it wears off. Patients like OnabotulinumtoxinA because it is a short office procedure, works well, and allows them to stop OA medications.

Sacroneuromodulation (SNM) is a sacral nerve stimulator that involves placing a wire lead with four electrodes into the spinal column at the S3 level to stimulate the sacral nerve roots and an internal battery. It can be used to treat overactive bladder, urinary retention, and fecal incontinence. For overactive bladder, 60% of patients had at least a 50% improvement in symptoms at 6 months after placement, and 50% at 2 years after placement. A subset of patients (20%) had >75% improvement at 6 months.
and 2 years [11]. For urinary retention, 80% of patients had >50% improvement at 2 years after placement with increased voids and low residuals [12]. For fecal incontinence, 88% of patients had >50% improvement in their symptoms at 3 years after implant with a significant improvement in quality of life [13]. I often suggest SNM for patients with dual urinary and fecal incontinence. For fecal incontinence and urinary retention, we have a few other SNM treatment options to help patients improve their quality of life.

Patients can now opt for a 5-year non-rechargeable SNM or a 15-year rechargeable device. MRI compatibility and the longer life of a rechargeable system allows more patients to be eligible and for it to last longer prior to battery replacement. We place the SNM device in two stages. In the first stage, the sacral lead can be placed in the office under local anesthesia or in the operating room. If the lead is placed in the office, it is attached to an external battery and the patient tests the effect for one week. If it is placed in the operating room, the lead can be tested for two weeks. During the second stage, if at the end of the testing period the patient has more than 50% improvement in their symptoms, we implant the battery into the patient’s buttock at a place they can access. The patient controls the program with a smartphone-like device that connects to the battery via Bluetooth technology. Both the patient and their provider can adjust the settings over time to achieve the optimal stimulation.

When medications aren’t enough

<table>
<thead>
<tr>
<th>Treatment</th>
<th>Helps patients with</th>
<th>Pros</th>
<th>Cons</th>
</tr>
</thead>
<tbody>
<tr>
<td>PTNS</td>
<td>Synergistic with OAB meds</td>
<td>Low risk, External procedure</td>
<td>Weekly office visits x 12, then monthly</td>
</tr>
<tr>
<td>Botox*</td>
<td>Urge incontinence, Neurogenic bladder</td>
<td>Can stop OAB meds, Short office procedure</td>
<td>If over-corrected, may need temporary self-catheterization, Lasts 9-14 months</td>
</tr>
<tr>
<td>SNM</td>
<td>Urinary retention, Combined urinary &amp; fecal incontinence</td>
<td>New 15-year rechargeable device, Adjust settings over time</td>
<td>May need to go to OR for placement</td>
</tr>
</tbody>
</table>

Summary

Overactive bladder is a common problem that affects many patients. There are treatments available that can help patients limit bathroom trips, stop using incontinence products, and greatly improve their social and overall quality of life. We have four fellowship-trained, board-certified urogynecologists and one nurse practitioner who see patients at our main campus and at several off-site locations.

REFERENCES

UROGYNECOLOGY
AND PELVIC
RECONSTRUCTIVE
SURGERY

Referrals and consultations
Call 773-702-6118
Email womenshealth@uchospitals.edu

LOCATIONS

Hyde Park
Duchossois Center for
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5758 S. Maryland Ave.
Third Floor
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Hinsdale
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Salt Creek Suite 106
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Orland Park
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