Activation and inhibition of the micturition reflex by penile afferents in the cat

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Wook JP, Yoo PB, Grill WM. Activation and inhibition of the micturition reflex by penile afferents in the cat. Am J Physiol Regul Integr Comp Physiol 294: R1880–R1889, 2008. First published April 23, 2008; doi:10.1152/ajpregu.00029.2008.—Coordination of the urinary bladder and the external urethral sphincter is controlled by descending projections from the pons and is also subject to modulation by segmental afferents. We quantified the effects on the micturition reflex of sensory inputs from genital afferents traveling in the penile component of the somatic pudendal nerve by electrical stimulation of the dorsal nerve of the penis (DNP) in α-chloralose anesthetized male cats. Depending on the frequency of stimulation (range, 1–40 Hz), activation of penile afferents either inhibited contractions of the bladder and promoted urine storage or activated the bladder and produced micturition. Stimulation of the DNP at 5–10 Hz inhibited distension-evoked contractions and increased the maximum bladder capacity before incontinence. Conversely, stimulation at 33 and 40 Hz augmented distension-evoked contractions. When the bladder was filled above a threshold volume (70% of the volume necessary for distension-evoked contractions), stimulation at 20–40 Hz activated de novo the micturition reflex and elicited detrusor contractions that increased voiding efficiency compared with distension-evoked voiding. Electrical stimulation of the DNP with a cuff electrode or percutaneous wire electrode produced similar results. The ability to evoke detrusor contractions by activation of the DNP was preserved following acute spinal cord transection. These results demonstrate a clear role of genital afferents in modulating the micturition reflex and suggest the DNP as a potential target for functional restoration of bladder control using electrical stimulation.

electrical stimulation; spinal cord injury; dorsal nerve of the penis; frequency dependence

Reciprocal coordination of the urinary bladder and the external urethral sphincter (EUS) to maintain continence and produce micturition is controlled by the pontine micturition center (2, 3, 24). The spino-bulbospinal micturition reflex is also subject to modulation by peripheral afferent activity (29, 30) that can influence voiding efficiency (41). The objective of the present study was to quantify the effects of genital afferents traveling in the penile component of the somatic pudendal nerve on the micturition reflex in the cat.

Activation of pudendal afferent nerve fibers can engage spinal (5) and spino-bulbospinal (2, 3) reflexes that are integral to the regulation of bladder function. Fluid flow through the urethra activates pudendal afferents (56), and this flow-driven activation can elicit detrusor contractions and facilitate micturition (2, 44, 48). Furthermore, electrical stimulation of pudendal urethral afferents in the cat (5, 49) and human (21) elicits detrusor contraction, and these responses are preserved following decerebration and acute and chronic spinal cord transection (SCT; 5, 7, 49, 52). Conversely, activation of pudendal afferents can also inhibit detrusor contractions (33, 50), and electrical stimulation of genital afferents inhibits the micturition reflex in humans with spinal cord injury (SCI), multiple sclerosis, Parkinson’s disease, and other conditions resulting in neurogenic detrusor overactivity (19, 22, 28, 31, 39, 58, 62).

Recent experiments in cats revealed that the frequency of electrical stimulation of afferents in the compound pudendal nerve determines their effect on the micturition reflex, with frequencies <20 Hz producing inhibition and frequencies >20 Hz producing activation of the micturition reflex (7, 52). However, the effects of genital afferent inputs on the micturition reflex and whether they exhibit similar frequency dependence is unknown. The present results demonstrate that, similar to effects produced by other pudendal afferents, activation of genital afferents produced either activation or inhibition of the micturition reflex in the α-chloralose anesthetized male cat, depending on the frequency of electrical stimulation of the afferents. The results reveal that genital afferent stimulation can activate both continence and micturition-like neural pathways, challenging the perception that the effect of stimulation of the genital afferents is solely inhibition of the micturition reflex.

Methods

All animal care and experimental procedures were followed according to the National Institutes of Health guidelines and were approved by the Institutional Animal Care and Use Committee at Duke University.

Preparation. Twenty sexually intact, adult male cats weighing 2.5–5.5 kg were anesthetized with ketamine-HCl (Ketaset; 35 mg/kg IM) and anesthesia was maintained with α-chloralose (65 mg/kg IV, supplemented at 15 mg/kg; Sigma-Aldrich). The animals were intubated and respired artificially to maintain end-tidal CO2 between 3.3 and 4.5%. Blood pressure was monitored via a catheter in the carotid artery. Core body temperature was maintained at 38° with a thermostatic heating pad, and intravenous fluids were administered (saline or lactated Ringers solution at 15 cc·kg⁻¹·h⁻¹).

The bladder was exposed through a midline abdominal incision, and the ureters were isolated, ligated, and cut proximal to the ligation. A suprapubic catheter (3.5 French) was inserted into the bladder dome and secured with a purse-string suture, and the abdominal incision was closed in layers. Intravesical pressures were measured with a solid-state pressure transducer (Deltran; Utah Medical) connected to the catheter and recorded (sampling rate = 12.5–20 kHz, Astromed8Xe; Astro-Med). For isovolumetric experiments, the urethra was occluded with a 3.5 French or 5 French catheter. In five cats, the catheter had three platinum ring electrodes positioned 6, 7, and 8 cm from the tip. The costs of publication of this article were defrayed in part by the payment of page charges. The article must therefore be hereby marked “advertisement” in accordance with 18 U.S.C. Section 1734 solely to indicate this fact.

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catheter was inserted so that two of the electrodes were within the postprostatic urethra and bulbourethra (3–5 cm from the urethral meatus (60)) and used to measure the activity of the periurethral musculature. This electromyogram (EMG) recording has been defined as the periurethral EMG (PU EMG) to reflect that it may include contributions from multiple active muscles, including the EUS. Wire electrodes were also inserted into the external anal sphincter (EAS) to record the anal sphincter EMG. The EMG signals were amplified (gain = 1,000), filtered (10–2 kHz), and sampled.

In two cats, the spinal cord was transected and responses to genital afferent stimulation were measured 8–15 h after SCT. The spinal cord was exposed at the T10 vertebral level via laminectomy, the dura was incised, and lidocaine was administered to the exposed cord. The spinal cord was elevated and transected, and Surgicel was packed between the transected ends of the spinal cord.

Nerve stimulation. The dorsal nerve of the penis (DNP) was stimulated unilaterally either by placing a cuff electrode directly around the nerve (n = 13) or by percutaneous insertion of a wire electrode adjacent to the nerve (n = 7). With the animal in a supine position, a midline incision was made from the caudal border of the gracilis muscle to a few millimeters cranial to the caudal end of the prepuce. For direct stimulation, the DNP was dissected free from the body of the penis caudal to the bulb of the penis, and a monopolar cuff electrode (platinum contact embedded in a silicone elastomeric cuff) was placed around the nerve. Indirect stimulation was delivered through an insulated stainless steel wire inserted via a 22-gauge needle between the prepucce and the glans penis and directed along the dorsolateral body of the penis 2.5–3.0 cm from the tip of the glans penis. The percutaneous electrode was inserted so that the DNP was likely activated unilaterally, and no characteristics of the observed responses suggested otherwise. A 20-gauge stainless steel needle was inserted into the ipsilateral leg of the animal and served as the anode during percutaneous and cuff electrode stimulation. Stimuli were 20- to 30-s trains of monophasic constant current pulses (100–μs pulse width) delivered at varying amplitudes (direct stimulation: 10 μA–1 mA; percutaneous stimulation: 100 μA–8 mA) and frequencies (1–40 Hz). The train lengths and frequency range were chosen based on previous data on the effect of stimulus parameters on bladder response (7).

Experimental design. Experiments were performed either with the urethra occluded (isovolumetric experiments) or unobstructed. All artificial bladder filling was performed with room temperature saline. Isovolumetric experiments were performed on 18 cats, of which 18 cats included stimulation delivered when the detrusor was relaxed (direct stimulation in 11 cats, percutaneous stimulation in 7 cats) and 16 cats included stimulation delivered during distension-evoked detrusor contractions (direct stimulation in 9 cats, percutaneous stimulation in 7 cats).

Volume thresholds were investigated systematically in eight cats. The bladder was filled in discrete 1-ml increments, and stimulation was applied during a 2- to 3-min interval between filling. The minimum volume at which electrical stimulation of the DNP could evoke detrusor contractions was defined as the stimulation threshold volume (STV). The minimum volume at which reflex bladder contractions (≥10 cmH2O) occurred was defined as the distension threshold volume (DTV). For a detected distension-evoked contraction, the pressure baseline was defined as the 2 s preceding the onset of the distension-evoked contractions, and all PTPs were computed from the intravesical pressure (minus baseline) for 20 s following onset of the distension-evoked contractions. Distension-evoked detrusor contractions were only included in the analysis if they lasted at least 10 s in the absence of stimulation, and stimulation trials were only included if stimulation began within 10 s of the onset of the distension-evoked contraction. Trials that occurred in the same animal, at the same volume, and within 3 min of one another were grouped together. Each group of trials included one or more distension-evoked contractions during which stimulation did not occur. The PTPs in each group were normalized by dividing by the average PTP of the distension-evoked contractions.

Direct DNP stimulation was performed in four cats with an unobstructed urethra to investigate the effects of genital afferent activity on bladder storage and voiding. The bladder was filled continuously at 1 ml/min, and failure of urine storage was defined as the point when urine leakage was observed or a sustained distension-evoked contraction occurred (defined as a contraction lasting at least 20 s and having a pressure increase ≥15 cmH2O). In the urine storage trials, continuous stimulation was applied starting at 50–80% of the previously determined volume at which failure of storage occurred (in absence of stimulation) and the stimulation was stopped when failure of storage occurred. Distension-evoked voiding was measured by stopping bladder filling when failure of storage occurred and allowing distension-evoked voiding to complete (3–5 min after the last volume was voided). Stimulation-evoked voiding was measured by stopping filling when failure of urine storage occurred and immediately applying stimulation. Stimulation was applied at a single frequency and at varying amplitudes until stimulus-evoked contractions (which resulted in voiding) could no longer be elicited. For each cat, a distension-evoked voiding/storage trial was performed first, and subsequent trials for the storage and voiding studies were randomized.

Voiding efficiencies were defined by dividing the difference between the initial bladder volume and the postvoid residual volume by the initial bladder volume

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\text{Voiding efficiency} \% = \frac{V_{\text{initial}} - V_{\text{residual}}}{V_{\text{initial}}} \times 100
\]

Statistical analysis. Statistical analysis of the dependence of detrusor responses on frequency was performed using a one-way Kruskal-Wallis test with the null hypothesis that the percentage of trials showing a specific response did not vary across stimulus frequencies. Post hoc paired comparisons between individual frequencies were made using Bonferroni inequalities. Comparison of mean contraction amplitudes (across frequencies, across cats, and across frequencies for individual cats) was made using a one-way Kruskal-Wallis test or a Wilcoxon rank sum test (for individual cats if contraction occurred at only 2 frequencies). Statistical analysis of the normalized PTPs computed for distension-evoked contractions with and without stimulation was done by a one-way Kruskal-Wallis test and post hoc paired comparisons were made using Bonferroni inequalities. Continent and voided volumes in the presence and absence of stimulation...
were compared separately for each cat using a Welch two-sample t-test. All reported values are means ± SD.

RESULTS

The DNP originated from the sensory branch of the pudendal nerve, passed superficially along the ventral insertion of the ischiocavernosus muscle into the bulb of the penis, and continued toward the glans penis. As it coursed along the body of the penis, the DNP branched into three discernible fiber populations (Fig. 1). Superficial fibers traveled along the dorsal aspect of the body of the penis and innervated the glans and the prepuce. Lateral fibers branched off deeper into the penile body and innervated the glans and the prepuce. Lateral fibers branched off deeper into the penile body and had cutaneous terminations. Similar DNP anatomy was observed in all 13 cats in which the DNP was isolated for stimulation with a cuff electrode. Some variability was observed in the distance caudal of the bulb of the penis before the branches of the DNP could be distinguished from one another.

The cuff electrode was placed on the DNP proximal to any branching unless otherwise specified.

Activation of penile afferents in the DNP elicited stimulation-frequency dependent activation or inhibition of the micturition reflex. Detrusor contraction was elicited by direct or percutaneous stimulation of the DNP in 17 of 20 cats, and inhibition of distension-evoked detrusor contractions was elicited in 16 of 18 cats. No response to DNP stimulation was observed in two cats, and only activation (detrusor contraction) or only inhibition (detrusor relaxation) was observed in one cat each (these cats were omitted from further analysis). Three cats were excluded from quantitative analysis because urethral leakage occurred during detrusor contractions despite the presence of the urethral catheter. Transecting the DNP cranial to the stimulating electrode (3 cats) or transecting the compound pudendal nerve (3 cats) abolished both the detrusor and EMG responses to DNP stimulation. The ability to elicit detrusor activation by electrical stimulation of the DNP was dependent on the bladder volume. STVs (3–10 ml, n = 8 cats in which thresholds were investigated in 1-ml increments) were less than distention threshold volumes (4–13 ml), and STVs averaged 70% ± 7% of DTVs (range = 61–80%).

Stimulation at bladder volumes above the STV elicited frequency-dependent activation of the micturition reflex. Direct electrical stimulation of the DNP activated the micturition reflex and elicited detrusor contractions in nine of eleven cats (Fig. 2A). The effect of stimulation frequency was examined systematically in seven cats. The ability to elicit sustained detrusor contractions depended on stimulation frequency (Fig. 2B, P < 0.001, Kruskal-Wallis test, n = 453 trials across seven cats, stimulation amplitude at 2–4× threshold). Stimulation at 20, 33, and 40 Hz elicited contractions more consistently than stimulation at any of the lower frequencies (≤10 Hz) (P < 0.01, Bonferroni inequalities). However, the response to stimulation at 20 Hz was not consistent across all cats. In five of nine cats, 20-Hz stimulation elicited contractile responses (contraction occurred in >80% of the trials), while in four of nine cats it did not elicit contractile responses (contraction occurred in <15% of the trials). The lower percentage of trials in which 20-Hz stimulation evoked contractions compared with 33 and 40 Hz is a reflection of this interanimal variability and not the effectiveness of 20 Hz stimulation within individual cats. The threshold stimulation amplitudes for eliciting detrusor responses ranged from 50 to 300 μA. The ability to elicit contractions and contraction magnitudes increased when stimulation amplitude was increased from threshold to 2× threshold, but stimulation at 2–4× threshold evoked similar detrusor contractions.

The mean increase in intravesical pressure evoked by stimulation at 20, 33, and 40 Hz was 33.5 cmH2O (±15.4 cmH2O, n = 283 contractions across seven cats, stimulation amplitude at 2–4× threshold), and did not vary significantly across these frequencies (P = 0.207, Kruskal-Wallis). While the magnitude of stimulus-evoked contractions varied from cat to cat (P < 0.01, Kruskal-Wallis), there was no significant difference between contraction amplitudes elicited by stimulation at 20, 33, or 40 Hz in six of the seven cats (P > 0.05, Kruskal-Wallis, n = 24–67 contractions per cat). In one cat, stimulation at 40 Hz elicited larger contractions than stimulation at 33 Hz (32.5 ± 8.0 vs. 27.1 ± 8.5 cmH2O, P < 0.05, Wilcoxon rank
were 2–4 / H11003 DTVs. Stimulus amplitudes ranged from 150
0.01, Bonferroni inequalities). Bladder volumes were above STVs and below
bar is the number of trials.

A

![Intravesical Pressure](image)

**Fig. 2.** Frequency-dependent bladder responses to direct electrical stimulation of the DNP. A: direct stimulation of the DNP is shown evoking detrusor contractions at bladder volumes between the stimulation threshold volume (STV) and distension threshold volume (DTV) in 2 different cats. Contractions were generated within 5 s of the onset of high-frequency stimulation (33 and 40 Hz in the top trace, 20–40 Hz in the bottom trace) and ended with the termination of stimulation or shortly thereafter. Stimulation at low frequencies (≤10 Hz) did not elicit contractions. Black bars indicate the duration of stimulation, which consisted of 20- or 30-s trains at the frequency and amplitude above each bar. B: percentage of trials in which direct stimulation of the DNP elicited detrusor contractions at different stimulus frequencies. The ability to elicit detrusor contraction by stimulation of the DNP was dependent on stimulation frequency (P < 0.001, Kruskal-Wallis test, n = 453 trials across 7 cats). Stimulation at 33 and 40 Hz consistently evoked detrusor contractions when stimulation was applied at appropriate bladder volumes and stimulus amplitude. Stimulation at 20 Hz evoked contractions in 4 of the 7 cats represented. Stimulation at 20, 33, and 40 Hz elicited contractions in a significantly greater percentage of trials than stimulation at 2–10 Hz (∗∗P < 0.01, Bonferroni inequalities). Bladder volumes were above STVs and below DTVs. Stimulus amplitudes ranged from 150 μA to 600 μA (all amplitudes were 2–4× the threshold to elic a bladder response). The number above each bar is the number of trials.

sum test, n = 60 contractions) while 20-Hz stimulation did not
evoke detrusor contraction.

**Stimulation at volumes above the DTV elicited frequency-
dependent detrusor relaxation and contraction.** At bladder volumes above the DTV, direct stimulation of the DNP elicited detrusor contraction and relaxation in eight of nine cats (Fig. 3, A and B). The effect of stimulation frequency was examined systematically in seven cats, and the detrusor response depended on stimulation frequency (Fig. 3C, P < 0.001, Kruskal-Wallis, n = 212 trials across 7 cats). Stimulation at 5–10 Hz consistently inhibited distension-evoked detrusor contractions, and stimulation at these frequencies inhibited distension-evoked detrusor contractions more consistently than stimulation at a lower frequency (2 Hz) and higher frequencies (33–40 Hz) (P < 0.01, Bonferroni inequalities). Stimulation during distension-evoked contractions at 20 Hz inhibited detrusor contractions in three of eight cats, augmented detrusor contractions in four cats, and did not elicit either response in one cat. Detrusor relaxation was evoked at threshold amplitudes of 80–400 μA, but was elicited more consistently at 2–4× the amplitude threshold for relaxation.

At volumes above the DTV, stimulation at higher frequencies elicited detrusor contractions when delivered during distension-evoked contractions and augmented detrusor contractions when applied during distension-evoked detrusor contractions (Fig. 3, A and B). The ability to augment distension-evoked detrusor contractions was dependent on stimulation frequency (Fig. 3C, P < 0.001, Kruskal-Wallis test, n = 212 trials across 6 cats). Stimulation at 33 and 40 Hz augmented contractions in a greater percentage of trials than stimulation at 2–20 Hz (P < 0.05, Bonferroni inequalities). Augmentation of distension-evoked contractions resulted in maximum intravesical pressures similar to maximum stimulation-evoked pressures (in absence of distension-evoked contractions), but in some cases, stimulation elicited a brief period of relaxation prior to contraction (Fig. 3B).

The effect of DNP stimulation on distension-evoked contractions was investigated systematically in six cats. The average PTP of detrusor contraction differed significantly across stimulation groups (Fig. 4, Kruskal-Wallis, P < 0.001). Inhibition of a contraction resulted in a mean PTP of 0.47 (±0.017), which was significantly less than that of the unstimulated contractions (1.0 ± 0.07) and the augmented contractions (1.47 ± 0.29) (P < 0.01, Bonferroni inequalities). Also, the augmented contraction had significantly larger relative PTPs than the unstimulated contractions (P < 0.01, Bonferroni inequalities).

**The reflex response of perirethral musculature to direct stimulation of the DNP.** Direct stimulation of the DNP did not directly activate the perirethral musculature but elicited reflex PU EMG response in four of five intact cats. Whether electrical stimulation of the DNP elicited PU EMG responses depended significantly on stimulation frequency (Fig. 5). Low-frequency stimulation (1–5 Hz) elicited reflex responses in the bulbourethra with reflex latencies of ~10–12 ms. The persistence of the reflex response throughout the trial diminished at higher frequencies (7.5–10 Hz), and reflex responses were elicited only by the first 1–4 stimulation pulses at high frequencies (20–40 Hz). The magnitude of reflex responses appeared to decrease with increasing bladder volume and over the course of repeated stimulation during the experiment. Stimulation of the DNP also evoked a reflex response in the EAS in four of five cats at a latency of 8–9 ms. Similar frequency dependence was observed in the EAS reflex response to DNP stimulation (Fig. 5).

**Stimulation-controlled urine storage and voiding.** Urine storage and voiding were elicited by direct stimulation of the DNP in four cats (Fig. 6). Bladder volumes at incontinence in the presence of low-frequency (5–10 Hz) stimulation were
21 ± 10 ml compared with bladder volumes at incontinence in the absence of stimulation of 15 ± 7 ml (P < 0.02 for each cat, n = 25 trials across four cats with at least three trials of each type per cat, Welch two-sample t-test). Direct stimulation of the DNP at higher frequencies (33 and 40 Hz) begun at the volume at which continence was lost resulted in an increase in percent bladder voiding compared with distension-evoked percent bladder voiding in four of four cats (P < 0.05, n = 27 trials across four cats with at least three trials of each type per cat, Welch two-sample t-test). Distension-evoked voiding resulted in 37% (±13.5%) bladder voiding (19–60%), while stimulation-evoked voiding resulted in 64% (±12%) voiding (49%–84%).

Percutaneous stimulation of the DNP elicited frequency-dependent detrusor responses. Electrical stimulation of the DNP with a percutaneous wire electrode evoked activation and inhibition of the micturition reflex in six of seven cats. These responses exhibited the same characteristics as those evoked by direct stimulation of the DNP with a cuff electrode. The threshold amplitude for eliciting responses ranged from 1.5 to 4 mA. Whether stimulation evoked detrusor contraction or relaxation depended on the stimulation frequency (P < 0.001 for augmentation and inhibition, Kruskal-Wallis test, n = 520 trials across 6 cats). Stimulation at 20–40 Hz elicited detrusor contraction more effectively than stimulation at 2–10 Hz (P < 0.01, Bonferroni inequalities). Stimulation at 7.5 and 10 Hz inhibited distension-evoked contractions in a greater percentage of trials than stimulation at 33 and 40 Hz (P < 0.01, Bonferroni inequalities).

Selective stimulation of the penile body branches of the DNP elicits detrusor contractions. The cutaneous branch of the DNP, observed to branch off of the penile body within a centimeter of the bulb (Fig. 1), was stimulated selectively, separate from the two larger branches of the DNP that continued to course along the penile body. No EAS or detrusor responses were elicited by stimulation of the cutaneous branch in six of six cats. Conversely, costimulation of the two penile branches (separately from the cutaneous branch) elicited detrusor contractions and EAS responses comparable to those evoked by stimulation of the DNP in five of six cats. In three of four cats in which distension-evoked contractions occurred during stimulation of the divisions of the DNP, inhibition of distension-evoked contractions was elicited by low-frequency costimulation of the two penile branches.

Responses to DNP stimulation were preserved after acute SCT. At 8–15 h following SCT, direct stimulation of the DNP elicited detrusor contractions in two of two cats (Fig. 7B). Detrusor contractions were evoked at stimulation frequencies of 20, 33, and 40 Hz, while stimulation at lower frequencies failed to elicit detrusor contractions (Fig. 7C). The ability to elicit detrusor contractions depended on stimulation frequency.

Fig. 3. Direct stimulation of the DNP inhibited or augmented distension-evoked detrusor contractions dependent on the stimulation frequency. A: intravesical pressure in 2 cats shows inhibition and augmentation of distension-evoked detrusor contractions by stimulation of the DNP. Stimulation at 10 Hz (and 20 Hz in the bottom trace) inhibited distension-evoked contractions, while stimulation at 33 and 40 Hz augmented distension-evoked contractions. Bladder volumes were above the DTVs. The black bars indicate the duration of the DNP stimulation, which consisted of 20- or 30-s trains at the frequency and amplitude above each bar. B: a more detailed view of the detrusor inhibition (left) and augmentation (right) from the bottom trace in A. Stimulation at 10 and 20 Hz after the onset of distension-evoked contractions caused the intravesical pressure to rapidly return to baseline. Stimulation at 33 Hz during a distension-evoked contraction caused a rapid rise (after a brief decrease) in the intravesical pressure. C: percentage of trials in which direct stimulation inhibited or augmented distension-evoked detrusor contractions at different stimulus frequencies. Stimulation at 5–10 Hz inhibited detrusor contractions in a greater percentage of trials than stimulation at 2, 20, 33, and 40 Hz (***P < 0.01, Bonferroni inequalities, n = 7 cats). Bladder volumes were above the DTV and stimulus amplitudes ranged from 160 µA to 800 µA (all amplitudes were ±4× threshold). Numbers above each bar are the number of trials at each frequency.
Activation of the detrusor by stimulation of the DNP occurred only when there was a sufficient volume of fluid in the bladder, both before and after acute SCT, as observed with stimulation of the compound pudendal nerve and urethral and perineal pudendal afferents in cat (5, 7, 49) and human (21, 66) and with contractions evoked by urethral fluid flow (9, 44). The bladder volume dependence of the detrusor response is due to a neural mechanism rather than the length-tension properties of the bladder (5), and the ability of urethral afferent activity to elicit reflex discharges in bladder pelvic nerves is similarly dependent on the level of background facilitation of the micturition reflex (37). That detrusor activation was evoked only if there was a sufficient volume in the bladder implies a convergence between bladder (pelvic) and pudendal afferents in the spinal cord, as the responses were preserved following acute SCT. In addition to the volume dependence of the bladder response to DNP stimulation, studies have shown that pudendal stimulation evoked inhibition of bladder activity in chronic spinalized cats and dorsal genital nerve (DGN) stimulation evoked inhibition of the pelvic c-fiber bladder reflex in acute spinalized cats can be enhanced by increasing the stimulation intensity at inhibitory stimulation frequencies (36, 52). The dependence of DNP stimulation evoked bladder inhibition on stimulus amplitude has also been illustrated in humans with SCI (42).

Detrusor contraction elicited by activation of pudendal afferents occurs via both supraspinal and spinal pathways (2, 5). The presence of a pudendal afferent-driven spinal bladder reflex has been confirmed in neonatal and chronic SCI cats (14, 52), but this pathway is masked by supraspinal serotonergic inhibition in the adult cat (55). Contraction of the detrusor evoked by penile afferent activation in the intact cat could reflect activation of a spinal-bulbo-spinal excitatory pathway, a spinal-pudendo-vesical reflex, or the simultaneous inhibition of the supraspinal serotonergic inhibition of the spinal-pudendo-vesical reflex and activation of the spinal reflex. In the acute SCT cat, the activation is through a spinal reflex, which may only be accessible after removal of supraspinal inhibitory inputs by SCT.

Activation of urethral afferents influences the bladder through an apparent convergence with bladder afferents in the spinal cord (5, 54). Pelvic and pudendal afferent projections overlap in the lateral dorsal horn and the dorsal gray commissure of the lumbar-sacral spinal cord in the rat, cat, and macaque monkey (12, 27, 40, 43, 45, 57). Similarly, afferent fibers from the DNP of the rat project to the dorsal horn, the dorsal gray commissure, and the sacral parasympathetic nucleus (40, 43).

DISCUSSION

Stimulation of penile afferents in the DNP activated reflexes that elicited either contraction or relaxation of the detrusor. Detrusor activation and inhibition were evoked differentially by electrical stimulation of the DNP dependent on the stimulation frequency. Low-frequency (5–10 Hz) stimulation inhibited distension-evoked detrusor contractions and promoted urine storage, while high-frequency stimulation (33–40 Hz) elicited detrusor contractions, augmented distension-evoked contractions, and produced bladder voiding. These results demonstrate that a somatic sensory input, heretofore asserted to only cause detrusor inhibition, can also cause detrusor contraction, and that the evoked response is strongly dependent on the frequency of electrical stimulation of the afferent inputs.

Activation of the detrusor by stimulation of the DNP occurred only when there was a sufficient volume of fluid in the bladder, both before and after acute SCT, as observed with stimulation of the compound pudendal nerve and urethral and perineal pudendal afferents in cat (5, 7, 49) and human (21, 66) and with contractions evoked by urethral fluid flow (9, 44). The bladder volume dependence of the detrusor response is due to a neural mechanism rather than the length-tension properties of the bladder (5), and the ability of urethral afferent activity to elicit reflex discharges in bladder pelvic nerves is similarly dependent on the level of background facilitation of the micturition reflex (37). That detrusor activation was evoked only if there was a sufficient volume in the bladder implies a convergence between bladder (pelvic) and pudendal afferents in the spinal cord, as the responses were preserved following acute SCT. In addition to the volume dependence of the bladder response to DNP stimulation, studies have shown that pudendal stimulation evoked inhibition of bladder activity in chronic spinalized cats and dorsal genital nerve (DGN) stimulation evoked inhibition of the pelvic c-fiber bladder reflex in acute spinalized cats can be enhanced by increasing the stimulation intensity at inhibitory stimulation frequencies (36, 52). The dependence of DNP stimulation evoked bladder inhibition on stimulus amplitude has also been illustrated in humans with SCI (42).

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and clitoral afferents in the cat terminate in the dorsal gray commissure (27). Activation of penile afferents results in Fos labeling of neurons in the sacral parasympathetic nucleus after SCT, consistent with supraspinal inhibition of DNP-activated spinal neurons (43). There is also evidence of supraspinal convergence of pelvic and pudendal afferents in regions associated with control of lower urinary tract function. Penile afferents project to the paragigantocellularis region of the medial reticular formation (20), and afferents activated by DNP stimulation, urethral infusion, and bladder distension converge on neurons in the medullary reticular formation (26). The ventral medullary gigantocellular reticular nuclei (which consists partially of the lateral paragigantocellular nucleus) projects to the intermediate gray and sacral parasympathetic nucleus of the sacral spinal cord (23). The convergence of penile somatic and bladder parasympathetic pathways is not surprising considering that DNP afferents and parasympathetic efferents are involved in erectile function (43). The present results extend the role of DNP afferents to urinary function, as well.

Stimulation of the DNP caused detrusor contraction at high frequencies (33 and 40 Hz) and detrusor relaxation at 5–10 Hz, similar to the frequency dependence for detrusor contraction and relaxation evoked by stimulation of afferents in the compound pudendal nerve (7, 52, 65). No studies have provided evidence to suggest a particular mechanism for this frequency dependence. The frequency dependence of the detrusor response to penile afferent stimulation may be due to frequency-dependent interactions between DNP afferents and parasympathetic bladder afferents and efferents. Low-frequency (5–10 Hz) activity in DNP afferents may evoke primary afferent depolarization of bladder afferents (1) or postsynaptic inhibition of interneurons activated by bladder afferents. At higher stimulation frequencies a second pathway may be excited by activation of an interneuron via temporal summation of the DNP afferent activity. This pathway would result in the inhibit...
Detrusor relaxation was likely caused by inhibition of the genital afferent stimulation in the human (19, 22, 28, 58, 62). Increases (22–366%) in continent bladder volumes result from previous study in intact cats with pudendal afferent stimulation of continent bladder volume of 35–77% over no stimulation. A sympathetic preganglionic neurons and sacral interneurons. Recordings from parasympathetic preganglionic neurons and sacral spinal interneurons reveal the existence of inhibitory and excitatory inputs from pudendal and pelvic afferents (15, 34).

Direct stimulation of the DNP resulted in increases in the continent bladder volume of 35–77% over no stimulation. A previous study in intact cats with pudendal afferent stimulation found volume increases of 18% over no stimulation (61), and in chronic spinalized cats the increase in continent volume during pudendal nerve stimulation was 147% (53). Similar increases (22–366%) in continent bladder volumes result from genital afferent stimulation in the human (19, 22, 28, 58, 62). Detrusor relaxation was likely caused by inhibition of the Aδ-fiber-mediated micturition reflex via reflex activation of sympathetic hypogastric efferents (at low intravesical pressure) or inhibition of pelvic efferents (at high intravesical pressure) (18, 32). The micturition reflex after chronic SCI is mainly mediated by c-fiber bladder afferents (8, 13, 16), but stimulation of the DGN can inhibit both the Aδ- and the c-fiber-mediated reflexes (36).

Electrical stimulation of the DNP at high frequencies (33 and 40 Hz) improved the voiding efficiency from 37% (±14%, no stimulation) to 64% (±12%, DNP stimulation). Voiding via stimulation of the DNP was achieved with continuous stimulation and without transection of any nerves via reflex activation of parasympathetic pelvic efferents (2, 3). The stimulation-evoked voiding efficiencies are comparable with voiding percentages reported from stimulation of perineal afferents of the pudendal nerve (63% ± 20%) and the compound pudendal nerve (64% ± 14%) in the α-chloralose anesthetized male cat (5, 6). Incomplete voiding and the consistent postvoid residual bladder volume may be due to the effects of the α-chloralose (46) or the reduced drive to the micturition reflex from the bladder afferents in the pelvic nerve when the bladder volume decreases as voiding occurs.

The ability to elicit voiding with continuous DNP stimulation demonstrates an apparent absence of stimulation-induced detrusor-sphincter dyssynergia (DSD), which is consistent with the present PU EMG recordings and previous EUS recordings during pudendal afferent stimulation (5, 7, 49). The PU EMG response was strongly dependent on stimulation frequency and did not persist at stimulus frequencies ≥ 20 Hz, consistent with previous findings (7, 10, 11). The lack of direct activation of the urethral sphincter as measured in the PU EMG, combined with the ability to void during continuous stimulation, suggests that stimulation of the DNP in the intact cat does not result in DSD. However, the bulbocavernous reflex and EUS response to DNP stimulation during detrusor contraction are enhanced in patients with DSD (17, 59), so the effect of high-frequency DNP stimulation evoked detrusor contractions on already present DSD is unclear.

While reorganization of the sacral spinal components of the micturition reflex pathway (4, 13) may affect the ability to elicit detrusor contractions by electrical stimulation of the DNP, recent results suggest that pudendal afferent mediated reflex activation of the bladder is preserved in humans with chronic SCI (21, 66). Also, perigenital electrical stimulation in the chronic spinal transected cat can activate and inhibit the bladder at similar frequencies as DNP stimulation (51). This method of stimulation in the female cat is likely activating the dorsal clitoral nerve, the female analog of the DNP, suggesting that the frequency-dependent bladder responses to electrical stimulation of the DNP observed in the intact and acute cat are still present in the chronic SCI cat.

**Perspectives and Significance**

Stimulation of the DNP at high frequencies (20–40 Hz) elicited detrusor contractions and augmentation of distension-evoked detrusor contractions, while stimulation at 5–10 Hz consistently inhibited distension-evoked detrusor contractions. These findings may find application in electrical restoration of bladder function, both storage and emptying, and electrical stimulation of the DNP is an especially compelling approach because of the ease of access to the dorsal genital nerves in human (63). Access to the DGN for electrical stimulation can be achieved more readily than access to the compound pudendal nerve or other urethral pudendal afferents (35, 47, 64), and the proximity of the DGN to the skin makes it an ideal target for percutaneous stimulation (64). Percutaneous stimulation of the DNP produced bladder responses equivalent to those generated by direct nerve stimulation. In addition, transection or block of the nerve distal to the stimulating electrode is not necessary because the DGN does not innervate the EUS (35). However, DNP stimulation may only be applicable to a limited population as certain individuals may be unable to tolerate stimulation at an adequate intensity (19). Furthermore, the excitatory bladder response to DNP stimulation has only been demonstrated in cats, and the existence of a similar excitatory bladder pathway from penile afferents in humans is unknown.

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**REFERENCES**

ACTIVATION AND INHIBITION OF THE MICTURITION REFLEX


