Advances in the role of sacral nerve neuromodulation in lower urinary tract symptoms

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Abstract Sacral neuromodulation has been developed to treat chronic lower urinary tract symptoms, resistant to classical conservative therapy. The suspected mechanisms of action include afferent stimulation of the central nervous system and modulation of activity at the level of the brain. Typical neuromodulation is indicated both in overactivity and in underactivity of the lower urinary tract. In the majority of patients, a unilateral electrode in a sacral foramen and connected to a pulse generator is sufficient to achieve significant clinical results also on long term. In recent years, other urological indications have been explored.

Keywords Lower urinary tract symptoms (LUTS) · Sacral neuromodulation · Bladder pacemaker

Introduction

Chronic types of lower urinary tract dysfunction, including urge incontinence, urgency–frequency (both included in the overactive bladder syndrome or OAB together eventually with nocturia) and non-obstructive urinary retention, still present a therapeutic challenge. Most patients are initially treated with conservative therapies including bladder retraining, pelvic floor exercises, and biofeedback. In the majority of patients, this standard regimen is supported with pharmacological therapy (anticholinergics). However, approximately 40% either do not achieve an acceptable level of therapeutic benefit or remain completely refractory to treatment. Alternative surgical procedures such as bladder transsection, transvesical phenol injection of the pelvic plexus, augmentation cystoplasty, and even urinary diversion have been advocated for these chronic conditions. However, these procedures have variable efficacy and have been associated with significant morbidity and risk. Therefore, research into the use of electrical current for the treatment of lower urinary tract dysfunction has been initiated.

In 1878, Saxtorph reported intravesical electrostimulation in patients with a contractile bladder and complete urinary retention [1]. He inserted a special catheter with a metal electrode transurethrally. After him, Katona, Ascoli, and Federici applied electrostimulation in patients with chronic neurogenic retention and hyperreflexia [2, 3]. In the field of urology, electric currents were and are used particularly in the bladder, the pelvic floor muscles, and the sacral roots [4–6]. A publication by Nashold et al. in 1971 reported on a successful implantation of a neural prosthesis in the sacral segment of the spinal cord [7]. The implant was used to activate voiding in a patient with spinal cord injury. Jonas and Tanagho tried to improve this prosthesis because during stimulation not only the bladder contracted but the urinary sphincter as well [8, 9]. Later Tanagho and Schmidt demonstrated that the stimulation of sacral root S3 generally modulates detrusor and sphincter action and could be used in clinical practice [10–13]. After two decades of experimentation with sacral root stimulation, finally in October of 1997, sacral neuromodulation for treatment of refractory urge incontinence was approved by the Food and Drug Administration in the United States. More than 25,000 patients underwent sacral nerve stimulation (SNS) since the FDA approval.

Mechanism of action

The stimulation of afferent nerve fibers by electrical current modulates reflex pathways involved in the filling and
evacuation phase of the micturition cycle through spinal circuits mediating somato-visceral interactions within the sacral spinal cord. SNS is proposed to activate or “reset” the somatic afferent inputs that play a pivotal role in the modulation of sensory processing and micturition reflex pathways in the spinal cord [14, 15]. Because beneficial effects can be demonstrated at intensities of stimulation that do not activate movements of striated muscle, the afferent system is the most likely effected [16].

Urinary retention and dysfunctional voiding can be resolved by inhibition of the guarding reflexes. Detrusor overactivity can be suppressed by one or more pathways, i.e., direct inhibition of bladder preganglionic neurons, as well as inhibition of interneuronal transmission in the afferent limb of the micturition reflex [15].

Recent research with PET scanning indicates that at the level of the brain, the activity of centers in the paraventricular grey involved in activation or inhibition of the micturition reflex, can be enhanced or reduced by sacral nerve stimulation and results in up- or downgrading of lower urinary tract activity [17–19]. Blok et al. reported on the acute and chronic effects of SNS for urge incontinence on the brain. They registered differences between newly and chronically implanted patients in brain areas involved in sensory and motor learning. No differences were seen in regional cerebral blood flow (rCBF) in areas that are part of the micturition reflex. Changes in rCBF were seen in specific areas: areas known to be involved in micturition and areas involved in awareness and awakeness. Acute SNS modulates sensorimotor learning areas and these become less active during chronic SNS [20].

Selecting patients for SNS

All patients who have symptoms of voiding dysfunction and who cannot be helped by other measures should be considered for SNS. Patient selection begins with a careful history, physical examination, routine urine tests, and, very important, the voiding diaries. Voiding diaries are a valuable instrument during the selection and have to be filled in carefully. Urodynamics are used to identify the patients with detrusor overactivity with or without urinary leakage or urinary retention. Koldewijn et al. studied predictors of success in 100 test stimulation patients and did not find any [21]. Scheepens et al. studied the data from 211 patients who underwent a trial stimulation (percutaneous nerve evaluation [PNE]) to determine the clinical parameters that can enhance the prediction of PNE success. They found that intervertebral disk prolapse, duration of complaints, neurogenic bladder dysfunction, and urge incontinence were found to be significant predictive factors. However, a PNE remains necessary to evaluate a patient’s chance of permanent implant success objectively [22].

Cohen et al. recently published a study on motor versus sensory response. They concluded that a good motor response during implantation was a predictive factor (in 95% of successfully treated patients) for success while a sensory response was not. All these patients were implanted under local anesthesia but with intravenous sedation and therefore the sensory perception of these patients may be unreliable [23].

Although not clearly reported before, it is known that a substantial part of the patients selected for SNS therapy have a history of psychological dysfunction and/or sexual abuse in the past. Weil et al. reported that special attention is needed for this group of patients [24]. They noted that patients with a history of psychological disorders, who had a good response during temporary test stimulation, had a far greater chance of lack of maintaining effect after permanent implantation. Of these patients, 82% showed a poor result after definitive implantation compared to 28% of the patients without a history of psychological disorders. Besides this lack of effect, 25% of the reoperations was done in this group, most of them with no effect. Psychological testing or psychiatric evaluation in case of doubt was advised before implantation of a permanent system.

A study by Everaert et al. showed similar findings [25]. In this study, the two-stage procedure was compared with the single-stage procedure. In the two-stage implant group, there were no failures during the first stage, while in the single-stage procedure, three patients had an immediate failure. They suggested that these results might be strongly influenced by psychological factors. Mental disorders were not related to objective or subjective success but these cofactors surely interfere with symptomatology and therewith co-influencing the results of therapy.

During the MDT-103 trial (to get the FDA approval) in 89 patients, depression and health-related quality of life (HRQOL) were assessed [26]. Patients were divided in a direct implant group and a delayed implant group. At baseline, they noted detectable levels of depression in 73% of all patients. After 3 months, patients in the implanted group had a significant improvement in depression scores. These improved scores remained at the 6- and 12-month visits. The scores on the SF-36 questionnaire, a questionnaire to investigate pain, vitality, physical functioning, social functioning, and mental health, increased in the implant group for role physical, pain, and social functioning. This study demonstrated the serious impact that unresolved voiding dysfunction has on quality of life. SNS was associated with significant improvement in depression and HRQOL.
Technique of sacral neuromodulation

Although sacral neuromodulation is planned as a long-term treatment, the therapy incorporates a unique temporary test stimulation procedure that allows patients and physicians to assess sacral nerve stimulation over a trial period [27]. Results of the trial are used by the physician to evaluate the viability of a permanent surgical implant. This test stimulation is conducted as an outpatient procedure preferably under local anesthetic and comprises of two steps, acute testing and the home evaluation phase. This testing phase is used to be performed as a PNE test in an outpatient clinic setting. The original technique was described by Schmidt et al. and is a simple procedure to evaluate the effect of sacral neuromodulation [27]. A test needle is inserted into the third sacral foramen to stimulate the sacral root. Lead migration is a known complication of this test, other complications are technical failures or pain [28]. Some patients who fail a PNE test are still good candidates for SNS therapy. For these reasons, a two-stage implant technique was developed [29]. With this technique, a permanent electrode is implanted and connected to an external stimulator. Less lead migration and a longer test period made it possible for clinicians to separate non-responders from technical failures or pain. Results of the trial are used by the physician to evaluate the effects of therapy on the voiding variables. If the patient’s symptoms under the existing list of indications for SNS improve at least more than 50% then the patient is a candidate to undergo the stage II or permanent step in which the permanent implantable pulse generator (IPG) unit is implanted in the soft tissue of the buttock of the patient.

During the second stage, the implanted tined lead is connected to the implantable pulse generator with a connecting cable that is passed subcutaneously. Relatively low amplitudes (0–3.0 V) are sufficient for stimulation of the somatic nerve fibers and to minimize the potential for nerve damage due to overstimulation. Within the recommended stimulation parameters (210 μS, 10–16 Hz), continuous stimulation is possible without pain sensation.

Unilateral or bilateral stimulation

Although temporary and chronic SNS can result in significant permanent clinical improvement, some patients improve only partially or temporarily [34, 35]. For these patients, several methods have been developed to improve the results [30–32]. The most widely accepted method to test a patient for SNS therapy is unilateral stimulation. In some clinics, bilateral stimulation has been suggested as a method to obtain better results [36, 37]. The bilateral innervation of the bladder is the basis for this type of intervention [38, 39]. Animal studies were performed to find a scientific basis for the application of bilateral neuromodulation. Animal studies by Schultz-Lampel et al. suggest that bilateral sacral neuromodulation can be a more effective technique for voiding dysfunction [40]. They conclude that bilateral stimulation may be more effective at lower stimulation intensities with positive side effects as longer stimulator-battery life and less potential nerve damage.
The only prospective randomized crossover trial to compare unilateral approach with bilateral sacral nerve stimulation is performed by Scheepens et al. [41]. In this study, 33 patients with chronic voiding dysfunction underwent unilateral as well as bilateral test stimulation to assess the possible advantages of bilateral stimulation. All patients were stimulated during at least 72 h in a unilateral and a bilateral setting with a washout period of at least 48 h between these two test periods. Standardized voiding records were used, and urine was measured using standard measuring cups. They analyzed results for 12 patients with urge incontinence and for 13 patients with non-obstructive urinary retention. They did not find any significant differences comparing the results for unilateral with bilateral stimulation. Although two patients of the retention group started voiding during bilateral stimulation, during unilateral stimulation they were still in complete retention. The reason for this remarkable result could be that with bilateral stimulation sufficient sacral nerve afferents are stimulated to achieve marked effect at central level.

In conclusion, unilateral stimulation should be performed before bilateral sacral stimulation is considered. However, a bilateral test stimulation could be indicated when a unilateral test fails [41, 42]. Further research with clinical follow-up could identify suitable patients for bilateral sacral nerve stimulation.

Clinical results and complications of SNS

In 1999, a prospective randomized study was published which evaluated the results of SNS therapy for urge incontinence (UI) were evaluated [43]. In total, 76 patients were treated in a multicenter trial; 34 patients were implanted and received chronic stimulation for 6 months; after these 6 months, they completed a therapy evaluation test (on versus off), 42 patients in a delay group were treated with standard medical therapy for 6 months and were offered implantation after this period. After 6 months, the number of daily incontinence episodes, the number of daily replaced diapers, and the severity of incontinence was significantly reduced in the stimulation group. In the stimulation group, 16 patients (47%) were completely dry and ten patients (29%) showed a greater than 50% reduction in incontinence episodes. After 18 months, the efficacy appeared to be sustained. During the therapy evaluation at 6 months, the stimulation group returned to baseline symptoms when stimulation was stopped.

Hassouna et al. reported in 2000 on the treatment of urgency–frequency (UF) symptoms with SNS therapy [44]. In total 51 patients, a stimulation group of 25 patients and a control group of 26 patients enrolled in this multicenter trial. All these patients had been tested with a PNE test and showed satisfactory responses. The stimulation group was implanted directly after this test, the control group was implanted after 6 months delay period. Statistically significant improvements were seen in the stimulation group for diary parameters as: number of voids daily (16.9±9.7–9.3±5.1), volume per void (118±74–226±124 ml), and degree of urgency (rank 2.2±0.6–1.6±0.9). In the control group, no significant changes were seen. After 6 months, the stimulation group had an evaluation test and urinary symptoms returned to baseline when stimulation was turned off. After reactivation of the stimulation, sustained efficacy was seen at 12 and 24 months.

A report of use of SNS in urinary retention was published in 2001 by Jonas et al. One hundred seventy-seven patients with urinary retention refractory to conservative therapy were enrolled in this multicenter trial between 1993 and 1998 [45]. Thirty-seven patients were assigned to treatment and 31 to the control group. At 6 months, the stimulation group showed 69% elimination of catheterization and an additional 14% with greater than 50% reduction in catheter volume per catheterization. Temporary inactivation (3 days) of SNS therapy resulted in significant increase in residual volume. The effectiveness of SNS therapy was sustained for 18 months after implantation. The first long-term follow-up results of the above-mentioned patient series were published in 2000 [28]. Results were reported for 1.5 to 3 years of follow-up. Of 41 UI patients, 59% showed a greater than 50% reduction in leaking episodes with 46% of these patients being completely dry after 3 years. After 2 years of follow-up, 56% of the UF patients showed a greater than 50% reduction in voids per day. In the retention group, 70% of 42 patients showed a greater than 50% reduction in catheter volume per catheterization.

Recently, the 5-year follow-up results of patients included in the trial, in order to get FDA approval, were analyzed. Of 163 patients enrolled, 152 have been implanted. Of the 152 implanted patients, 96 (63.2%) had UI, 25 (16.4%) UF, and 31 (20.4%) retention. Voiding diaries were collected annually over 5 years. For UI patients, the mean number of leaking episodes per day declined from 9.6±6.0 to 3.9±4.0. For UF patients, mean voids per day decreased from 19.3±7.0 to 14.8±7.6. Mean volume voided per void increased from 92.3±52.8 to 165.2±147.7 ml. No life-threatening or irreversible adverse events occurred. Of 152 patients, 102 experienced 33 device-related and 246 therapy-related adverse events. At 5 years, 68% of UI, 56% of UF, and 71% of retention patients had successful outcomes. However, an important finding in this study is the high correlation between the 1- and 5-year success rates. Of the implanted patients, 84% with UI, 71% with UF, and 78% with retention continued to have a successful outcome at 5-year follow-up if successful at 1 year [46]. Different clinics have published their long-term results in the previous years.
July 2002 and September 2004, Hijaz et al. described the complications of the implanted system. These three patients had good results afterwards. One reoperation to reposition the IPG after complaints of pain. One or two reprogramming sessions. Three patients had a decrease in most severe pain scores; after a median follow-up of 19 months, six out of ten patients reported significant improvement in pain symptomatology.

All the above-mentioned studies reported on complications during SNS. Siegel et al. summarized the complications in patients with refractory urge incontinence, urgency–frequency, and urinary retention that were included in the original trials of SNS [28]. The complications were divided in PNE-related complications and implant-related problems. Of the 914 test stimulation procedures done on the 581 patients, 181 adverse events occurred in 166 of these procedures (18.2% of the 914 procedures). The vast majority of complications were related to lead migration (108 events, 11.8% of procedures). Technical problems and pain represented 2.6% and 2.1% of the adverse events. For the 219 patient who underwent implantation of the permanent system, the following adverse events were seen during follow-up: pain at neurostimulator site (15.3%), new pain (9%), suspected lead migration (8.4%), transient electric shock (5.5%), pain at lead site (5.4%), adverse change in bowel function (3.0%), and some less frequent events like technical problems, device problems, change in menstrual cycle, and others. Surgical revisions of the implanted neurostimulator or lead system were performed in 33.3% of cases (73 of 219 patients) to resolve an adverse event. Mostly, this was done to relocate the stimulator because of pain or because of suspected lead migration. No serious adverse events, side effects, or permanent injury was reported.

Recently our long-term follow-up results with complication rates were published [47]. Of 149 patients analyzed, 107 had overactive bladder symptoms and 42 had urinary retention. Mean follow-up was 64.2 (SD=38.5) months. In the whole group, 194 adverse events occurred. Six patients had infection in their implanted system, one was explanted for infection. Most events could be solved by giving advice or by reprogramming the stimulator. One hundred twenty-nine reoperations have been performed and 21 patients had their system explanted. Analysis of the data shows a striking difference in the incidence of reoperations, but small differences in subjective results in the groups of patients implanted before or after 1996, suggesting that a proactive approach towards adverse events is worthwhile.

In our experience with the tined lead implantation, we see a clear decrease in reoperation rate [50]. Of 39 patients implanted with the tined lead, Voskuilen et al. described seven severe adverse events on medium term, three of these needed a reoperation. Three patients could be treated with one or two reprogramming sessions. Three patients had a reoperation to reposition the IPG after complaints of pain. These three patients had good results afterwards. One patient with an incomplete spinal cord lesion has no benefit of the implanted system.

Of 161 patients implanted with the tined lead between July 2002 and September 2004, Hijaz et al. described the complications seen in their institute [51]. They had three categories for complications: infections, mechanical problems, and response-related dysfunction. In total, they reported 17 explantations (10.5%). Eight explantations were done due to infection and seven due to loss of effect. In 26 (16.1%) patients, they performed a revision after these patients presented with a decrease in clinical response. The reasons for revision were: mechanical (lead) problems, IPG site discomfort, lead migration, and infectious causes. These complication rates show a decrease over the years mainly due to technical and procedural improvements. Gaynor-Krupnick et al. as well as Hijaz and Vasada presented an algorithm for evaluation and managing of a malfunctioning neuromodulation system [52, 53].

**Expanding indications**

With the widespread use, incidental improvements were published for other pathological conditions. Use of SNS for other off-label applications has been reported for treatment of interstitial cystitis, chronic pelvic pain, pediatric voiding dysfunction, and neurogenic lower urinary dysfunction seen in multiple sclerosis.

In 2000, the first papers were published with positive results with the use of SNS in interstitial cystitis [54, 55]. Comiter evaluated the effect of SNS therapy for interstitial cystitis in a prospective study in 2003 [56]. Seventeen out of 25 patients were implanted with a permanent system. After a mean follow-up of 14 months, there were significant improvements in daytime frequency and nocturia improved from 17.1 to 8.7 and 4.5 to 1.1, respectively (p<0.01). Mean voided volume increased from 111 to 264 ml (p<0.01). Average pain score decreased from 5.8 to 1.6 points on a scale of 0 to 10 (p<0.01). Interstitial cystitis symptom and problem index scores decreased from 16.5 to 6.8 and 14.5 to 5.4, respectively (p<0.01).

Chronic pelvic pain or genitourinary pain is a hard to treat condition. Over the years, some cases were presented of patients implanted with a SNS system for these conditions. In 2001, Siegel et al. implanted ten patients with a permanent system [57]. After 9 months of follow-up, nine of ten patients had a decrease in most severe pain scores; after a median follow-up of 19 months, six out of ten patients reported significant improvement in pain symptomatology.

After the clinical implication of SNS therapy for voiding dysfunction, Matzel together with Schmidt and Tanagho started to investigate SNS therapy in bowel dysfunction [58, 59]. In a prospective non-randomized multicentre study, 37 patients underwent test stimulation with SNS therapy for fecal incontinence [60]. Thirty-four patients were implanted with a permanent system. The effect on incontinence was assessed by daily bowel habit diaries and...
a disease-specific quality of life questionnaire. The frequency of incontinent episodes per week decreased from 16.4 to 3.1 at 12 months and to 2.0 at 24 months for both urge and passive incontinence. The mean number of incontinence episodes per week, staining, and pad use declined significantly too. Quality of life improved significantly in ASCRS scales; in the SF-36 scales only social functioning improved significantly.

Jarrett et al. did a systematic review of SNS for fecal incontinence and constipation [61]. They reported total continence in 41–75% of the patients, 75–100% experienced improvement in the incontinence symptoms. The results for patients treated with SNS for constipation discussed in this review seem promising but limited data is available by this time.

The results of SNS therapy in children with neurogenic bladder dysfunction is described by Guys et al. [62]. In total, 42 children with neurogenic bladder dysfunction, mainly due to spina bifida, enrolled in this prospective randomized controlled trial. Twenty-one patients were treated conservatively, while the other 21 patients were treated with SNS therapy. After 12 months, no significant better results were seen in the group treated with SNS. The authors stated that probably the intervention group was too small or the bladder dysfunction in these patients too severe.

Sexuality

During routine follow-up, patients may report improved sexual functioning after implant. Pauls et al. recently reported a pilot study to determine if sacral neuromodulation has an effect on the patient’s subsequent sexual function [63]. Eleven patients with a permanent system implanted were questioned about their sexual function before and after implantation. With SNS therapy, sexual frequency and female sexual function index (FSFI) increased significantly. No correlation was found between improvement in urinary symptoms and FSFI scores.

Conclusions

After years of experimental therapy, initiated by Tanagho and Schmidt, sacral nerve stimulation is nowadays a widely used therapy. Although the mechanism for this therapy is still not fully understood, the therapy has been proven effective on the long term. Due to the less invasive technique and other technical improvements, it is expected that complication rates will further decrease within the coming years. The expanding use of SNS therapy in fields other than urology will probably result in FDA approval for gastrointestinal indications.

Further research, possible with the help of animal models, has to be performed to understand in a more precise way the mechanism of SNS therapy. Other goals in research could be: patient selection (finding ways to identify more appropriate candidates), the effect of sacral neuromodulation in combined (urology–gynecology–GI) pathology, the effect of bilateral versus unilateral stimulation.

Conflicts of interest The author is advisor and lecturer for Medtronic Inc.

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