Medicare Coverage Policy ~ Decisions

Pelvic Floor Electrical Stimulation for Treatment of Urinary Incontinence (#CAG-00021N)

Exclusion Tables

Study/year	Patient characteristics	Study Design	Treatment	Dropouts	Outcome Measures	Possible threats to validity
Bent et al.	45 patients with stress incontinence (14) with detrusor instability (10), or mixed (21). Age range 25-80 years	Case series	Patients had self- administered therapy for 15 minutes, separated by at least 4 hrs, twice a day for 6 weeks. 20Hz applied for detrusor instability, 50Hz for stress incontinence.		Leakage episodes Pad use Pad test Standing stress test Standing CMG Resting/dynamic urethral closure pressure profiles Questionnaire	Potential for selection bias Exclusion criteria included "poor medical health" without further specification
Bratt, et al 1998	48 women with unstable detrusor and urge incontinence originally enrolled in a short-term study in 1989. 30 women located/surveyed Mean age of respondents 62 years (age range of original 48 women 15-79 years)	Questionnaire	Patients received 5-10Hz for 20 minutes, twice a week, at least five times	18/48 surveyed 10 deceased 2 disabled secondary to stroke, 5 lost to follow-up, 1 counted twice in the original study	6 close-ended questions including: Do you have any leakage of urine? If you have leakage, how often does it happen? IF you do have leakage, what is the worse problem? Were you satisfied with the electrostimulation as a treatment method? Would you recommend the treatment to a good friend?	Potential for selection bias, performance bias. Authors quote 90% response rate, however, 18 people could not be surveyed of the original 48. Lack of rigor of survey instrument.
Caputo, et al	76 women- 19 patients with	Case series	Patients received electrical stimulation for 15		Urinary incontinent episodes	Potential for selection bias

Table 1a. Exclusion articles- methodologic features

al	19 patients with SI,	minutes at 20 Hz, once a week for 6	episodes	selection bias
1993	30 patients with	weeks.	Voiding frequency	Numerous confounders
	DI,	Patients were		
	27 with mixed.	taught Kegel exercises and asked to perform exercise 50 times	(Measured by bladder diary)	
	Average age: 52.6 years	daily.		

Table 1a. Exclusion articles- methodologic features (cont'd)

Study/year	Patient characteristics	Study Design	Treatment	Dropouts	Outcome Measures	Possible threats to validity
Dumoulin, et al 1995	8 female patients (out of 10 volunteers) with GSI, persisting more than 3 months after delivery Average age 32 years	Case series	Subjects received nine treatment sessions over 3 weeks, consisting of two 15-minute sessions of neuromuscular electrical stimulation, followed by a 15 minute pelvic floor		Maximum muscle contractions (pressure using perineometry) Urine loss (pad test) Frequency of incontinence (self diary)	Small sample size (n=8) and short follow-up time (3 weeks) Confounders include that patients received both electrical stimulation and an exercise program, and therefore it is unclear what
			muscle exercise program (50 Hz applied)			benefit can be attributed to which element of therapy.
Eriksen, et al 1989	55 women with urinary stress incontinence awaiting surgical repair Average age 49 years	Case series	Chronic stimulation (25 Hz) applied anally or vaginally for a median of 5.4 months (Range 0.5 - 29 months). Pts instructed to use stimulator		Urethral closing pressure Amount of leakage	Potential for selection bias Potential for attrition bias- little data provided about the use of electrical stimulation
			regularly and as much as possible every day for at least 3 months before effect was evaluated. If improved, they were			No standardization of protocol. Large range of treatment time – 15 days to 29 months.
			encouraged to continue therapy. If no effect, colposuspension urethropexy was recommended.			Outcome measures were used that are not typically reported
Fossberg et al	91 patients (11 males and 80 females) with unstable	Case series	Patients received 12 treatments of 5-10 Hz for 20	17 dropouts	Frequency/volume charts	Potential for selection and attrition bias

	detrusor, frequency and urge incontinence		minutes	females 2 males	Cystometry Flowmetry	
	Mean age 53 years (20-78 years)				Subjective assessment	
Kralj, B 1999	111 women with moderate SI	Case series	Patients received electrical stimulation 20 Hz for 1.5-2 hours daily for 3 months		Pad test [Outcomes assessed six months after beginning of treatment; 3 months after terminating	Potential for selection bias. Little data provided on patient characteristics, as well as inclusion/exclusion
Miller et al 1998	31 women with GSI Average age: early 50's	Case series	15 patients treated daily 16 treated every other day Used device for 15 minutes twice a day or every other day for 20 weeks	3/31	Modified pad test QOL questionnaire Total number of incontinent episodes/ 3 day period as recorded in voiding diary	Potential for selection bias and performance bias.
Study/year	Patient	Study Design	Treatment	Dropouts	Outcome Measures	Possible threats
Study/year Richardson, et al 1996	Patient characteristics 31 women with GSI Average age 50	Study Design Case series	Study conducted at 6 sites.	Dropouts 10/31 one year follow-up	Outcome Measures Leakage episodes over 3 days (voiding diary)	Possible threats to validity Potential for selection bias and performance bias.
Study/year Richardson, et al 1996	Patient characteristics 31 women with GSI Average age 50 years Average years incontinent 6.5	Study Design Case series	Treatment Study conducted at 6 sites. Patients were assigned consecutively to either daily or every-other-day pelvic floor electrical stimulation for 15 minutes twice a day [50Hz applied]	Dropouts 10/31 one year follow-up	Outcome Measures Leakage episodes over 3 days (voiding diary) Pad count Leakage amount Subjective assessment and quality of life	Possible threats to validity Potential for selection bias and performance bias.
Study/year Richardson, et al 1996 Sand PK 1996	Patient characteristics 31 women with GSI Average age 50 years Average years incontinent 6.5 26 women with mixed incontinence complicated by a low-pressure urethra	Study Design Case series Case series (Retrospective)	Treatment Study conducted at 6 sites. Patients were assigned consecutively to either daily or every-other-day pelvic floor electrical stimulation for 15 minutes twice a day [50Hz applied] Patients used device twice a day [20 Hz] for 15 minutes for 8 weeks	Dropouts 10/31 one year follow-up 5/26	Outcome Measures Leakage episodes over 3 days (voiding diary) Pad count Leakage amount Subjective assessment and quality of life Visual analogue symptom scales Weekly incontinent epsidoes	Possible threats to validity Potential for selection bias and performance bias. Potential for selection bias. Nearly 25% dropout.

Siegel, et al	72 patients at 8	Case series	33 patients	4/72	Leakage episodes,	Potential for
	study sites. 66		treated daily,		nocturnal	selection bias
1007	completed 20		and 35 patients		episodes, voiding	
1997	week protocol.		treated every-		frequency, total	Brood evolution
			other-day with		voids, pad count	Broad exclusion
			either 12.5 Hz		, ,	criteria
	36 patients with		or 50Hz for 15			
	urge,		minutes twice a		Patient subjective	
			dav		assessment and	
	30 with stress				quality of life	
	50 With Stress.					
			Subjects agreed			
	Average age 53		to no other			
	years (34-82)		incontinence			
			treatments			
	A		during duration			
	Average years		of study			
	incontinent 9.7-					
	10.3					
Zollner-	38 female	Case series	Patients treated	7/38	Mean bladder	Potential for
Nielsen, et	patients with		for 20 minutes,		volume	selection bias
al	frequency,		twice a week			
	urgency, or				Number	Determinal form
1000	urge		Descional F 1F		Number of	
1992	incontinence		Received 5-15		micturitions	performance blas
			treatments			
					Ouestionnaire	
	Median age /1					
	years (35-90)					
	74% of patients					
	were 60 years					
	or older					
		<u> </u>		1	1	
Table 1b.	Exclusion a	articies– ou	itcomes			

Study/year	Pt recorded diaries	Pad test (grams)	Comments
	% % pts	% % pts %	
	Measure Pre- Post- change ¹ improv ² %cure ³	Pre- Post- change ¹ improv ² cure ³	
Bent et al	Not reported	Not reported	Objective criteria defined by authors did not show
1993	However, authors state that incontinent episodes decreased in 6 patients with SI, 1 patient with DI, 6 patients with mixed.	However, authors state that pad test results improved for 8 patients with SI and 5 patients with DI.	treatment. Subjective measures, based on a questionnaire, did demonstrate success but it is unclear as to the
	p=NS	p=NS	significance and reproducibility of survey results. Authors state that results were statistically significant but do not provide specific data.
			Short study period – 6 weeks
Bratt et al, 1998	Not measured 21 women (78%) reported symptoms of urge incontinence, 13 women having symptoms daily. 19 women (70%) reported symptoms of stress incontinence.	Not measured	Limited conclusions can be made since data is subjective. No objective data reported. Of note, a large number of women were suffering from stress incontinence that ten years earlier were treated for urge. The prevalence of

	21 women would recommend maximal stimulation to a friend.		urge incontinence was reported to be higher in the follow-up than the original study.
Caputo, et al	Not reported	Not measured	No statistical analyses provided.
1993	Overall objective improvement 76% ; 89% for GSI, 73% for DI, 70% for mixed.		Stimulation applied once a week – appears to be a departure from practice in most other studies.
	[objective improvement defined as a reduction in urinary incontinence episodes by 50%, or reduction in voiding frequency by 50%, or to 10 or fewer voids per 24 hours.]		Study lasted 6 weeks. Achieved up to one-year follow-up in only 40% of patients [avg 6.4 months]. The fact that only 15% relapsed is difficult to interpret due to attrition bias.
			Of note, authors state that controlled clinical trials are needed to determine its efficacy and standardize stimulation protocols before its widespread use.
Dumoulin, et al	Leaks/week	74.4 24.4 67% NR NR	Authors provide little raw data for results to be
1995	16.3 4.0 75% NR NR	p= 0.012	Teassesseu.
			Broad exclusion criteria, including patients with diabetes and heart disease.
			Authors do note that "further studies are needed to validate thisprotocol."

Table 1b. Exclusion articles- outcomes (cont'd)

Study/year	Pt recorded diaries	Pad test (grams)	Comments
	% % pts %	% % pts %	
	Measure Pre- Post- change ¹ improv ² cure ³	Pre- Post- change ¹ improv ² cure ³	
Eriksen et al	Not measured	Not measured	Data analysis minimal.
1989	Authors state that 68% of patients were continent or had improved. (Cure defined as positive urethral closure pressure found, and no leakage observed during the stress provocation tests)		Authors only provide data at 2 year follow-up, although outcome measures were apparently obtained at 3 months.
	At 2 year follow-up, success rate was reduced to 56%.		Data is not provided on all 55 women initially enrolled.

			Authors do not define "improved"
Fossberg, et al 1990	Leaks/day at 6 weeks 1. 8.0 12 % NR	Not measured	Study conducted in Norway with a device that is only slightly analogous to that used in the US
	p=0.003		Short study time
	Leaks/night at 6 weeks		Are reductions in micturition clinically significant?
	1.6 1.1 31% NR NR		At 6 weeks post treatment, almost half of patients felt their condition was unchanged.
Kralj, B	Not measured	Not reported	No statistical analysis provided.
1999	Authors state that 50.5 % of patients were cured, 23.4% improved, 26.1% failed. [Cure defined as no subjective complaints and pad tests were negative; improved defined as no subjective complaints, and pad test not negative]		Authors conclude by stating that "the efficacy of treatment depends on the patient selection, parameters of electrical stimulation, stimulator of the pelvic floor muscles, mode of stimulation and on motivation of the patient." Authors, however, provide no such guidance.
			Definitions of cure and improvement are not standard.
Miller et al 1998	Leaks/3day at 20 weeks Responders (n=19) 7.6 1.71 78 % NR NR	Not reported	Study designed to have adequate power to detect a reduction of 2.1 leakage episodes over 3 days. Is this clinically significant?
	Nonresponders (n=9) 7.8 9.51 - 22% NR NR		Data presentation unusual. There were two groups, yet no data was presented for those two separate groups. Instead, data stratified by those patients who showed response [defined as 50% decrease in total number of leakage episodes] vs those who did not respond. P values not reported.
Study/year	Pt recorded diaries	<u>Pad test (grams)</u>	Comments
	% % pts %	% % pts %	
	Measure Pre- Post- change ¹ improv ² cure ³	Pre- Post- change ¹ improv ² cure ³	
Richardson, et al	Leaks/3 day at 20 weeks		Since there is no direct comparison between this
1996	Daily (n=13)	Daily (n=13)	therapy and conventional therapy, it is difficult to discern that similar results
	0 C E C 2EN. 2001 2201	10 7 26 7 2E0/ ND	would have not been

	8.6 5.6 35% 39% 23%	48.7 36.7 25% NR NR	would have not been reported for
	p=0.06	p=0.11	Kegel/ bioreedback.
	Every-other-day (n=15)	Everv-other-dav	Short study time (20 weeks)
	6.9 3.0 56% 20% 53%	(n=15)	Cure improvement rates
	p=0.04	12.5 6.9 45% NR NR	were not statistically significant.
	Leaks/3 day at one year	p=0.38	Only total leakage episodes for daily users was
	Users (n=10)		statistically significant.
	9.2 2.0 78% NR NR		Little data is provided on one-year follow-up. At one
	p=0.009		year, ¹ / ₂ of users were performing Kegel and some started bladder training
	Nonusers (n=11)		confounders.
	5.8 4.6 21% NR NR		
	p=0.06		
Sand PK	Not measured	Not measured	Although there was
1996			subjective improvement in voiding frequency, urgency and stress incontinence, there is question of clinical significance. For example, between therapy, pts had 1.5 hrs between voids. After therapy, time between voids was 2.08 hrs. Is 30 minutes clinically significant? The actual number of incontinent episodes showed no difference between pre and post treatment.
			Authors do not define cure/improvement.
Siegel, et al	Leaks/3day	Not measured	Authors state that there was no difference in data
1997	Urge (n=35)		between daily and every- other-day treatment,
	9.6 5.3 45% NR NR		are not provided.
	p<0.001		Short study time – 20 weeks.
	Mixed (n=33)		Logistic regression
	9.6 3.9 59% NR NR		demonstrated lack of response associated with number of previous
	p<0.001		therapies.

Study/year	Pt recorded diaries	Pad test (grams)	Comments
	% % pts %	% % pts %	
	Measure Pre- Post- change ¹ improv ² cure ³	Pre- Post- change ¹ improv ² cure ³	
Zollner- Nielsen	Leaks/48 hrs	Not measured	Incomplete data provided.
1992	7.4 5.9 20% NR NR		Authors combine cured/improved with no p
	p<0.01		values.
			Study included 8 patients with neurologic causes of UI.
			No explanation of dropouts.
			No description of instrument used to assess subjective opinions.
			Results were similar for patients < 60 years and > 60 years.

¹ % change – Defined as the percent decrease in the frequency of incontinence over a specified time period, calculated by the following equation:

pretreatment episodes/period - posttreatment episodes/period X 100

pretreatment episodes/period

 2 % pts improv – Defined as the percentage of patients with 50% or greater decrease in the frequency of incontinence, as calculated by the previous equation.

³ % cure – Defined as the percentage of patients with 100% decrease in frequency of incontinence, i.e., no incontinent episodes over the specified time period.

⁴ % change – Defined as the percent decrease in the amount of urine lost in grams, following provocative maneuvers, calculated by the following equation:

pretreatment pad weight difference - posttreatment pad weight difference x 100

pretreatment pad weight difference

⁵ % pts improv – Defined as the percentage of patients with 50% or greater decrease in the amount of urine lost in grams following provocative maneuvers.

 6 % cure – Defined as the percentage of patients with 100% decrease urine loss, ie no urine lost following the provocative maneuvers.

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Key to Tables

ICS International Continence Society

MI Mixed incontinence (stress and urge incontinence)

%change Percent change in incontinence (frequency by pt recorded diary or urine loss on pad test)

%cure Percent of patients with no further incontinence

% pts improv Percent of patients with >50% decrease in incontinence (frequency by pt recorded diary or urine loss on pad test)

PFES Pelvic floor electrical stimulation

PME Pelvic floor muscle exercise

SI Stress incontinence

UI Urge incontinence

Selection bias Imbalances in patient characteristics between groups with potential for differences to affect outcomes

Performance bias Inequality in the intensity of treatment given between groups

Attrition bias Significant number of dropouts in one or more study arms, not taken into account in the statistical analysis